Kinetics of Photoelectrochemical Oxidation of Methanol on Hematite Photoanodes

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Supporting Information

ABSTRACT: The kinetics of photoelectrochemical (PEC) oxidation of methanol, as a model organic substrate, on α-Fe2O3 photoanodes are studied using photoinduced absorption spectroscopy and transient photocurrent measurements. Methanol is oxidized on α-Fe2O3 to formaldehyde with near unity Faradaic efficiency. A rate law analysis under quasi-steady-state conditions of PEC methanol oxidation indicates that rate of reaction is second order in the density of surface holes on hematite and independent of the applied potential. Analogous data on anatase TiO2 photoanodes indicate similar second-order kinetics for methanol oxidation with a second-order rate constant 2 orders of magnitude higher than that on α-Fe2O3. Kinetic isotope effect studies determine that the rate constant for methanol oxidation on α-Fe2O3 is retarded ~20-fold by H/D substitution. Employing these data, we propose a mechanism for methanol oxidation under 1 sun irradiation on these metal oxide surfaces and discuss the implications for the efficient PEC methanol oxidation to formaldehyde and concomitant hydrogen evolution.

INTRODUCTION

Electrochemical and photoelectrochemical (PEC) processes are widely used to drive organic oxidation reactions, with applications including molecular syntheses, photocatalytic pollutant destruction, and photoelectrochemical hydrogen generation (i.e., water splitting). For example, in PEC hydrogen generation, organic substrate oxidation can replace water oxidation as a source of electrons for proton reduction. In such systems, oxidation of sacrificial organic molecules has been shown to increase hydrogen generation yields, avoiding the kinetic limitations of water oxidation.1-3 Additionally, selective PEC oxidation of industrial byproducts can be used to synthesize higher value-added products. Different cases include the selective oxidation of glycerol to produce dihydroxyacetone,4 the synthesis of hydrogen and aldehydes or ketones from biomass,5 and epoxides from alkynes.6 For example, Yao and co-workers7 have recently reported a highly selective oxidation of various benzyl alcohols on H-titanate nanotubes. However, using photogenerated charge carriers to drive these processes can be challenging in terms of production yields and selectivity, often with only limited understanding of reaction mechanisms.7

In the particular case of PEC hydrogen evolution, organic substrates such as methanol or ethanol have been used to scavenge holes in metal oxide photoanodes such as TiO2,8,9 and α-Fe2O3.10 (Photo)electrochemical oxidation of methanol to formaldehyde has been reported, although with only a low Faradaic efficiency.11,12 With commercial formaldehyde synthesis from methanol primarily being achieved at high temperatures on iron molybdate catalysts,13 however, very little consideration has been given to the key kinetic and thermodynamic parameters controlling the rate-limiting step (RLS) of PEC oxidation of methanol and their implications in the system efficiency, such as rate and yield of reaction.

This paper focuses on methanol oxidation, as a model oxidation reaction, on a widely studied photoanode material, α-Fe2O3, and a comparison of the kinetics of this reaction on an alternative photoanode material, TiO2. The use of hole scavengers such as methanol has been shown to be an effective strategy to reduce electron/hole recombination losses in such metal oxides, as an alternative to the application of anodic potentials.10,14 Methanol oxidation studies on semiconductors such as ZnO and TiO2 have indicated a methanol adsorption process followed by the formation of a CH3O radical and its subsequent oxidation with a valence band hole. Other studies have provided evidence for a photocurrent doubling mechanism where methanol scavenging results in the formation of two long-lived conduction band electrons per scavenged hole. However, kinetic and mechanistic studies under operating conditions have received relatively little attention to date and are the subject of this paper.

In this study, we employ a rate law analysis to determine the key factors involved in the RLS of the methanol oxidation reaction (MOR). Such factors include different surfaces (metal oxides), the density of surface holes, and the kinetic isotope effect of deuterium (CD3OD). The approach, following that recently employed by Le Formal et al. for water oxidation,15 employs photoinduced absorption (PIA) spectroscopy to

Received: May 19, 2017
Published: July 22, 2017

DOI: 10.1021/jacs.7b05184
J. Am. Chem. Soc. 2017, 139, 11537–11543

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11537
determine the density of surface holes and correlates this density with transient photocurrent (TPC) measurements under quasi-steady-state conditions of PEC methanol oxidation using long (5 s) pulsed irradiation. We provide evidence, from rate law analyses, that the kinetics of the oxidation of methanol on \(\alpha\)-Fe\(_2\)O\(_3\) and TiO\(_2\) are independent of the band bending at the semiconductor–liquid junction but are instead sensitive to the choice of semiconductor and to the density of surface holes. The results presented herein allow us to propose a kinetic model and a plausible mechanism for the MOR on \(\alpha\)-Fe\(_2\)O\(_3\) that serves as a model for this oxidation reaction on such metal oxide photoanodes.

### EXPERIMENTAL SECTION

**Preparation of the Semiconductor Films.** Silicon-doped \(\alpha\)-Fe\(_2\)O\(_3\) photoanodes were prepared by atmospheric pressure chemical vapor deposition (APCVD), by a procedure detailed elsewhere.\(^{16}\) These nanostructured 400 nm thick \(\alpha\)-Fe\(_2\)O\(_3\) films show a dendritic structure with a feature size of 5–10 nm at the surface and roughness factor of 21.

Mesoporous TiO\(_2\) photoanodes were grown from a colloidal anatase paste, according to the method developed by Xiao-e et al.\(^{17}\) Mesoporous films were produced on FTO glass by doctor-blading the anatase using a k-bar followed by heat treatment at 450 °C in air for 30 min. These films were approximately 1 µm thick with crystallites of average ~40 nm wide and roughness factor of 120.\(^{14}\)

**Photoelectrochemical Setup.** A three-electrode cell was used for photoelectrochemical, photinduced absorption spectroscopy, and transient photocurrent measurements. The electrolyte solution contained typically 0.1 M NaOH and 4% (for TiO\(_2\)) and 95% (for \(\alpha\)-Fe\(_2\)O\(_3\)) volume methanol in deionized water. For the kinetic isotope effect (KIE) study, the electrolyte solution contained 95% CD\(_2\)OD in 0.1 M NaOD in D\(_2\)O. No concentrations higher than 98% methanol were tested due to insolubility of NaOH in such high methanol concentrations at room temperature. A Pt mesh was used as the counter electrode and a silver/silver chloride (Ag/AgCl) saturated with KCl (\(E^\circ\) = +0.197 V vs NHE) as the reference electrode. All potentials are reported against this Ag/AgCl electrode as the conversion of the potentials from Ag/AgCl to the reversible hydrogen electrode in highly concentrated organic aqueous solutions might not be accurate.

Linear sweep voltammograms were measured in the dark and under electrode–electrolyte (EE; front side) illumination conditions with a photon flux equivalent to approximately 100 mW cm\(^{-2}\) (1 sun) provided by two 365 nm light-emitting diodes (LEDs) (LZ1-10U600, LedEngin Inc.). The scan speed was 20 mV s\(^{-1}\), and the light was chopped at a frequency of 0.4 Hz.

**Optoelectronic Setup.** Photinduced absorption spectroscopy allows long-lived photogenerated species to be monitored under pseudo-steady-state conditions. The PIA signal is proportional to the density of holes accumulated at the surface. Simultaneously, the transient photocurrent signal is measured in the PEC cell, by converting the potential difference between the photoanode and the counter electrode (as measured across a 98.7 Ohm resistor) into current using Ohm’s law. Therefore, the TPC signal provides information on the extraction of electrons through the external circuit. The PIA and TPC signals were measured simultaneously for a 10 s period, with a 5 s on/5 s off of 365 nm LED pulse. The PIA signal was measured by registering the change in the optical density (absorption) of the hematite after excitation by the UV light. The light intensity of the LEDs was varied between 0.5 and 70 mW cm\(^{-2}\) by applying a fixed current (from 0.05 to 0.70 A). This is equivalent to a photon flux of 0.06–2.7 and 0.55–5.4 suns for hematite and anatase, respectively, as calculated by Ma et al.\(^{16}\) by integrating the solar power flux from the lowest limit of the measured solar spectrum to the typical absorption edge of hematite (600 nm) and anatase (380 nm).

### RESULTS

**Formaldehyde Quantiﬁcation.** Formaldehyde, as methanol oxidation product, was quantified by spectrophotometric measurements. A violet color is developed by reaction between formaldehyde and 4-amino-3-hydrazino-5-mercapto-1,2,4-triazole, 4-amino-5-hydrazino-1,2,4-triazole-3-thiol (Purpald) \(\geq99\%\), Sigma-Aldrich. A calibration curve was prepared from a concentration of 0 to 5 ppm of formaldehyde, ACS reagent, 37% W, Sigma-Aldrich, following a method developed by Jacobsen et al.\(^{19}\) The quantification was carried out in a PerkinElmer Lambda 25 spectrophotometer, measuring at 549 nm. The method is sensitive also to acetaldehyde, propionaldehyde, butyraldehyde, and benzaldehyde at different wavelengths; however, the only aldehyde expected from the oxidation of methanol is formaldehyde.

**Figure 1A** shows a typical current/potential (\(J−V\)) response of a nanostructured Si-doped hematite (APCVD \(\alpha\)-Fe\(_2\)O\(_3\)) in 0.1 M NaOH and in 95% methanol in 0.1 M NaOH (see Supporting Information (SI), Figure S1, for the photocurrent response on both \(\alpha\)-Fe\(_2\)O\(_3\) and TiO\(_2\) photoanodes, as a function of methanol concentration). Under simulated 1 sun illumination conditions, the photocurrent onset for the 95% methanol electrolyte, assigned below to the methanol oxidation reaction, requires approximately 270 mV less oxidative potential than that for water oxidation, consistent with previous studies.\(^{10,11,20}\)
Additionally, the oxidation of methanol produces 3.9 mA·cm⁻² photocurrent at strong anodic potential (0.55 V\textsubscript{Ag/AgCl}) compared to 2.6 mA·cm⁻² obtained from water oxidation at the same applied potential. Both the shift in the onset potential and enhancement in the plateau photocurrent are likely due to the more facile oxidation of methanol.\textsuperscript{20}

Figure 1B presents comparative \(J-V\) curves under chopped illumination for the oxidation of methanol on nanocrystalline \(\alpha\)-Fe\textsubscript{2}O\textsubscript{3} versus that on mesoporous anatase TiO\textsubscript{2}. At high applied potentials, methanol oxidation on hematite produces an order of magnitude more photocurrent than on anatase, most likely due to better light absorption by hematite relative to titania (band gaps of 2.1 and 3.1 eV, respectively). On the other hand, titania shows a photocurrent onset approximately 500 mV cathodic of that for hematite, in accordance with their difference in the valence band edge (2.6 and 2.1 V\textsubscript{NHE} at pH 14, respectively).\textsuperscript{21,22}

The product of methanol oxidation on hematite was determined by spectrophotometric titration. Formaldehyde was formed with a 96% Faradaic efficiency, as shown in Figure 2 (see Figures S2 and S3 for further details on the bulk electrolysis and the calibration curve). This high Faradaic efficiency indicates that formaldehyde is not further oxidized to formic acid or carbon dioxide under these experimental conditions. A general equation corresponding to the oxidation can be written as follows:

\[
\text{CH}_3\text{OH} + 2h^+ \rightarrow \text{H}_2\text{CO} + 2\text{H}^+ 
\]

where \(h^+\) represents a surface hole on the photoanode. The strikingly high Faradaic efficiency of this reaction on hematite compared to that of an electrochemical route on Pt (81%)\textsuperscript{23} and a photochemical process on TiO\textsubscript{2}, in methanol concentrations under 1% (30%),\textsuperscript{11} makes this an appealing route for PEC formaldehyde synthesis.

In order to analyze the kinetics of methanol oxidation on our hematite and titania photoanodes, PIA spectroscopy and TPC measurements were conducted employing variable intensities for a duration of 5 s at 365 nm, as detailed in the Experimental Section. In these studies, the PIA signal is employed to monitor the absorbance and therefore the density of long-lived photogenerated holes, while the photocurrent density monitors the net flux of holes transferred to the electrolyte in quasi-steady-state conditions (i.e., the rate of methanol oxidation). This approach follows that previously reported by Le Formal et al., where we have demonstrated that this PIA approach allows us specifically to probe the accumulation of long-lived holes within the depletion region at the photoanode surface and therefore assay the surface density of holes driving surface electrochemistry.\textsuperscript{15} For the PIA data, probe wavelengths of 650 and 500 nm were employed for hematite and titania photoanodes, respectively, corresponding to their valence band hole photoinduced absorption maxima.\textsuperscript{9,15}

Figure 3 shows the PIA (Figure 3A) and TPC (Figure 3B) responses for the oxidation of 95% methanol in 0.1 M NaOH on \(\alpha\)-Fe\textsubscript{2}O\textsubscript{3} with the photoanode held at 0.55 V\textsubscript{Ag/AgCl}. This strongly anodic applied potential minimizes recombination in the photoanode.\textsuperscript{24,25} The PIA signal measured at 650 nm and (b) transient photocurrent measured simultaneously.
Further data analogous to those shown in Figure 3 were collected at 0.00 V \text{Ag/AgCl} (see Supporting Information Figure S4A,B). Under these modest potential conditions, the attenuated space charge layer width results in less band bending (as the space charge layer is smaller than the particle size in these APCVD α-Fe2O3 films).27 As expected, due to the resulting more severe recombination losses, higher light intensities were required to generate PIA and photocurrent signals comparable to those obtained at 0.55 V \text{Ag/AgCl}. Confirming this, Figure S5 shows that at low applied potentials, back electron/hole recombination is not completely turned off and accelerates the decay kinetics of the holes accumulated at the surface of the photoanode. Furthermore, an analogous study was conducted using TiO2 as the photoanode in 4% methanol in 0.1 M NaOH electrolyte (see Supporting Information Figure S6A,B). A lower methanol concentration was used as the photocurrent increases following methanol addition saturating at this concentration of 4% (see Figure S1). This enabled us also to monitor the titania surface hole accumulation under conditions of quasi-steady-state methanol oxidation. Before undertaking a quantitative comparison of these data, we first present the kinetic model used for their analysis.

Based on Le Formal et al.,15 we turn now to a simple kinetic model for the PEC oxidation of methanol on the photoanodes studied herein. The model is considered under steady-state conditions when the change in surface hole density, \(dp_\text{sur}/dt\), with the time, \(dt\), is zero (see eq 2), and the flux of photogenerated holes to the surface, \(J_\text{fl} \approx \text{Jp} + \text{kobs} \cdot \text{p}^{\alpha}\), is equivalent to the photocurrent, \(J_\text{V}\) (see eq 3).

\[
\frac{dp_\text{sur}}{dt} = 0 = \text{Jfl} = \text{kobs} \cdot \text{p}^{\alpha}
\]  

\[
\log(J_\text{V}) = \alpha \cdot \log(p_\text{obs}) + \log(k_{\text{obs}})
\]  

where \(k_{\text{obs}}\) is the observed rate constant for methanol oxidation and \(\alpha\) is the order of the methanol oxidation reaction with respect to the density of surface accumulated holes, \(p_\text{obs}\). As such, a plot of \(\log(J_\text{V})\) versus \(\log(p_\text{obs})\) will have a gradient equivalent to the reaction order \(\alpha\), \(J_\text{V}\) can be determined from the current densities (e.g., Figure 3B) measured at 5 s after light on (i.e., quasi-steady-state conditions). The surface density of holes, \(p_\text{obs}\), can be determined at the same time from the PIA (e.g., Figure 3A) using the Beer–Lambert law from measured hole extinction coefficients at the probe wavelengths used (640 \text{M}^{-1} \text{cm}^{-1} for α-Fe2O3,15 and 2000 \text{M}^{-1} \text{cm}^{-1} for TiO2).28

Figure 4 shows plots of the photocurrent density, \(J_\text{V}\), as a function of the surface hole density, \(p_\text{obs}\), for the oxidation of methanol on TiO2 and α-Fe2O3, employing the PIA and TPC data shown in Figure 3 and Figures S4 and S6. The data are plotted in units of \text{nm}^{-2}, correcting for surface roughness of the two electrodes. For all data sets, the gradients of \(\log(J_\text{V})\) versus \(\log(p_\text{obs})\) are ~2 (within the range 1.88 to 2.13; see Figure 4), indicating that in all cases the oxidation of methanol is second order with respect to surface-accumulated holes. This second-order behavior is further supported by an initial rates law analysis (see Figure S7) of the PIA decay kinetics for methanol oxidation on hematite at 0.55 V \text{Ag/AgCl}. From eq 3, we obtain second-order rate constants of 15 000 and 33 holes nm\(^{-2}\) s\(^{-1}\) for TiO2 and α-Fe2O3, respectively, independent of the applied potential. For α-Fe2O3 at 0.55 V \text{Ag/AgCl} under conditions of approximately 1 sun irradiation (~4 mA cm\(^{-2}\)), \(p_\text{obs}\) is ~0.5 holes nm\(^{-2}\), leading to a hole flux to the electrolyte of ~10 holes nm\(^{-2}\) s\(^{-1}\), corresponding to a "turnover frequency" per hole of ~20 s\(^{-1}\). We further note that we obtain indistinguishable rate constants and rate laws for methanol oxidation on hematite at 0.00 and 0.55 V \text{Ag/AgCl} despite the large difference in band bending and recombination losses between these two conditions; a point we discuss in further detail below.

We undertook a kinetic isotope effect study to further analyze the kinetics and the second-order dependence of the reaction with respect to the density of accumulated holes at the surface. Therefore, we collected data analogous to that shown in Figure 3 (see Supporting Information Figure S8A,B) using deuterated 95% methanol-\text{d}_4 in 0.1 M NaOD in D\(_2\)O as electrolyte. Figure 5 shows the resulting rate law analysis comparing the oxidation of CH\(_3\)OH versus CD\(_3\)OD on α-Fe2O3. For the methanol-\text{d}_4 electrolyte the gradient of \(\log(J_\text{V})\) versus \(\log(p_\text{obs})\) is also ~2 but showed a 20-fold reduction in current, compared with CH\(_3\)OH, at equivalent surface hole densities.
densities. The corresponding second-order rate constant for CD$_3$OD oxidation (1.35 holes$^{-1}$ nm$^{-2}$ s$^{-1}$) gives a KIE of $\sim$20, decreasing the turnover frequency per hole from $\sim$20 to $\sim$1 s$^{-1}$ under conditions of approximately 1 sun irradiation. These slower kinetics for CD$_3$OD oxidation indicate that the rate-limiting step of the reaction is a chemical step and involves the breaking of a C–H bond, as we discuss further below.

## DISCUSSION

We have shown that, under operating photoelectrochemical oxidation conditions, methanol is fully oxidized to formaldehyde on both TiO$_2$ and α-Fe$_2$O$_3$ with a rate that depends on the square of the density of surface-accumulated holes, by quasi-steady-state kinetic analysis of the reaction. We note this result differs from our analysis of water oxidation on identical α-Fe$_2$O$_3$ (see Figure S9), where we observed first-order behavior with respect to $p_h$ at low surface hole densities transitioning to third-order behavior at high hole densities. More significantly, the kinetics of the methanol oxidation reaction depend upon the metal oxide surface chemistry, as well as the presence of deuterium in the electrolyte, but are independent of the applied potential.

We first focus on the hematite data collected at two different applied potentials. It is striking that our plots of $j_F$ versus $p_h$ collected at 0.00 and 0.55 V$_{Ag/AgCl}$ overlap each other, showing the same reaction order and rate constant with respect to surface hole density. These reaction conditions are very different, with severe surface electron/hole recombination losses at 0.00 V$_{Ag/AgCl}$ but no surface recombination at 0.55 V$_{Ag/AgCl}$. Although water oxidation does not occur at 0.00 V$_{Ag/AgCl}$, it is possible that it becomes competitive with methanol oxidation at high applied potentials; however, this is ruled out by Supporting Information Figure S9. The agreement between 0.00 and 0.55 V$_{Ag/AgCl}$ data confirms the validity of our experimental protocol and that our analysis does indeed address the kinetics of methanol oxidation at the semiconductor/electrolyte interface. We observe that at equivalent surface hole densities, the kinetics of methanol oxidation are independent of applied potential, clearly demonstrating that the kinetics of the reaction are not determined by the electrode Fermi level or band bending. Rather this observation indicates that these kinetics are simply determined by the density of holes accumulated at the electrode surface, with an energy determined by the valence band edge. This situation is consistent with the semiconducting nature of hematite and contrasts the behavior of metal electrodes, where changing the applied potential changes the free energy driving the reaction.

Turning now to the comparison of titania and hematite shown in Figure 4, it is apparent that titania shows methanol oxidation kinetics $\sim$500-fold faster than that of hematite, despite the lower “photocurrent-saturating” methanol concentration (4% in titania compared to 95% in hematite, as shown in Figure S1). This difference in the concentration of methanol needed to reach the maximum photocurrent densities has been suggested to depend on a competitive mechanism of adsorption between water and methanol on anatase, compared to hematite where a strong chemisorption of methanol (as methoxide) has been reported. This difference in concentration dependence may also be related to the lower Faradaic efficiencies reported for methanol oxidation on TiO$_2$ (e.g., 30% reported by Wahl et al. for the oxidation of 0.4% methanol in 0.1 M NaOH). Despite these differences, it is striking that methanol oxidation on titania also exhibits second-order behavior as a function of surface hole density, suggesting some similarity in the reaction mechanism. A full analysis of methanol oxidation on titania and its dependence on, for example, methanol concentration is beyond the scope of this study. Nevertheless, it is clear that methanol oxidation on titania is at least 2 orders of magnitude faster than that on hematite. These faster kinetics can be most obviously assigned to the deeper valence band edge of TiO$_2$ relative to α-Fe$_2$O$_3$ (2.6 and 2.1 V$_{NHE}$ at pH 14, respectively), providing a larger energy offset to drive the methanol oxidation reaction, although we note that differences in methanol surface adsorption may also be important.

A two-step methanol oxidation mechanism on oxide surfaces has been proposed previously in the context of observations of photocurrent doubling for similar systems. However, we note that Schoenmakers et al. have reported a reduction in the quantum efficiency of methanol oxidation on ZnO from 2 to 1 when increasing the light intensity by $\sim$0.005 to 5 suns. The results we report here provide an explanation for Schoenmakers et al.’s observation and indicate that under $\sim$1 sun irradiation conditions, where there is significant hole accumulation at the oxide surface, both steps of methanol oxidation are driven by photogenerated valence band holes. These results in the observed second-order dependence on surface hole density and no significant current doubling effect are in agreement with previous reports operating at $\sim$1 sun conditions.

We finally focus on the KIE shown in Figure 5. The aforementioned methanol adsorption on the photoanode surface leads to the loss of the H atom from the O–H bond. Therefore, our observation of a KIE of $\sim$20 is most obviously assigned to a C–H bond breaking in the rate-limiting step of methanol oxidation. We note that this would be an unusually high KIE value for a C–H bond breaking considering only the stretching mode of this bond (KIE$\text{expected} \sim 7$). However, C–H bond breaking of surface-bound CH$_2$ species will be associated with significant structural changes and specifically a change in the carbon hybridization from sp$^3$ to sp$^2$. This rehybridization can be expected to lead to considerable differences in the zero-point energy of the transition state due to the loss of stretching as well as bending modes. Therefore, we conclude that the rate-limiting step in the oxidation of methanol under conditions of $\sim$1 sun illumination and alkaline electrolyte involves the breaking of a C–H bond and an associated rehybridization of the carbon.

Based upon these data and previous literature studies, we propose a mechanism for methanol oxidation on hematite and titania photoanodes, as shown in Figure 6. We note that Grassian and co-workers have determined the “saturating” methanol-adsorbed surface density on α-Fe$_2$O$_3$ to be $\sim$2 $\times$ 10$^{13}$ molecules-cm$^{-2}$, corresponding to one molecule of methanol adsorbed every 5 nm$^2$ on α-Fe$_2$O$_3$, which is comparable to our measured surface hole densities.
species; methanol oxidation requires two of these species to react with surface valence band holes, which have been assigned previously to Fe(IV) species on hematite.23,31,38 Surface valence band hole density has been reported to be significant at Fe(IV) species on hematite.11,23,31,38 Surface valence band holes (Fe(IV) species on hematite) have been reported previously to be significant under conditions of 1 sun illumination. Our observation of second-order behavior also has important implications for technological applications, as it implies that the kinetics and therefore potentially the efficiency of this widely used reaction will be superlinearly dependent upon the surface density of accumulated holes, with implications for photoanode design (e.g., surface area) and optimum operational light intensities.

## CONCLUSIONS

The use of organic oxidation substrates can substantially reduce the requirement of strong anodic potentials for PEC hydrogen evolution, as well as potentially enable the synthesis of useful organic compounds. As a study case, we have reported kinetic and mechanistic analyses for the selective oxidation of a model substrate, methanol to formaldehyde, on titania and hematite photoanodes under PEC working conditions. The methanol oxidation reaction was found to be second order with respect to the density of surface holes, indicating a reaction mechanism where both steps of methanol oxidation are driven by valence band holes. The second oxidation, involving a C–H bond breaking, is the rate-limiting step. Remarkably, this oxidation is observed to proceed with near unity Faradaic efficiency, suggesting a potentially attractive route to formaldehyde synthesis from methanol. Our observation of second-order behavior also has important implications for technological applications, as it implies that the kinetics and therefore potentially the efficiency of this widely used reaction will be superlinearly dependent upon the surface density of accumulated holes, with implications for photoanode design (e.g., surface area) and optimum operational light intensities.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.7b05184.

Current–voltage curves for different concentrations of methanol in 0.1 M NaOH on α-Fe2O3 and TiO2; formaldehyde quantification details; PIA and TPC signals measured at 0.00 V for hematite and –0.80 V for anatase, as well as PIA and TPC signals for the oxidation of CD3OD on hematite at 0.55 V; effect of recombination on PIA decays, an initial rates analysis for methanol oxidation, and a comparison of rate law analysis for water and methanol oxidation (PDF). 

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#### Notes

The authors declare no competing financial interest.

#### ACKNOWLEDGMENTS

We acknowledge Dr. Robert Godin for helpful discussions and financial support from the European Research Council (project Intersolar 291482), Swiss National Science Foundation (project 140709) and Swiss Federal Office for Energy (project PECHouse 3, contract number SI/500090-03). C.A.M. thanks COLCIENCIAS for funding; L.F. thanks the EU for a Marie Curie fellowship (658270), and E.P. thanks the EPRSC for a DTP scholarship.

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