**Catalysis
Science & Technolog** Technology (SS)

PAPER

Cite this: Catal. Sci. Technol., 2020, 10, 8203

Received 9th June 2020, Accepted 15th October 2020

DOI: 10.1039/d0cy01163k

rsc.li/catalysis

Introduction

The direct synthesis of hydrogen peroxide (H_2O_2) from molecular hydrogen and oxygen offers an attractive alternative to the current means of production, the anthraquinone oxidation or in-direct process. The direct route to H_2O_2 has been carried out in a wide range of reaction media, including water. $1-9$ For many applications the most convenient solvent for the reaction would be water as it is readily available, nontoxic, non-flammable and completely miscible with H_2O_2 . Indeed a large number of industrial processes utilise aqueous $H₂O₂$, in particular for application as a bleaching agent in the pulp and textiles sector (accounting for nearly 50% of annual usage¹⁰) H₂O₂ diluted in water is favoured.¹¹ However, a major drawback of using H_2O as a solvent is the low solubility of reagent gases H₂ and O₂, (1.62 mg L⁻¹ and 40 mg L⁻¹ respectively, at room temperature)¹² which has been shown to limit H_2O_2 synthesis rates, compared to that observed when using alcohol/water co-solvent systems.^{1,2} To overcome these limitations solvents other than water have been investigated including supercritical $\mathrm{CO_2}^{\,13\mathrm{-}16}$ and halogenated solvents $^{17\mathrm{-}19}$ as well as short chain alcohols,^{9,20,21} with Paunovic et al.²² providing a comprehensive study on the role of the co-solvent

The direct synthesis of hydrogen peroxide using a combination of a hydrophobic solvent and water†

Adeeba Akram,^a [Gre](http://orcid.org/0000-0002-6571-5731)g Shaw,^a Richard J. Lewis, D^a Marco Piccinini,^a David J. Morgan, ¹ [Th](http://orcid.org/0000-0003-4089-2827)omas E. Davies,^a Simon J. Freakley,^b Jennifer K. Edwards, \mathbf{D}^a Jacob A. Moulijn^a and Graham J. Hutchings \mathbf{D}^{*a}

The direct synthesis of hydrogen peroxide (H_2O_2) has been studied using a solvent system comprising a hydrophobic alcohol (decan-1-ol) and water. It is demonstrated that, with the optimum combination of solvent and catalyst the contribution of H_2O_2 degradation pathways can be minimised to achieve industrially acceptable H_2O_2 concentrations under moderate conditions. This is achieved through the use of a catalyst that is retained by the organic component and the extraction of synthesised H₂O₂ into the aqueous phase, consequently limiting contact between the synthesised H_2O_2 , catalyst and reactant gases, resulting in an improved selectivity towards H₂O₂. Investigation of the reaction parameters provides an insight into the proposed solvent system, and optimised conditions to produce H_2O_2 from molecular H_2 and $O₂$ have been identified. Through this optimisation $H₂O₂$ concentrations up to 1.9 wt% have been achieved via sequential gas replacement experiments. **PAPER**
 (a) Check for undates
 (a) Check for undates
 EVALUATE CONSTRANT CO

in the direct synthesis reaction. It has been reported that the rate of H_2O_2 production in short chain alcohols is much higher than that in aqueous media due to higher gas solubility.^{20,23} The solubility of H_2 in short chain alcohols, in particular methanol, has been reported to be 4–5 times higher than in water whereas that of O_2 may increase up to eightfold.24,25 Furthermore, additional studies have reported that using a methanol co-solvent can lead to the suppression of H_2O_2 decomposition activity, likely due to the ability of methanol to act as a hydroxyl radical scavenger.²⁶

Despite this approach, the presence of subsequent reactions in the H_2O_2 direct synthesis process (Scheme 1) suggests that it will remain challenging to achieve high H_2O_2 concentrations, due to the thermodynamically favoured H_2O_2 degradation pathways (hydrogenation and decomposition). Typically, these unwanted reaction pathways have been inhibited through the use of a combination of sub-ambient reaction temperatures,²⁷ acidic promoters,^{28–30} halide

Scheme 1 Reaction pathways associated with the direct synthesis of $H₂O₂$ from $H₂$ and $O₂$.

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

Building, Park Place, Cardiff, CF10 3AT, UK. E-mail: hutch@cardiff.ac.uk

 b Department of Chemistry, University of Bath, Claverton Down, Bath, BA2 7AY, UK † Electronic supplementary information (ESI) available. See DOI: 10.1039/ d0cy01163k

stabilizing agents $31,32$ (with obvious drawbacks associated with reactor corrosion and the need for their down-stream removal) or complicated catalyst design, such as pretreatment of catalyst supports prior to metal immobilisation 33 or sequential heat treatment cycles. 34 While a growing number of catalysts have been reported that are capable of achieving high H_2O_2 selectivity,^{33–35} in the absence of stabilizing agents, the consecutive reaction pathways still prohibit the production of H_2O_2 concentrations on an industrial scale, with H_2O_2 concentrations between 0.6-1.8 wt% typically produced in initial stages of the industrial process, prior to concentration and shipping.³⁶

With the first order dependence of H_2O_2 synthesis on H_2 partial pressure,³⁷ increasing H_2 concentration at the catalyst surface through application of a short chain alcohol as solvent or increasing reaction pressure (which is not attractive for a commercial process) can inherently be related to a proportional increase in the rate of H_2O_2 production. A bi-phasic solvent system consisting of long chain alcohols and water could provide both high solubility of gases and the ability to extract H_2O_2 into the aqueous phase under moderate conditions. Ideally in this system, the catalyst and gases will be held in the organic phase where H_2O_2 will be produced and subsequently extracted in situ into the aqueous phase due to the higher solubility of H_2O_2 in water than the organic solvent. The use of a bi-phasic solvent system has the potential to lead to enhanced selectivity towards H_2O_2 , through minimising contact time between H_2O_2 , the catalyst and reactant gases, which in turn leads to a reduction in the subsequent H_2O_2 degradation reactions and, thus, an **Paper**
 Catalysis Science & Technology

valis reader corresponds and the need for their downloades are been received been represented to the production of the

common paper are expected to the production of the common

increase in H_2O_2 concentration. Indeed similar approaches have been recently been reported for the production of H_2O_2 *via* photocatalytic processes³⁸⁻⁴⁰ as well as in the biomass conversion literature, where bi-phasic solvent systems often lead to higher product yields than with mono-phasic aqueous systems due to the higher solubility of products in the organic phase. 41 Furthermore, the use of multi-phasic systems has been well established in processes that utilise homogeneous catalysts.^{42,43} In the case of H_2O_2 synthesis, when selecting a suitable candidate for the organic phase two key features are required, an ability to produce H_2O_2 in the solvent, and a low water solubility. In order to narrow down the search for a suitable organic solvent only primary alcohols were considered in this study. Water was selected as the co-solvent due to it being low cost, readily available and its full miscibility with H_2O_2 .

The choice of the catalyst support is also an important parameter to consider, with the ability of the catalyst to be retained in the hydrophobic solvent, during the course of the reaction, crucial to inhibiting H_2O_2 degradation. In this optimum scenario (Scheme 2a) the contact between catalyst and reagent gases would be maximized, while also isolating the formed H_2O_2 and inhibiting degradation pathways. On the other hand, the use of a catalyst that is retained primarily in the aqueous phase (Scheme 2b), or indeed the use of a single phase solvent system, such as water/ethanol⁴⁴ or water/methanol,^{35,45,46} which are both commonly studied in the literature, (Scheme 2c) would not limit contact between synthesised H_2O_2 , catalyst and reagent gases, leading to increased rates of H_2O_2 degradation. With the application of

Scheme 2 The effect of solvent and catalyst support on promoting H_2O_2 selectivity in (a) a non-miscible solvent system where the catalyst is retained in the organic component of the solvent, (b) a non-miscible solvent system where the catalyst is retained in the aqueous component of the solvent and (c) a completely miscible solvent system.

stabilizing agents, to inhibit catalytic degradation of H_2O_2 , typical in these miscible solvent systems.

This paper outlines the investigation of the direct synthesis of H_2O_2 based on the described hydrophobic alcohol and water solvent system. Reaction parameters have been varied, namely water concentration, catalyst support, solvent mass and reaction time to deduce the feasibility of the outlined solvent system.

Experimental

Preparation of AuPd supported catalysts by wet impregnation

AuPd bimetallic catalysts were prepared via a wet coimpregnation onto a range of supports; Darco G60 carbon (C), Aeroxide TiO₂ (Degussa, Aeroxide, P25), SiO₂ and CeO₂. The standard preparation of 2.5 wt% Au–2.5 wt% Pd supported catalyst is described as follows (all quantities stated are per 1 g of catalyst). PdCl₂ (0.0417 g, Sigma Aldrich) was added to HAuCl₄·3H₂O (2.04 mL, 12.25 mg mL $^{-1}$, Strem Chemicals). The solution was stirred and heated (80 °C) until the $PdCl₂$ dissolved completely to form a homogeneous solution. The appropriate support (0.95 g) was added to the solution and stirred until a paste was formed. The resultant material was dried in an oven (110 °C, 16 h) before being ground and calcined in static air (400 °C, 3 h, 20 °C min $^{-1}$).

Characterisation

The volumetric Karl Fischer method was performed using a TitroLine® 7500 KF volumetric Karl Fischer titrator where the titrating agent was accurately added through a piston burette. The standard procedure for each titration was as follows. The burette was filled with the titrating agent (Hydranal®- Composite 5). The working medium (Hydranal®- CompoSolver E) was added to the titration vessel, which was titrated to dryness with the titrating agent. The organic solvent sample was added (between 0.1–1.0 g depending on expected water concentration) and the sample was titrated with the titrating agent to determine the water concentration.

¹H NMR spectra were recorded on a Bruker Ultrashield 500 MHz spectrometer, using a $H₂O$ solvent suppression program. Filtered solvent (0.7 mL) was added to an NMR tube containing D_2O (0.1 mL).

Investigation of the bulk structure of the crystalline materials was carried out using a $(\theta-\theta)$ PANalytical X'pert Pro powder diffractometer using a Cu K_{α} radiation source, operating at 40 KeV and 40 mA. Standard analysis was carried out using a 40 min run with a back filled sample, between 2θ values of 10–80°. Phase identification was carried out using the International Centre for Diffraction Data (ICDD).

X-ray photoelectron spectroscopy (XPS) analyses were made on a Kratos Axis Ultra DLD spectrometer. Samples were mounted using double-sided adhesive tape and binding energies were referenced to the C (1s) binding energy of adventitious carbon contamination that was taken to be 284.7 eV. Monochromatic AlK_{α} radiation was used for all measurements; an analyser pass energy of 160 eV was used for survey scans while 40 eV was employed for detailed regional scans. The intensities of the Au (4f) and Pd (3d) features were used to derive the Au/Pd surface ratios.

To allow for quantification of total metal loading catalysts were digested via an aqua-regia assisted, microwave digestion method using a Milestone Connect Ethos UP microwave with an SK15 sample rotor. Digested samples were analysed using an Agilent 7900 ICP-MS equipped with I-AS auto-sampler. All samples were diluted by a factor of 10 using HPLC grade H_2O $(1\%$ HNO₃ and 0.5% HCl matrix). All calibrants were matrix matched and measured against a five-point calibration using certified reference materials purchased from Perkin Elmer and certified internal standards acquired from Agilent.

Direct synthesis of H_2O_2

Catalyst testing was performed using a Parr Instruments stainless steel autoclave which had a nominal volume of 100 mL and a maximum working pressure of 14 MPa. The autoclave was equipped with an overhead stirrer (0–2000 rpm) and provision for measurement of temperature and pressure. To test the direct synthesis of H_2O_2 , the autoclave was typically charged with catalyst (0.01 g) and saturated decan-1-ol $(0.39 \text{ g H}_2\text{O}$ and 8.11 g decan-1-ol). The charged autoclave was purged three times with 5% H_2/CO_2 (100 psi) and then filled with 5% H_2/CO_2 (420 psi) and 25% O_2/CO_2 (160 psi) to give a H_2 : O₂ ratio of 0.5 and a total working pressure of 580 psi, with no continual introduction of gas. The reaction mixture was allowed to stabilise at the desired temperature (25 °C) after which stirring commenced (1200 rpm) and experiments were carried out for 30 minutes, unless stated otherwise. H_2O_2 yield was determined by titrating all of the final solution (unless stated otherwise) with acidified $Ce(SO_4)_2$. The $Ce(SO_4)_2$ solutions were standardised against $(NH_4)_2Fe(SO_4)_2.6H_2O$ using ferroin as an indicator. These reaction conditions were systematically varied in this study. Catalysis Science & Technology

stabilizing agents, to inhibit catalytic degradation of H4O₂, incessure which as 0 V was uncertified to the control of the control of the sumplements are also that the control of the cont

> In this paper, results have been primarily obtained with saturated (with water) decan-1-ol, with the amount of hydrogen peroxide discussed in terms of the theoretical H_2O_2 concentration in water (wt%). The H_2O_2 concentration is expressed assuming facile separation of decan-1-ol and water can be achieved, with no H_2O_2 retained in the organic solvent.

Time-on-line analysis for the direct synthesis of H_2O_2

An identical procedure to that outlined above for the direct synthesis of H_2O_2 is followed for the desired reaction time. It should be noted that individual experiments are carried out and the reaction mixture is not sampled on-line.

Gas replacement experiments for the direct synthesis of H_2O_2

An identical procedure to that outlined above for the direct synthesis of H_2O_2 is followed for a reaction time of 0.5 h. After this, stirring is stopped and the reactant gas mixture is vented prior to replacement with the standard pressures of

5% H_2/CO_2 (420 psi) and 25% O_2/CO_2 (160 psi). The reaction is then stirred (1200 rpm) for a further 0.5 h.

Degradation of H_2O_2

Catalytic activity towards H_2O_2 degradation was determined in a similar manner to the direct synthesis activity of a catalyst. The autoclave was typically charged with catalyst (0.01 g) and reaction solution (0.39 g 4 wt% H_2O_2 and 8.11 g organic solvent). The charged autoclave was purged three times with 5% H_2/CO_2 (100 psi) and then filled with 5% H_2/CO_2 (420 psi). The reaction mixture was allowed to stabilise at the desired temperature (25 °C) after which stirring commenced (1200 rpm) and experiments were carried out for 30 minutes. H_2O_2 degraded was determined by titrating all of the final solution with acidified $Ce(SO₄)₂$. The $Ce(SO₄)₂$ solutions were standardised against $(NH_4)_2Fe(SO_4)_2·6H_2O$ using ferroin as an indicator. Catalytic activity towards H_2O_2 degradation is reported herein in terms of rate $(mol_{H_2O_2}$ kg_{cat}⁻¹ h⁻¹) to better allow for comparison to the literature and accounts for hydrogenation and decomposition pathways.

Note: within this work reactant gases have been diluted with $CO₂$ to ensure that at no time do mixtures of $H₂$ and $O₂$ enter the explosive region (4–94 mol%).

Results and discussion

Within this study we evaluate the catalytic performance of a range of previously studied supported AuPd catalysts prepared by a wet co-impregnation methodology, (5% AuPd/ $C³³$ 5% AuPd/TiO₂,⁶ 5% AuPd/CeO₂ (ref. 47) and 5% AuPd/ $SiO₂$ (ref. 27)) (actual metal loading, as determined by *aqua*regia assisted microwave digestion reported in Table S.1†) towards H_2O_2 synthesis, using a solvent system consisting of a hydrophobic alcohol and water. We aim to address H_2O_2 selectivity through minimizing contact between catalyst and the synthesised H_2O_2 , with the choice of solvent ensuring the former is preferentially retained in the hydrophobic alcohol while the latter extracted into the aqueous phase.

We have previously demonstrated that catalytic selectivity of the materials studied within this work correlates strongly with the surface charge of the support, with catalysts synthesised on supports such as C and $SiO₂$, which have low isoelectric points, offering the greatest rates of H_2O_2 synthesis.⁴⁸ Although other parameters such as nanoparticle dispersion,⁴⁹ the degree of alloying⁵⁰ and Pd oxidation state^{51,52} are clearly also strongly related to catalytic performance.

With this in mind we initially investigated the as-prepared catalysts via XRD (Fig. S.1†), with reflections observed at 38, 44, 66 and 78° corresponding to Au (111), (200) (220) and (311) planes in all samples, in addition to reflections at 65 and 79° corresponding to the Pd (220) and (311) planes observed in the $AuPd/SiO₂$ catalyst. This is indicative of the supported nanoparticles generally being large or poorly dispersed. This is perhaps unsurprising, with catalysts prepared via a wet-impregnation procedure well known to display a bimodal distribution of nanoparticle size.⁵³ Indeed

detailed STEM-XEDS analysis of oxide supported AuPd nanoparticles has previously revealed a distinct relationship between particle size and elemental composition, with larger nanoparticles found to be Au-rich, while smaller nanoparticles are Pd-rich.⁵⁴ In a similar manner we have demonstrated that analogously prepared carbon supported catalysts also adopt a size dependent nanoparticlecomposition. However, unlike with catalysts prepared on oxide supports, which adopt a Au-core Pd-shell morphology after exposure to an oxidative heat treatment, analogous catalysts prepared on carbon have been found to maintain the random alloy morphology observed prior to calcination.⁵⁵

 $We⁵⁵$ and others^{56,57} have extensively studied the synergistic effect achieved through the incorporation of Au into supported Pd catalysts with the reduction in contiguous Pd ensembles often attributed as the cause for the enhanced catalytic performance observed compared to monometallic analogues. Further studies have highlighted the role Au plays in electronically modifying Pd, with Au incorporation supressing O–O bond cleavage, inhibiting the formation of $H₂O$, as well as promoting $H₂O₂$ desorption through weakening the interaction between the as-synthesised H_2O_2 and the catalyst surface.⁵⁸⁻⁶⁰ In particular the formation of Au-core Pd-shell nanoparticle morphology, commonly adopted upon the calcination of AuPd nanoparticles supported on oxide supports, has been widely reported to result in enhanced catalytic performance. Analysis of the various AuPd catalyst (Table S.2 and Fig. S.2†) via XPS reveals that the observed Pd : Au surface atomic ratios for the oxide supported catalysts are markedly higher than the nominal compositions, with these observations consistent with a Aucore Pd-shell nanoparticle morphology. By comparison, the Au : Pd surface ratio for the carbon supported catalyst is more in keeping with the formation of a homogeneous alloy, again with this aligning well with our previous studies.^{54,55} **Paper** Catalysis Science & Technology

So Hoto (120 psi) and 25% 0.070, (100 psi). The reaction detailed site procedure as procedure as procedure and distinct relationships are proported under the site of the site of the

> The as-prepared catalysts were investigated for their affinity to be retained in a hydrophobic solvent. Samples were added to a solvent with two distinct phases, decan-1-ol and water. After vigorous shaking it was observed (Fig. 1) that only the 5% AuPd/C remained in the alcohol phase, whereas the analogous $TiO₂$, $CeO₂$ and $SiO₂$ supported catalysts were predominately present in the aqueous phase. These results are not unexpected as $TiO₂$, CeO₂ and $SiO₂$ are more polar than C.

> We have previously investigated the ability of these catalysts to synthesise H_2O_2 under conditions optimised to promote H_2O_2 stability, namely the use of a methanol-water solvent system and sub-ambient temperature¹ (see Table S.3[†] for H_2O_2 synthesis and degradation under these conditions). Subsequently we have investigated the catalytic activity of these bi-metallic catalysts towards H_2O_2 degradation using a water/decan-1-ol solvent system, (Table S.4†) with the 5% AuPd/C catalyst demonstrating significantly lower rates of $H₂O₂$ degradation compared to oxide supported analogues, presumably due to the greater retention of the catalyst in the organic phase and decreased contact between catalyst and synthesised H_2O_2 .

Fig. 1 Images of 5% AuPd/supported catalysts (carbon, SiO₂, TiO₂ and CeO₂) in a two-phase solvent system consisting of decan-1-ol (top layer) and water (bottom layer).

We have previously reported that the 5% AuPd/C catalyst, offers a reasonably high activity towards H_2O_2 synthesis using a methanol–water co-solvent system under reaction conditions that are similar to those reported herein (98 $mol_{\text{H}_2\text{O}_2}$ kg_{cat}⁻¹ h⁻¹).³

We next investigated the efficacy of the 5% AuPd/C catalyst towards the direct synthesis of H_2O_2 in a range of straight chain alcohols (containing no aqueous co-solvent) as solvent (Table 1). Previous studies investigating the direct synthesis of $H₂O₂$ in short chain alcohol solvents have typically utilised acidic or halide promoters (or a combination thereof) to achieve high selectivity towards H_2O_2 .^{20,61} Under our reaction conditions, where no halide or acidic promoters are utilised, low concentrations of H_2O_2 were produced in all alcohols studied with H_2O_2 concentration decreasing with increasing carbon chain length, with this possibly resulting from a number of factors including (i) decreased $O₂$ solubility as a function of alcohol carbon-chain length, 62 or (ii) increasing H_2O_2 degradation due to improved H_2 solubility.²⁴ The ability to produce H_2O_2 therefore strongly depends on the overall rate of mass transfer of gaseous reactants to the catalyst surface. Thus, the decreasing concentration of observed H_2O_2 is likely to be due to a mixture of both chemical and physical properties of the solvent such as higher viscosity and surface tension as the alcohol carbon chain length increases. A larger pressure drop of the reagent gases was also observed during reaction (Table S.5†), indicating increased solubility of reactant gases, although the possibility for increased solubility of the $CO₂$ diluent should not be ruled out, in keeping with observations by Francesconi et al^{24} and Wainwright et al^{25} who have shown that H_2 solubility in the C1–4 primary alcohol series increases with carbon chain length and therefore it is reasonable to expect H_2 to dissolve better in decan-1-ol rather than methanol.

Another important consideration when selecting the organic solvent was the requirement to have low water solubility in the alcohol phase in order to create a bi-phasic solvent system, which would allow for ease of separation of $H₂O₂$. Table 1 shows that the use of a methanol-only solvent results in the highest H_2O_2 concentration (250 μmol). However, due to the complete miscibility of methanol with water, it is not a suitable solvent choice for this investigation. Higher carbon chain alcohols (>C6) have lower water solubility and the addition of excess water can create a biphasic solvent system.⁶³ The saturation point of the alcohols in water has been determined experimentally by Karl Fischer titration (Table 2). As decan-1-ol was determined to have the lowest water solubility (46 g_{H_2O} kg_{decan-1-ol}⁻¹), while also allowing for the formation of H_2O_2 , it was chosen as a suitable solvent for the hydrophobic solvent layer of the proposed twophase system. Furthermore, decan-1-ol has been shown to be stable under reaction conditions by NMR analysis (Fig. S.3†). Catalysis Science & Technology

Decar-1-of
 $\frac{1}{2}$
 $\frac{1}{2}$

The effect of solvent composition on the direct synthesis of $H₂O₂$, by varying the water concentration in decan-1-ol, from 0 to 590 g_{H_2O} kg_{decan-1-ol}⁻¹, was investigated (Fig. 2), while maintaining all other reaction variables, including total solvent mass which was maintained at 8.5 g. Increasing water content is observed to lead to an increase in H_2O_2 concentration, with the maximum H_2O_2 concentration (0.17) wt%) being observed at the saturation point of decan-1-ol; 46 g_{H_2O} kg_{decan-1-ol}⁻¹ (0.39 g of H₂O and 8.11 g of decan-1-ol). This is ascribed to the inhibition of H_2O_2 degradation (via hydrogenation and decomposition pathways) through the

	Table 1 Catalytic activity of 5% AuPd/C towards H_2O_2 synthesis as a function of alcohol chain length and the solubility of water in various alcohols						
Solvent		H_2O_2 productivity (mol _{H,O,} kg _{cat} ⁻¹ h ⁻¹)	$[H_2O_2]$ (wt%)	H_2O_2 produced (µmol)	H_2O solubility ^{<i>a</i>} ($g_{H,O}$ kg _{solvent} ⁻¹)		
Water		10	0.016	50			
Methanol		50	0.094	250			
Propan-1-ol		32	0.063	160			
Butan-1-ol		26	0.030	130			
Hexan-1-ol		4.4	0.008	22	94		
Octan-1-ol		1.4	0.003		53		
Decan-1-ol		0.8	0.002	4.2	46		

 $H₂O₂$ synthesis reaction conditions: 5% AuPd/C (0.01 g), total pressure (580 psi), $H₂/O₂$ (0.5), 1200 rpm, 8.5 g total solvent, 25 °C, 30 min (note: no H_2O added). a As measured by Karl Fisher titration.

Table 2 The effect of total reaction pressure on catalytic activity towards H_2O_2 formation using a bi-phasic H_2O /decan-1-ol solvent system using a 5% AuPd/C catalyst

Entry	Total pressure/psi	$[H_2O_2]$ /wt%	H_2O_2 productivity/ $mol_{H_2O_2}$ kg _{cat} ⁻¹ h ⁻¹
1 ^a	580	0.17	3.3
2	580	0.45	2.0
3	700	0.81	3.7

 $H₂O₂$ synthesis reaction conditions: 5% AuPd/C (0.02 g), total pressure (580–700 psi), H_2/O_2 (0.5), 1200 rpm, 4 g total solvent (39 g_{H₂O kg_{decan-1-ol}⁻¹), 25 °C, 30 min. ^{*a*} Non-optimised reaction} conditions. 5% AuPd/C (0.01 g), total pressure (580 psi), H_2/O_2 (0.5), 1200 rpm, 8.5 g total solvent (39 g_{H_2O} kg_{decan-1-ol}⁻¹), 25 °C, 30 min.

separation of H_2O_2 from the catalyst and the *in situ* formation of carbonic acid in the aqueous phase, through solvation of the $CO₂$ gaseous diluent, with the improved stability of $H₂O₂$ under acidic conditions well known.^{64,65} Indeed similar improvements in H_2O_2 concentrations have been observed through the introduction of water into methanol solvent systems.¹⁻³ The sharp increase in the H_2O_2 concentration up to 46 $g_{H₂O}$ kg_{decan-1-ol}⁻¹ demonstrates that the presence of water has a significant effect on the concentrations of H_2O_2 that can be achieved over the 5% AuPd/C catalyst. On further increasing the water content, to form a bi-phasic solvent, there was little effect on the H_2O_2 concentration up to 120 g_{H_2O} kg_{decan-1-ol}⁻¹. Increasing H₂O content beyond 150 g_{H_2O} kg_{decan-1-ol}⁻¹ lead to a significant decrease in H_2O_2 concentration, with this attributed to a combination of decreased H_2 availability, with increasing water content, and the inability of the decan-1-ol phase to retain the totality of the catalyst, leading to increased contact between catalyst and synthesised H_2O_2 and in turn increased H_2O_2 degradation. **Paper**
 Catalysis Science & Technology

Table The distribution in the square on consideration in the square on the comparison in the square of the state of the state

With water concentrations above 120 $g_{H,O}$ kg_{decan-1-ol}⁻¹ $(1 \text{ g } H_2O \text{ and } 7.5 \text{ g }$ decan-1-ol) it was possible to separate the

water and decan-1-ol in order to directly calculate the H_2O_2 concentration in the aqueous phase. Comparing these extracted H_2O_2 concentrations with the results obtained from titrating the entire decan-1-ol and water solvent are shown in Fig. 3. When 120 $g_{H,O}$ kg_{decan-1-ol}⁻¹ (1 g H₂O and 7.5 g decan-1-ol) and 235 g_{H_2O} kg_{decan-1-ol}⁻¹ (2 g H₂O and 6.5 g decan-1-ol) were added the difference between the results obtained through analysis of the total reaction solution or the aqueous phase only is $16-17\%$, suggesting that at low H_2O extraction of H_2O_2 is incomplete. The difference between the two measures of H_2O_2 content was subsequently decreased to 6% when the water phase was more easily separated from the decan-1-ol at 590 $g_{H,O}$ kg_{decan-1-ol}⁻¹ (5 g H₂O and 3.5 g decan-1-ol). This indicates that the efficiency of separating the two solvents will be a critical step to maintaining the formed $H₂O₂$ concentrations. All preceding results were obtained from titrating the whole solvent mixture to allow for accurate determination of H_2O_2 concentrations.

The solvent system was next investigated for its ability to suppress H_2O_2 degradation, under an atmosphere of 5% $H₂/CO₂$ (Fig. 4). Reactions were carried out with increasing amounts of pre-formed 4 wt% aqueous H_2O_2 added to decan-1-ol; from a saturated decan-1-ol solution (0.39 g) to a truly bi-phasic solvent system (3.0 g), keeping the total solvent mass constant (8.5 g). When 0.39 g of 4 wt% H_2O_2 , was added to decan-1-ol (8.11 g), to achieve a saturated decan-1-ol solution, a degradation activity of 29 mol_{H,O2} kg_{cat}⁻¹ h⁻¹ was observed. Catalytic activity towards H_2O_2 degradation was seen to increase to a value of 113 mol_{H₂O₂, kg_{cat}⁻¹ h⁻¹ when using 1 g of 4 wt% H₂O₂,} before plateauing when a higher volume, 3.0 g of 4 wt% $H₂O₂$ was added to the decan-1-ol solvent. With this plateau ascribed to the ability of H_2O_2 to catalyse the oxidation of Pd, as previously reported by Choudhary et al.,⁶⁴ with Pd^{2+}

Fig. 2 The effect of H_2O content in a H_2O /decan-1-ol solvent system on the direct synthesis of H_2O_2 . H_2O_2 concentration (black squares). H2O2 synthesis reaction conditions: 5% AuPd/C (0.01 g), total pressure (580 psi), H_2/O_2 (0.5), 1200 rpm, 8.5 g total solvent (decan-1-ol + water), 25 °C, 30 min. (-) $H₂O$ saturation point of decan-1-ol.

Fig. 3 Calculation of the H_2O_2 concentration in a mixture of decan-1ol and water (black bars) and the separated water phase (red bars). H₂O₂ synthesis reaction conditions: 5% AuPd/C (0.01 g), total pressure (580 psi), H_2/O_2 (0.5), 1200 rpm, 8.5 g total solvent (decan-1-ol + H₂O), 25 °C, 30 min.

Fig. 4 Catalytic activity of a 5% AuPd/C catalyst towards H_2O_2 degradation with increasing amounts of aqueous 4 wt% H_2O_2 . H_2O_2 synthesis reaction conditions: 5% AuPd/C (0.01 g), total pressure (580 psi), H_2/O_2 (0.5), 1200 rpm, 8.5 g total solvent decan-1-ol + X g 4 wt% H₂O₂, 25 °C, 30 min.

species known to be far less active towards H_2O_2 degradation than Pd⁰.^{66,67} The observed degradation activity, even at low amounts of H_2O_2 suggests that there is still some contact between catalyst and H_2O_2 , which may be expected given the shear forces that result from high stirring speed used in this study it is promising that the amount of H_2O_2 degraded is relatively low. Indeed, it should be noted that the degradation activity observed using the proposed solvent system is far less than that previously reported over the same catalyst when using a miscible, H2O/methanol solvent system, under identical reaction conditions (352 mol_{H₂O₂ kg_{cat}⁻¹ h⁻¹), although there are clear} differences in H_2 solubility between $H_2O/methanol$ solvent systems and those used within this study.³

The effect of solvent mass, while maintaining the mass of the 5% AuPd/C catalyst at 0.01 g, was next investigated (Fig. 5). As expected these results demonstrate that it is possible to enhance the concentration of H_2O_2 by decreasing the total solvent mass, this is mainly due to a dilution effect, producing the same amount of H_2O_2 in a smaller volume and increasing H_2 partial pressure relative to the solvent, due to increased reactor head space.

The effect of varying catalyst mass from 0.005 to 0.04 g, whilst keeping the solvent mass constant at 8.5 g, was next investigated (Fig. 6). It can be seen that with increasing mass of catalyst, H_2O_2 concentration increases in a non-linear manner, to a value of 0.25 wt% when using 0.02 g of catalyst, beyond which the increase in H_2O_2 concentration is far less pronounced. It is likely that the observed plateau in H_2O_2 concentration can be attributed to a combination of factors; (i) limitations associated with diffusion of reactant gas; (ii) the inability of the decan-1-ol to retain the catalyst forcing a higher concentration of catalyst into the aqueous phase, thus further increasing contribution of H_2O_2 degradation.

Time-on-line analysis was next carried out, with the vast majority of H_2O_2 seen to be produced over 30 minutes (0.17) wt%), although a steady rise in H_2O_2 concentration is observed over 60 minutes (0.19 wt%) (Fig. 7). Beyond 60 minutes there is a minor decrease in H_2O_2 concentration, presumably, this is due to a depletion in reagent gas

H₂O₂ synthesis. H₂O₂ synthesis reaction conditions: 5% AuPd/C (0.01 g), total pressure (580 psi), H_2/O_2 (0.5), 1200 rpm, X g total solvent (46

g_{H₂O} kg_{decan-1-ol}⁻¹), 25 °C, 30 min.

availability, in particular, H_2 and the rate of H_2O_2 degradation exceeding that of H_2O_2 synthesis, with similar observations previously reported by Crole et al. in a range of solvents.² However, it is promising that H_2O_2 concentrations can be generally retained, even at extended reaction times.

In an attempt to further enhance H_2O_2 concentration, optimised conditions were combined from the previous experiments carried out above (Fig. 2–7); 0.02 g catalyst, 4 g

Fig. 6 The effect of catalyst mass on catalytic activity towards the direct synthesis of H_2O_2 . H_2O_2 synthesis reaction conditions: 5% AuPd/ C (X mg), total pressure (580 psi), H_2/O_2 (0.5), 1200 rpm, 8.5 g total solvent (46 g_{H_2O} kg_{decan-1-ol}⁻¹), 25 °C, 30 min.

Fig. 7 The effect of reaction time on catalytic activity towards the direct synthesis of H_2O_2 . H_2O_2 synthesis reaction conditions: 5% AuPd/C (0.01 g), total pressure (580 psi), H₂/O₂ (0.5), 1200 rpm, 8.5 g total solvent (46 g $_{\mathsf{H}_2\mathsf{O}}$ kg_{decan-1-ol} $^{-1}$), 25 °C, X min.

total solvent with 46 $g_{H,O}$ kg_{decan-1-ol}⁻¹ and a reaction time of 30 minutes. It can be seen (Table 2, entry 2) that when the direct synthesis reaction was carried out with a total reactant gas pressure of 580 psi $(H_2: O_2 = 0.5)$ it resulted in an H_2O_2 concentration of 0.45 wt%, over 2.5 times greater than that observed over non-optimised conditions (0.17 wt%). Further increasing total reaction pressure to 700 psi, while maintaining optimised reaction conditions (Table 2, entry 3) is observed to result in a 4-fold increase in H_2O_2 concentration (0.81 wt%) over standard conditions.

Finally, with the plateau in H_2O_2 concentration, previously observed (Fig. 7) we carried out a series of sequential H_2O_2 direct synthesis reactions under our optimised reaction conditions (Fig. 8). We observe a steady increase in H_2O_2 concentration up to 1.9 wt% after 5 sequential reactions.

Fig. 8 Sequential H_2O_2 synthesis reactions, under optimised reaction conditions using a 5% AuPd/C catalyst. H_2O_2 synthesis reaction conditions: 5% AuPd/C (0.02 g), total pressure (700 psi), $H₂/O₂$ (0.5), 1200 rpm, 4 g total solvent (46 g_{H2}o kg_{decan-1-ol} $^{-1}$), 25 °C, 30 min.

Although a limited volume of H_2O_2 is reported (0.16 ml of 1.9 wt% H_2O_2) at low productivities, the concentration achieved highlights the advantages of using a bi-phasic solvent for the direct synthesis of H_2O_2 . If facile separation of the organic and aqueous phases can be achieved, the H_2O_2 concentration is comparable to that achieved in the initial stages of the current indirect method of industrial H_2O_2 production, prior to the use of multiple distillation steps to raise H_2O_2 concentrations to exceed ∼70 wt%.36

Conclusion

In this paper the direct synthesis of H_2O_2 from H_2 and O_2 over a 5% AuPd/C catalyst has been investigated using a solvent system containing decan-1-ol and water. As a result of the preference for the catalyst to be retained in the hydrophobic, organic component, thus limiting contact between the synthesised H_2O_2 and catalyst a significant improvement in catalytic efficacy is observed, compared to the use of a miscible solvent mixture. Through optimisation of the reaction conditions, and assuming facile separation of the organic and aqueous solvent, we demonstrate that it is possible to reach concentrations of H_2O_2 comparable to that produced in the initial stages of H_2O_2 production, on an industrial scale, with a $H₂O₂$ concentration of 1.9 wt% reported. Perhaps more importantly we demonstrate the feasibility of a bi-phasic solvent system in allowing simultaneous reaction and extraction, minimising H_2O_2 degradation rates, resulting in increased concentrations of H_2O_2 . We consider that this work represents a promising basis for further exploration of a wider set of solvents and catalysts for use in the direct synthesis of $H₂O₂$. In particular we believe that, given the high desirability for a continuous production of H_2O_2 , this approach would lend itself well to a flow regime. **Catalysis Science Articles**
 $\frac{1}{2}$ **Catalysis Articles** Articles Article

Conflicts of interest

The authors declare no conflict of interests.

Acknowledgements

The authors wish to acknowledge the financial support of and research discussion with Solvay S. A.

References

- 1 A. Santos, R. J. Lewis, G. Malta, A. G. R. Howe, D. J. Morgan, E. Hampton, P. Gaskin and G. J. Hutchings, Ind. Eng. Chem. Res., 2019, 58, 12623–12631.
- 2 D. A. Crole, S. J. Freakley, J. K. Edwards and G. J. Hutchings, Proc. R. Soc. London, Ser. A, 2016, 472, 20160156.
- 3 S. J. Freakley, R. J. Lewis, D. J. Morgan, J. K. Edwards and G. J. Hutchings, Catal. Today, 2015, 248, 10–17.
- 4 T. Deguchi and M. Iwamoto, J. Catal., 2011, 280, 239–246.
- 5 E. N. Ntainjua, M. Piccinini, S. J. Freakley, J. C. Pritchard, J. K. Edwards, A. F. Carley and G. J. Hutchings, Green Chem., 2012, 14, 170.

Catalysis Science & Technology Paper

- 6 J. Edwards, B. Solsona, P. Landon, A. Carley, A. Herzing, C. Kiely and G. Hutchings, J. Catal., 2005, 236, 69–79.
- 7 M. Piccinini, E. Ntainjua, J. K. Edwards, A. F. Carley, J. A. Moulijn and G. J. Hutchings, Phys. Chem. Chem. Phys., 2010, 12, 2488–2492.
- 8 V. V. Krishnan, A. G. Dokoutchaev and M. E. Thompson, J. Catal., 2000, 196, 366–374.
- 9 R. Burch and P. R. Ellis, Appl. Catal., B, 2003, 42, 203–211.
- 10 R. J. Lewis and G. J. Hutchings, ChemCatChem, 2019, 11, 298–308.
- 11 R. E. Brooks and S. B. Moore, Cellulose, 2000, 7, 263–286.
- 12 C. Samanta, Appl. Catal., A, 2008, 350, 133–149.
- 13 P. Landon, P. J. Collier, A. F. Carley, D. Chadwick, A. J. Papworth, A. Burrows, C. J. Kiely and G. J. Hutchings, Phys. Chem. Chem. Phys., 2003, 5, 1917–1923.
- 14 C. M. Piqueras, J. García-Serna and M. J. Cocero, J. Supercrit. Fluids, 2011, 56, 33–40.
- 15 T. Moreno, J. García-Serna and M. J. Cocero, Green Chem., 2010, 12, 282–289.
- 16 Q. Chen and E. J. Beckman, Green Chem., 2007, 9, 802–808.
- 17 F. Moseley and P. N. Dyer, US Pat., 4336240, 1982.
- 18 I. T. Horváth, Acc. Chem. Res., 1998, 31, 641–650.
- 19 M. Kawakami, Y. Ishiuchi, H. Nagashima, T. Tomita and Y. Hiramatsu, US Pat., 5399334, 1995.
- 20 Q. Liu, J. C. Bauer, R. E. Schaak and J. H. Lunsford, Appl. Catal., A, 2008, 339, 130–136.
- 21 S. Melada, F. Pinna, G. Strukul, S. Perathoner and G. Centi, J. Catal., 2006, 237, 213–219.
- 22 V. Paunovic, V. V. Ordomsky, V. L. Sushkevich, J. C. Schouten and T. A. Nijhuis, ChemCatChem, 2015, 7, 1161–1176.
- 23 E. J. Beckman and J. Supercrit, Fluids, 2004, 28, 121–191.
- 24 J. V. H. d'Angelo and A. Z. Francesconi, J. Chem. Eng. Data, 2001, 46, 671–674.
- 25 M. S. Wainwright, T. Ahn, D. L. Trimm and N. W. Cant, J. Chem. Eng. Data, 1987, 32, 22–24.
- 26 Z. Laughrey, E. Bear, R. Jones and M. A. Tarr, Ultrason. Sonochem., 2001, 8, 353–357.
- 27 R. J. Lewis, K. Ueura, Y. Fukuta, S. J. Freakley, L. Kang, R. Wang, Q. He, J. K. Edwards, D. J. Morgan, Y. Yamamoto and G. J. Hutchings, ChemCatChem, 2019, 11, 1673–1680.
- 28 T. Pospelova and N. Kobozev, Russ. J. Phys. Chem., 1961, 35, 584–587.
- 29 E. N. Ntainjua, M. Piccinini, J. C. Pritchard, J. K. Edwards, A. F. Carley, J. A. Moulijn and G. J. Hutchings, ChemSusChem, 2009, 2, 575–580.
- 30 S. Abate, P. Lanzafame, S. Perathoner and G. Centi, Catal. Today, 2011, 169, 167–174.
- 31 S. Melada, F. Pinna, G. Strukul, S. Perathoner and G. Centi, J. Catal., 2005, 235, 241–248.
- 32 E. N. Ntainjua, M. Piccinini, J. C. Pritchard, Q. He, J. K. Edwards, A. F. Carley, J. A. Moulijn, C. J. Kiely and G. J. Hutchings, ChemCatChem, 2009, 1, 479–484.
- 33 J. K. Edwards, B. Solsona, E. N. Ntainjua, A. F. Carley, A. A. Herzing, C. J. Kiely and G. J. Hutchings, Science, 2009, 323, 1037–1041.
- 34 S. J. Freakley, Q. He, J. H. Harrhy, L. Lu, D. A. Crole, D. J. Morgan, E. N. Ntainjua, J. K. Edwards, A. F. Carley, A. Y. Borisevich, C. J. Kiely and G. J. Hutchings, Science, 2016, 351, 965–968. Catalysis Science & Technology
 θ Lebesta, A. Statey, A. Herring, C. 34 S. J. Neulig, A. Evaluation, L. This article, L. C. This article is licensed to Figure 2020, A. States Articles. Article is licensed to Figure 202
	- 35 N. M. Wilson, J. Schröder, P. Priyadarshini, D. T. Bregante, S. Kunz and D. W. Flaherty, J. Catal., 2018, 368, 261–274.
	- 36 H. Li, B. Zheng, Z. Pan, B. Zong and M. Qiao, Front. Chem. Sci. Eng., 2018, 12, 124–131.
	- 37 Q. Liu and J. H. Lunsford, Appl. Catal., A, 2006, 314, 94–100.
	- 38 Y. Kawase, Y. Isaka, Y. Kuwahara, K. Mori and H. Yamashita, Chem. Commun., 2019, 55, 6743–6746.
	- 39 Y. Isaka, Y. Kawase, Y. Kuwahara, K. Mori and H. Yamashita, Angew. Chem., Int. Ed., 2019, 58, 5402–5406.
	- 40 X. Chen, Y. Kuwahara, K. Mori, C. Louis and H. Yamashita, J. Mater. Chem. A, 2020, 8, 1904–1910.
	- 41 J. E. Romo, N. V. Bollar, C. J. Zimmermann and S. G. Wettstein, ChemCatChem, 2018, 10, 4805–4816.
	- 42 V. M. Blasucci, Z. A. Husain, A. Z. Fadhel, M. E. Donaldson, E. Vyhmeister, P. Pollet, C. L. Liotta and C. A. Eckert, J. Phys. Chem. A, 2010, 114, 3932–3938.
	- 43 X. Jin, J. Feng, H. Song, J. Yao, Q. Ma, M. Zhang, C. Yu, S. Li and S. Yu, Green Chem., 2019, 21, 3583–3596.
	- 44 G.-H. Han, X. Xiao, J. Hong, K.-J. Lee, S. Park, J.-P. Ahn, K.-Y. Lee and T. Yu, ACS Appl. Mater. Interfaces, 2020, 12, 6328–6335.
	- 45 Z. Cheng, R. Lippi, C. E. Li, Y. Yang, L. Tang, S. Huang, W. J. Lee, S. Lim, X. Ma and J. Patel, Ind. Eng. Chem. Res., 2019, 58, 20573–20584.
	- 46 R. J. Lewis, J. K. Edwards, S. J. Freakley and G. J. Hutchings, Ind. Eng. Chem. Res., 2017, 56, 13287–13293.
	- 47 E. N. Ntainjua, M. Piccinini, J. C. Pritchard, J. K. Edwards, A. F. Carley, C. J. Kiely and G. J. Hutchings, Catal. Today, 2011, 178, 47–50.
	- 48 E. N. Ntainjua, J. K. Edwards, A. F. Carley, J. A. Lopez-Sanchez, J. A. Moulijn, A. A. Herzing, C. J. Kiely and G. J. Hutchings, Green Chem., 2008, 10, 1162.
	- 49 P. Tian, L. Ouyang, X. Xu, C. Ao, X. Xu, R. Si, X. Shen, M. Lin, J. Xu and Y.-F. Han, J. Catal., 2017, 349, 30–40.
	- 50 A. Cybula, J. B. Priebe, M.-M. Pohl, J. W. Sobczak, M. Schneider, A. Zielińska-Jurek, A. Brückner and A. Zaleska, Appl. Catal., B, 2014, 152–153, 202–211.
	- 51 X. Gong, R. J. Lewis, S. Zhou, D. J. Morgan, T. E. Davies, X. Liu, C. J. Kiely, B. Zong and G. J. Hutchings, Catal. Sci. Technol., 2020, 10, 4635–4644.
	- 52 L. Ouyang, P.-F. Tian, G.-J. Da, X.-C. Xu, C. Ao, T.-Y. Chen, R. Si, J. Xu and Y.-F. Han, J. Catal., 2015, 321, 70–80.
	- 53 C. M. Crombie, R. J. Lewis, D. Kovačič, D. J. Morgan, T. E. Davies, J. K. Edwards, M. S. Skjøth-Rasmussen and G. J. Hutchings, Catal. Lett., 2020, DOI: 10.1007/s10562- 020-03281-1.
	- 54 J. K. Edwards, A. F. Carley, A. A. Herzing, C. J. Kiely and G. J. Hutchings, Faraday Discuss., 2008, 138, 225.
	- 55 J. K. Edwards, A. Thomas, A. F. Carley, A. A. Herzing, C. J. Kiely and G. J. Hutchings, Green Chem., 2008, 10, 388.
	- 56 S. Kanungo, V. Paunovic, J. C. Schouten and M. F. Neira D'Angelo, Nano Lett., 2017, 17, 6481–6486.

- 57 S. Kanungo, L. van Haandel, E. J. M. Hensen, J. C. Schouten and M. F. Neira d'Angelo, J. Catal., 2019, 370, 200–209.
- 58 J. Li, T. Ishihara and K. Yoshizawa, J. Phys. Chem. C, 2011, 115, 25359–25367.
- 59 N. M. Wilson, P. Priyadarshini, S. Kunz and D. W. Flaherty, J. Catal., 2018, 357, 163–175.
- 60 A. V. Beletskaya, D. A. Pichugina, A. F. Shestakov and N. E. Kuz'menko, J. Phys. Chem. A, 2013, 117, 6817–6826.
- 61 F. Menegazzo, P. Burti, M. Signoretto, M. Manzoli, S. Vankova, F. Boccuzzi, F. Pinna and G. Strukul, J. Catal., 2008, 257, 369–381. **Catalysis Science & Technology**

ST S. Rammage, L. Tea Handel, *C. J. D.* H. This article. Company, C. C. J. D. J
	- 62 S. Bo, R. Battino and E. Wilhelm, J. Chem. Eng. Data, 1996, 41, 644.
	- 63 N. Šegatin and C. Klofutar, Monatsh. Chem., 2004, 135, 241–248.
	- 64 V. R. Choudhary, C. Samanta and T. V. Choudhary, J. Mol. Catal. A: Chem., 2006, 260, 115–120.
	- 65 V. R. Choudhary, C. Samanta and P. Jana, Appl. Catal., A, 2007, 317, 234–243.
	- 66 A. G. Gaikwad, S. D. Sansare and V. R. Choudhary, J. Mol. Catal. A: Chem., 2002, 181, 143–149.
	- 67 G. Blanco-Brieva, E. Cano-Serrano, J. M. Campos-Martin and J. L. G. Fierro, Chem. Commun., 2004, 1184–1185.