Aerobic Conditions Enhance the Photocatalytic Stability of CdS/CdOₓ Quantum Dots

David W. Wakerley, Khoa H. Ly, Nikolay Kornienko, Katherine L. Orchard, Moritz F. Kuehnel, and Erwin Reisner

Abstract: Photocatalytic H₂ production through water splitting represents an attractive route to generate a renewable fuel. These systems are typically limited to anaerobic conditions due to the inhibiting effects of O₂. Here, we report that sacrificial H₂ evolution with CdS quantum dots does not necessarily suffer from O₂ inhibition and can even be stabilised under aerobic conditions. The introduction of O₂ prevents a key back-electron transfer to CdS (over-accumulation of metallic Cd and particle agglomeration) and thereby affords particles with higher stability. These findings represent a possibility to exploit the O₂ reduction reaction to inhibit deactivation, rather than catalyst, offering a strategy to stabilise photocatalysts that suffer from similar degradation reactions.

Clean-burning, renewable H₂ fuel can in principle be generated effectively through solar-driven proton reduction coupled to water oxidation as an abundant source of electrons. Alternatively, this reaction can be undertaken through the oxidation of organic species, either in the form of biomass-derived substrates, such as EtOH, MeOH, glucose or lignocellulose or through selective organic oxidation reactions to generate higher-value products. Semiconductor particles are particularly well-suited to perform the underlying reactions behind artificial photosynthesis and as such, rapid light-driven H₂ evolution has been reported for numerous metal oxide, sulfide, selenide and nitride-based semiconductors.

Due to the ubiquity of O₂ in the atmosphere, as well as its production in the water-splitting reaction, a proton-reduction catalyst must be able to tolerate its presence during activity. To date, little research has considered the effect of O₂ on semiconductor-driven H₂ evolution and only few reports are available on O₂-tolerant molecular proton-reduction catalysts. Several strategies have therefore sought to protect proton-reduction photocatalysts from O₂ to allow catalysis to proceed. For example, deposition of thin layers of metal oxides, such as Cr₂O₃ and SiO₂/TiO₂, on the surface of a proton reduction catalyst can selectively prevent diffusion of O₂ to the catalyst, albeit under low levels of O₂ (<1 atm of pressure).

Previously reported systems have shown that proton reduction catalysts fall into two groups: O₂ sensitive, where a catalyst is irreversibly damaged by O₂, or O₂ tolerant, where a catalyst is able to function under O₂ but at a reduced rate (Scheme 1). Nevertheless, the intrinsic oxidising nature of O₂ does not need to be considered exclusively as a disadvantage and methods that use O₂ to stabilise activity can be envisioned. O₂ reduction is thermodynamically more facile than proton reduction and its presence in solution therefore offers a route to prevent a photocatalyst from a reductive deactivation pathway. Cu₃RhO₃ and CuFe₄O₅ delafossite-structured H₂-evolving photocathodes were previously demonstrated to operate most effectively under air using this strategy. In these examples, CuO accumulates under inert conditions, which can be avoided through the introduction of O₂, thereby increasing the electrode stability.

In this study, we demonstrate that O₂ can be used to stabilise activity in colloidal “one-pot” photocatalytic schemes and that even an improvement in catalytic H₂ evolution performance can be achieved with CdS quantum dots (QDs). CdS...
QDs are nanocrystals below 10 nm in diameter that have previously demonstrated excellent photophysical properties for light-driven proton reduction in the presence of sacrificial electron donors, catalysing this reaction at benchmark rates.\(^{29}\)

The photocatalytic \(H_2\) evolution activity of CdS has been reported to drop by 20% under 21% \(O_2\) when compared to anaerobic conditions.\(^{20}\) This observation can be assigned to the competitive reduction of \(O_2\) versus protons, as seen for other \(O_2\)-tolerant catalysts (Scheme 1); however, we show that by encouraging sufficiently fast \(H_2\) evolution, over-accumulation of reduced \(Cd^0\) at the particle occurs. Addition of \(O_2\) to this system precludes \(Cd^0\) formation and affords rapid and stable light-driven \(H_2\) evolution.

Capping-ligand-free CdS QDs were synthesised with a diameter of 4–5 nm,\(^{21}\) as confirmed by transmission electron microscopy (TEM, Figure 1a, see Supporting Information for experimental details).\(^{3}\) Photocatalytic experiments were undertaken by combining the QDs with Co(BF\(_4\))\(_2\) (0.25 mm), as a cocatalyst for the proton-reduction reaction, and MeOH (10 mM), as a sacrificial electron donor, in a sealed photoreactor that was irradiated with simulated solar light (AM 1.5 G, 100 mW cm\(^{-2}\)) at 25°C. The production of \(H_2\) was analysed at designated time intervals by headspace gas chromatography (Table S1 in the Supporting Information). Control experiments showed that no \(H_2\) was produced in the dark or in the absence of CdS (Table S2).

In highly-alkaline solutions (>1 M KOH), a surface layer comprised of \(CdO/Cd(OH)_2\) (\(CdO_x\), Figure 1b), forms on the particles, creating \(CdS/CdO_x\) QDs.\(^{3}\) This layer is believed to enhance the rate of photocatalysis. As such, in anaerobic conditions (Figure 1c, solid lines), CdS QDs displayed the highest rate of \(H_2\) evolution in 5 M KOH, as previously reported.\(^{22}\) However, the activity is not stable and slows down after only a few hours (Figure 1c). This coincides with the formation of a black precipitate in the solution and a loss of the \(CdS\) absorption peak in the UV/Vis spectrum (Figure 1d). In contrast, in 0.5 M and 0.05 M KOH the rate of \(H_2\) evolution was slower, but did not drop significantly over time and the solution remained yellow. Electron transfer from photoexcited CdX semiconductors \((X=S, Se)\) has previously been proposed to originate from surface \(Cd^0\) sites,\(^{23}\) and the black colour was therefore assigned to over-formation of \(Cd^0\) (see below for characterisation).

Photocatalysis was subsequently performed in a closed vessel with an aerobic headspace to determine whether the extent of \(Cd^0\) formation could be reduced using \(O_2\). Depending on the rate of \(H_2\) evolution, the presence of \(O_2\) had differing effects (Figure 1c, dashed lines). In 0.5 M and 0.05 M KOH, the introduction of \(O_2\) led to a decrease in photocatalytic performance by a factor of 48 and 82%, respectively. This effect is expected, as the \(O_2\) reduction reaction competes with the desired proton reduction reaction (Scheme 1). In 5 M KOH, the aerobic activity was surprisingly enhanced relative to anaerobic conditions, with a reduced formation of the black precipitate. Figure 1e illustrates the change in the UV/Vis spectrum of an aerobic sample over time, showing the retention of the \(CdS\) absorption band over 3 h of photocatalysis. In this sample, an eventual slowdown of the catalysis was visible after 6 h (Figure 1c), which was a result of consumption of \(O_2\) within the vessel headspace.

Transient absorption (TA) and Raman spectroscopy were subsequently employed to gain further understanding of the processes. The relationship between the pH and electron/hole dynamics was first probed under aerobic conditions by TA. Figure 2a shows the TA spectrum of \(CdS/CdO_x\) QDs in 10 M, 0.1 M and 0 M KOH with EtOH as an electron donor at a 1.5 ps delay, normalised to unity. The band-edge bleach of the \(CdS/CdO_x\) QDs appears at \(\approx 488\) nm, arising from electrons being excited to the conduction band. The spectra also display a second negative peak at 600–700 nm, which is tentatively assigned to surface-state traps.\(^{22}\)

Given the lower magnitude of the \(CdS\) signal in 0 M and 0.1 M KOH relative to \(CdS\) in 10 M KOH (see Figure S4 for absolute absorbance data), we propose that fewer photogenerated electrons are available due to ultrafast trapping and recombination pathways in the less alkaline conditions. In addition, a proportionally stronger bleach signal at 675 nm indicates that a larger fraction of electrons is trapped in the surface states rather than the conduction band at lower pH. The growth of a \(CdO_x\) layer on the \(CdS\) surface may therefore increase the

---

**Figure 1.** (a) TEM image of ligand-free CdS QDs. (b) Illustration of \(CdS/CdO_x\) formation from particles of ligand-free \(CdS/\text{BF}_4\). (c) Photocatalytic \(H_2\) production (AM 1.5 G, 100 mW cm\(^{-2}\)) at 25°C from \(CdS\) QDs (0.5 μm) in various concentrations of aqueous KOH (2 mL) containing MeOH (10 mM) in anaerobic (solid traces) or aerobic (dashed traces) conditions with 0.25 mm Co(BF\(_4\))\(_2\). (d) UV/Vis spectra of \(CdS/CdO_x\) QDs at designated intervals after photocatalysis in aqueous KOH (2 mL, 5 M) containing MeOH (10 mM) in the presence of Co(BF\(_4\))\(_2\) (0.25 mM) under anaerobic conditions (N\(_2\)). (e) The aerobic equivalent of the experiment in (d).
efficiency of photocatalysis by ensuring that electrons remain in the conduction band, rather than other trap states. The greater accumulation of electrons in the conduction band at high pH is likely to lead to a greater propensity for proton reduction, as well as the self-reduction of CdS to Cd\(^0\).

Raman spectroscopy under anaerobic conditions supports the proposed formation of Cd\(^0\) after electron accumulation. QD solutions in 10 mM KOH with EtOH were irradiated with a 413 nm laser (1 mW) over various time intervals and spectra were recorded using a 514 nm excitation beam (Figure 2b). The QDs show two bands in all cases, corresponding to the first and second overtone of the longitudinal optical phonon (LO) of CdS at 305 and 605 cm\(^{-1}\), respectively.\([24]\) Shoulders on either side of the LO peak were observed due to CdO on the CdS surface at high pH, consistent with previous reports.\([3, 25]\) Although Cd\(^0\) does not show Raman bands, Cd nanoparticles around 5 nm in size (as well as analogous Ag-based clusters) exhibit collective vibrations that give rise to appreciable bands in the low frequency region at 115 cm\(^{-1}\).\([26–29]\) Such a peak is observed after 2 min of irradiation using CdS/CdO QDs, which is believed to arise from Cd\(^0\) localised at the particle surface (Figure 2c, note that the low resolution of this peak is due to its location at the edge of the spectral window).\([21]\) At 5 and 10 min of irradiation, the peak is less pronounced, which is assigned to agglomeration after photocatalysis. This agrees with the observation of particle agglomerates in TEM images (Figure 2d).

Based on these experiments, the route to O\(_2\)-stabilisation in this system is summarised in Figure 2e. The consecutive processes are illustrated as (1) light absorption, (2) donor oxidation and (3) proton reduction. As reactions (1) and (2) are substantially faster than (3) in strongly alkaline conditions, excited electrons can accumulate on the particle surface in the form of Cd\(^0\) sites. Formation of Cd\(^0\) causes the QDs to agglomerate, which significantly lowers activity. O\(_2\) provides an easily reduced secondary substrate in the photoreactor that precludes the accumulation of Cd\(^0\) and thereby maintains the stability of the particle during photocatalysis. Note that this mechanism consumes a portion of electron donor without concomitant release of H\(_2\).

Taking this mechanism into account, a photocatalytic system was designed where the continued presence of O\(_2\) was exploited to stabilise the rate of photocatalysis. A constant flow of air was introduced into a photoreactor containing QDs in 5 mM KOH with EtOH (7.5 mM) as an electron donor and a cobalt co-catalyst. The vial was irradiated and the outlet gas-stream was continuously analysed by gas chromatography. The rate of H\(_2\) evolution reached a maximum activity of 432 mmol\(_{H_2}\) g\(^{-1}\) h\(^{-1}\), the highest reported rate for photocatalysis driven by an organic oxidation reaction on CdS under AM 1.5 G, 100 mW cm\(^{-2}\) irradiation (to the best of our knowledge higher activity has only been reported when using a S\(^2^-\) donor).\([19]\) Here, the constant influx of O\(_2\) was able to stabilise H\(_2\) production relative to an N\(_2\)-purged equivalent that operated at only 202 mmol\(_{H_2}\) g\(^{-1}\) h\(^{-1}\) (Figure 3a). MeOH-promoted H\(_2\) evolution was similarly high, exhibiting a rate of 165 mmol\(_{H_2}\) g\(^{-1}\) h\(^{-1}\) under air (Figure 3b).

In summary, the presented system demonstrates how photoredox reactions can be tuned to ensure discharging of potentially inhibiting deactivation reactions. This counter-intuitive strategy employs O\(_2\) to regenerate/stabilise a damaged...
photocatalyst and has achieved unmatched rates of proton reduction driven by photoreforming of an organic substrate. Consideration of aerobic strategies to benefit photocatalysis may therefore be vital in attaining both fast and stable systems in solar fuel and organic photoredox catalysis, where the presence of O₂ is often seen as a source of inhibition.

Acknowledgements

The Christian Doppler Research Association (Austrian Federal Ministry of Science, Research and Economy and the National Foundation for Research, Technology and Development), the OMV Group (E.R., M.F.K., D.W.W.), the EPSRC for a PhD studentship (D.W.W.) and Follow-on-Fund (M.F.K., E.R.), the World Premier Institute Research Center Initiative (WPI), MEXT, Japan (K.L.O.), the European Commission for a Marie Curie fellowship (K.H.L., GAN701192 - VSHER) and the Royal Society for a Newton Fellowship (N.K.). We would like to thank Prof. Peter Hildebrandt (TU Berlin, Germany) for providing access to his laboratory for Raman spectra. We thank Adam Schwartzberg (Lawrence Berkeley Laboratory) for assistance with TA experiments in the Molecular Foundry (supported by the Office of Science, Office of Basic Energy Sciences, of the US DOE under Contract DE-AC02-05CH11231).

Conflict of interest

The authors declare no conflict of interest.

Keywords: hydrogen · oxygen inhibition · oxygen tolerance · photocatalysis · quantum dots


Manuscript received: May 10, 2018
Accepted manuscript online: May 11, 2018
Version of record online: June 27, 2018