NJC

PAPER

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Cite this: New J. Chem., 2019, 43, 13985

Received 24th April 2019, Accepted 1st August 2019

DOI: 10.1039/c9nj02114k

rsc.li/njc

Introduction

The growing demand for natural gas and petroleum based feedstock for power generation, transport and chemicals production has, in recent decades, resulted in an unsustainable increase in CO₂ emissions.^{1,2} Irreversible climate change, diminishing conventional fossil fuel reserves and the complexities involved in extracting unconventional fossil reserves, are driving the development of sustainable, renewable and non-fossil-based feedstock for the production of chemicals and fuels. In spite of its thermodynamic stability, carbon dioxide has become an attractive C-1 feedstock for the production of chemicals and fuels.³⁻⁵ With the recent advances in carbon capture technologies, cost-competitive pure CO₂ could become available for its conversion to valueadded chemicals.⁶ Economically sustainable large-scale CO₂ conversion could help in achieving the emission targets set out in the Paris Agreement⁷ while increasing the production of chemicals and fuels.

Commercially, CO_2 utilisation is thus far limited to the production of a few chemicals, including urea⁸ (for nitrogen fertilizers and plastics), polycarbonates⁹ (for plastics), salicylic acid (a pharmaceutical ingredient)¹⁰ and methanol.⁵ Among all

Liquid phase hydrogenation of CO₂ to formate using palladium and ruthenium nanoparticles supported on molybdenum carbide[†]

Claire E. Mitchell, ២ ^a Umberto Terranova, ២ ^a Ihfaf Alshibane, ^b David J. Morgan, ២ ^a Thomas E. Davies, ២ ^a Qian He, ^a Justin S. J. Hargreaves, ^b Meenakshisundaram Sankar 🕩 * ^a and Nora H. de Leeuw ២ * ^a

We report the development of palladium nanoparticles supported on Mo₂C as an active catalyst for the liquid-phase hydrogenation of CO₂ to formate under mild reaction conditions (100 °C and 2.0 MPa of a $1:1 \text{ CO}_2:\text{H}_2$ mixture). A series of Pd/Mo₂C catalysts were synthesised *via* the modified wet-impregnation (MIm) and sol-immobilization (SIm) techniques and evaluated for CO₂ hydrogenation, in an aqueous 1 M NaOH solution. MIm catalysts synthesised using PdCl₂ dissolved in a 2 M HCl solution gave the highest formate yield with turnover numbers of up to 109 after 19 h. We further report the crucial role of base and the pH of the reaction medium for the hydrogenation of CO₂ to formate. Based on stability studies, electron microscopic characterisation and density functional theory calculations we found that Ru has a stronger affinity than Pd to Mo₂C resulting in the development of a stable bimetallic RuPd/Mo₂C catalyst for the hydrogenation of CO₂ to formate.

the possible transformations, CO_2 hydrogenation to acids, alcohols and hydrocarbons using H_2 derived from non-fossil feedstock (*e.g.* water splitting using electricity derived from a renewable source) is the most promising strategy.¹¹ Here, it is important to note that recently tremendous progress has been made in the production of H_2 *via* water-splitting using electricity.¹² CO_2 hydrogenation is challenging because of the inherent thermodynamic stability of CO_2 along with the lack of active catalysts that can activate CO_2 under reasonable reaction conditions.⁴ However, the potential of such an abundant feedstock for producing organic compounds, coupled with the possibility of balancing CO_2 emissions make CO_2 hydrogenation an exciting and challenging topic for research.

An important product that is formed from the hydrogenation of CO_2 is formic acid, which is a liquid at room temperature and contains 4.4 wt% of hydrogen.¹³ Current annual production of formic acid is around 600 000 tons and this is projected to grow by 22% annually.¹⁴ Formic acid is a vital intermediate in many industries including dyeing, leather, food, agrochemical and many more.^{15–17} It is also used to promote the fermentation of acetic acid, as a coagulant in rubber synthesis, an antibacterial agent in animal feed and as a de-icing agent in many manufacturing industries.¹⁸ Using appropriate catalysts, formic acid can be controllably converted to CO_2 and H_2 even at room temperature.¹⁹ Hence, hydrogenation of CO_2 to formic acid is considered as an effective strategy to store H_2 chemically. Current technology for the production of formic acid involves

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^a Cardiff Catalysis Institute, School of Chemistry, Cardiff University,

Cardiff CF10 4AT, UK. E-mail: Sankar@cardiff.ac.uk, deleeuwn@cardiff.ac.uk

^b School of Chemistry, Glasgow University, Glasgow G12 8QQ, UK

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/c9nj02114k

the hydrolysis of methyl formate, with a capacity of *ca.* 770 kilotons per annum in 2014.²⁰ Limitations of this process include the use of fossil fuel based feedstock, slow reaction rate, undesirable by-products and high cost. Therefore, it is not surprising that utilization of CO_2 is gaining momentum in the scientific community in order to shift from conventional fossil-based processes toward environment-friendly direct hydrogenation of CO_2 to formic acid.

Most of the reported catalysts for the hydrogenation of CO₂ to formic acid are homogeneous metal (Rh, Ru, Ir and Fe) complexes containing sophisticated ligands (N-heterocyclic carbenes, pincer ligands and phosphines) that require complicated synthesis and/or handling procedures.²¹⁻²⁴ Although these homogeneous catalysts are highly active, heterogeneous catalysts are preferred for a number of reasons, e.g. catalyst separation and scalability. Transition metal carbides have been reported to exhibit catalytic activities for a number of chemical reactions, including CO hydrogenation,25 the water gas shift reaction,²⁶ hydrodesulphurization²⁷ and methane reforming.²⁸ Recently, transition metal carbides have also been reported as catalysts for CO₂ hydrogenation²⁹ including by Dubois et al.³⁰ and Vidal et al.³¹ All these reactions were carried out at relatively high temperatures (220-320 °C), and mostly in the gas-phase, where the most interesting products like CH₃OH or HCOOH are thermodynamically not favoured.^{32,33} Hence, it is desirable to perform the hydrogenation of CO₂ at lower temperatures where CO (a product from an endothermic reaction) is not favoured.

Chen et al. demonstrated this by performing CO₂ hydrogenation at 200 °C in the liquid phase for the production of MeOH, they reported catalysis using metal nanoparticles supported on Mo₂C using 1,4-dioxane as the solvent.^{34,35} They synthesized a series of M/Mo₂C (M = Pd, Cu, Co and Fe) catalysts, for the formation of CH₃OH, C₂H₅OH and C₂₊ hydrocarbons. 1,4-Dioxane however, is not an environmentally-benign solvent of choice,³⁶ other organic solvents have also been reported for the liquid phase hydrogenation of CO₂, including ethanol.³⁷ Organic solvents have the disadvantage of being involved in the reaction itself, and may play a role in the synthesis of the targeted products; hence, it is better to avoid using them. It has been reported that addition of water is effective for improving CO₂ hydrogenation to formic acid.^{38,39} It is proposed that the hydrogen-bonding between the H₂O molecule and CO₂ improves the electrophilicity of the carbon atom on the CO₂ molecule,³⁹ thus reducing the reaction barrier for CO₂ activation. The same group reported Mo₂C acted as a co-catalyst as well as a support as it performed CO2 conversion without any other metals present. Posada-Perez et al. compliment this by explaining that the specific carbon/ metal ratio of Mo₂C is responsible for the high reactivity resulting in the breaking of both C–O bonds in CO₂ before hydrogenation.⁴⁰ Mori et al. reported the formation of formic acid from CO₂ hydrogenation using a ruthenium-based catalyst, in water under alkaline conditions at 100 °C,⁴¹ Song et al. also reported hydrogenation of CO₂ to formic acid in water using a Na₂CO₃ base, catalysed by palladium supported on chitin.42 These alkaline conditions are important in improving the CO₂ solubility in water. CO₂ has poor solubility in pure water $(0.0693 \text{ mol kg}^{-1} \text{ at } 1 \text{ bar at } 30 \degree \text{C})^{43}$ and the solubility decreases with increasing temperature. Carbonic acid (H_2CO_3), formed when CO_2 dissolves in water, deprotonates sequentially to form the carbonate ion (eqn (R1)–(R3)).

$$CO_{2(aq)} + H_2O \leftrightarrow H_2CO_{3(aq)} \quad pH = 4-6$$
 (R1)

$$H_2CO_{3(aq)} \leftrightarrow H^+_{(aq)} + HCO_{3(aq)} pH = 6-10$$
 (R2)

$$\mathrm{HCO}_{3(\mathrm{aq})}^{-} \leftrightarrow \mathrm{H^{+}_{(\mathrm{aq})}} + \mathrm{CO}_{3(\mathrm{aq})}^{2-} \quad \mathrm{pH} > 10 \qquad (\mathrm{R3})$$

Because of these equilibria, different species dominate the aqueous medium at different pH,⁴⁴ as noted above. Consequently, pH is an important parameter in this reaction, especially for the dissolution of CO_2 in the reaction medium. There are very few other reported catalysts for CO_2 hydrogenation to formic acid under aqueous alkaline conditions, these have been listed in the ESI† (Table S1). In the literature, NaHCO₃ and Na₂CO₃ have been reported to increase the pH and hence improve the dissolution of CO_2 into the aqueous medium.^{41,45–47} However, NaHCO₃ alone, without any added CO_2 , produces formic acid under hydrogenation conditions (*vide infra* Fig. 2). Hence, it is very important to choose the most appropriate base.

Plausible mechanisms for formation of formic acid from CO_2 hydrogenation has been widely investigated.⁴⁸⁻⁵² H₂ is a non-polar molecule, therefore is not very soluble in water as it does not readily form hydrogen bonds with water. It has been widely reported that Pd readily adsorbs and dissociates H₂ gaseous molecules.^{53–55} In fact, Wang et al. found that H₂ chemisorption increased when Pd was supported on Mo₂C in comparison with Pd supported on Al2O3 due to additional chemisorption onto the Mo₂C support⁵⁶ likely due to the strong electronic interaction between the β -Mo and H₂ (-0.67 eV).⁵⁷ Wang et al. studied palladium nanoparticles supported on nitrogen-doped mesoporous carbon in aqueous conditions.⁴⁹ Here they report the metallic palladium nanoclusters help the dissociative adsorption of H from H2. The resulting Pd-H bond activates the adsorbed bicarbonate by inserting the adsorbed H into the C-OH group of the bicarbonate, replacing the -OH which then binds with the remaining adsorbed H to form water. These findings are within good agreement with He et al.⁵⁰ and also Mori and co-workers work⁵¹ all within aqueous conditions. Using systematic calculations, both of these papers go on to explain that the reduction of HCO₃⁻ through the attack of the adsorbed H atom to the C atom of HCO₃⁻ leading to formate formation is much more energetically favoured rather than hydrogenation at its O atoms leading to carboxyl formation.⁴⁸ He and Mori's kinetic investigations demonstrate that the attack of the adsorbed H onto the HCO₃⁻ species is the rate determining step rather than the dissociation of H₂.

Here, we report palladium nanoparticles supported on molybdenum carbide, and a synthesis method to produce a more active Pd/Mo₂C catalyst for the liquid phase hydrogenation of CO₂ to formate, as the reaction solution is in basic conditions. The catalytic activities of four Pd/Mo₂C monometallic catalysts were studied and compared at mild reaction conditions (100 °C, 2 MPa (CO₂:H₂ 1:1)), with the most active catalyst producing a significant amount of formate salt with a turnover number (TON) of 109. Palladium is known as an active metal for H₂ activation as well as CO₂ activation,⁵⁸ Chen *et al.* reported Pd/Mo₂C as the most active catalyst for CO₂ hydrogenation to methanol *albeit* achieving a TON of 4.25, in an environmentally benign solvent (1,4-dioxane), at higher temperatures (135 °C) and pressure (40 bar).³⁵ Finally, a bimetallic RuPd/Mo₂C catalyst was also developed and tested for the liquid phase hydrogenation of CO₂. All monometallic and bimetallic catalysts were tested for their stability and the results were rationalised using a combination of high-resolution transmission electron microscopy (>HR-TEM), field emission gun scanning electron microscopy (FEG-SEM) and density functional theory (DFT) calculations.

Materials and methods

Mo₂C

Two different Mo₂C materials have been used in this work. Commercial β -Mo₂C was purchased from Alfa Aesar (99.5% purity) and was used without any further modification. β -Mo₂C was also synthesized *via* the procedure reported by Volpe *et al.*⁵⁹ Briefly, β -Mo₂C was prepared by the direct carburisation of MoO₃ (0.5 g, B.D.H.) using a flow of 12 ml min⁻¹ of 20 vol% CH₄ in H₂ (BOC, 99.98%) at 800 °C for 2 hours with a heating rate of 6 °C min⁻¹ till 350 °C followed by a 1 °C min⁻¹ heating rate till 800 °C. The resultant material was characterised by powder XRD (ESI,† Fig. S1). Thus prepared β -Mo₂C was used as support for the synthesis of Pd/Mo₂C catalyst and was used in the hydrogenation reaction without any further modification.

M/Mo₂C synthesis

Two different synthesis methods were used for the preparation of monometallic Pd nanoparticles supported on Mo₂C (commercial and synthesised) *i.e.* modified wet-impregnation method^{60,61} and sol-immobilisation method.⁶² Bimetallic RuPd/Mo₂C was prepared using a modified wet-impregnation method. A brief description of these methods is given below.

Modified wet-impregnation (MIm) method:

In a typical synthesis, of 2 g of 1 wt% M/Mo₂C catalyst, the requisite amount of the aqueous metal precursor (metal chloride) solution (equivalent to 0.02 g of metal; in the case of bimetallic catalyst, the two metals were taken in a equimolar ratio) was added to 16 ml of deionized water in a 50 ml glass round bottom flask with vigorous stirring. To this precursor solution, 1.98 g of Mo₂C was added slowly and steadily with constant stirring at 25 °C. After the completion of the addition of Mo_2C , the temperature of the stirring slurry was raised to 60 °C and stirred for 30 minutes. Finally, the temperature was raised to 95 °C and left overnight (16 hours) for the complete evaporation of water. After 16 h, the remaining dry, dark-grey solid was ground thoroughly and this dried material was reduced under 5 vol% H_2 in Ar at 400 °C for 4 hours with a heating rate of 10 $^{\circ}$ C min⁻¹. The reduced catalyst was used in the liquid phase reduction of CO₂ without any further modification unless specified otherwise.

Sol-immobilization (SIm) method

For the synthesis of 1 g of 1 wt% Pd/Mo_2C catalyst, an aqueous solution of $PdCl_2$ (Sigma Aldrich) was prepared. Polyvinyl

alcohol (PVA) (1 wt% aqueous solution, Aldrich, $M_W = 10000$, 80% hydrolyzed) and an aqueous solution of NaBH₄ (0.1 M) were also freshly prepared. To the requisite amount of an aqueous solution of PdCl₂, the required amount of a freshly prepared PVA solution (1 wt%) (PVA/(Au + Pd) (w/w) = 1.3) was added. A freshly prepared solution of NaBH₄ (0.1 M, NaBH₄/ (Pd) (mol/mol) = 5) was then added to form a dark-brown metallic sol. After 30 min of sol-generation, the colloid was immobilized by adding the support material (Mo₂C (0.99 g) Alfa Aesar). 5 drops of concentrated H₂SO₄ was added under vigorous stirring. After 2 h, the slurry was filtered, and the catalyst was washed thoroughly with 2 L of distilled water (until the mother liquor was neutral) and then dried at 120 °C overnight under static air in an oven.

All the above-mentioned catalysts were used in the hydrogenation reaction without any further modification or activation.

Catalyst characterisation

XRD

The bulk crystalline structures were characterised using X-ray diffraction. Conventional powder X-ray diffraction (PXRD) analysis of the materials was performed on a (θ – θ) PANalytical X'pert Pro powder diffractometer with a Ni filtered CuK α radiation source operating at 40 keV and 40 mA. Patterns were recorded over the 2θ angular range 10–80° using a step size of 0.016°.

XPS

X-ray photoelectron spectroscopy (XPS) was performed on a Thermo Fisher Scientific K-alpha+ spectrometer. Samples were analysed using a micro-focused monochromatic Al X-ray source (72 W) over an elliptical area of approximately 400 μ m. Data were recorded at pass energies of 150 eV for survey scans and 40 eV for high resolution scan with 1 eV and 0.1 eV step sizes respectively. Charge neutralisation of the sample was achieved using a combination of both low energy electrons and argon ions. Data analysis was performed in CasaXPS using a Shirley type background and Scofield cross sections, with an energy dependence of -0.6.

Microscopic studies

Transmission electron microscopy (TEM) were performed on a JEOL JEM-2100 operating at 200 kV. Energy dispersive X-ray analysis (EDX) was done using an Oxford Instruments X-MaxN 80 detector and the data analysed using the Aztec software. Samples were prepared by dispersion in ethanol by sonication and deposited on 300 mesh copper grids coated with holey carbon film. Scanning electron microscopy was performed on a Tescan Maia3 field emission gun scanning electron microscope (FEG-SEM) fitted with an Oxford Instruments X-MaxN 80 energy dispersive X-ray detector (EDX). Images were acquired using the secondary electron and backscattered electron detectors. Samples were prepared by dispersion in ethanol by sonication and deposited on 300 mesh copper grids coated with holey carbon film.

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Catalytic CO₂ hydrogenation reaction and products analyses

The hydrogenation of CO₂ to formate was carried out in a highpressure stainless steel Parr autoclave (50 ml) reactor fitted with an overhead stirrer. In a typical run, 150 mg of the catalyst was charged into a Teflon liner containing 15 ml of 1 M aqueous NaOH (Sigma Aldrich) solution. Then the Teflon liner was placed inside the autoclave reactor before the reactor was closed airtight. The reactor with its contents was first purged with N_2 (3 times) and then with CO_2 (3 times) to remove traces of air or oxygen from the system and then finally charged with CO_2 (10 bar) and H_2 (10 bar) at 25 °C. Then the reactor was heated to the reaction temperature (100 °C) while stirring at 800 rpm, when the reaction temperature is stabilised, the reaction pressure reaches approximately 26 bar. After 19 h of the reaction time the reactor was cooled to <10 °C using an ice bath, the gas-phase was collected in a gas-bag and the liquid sample was collected and the solid catalyst was removed via centrifugation followed by filtration using a syringe filter fitted with a 45 µl filter tip.

The gas phase products were analysed by gas chromatography (Varian 450-GC with a CPSil5 column 50 m \times 0.32 mm \times 5 µl, fitted with FID), while the liquid products were analysed by high performance liquid chromatography (HPLC) (Agilent 1260 infinity), injected into an Agilent Metacarb 67H column fitted with a refractive index detector. The method ran at 25 °C with a flow rate of 0.25 ml min⁻¹ with dilute sulfuric acid (0.1 vol%) as the mobile phase. The identity of the products (HCOOH) were confirmed using ¹H-NMR. A series of known standard solutions of formic acid were prepared to generate a calibration curve and response factors, which were used for quantitative analyses of the reaction mixtures.

When stability tests were conducted; 4 identical reactions were run using the procedure above, the solvent was removed by filtration and the catalyst was washed twice with H_2O and once again with acetone, and left to dry in a vacuum oven. The remaining catalyst was reused using the same reaction procedure as above, washed and dried again to continue the study. The turn over number (TON) was determined as the number of moles of sodium formate produced per mole of metal, calculated using microwave plasma atomic emission spectroscopy (MP-AES) (Agilent 4100), present in the catalyst used. Catalyst was digested in aqua regia first (HCl/HNO₃ = 3 : 1) until all solid was dissolved, before being diluted with water and manually submitted into the instrument.

Density functional theory calculations

We have modelled the orthorhombic phase of Mo₂C (a = 4.732, b = 6.037, $c = 5.204^{63}$) and its low Miller-index surfaces. The (001), (010), (011) and (101) surface slabs contained, respectively, 4, 6, 8 and 6 layers of Mo₂C units, and were separated by a vacuum region of at least 12 Å along the normal to avoid spurious interactions. We performed all geometry optimisations with the VASP 5.3 code,^{64,65} using the PBE functional,⁶⁶ spin polarisation and the projector augmented wave method,⁶⁷ treating explicitly the 5s and 4d electrons of Pd, Ru and Mo, and the 2s and 2p of C.

We adopted a plane wave cut off of 400 eV and scaled the bulk converged 5 \times 5 \times 5 Monkhorst–Pack⁶⁸ grid to 5 \times 5 \times 1 for surfaces.⁶⁹ Optimisations stopped when forces acting on ions were less than 2 \times 10⁻² eV Å⁻¹.

Results and discussion

Monometallic Au, Pd and Ru nanoparticles supported on Mo₂C were prepared *via* the modified wet-impregnation (MIm) method and tested for the CO₂ hydrogenation reaction in liquid phase using an aqueous NaHCO₃ solution as solvent. Results presented in Fig. 1 show that Pd/Mo₂C unsurprisingly is the most active catalyst for CO₂ hydrogenation, followed by Ru/Mo₂C and Au/Mo₂C. Based on this result, Pd/Mo₂C was used as the catalyst for further tests. It should be noted that Mo₂C alone managed to produce formate, with no supported metal, confirming that although only a small amount of product was formed (0.0049 mmol), the support material holds catalytic activity. Therefore, when acting as a support for metal nanoparticles, it also acts as a co-catalyst for the reaction.

As discussed in the Introduction section, for CO₂ hydrogenation reactions, an alkaline solution is vital to improve the solubility and reactivity of CO₂ and thereby increase the yield of desired product(s). In order to choose the most appropriate base, the hydrogenation reaction was carried out using different bases (NaHCO₃, Na₂CO₃ and NaOH) with and without CO₂, using 1% Pd/Mo₂C catalyst at 100 °C for 19 h. Many previous reports in the literature have used NaHCO₃ for CO₂ hydrogenation reaction;^{41,45–47} while HCO₃⁻ from the sodium bicarbonate is able to form formate in the absence of CO₂, it has been reported that the presence of CO₂ increases the formate productivity.⁷⁰ However, under our reaction conditions more formate was formed from an aqueous NaHCO₃ solution without CO₂ (1.91 mmol) compared to the reaction with CO₂ (1.18 mmol); a decrease of approximately 38% (Fig. 2).



Fig. 1 Screening of different monometallic nanoparticles supported on Mo₂C for CO₂ hydrogenation reaction. Reaction conditions: 1 wt% M/Mo₂C: 200 mg (Pd: 0.018 mmol; Ru: 0.019 mmol; Au: 0.01 mmol); stirring speed: 800 rpm; 1 M aqueous NaHCO₃ solution: 20 ml; pCO_2 : 10 bar (at 25 °C); pH_2 : 10 bar (at 25 °C); reaction temperature: 100 °C; reaction time: 24 h.

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Fig. 2 Production of formate using different bases (NaHCO₃, Na₂CO₃ and NaOH) with and without CO₂. Reaction conditions: 1% Pd/Mo₂C: 150 mg (Pd: 0.014 mmol); 1 M aqueous base: 15 ml; pCO_2 : 10 bar (25 °C); pH_2 : 10 bar (25 °C) for reaction with CO₂; pN_2 10 bar (25 °C) pH_2 : 10 bar (25 °C) only for reaction without CO₂; reaction temperature: 100 °C; reaction time: 19 h.

This effect may be due to the dissolution of CO_2 affecting the equilibrium of the carbonate species, reducing the amount of HCO_3^- available in solution for further hydrogenation. Na₂CO₃ again produces formate in the absence of CO_2 , but more formate was formed in the presence of CO_2 (0.75 mmol) than in the absence of CO_2 (0.52 mmol). When NaOH was used, formate was formed only in the presence of CO_2 (0.82 mmol). When CO_2 was dissolved in 1 M NaOH aqueous reaction mixture, the pH of the solution started decreasing and in 10 min, it stabilized at 8.0 until the end of the reaction. It is well established that at pH 8 the dissolved CO_2 forms HCO_3^- species, hence it is considered as the intermediate in the formation of formate.⁴⁴ During the course of the reaction there is always an equilibrium between CO_2 , HCO_3^- and $HCOO^-$. By choosing NaOH as the base we are confident that the C in $HCOO^-$ comes from CO_2 only.

The catalytic properties of any supported metal catalyst depend on its structural properties, such as particle size and morphology for monometallic catalysts and additional composition and nanostructure for bimetallic catalysts.⁷¹⁻⁷⁵ We have developed a number of synthesis strategies to control these structural properties of monometallic and bimetallic supported Pd catalysts. In an effort to tune the structural properties of the Pd/Mo₂C catalyst, four different Pd/Mo₂C structures were synthesised using modified wet-impregnation and sol-immobilisation methods.⁶⁰⁻⁶² 1% Pd/Mo₂C and 5% Pd/Mo₂C were prepared using an aqueous solution of Pd precursor, where PdCl₂ is dissolved in 0.58 M HCl solution; for the purpose of simplicity these will be labelled 1% Pd/ Mo₂C-MIm (0.58 M) and 5% Pd/Mo₂C-MIm, respectively, for the rest of this paper. Another 1% Pd/Mo2C was prepared using another precursor solution where PdCl₂ was dissolved in a 2 M HCl solution, we will label this 1% Pd/Mo₂C-MIm (2 M). We have reported previously that addition of excess of Cl⁻ ions (via the addition of either HCl or NaCl) during the wet-impregnation

procedure controls the particle size and morphology of supported AuPd catalysts.⁶⁰ Recently Li *et al.* reported the beneficial effect of the addition of excess chloride during the preparation of bimetallic PdRe catalyst for glycerol hydrogenolysis.⁷⁶ The fourth catalyst was synthesised using sol-immobilization, which will be labelled 1% Pd/Mo₂C-SIm. To confirm Mo₂C was an appropriate support material for this reaction, Pd supported on CeO₂ (1 wt%) was also tested and found that it was almost 20× less active than 1% Pd/Mo₂C, forming only 0.063 mmol of formate (ESI,† Table S2).

All the four Pd catalysts were tested for CO₂ hydrogenation and the formate yields are presented in the ESI⁺ Table S2. As expected the 1% Pd/Mo₂C-MIm (2 M), prepared from the precursor in 2 M HCl was found to be much more active (1.53 mmol) in comparison to the 1% Pd/Mo₂C-MIm (0.58 M) prepared from 0.58 M HCl (1.09 mmol). However, 5% Pd/Mo₂C-MIm catalyst gave only 2.14 mmol of formate under the same reaction conditions. For this catalyst, in spite of a 5-fold increase in Pd content compared to 1% Pd/Mo₂C-MIm (0.58 M) catalyst, the increase in formate yield is less than 2-fold. To normalise the formate productivity with Pd loading, the amounts of Pd present in all these catalysts were determined by MP-AES (ESI,† Table S2), Pd surface content analysis was attempted via CO chemisorption, although this proved unsuccessful, possibly due to the low weight loading and low surface area of the large nanoparticles. Actual Pd contents of all the wet-impregnation catalysts were found to be closer to the nominal loading (calculated from the amount of Pd precursor added during the catalyst preparation). The actual loading of the sol-immobilisation catalyst was found to be less than the expected nominal loading, possibly due to inefficient immobilisation of Pd nanoparticles onto the support and their subsequent loss during the washing of 1% Pd/Mo2C-SIm catalyst. The TON of all the catalysts, calculated based on the actual amount of Pd (Fig. 3), indicate that 1% Pd/Mo₂C-MIm (2 M) catalyst prepared from a 2 M HCl solution is



Fig. 3 Comparison of TON for different Pd/Mo₂C catalysts for the hydrogenation of CO₂ to produce formate. TON is the mol of formate produced per mol of Pd calculated from MP-AES. Reaction condition: Pd/Mo₂C: 150 mg (1 wt% = 0.014 mmol Pd, 5 wt% = 0.7 mmol Pd); 1 M aqueous NaOH: 15 ml; pCO_2 : 10 bar (at 25 °C); pH_2 : 10 bar (at 25 °C); reaction temperature: 100 °C; reaction time: 19 h.

the most active catalyst exhibiting a TON of 109. Chloride ions have long been considered a poison for noble metal catalysts and chloride precursors have therefore been avoided in the preparation of supported noble metal catalysts. Chen et al., who have previously reported M/Mo₂C for CO₂ hydrogenation, have avoided the use of PdCl₂ precursor as they report that the chlorine ions poison the catalyst surface and consequently reduce their catalytic activities,³⁴ which is not the first time Cl⁻ ions has considered as a poison.^{77,78} However, contradicting reports found Cl⁻ ions aid the dispersion of metal nanoparticles on the support surface.⁶⁰ Our current results clearly suggest that the addition of an excess of Cl⁻ ions, during the preparation of 1% Pd/Mo₂C-MIm (2 M) catalyst, is beneficial and 1% Pd/Mo₂C-MIm (2 M) was thus selected for the continuing studies in this paper. We have studied the effect of adding excess of chloride ions during the preparation of Pd based bimetallic catalysts by wet impregnation method in detail.55,56

Time on line data (Fig. 4a) for 1% Pd/Mo₂C-MIm (2 M) catalyst, show that the formation of formate increases steadily with time until 19 h (1.53 mmol of formate) after which time the productivity starts to plateau (1.58 mmol after 24 h). The hydrogenation of CO₂ was also tested at different temperatures (75, 100, 125 and 150 °C) and the formate productivity increases (Fig. 4b) with increasing reaction temperature, which is in line with many reported trends.^{79,80} At 75 °C 0.81 mmol of formate was produced and increasing the temperature to 125 °C, we see a remarkable increase in formate productivity, achieving 4.30 mmol of formate, with a TON of 307. At 150 °C formate production increases again to 5.26 mmol. The initial activity (turn over frequency) of 1% Pd/Mo₂C-MIm (2 M) for the CO₂ hydrogenation at 100 °C is ~14.1 h⁻¹ during the first initial 2 hours (calculated from ESI,† Fig. S5). It then decreases steadily in an approximately exponential way until it reaches a final activity of 4.6 h^{-1} after 19 hours. For a 125 °C reaction we see a TOF of approximately 42.5 h^{-1} during the first 2 hours

(Fig. S5, ESI[†]), and after 19 hours this drops to 16.2 h⁻¹. This result is comparable to Mori *et al.*'s Pd/TiO₂ and PdAg/TiO₂ catalysts, achieving TOFs of 12 and 31 h⁻¹ respectively,⁵¹ also their Ru/LDH catalyst, with TOF of 29. However, this catalyst indeed has a lot of competition and achieves TOFs less than Song *et al.*'s Pd/Chitin (TOF = 257 h⁻¹)⁴² and Maru *et al.*'s Pd/g-C₃N₄ (TOF = 660 h⁻¹)⁸¹ both achieving results at higher pressure but lower temperatures. A comparison of recent CO₂ hydrogenation catalysts, their TOFs and reaction conditions can be found in the ESI[†] (Table S1).

The pH of the reaction medium is an important parameter for the dissolution, activation and hence the hydrogenation of CO₂. As mentioned previously, upon CO₂ dissolution the pH of 1 M NaOH solution (pH = 14) dropped to pH 8 within 10 minutes as bicarbonate species formed in solution. Hence the hydrogenation of CO₂ to formate can be represented as $HCO_3^- + H_2 \rightarrow HCO_2^- +$ H₂O. Reducing the concentration of NaOH by half (0.5 M), halved the amount of HCOOH formed and the same effect occurred with 0.1 and 0.05 M NaOH solutions (Fig. 5); when no base is added to the reaction solution, the formate production is negligible, resulting in direct correlation between NaOH concentration and formate production.

After optimizing the reaction conditions, we studied the heterogeneous nature of the 1% Pd/Mo₂C-MIm (2 M) catalyst using the hot filtration method. The catalyst was removed by filtration after 4 h of the reaction (0.6 mmol of formate) and the reaction was continued without any catalyst. There was no increase in the formate yield even after 20 h of further reaction without catalyst (Fig. 6) which proves the heterogeneous nature of the 1% Pd/Mo₂C-MIm (2 M) catalyst. We further tested the stability and reusability of the Pd/Mo₂C catalyst by recycling the catalyst two times for CO₂ hydrogenation reactions (Fig. 7a) under identical reaction conditions. The 1% Pd/Mo₂C-MIm (2 M) catalyst displayed a large drop in the catalytic activity from 1.54 mmol of formate for the fresh catalyst to the



Fig. 4 (a) Time on line evolution of formate over 1% Pd/Mo₂C catalyst. Reaction condition: 1% Pd/Mo₂C-MIm (2 M): 150 mg (0.014 mmol Pd); 1 M aqueous NaOH: 15 ml; pCO_2 : 10 bar (at 25 °C); pH_2 : 10 bar (at 25 °C); reaction temperature: 100 °C. (b) Effect of temperature on the production of formate. Reaction condition: 1% Pd/Mo₂C-MIm (2 M): 150 mg (0.014 mmol Pd); 1 M aqueous NaOH: 15 ml; pCO_2 : 10 bar (at 25 °C); pH_2 : 10 bar (at 25 °C); reaction time: 19 h.



Fig. 5 Effect of NaOH concentration on the production of formate. Reaction conditions: 1% Pd/Mo₂C-MIm (2 M): 150 mg (0.014 mmol Pd); aqueous alkaline solution of various NaOH concentration: 15 ml; *p*CO₂: 10 bar (at 25 °C); *p*H₂: 10 bar (at 25 °C); temperature: 100 °C; reaction time: 19 h.



Fig. 6 Catalyst filtration study to show that leaching of Pd is not responsible for the catalytic activity. 1% Pd/Mo₂C catalyst was filtered after 4 h of the reaction and the reaction was continued with the filtrate until the overall reaction time reaches 24 h (circle). A time-on-line profile of the hydrogenation of CO₂ in the presence of the catalyst is also given for comparison (square). Reaction conditions: 1% Pd/Mo₂C-MIm (2 M): 150 mg (0.014 mmol Pd); 1 M NaOH: 15 ml; pCO₂: 10 bar (at 25 °C); pH₂: 10 bar (at 25 °C); reaction temperature: 100 °C.

0.28 mmol after the 2nd reuse; an 82% drop in the catalytic activity (Fig. 7a).

Commercial Mo₂C was found to have very low surface area $(<1 \text{ m}^2 \text{ g}^{-1})$ as BET and CO chemisorption of Pd/Mo₂C did not give any reasonable data. It has been reported that synthesised Mo₂C can have a much higher surface area of 151 m² g⁻¹.³⁴ Therefore, increasing the support surface area would be expected to decrease the size of the supported metal nanoparticles and thus improve the catalyst stability. In an effort to improve the stability of the Pd/Mo₂C catalyst, a fresh batch of β -Mo₂C was synthesized in the lab. These materials were found to

have a slightly improved surface area of approximately 13 m² g⁻¹ compared to commercial source and was used as the support for the synthesis of 1% Pd/Mo₂C catalyst. For the purpose of simplicity, this catalyst will be labelled 1% Pd/Mo₂C-Sy. Another approach to improve catalytic stability was to synthesize a Pd-based bimetallic catalyst. It is well known that the addition of a second metal to Pd can dramatically increase the stability of supported Pd catalysts.^{33,82–84} Recently, we reported supported bimetallic RuPd nanoparticles for the hydrodeoxygenation of levulinic acid to y-valerolactone with enhanced stability.61,85 Two Pd catalysts were prepared to improve the stability of Pd/Mo₂C catalyst based on the two approaches mentioned above. 1% Pd/Mo₂C-Sy was prepared using homemade β-Mo₂C and a bimetallic 1% RuPd/Mo₂C (commercially sourced Mo₂C) and both these catalysts were tested for catalytic activity and stability. Lab-synthesised β -Mo₂C support neither improved the catalytic activity nor the stability of 1% Pd/Mo₂C (Fig. 7d). However, the bimetallic 1% RuPd/Mo₂C displayed a much better stability compared to the monometallic 1% Pd/Mo₂C-MIm (2 M) catalyst, we did not load the lab synthesised β-Mo₂C with RuPd bimetallic nanoparticles as the commercial and synthesised supports produced the similar results when loaded with Pd monometallic nanoparticles. For the monometallic catalyst, during the 2nd reuse, an 82% reduction in activity was observed, whereas for the bimetallic RuPd catalyst only a 30% drop in activity was observed (Fig. 7c). The total Pd contents of fresh and spent catalysts were analysed by MP-AES (ESI,[†] Table S3), which showed that there is no decrease in Pd content, suggesting there is no leaching of Pd which was further confirmed by the absence of Pd in the reaction mixture.

In an effort to understand the reason for this deactivation, the Pd metal particle size of fresh and used catalysts, all fresh and used catalysts were analysed by HR-TEM and FEG-SEM. When observing catalysts synthesised using a commercially sourced Mo₂C, HR-TEM image (Fig. 8a), showed large, highly crystalline Mo₂C particles, which implies that the support has a very small surface area and few defects on the surface for Pd nanoparticles to bind to, resulting in large Pd particles on the surface. Due to the density of the Mo₂C support, the HR-TEM images give us minimal information as the material was blocking any transmission through the bulk of the material. FEG-SEM provides us with more detail with respect to the size of the metal particles and the nature of the support surface. The LE-BSE detector was used for distinguishing the palladium from the molybdenum surface. The BSE is strongly related to atomic number, with heavier elements backscattering more electrons, thereby contributing to a brighter signal. Large Pd nanoparticles can be observed (Fig. 8b) with a size range of approximately 50-70 nm and a good dispersion. When analysing the elemental composition of the Mo₂C surface, EDX showed a Pd wt% of 0.5-0.8%, which is a good result for a 1 wt% loading. Characterising the used Pd/Mo₂C sample we see a less even particle distribution and large agglomerations of palladium particles with sizes of approx. 1 µm (Fig. 8c), phase separated agglomerations were also found isolated from the support (not shown). This implies that during reaction, due to a weak



Fig. 7 Stability studies of Pd/Mo₂C and RuPd/Mo₂C catalysts. All catalysts were reused 2×. (a) Formate formed with 1% Pd/Mo₂C-Mlm (2 M), (b) formate formed with Pd/Mo₂C-Mlm (0.58 M), (c) formate formed with 1% RuPd/Mo₂C, (d) TON of formate production from 1% Pd/Mo₂C-Mlm (2 M) with commercially sourced Mo₂C (patterned bar) and 1% Pd/Mo₂C-Sy with lab synthesised Mo₂C (solid bar). Reaction conditions (a – c + d (patterned bar)): M/Mo₂C: 150 mg (monometallic: 0.014 mmol Pd) (bimetallic 0.007 mmol (Pd,Ru)); 1 M NaOH: 15 ml; pCO_2 : 10 bar (at 25 °C); pH_2 : 10 bar (at 25 °C); reaction time: 19 h. Reaction conditions (d (solid bar)):10 ml stainless steel autoclave: M/Mo₂C: 40 mg; 1 M NaOH: 4 ml; pCO_2 : 10 bar (at 25 °C); reaction temperature: 100 °C; reaction time: 19 h.



Fig. 8 (a) HR-TEM images of fresh 1% Pd/Mo₂C-MIm (2 M). FEG-SEM images of (b) fresh 1% Pd/Mo₂C-MIm (2 M) LE-BSE (c) used 1% Pd/Mo₂C-MIm (2 M), Pd cluster LE-BSE.

metal-support interaction and surface decomposition, palladium nanoparticles sinter, and in turn, are lost from the support as heterogeneous particles, explaining the drop in catalytic activity. This did not show in the leaching studies as the particles are too large to be retained in the filtrate solution. Tests were run to confirm that these Pd particles were not the catalyst for formate production; PdCl₂ and Pd colloids were catalytically tested and compared to 1% Pd/Mo₂C-MIm (2 M), using the same palladium weight and reaction conditions. Formate productivities of 0.06 and 0.09 mmol were observed for PdCl₂ solution and Pd colloids respectively (ESI,† Fig. S2), compared to 0.42 mmol produced from 1% Pd/Mo₂C-MIm (2 M). This confirms that the metalsupport interactions are indeed causing an increase in catalytic activity of the palladium. Observing fresh bimetallic 1% RuPd/ Mo₂C under HR-TEM and FEG-SEM shows a larger range of metal nanoparticle sizes, ca.10-85 nm, with an even dispersion over the support (Fig. 9a). Analysing recycled 1% RuPd/Mo₂C, it was found that the metal particles had increased in size (Fig. 9c), exceeding 100 nm. However, no metal particles, palladium or ruthenium, were found isolated from the support. This implies that, although there was sintering along the support surface during reaction, the addition of the ruthenium creates a stronger metal-support interaction and as a result, no metal particles are lost from the support, explaining the improved stability compared to the monometallic 1% Pd/Mo₂C.

We have used atomistic simulations based on DFT to shed light on the improved stability of the bimetallic nanoparticles. In order to have a model for the support, we have first calculated the surface energies, γ , of the low index (001), (010), (011) and



Fig. 9 (a) HR-TEM image of fresh 1% RuPd/Mo₂C; FEG-SEM images: (b) fresh 1% RuPd/Mo₂C LE-BSE, (c) used 1% RuPd/Mo₂C LE-BSE.

(101) surfaces of orthorhombic $Mo_2C,$ shown in ESI,† Fig. S3, according to the formula:

$$\gamma = \frac{1}{2A} \left(E_{\text{slab}} - N E_{\text{Mo}_2 \text{C}} \right), \tag{1}$$

where E_{slab} is the energy of the slab with surface area *A* and containing *N* Mo₂C units, $E_{\text{Mo}_2\text{C}}$ is the energy per formula unit of bulk Mo₂C, and the factor 2 takes into account the two symmetric terminations of the surface slabs. Note that we have not considered the polar (100), (110) and (111) surfaces as they would undergo a significant reconstruction to eliminate the electric dipole moment⁸⁶ and are generally less stable.²⁹

We report in ESI,[†] Table S4 the surface energies of the different surfaces. Our calculations indicate that the (101) is the most stable face of Mo_2C , in agreement with the previous literature,⁸⁷ followed by the (010), (011) and (001). In view of these results, we have used the (101) and (010) surfaces as substrate models for the adsorption of single Pd and Ru atoms.

Fig. 10 illustrates the adsorption sites obtained after relaxing a number of different initial positions. On the (101) surface, aand b are sixfold hollow sites. Here, the metal adatoms coordinate to one C and four Mo atoms of the top layer, in addition to one Mo atom of the second layer. In c, which is a fourfold hollow site, the adatoms coordinate only to topmost Mo atoms. Site d is threefold, and the adatoms form a bridging bond with two Mo besides bonding a third C atom. Considering the (010) surface, site e is above a C atom of the topmost layer, while f is a sixfold hollow site. Finally, g and h are different bridging sites.



Fig. 10 Adsorption sites on the (101) (a) and (010) (b) surfaces of Mo₂C. Top and bottom panels show top and side views, respectively. Only the atoms of the topmost Mo₂C layers are depicted in the top views. Colour code: Mo – large purple spheres, C – small brown spheres.

Surface	Site	ΔE (eV per atom)		
		Pd	Ru	
(101)	а	-4.25	-6.52	
	b	-4.03	-6.17	
	с	-4.34	-6.05	
	d	-3.06	-4.77	
(010)	e	-3.01	-4.35	
	f	-3.05	-4.54	
	g	-4.19	-6.40	
	ĥ	-3.71	-5.36	

We list in Table 1 the adsorption energies, ΔE , of Pd and Ru atoms at the sites described above, which have been calculated according to the formula:

$$\Delta E = \frac{1}{2} \left(E_{2\mathrm{M/slab}} - E_{\mathrm{slab}} - 2E_{\mathrm{M}} \right), \tag{2}$$

where $E_{2M/\text{slab}}$ and E_{slab} are the energies of the slab with and without the metal adatoms M, E_M is the energy of M in the gas phase and the factor 2 takes into account the fact that adsorptions occur on symmetric terminations. For all sites of both surfaces, ΔE is more negative for Ru compared to Pd, with a difference which ranges from 1.3 to 2.3 eV. These results represent evidence that Ru atoms have significantly better affinities than Pd to the Mo₂C support adopted in the experiment. We propose that the Ru atoms act like a link between Pd and the Mo₂C support and thereby increase the stability and reusability of the bimetallic RuPd catalyst.

Pd/Mo₂C and RuPd/Mo₂C catalysts were characterised by XPS. From previous XPS studies of Mo₂C catalysts, Moon *et al.* concluded that the deactivation of a Mo₂C catalyst is caused by the transformation of Mo₂C (Mo^{2+}) to MoO₃ (Mo^{6+}) on the

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Fig. 11 Pd 3d XPS spectra of 1% Pd/Mo₂C-MIm (2 M) (a) fresh sample (b) $3 \times$ used sample, and 1% RuPd/Mo₂C (c) fresh sample (d) $3 \times$ used sample; displaying Pd(0) (green) and Pd(II) (red). Ru 3d XPS spectra of 1% RuPd/Mo₂C (e) fresh sample (f) $3 \times$ used sample, displaying RuO₂ (blue) species.

surface of the catalyst in the presence of H₂O during WGS reaction.⁸⁸ X-ray photoelectron spectroscopic data (Mo 3d) of 1% Pd/Mo₂C-MIm (2 M) (ESI,† Fig. S4) reveals that the support is predominantly oxidised with several oxidation states of molybdenum ranging from Mo^(δ +) to Mo⁽⁶⁺⁾ (1 < δ < 4). However, these oxidised species do not show much deviation between the fresh and used samples. In respect of the metal, palladium is observed as a well resolved doublet, with the Pd3d_{5/2} centred at 335.7 eV, and exhibiting shake-up structure

to higher binding energies indicative of metallic Pd^{56} and $Pd3d_{5/2}$ centred at 337 eV indicative of Pd(II) (Fig. 11). Post- reaction we only see Pd(0) present in the sample possibly accounting to the hydrogenation conditions of the reaction process.

In the XPS of bimetallic RuPd samples, Ru peak (Fig. 11) is observed as an asymmetric doublet with $Ru3d_{5/2}$ centred at *ca*. 281 eV indicative of RuO_2 .⁸⁹ The amount of Pd(n) increases postreaction (Fig. 11d) which may be a result of some charge transfer from Pd to Ru. This is in line with Zhang *et al.*, who reported that

Sample	Pd(0)		Pd(II)		Ru(rv)	
	3d _{5/2} , eV	3d _{3/2} , eV	3d _{5/2} , eV	3d _{3/2} , eV	3d _{5/2} , eV	3d _{3/2} , eV
1% Pd/Mo ₂ C-MIm (2 M) fresh	335.73	341.03	337.25	342.55	_	_
$1\% \text{ Pd/Mo}_2\text{C-MIm}(2 \text{ M}) \text{ used}$	335.84	341.14	_	_	_	_
1% RuPd/Mo ₂ C fresh	336.20	341.50	_	_	281.03	285.23
1% RuPd/Mo ₂ C used	336.11	341.41	337.06	342.36	280.71	284.91

for PdRu bimetallic samples the addition of Ru increases Pd(π) species.⁹⁰ Ru(π v) has a higher reduction potential (+1.12 E°/V) than Pd(π) (+0.95 E°/V) causing an electron transfer from Pd to Ru resulting in an increase in Pd(π) species, which could be one of the reasons for the deactivation of this bimetallic catalyst.

A positive shift (+0.5eV) of the Pd 3d binding energies is observed for the RuPd bimetallic catalyst compared to the monometallic Pd catalyst (Table 2), indicating that a change in the electronic properties of Pd is modified upon alloying with Ru.⁹⁰ Comparing the fresh and used RuPd bimetallic catalyst, shifts for the Pd 3d binding energies are within experimental error, however a negative shift (-0.3 eV) for the Ru 3d binding energy in the post reaction catalyst is observed, suggesting that there are still some alloy properties even after 3 uses, but it has decreased. It is this alloying affect that may be the reason of the bimetallic increased stability, as this metalmetal interaction is contributing to stronger metal-support interaction for Pd. Unfortunately, due to the similar atomic number of Pd, Ru and Mo it has proved very difficult to confirm this alloying through EDX.

The initial rate of formation of formate were calculated at different temperatures (75, 100 & 125 °C) for both monometallic Pd/Mo₂C and bimetallic RuPd/Mo₂C catalysts under standard conditions for the first 2 hours of the reaction (ESI,† Fig. S5 + S6). Using this rate data, the apparent activation energies for 1% Pd/Mo₂C and 1% RuPd/Mo₂C were calculated as +36 kJ mol⁻¹ and +43 kJ mol⁻¹ respectively. Luo *et al.*⁶¹ reported that alloying RuPd results in Ru diluting and isolating certain active sites of Pd, which may result in this increase in activation energy as well as creating stabilising effects. In comparison with the literature, these results are lower than Chen et al.'s work, reporting CO₂ hydrogenation to methanol Fe and Cu supported on Mo₂C within an organic solvent, with activation energies of +96 and +105 kJ mol⁻¹ respectively.³⁴ In other catalytic systems known for hydrogenation of CO₂ and HCO₃⁻ salts, the activation energy values are larger than our Pd/Mo₂C catalyst, for example +54.3 kJ mol⁻¹ for Ru/LDH,⁴¹ +39 kJ mol⁻¹ with Pd/C catalyst.^{91,92} The activation energy is similar to certain homogeneous catalysts [RhCl(TPPMS)₃]⁹³ with +36 kJ mol⁻¹. But higher than others for example with K[RuCl(EDTA-H)]⁹⁴ it is +31 kJ mol⁻¹, and $[RhCl(TPPTS)_3]^{95}$ it is 25 kJ mol⁻¹.

Conclusions

Here we report 1% Pd/Mo₂C as an active catalyst for the liquid phase hydrogenation of CO₂ to formate under relatively mild reaction conditions. It was found that increasing the HCl

concentration from 0.58 M to 2 M for the preparation of PdCl₂ precursor solution for catalyst preparation increased the catalytic activity by more than 70%, producing 1.5 mmol of formate with a TON of 109 and activation energy of +36 kJ. Our study found that compared to NaHCO3 and Na2CO3, a 1 M NaOH basic solvent proved the most effective as it allowed 100% of formate formed to be derived from the CO₂. Stability studies and electron microscopy data found that 1% Pd/Mo2C has a poor reusability as Pd nanoparticles easily sinter and can be partially lost from the support. However, when introducing ruthenium in a bimetallic 1% RuPd/Mo₂C catalyst, although less active, greatly improves its reusability. Simulations based on DFT have found that the adsorption energies of Ru on the Mo₂C surfaces are significantly lower than Pd. As such, Ru atoms have a stronger affinity towards the Mo₂C support. XPS data suggests close interaction between Pd and Ru, thus, in the bimetallic catalyst the Ru atoms act like a link between the Pd and Mo₂C surface, resulting in an increased stability of the RuPd/Mo₂C catalyst.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

We thank the Cardiff University electron microscopy facility for the transmission (TEM) and scanning electron microscopy (SEM). XPS data collection was performed at the EPSRC National Facility for XPS ('HarwellXPS'), operated by Cardiff University and UCL, under contract No. PR16195. *Via* our membership of the UK's HEC Materials Chemistry Consortium, which is funded by EPSRC (EP/L000202), this work used the ARCHER UK National Supercomputing Service (http://www.archer.ac.uk). All data created during this research are openly available from the Cardiff University Research Portal at http://doi.org/10.6084/m9.figshare.9586913.

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