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Introduction

Water electrolysis is the most popular alternative to the production of hydrogen from fossil fuels as it is the only route that permits the use of renewable (*e.g.*, photovoltaic, wind, biomass, geothermal) energy sources combined with the production of 99.999% pure hydrogen.¹⁻³ Currently, only a small proportion of the world's hydrogen production (circa 4%) comes from electrolytic water splitting.¹ In fact, although water electrolysis is a well-known and consolidated process it does not have a significant commercial impact owing to its high-energy consumption, which, ultimately, makes it economically unattractive. The U.S. Department of Energy (DOE) has highlighted this drawback. Indeed, in 2011, the DOE has set a target that the electrical energy input to an electrolyzer stack should drop from 45 to 43 kW h kg⁻¹ H₂ by 2020.⁴

The three most common low temperature water electrolysis technologies may be categorized as follows;⁵

Carbon supported Rh nanoparticles for the production of hydrogen and chemicals by the electroreforming of biomass-derived alcohols[†]

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Electroreforming is a low energy cost technology that combines the production of valuable chemicals from biomass-derived alcohols with the evolution of clean hydrogen at low temperature and atmospheric pressure. The selectivity for the desired chemicals is governed by the nature of the anode catalyst. Here we report the synthesis and characterization of a carbon supported nanostructured Rh electrocatalyst. The Rh nanoparticles are shown to be highly dispersed (2.2 nm) and a complete electrochemical study is reported. This Rh/C catalyst exhibits high activity for alcohol electrooxidation (e.g. 5700 A g_{Rh} for EG at 80 °C) and when employed with an anion exchange membrane and Pt/C cathode in an electroreformer produces high volumes of hydrogen at low electrical energy input (e.g. 500 mA cm⁻² at 0.7 V_{cell} and $E_{cost} = 9.6$ kW h kg_{H₂}⁻¹). A complete analysis of the alcohol oxidation products from several renewable alcohols (ethanol, ethylene glycol, glycerol and 1,2-propandiol) shows a selectivity in the formation of valuable chemicals such as lactate and glycolate.

(i) Alkaline electrolysis with a liquid alkaline electrolyte (typically aqueous KOH).

(ii) Zero gap or advanced alkaline configuration.

(iii) Acidic PEM electrolysis with a proton-conducting polymer electrolyte membrane.

In alkaline electrolyte, water electrolysis proceeds as follows:

Anode reaction:
$$2OH^- \rightarrow \frac{1}{2}O_2 + H_2O + 2e^ E_a^0 = 0.404$$
 V (1)

Cathode reaction: $2H_2O + 2e^- \rightarrow H_2 + 2OH^ E_c^0 = -0.826$ V (2)

In acid electrolyte:

Anode reaction:
$$H_2O \rightarrow \frac{1}{2}O_2 + 2H^+ + 2e^ E_a^0 = 1.23$$
 V (3)

Cathode reaction:
$$2H^+ + 2e^- \rightarrow H_2$$
 $E_c^0 = 0.0$ V (4)

The standard reaction potential for water splitting is 1.23 V, meaning that the process is thermodynamically a non-spontaneous reaction. In practice, to get electrolysis current densities in the range of 1-2 A cm⁻² the electrolysis cell potential usually ranges between 1.6 and 2 V.⁶ Using 1.8 V as a reasonable value, we can calculate that 68.3% of the energy input is consumed by overcoming thermodynamics, while

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kinetic factors account for only 31.7%. Replacing anodic oxygen evolution with the oxidation of much more readily oxidizable species leads to a significant reduction of the potential required to produce hydrogen. Researchers have demonstrated that this strategy works using so-called sacrificial compounds such as ammonia,^{7,8} methanol,⁹ ethanol,¹⁰⁻¹³ glycerol^{11,12,14,15} and urea.¹⁶ This electrolysis technology combining the oxidation of such substrates with the generation of hydrogen at the cathode are often indicated as "Electrochemical Reforming" or "electroreforming". Electrolysis occurs at cell potentials lower than 1 V, leading to electrical power savings as compared to conventional electrolytic water splitting. Renewable biomass derived alcohols are attractive substrates for application in electrochemical reforming due to their low toxicity and their potential as part of carbon neutral energy transformation processes.^{17,18} Included in this group are ethylene glycol^{19,20} (EG) and glycerol^{21,22} (G). Ethylene glycol has a volumetric energy density of 5.9 kW h L^{-1} and can be produced by the heterogeneous hydrogenation of cellulose derivatives.^{23,24} Glycerol has a volumetric energy density of 6.3 kW h L^{-1} , and is a by-product of biodiesel production and, as such, is inexpensive (0.3 US kg⁻¹) and

readily available (2.4 million tonnes produced per year).²⁵ Eqn (5)–(7) describe the anode, cathode and overall electrochemical reforming reactions and standard electrode potentials (RHE) respectively for ethanol under alkaline conditions. With ethanol as fuel, the standard reaction potential for hydrogen production is 0.106 V. Therefore, hydrogen evolution is much more favorable when compared to water electrolysis where the standard potential is 1.23 V (RHE).

$$CH_3CH_2OH + 5OH^- \rightarrow CH_3COO^- + 4e^- + 4H_2O$$
(5)

 $E_{
m a}^0 = -0.72~{
m V}$

 $4H_2O + 4e^- \rightarrow 2H_2 + 4OH^ E_c^0 = -0.826$ V (6)

 $CH_3CH_2OH + OH^- \rightarrow CH_3COO^- + 2H_2 \quad E_{tot}^0 = 0.106 \text{ V}$ (7)

Other poly-alcohols can be oxidized provide not only hydrogen but also valuable chemicals from the formation of partially oxidized intermediates.^{26–29} For example, 1,2-propandiol can be employed in electroreforming to produce lactate, an industrially relevant chemical.^{30–33}

The efficiency of an electrolyzer is directly related to the electrode materials, in particular to the catalysts deposited onto the surface of the electrodes. The function of these catalysts is to reduce the activation energy for both the anode and cathode reactions. Therefore, a crucial role in improving the effective energetic efficiency of an electrolyzer is played by the electrode materials, because they determine both the energy consumption (at a given reaction rate) and the maximum reaction rate in the cell.

In this paper, we report an electroreformer that couples the partial oxidation of renewable alcohols with hydrogen evolution for the simultaneous and sustainable production of carboxylic compounds and high-purity hydrogen. The electroreformer is assembled with an anode containing nanostructured Rh/C electrocatalyst. This catalyst is composed of highly dispersed Rh nanoparticles supported on carbon, which shows high activity for alcohol electrooxidation. The employment of this catalyst enables high hydrogen production levels at very low energy cost. The paper also reports a net energy analysis of hydrogen production using bio-ethanol (as example) in order to clearly demonstrate the potential of electroreforming technology.

Experimental

Materials

Carbon black (Vulcan XC-72) was purchased from Cabot Corp., USA. The alkaline anion exchange membrane used was the A201 obtained from Tokuyama Corp. (Japan). All metal salts and reagents were purchased from Aldrich and used without further purification. All the solutions were freshly prepared with doubly distilled deionized water.

Synthesis of Rh/C anode catalyst

3.8 g of carbon powder (Vulcan XC-72, Cabot corp., USA) was suspended in 640 mL of ethylene glycol and sonicated for 80 min in a 2 L round-bottomed flask. Then an aqueous solution of H₂O (160 mL) containing 740 mg of dissolved RhCl₃- 6 H₂O (Aldrich), was added dropwise to the suspension under stirring. After this an alkaline solution of NaOH (15.8 g) in H₂O (80 mL) and ethylene glycol (200 mL) was introduced to the reactor which was then heated at 120 °C for 3 h under a N₂ atmosphere, under vigorous stirring. The mixture was then cooled to room temperature and the solid product was filtered off and washed thoroughly with H₂O to neutral pH. The solid product thus obtained was dried under vacuum at 40 °C. Yield 3.67 g. ICP-AES analysis: Rh content = 6 wt% (ICP-AES).

Physical characterization

Transmission electron microscopy (TEM) was performed using a Philips CM12 microscope at an accelerating voltage of 100 kV. The microscope was equipped with an EDAX energy dispersive microanalysis system.

The BET surface area was determined using an ASAP 2020C Instrument (Micromeritics Corp.).

X-ray photoelectron Spectroscopy (XPS) measurements were performed in a system equipped with a VSW HAC 5000 hemispherical electron energy analyzer and a non-monochromated Mg K α X-ray source (1253.6 eV). Photoelectron spectra were acquired in the constant-pass-energy mode at $E_{pas} = 44$ eV, and the overall energy resolution was 1.2 eV measured as a fullwidth at half maximum (FWHM) of the Ag 3d_{5/2} line of a pure silver reference. The pressure during the experiment was kept below 2 × 10⁻⁹ Torr. No neutralizer was utilized and the spectra energy scale was corrected using the Au 4f_{7/2} peak. The recorded spectra were fitted using XPS Peak 4.1 software. Gauss–Lorentz curves were used to fit the data after subtraction of a Shirleytype background.

Electrochemical characterization

The electrochemical measurements were performed using a Parstat 2273 potentiostat-galvanostat (Princeton Applied

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Research) equipped with a Model 616 rotating disk electrode RDE (PAR-Ametek). A 5 mm (A = 0.1963 cm²) Teflon-potted glassy-carbon disk electrode tip (PINETM) was used as substrate for the deposition of the catalyst ink. Before the deposition, the glassy-carbon surface was polished with different CT diamond suspension with a progressively smaller particle size (1 µm, 0.25 µm, 0.1 µm) and finally washed with distilled water. The catalyst ink was prepared by mixing the Rh/ C catalyst and distilled water in order to obtain 1 wt% suspension. The ink was sonicated for 1 h in a FALC sonicating bath to obtain a uniform suspension. The catalyst film was prepared by dispersing 8 µL of the catalyst ink on the glassy-carbon electrode. The exact amount of dry catalyst deposited was determined using an analytical balance. The final Rh loading on the electrode was between 5 and 6 μ g (25–30 μ g cm⁻²). Each electrode was dried for 30 min before the addition of 2.5 µL of a 0.4 wt% Tokuyama[™] OH-type anion exchange resin alcohol solution (AS4). After drying, the catalyst-ionomer coated disk was then mounted on the rotating disk electrode shaft and immersed into the electrolyte solution. The counter electrode was a platinum gauze enclosed in a glass tube with porous bottom.

The reference electrode used was commercial Ag/AgCl/KCl_{sat} (Princeton Applied Research). All cyclic voltammetry potentials are reported against reversible hydrogen electrode (RHE) calculated taking into account the potential-temperature dependence, as described in the ESI.† The electrochemical experiments were conducted at three temperatures (25, 60 and 80 °C). The CV experiments were undertaken in 2 M aqueous KOH (Sigma-Aldrich, 99.8%) while the alcohol electrooxidation studies were performed in aqueous solutions of 2 M KOH and 2 M alcohol. All the solutions were prepared with Millipore water (18 M Ω cm⁻¹) provided by a Milli-Q Labo apparatus (Nihon Millipore Ltd.). Solutions were deaerated by bubbling high-purity N₂ for 30 min before each measurement.

CO stripping voltammograms were performed in aqueous 2 M KOH. As first step, the KOH solution was saturated with bubbling CO for 20 min, after which a fixed voltage of 70 mV (RHE) was applied for 15 min to the working electrode to allow the complete adsorption of a monolayer of CO onto the surface of the catalyst. Then the excess CO in the electrolyte was purged out with bubbling N₂ for 20 min. The amount of CO_{ads} was calculated integrating the respective stripping peak and the corresponding charge was used to extrapolate and calculate the ECSA of the Rh/C catalyst.

Electroreformer cell testing

The MEAs (membrane electrode assemblies) prepared for the electroreformer cell consist of the Rh/C catalyst as anode, a commercial Tokuyama A-201 anion-exchange membrane, and a commercial 40 wt% Pt/C catalyst (Aldrich) as cathode.

A dense anode ink was prepared by mixing the desired quantity of powdered catalyst Rh/C with an aqueous dispersion of PTFE (60 wt% PTFE dispersion in H_2O obtained from Aldrich) and distilled water. The amount of PTFE added was 5 wt% relative to the dry catalyst weight. The catalyst paste was applied uniformly to a porous nickel foam support (110 PPI

HZTYKJ from Heze Tianyu Technology Development Co., Ltd. dimensions 5 cm²). The cathodic ink was prepared in a 5 mL high density polyethylene vial, mixing 200 mg of the commercial Pt (40 wt%)/C in 450 mg of distilled water, 790 mg of 1-propanol and 1.56 g of the ionomer Nafion® (5 wt% in 2-propanol). The mixture was suspended with three pulses of ultrasound, 20 W power at the frequency of 20 kHz (Bandelin Sonor pulse UW 2200 SERIES). Finally this paste was spread onto a carbon cloth W1S1005 (CeTech Co. Ltd.) gas diffusion layer, with a Meyer rod (no. 150) obtaining a 0.4 mg cm⁻² Pt loading.

The electroreformer test cell fixture was purchased from Scribner-Associates (USA). The MEAs were assembled by mechanically pressing together the anode, cathode and membrane described above within the cell hardware. The cell temperature was regulated at 60 °C using a Scribner 805e fuel cell station. The aqueous fuel solution (30 cc of 2 M KOH and 2 M alcohol) was delivered to the anode at 1 mL min⁻¹. Voltage scans and galvanostatic experiments were carried out using an ARBIN BT-2000 5A 4-channel instrument. Polarization experiments were recorded by applying a linear voltage ramp with a 10 mV s⁻¹ scan rate between 0.2 to 0.8 V.

Alcohol electrooxidation selectivity study

Chronopotentiometry experiments were performed by applying a constant electrolysis current of 125 mA to the cell at 60 °C until the voltage reached the cut off value of 0.65 V. The hydrogen produced at the cathode was monitored by a Bronkhorst massspecific hydrogen-calibrated flowmeter. The fuel solutions after the constant current experiments were quantitatively and qualitatively analyzed by ${}^{13}C{}^{1}H$ NMR spectroscopy and HPLC. A UFLC Shimadzu Chromatograph equipped with refraction index detector (RID) was used; the column is a GRACE-Alltech OA-1000 Organic Acids (300 mm \times 6.5 mm), thermostated at 35 °C. The eluent is 0.01 N H₂SO₄; and the eluent flow is 0.8 mL min^{-1} . NMR spectra were acquired with a with a Bruker Advance DRX 400 spectrometer. Chemical shifts (δ) are reported in ppm relative to TMS (¹H and ¹³C NMR spectra). Deuterated solvents (Sigma-Aldrich) used for NMR measurements were dried with activated molecular sieves; 1,4-dioxane was used as internal standard for product quantification.

Results and discussion

Catalyst synthesis and characterization

The Rh/C catalyst was prepared by electro-less reduction in aqueous media of Vulcan XC-72-impregnated with RhCl₃ using ethylene glycol at 120 °C for 3 h under inert atmosphere. The morphology and distribution of the supported metal particles was evaluated by transmission electron microscopy (TEM in Fig. 1). The particle size distribution of the Rh/C catalyst is also shown and has a mean diameter value of 2.2 nm.

The crystalline structure of the catalyst was also investigated by X-ray powder diffraction (XRD). The XRD pattern (Fig. SI[†]) does not reveal the presence of a metallic Rh phase, consistent with the formation of small nanoparticles of Rh in agreement



Fig. 1 TEM image and histogram of particle distribution versus diameter for Rh/C.

with the TEM analysis. The Brunauer–Emmett–Teller (BET) surface area was calculated as 195 m² g⁻¹.

The surface composition of the Rh/C sample was also studied using X-ray photoelectron spectroscopy (XPS). The fitting of the data obtained shows the presence of three peaks (from the binding energy values) attributable to Rh(0) 307.3 eV, Rh₂O₃ 308.8 eV (ref. 34–36) and a satellite peak of Rh₂O₃ 311.5 eV (Fig. 2). The latter species was recently assigned by Boronin and coworkers.³⁷ This analysis shows that rhodium oxide is the major component of the surface of the nanoparticles with Rh metal accounting for only around 35%. Such a structure would lead to the XRD pattern that showed no visible trace for metallic Rh crystallites.

Electrochemical properties

The electrochemical activity of Rh/C for alcohol electrooxidation was investigated by linear sweep voltammetry in electrochemical cells maintained at three different temperatures (25, 60 and 80 °C). For all experiments, the Rh loading on the glassy carbon substrate was between 25 and 35 μ g_{Rh} cm⁻². We first studied the electrochemical behavior by cyclic voltammetry (CV) in 2 M KOH purged with N₂ (Fig. 3).



Fig. 2 XPS spectra in the Rh 3d region. Spectra are referenced with respect to adventitious carbon at 284.8 eV.



Fig. 3 CVs of Rh/C in N₂ saturated 2 M KOH. Scan rate: 10 mV s⁻¹.

As reported in the literature the CV exhibits a number of anodic and cathodic transitions that can be assigned to Rh centered transitions.^{38,39} On the positive potential scan (0 to 1.2 V), peak A' represents the oxidative desorption of hydrogen; peak A'' relates to the formation of adsorbed OH species on the Rh nanoparticle surface and peak A''' is the onset of the formation of Rh oxide species. On the cathodic return scan peak C' corresponds to the electroreduction of the oxide layer, and peak C'' to the electrosorption of H-adatoms.

CVs obtained at increasing temperatures (60 and 80 °C) show significant changes to the intensity and potential of these transitions. Peak A' shifts by approximately 0.1 V towards more negative potentials, while decreasing in intensity. Peak A'' increases in intensity and shifts about 75 mV towards more negative potentials. This behavior indicates an increased ability for the electrooxidation of alcohols, as it shows that RhOH_{ads} species form at lower potentials (A''). The RhOH_{ads} species are those responsible for alcohol electrooxidation, analogously to the PdOH_{ads} species formed onto palladium electrocatalysts, albeit formed at lower potential for Rh.

The electrochemically active surface area (ECSA) of the Rh nanoparticles was determined by integrating the charge of the

electrooxidation peak of an adsorbed monolayer of CO_{ads} in CO stripping voltammograms (Fig. 4).40 The ECSA normalized by the mass of Rh was calculated as 75 m² g_{Rh}^{-1} .

The electrochemical activity of Rh/C towards the oxidation of alcohols in alkaline media was investigated by cyclic voltammetry at 25 °C, 60 °C and 80 °C, in degassed 2 M aqueous KOH and 2 M of either ethanol (EtOH), 1,2-propandiol (1,2-P), glycerol (G) or ethylene glycol (EG) (Fig. 5). Relevant electrochemical data is also listed in Table 1. In Fig. 5, we show the onset region (0.2-0.6 V RHE) of the forward scan for each alcohol at all three temperatures (25, 60 and 80 °C).

From Fig. 5 and Table 1, we can observe low oxidation onset potentials, which generally decrease as the temperature increases from 25 °C to 80 °C, whereas the specific current densities dramatically increase with temperature for all alcohols, up to 35 times for EG. The intensity of the forward peaks in the CVs is determined by the progressive coverage of the Rh surface with oxide or partially oxidized poisoning species. In the complete CVs, (Fig. S2[†]) for ethanol a change of slope at 0.7 V is observed, caused by different adsorption types of ethanol on Rh.41 The enhanced electrocatalytic activity observed at higher temperatures (60-80 °C) can be ascribed to the accelerated kinetics of alcohols oxidation, which it is also favored by an

0.12 first cycle 0.1 second cycle 0.08 Specific activity / mA µg_{Rh}⁻¹ 0.06 0.04 0.02 0 -0.02 -0.04

Fig. 4 CO-stripping voltammograms of Rh/C in 2 M KOH. Scan rate: 10 mV s^{-1} .

anticipated formation of RhOH_{ads} species at lower potentials as observed in Fig. 3.

Electrochemical-reforming experiments

Nickel foam (5 cm²) coated with Rh/C and using PTFE as binder was used as anode electrode (1 $mg_{Rh} cm^{-2}$) in the electroreformer (ER) cell equipped with an anion exchange membrane (Tokuyama A201) and a Pt/C on carbon cloth cathode (0.4 mg_{Pt} cm^{-2}). The anode compartment of the cell was fed by recycling 30 mL of aqueous solutions of alcohol 2 M and 2 M KOH. At a cell temperature of 60 °C the performance was measured using potentiodynamic scans and galvanostatic experiments. For each alcohol, three batches of fuel were run consecutively in the same cell.

Polarization curves were obtained in the potential range from 0.2 to 0.7 V at 10 mVs⁻¹. Fig. 6a and Table 2 show that the electroreformer performs best with EtOH and EG reaching current densities of 492 mA cm⁻² and 416 mA cm⁻², respectively at a cell voltage of 0.7. With the longer chain alcohols, 1,2-P and G, the current density obtained at 0.7 V were 367 mA cm $^{-2}$ and 113 mA cm^{-2} respectively.

In all cases, the cell performance was stable over the three batch experiments indicating no loss in performance during

Table 1 Relevant electrochemical parameters for alcohol electro-

Eforward peak

(V vs. RHE)

J_{peak}^a

45

95

167

103

320

521

2728

287

500

886

5767

1839

 $(\text{mA mg}_{\text{Rh}}^{-1})$

Jpeak/ECSA $(mA cm^{-2})$

0.060

0.127

0.223

0.137

0.427

0.695

3.637

0.383

0.667

1.155

7.689

2.452

oxidation at 25, 60 and 80 °C on electrodes coated with Rh/C

0.56

0.55

0.59

0.57

0.56

0.53

0.57

0.54

0.59

0.49

0.57

0.55

Eonset

0.34

0.35

0.33

0.46

0.32

0.33

0.32

0.44

0.30

0.31

0.28

0.31

(V vs. RHE)

 $T(^{\circ}C)$

25

25

25

25

60

60

60

60

80

80

80

80

Fuel

EtOH

1,2-P

EtOH

1,2-P

EtOH

1,2-P

EG

G

EG

G

EG

G



^a Current normalized on total Rh metal loading. ^b Current normalized on Rh electrochemical surface area (ECSA) i.e. 75 m² g_{Rh}⁻



CVs of Rh/C in N₂ sat. 2 M KOH and 2 M alcohol solutions: (a) 25 °C, (b) 60 °C and (c) 80 °C (0.2–0.6 V RHE). Scan rate: 10 mV s⁻¹. Fia. 5



Fig. 6 (a) Potentiodynamic cycles of the electroreformer fueled with: 2 M aqueous alcohol solutions in 2 M KOH. Linear potential ramp, 10 mV s^{-1} . Cell temperature 60 °C; (b) galvanostatic curves at 125 mA, fueled with 30 cc of 2 M aqueous alcohol solutions in 2 M KOH. Cell temperature 60 °C.

Table 2 Catalytic data of electroreforming with Rh/C at the anode at 60 °C

Fuel	$J_{ m max}$ (a) 700 mV (mA cm ⁻²)	Conversion (%) [mmol]	H_2 production (Nm ³ m ⁻²)	Energy consumption (kW h kg ⁻¹ H ₂)	Selectivity%
EtOH	492	57 [34.2]	3	9.6	Acetate (100)
1,2-P	367	55 [33.0]	2.8	14.4	Lactate (98) Carbonate (1) Acetate (1)
EG	416	89 [53.4]	5.4	11.0	Glycolate (76) Carbonate (16) Oxalate (5) Formate (3)
G	113	72 [43.2]	4.3	12.6	Glycerate (46) Tartronate (23) Carbonate (18) Formate (8) Glycolate (3) Oxalate (2)

electrocatalysis. In order to study both the stability and the electrocatalytic activity over time as well as the electrooxidation product selectivity, the electroreformer was subjected to galvanostatic experiments where 30 mL of solutions of 2 M alcohol and 2 M KOH were recycled at the anode electrode and the cell was set at a constant total current of 125 mA. Each batch was stopped when the cell potential reached the value of 0.65 V. The same set-up was run with three consecutive batches of fresh alcohol solutions. No decrease in activity was observed over the course of the three batches, indicating the stability of the catalyst over time. Fig. 6b and Table 2 summarize the data obtained at 60 $^{\circ}$ C.

The fuel exhausts were quantitatively and qualitatively analyzed by ${}^{13}C{}^{1}H$ NMR spectroscopy and HPLC. The results summarized in Table 2 show the fuel conversion and the distribution of the oxidation products identified and quantified by NMR and HPLC analysis.

With ethanol, the only product obtained is acetate with no evidence of C–C bond breaking and formation of carbonate. The cells fed with EG and 1,2-P yielded circa 76% glycolate and 98% of lactate, respectively. Glycerol oxidation leads to significant C–C bond scission, resulting in a mixture of products including glycerate, tartronate and also oxalate, glycolate, formate and carbonate, all from C–C bond breaking reactions. No trace of the secondary alcohol oxidation products of G such as dihydroxyacetone, hydroxypiruvate, or mesoxalate was observed. As expected, at the cell working potentials no oxygen evolution at the anode was detected.

The amount of hydrogen evolved during each experiment was calculated from the number of moles of electrons exchanged in the electroreformer (see ESI[†]) and measured experimentally with an hydrogen-calibrated flowmeter; the values were consistent indicating a 100% faradaic efficiency for the cathodic process. The energy consumption was then evaluated from the integration of the instantaneous charging power over the experimental duration time. This value, expressed in kW h, is reported with respect to the amount of hydrogen produced in kilograms (*i.e.* kW h kg_{H₂}⁻¹). Table 2 shows that the amount of electrical energy required for H₂ production for all alcohols is in the range of 10–14 kW h kg_{H₂}⁻¹.

Our recent reports of alcohol electroreforming with Pd based anode catalysts showed electrical energy consumption for the same levels (*i.e.* same current density) of H₂ production in the region of 18–24 kW h kg⁻¹.^{30,31} Hence, the use of Rh/C as anode catalyst significantly reduces the energy cost of electroreforming. In particular, we can calculate that the use of ethanol (9.6 kW h kg⁻¹ H₂) results in an electrical energy saving of 35.4 kW h kg⁻¹ H₂ as compared with that reported by the DOE for PEM water electrolyzer stacks as of 2011 (45 kW h kg⁻¹ H₂).⁴ The reason of this drop in energy consumption is due to the lower operating potentials for alcohol electrooxidation caused by the cathodic shift of the onset of the formation of RhOH_{ads} species, that are responsible for the alcohol electrooxidation process itself.

One may now consider if from an overall energetic point of view the exploitation of alcohols such as ethanol for hydrogen production by electroreforming is actually worthy of consideration. According to the DOE the output energy density for bioethanol is 7.4 kW h kg_{EtOH}⁻¹. From our calculations reported in the SI, we conclude that the production of bioethanol with an Energy Return of the Energy Invested (EROEI) larger than 3.1 is required in order to obtain a net energy saving with respect to current water electrolysis technology (47 kW h kg⁻¹ H₂). The DOE target for electrolyzer stack electric energy input for 2020,4 43 kW h kg⁻¹ H₂, is instead met for a bio-ethanol EROEI larger than 3.6. Hence, hydrogen production by the electroreforming of bio-ethanol is energetically convenient depending on the production source of the alcohol e.g. using sugarcane that is commonly reported to have an EROEI of 8 and higher⁴² from cellulose with an EROEI potentially up to 35 depending on production methods.43

Conclusions

In this paper, examples of electrooxidation of biomass-derived alcohols such as EtOH, EG, G and 1,2-P have been studied in an electroreformer containing Rh nanoparticles supported on carbon as the anode electrocatalyst, equipped with an anion exchange membrane and a Pt/C on carbon cloth cathode. The oxidation of alcohols was first investigated in electrochemical half-cells at room temperature and at 60–80 °C in alkaline media. The results highlighted the excellent activity of Rh/C in terms of peak current densities (as high as 5700 A g_{Rh}^{-1} for EG at 80 °C) and low onset of potentials.

When employed as anode catalyst in electroreformer cells high current densities were obtained at cell potential well below 1 V, resulting in significant energy saving when compared to traditional water electrolyzers. The amount of electrical energy required for H₂ production for all alcohols is in the range of 10– 14 kW h kg_{H₂}⁻¹. These values are the lowest compared to those reported up to now in the literature. In particular, 9.6 kW h kg_{H₂}⁻¹ from EtOH results in an electrical energy saving of 35.4 kW h kg_{H₂}⁻¹ as compared with that reported by the DOE for PEM electrolyzer.

In terms of oxidation product selectivity, this Rh/C catalyst was selective for EtOH electrooxidation to acetate (100%) and 1,2-P oxidation to lactate (98%). From EG and G a more complex

mixture of partial oxidation products was obtained. In summary, the electroreformer reported in this paper provides not only hydrogen production but also valuable chemicals from the large variety of products that can be obtained by the electrooxidation of renewable alcohols. The remarkable electrocatalytic activity of Rh/C employed as anode catalyst can be attributed to both the high dispersion of the metal nanoparticles as well as the intrinsic properties of nanostructured metallic Rh that shows low onset potentials for alcohol electrooxidation in half cells and low cell working potential during electroreforming.

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References

- 1 J. Mergel, M. Carmo and D. Fritz, *Transition Renewable Energy Syst.*, 2013, 425–450, DOI: 10.1002/9783527673872.
- 2 A. Ursua, L. M. Gandia and P. Sanchis, *Proc. IEEE*, 2012, **100**, 410–426.
- 3 J. A. Turner, Science, 2004, 305, 972-974.
- 4 Fuel Cell Technologies Office, Fuel cells, Fuel Cell Technologies Office Multi-year Research, Development and Demonstration Plan of the US Department of Energy, 2011.
- 5 S. Marini, P. Salvi, P. Nelli, R. Pesenti, M. Villa, M. Berrettoni,
 G. Zangari and Y. Kiros, *Electrochim. Acta*, 2012, 82, 384–391.
- 6 M. Carmo, D. L. Fritz, J. Merge and D. Stolten, *Int. J. Hydrogen Energy*, 2013, **38**, 4901–4934.
- 7 M. Schalenbach, M. Carmo, D. L. Fritz, J. Mergel and D. Stolten, *Int. J. Hydrogen Energy*, 2013, 38, 14921–14933.
- 8 F. Vitse, M. Cooper and G. G. Botte, *J. Power Sources*, 2005, **142**, 18–26.
- 9 T. Take, K. Tsurutani and M. Umeda, *J. Power Sources*, 2007, **164**, 9–16.
- 10 C. Lamy, T. Jaubert, S. Baranton and C. Coutanceau, *J. Power Sources*, 2014, 245, 927–936.
- V. Bambagioni, M. Bevilacqua, C. Bianchini, J. Filippi, A. Lavacchi, A. Marchionni, F. Vizza and P. K. Shen, *ChemSusChem*, 2010, 3, 851–855.
- A. Caravaca, F. M. Sapountzi, A. de Lucas-Consuegra, C. Molina-Mora, F. Dorado and J. L. Valverde, *Int. J. Hydrogen Energy*, 2012, 37, 9504–9513.
- 13 A. Caravaca, A. de Lucas-Consuegra, A. B. Calcerrada,
 J. Lobato, J. L. Valverde and F. Dorado, *Appl. Catal., B*, 2013, 134, 302–309.
- 14 A. T. Marshall and R. G. Haverkamp, *Int. J. Hydrogen Energy*, 2008, **33**, 4649–4654.
- 15 S. Kongjao, S. Damronglerd and M. Hunsom, *J. Appl. Electrochem.*, 2011, **41**, 215–222.
- 16 W. Yan, D. Wang and G. G. Botte, *Appl. Catal., B*, 2012, **127**, 221–226.
- 17 A. Brouzgou, A. Podias and P. Tsiakaras, *J. Appl. Electrochem.*, 2013, **43**, 119–136.

- 18 E. H. Yu, U. Krewer and K. Scott, *Energies*, 2010, 3, 1499–1528.
- 19 V. Livshits, A. Philosoph and E. Peled, *J. Power Sources*, 2008, **178**, 687–691.
- 20 D. Kaplan, L. Burstein, Y. Rosenberg and E. Peled, *J. Power Sources*, 2011, **196**, 8286–8292.
- 21 L. Xin, Z. Y. Zhang, Z. C. Wang and W. Z. Li, *ChemCatChem*, 2012, 4, 1105–1114.
- 22 A. Marchionni, M. Bevilacqua, C. Bianchini, Y. X. Chen, J. Filippi, P. Fornasiero, A. Lavacchi, H. Miller, L. Q. Wang and F. Vizza, *ChemSusChem*, 2013, 6, 518–528.
- 23 N. Ji, T. Zhang, M. Y. Zheng, A. Q. Wang, H. Wang, X. D. Wang and J. G. G. Chen, *Angew. Chem., Int. Ed.*, 2008, 47, 8510–8513.
- 24 M. Y. Zheng, A. Q. Wang, N. Ji, J. F. Pang, X. D. Wang and T. Zhang, *ChemSusChem*, 2010, 3, 63–66.
- 25 J. Van Gerpen, Fuel Process. Technol., 2005, 86, 1097-1107.
- 26 V. Bambagioni, M. Bevilacqua, C. Bianchini, J. Filippi, A. Lavacchi, A. Marchionni, F. Vizza and P. K. Shen, *ChemSusChem*, 2010, 3, 851–855.
- 27 J. de Paula, D. Nascimento and J. Linares, *J. Appl. Electrochem.*, 2015, **45**, 689–700.
- 28 S. Gonzalez-Cobos, S. Baranton and C. Coutanceau, *ChemElectroChem*, 2016, 3, 1694–1709.
- 29 Z. Y. Zhang, L. Xin, J. Qi, D. J. Chadderdon, K. Sun, K. M. Warsko and W. Z. Li, *Appl. Catal.*, *B*, 2014, 147, 871– 878.
- 30 Y. X. Chen, A. Lavacchi, H. A. Miller, M. Bevilacqua, J. Filippi,
 M. Innocenti, A. Marchionni, W. Oberhauser, L. Wang and
 F. Vizza, *Nat. Commun.*, 2014, 5, 4036–4041.

- 31 H. A. Miller, M. Bellini, F. Vizza, C. Hasenohrl and R. D. Tilley, *Catal. Sci. Technol.*, 2016, 6, 6870–6878.
- 32 D. J. Chadderdon, L. Xin, J. Qi, B. Brady, J. A. Miller, K. Sun, M. J. Janik and W. Z. Li, ACS Catal., 2015, 5, 6926–6936.
- 33 M. Bellini, M. Bevilacqua, J. Filippi, A. Lavacchi,
 A. Marchionni, H. A. Miller, W. Oberhauser, F. Vizza,
 S. P. Annen and H. Grutzmacher, *ChemSusChem*, 2014, 7, 2432–2435.
- 34 M. H. M. T. Assumpção, R. M. Piasentin, P. Hammer, R. F. B. De Souza, G. S. Buzzo, M. C. Santos, E. V. Spinacé, A. O. Neto and J. C. M. Silva, *Appl. Catal.*, B, 2015, 174–175, 136–144.
- 35 Q. He, S. Mukerjee, B. Shyam, D. Ramaker, S. Parres-Esclapez, M. J. Illán-Gómez and A. Bueno-López, *J. Power Sources*, 2009, **193**, 408–415.
- 36 M. A. Montero, J. L. Fernández, M. R. Gennero de Chialvo and A. C. Chialvo, *J. Power Sources*, 2014, **254**, 218–223.
- 37 L. S. Kibis, A. I. Stadnichenko, S. V. Koscheev, V. I. Zaikovskii and A. I. Boronin, *J. Phys. Chem. C*, 2016, **120**, 19142–19150.
- 38 Z. Cataldi, R. O. Lezna, M. C. Giordano and A. J. Arvia, J. Electroanal. Chem. Interfacial Electrochem., 1989, 261, 61–75.
- 39 M. Łukaszewski, H. Siwek and A. Czerwiński, *Electrochim. Acta*, 2007, **52**, 4560–4565.
- 40 M. A. Montero, M. R. Gennero de Chialvo and A. C. Chialvo, *J. Power Sources*, 2015, **283**, 181–186.
- 41 S. Y. Shen, T. S. Zhao and J. B. Xu, *Int. J. Hydrogen Energy*, 2010, 35, 12911–12917.
- 42 J. Goldemberg, Science, 2007, 315, 808-810.
- 43 C. A. S. Hall, B. E. Dale and D. Pimentel, *Sustainability*, 2011, 3, 2413–2432.