# Chemical Science



# **EDGE ARTICLE**

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# Water oxidation kinetics of nanoporous BiVO<sub>4</sub> photoanodes functionalised with nickel/iron oxyhydroxide electrocatalysts†

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In this work, spectroelectrochemical techniques are employed to analyse the catalytic water oxidation performance of a series of three nickel/iron oxyhydroxide electrocatalysts deposited on FTO and BiVO<sub>4</sub>, at neutral pH. Similar electrochemical water oxidation performance is observed for each of the FeOOH, Ni(Fe)OOH and FeOOHNiOOH electrocatalysts studied, which is found to result from a balance between degree of charge accumulation and rate of water oxidation. Once added onto BiVO<sub>4</sub> photoanodes, a large enhancement in the water oxidation photoelectrochemical performance is observed in comparison to the un-modified BiVO<sub>4</sub>. To understand the origin of this enhancement, the films were evaluated through time-resolved optical spectroscopic techniques, allowing comparisons between electrochemical and photoelectrochemical water oxidation. For all three catalysts, fast hole transfer from BiVO<sub>4</sub> to the catalyst is observed in the transient absorption data. Using operando photoinduced absorption measurements, we find that water oxidation is driven by oxidised states within the catalyst layer, following hole transfer from BiVO<sub>4</sub>. This charge transfer is correlated with a suppression of recombination losses which result in remarkably enhanced water oxidation performance relative to unmodified BiVO<sub>4</sub>. Moreover, despite similar electrocatalytic behaviour of all three electrocatalysts, we show that variations in water oxidation performance observed among the BiVO<sub>4</sub>/MOOH photoanodes stem from differences in photoelectrochemical and electrochemical charge accumulation in the catalyst layers. Under illumination, the amount of accumulated charge in the catalyst is driven by the injection of photogenerated holes from BiVO<sub>4</sub>, which is further affected by the recombination loss at the BiVO<sub>4</sub>/ MOOH interface, and thus leads to deviations from their behaviour as standalone electrocatalysts.

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## Introduction

Artificial photosynthesis employing photoelectrochemical or photocatalytic systems is attracting increasing interest for the sustainable synthesis of molecular fuels and chemicals.<sup>1-4</sup> Metal

oxides are widely studied as light absorbing materials for such solar-to-fuel systems, particularly for the challenging water oxidation reaction.5,6 This interest is derived particularly from their stability under the highly oxidising conditions required to drive this reaction. However, the performance of such metal oxide-based photoelectrodes is generally limited by recombination losses; in particular, competing surface recombination losses (also known as back electron/hole recombination) due to the slow rate of water oxidation, can be very significant.<sup>7-12</sup> One widely used approach to reduce such losses is the addition of water oxidation catalyst layers on the photoanode surface, which can lead to an enhancement in performance. 12-16 However, the selection of suitable catalysts is challenging as the requirements for efficient photoanode performance is likely to be different from the requirements for efficient, dark electrocatalysis. This is because, unlike electrocatalytic systems, driven by a voltage source, photoanodes utilise photogenerated holes, which are subject to electron-hole recombination losses at the photoanode/electrocatalyst/electrolyte interfaces, substantially affecting water oxidation currents. 11,13,17-19 Thus, even for a well-

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understood water oxidation catalyst used as an electrocatalyst, understanding the catalyst–photoanode interaction is critical to enhance photocurrent yields in metal oxide/electrocatalyst photoanodes.

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Several earth abundant and inexpensive electrocatalysts have garnered extensive attention for water oxidation, such as cobalt based CoPi,20 and Co-Fe Prussian Blue.14 Transition metal oxyhydroxides (referred to herein as MOOH) like NiOOH and FeOOH have also been coupled with BiVO₄ and some of these combinations have resulted in the construction of state-of-theart BiVO<sub>4</sub> photoanodes, 13,15,21 exhibiting remarkable photocurrent densities of 5.87 mA cm<sup>-2</sup> at 1.23 V<sub>RHE</sub>. However, while the presence of MOOH considerably enhances the water oxidation performance of BiVO<sub>4</sub>, most notably for nanoporous electrodes, the origin of this enhancement has not been clearly elucidated to date. Crucially, it has also been observed that the performance enhancement of the BiVO<sub>4</sub>/MOOH system using either FeOOH or NiOOH for water oxidation does not correlate with the performance of FeOOH or NiOOH, respectively, when used as electrocatalysts on FTO.13 This discrepancy complicates the optimisation of photoelectrochemical devices and highlights the need to improve our understanding of photoanode/ electrocatalyst interface.13

In this study, transient optical and spectroelectrochemical techniques are utilised to: (1) compare performances of FeOOH, NiOOH, FeOOHNiOOH in terms of when they are used as an electrocatalyst on a conducting substrate in neutral electrolyte; and (2) investigate how their performance changes when they are placed on BiVO4 nanoporous photoanodes and need to utilise photogenerated holes for water oxidation under the same electrolyte conditions. We employ a population model to study the kinetics and the charge accumulation of our systems as it offers a direct pathway to compare electrocatalysts and photoelectrodes, and highlights the role of charge accumulation in driving reaction kinetics.22,23 This model has previously been employed to study water oxidation,24-28 proton reduction,29 and organic oxidations30 across electrocatalysts and photoelectrodes. The results obtained from this study provide insight into the role of MOOH on BiVO4 in enhancing water oxidation photocurrent and in the performance differences of the three MOOH catalysts on a conducting substrate and on a BiVO<sub>4</sub> photoelectrode.

## Results and discussion

Nanoporous BiVO<sub>4</sub> photoanodes used in this study were prepared using previously reported methods with one modification (see ESI† for details), <sup>13</sup> which was decreasing the film thickness (from  $\sim$ 750 to  $\sim$ 450–500 nm). This was necessary to increase the transparency of the BiVO<sub>4</sub> photoanodes, which is critical for the spectroscopic methods used in this study. As a result, the photocurrent obtained in this study with thinner BiVO<sub>4</sub> photoanodes is lower than that previously reported. <sup>13</sup> However, this decrease did not affect our comparison of the interactions of BiVO<sub>4</sub> with various MOOH electrocatalysts.

The FeOOH, NiOOH, FeOOHNiOOH electrocatalysts were synthesised as previously reported by anodic deposition on FTO

to examine electrocatalytic function<sup>27</sup> and on BiVO<sub>4</sub> to examine photoelectrocatalytic function.<sup>13</sup> The trend in the photoelectrochemical enhancements between the three electrocatalysts upon deposition on nanoporous BiVO<sub>4</sub> has previously been reported.<sup>13</sup> The thickness of the thinnest possible NiOOH layer that could be reproducibly deposited on FTO with a uniform coverage was ~80 nm. Therefore, the same thickness was used to prepare FeOOH on FTO. The thickness of FeOOH-NiOOH, which was prepared by sequential deposition of FeOOH and NiOOH layers, was not ~160 nm, but ~120 nm.27 This is because the underlying FeOOH is not dense but porous, allowing for the facile infiltration of NiOOH into the FeOOH layer and resulting in an atomic-level mixing of FeOOH and NiOOH.27 The thickness of the catalyst layer on BiVO4 was <5 nm for FeOOH or NiOOH and <10 nm for FeOOHNiOOH. These thicknesses were optimised to maximise photocurrent generation of BiVO4 prepared as thin photoelectrodes suitable for transient absorption spectroscopy studies. (The use of a catalyst layer that is thicker than the optimal thickness on a photoelectrode results in an increase in electron-hole recombination and is not favourable for photocurrent generation.13) We note that it would have been more ideal if the catalyst layers of the same thickness could be used on FTO and BiVO<sub>4</sub>. However, the thickness of the catalysts on FTO cannot be further reduced, and using thick catalyst layers on BiVO<sub>4</sub> for the sake of using the same catalyst thickness makes it difficult to study optimal BiVO<sub>4</sub>-catalyst interactions. Thus, we chose to use catalysts with an optimised thickness on BiVO4 to obtain a more meaningful and useful understanding of the system. We confirmed using electrochemical impedance spectroscopy that the catalysts on FTO are still thin enough not to crease a serious resistivity issue (see ESI† for details). We note that NiOOH in this study was deposited, as previously, in an electrolyte that was not rigorously free of Fe impurities, and the NiOOH contained a trace amount of Fe (~1 at%).27 Thus, hereafter, NiOOH will be denoted as Ni(Fe)OOH.

All investigations presented herein were performed at pH 7 as the current work aims to better understand the high performance of BiVO<sub>4</sub>/MOOH photoanodes reported previously at this pH.<sup>13</sup> While MOOH electrocatalysts are typically used in basic conditions, MOOH-coupled photoanodes that are not stable in strongly basic conditions, such as BiVO<sub>4</sub>, are often used in (near) neutral conditions to optimise system stability.

At pH 7, the three MOOH electrocatalysts studied herein exhibit similar electrochemical water oxidation performances, as can be seen in the steady-state current voltage (*J-V*) plots shown in Fig. 1a. The similar overall performance at pH 7 differs from that observed under alkaline media (pH 13), where the catalytic onset potential for FeOOH is anodically shifted by 70 mV compared with Ni(Fe)OOH and FeOOHNiOOH.<sup>27</sup> All the presented studies for NiOOH and FeOOHNiOOH and their photoanode counterparts were obtained after the *in situ* regeneration of NiOOH from Ni(OH)<sub>2</sub>, using an activation procedure detailed previously,<sup>27</sup> as Ni(OH)<sub>2</sub> forms over time when the electrodes are not in use (see ESI† for details). This activation process is referred to herein as the (0/+) oxidation process.

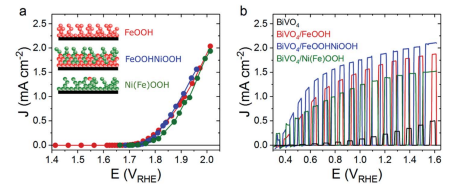


Fig. 1 Electrochemical and photoelectrochemical performance of FeOOH, FeOOHNiOOH and Ni(Fe)OOH as electrocatalyst and when functionalised on BiVO<sub>4</sub> photoanodes at pH 7. (a) Steady state J-V curves for FeOOH (red), FeOOHNiOOH (blue) and Ni(Fe)OOH (green); these were recorded by measuring the steady state current at different applied potential under atmospheric conditions. Inset: schematic representation of the different electrocatalyst samples studied in this work. Red represents a Fe-based group, and green represents a Ni-based group. (b) Linear Sweep Voltammetry (LSV) under chopped light (1 sun equivalent, 365 nm LED, back irradiation) for BiVO<sub>4</sub>/FeOOH (red), BiVO<sub>4</sub>/FeOOHNiOOH (blue), BiVO<sub>4</sub>/Ni(Fe)OOH (green) and un-modified BiVO<sub>4</sub> (black). All measurements were conducted in 0.1 M phosphate buffer. The J-V curves were recorded using a 10 mV s<sup>-1</sup> scan rate.

When these catalysts are deposited on top of BiVO $_4$  photo-anodes remarkable enhancements in the photoelectrochemical water oxidation performance are observed in comparison to the unmodified nanoporous BiVO $_4$  photoanode (Fig. 1b). To understand the origin of this performance enhancement upon MOOH addition, transient optical and spectroelectrochemical techniques were employed to compare the water oxidation and recombination kinetics of the Ni/Fe oxyhydroxide water oxidation electrocatalysts when used either as electrocatalyst or when deposited on BiVO $_4$  photoanodes.

Firstly, the optical and kinetic characteristics of the electrocatalysts when deposited on FTO were analysed. The optical changes during the sample activation (0/+) oxidation process to convert Ni(OH)<sub>2</sub> to NiOOH for both Ni(Fe)OOH and FeOOH-NiOOH electrocatalyst samples are shown in Fig. S2.† $^{27,31-33}$  It is observed that Ni(Fe)OOH accumulated more (0/+) compared with FeOOHNiOOH. Since Ni(Fe)OOH and FeOOHNiOOH both contain similar amounts of NiOOH, $^{27}$  this result suggests that as the Fe content increases, the oxidation of Ni(OH)<sub>2</sub> to NiOOH becomes more difficult. This is consistent with previous literature where the shift in the oxidation of Ni(OH)<sub>2</sub>/NiOOH to more anodic potentials has been correlated with the increase in Fe content. $^{32,34}$ 

Fig. 2a shows the optical response of the electrocatalysts. At applied potentials greater than the water oxidation catalytic onset, growth of new spectral features can be observed for each of the electrocatalysts, as shown in Fig. 2a (and Fig. S3† for spectra as a function of applied potential). These spectral features are assigned to the generation of oxidised states that accumulate under water oxidation catalysis, referred to as MOOH(++) herein. As can be observed in Fig. 2a (blue and red traces), the spectra of the FeOOH(++) and the FeOOH-NiOOH(++) species are almost indistinguishable, with an absorption maximum between 400–425 nm, suggestive of the similar nature of these oxidised states. In contrast, the Ni(Fe) OOH(++) species shown in Fig. 2a (green trace) exhibit a peak

centred at  $\sim$ 650 nm, indicative of a different nature of the oxidised state that forms in Ni(Fe)OOH. A similar trend for the respective spectral features of the MOOH(+) and MOOH(++) species of these three electrocatalysts, is also present at pH 13,  $^{27}$  suggesting that the catalytically active species observed under neutral and alkaline conditions are similar in nature. It is worth noting that the Ni(Fe)OOH samples were found to be relatively unstable when potentials greater than 2  $V_{\rm RHE}$  were applied, leading to a loss in the optical signal and performance. This effect can be associated with the gradual dissolution of NiOOH at pH 7 under anodic bias required for water oxidation (see ESI for more details, Fig. S5 and S6†).  $^{35}$ 

To allow quantitative comparison of the oxidation of these electrocatalysts and the associated water oxidation kinetics, the extinction coefficients ( $\varepsilon$ ) of the MOOH(++) oxidised species for each of the catalysts shown in Fig. 2a were estimated using steppotential spectroelectrochemistry (SP-SEC), (see Section D in ESI† for description of the method and additional data). Extinction coefficient ( $\varepsilon$ ) values of  $\sim$ 3500 and  $\sim$ 3100 M<sup>-1</sup> cm<sup>-1</sup> were obtained at wavelengths of 550 nm for FeOOH and FeOOHNiOOH, and a value of  $\sim$ 3100 M<sup>-1</sup> cm<sup>-1</sup> was obtained at 650 nm for Ni(Fe)OOH. Using the estimated values of  $\varepsilon$  for each of the samples, the optical signal ( $\Delta$ OD) shown in Fig. S3† can be correlated with the density of accumulated MOOH(++) species as a function of applied potential (Fig. S4†). When combined with current densities, these data yield kinetic information on the water oxidation process, allowing the estimation of reaction time constant  $(\tau)$  or turn over frequency (TOF) for O<sub>2</sub> evolution, as a function of MOOH(++) species. These analyses have been described in Section G in the ESI† and have been reported previously.27 The water oxidation kinetic data are plotted as a function of MOOH(++) accumulated species (Fig. 2b) and also as a function of applied potential (Fig. 2c).

As can be observed in Fig. 2b (empty circles, bottom panel), the density of the MOOH(++) accumulated species increases

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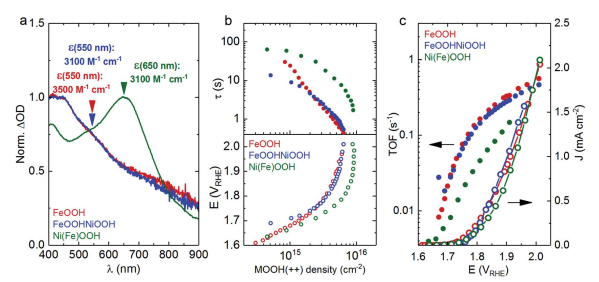


Fig. 2 Kinetics of water oxidation under electrochemical conditions by the FeOOH, FeOOHNiOOH and Ni(Fe)OOH electrocatalysts. (a) Normalised  $\Delta$ OD spectra of the MOOH(++) species and their corresponding extinction coefficient for FeOOH (red), FeOOHNiOOH (blue) and Ni(Fe)OOH (green). The spectra are obtained by subtracting the spectra after activation from the spectrum at 1.81 V<sub>RHE</sub>. (b) Water oxidation reaction time constants ( $\tau$ , solid circles, top panel) and the applied potential (bottom panel) as a function of the accumulated MOOH(++) density. (c) The TOF for O<sub>2</sub> evolution (solid circles, left axis) against the applied potential, plotted together with the current density (open circles, right axis). All measurements were taken in 0.1 M phosphate buffer (pH 7). Same colours used in (a), (b) and (c).

with the applied potential for each of the electrocatalysts, resulting in an acceleration of the water oxidation  $\tau$  by two orders of magnitude (solid circles, top panel). For a given potential, it is also observed that Ni(Fe)OOH accumulates more MOOH(++) species than FeOOH and FeOOHNiOOH. Despite the difference in thickness between the FeOOH and FeOOH-NiOOH electrodes, both electrocatalyst electrodes display negligible differences in charge accumulation densities during catalysis. Whilst it is apparent that both FeOOH and FeOOH-NiOOH exhibit similar  $\tau$ , Ni(Fe)OOH(++) states oxidise water circa an order of magnitude more slowly, when compared at equivalent densities of MOOH(++) species. The slower water oxidation kinetics of Ni(Fe)OOH(++) is also reflected in the plot of TOF (s<sup>-1</sup>) vs. applied potential (Fig. 2c) which shows that at a given potential, FeOOH(++) and FeOOHNiOOH(++) both oxidise water at a similar rate, whereas the TOF (s<sup>-1</sup>) for Ni(Fe) OOH(++) is approximately halved. Overall, while Ni(Fe)OOH(++) is able to accumulate a greater density of MOOH(++) species than FeOOH and FeOOHNiOOH, this gain is offset by slower water oxidation kinetics. As a result, the electrocatalytic performance for water oxidation is relatively invariant for all three electrocatalyst as shown in Fig. 1a and 2c. These trends are markedly different to those observed at pH 13 where Ni(Fe) OOH shows the fastest water oxidation kinetics amongst the three and, in comparison to FeOOHNiOOH, FeOOH starts accumulating charge at more anodic potentials, consistent with a later catalytic onset.<sup>27</sup> However the origin of these differences with pH is beyond the scope of this study.

Next we look at the interaction between these electrocatalyst and BiVO<sub>4</sub> photoanodes. The poor performance of the unmodified nanoporous BiVO<sub>4</sub> has previously been assigned to

holes accumulating at the BiVO<sub>4</sub> surface undergoing extensive losses due to water oxidation kinetics on BiVO<sub>4</sub> being slower than surface recombination.<sup>13</sup> However it was not clear if the performance enhancement observed after addition of MOOH results primarily from faster water oxidation kinetics or slower surface recombination kinetics.

The enhancement observed in Fig. 1b is consistent with that reported previously for these BiVO<sub>4</sub>/MOOH combinations<sup>13,15</sup> and is much greater than that typically found when electrocatalysts (e.g. CoPi) are deposited on dense planar BiVO<sub>4</sub> electrodes;11,36 such comparisons between CoPi and MOOH electrocatalysts are discussed later. Interestingly, while the three MOOH electrocatalysts show comparable performances, the BiVO<sub>4</sub>/MOOH photoelectrodes exhibit differences in their water oxidation photocurrents (Fig. 1b). In order to unravel the origin of the differences in performance enhancement upon the addition of MOOH electrocatalysts to BiVO<sub>4</sub>, two different timeresolved optical and photoelectrochemical techniques are employed: transient absorption spectroscopy (TAS) employing short laser pulse excitation, and spectroelectrochemical photoinduced absorption spectroscopy (PIAS), employing quasisteady state LED excitation.

To investigate if charge transfer occurs following bandgap excitation of  $BiVO_4$ , TAS was employed under an applied potential of  $1.4~V_{\rm RHE}$ . Fig. 3a and b presents the TA spectra for  $BiVO_4$  and  $BiVO_4/Ni(Fe)OOH$  and a comparison of the normalised charge carrier kinetics between un-modified  $BiVO_4$  and the  $BiVO_4/Ni(Fe)OOH$  photoanodes. Equivalent TA spectra for the other  $BiVO_4/MOOH$  configurations are included in the ESI (Fig. S8†). The data for  $BiVO_4$  are similar to that reported previously, and are associated with holes accumulated at the

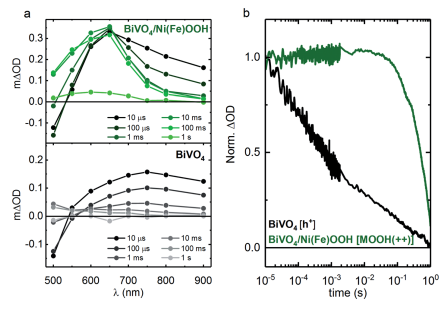


Fig. 3 Transient absorption spectra and kinetics of BiVO<sub>4</sub> and BiVO<sub>4</sub>/Ni(Fe)OOH photoanodes. (a) The transient absorption spectra showing the spectral signature of Ni(Fe)OOH(++) species in BiVO<sub>4</sub>/Ni(Fe)OOH (top panel) and BiVO<sub>4</sub> (bottom panel). (b) Normalised transient absorption kinetics of BiVO<sub>4</sub> (black, 650 nm, h<sup>+</sup> in BiVO<sub>4</sub>) and BiVO<sub>4</sub>/Ni(Fe)OOH (green, 650 nm, Ni(Fe)OOH(++)). All data were obtained under an applied potential of 1.4  $V_{RHE}$ , using 355 nm excitation (300  $\mu$ J cm<sup>-2</sup>), in 0.1 M phosphate buffer.

surface.³ The optical signal observed at shorter wavelengths ( $\sim$ <500 nm) across all BiVO<sub>4</sub> and BiVO<sub>4</sub>/MOOH photoanodes is dominated by a short-lived transient bleach and is indicative of the trapping of conduction band electrons into oxygen vacancy states as reported previously (we note that the magnitude of this bleaching varies between samples, mostly likely resulting from differences in oxygen vacancy density).³ The recovery of the bleach in the optical signal on the 1 ms timescale is concomitant with the electron extraction from the photoanodes, after which only the longer lived optical signal for BiVO<sub>4</sub> holes or MOOH(++) species can be observed.³

In the case of BiVO<sub>4</sub>/Ni(Fe)OOH (Fig. 3a, top panel), a peak at 650 nm can be observed from the 10 µs TA spectra onwards, which is reminiscent of the Ni(Fe)OOH(++) species observed under electrocatalytic conditions shown in Fig. 2a. This similarity in the spectral features at 650 nm suggest that both electrical and photo-induced oxidation (via surface holes from BiVO<sub>4</sub>) generate similar Ni(Fe)OOH(++) species. The observation of this feature in the spectrum obtained at 10 µs is indicative of a significant fraction of fast hole transfer from BiVO4 to the MOOH catalyst, which occurs within the time resolution of the measurements (<10 µs). The spectral shape in Fig. 3a (top panel) is clearly distinct from un-modified BiVO4 (Fig. 3a, bottom panel) at all timescales, indicating that holes primarily reside in the Ni(Fe)OOH catalyst layer over the timescales studied herein. Comparison of the kinetic traces presented in Fig. 3b shows that Ni(Fe)OOH(++) species have a much longer lifetime than the holes in BiVO<sub>4</sub>. Following hole transfer from BiVO<sub>4</sub> to the Ni(Fe)OOH layer, the accumulated Ni(Fe)OOH(++) species are spatially separated from photogenerated electrons in BiVO<sub>4</sub> and therefore, surface recombination in BiVO<sub>4</sub>/Ni(Fe)

OOH is reduced in comparison to un-modified BiVO<sub>4</sub>. Analogous conclusions can be drawn for the cases of BiVO<sub>4</sub>/FeOOH and BiVO<sub>4</sub>/FeOOHNiOOH, where the spectral profiles of the MOOH(++) species can be observed in the TA spectra (millisecond onwards), although charge transfer is less clear than the case of BiVO<sub>4</sub>/Ni(Fe)OOH due to the similarity in the spectral signatures of holes in BiVO<sub>4</sub> and MOOH(++) species in FeOOH and FeOOHNiOOH within the probed region. In addition, the optical signature of MOOH(++) species for FeOOH and FeOOHNiOOH at early timescales (µs to ms) also overlaps with the transient bleach observed due to electron trapping in BiVO<sub>4</sub>, further obscuring the detection of charge transfer at early timescales for these two cases. Nonetheless, all three catalystfunctionalised samples exhibit much longer lived signals than BiVO<sub>4</sub> alone (as seen in the TA spectra, Fig. 3 and S8 in the ESI†), indicative of catalyst deposition resulting in a substantial extension of the lifetime of photogenerated charges, attributed to slower recombination kinetics.

At early timescales (between 10  $\mu$ s and 1 ms), an absorption tail at longer wavelengths ( $\sim$ 700–900 nm) is present in all samples (Fig. 3a and S8†). As this spectral signal is also present in the un-modified BiVO<sub>4</sub> photoanode, it may be indicative of photogenerated charges within BiVO<sub>4</sub>. In the case of the catalyst modified samples (BiVO<sub>4</sub>/MOOH), this tail absorption decays by  $\sim$ 1 ms without any significant concomitant change in the signal observed for the MOOH(++) species, most clearly visible for the case of BiVO<sub>4</sub>/Ni(Fe)OOH (Fig. 3a). Since the extinction coefficients of the MOOH(++) species are  $\sim$ 7–8 times larger than the hole  $\epsilon$  of BiVO<sub>4</sub> (420 M $^{-1}$  cm $^{-1}$ ), <sup>36</sup> any further hole transfer from BiVO<sub>4</sub> to the catalyst layer within 10  $\mu$ s and 1 ms would be expected to give rise to a much larger signal observed for the

MOOH(++) species. Since no increase in signal is observed, this may be indicative of recombination of photogenerated species in BiVO<sub>4</sub> that have not transferred to the catalyst layer by  $\sim$ 10

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It is intriguing that the FeOOHNiOOH functionalised BiVO<sub>4</sub> photoanode performs better than the FeOOH counterpart (Fig. 1b), despite exhibiting almost invariant kinetics for water oxidation under electrocatalytic conditions (Fig. 2b). From the TA spectra (Fig. S8†), it appears that the signal amplitude for BiVO<sub>4</sub>/FeOOHNiOOH is slightly larger than it is for BiVO<sub>4</sub>/ FeOOH. Taking the respective extinction coefficients of the MOOH(++) species into consideration, this would indicate that a greater amount of holes successfully transfer from BiVO4 to FeOOHNiOOH in comparison to FeOOH, thus suggestive of FeOOHNiOOH being able to accumulate a greater density of MOOH(++) in comparison to FeOOH alone, when deposited on BiVO<sub>4</sub>. On the other hand, the lower performance of BiVO<sub>4</sub>/ Ni(Fe)OOH can be explained by the slower kinetics of the Ni(Fe) OOH(++) species (as shown in Fig. 2b) for the same accumulated charges. For electrochemical water oxidation, this is compensated by the accumulation of a greater density of MOOH(++) species. However, under illumination when BiVO<sub>4</sub>/Ni(Fe)OOH is used, the maximum number of charges accumulated on the Ni(Fe)OOH layer will be determined primarily by the number of photogenerated holes that are transferred to the catalyst layer. As observed in the TA spectra at early timescales, the amount of MOOH(++) accumulated on the MOOH layer is within the same order of magnitude between the three catalysts. Since water

oxidation is slower on Ni(Fe)OOH in comparison to the other two catalysts for a given MOOH(++) density, there is increased scope for competing recombination processes at the  $BiVO_4/Ni(Fe)OOH$  interface, as reported previously. <sup>18</sup>

To examine the photoelectrochemical processes under steady-state conditions, spectroelectrochemical photoinduced absorption spectroscopy (PIAS) was employed. Unfortunately, the instability of BiVO<sub>4</sub>/Ni(Fe)OOH samples (Fig. S10†) under oxidative conditions over prolonged periods of time required for PIAS measurements precludes them from these analyses. The normalised PIAS spectra of BiVO4, BiVO4/FeOOH and BiVO<sub>4</sub>/FeOOHNiOOH under an applied potential of 1.4 V<sub>RHE</sub> and 1 sun illumination are shown in Fig. 4, along with the normalised spectra of the respective MOOH(++) species obtained under electrochemical conditions. The resemblance of the spectral shapes indicate that similar species accumulate under the two catalytic conditions, which, in conjunction with the TAS data discussed above, further confirms that photoexcitation of BiVO4 leads to the oxidation of the catalyst, also under PEC operation. Similar to the TA spectra, the difference in signal between PIA and SEC at shorter wavelengths (<550 nm) is most likely due to electron trapping within BiVO<sub>4</sub>.38

The optical signals obtained through PIAS and SP-SEC for the photoanodes and electrocatalyst electrodes, respectively, under catalytic conditions, were converted to the amount of oxidising equivalents for water oxidation, *i.e.* holes in BiVO<sub>4</sub> and MOOH(++) species in the case of the catalysts. By relating the optical signal to the concomitant current transient signals,

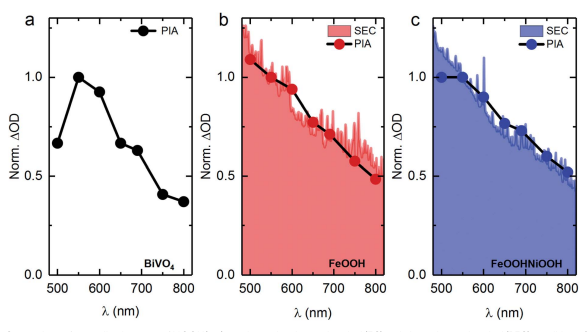


Fig. 4 Comparison of normalised spectra of MOOH(++) species under electrochemical (EC) and photoelectrochemical (PEC) conditions. (a) The PIA spectrum of BiVO $_4$ . (b) The SEC spectrum of FeOOH electrocatalyst (red, shaded) and the PIA spectrum of BiVO $_4$ /FeOOH (solid red circle, black line). (c) The SEC spectrum of FeOOHNiOOH electrocatalyst (blue, shaded) and the PIA spectrum of BiVO $_4$ /FeOOHNiOOH (solid blue circle, black line). All SEC spectra under electrochemical conditions were obtained under an applied potential of 2  $V_{RHE}$ , and all PIA spectra were obtained under 1 sun equivalent illumination (365 nm LED) under 1.4  $V_{RHE}$ . All spectra were normalised at 550 nm. All measurements were taken in 0.1 M phosphate buffer (pH 7).

kinetic information associated with water oxidation reaction can be analysed using rate law plots according to the following equation:

$$\log J = \log k_{WO} + \beta \log(\text{MOOH}(++)) \tag{1}$$

where J is the (photo)electrochemical steady-state current density (water oxidation flux), (MOOH(++)) is the density of the MOOH(++) species (in the case of un-modified BiVO<sub>4</sub>, it is the surface h<sup>+</sup> density),  $k_{\rm WO}$  is the water oxidation rate constant and  $\beta$  is the order of the reaction with respect to the surface density of MOOH(++) (or h<sup>+</sup> density in BiVO<sub>4</sub>). This analysis approach has been employed to investigate reaction kinetics on metal oxide surfaces<sup>24–26</sup> and electrocatalytic systems previously.<sup>27,29</sup> This analysis assumes that  $k_{\rm WO}$  is independent of applied potential or the density of the MOOH(++) species, more discussion and justification of this assumption is presented elsewhere.<sup>39,40</sup>

Firstly, from Fig. 5a, we can observe that photogenerated holes in BiVO<sub>4</sub> follow a reaction order ( $\beta$ ) of 3 for water oxidation, consistent with previous reports, <sup>25,26</sup> whereas both FeOOH and FeOOHNiOOH operate with a  $\beta$  value of  $\sim$ 2 under both photoelectrochemical and electrochemical conditions at pH 7. A reaction order of  $\sim$ 2 for these electrocatalysts differs from that observed under alkaline conditions ( $\beta \sim$  4 at pH 13),<sup>27</sup> and is indicative of differences in reaction mechanisms in neutral and alkaline media. Surface protonation as well as the reactivity and

stability of reaction intermediates can result in different reaction mechanisms being observed at neutral and alkaline pH.41 The effect of pH on reaction order has also been observed for TiO2.42 More importantly, at pH 7, the different reaction orders between BiVO<sub>4</sub> and BiVO<sub>4</sub>/MOOH photoanodes indicates that water oxidation proceeds via different active species and consequently through different mechanisms. Whilst in un-modified BiVO<sub>4</sub> photoanodes water oxidation is driven directly by photogenerated BiVO<sub>4</sub> holes, in the case of BiVO<sub>4</sub>/MOOH(++) photoanodes, the oxidised MOOH(++) species drive water oxidation in the catalyst layer, following hole transfer from BiVO<sub>4</sub>. Furthermore, it is apparent that similar orders of magnitude of MOOH(++) species are observed optically under both electrochemical (EC) and photoelectrochemical (PEC) conditions, consistent with the MOOH(++) species driving water oxidation in both systems, although differences in film morphology between the EC and PEC samples prevent quantitative comparison. A comparison of the reaction time constants for PEC water oxidation, shown in Fig. 5b for BiVO<sub>4</sub> and BiVO<sub>4</sub>/MOOH photoelectrodes, indicate that no significant enhancement in water oxidation timescale is gained upon electrocatalyst deposition.

It is also striking that catalysts often employed for water oxidation, where water oxidation is observed to take place *via* oxidised states in the catalyst (*i.e.* Co–Fe based Prussian Blue), display time constants on the order of seconds.<sup>43</sup> Even the popular Ni(Fe)OOH catalysts investigated under optimal alkaline conditions<sup>27</sup> operate with time constants that are

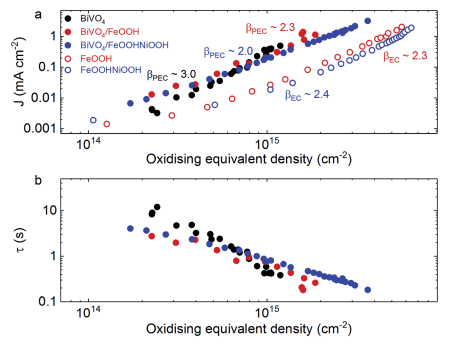
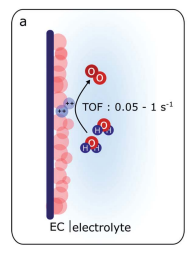
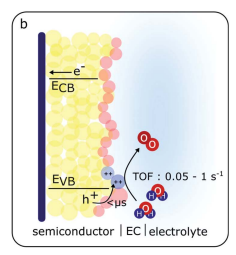


Fig. 5 Comparison of kinetics under electrochemical and photoelectrochemical conditions. (a) The density of oxidising equivalents ( $h^+$  for BiVO<sub>4</sub>, or MOOH(++) species for catalysts) under catalytic conditions plotted against the (photo)current obtained. The reaction orders ( $\beta$ ) for the respective electrochemical and photoelectrochemical water oxidation reactions are also shown (determined from the slope). (b) The density of oxidising equivalents plotted against the water reaction time constants ( $\tau$ ) under photoelectrochemical conditions for BiVO<sub>4</sub> and BiVO<sub>4</sub>/MOOH photoanodes. Same colours used in (a) and (b): BiVO<sub>4</sub> (black solid circles); BiVO<sub>4</sub>/FeOOH (red solid circles); BiVO<sub>4</sub>/FeOOHNiOOH (blue solid circles); FeOOH (red empty circles); FeOOHNiOOH (blue empty circles). All measurements were taken in 0.1 M phosphate buffer.





Scheme 1 Schematic illustration of electrochemical and photoelectrochemical water oxidation in MOOH electrodes and  $BiVO_4/MOOH$  photoelectrodes. (a) The electrochemically generated MOOH(++) species (blue spheres) in the activated electrocatalyst accumulate and drive water oxidation. (b) The photogenerated holes in the  $BiVO_4$  transfer to the activated catalyst, generating MOOH(++) species (blue spheres) which then drive water oxidation. The electrons in the conduction band are collected at the back contact.

comparable to water oxidation by metal oxides such as  $\mathrm{BiVO_4}$ , and slower than that observed for  $\mathrm{WO_3}.^{25,44}$  We note however that these electrocatalysts operate at much lower overpotentials for matched charge densities. Therefore, it appears that the significant enhancement in photoelectrochemical performance of  $\mathrm{BiVO_4}$  photoanodes decorated with these electrocatalysts does not originate from faster kinetics of water oxidation relative to  $\mathrm{BiVO_4}$ . Rather, the results obtained in this study suggest that the role of the catalyst layer is to decrease the kinetics of surface recombination and effectively decrease the recombination losses via fast hole transfer from  $\mathrm{BiVO_4}$  to the catalyst layer. This spatially separates charges, and retards surface recombination losses that compete with water oxidation.

The same observations were found for Co-Fe Prussian Blue, where fast hole transfer to the catalyst layer followed by fast water oxidation kinetics led to enhanced performance.43 This is however, markedly different to CoPi, which exhibits water oxidation time constants ( $\tau$ ) that are two to three orders of magnitude slower than what is observed for BiVO<sub>4</sub> alone, where the enhancement in performance has been attributed to increased band-bending within the BiVO4 that drives charge separation. 17,36 It is observed that the amount of charge accumulated in CoPi under PEC conditions (when deposited on BiVO<sub>4</sub>) is significantly lower (by two orders of magnitude) than that observed in the electrocatalyst under electrochemical water oxidation conditions at matched current densities.36 Therefore, it is concluded that the amount of charge that accumulates in CoPi from BiVO<sub>4</sub> is insufficient to drive water oxidation. This conclusion is also supported by intensity-modulated photocurrent spectroscopy (IMPS) studies which indicate that CoPi mainly serves to passivate surface defects, with no significant enhancement of water oxidation kinetics when deposited on BiVO<sub>4</sub> or α-Fe<sub>2</sub>O<sub>3</sub>.9,11 However, as conflicting studies based on dual working electrode measurements that assign water

oxidation to the CoPi catalyst overlayers also have been reported,  $^{45}$  this makes the role of CoPi as a co-catalyst overlayer on semiconductor photoelectrodes difficult to determine. We note that factors such as the porosity of the semiconductor and the ion permeability of catalysts can also affect the nature of the interface between the semiconductor/catalyst. For example, the ability of some catalysts to form Schottky junctions has been proposed to increase the efficiency of a semiconductor/catalyst system. The nature of the interface playing a critical role is further demonstrated by  $\text{IrO}_x$  electrocatalysts, where despite  $\text{IrO}_x$  being an excellent water oxidation electrocatalyst, it has been reported as an ineffective co-catalyst when deposited on  $\text{BiVO}_4$ .

For the case of the MOOH electrocatalysts studied herein, although a slight discrepancy in the amount of MOOH(++) species is also observed between the electrocatalysts and the BiVO<sub>4</sub>/MOOH photoelectrodes, this difference (within an order of magnitude) is less dramatic than the case of CoPi,36 and is more similar to the case of RuOx catalysts operating as electrocatalysts and co-catalysts on photoelectrochemical systems.<sup>29</sup> The charge transfer observed from BiVO<sub>4</sub> to the catalyst layer to generate MOOH(++) species is seen both in the TA spectra and the photo-induced absorption spectra under steady-state conditions. Additionally, as the reaction order for water oxidation for BiVO<sub>4</sub>/MOOH photoanodes also differs from that of BiVO<sub>4</sub> photoanodes, it strongly indicates that water oxidation in the catalyst-decorated photoanodes studied herein is driven via MOOH(++) species accumulated in the catalyst layer (as illustrated in Scheme 1).

#### Conclusions

In this work, the electrochemical and photoelectrochemical performance of FeOOH, FeOOHNiOOH, and Ni(Fe)OOH catalysts are studied at pH 7 as standalone electrocatalysts and

when added on the surface of BiVO<sub>4</sub> photoanodes. Our results demonstrate that electrocatalytic performance results from both the intrinsic reaction kinetics and the potential dependence of charge accumulation, where the latter consideration is not important under photoelectrochemical conditions. We show that while FeOOH and FeOOHNiOOH exhibit faster kinetics for water oxidation than Ni(Fe)OOH, this nickel-based catalyst is able to accumulate a larger density of oxidative MOOH(++). As a result, the three catalyst show a similar electrocatalytic water oxidation performances. This trade-off between accumulation and reaction kinetics hinders the selection of electrocatalysts to make junctions with photoanodes, through standard *J–V* characterisations.

Once deposited onto the surface of nanoporous BiVO4 phosubstantial improvements in the phototoanodes. electrochemical performance are observed with respect to unmodified BiVO<sub>4</sub>, attributed to fast charge transfer (<10 µs) from BiVO<sub>4</sub> to the catalyst layer, and the resultant suppressed recombination losses. However, rather surprisingly, no significant enhancement in the water oxidation time constant is observed for the BiVO<sub>4</sub>/MOOH photoanodes in comparison to BiVO<sub>4</sub>. Moreover, differences between the catalysts emerge in the BiVO<sub>4</sub>/MOOH photoanodes, stemming from the charge accumulation within the catalyst layer being governed by the transfer of photogenerated holes from BiVO<sub>4</sub>. This significantly impedes the ability of Ni(Fe)OOH to accumulate a greater amount of Ni(Fe)OOH(++) needed to offset its slower water oxidation kinetics, which leads to a lower performance of BiVO<sub>4</sub>/Ni(Fe)OOH photoanodes in comparison to the other two configurations. On the other hand, when NiOOH is added to FeOOH as a dual-layer catalyst on BiVO4, it helps accumulate more MOOH(++) species within the catalyst layer than FeOOH alone, resulting in a greater enhancement in the photocurrent generated by BiVO<sub>4</sub>/FeOOHNiOOH photoanodes. Overall, this study highlights the two different factors that govern the performance BiVO<sub>4</sub>/MOOH photoanodes and need to be considered when selecting electrocatalysts for applications in photoelectrochemical systems: (1) the number of charges that can be accumulated within the catalyst layer; and (2) how fast these charges can oxidise water to compete with recombination processes.

#### Author contributions

J. D. and K.-S. C. contributed to the conceptualisation. L. F., S. S., S. C. and D. L. contributed to the investigation and formal analysis. C. A. M. and E. P. contributed to the development of methodology and analysis.

#### Conflicts of interest

Authors declare no conflict of interest.

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