

Wireless Rotating Disk Electrode (wRDE) for assessing Heterogeneous Water Oxidation Catalysts (WOCs)

Mills, A., & O'Rourke, C. (2016). Wireless Rotating Disk Electrode (wRDE) for assessing Heterogeneous Water Oxidation Catalysts (WOCs). Chemical Communications, (49), 7727-7730. DOI: 10.1039/c6cc03356c, DOI: 10.1039/c6cc03356c

Published in:

Chemical Communications

Document Version:

Peer reviewed version

Queen's University Belfast - Research Portal:

Link to publication record in Queen's University Belfast Research Portal

Publisher rights

© The Royal Society of Chemistry 2016

General rights

Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy

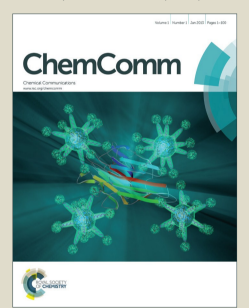
The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.



ChemComm

Accepted Manuscript

This article can be cited before page numbers have been issued, to do this please use: A. Mills and C. O'Rourke, *Chem. Commun.*, 2016, DOI: 10.1039/C6CC03356C.



This is an *Accepted Manuscript*, which has been through the Royal Society of Chemistry peer review process and has been accepted for publication.

Accepted Manuscripts are published online shortly after acceptance, before technical editing, formatting and proof reading. Using this free service, authors can make their results available to the community, in citable form, before we publish the edited article. We will replace this Accepted Manuscript with the edited and formatted Advance Article as soon as it is available.

You can find more information about *Accepted Manuscripts* in the **Information for Authors**.

Please note that technical editing may introduce minor changes to the text and/or graphics, which may alter content. The journal's standard <u>Terms & Conditions</u> and the <u>Ethical guidelines</u> still apply. In no event shall the Royal Society of Chemistry be held responsible for any errors or omissions in this *Accepted Manuscript* or any consequences arising from the use of any information it contains.



Published on 23 May 2016. Downloaded by Queen's Library on 26/05/2016 12:51:47

DOI: 10.1039/C6CC03356C



Journal Name

COMMUNICATION

Wireless Rotating Disk Electrode (wRDE) for assessing Heterogeneous Water Oxidation Catalysts (WOCs)

Received 00th January 20xx, Accepted 00th January 20xx

A. Mills^a* and C. O'Rourke^a

DOI: 10.1039/x0xx00000x

www.rsc.org/

A novel method for assessing the activity of a powdered water oxidation catalyst (WOC) is described, utilising an easily-prepared wireless rotating disc electrode of the WOC, thereby allowing its activity to be probed, via the observed kinetics of water oxidation by Ce(IV) ions, and so provide invaluable electrochemical information.

There is a great deal of interest in the development of efficient, long-lived, low-cost, solar-driven photosystems for the cleavage of water into hydrogen and oxygen¹⁻³. One of the great challenges in this work is the identification of a suitable redox catalyst to mediate the oxidation of water, i.e. a water oxidation catalyst, WOC, by the photo-generated oxidant, Ox, which is usually a valance-band hole in a semiconductor photocatalyst^{4,5}. The challenge arises because the oxidation of water is an electrochemically irreversible process, so that most WOCs require a significant overpotential, η , to drive the reaction at the current densities (ca. 20 mA cm⁻²)⁴ that are possible in a very efficient, solar-driven, water-splitting cell⁴. The best WOCs are the oxides of Ru or Ir^6 but, the latter are not abundant⁴ and so there has been a great deal of effort in recent years to identify new, alternative, inexpensive, i.e. Earth-abundant, highly active $WOCs^3$.

As part of this effort, many new *WOCs* are generated in powder form and then screened for activity using a simple redox reaction involving the oxidation of water, i.e. reaction (1).

$$NOX + 2H_2O \longrightarrow nRed + 4H^{+} + O_2 \uparrow$$
 (1

Where Ox is the oxidant, Red is the reduced form of Ox, and n is the number of equivalents necessary to consume 4 electrons from water^{3,7,8}; obviously, Ox must have an oxidation potential greater than that of water, i.e. $E(Ox/Red) > \{E(O_2/H_2O) = 1.23 - 0.059pH\}$. In such screening studies^{3,8}, the powdered WOC is commonly dispersed in solution and, upon addition (or photochemical generation) of the oxidising chemical species, Ox, the rate of

reaction (1) is measured, usually spectrophotometrically, via its rate of decrease in [Ox] with time; this rate is taken as a measure of the activity of the $WOC^{3,7\cdot9}$. A common chemical oxidant for such work is Ce(IV), since E(Ce(IV)/Ce(III)) = 1.44, 1.61 and 1.70 in 1 N H₂SO₄, HNO₃ and HClO₄ acid, respectively¹⁰ (see (ESI†: S1)); and, in the absence of a WOC, it is very stable^{7,8}. On most conducting electrode materials, the Ce(IV)/Ce(III) redox couple acts a highly reversible (i.e. Nernstian) couple^{11,12}, whereas that for water oxidation is highly irreversible⁶.

Reaction (1) is generally used by most to provide only a crude assessment of the activity of the *WOC* under test, such as initial rate and/or turnover frequency^{13,14}. Thus, in order to probe the electrocatalytic properties of a powder-based *WOC*, a knowledge of which is key if it is to be used in a final, solar-driven, water-splitting device, the powder-based *WOC* must be fabricated as an anode in what is often a poorly defined, non-trivial and time-consuming process¹⁵. However, the latter exercise may be unnecessary, since the kinetics of reaction (1) can allow the extraction of useful electrochemical information that relates to the electrocatalytic properties of the powder-based *WOC*. This option arises because, in most cases, the powder particles of the *WOC* act simply as microelectrodes that couple, the electrochemical reduction of the oxidant, such as Ce(IV), with the oxidation of water¹⁶.

In order to extract out electrochemical information via the kinetics of decay of Ox, e.g. Ce(IV) in reaction (1), it is essential that the powder dispersion remains unchanged, i.e. doesn't corrode nor aggregate, during the course of the reaction. The latter condition is often difficult to achieve, especially with low activity WOCs, which exhibit a long reaction time and so provide ample opportunity for extensive aggregation (and often corrosion) to occur. Although the addition of an anti-flocculant can help reduce this effect, such additives will necessarily affect the intrinsic electrochemical characteristics of the WOC under test that are being probed 17 .

Herein we describe a simple, quick method for fabricating a wireless, i.e. no external electrical contact, rotating disc electrode, wRDE, of a powdered WOC, here, primarily, RuO₂(Adams), which allows the electrochemical characteristics of the WOC to be readily probed, via the measured kinetics of decay of Ce(IV) in reaction (1).

^{a.} School of Chemistry and Chemical Engineering, Queen's University Belfast, Stranmillis Road, Belfast, BT9 5AG, UK. E-mail: andrew.mills@qub.ac.uk; Fax: +44 (0)28 9097 6524; Tel: +44 (0)28 9097 4339

 $[\]mbox{\ensuremath{^{\dagger}}}$ Electronic supplementary information (ESI) available. See DOI:

DOI: 10.1039/C6CC03356C

ARTICLE Journal Name

In this work, the powdered WOC under test was the black-coloured powder, $RuO_2(Adams)$, the preparation of which has been described elsewhere 18 . A schematic illustration of the general method of preparation of the wRDE of this WOC is illustrated in figure 1 and full details, and photographs of the final product, are given in (ESI $^+$: S2).

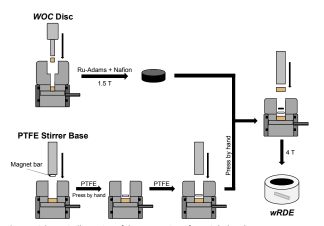


Figure 1 Schematic illustration of the preparation of a $RuO_2(Adams)$ wRDE.

Published on 23 May 2016. Downloaded by Queen's Library on 26/05/2016 12:51:47

The electrochemical properties of the RuO₂(Adams) wRDE were probed by placing it in a spectrophotometer cuvette containing 1.5 ml 0.5 M H₂SO₄ solution, thermostatted at 30°C, and spinning it continuously at 1000 rpm using a magnetic stirrer. The water oxidation reaction (1) was initiated by injecting, into the solution in the cuvette, 54 µL of a 0.1 M Ce(IV) sulfate in 0.1 N H₂SO₄ (Aldrich Chemicals). The subsequent change in the absorbance, at 430 nm, of the reaction solution was measured as a function of time, using a Cary 60 spectrophotometer. A typical set of absorption spectral changes, derived from such a typical kinetic run, are illustrated in figure 2. The insert diagram shows the measured decays in absorbance, due to Ce(IV) for two serial injections of Ce(IV). The two decays are identical and near perfect 1st order with respect to Ce(IV) concentration, [Ce(IV)]t, yielding a value for the first order rate constant, $k_1 = 0.0149 \text{ min}^{-1}$, over 3 half-lives, and correlation coefficient > 0.9999. Previous work carried out using this WOC has established the generation of stoichiometric amounts of O2 via reaction (1) under the same conditions¹⁶. In this system a value of the molar Ce:Ru ratio, θ , for just one injection of Ce(IV) 9.6x10⁻⁵ can be calculated 19 which suggests that it is operating under catalytic conditions. Details of this calculation can be found in (ESI+: S3).

From the identical natures of the two decay profiles illustrated in figure 2 it is clear that the rate of reaction (1) is independent of the concentration of Ce(III), i.e. [Ce(III)]_t, generated during the course of reaction, at reaction time t. This suggests that the rate is independent of the redox potential of the Ce(IV)/Ce(III) couple in solution, which, according to the Nernst equation, would change, as the [Ce(IV)]_t:[Ce(III)]_t ratio changes, during the course of reaction (1).

Many researchers working on WOCs favour using cerium ammonium nitrate (CAN) as the oxidant and 1 M HNO₃ as the acid. Thus, (i) to show that such a medium is appropriate for work with a wRDE, (ii) provide further support for the electrochemical model presented here and (iii) to emphasise further the catalytic nature of

the RuO₂(Adams), the CAN/1M HNO₃ system was used to test the RuO₂(Adams) *wRDE*. The results from this work revealed the same 1st order kinetics of Ce(IV) decay, using either 3.6×10^{-3} or 0.1 M CAN, in 1 M HNO₃, as illustrated by the decay profiles and photographs in (ESI†: S4).

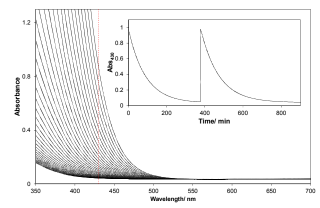


Figure 2 Change in the UV/Vis spectra upon the 1st injection of 54 μ L of 0.1 M Ce(SO₄)₂ into 1.5 mL of 0.5 M H₂SO₄. The red broken line indicates the wavelength at which the reduction of Ce(IV) is monitored, i.e. 430 nm. The inset shows a plot of the decay of Ce(IV) over 2 serial injections. In all cases the wRDE rotation speed = 1000 rpm and reaction temperature = 30°C.

At a conventional, i.e. wired, *RDE*, when the rate of reduction of a oxidising species, such as Ce(IV), is independent of applied potential, the reaction is diffusion-controlled and the measure current obeys the Levich equation ^{11,12}:

$$i_d = 0.62.n.$$
F. $A.$ **D**^{2/3} $\omega^{1/2}$ $v^{-1/6}$ [Ox] (2)

where, i_d = diffusion limited current (A), n = number of electrons transferred during the reduction reaction, \mathbf{F} = Faraday's constant (C/mol), A = electrode area (cm²), \mathbf{D} = diffusion coefficient for Ox (cm²/s), ω = rotation speed angular rate of rotation of the electrode (rads/s), υ = kinematic viscosity (cm²/s) and [Ox] = concentration of oxidant (mol/cm³). The observation of excellent 1st order kinetics for the Ce(IV) reduction on the RuO₂(Adams) wRDE, see figure 2, suggests that in this system the rate of reaction (1), with Ox = Ce(IV), is diffusion-controlled and, as such, should obey the following modified version of the Levich equation:

rate (M/s) =
$$k_d$$
[Ce(IV)]_t = 0.62. $A.D^{2/3}\omega^{1/2}v^{-1/6}$ [Ce(IV)]_t/ V (3)

where k_d is the measured value for the first order rate constant under these conditions, V is the reaction solution volume (cm³). Given $A = 0.196 \text{ cm}^2$, $V = 1.554 \text{ cm}^3$, $v = 0.01 \text{ cm}^2/\text{s}$, it follows that eqn(3) predicts the following relationship between k_d (i.e. the measured value of k_1 at 30°C) and the rotation speed of the wRDE, qr

$$k_d = 0.168. \mathbf{D}^{2/3} \omega^{1/2} \tag{4}$$

In order to test this predicted relationship, the same conditions as used to generate the data in figure 2 were employed, but with the rotation speed of the wRDE varied systematically (100-1300 rpm) in a series otherwise identical kinetic runs involving reaction (1), with Ox = Ce(IV). In all cases the kinetics of Ce(IV) decay were monitored

ChemComm Accepted Manuscript

spectrophometrically and yielded excellent 1^{st} order plots of the decay data over 3 half-lives ($r^2 > 0.9999$).

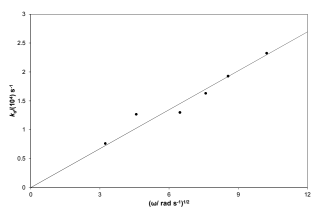


Figure 3 Plot of the 1st order rate, k_d , of reduction of Ce(IV) using the *wRDE* as a function of the square root of the rotation speed. Reaction conditions: otherwise as in figure 1.

A plot of the k_d values derived from this work versus $\omega^{1/2}$ is illustrated above in figure 3 and yields a gradient = 2.25×10^{-5} , from which a value for the diffusion coefficient, **D**, of 1.55×10^{-6} cm²/s was calculated using eqn (4). The latter appears low when compared to reported^{11,12} values for D for Ce(IV) on Pt electrodes $(3.4-5) \times 10^{-6}$ cm²/s, although this may be because the actual value for D was lower than the geometric value used in the calculations. The latter situation would occur if a significant part of the D0 in the D1 was not electro active, which would happen if, as well as naked D1 RuO2(Adams) particles, the surface comprised regions of pure Nafion, or Nafion-covered RuO2, particles. Initial optical and electron micrographs of the RuO2(Adams) wRDE provide some evidence for this.

Unfortunately, although the $RuO_2(Adams)$ wRDE appears to obey the Levich equation, this doesn't actually probe the electrocatalytic activity of the $RuO_2(Adams)$ as such, since the kinetics are diffusion-controlled. Instead, experimental conditions need to be used under which the kinetics is no longer purely diffusion controlled. In this work, this was simply achieved by running the same experiment as above, but at a much lower temperature, since the activation-controlled oxidation of water is much more temperature sensitive ($E_a = 38-55 \text{ kJ/mol})^6$ than the diffusion-controlled reduction of Ce(IV) ($E_a = ca.\ 15 \text{ kJ/mol})^{20}$, so that lower temperatures will favour activation-, rather than diffusion-, -controlled kinetics for reaction (1) using the $RuO_2(Adams)$ wRDE.

Thus, the absorbance decay plots illustrated in figure 4, were generated for three serial injections of Ce(IV), using the same reaction conditions as used in figure 1, but with a reaction temperature of 5°C, rather than 30°C. The results show that, unlike those recorded at 30°C, at 5°C the kinetics of reaction (1) are dependent upon the amount of Ce(III) generated, i.e. the reaction slows down as the redox potential, due to the Ce(IV)/Ce(III) couple, decreases with increasing reaction time, t.

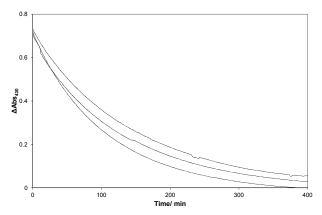


Figure 4 ΔAbs_{430} vs time upon 3 serial injections of 54 μL of 0.1 M Ce(SO₄)₂ into 1.5 mL of 0.5 M H₂SO₄ at 5°C. Rotation speed = 1000 rpm.

As commented earlier, in redox catalysis it is recognised²¹ that the role of the catalyst is often simply that of a conductor of electrons from one redox couple – in this case Ox/Red in reaction (1) – to another, i.e. O_2/H_2O in reaction (1). It follows that at any time during a Ce(IV) decay, such as illustrated in figure 4, each dispersed RuO₂(Adams) WOC particle is poised at a mixed potential, $E_{mix,tr}$, so as to allow a 'mixed' current, $i_{mix,tr}$ to flow through it, with a numerical value that is the same for both the anodic and cathodic electrochemical processes that are occurring concomitantly at separate sites on the redox catalyst. The Nernstian nature of the Ce(IV)/Ce(III) electrochemical process allows the value of $E_{mix,tr}$ at any time t during a decay, such as illustrated in figure 4, to be calculated 16 via:

$$E_{mix,t} = E'_{Ce} + (\mathbf{R}T/\mathbf{F}) \ln\{[-k_d[Ce(IV)]_t - r(t)]/[r(t) - k_d[Ce(III)]_t]\}$$
 (5)

where, $E'_{Ce} = 1.44 \text{ V}$ vs NHE in 0.5 M H₂SO₄, k_d = the diffusion-controlled rate constant (s⁻¹) and r(t) = the rate of Ce(IV) reduction at time t (= $d[\text{Ce}(IV)]_t/dt$). Since, on any RuO₂(Adams) particle, the electrochemical reduction of Ce(IV) is coupled to the oxidation of water, it follows that the latter, irreversible process, can be related to the rate of Ce(IV) reduction by the following expression:

$$r(t) = r_w \cdot \exp\{2.303(E_{mix.t} - E'_w)/b\}$$
 (6)

where, r_w = a constant that is related directly to the exchange current density for water oxidation, E'_w = the formal redox potential of the O_2/H_2O couple (here, taken as: 1.23 V vs NHE) and b = Tafel slope. It follows from eqn (6) that a plot of $E_{mix,t}$ vs $\log\{r(t)\}$ should yield a straight line with a gradient, b. At any reaction time t, for any of the decay curves illustrated in figure 4, it is possible to calculate appropriate values for: r(t), $[Ce(IV)]_t$, $[Ce(III)]_t$ and so a value for $E_{mix,t}$, using eqn (5). The results of this type of analysis of all the decay data illustrated in figure 4 are illustrated in figure 5 and reveal an approximate Tafel slope of 54 ± 2 mV/decade and an extrapolated value for $r(t) = 1.04 \times 10^{-8}$ M/min at $E_{mix,t} = 1.23$ V.

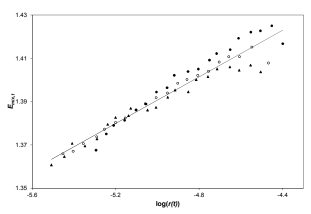
For RuO₂, the reported Tafel slope for oxygen evolution varies between 40 and 120 mV, depending how it is prepared⁶. A value of 65 mV/decade at 25°C in 0.5 M H_2SO_4 has been reported previously for RuO₂ films grown on Ru metal²². A value of 60 mV/decade at 25°C (\equiv 56 mV/decade at 5°C) may be derived from theory if the

DOI: 10.1039/C6CC03356C **ARTICLE** Journal Name

deprotonation of the reaction site S-OH is the rate determining step, as proposed by Krasil'shchikov^{6,23}, i.e.:

$$S-OH \longrightarrow S-O^{-} + H^{+}$$
 (7)

From the data in figure 5, given that at $E_{mix,t}$ = 1.23 V, $\equiv E'_{w}$, the value of r(t), $\equiv r_w$ in eqn(6), = 1.04x10⁻⁸ M/min, which in turn is \equiv 1.32 μ A/cm², given $A = 0.196 \text{ cm}^2$ and $V = 1.554 \text{ cm}^3$. It follows that, in order to oxidise water to O2 on the wRDE Ru(Adams) electrode at a current density = 20 mA/cm², an overpotential, = $(E_{mix,t} - E'_w)$ in eqn(6), of 226 mV will be required. This value is in reasonable agreement with the work²² referred to earlier on RuO₂ films grown on Ru metal, for which a value of η = 270 mV can be calculated.



Published on 23 May 2016. Downloaded by Queen's Library on 26/05/2016 12:51:47

Figure 5 Tafel plot of the data in figure 5 giving a slope of 54 mV per decade and a k_d value of 0.015 min⁻¹. The units of r(t) here are M/min.

Table 1 List of Tafel slopes, b, the rate at a potential of 1.23 V, r_w , and the calculated overpotential, η (at a current density of 20 mA/cm²), derived from the data in (ESI†: S5, figure S5).

Catalyst	b/ mV per decade	η/mV	r _w / (10 ⁹) M min ⁻¹
RuO ₂ (Adams)	54	226	10.4
IrO ₂	58	280	2.34
PtO₂‡	125	478	23.7

‡ Run in 1 N HNO₃ at 30°C, all other wRDE's were run in 1.5 mL of 1 N H₂SO₄ at 5°C.

Other wRDE's were prepared using commercial forms of IrO₂.2H₂O (Alfa Aesar) and PtO₂ (Sigma Aldrich) and tested using the Ce(IV) decay system yielding Tafel plots that are illustrated in (ESI†: S5), from which the values of b and η listed in Table 1 were derived. The difference in the η value for IrO₂, to that of RuO₂(Adams), (54 mV) is consistent with that (60 mV) reported by others recently for sputtered films of the same oxides 24 . PtO_2 has been reported previously as a WOC for reaction (1) in 1 N H₂SO₄, albeit of low activity²⁵, where Ox = Ce(IV), and so was studied here, as a wRDE, although under more oxidising conditions (i.e. in 1 M HNO3, vide supra) for which E_{Ce} is 1.61 V¹⁰, in order to generate a more easily measured decay. The values of b and η derived from this work for PtO₂, see Table 1, are in reasonable agreement with those reported by others for the same material and acid (i.e. b = 109 mV/decade; η $= 500 \text{ mV})^6$.

Conclusions

In summary, the construction of a wireless rotating disc electrode, wRDE, of a water oxidation catalyst, in this case RuO2(Adams), is reported. This method allows powdered materials, which might be able to act as WOCs, to be assessed for catalytic activity using a suitable reversible redox couple, such as Ce(IV)/Ce(III).

Acknowledgements

We thank DEL and the US/Ireland R&D Partnership programme (USI 049) for funding this work.

Notes and references

- N. S. Lewis and D. G. Nocera, Proc. Natl. Acad. Sci. USA, 2006, **103**, 15729-15735.
- H-J. Lewerenz, L. Peter, Photoelectrochemical Water Splitting: Materials, Processes and Architectures, RSC Publishing, Cambridge, 2013.
- J. R. Galán-Mascarós, ChemElectroChem, 2015, 2, 37–50.
- M. G. Walter, E. L. Warren, J. R. McKone, S. W. Boettcher, Q. Mi, E. A. Santori and N. S. Lewis, Chem. Rev., 2010, 110, 6446-6473.
- Y. W. Chen, J. D. Prange, S. Dühnen, Y. Park, M. Gunji, C. E. D. Chidsey and P. C. McIntyre, Nat. Mater., 2011, 10, 539-544.
- S. Trasatti and G. Lodi, Electrodes of Conductive Metallic Oxides, Part B, S. Trasatti (ed.), Elsevier, Amsterdam, 1981, ch. 10.
- A. R. Parent, R. H. Crabtree and G. W. Brudvig, Chem. Soc. Rev., 2013, 42, 2247-2252.
- A. Mills, Chem. Soc. Rev., 1989, 18, 285-316.
- B. Limburg, E. Bouwman and S. Bonnet, Coord. Chem. Rev., 2012. **256**. 1451-1467
- Ionic Equilibria in Analytical Chemistry, J-L. Burgot, Springer, New York, 2012, p. 395.
- T. H. Randle and A. T. Kuhn, J. Chem. Soc. Faraday Trans. 1, 1983, 79, 1741-1756.
- 12 R. Greef and H. Aulich, Electroanal. Chem. Interfac. Chem., 1968, **18**, 295-307.
- 13 F. Jiao and H. Frei, Energy Environ. Sci., 2010, 3, 1018-1027.
- 14 M. M. Najafpour, T. Ehrenberg, M. Wiechen and P. Kurz, Angew. Chemie, 2010, 49, 2233-2237.
- 15 J-I Jung, H. Y. Jeong, J-S Lee, M. G. Kim, J. Cho, Angew. Chem. Int. Ed., 2014, 53, 4582-4586.
- 16 A. Mills and H. Davies, *Electrochim. Acta*, 1992, **37**, 1217– 1225.
- 17 A. Mills and G. Meadows, J. Chem. Soc., Faraday Trans., 1993. 89. 3849-3850.
- 18 A. Mills, D. Hazafy, S. Elouali and C. O'Rourke, J. Materials Chem. A, 2016, 4, 2863-2872.
- 19 R.L. Doyle, I.J. Goodwin, M.P. Brandon and M.E.G. Lyons, Phys. Chem. Chem. Phys., 2013, 15, 13737-13783.
- 20 F. Wilkinson, Chemical Kinetics and Reaction Mechanisms, Van Nostrand Reinhold, London, 1981, p. 140.
- 21 M. Spiro, Chem. Soc. Rev., 1986, 15, 141-165.
- 22 J. Horkans and M. W. Shafer, J. Electrochem. Soc., 1977, 124, 1202-1207.
- 23 A.I. Krasil'shchikov, Zh. Fiz. Chim., 1963, 37, 273-275.
- 24 C.C.L. McCory, S. Jung, I.M. Ferrer, S. M. Chatman, J. C. Peters and T.F. Jaramillo, J. Am. Chem. Soc.,, 2015, 137, 4347-4357
- 25 J. Kiwi and M. Gratzel, Angew. Chem. Int. Engl. 1978, 17, 860-861.