A review of recent advances in the direct synthesis of H$_2$O$_2$.

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**Graphical abstract**

The direct synthesis of hydrogen peroxide represents an atom efficient way to make this important commodity chemical. In this mini review we discuss some of the latest advances for this reaction and also point out the challenge that remain to be solved so that this reaction can be considered ready for commercial exploitation; namely the need for increased catalyst activity so that more concentrated solutions can be synthesised.

**Key words:**

Hydrogen Peroxide  
Direct Synthesis  
Heterogeneous Catalysis  
Palladium  
In Situ Oxidation
Abstract

Hydrogen peroxide ($\text{H}_2\text{O}_2$) is a highly effective, green oxidant that has found application in sectors ranging from the synthesis of fine chemicals and waste stream treatment to the extraction of precious metals and the bleaching of paper pulp and textiles. The growing demand for $\text{H}_2\text{O}_2$ has seen it become one of the 100 most important chemicals in the world. The direct synthesis of $\text{H}_2\text{O}_2$ from $\text{H}_2$ and $\text{O}_2$ has been a challenge for the scientific community for over 100 years and represents an attractive alternative to the current means of production. Herein we discuss the historical perspective of the direct synthesis process, the recent literature regarding catalyst design and the role of additives as well as the application of $\text{H}_2\text{O}_2$ as an in situ oxidant. We discuss the key problems that remain and conclude that although there has been progress with respect to the selectivity of hydrogen utilisation, there is a need to now concentrate on catalyst activity as the key remaining problem requiring a solution is the concentration of $\text{H}_2\text{O}_2$ that can be achieved especially in flow reactors.

Introduction.

Hydrogen peroxide ($\text{H}_2\text{O}_2$) is a powerful, environmentally friendly, oxidant that can oxidise both inorganic and organic substrates, under mild conditions. The uses of $\text{H}_2\text{O}_2$ are predominantly in applications where its efficacy as a bleaching agent are required, or those that utilize the high active oxygen potential, such as the synthesis of fine chemicals. Unlike stoichiometric oxidants, such as tBuOOH, $\text{N}_2\text{O}$ or permanganate which produce large amounts of waste that requires separation from the desired product, $\text{H}_2\text{O}_2$ utilisation results only in the co-production of $\text{H}_2\text{O}$.

In recent years, global $\text{H}_2\text{O}_2$ production has exceeded 3 million tons per annum\cite{1} and is predicted to rise growing at a rate of approximately 4 % with demand forecast to reach 5.2 million tons per annum by 2020.\cite{2}

More niche uses of $\text{H}_2\text{O}_2$ include its use in the mining sector for the extraction of gold and uranium.\cite{10} Further demand for $\text{H}_2\text{O}_2$ is driven from its application in chemical synthesis (see Figure 2) with typical applications of $\text{H}_2\text{O}_2$ found in the integrated HPPO process\cite{12-19} and the ammonoximation of cyclohexanone to cyclohexanone oxime, a key intermediate in the formation of Nylong-6.\cite{20-23} These routes for chemical synthesis have, in-part, been driven by the discovery of

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Applications of $\text{H}_2\text{O}_2$}
\end{figure}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Schematic representation of oxidation reactions catalysed by $\text{H}_2\text{O}_2$ in conjunction with TS-1.}
\end{figure}
Current production of H₂O₂ on an industrial scale is limited to electrolytic production, the partial oxidation of isopropanol and the well-established anthraquinone oxidation (AO) or in-direct synthesis process. The AO process, which accounts for more than 95% of global H₂O₂ production was first developed by Riedl and Pfleiderer in BASF in 1939,[39] although the production of H₂O₂ can be traced back to its isolation in 1818 by L.J. Thenard.[39] The original process developed by Ridel and Pfleiderer has undergone continual improvement since 1939 and so is highly efficient, although the underlying chemistry has changed little and utilises H₂, O₂ and an anthraquinone derivative, with the latter reduced over a Pd based catalyst to produce the diol. The oxidation of the subsequent anthracinol reforms the original anthraquinone in tandem with H₂O₂ formation. This highly efficient process is able to produce H₂O₂ concentrations of 1-2 wt. %, through further purification and distillation it is possible to produce H₂O₂ concentrations in excess of 70 wt. % which can be shipped and stored prior to dilution at point of final use. Often on-site application of H₂O₂ will require dilution to a range of 1-10 wt. %

Although the anthraquinone process is highly efficient, there are some concerns regarding its carbon efficiency, with the unselective hydrogenation of the carrier molecule resulting in the need for its periodic replacement. This coupled with the overall complexity of the process, in particular the choice of appropriate solvent, requires production to occur on a large scale, often precluding the synthesis of H₂O₂ at point of use. Furthermore, the instability of H₂O₂, with its rapid decomposition to H₂O in the presence of relatively mild temperatures or weak bases requires the use of stabilizing agents, often acidic stabilizers such as; acetic acid, peracetic acid,[40] dipicolinic acid, quinolinic acid[41] or phosphoric acid[42] are utilised. However, the use of such stabilizing agents often lead to reactor corrosion as well increased costs associated with their downstream removal.

The direct synthesis of H₂O₂ as an alternative to the anthraquinone process.

The direct synthesis of H₂O₂ from molecular H₂ and O₂ has the potential to offer a more atomically efficient route than the current industrial process, where the non-selective hydrogenation of the quinone-derivative H₂ carrier molecule necessitates its periodic replacement. Pd based catalysts have been known to be highly active for the direct synthesis of H₂O₂ since 1914[43] and have received significant attention in the literature.[44-50] However, a major challenge associated with the direct synthesis of H₂O₂ is associated with catalyst selectivity; often catalysts that offer high activity towards the direct synthesis of H₂O₂ are also active to it’s degradation via over hydrogenation or decomposition to H₂O.[47, 51-53] The issue of catalyst selectivity can be understood as the formation of water from H₂ and O₂ is thermodynamically favourable in comparison to the formation of H₂O₂ as summarized by Equations 1-2.

1. H₂(g) + O₂(g) → H₂O₂(l)
   \[\Delta G_{298K} = -120.5 \text{ kJ / mol}\]
2. H₂(g) + ½ O₂(g) → H₂O(l)
   \[\Delta G_{298K} = -237.2 \text{ kJ / mol}\]
Furthermore, the undesired subsequent \( \text{H}_2\text{O}_2 \) decomposition and hydrogenation reactions are also thermodynamically favourable:

3. \( \text{H}_2\text{O}_2 (l) \rightarrow \text{H}_2\text{O} (l) + \frac{1}{2} \text{O}_2 (g) \)
   \( \Delta G^{298K} = -116.7 \text{ kJ} / \text{mol} \)

4. \( \text{H}_2\text{O}_2 (l) + \text{H}_2 (g) \rightarrow 2 \text{H}_2\text{O} (l) \)
   \( \Delta G^{298K} = -354.0 \text{ kJ} / \text{mol} \)

The direct synthesis of \( \text{H}_2\text{O}_2 \), therefore demonstrates the need for catalyst design to balance selectivity and activity carefully, as well as the selection of reaction conditions that inhibit the degradation of \( \text{H}_2\text{O}_2 \). It is known that \( \text{H}_2\text{O}_2 \) is highly unstable at high temperatures or in the presence of basic conditions and it has been demonstrated that through the use of low reaction temperatures it is possible to limit the thermodynamic favourability of the subsequent \( \text{H}_2\text{O}_2 \) degradation reactions.\[^{[54]}\]

Although experimental studies have demonstrated that the active site for the direct synthesis of \( \text{H}_2\text{O}_2 \) and its decomposition may be different the exact nature of sites responsible for both the formation of \( \text{H}_2\text{O}_2 \) and its subsequent degradation remains unclear.\[^{[51, 55]}\] It is widely accepted that the formation of \( \text{H}_2\text{O}_2 \) proceeds via the addition of hydrogen to \( \text{O}_2^- \), with \( \text{O}_2 \) isotope labelling experiments revealing that the irreversible cleavage of the \( \text{O}-\text{O} \) bond prevents the formation of \( \text{H}_2\text{O}_2 \), with a resulting production of \( \text{H}_2\text{O} \).\[^{[56, 57]}\] Thus perhaps the most significant challenge in terms of catalyst design is suppressing the formation of thermodynamically favoured intermediates that result in the production of \( \text{H}_2\text{O} \), namely \( \text{O}^- \) and \( \text{OH}^+ \).\[^{[58]}\]

There is significant debate within the literature around whether Pd as a metal or an oxide favours the formation of \( \text{H}_2\text{O}_2 \). Choudhary and co-workers\[^{[59]}\] have studied supported Pd catalysts for the direct synthesis of \( \text{H}_2\text{O}_2 \) and have reported that the degradation of \( \text{H}_2\text{O}_2 \) over supported Pd catalysts strongly depends on the presence of Pd\(^0\). Indeed Choudhary and co-workers\[^{[60, 61]}\] and others\[^{[62]}\] have reported that supported PdO catalysts offer much greater selectivity and activity for \( \text{H}_2\text{O}_2 \) synthesis than corresponding Pd\(^0\) catalysts, and the greater selectivity of PdO or Pd\(^2+\) compared to that of Pd\(^0\) can be attributed to the lower \( \text{H}_2\text{O}_2 \) decomposition activity of the PdO catalysts. Alternatively, Burch and Ellis\[^{[63]}\] have reported the reduction of supported PdO catalysts prior to use can enhance both H\(_2\) conversion and selectivity towards \( \text{H}_2\text{O}_2 \), with Liu et al. also reporting higher \( \text{H}_2\text{O}_2 \) yields over supported Pd\(^0\) catalysts.\[^{[64]}\] Although it should be noted that both Strukul and co-workers\[^{[65]}\] and Lunsford\[^{[66]}\] have reported a change in Pd oxidation state during the direct synthesis reaction. While studies by Ouyang et al. have revealed the production of \( \text{H}_2\text{O}_2 \) may take place at the interface between Pd and PdO indicating a strong dependence on both phases.\[^{[67, 68]}\] The work of Flaherty and co-workers also indicates the need for a proportion of nanoparticles to exist in the reduced state, with a significant induction time required for PdO nanoparticles to exhibit activity towards \( \text{H}_2\text{O}_2 \) formation, with no such induction period exists for Pd nanoparticles.\[^{[69]}\]

Although the nature of the active site responsible for the cleavage of the \( \text{O}-\text{O} \) bond and production of \( \text{H}_2\text{O} \) is unknown it is likely that \( \text{H}_2\text{O}_2 \) degradation is promoted by the high energy, low co-ordination Pd centres.\[^{[70]}\] Plauck et al. have, through a combination of density functional theory and reaction kinetic studies, reported that both the close-packed Pd (111) and more open Pd (100) facets can represent the active site for \( \text{H}_2\text{O}_2 \) decomposition for supported Pd nanoparticles.\[^{[71]}\] This is in keeping with the work of Kim et al.\[^{[49]}\] who reported enhanced activity of Pd of the Pd (111) facet towards \( \text{H}_2\text{O}_2 \) synthesis. However, computational studies by Yoshizawa and co-workers have revealed that the Pd (111) surface also has capability to catalyse the non-selective formation of \( \text{H}_2\text{O} \).\[^{[72]}\]

The role of additives in the direct synthesis of \( \text{H}_2\text{O}_2 \).

To overcome limitations around selectivity Pd supported catalysts have often relied on the presence of acid and halide promoters to suppress the routes responsible for \( \text{H}_2\text{O}_2 \) degradation. Pospelova et al.\[^{[73]}\] were the first to report an inhibition in \( \text{H}_2\text{O}_2 \) decomposition with the addition of inorganic acids, potentially by preventing the deprotonation of \( \text{H}_2\text{O}_2 \) to \( \text{OOH}^- \) and its subsequent decomposition.\[^{[74]}\] Lunsford and co-workers demonstrated the need for careful control of acid
concentrations to prevent leaching of supported Pd, resulting in a loss of catalyst stability and the development of a homogeneous catalytic component.\textsuperscript{[86, 57, 75]}

Despite the majority of research centering around the use of acids within the reaction solution, either in the form of oxyacids, such as phosphoric acid and sulphuric acid, or halogen acids, such as hydrochloric acid\textsuperscript{[76]} the use of solid acid additives such as Cs-exchanged tungstophosphoric acid has also shown promise in the direct synthesis of H\textsubscript{2}O\textsubscript{2}.\textsuperscript{[77]} Edwards et al. have also demonstrated that a beneficial effect can be achieved through the acid pre-treatment of both oxide\textsuperscript{[78, 79]} and carbon\textsuperscript{[80]} supports prior to metal deposition with a dramatic enhancement in catalytic selectivity over the AuPd / C catalyst in particular. This is ascribed to an enhancement in Au dispersion and a resulting increase in the proportion of smaller (2 to 5 nm) bi-metallic particles at the expense of large (> 50nm) Au-rich nanoparticles. Further investigations by Garcia et al.\textsuperscript{[81, 82]} have also demonstrated that enhanced catalytic selectivity can be achieved through the modification of a mesoporous carbon support, prior to precious metal impregnation, through a similar means to that previously reported by Edwards et al.\textsuperscript{[76, 77]}

Recently Wilson and Flaherty\textsuperscript{[83]} have completed a comprehensive mechanistic study of the direct synthesis of H\textsubscript{2}O\textsubscript{2} over a Pd supported catalyst and have demonstrated that the presence of protons are key for H\textsubscript{2}O\textsubscript{2} synthesis, with H\textsubscript{2}O\textsubscript{2} yields in protic solvents such as H\textsubscript{2}O and CH\textsubscript{3}OH much greater than in aprotic solvents, such as acetonitrile (Figure 3). In the same work they report that the presence of protons, from mineral acids can aid in the reduction of molecular O\textsubscript{2}, a key step in H\textsubscript{2}O\textsubscript{2} synthesis and make compelling arguments for the importance of the counterions, such as SO\textsubscript{4}\textsuperscript{2-} and Cl\textsuperscript{-} in determining selectivity towards H\textsubscript{2}O\textsubscript{2}. In particular they conclude the enhancement in H\textsubscript{2}O\textsubscript{2} selectivity can be related to a combination of electronic modification of the reaction solution at the liquid-solid interface by these anions as well as their adsorption onto the surface of Pd nanoparticles, which must be displaced prior to O-O bond cleavage. This explanation is in good agreement with the findings already discussed within the literature.\textsuperscript{[84, 85]}

\textbf{Figure 3.} The formation of H\textsubscript{2}O\textsubscript{2} as a function of time using protic (methanol (black squares), water (red circles)) or aprotic (dimethyl sulfoxide (green triangles), acetonitrile (blue inverted triangles), propylene carbonate (magenta diamonds)). Reproduced from ref. \textsuperscript{83}

Numerous studies have revealed the beneficial role of halides in the direct synthesis reaction, in particular bromide, either incorporated on the support\textsuperscript{[86-88]} or within the reaction solution, often in conjunction with acids.\textsuperscript{[89, 90]} It is generally considered that halides aid the inhibition of H\textsubscript{2}O\textsubscript{2} degradation to H\textsubscript{2}O and enhancing catalytic selectivity while the role of acids is to aid in H\textsubscript{2}O\textsubscript{2} stability by diminishing the base catalysed decomposition of H\textsubscript{2}O\textsubscript{2}. Previously halides had been purported to enhance catalytic selectivity through binding to sites responsible for the cleavage of the O-O bond.\textsuperscript{[52, 91]} resulting in the formation of H\textsubscript{2}O, recent studies have now revealed that the role of halides may be far more complex. Work by Biasi et al. has revealed that through post-synthesis catalytic treatment bromide can enhance catalytic activity through promoting a restructuring of the metal phase in addition to the previously suggested poisoning the highly active sites responsible for H\textsubscript{2}O\textsubscript{2} degradation.\textsuperscript{[92, 93]} Furthermore it has been suggested that the promotional effect associated with halide use can be attributed to an inhibition of electron back-donation to O\textsubscript{2} 2\pi* orbitals aiding in the maintenance of the O-O bond.\textsuperscript{[94]} It is therefore likely that a combination of site poisoning and electronic and structural modification all contribute to an enhancement in catalytic selectivity through the addition of halides. Although bromide has been found to be optimal for inhibiting sites responsible for O-O bond cleavage Choudhary et al. have found that the simultaneous use of fluoride or iodide in addition to
bromide is able to inhibit all three routes to H$_2$O formation (decomposition, hydrogenation and combustion). However, it should be noted that the addition of one halide, in particular chloride or bromide, only results in an inhibition of H$_2$O$_2$ decomposition and catalytic activity towards H$_2$O$_2$ hydrogenation remains. This observation implies that there may be significant differences between the active sites responsible for the decomposition and hydrogenation of H$_2$O$_2$. Finally, the use of increasing concentrations of halides has been reported to cause an increase in Pd particle size, possibly through the leaching and re-deposition of Pd with Tian et al. recently elucidating the effect of Pd particle size on activity towards H$_2$O$_2$ synthesis. Often the utilisation of increasing concentrations of halides, in particular chloride, results in an oxidation of metallic Pd and enhanced dissolution of the support resulting in the formation of soluble Pd$^{2+}$ complexes and a reduction in catalyst lifetime.

The role of secondary metals in the direct synthesis of H$_2$O$_2$.

Although Pd based catalysts are highly active towards the formation of H$_2$O$_2$ from molecular H$_2$ and O$_2$ as noted previously, they often display significant activity towards its subsequent degradation, through the hydrogenation and decomposition pathways. Landon et al. were the first to report the activity of Au supported catalysts for the direct synthesis of H$_2$O$_2$. Subsequently Haruta and co-workers reported the activity of Au catalysts supported on a range of supports with Ishihara et al. demonstrating that promising H$_2$O$_2$ yields could be achieved, in the absence of halide promoters over a Au / SiO$_2$ catalyst and that through Pd addition H$_2$O$_2$ synthesis rate could be greatly enhanced. With subsequent work by Todorovic and Meyer investigating the catalytic activity of Au, Pd and Pt crystal planes towards the direct synthesis and subsequent degradation of H$_2$O$_2$ via DFT calculations, Edwards et al. were the first to report increased catalytic activity towards H$_2$O$_2$ formation when alloying Au and Pd, building on these initial findings subsequent investigation of Pd-based, bi- and tri-metallic catalysts for the direct synthesis reaction with numerous combinations of catalysts since reported, including; Pd-Pt, Pd-Ru, Pd-Rh, Pd-Ir and Au-Pd-Pt.

A comprehensive study by Deguchi et al. has investigated the effect of precious metal (Au, Pt, Ru, Rh and Ir) addition to a Pd-polyvinylpyrrolidone colloid, with dramatic enhancements being observed with the addition of very low (0.5 at.%) concentrations of Pt or Ir doubling the formation rate of H$_2$O$_2$. Kinetic analysis, supported by DFT studies, indicated that the high H$_2$ activating abilities of Pt and Ir were responsible for the enhanced H$_2$O$_2$ synthesis reaction rate, while the subsequent H$_2$O$_2$ hydrogenation rate remained fairly constant with Pt and Ir incorporation, indicating that the H$_2^+$ activated on Ir or Pt played little role in the degradation of H$_2$O$_2$.

Hutchings and co-workers and others have extensively studied the synergistic effect achieved through the combination of Au and Pd. With the development of Au core- PdO shell nanoparticles morphology upon calcination often reported as key for improvement in catalytic selectivity, with Cybula et al. following the effect of calcination temperature on Au-Pd nanoparticles via HAADF microscopy combined with elemental mapping (Figure 4) and Tiruvalam comparing the nature of Au-Pd nanoparticle structure and morphology for the direct production of H$_2$O$_2$.

![Figure 4. HAADF images combined with elemental mapping of Au-Pd modified TiO$_2$.](image)

The means by which Au incorporation enhances catalytic selectivity is widely debated with electronic, structural and isolation effects all potential causes for the enhanced activity of Au-Pd supported catalysts. In many cases, the observed synergy is likely to be a combination these factors, but providing conclusive evidence on the nature of the enhancement is
exceptionally challenging. A number of theoretical studies by Yoshizawa and co-workers compared Pd and PdAu surfaces and provided evidence that the presence of Au inhibits the formation of H$_2$O, compared to that over a Pd (111) surface.$^{124, 125}$ This is in good agreement with the work of Han and Mullins who have recently probed the influence of the surface composition of Au-Pd catalysts for O-O bond dissociation, widely believed to be a key step for the formation of H$_2$O, and have reported that the extent of O$_2$ dissociation is proportional to Pd content, with the increasing development of Pd ensembles attributed as the cause for increased O-O bond cleavage and a resulting increase in H$_2$O formation.$^{126}$ A further study by Li and Yoshizawa$^{127}$ into the role of Au in bi-metallic AuPd systems has suggested a more direct involvement of Au in the production of H$_2$O$_2$ (Figure 5). They calculated the energetic favourability of O-O bond cleavage that exists over a pure Pd surface is reduced through the introduction of Au, to a state where the maintenance of the O-O bond is favoured, resulting in the high selectivity of AuPd systems reported experimentally in the literature. That is, the presence of Au in Pd-Au interface sites weakens the Pd-O interactions, leading to an enhanced selectivity towards the O-O bond.

![Figure 5](image5.png)

**Figure 5.** The case for the direct involvement of Au in the direct synthesis of H$_2$O$_2$ over AuPd supported catalysts as proposed by Li and Yoshizawa. Reproduced from ref. [127].

Until recently the modification of Pd with other precious metals, including Au, Pt, Rh, Ru and Ir have been the focus of many studies, with the modification of Pd by Au incorporation in particular widely reported. However, the choice incorporating secondary noble metals such as Au and Pt into supported Pd catalysts can significantly enhance costs, with both Au and Pt being more expensive than Pd which might prohibit the application of such catalysts on an industrial scale. As such replacing secondary noble metals with more abundant, cost effective alternatives is of great importance. To this end there is growing interest in the use of base metals as secondary metals in conjunction with Pd. Freakley et al.$^{128}$ recently reported a series of supported Pd-base metal catalysts, which are completely selective towards the direct synthesis of H$_2$O$_2$. Through successive calcination-reduction-calcination heat treatments of the catalyst it has been shown that the degradation of H$_2$O$_2$ can be completely inhibited. The detailed investigation of a Pd Sn/ TiO$_2$ catalyst that has been exposed to this optimized heat treatment cycle has yielded an approach that is believed to be applicable to a number of other Pd-base metal combinations, including Ni, Zn, Ga, In and Co.$^{128}$ It was proposed that as a result of this heat treatment the small Pd-rich nanoparticles responsible for H$_2$O$_2$ degradation are encapsulated into an oxide layer of the secondary metal, limiting catalytic activity towards H$_2$O$_2$ destruction possibly by reducing the availability of low coordination Pd edge sites (Figure 6).

![Figure 6](image6.png)

**Figure 6.** The role of the secondary metal in the inhibition of H$_2$O$_2$ degradation over small Pd-rich nanoparticles as proposed by Freakley et al. (A) Proposed mechanism, where the secondary metal encapsulates small Pd nanoparticles as a result of successive heat treatments. (B and C) STEM-EELS mapping of model 5% Pd / SnO$_2$ catalyst showing partial encapsulation of Pd nanoparticles (red) by SnOx (green). Reproduced from ref. [128].

Building on this work Li et al.$^{129}$ have demonstrated that through Sn incorporation supported Pd catalysts exposed to rapid thermal treatment can deliver complete inhibition of H$_2$O$_2$ decomposition and hydrogenation, with H$_2$O$_2$ synthesis activities approaching twice that reported by Freakley et al., with minimal catalyst deactivation over re-use and no structural changes observed via TEM or EDX analysis.

Through a combination of DFT calculations and Sabatier analysis Gao and co-workers have proposed that several base metal dopants, such as W, and Pb, are able to enhance both catalytic activity and selectivity...
of Pd-based and bi-metallic Pd-Au catalysts towards H\textsubscript{2}O\textsubscript{2}. They propose that electronegative elements in particular are able to inhibit O-O bond cleavage through the withdrawal of electron density from Pd.\textsuperscript{130} However, further experimental evidence is required to confirm these findings.

Additional investigations by Abate et al. revealed that N doping carbon nanotubes results in increased catalytic activity and stability through increased dispersion of Pd and enhanced acidity of the support.\textsuperscript{131, 132} While Melada et al. report an enhancement in catalytic activity through the introduction of sulphate and halide dopants into zirconia supported Pd catalysts.\textsuperscript{65} Although the use of dopants can enhance catalytic activity and selectivity in the direct synthesis reaction the leaching and sites favorable for H\textsubscript{2}O\textsubscript{2} formation. Table 1 below highlights the enhancement in catalytic selectivity towards H\textsubscript{2}O\textsubscript{2} that can be achieved through modification of Pd by secondary non-precious metals.

### Choice of support for precious metal catalysts.

The nature of the support is a key factor which can affect catalytic activity and selectivity towards H\textsubscript{2}O\textsubscript{2}. A range of zeolitic,\textsuperscript{138-142} oxide, resin,\textsuperscript{62, 108, 143, 144} heteropolyacid\textsuperscript{145-148} and carbon\textsuperscript{134, 149, 150} supports have been investigated to date.

Hutchings and co-workers have extensively studied the use of oxide supports for the direct synthesis of H\textsubscript{2}O\textsubscript{2} over bi-metallic AuPd catalysts and correlated catalytic activity with the iso-electric point of the support, with those more acidic supports beneficial for catalytic selectivity and hence net yield of H\textsubscript{2}O\textsubscript{2}.\textsuperscript{151, 152} Further study by Menegazzo et al.\textsuperscript{153} has reported that it is possible to control Pd nanoparticle size and in turn balance catalytic activity and selectivity towards H\textsubscript{2}O\textsubscript{2} through the choice of support. They reported that the use of SiO\textsubscript{2} is superior to that of either ZrO\textsubscript{2} or CeO\textsubscript{2} in tuning these two parameters of catalytic efficiency.

Due to their high acidity numerous studies have investigated heteropolyacids as both catalyst supports\textsuperscript{145, 148, 154-156} as well as solid acid additives\textsuperscript{177} for the direct synthesis of H\textsubscript{2}O\textsubscript{2}. Problems concerning their low surface area and high solubility in polar solvents can be overcome through the introduction of

Table 1. Comparison of catalytic selectivity towards H\textsubscript{2}O\textsubscript{2} as a function of secondary metal.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Catalyst</th>
<th>Reactor system</th>
<th>Temp. / °C</th>
<th>Pressure / Bar</th>
<th>Time / h</th>
<th>Solvent</th>
<th>Promoter</th>
<th>H\textsubscript{2}O\textsubscript{2} selectivity / %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gu [134]</td>
<td>1% Pd / C</td>
<td>Batch</td>
<td>2</td>
<td>30</td>
<td>0.25</td>
<td>MeOH</td>
<td>0.03M H\textsubscript{2}SO\textsubscript{4}</td>
<td>54</td>
</tr>
<tr>
<td></td>
<td>Pd-Ag(10) / C\textsuperscript{a}</td>
<td>Batch</td>
<td>2</td>
<td>30</td>
<td>0.25</td>
<td>MeOH</td>
<td>0.03M H\textsubscript{2}SO\textsubscript{4}</td>
<td>72</td>
</tr>
<tr>
<td>Wang [133]</td>
<td>1% Pd / γ-Al\textsubscript{2}O\textsubscript{3}</td>
<td>Batch</td>
<td>2</td>
<td>30</td>
<td>0.25</td>
<td>MeOH</td>
<td>0.03M H\textsubscript{2}SO\textsubscript{4}</td>
<td>64</td>
</tr>
<tr>
<td></td>
<td>1% Pd-5% Zn / γ-Al\textsubscript{2}O\textsubscript{3}</td>
<td>Batch</td>
<td>2</td>
<td>30</td>
<td>0.25</td>
<td>MeOH</td>
<td>0.03M H\textsubscript{2}SO\textsubscript{4}</td>
<td>79</td>
</tr>
<tr>
<td>Freakley [128]</td>
<td>1% Pd – 4% Sn / TiO\textsubscript{2}</td>
<td>Batch</td>
<td>2</td>
<td>40</td>
<td>0.5</td>
<td>H\textsubscript{2}O / MeOH</td>
<td>-</td>
<td>95</td>
</tr>
<tr>
<td></td>
<td>3% Pd – 2% Sn / TiO\textsubscript{2}</td>
<td>Batch</td>
<td>2</td>
<td>40</td>
<td>0.5</td>
<td>H\textsubscript{2}O / MeOH</td>
<td>-</td>
<td>96</td>
</tr>
<tr>
<td>Maltby [137]</td>
<td>Pd (unsupported)</td>
<td>Sembatch</td>
<td>10</td>
<td>1</td>
<td>1</td>
<td>H\textsubscript{2}O</td>
<td>0.1 M HCl + 0.01M KBr</td>
<td>82</td>
</tr>
<tr>
<td></td>
<td>Ni-Pd-x (unsupported)\textsuperscript{b}</td>
<td>Sembatch</td>
<td>10</td>
<td>1</td>
<td>1</td>
<td>H\textsubscript{2}O</td>
<td>0.1 M HCl + 0.01M KBr</td>
<td>95</td>
</tr>
<tr>
<td>Ding [136]</td>
<td>3 % Pd / TiO\textsubscript{2}</td>
<td>Sembatch</td>
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<td>1</td>
<td>0.25</td>
<td>EtOH + H\textsubscript{2}SO\textsubscript{4}</td>
<td>0.12 M H\textsubscript{2}SO\textsubscript{4}</td>
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<td>0.12 M H\textsubscript{2}SO\textsubscript{4}</td>
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\textsuperscript{a} Pd loading fixed at 1 wt.% with value in parentheses indicating Pd / Ag molar ratio. \textsuperscript{b} Subscript values indicate nominal Ni / Pd weight ratio.

\textsuperscript{a} Pd loading fixed at 3 wt.% with subscript value indicating Pd / Sb molar ratio. \textsuperscript{b} Subscript values indicate atomic Pd / Te ratio.
specific cations, such as Cs⁺, K⁺ and Rb⁺ into the structure of the heteropolyacid, while other studies have investigated palladium exchanged heteropolyacids immobilized onto mesoporous silica for the synthesis of H₂O₂. Interestingly these catalysts have been shown to outperform the analogous catalysts utilising a more conventional, less acidic, support under reaction conditions likely to be more favored by industry; (i.e. ambient temperature and a water only solvent) likely due to the increased selectivity imparted by the acidity of the support.\(^{145}\)

Mesoporous silicas such as MCM-41\(^{138,157,158}\) and SBA-15\(^{159-161}\) have seen growing interest as supports for the direct synthesis of H₂O₂ due to their high surface area, which allows for a high dispersion of active sites, and an ordered channel structure which can alleviate issues associated with mass transfer. In addition the large pore size of mesoporous silicas allows for the majority of active metal to be accommodated within the mesoporous framework. This aids in preventing agglomeration of metal nanoparticles during the course of the reaction and balances the need for high nanoparticle dispersion, resulting in high activity, and minimal O₂ dissociation which is necessary for high catalytic selectivity.\(^{159}\) Both Park et al.\(^{162}\) and Rodríguez-Gómez et al.\(^{119}\) have investigated the functionalisation of SBA-15 through the grafting of organic functional groups, with the introduction of amine groups in particular resulting in a significant enhancement in catalytic selectivity through the suppression of H₂O₂ decomposition.

Application of H₂O₂ as an in situ oxidant.

Propylene oxidation to propylene oxide. The selective oxidation of propylene to propylene oxide yields one of the major starting materials in industrial chemistry, with propylene oxide finding application in the production of surfactants, polyurethane and resins. Until recently propylene oxide has been manufactured on an industrial scale through the use of chlorohydric acid or hydroperoxides.\(^{14}\) The first process reacts propene with Cl₂ to produce chlorohydric acid, which is dehydrochlorinated using an aqueous alkali solution, producing an equimolar quantity of aqueous alkali metal chlorides along with the required product, which incurs significant removal and treatment costs. The use of hydroperoxides is more environmentally friendly than the use of chlorohydric acid but is greatly dependent on the market value of the co-product obtained alongside propylene oxide (styrene or tert-butanol).\(^{163}\) In recent years a new alternative, the HPPO process, utilising H₂O₂ as an oxidant, has offered significant environmental and economic advantages over the alternative processes. An appealing alternative to the HPPO process involves the in situ production of H₂O₂, either in the liquid or vapour phase, over supported precious metals, notably Pd, Au, and Pt, or combinations thereof. A range of supports have been explored within the literature including; TS-1\(^{164-166}\) Ti-MCM-41\(^{170-172}\) and oxides\(^{173-175}\) where H₂O₂ is activated at mild temperatures by isolated Ti(IV) sites present within the support structure. However, catalytic selectivity towards propylene oxide is still a concern with the use of promoters common place. In particular Uphade et al. have reported the use of CsCl as a promoter for Au / Ti-MCM-41 catalysts, leading to an enhancement in Au particle size, with a resulting 5 % increase in selectivity towards propene oxide.\(^{170}\) While Chen and Beckman have reported it is possible to achieve a four-fold increase in selectivity towards propylene oxide over a low loaded PdPt / TS-1 catalyst, through the use of ammonium acetate. They report that even at low concentrations ammonium acetate is able to suppress unwanted side reactions, including the acid catalysed hydrolysis of propylene oxide to propylene glycol. The focus of future research must now be placed on achieving high selectivity towards propylene oxide in the absence of promoters.

The Fenton process. The Fenton process, the catalytic formation of hydroxyl and hydroperoxy radicals by the disproportionation of hydrogen peroxide, can be considered one of the most efficient means for the degradation of organic pollutants in wastewater streams at low to moderate concentrations.\(^{176-178}\) Numerous studies have investigated the efficacy of H₂O₂ generated in situ over Pd-Fe based catalysts utilising a range of hydrogen sources such as formic acid, hydroxylamine and hydrazine.\(^{179-183}\) Indeed Underhill et al.\(^{184}\) have recently reported that superior rates of phenol conversion can be achieved from H₂O₂ generated in situ from molecular H₂ and O₂, compared to preformed
H$_2$O$_2$, with this superiority ascribed to the absence of stabilising compounds, such as phosphoric acid and acetaldehyde, found in commercial H$_2$O$_2$.\[242\]

**Methane valorisation to methanol.**

The selective oxidation of methane, a major component of natural gas, to methanol is an attractive means to produce a versatile chemical feedstock. However, the direct catalytic upgrading of methane is yet to be achieved under environmentally friendly conditions on a scale that is industrially viable. Methane conversion to methanol on an industrial scale currently utilises harsh conditions, with temperatures exceeding 800 °C, to produce synthesis gas which can then be converted into methanol. This approach although highly selective is associated with high energy and capital demands. As such significant economic and environmental advantages may be reached through the selective oxidation of methane at milder reaction conditions. Initial work focussed on the use of supported Pt catalysts but required the use of concentrated H$_2$SO$_4$ as the oxidant.\[186\] However a significant breakthrough occurred with the use of environmentally benign oxidants such as H$_2$O$_2$.\[186, 187\] The application of in situ generated H$_2$O$_2$ overcomes the significant drawbacks associated with the anthraquinone process, the means by which H$_2$O$_2$ is generated on an industrial scale and Lin et al.\[186\] were amongst the first to investigate the in situ generation of H$_2$O$_2$ for the oxidation of methane, starting from O$_2$, CO in a trifluoroacetic acid / H$_2$O solvent system in the presence of Pd-based catalyst. Building on this work numerous studies have since investigated the conversion of methane to more versatile chemical feedstocks based on the O$_2$/CO/H$_2$O system,\[189, 190\] as well as the synthesis of H$_2$O$_2$ directly from H$_2$ and O$_2$, avoiding the initial water gas shift reaction to produce H$_2$ in situ.\[191, 192\] Recently Rahim et al. have demonstrated that significant improvements in selectivity towards methanol can be achieved through the use of H$_2$O$_2$ generated in situ when compared to the use of pre-formed H$_2$O$_2$, under comparatively mild reaction conditions.\[193\]

Despite demonstration of the feasibility of the selective oxidation of short chain alkanes via H$_2$O$_2$ generated in situ further work is required for this approach to be considered industrially viable, with low productivities and a need for a reduction in the formation of the products of over oxidation, namely CO$_2$, significant challenges for future research.

**Cyclohexane oxidation and the production of KA oil.**

Cyclohexanone and cyclohexanol are important chemical feedstocks for the production of ε-caprolactam and adipic acid, key intermediates for the production of Nylon-6 and Nylon-6,6. In addition, cyclohexanol and cyclohexanone are also used as stabilising agents and homogenizers for synthetic detergents and soaps. The production of cyclohexanone and cyclohexanol (KA oil) on an industrial scale currently relies on the uncatalysed oxidation of cyclohexane with air at elevated temperatures.\[194-198\] To limit over oxidation to unwanted by-products conversion is limited to below 10% and additional steps are implemented to improve cyclohexanone:cyclohexanol ratio.\[199\]

Numerous studies have investigated the use of H$_2$O$_2$ as a replacement oxidant for molecular O$_2$. Due to its greater reactivity H$_2$O$_2$ allows for significantly lower temperatures to be utilised, with the current auto-oxidation process requiring temperatures in excess of 140 °C.\[200\] A range of supports have been explored including vanadium phosphorous oxide, which demonstrated near complete conversions of the substrate, however the catalyst displayed poor selectivity towards KA oil due to a high formation of cyclohexyl peroxide.\[201\] Spinace et al. was amongst the first to investigate the activity of TS-1 for the oxidation of cyclohexanone.\[202\] However, further work by the same group reported the ability of TS-1 to catalyse the further oxidation of cyclohexanol to unwanted by-products, which can lead to the deactivation of the catalyst through the blocking of the TS-1 pore system.\[200\] Expanding on this initial work Shi et al. developed a titanium silicate with a hollow structure (HTS) and enhanced Lewis acidity that offered higher activity towards KA oil compared to that observed for TS-1 alone.\[200\] Rezaei et al. have reported a reusable KIT-6 supported vanadium pyrophosphate catalyst for cyclohexane at low temperatures.\[204\] However the reported selectivity is still not sufficient for industrial application, even at low conversion rates.

The in situ generation of H$_2$O$_2$ for the oxidation of cyclohexanone has recently been studied by Kuznetsova and co-workers utilising Pt based catalysts. It is reported that the addition of small quantities of ionic
liquids containing Br- and HSO₄⁻ anions are able to stabilise the peroxide-intermediates, resulting in an enhanced yields of KA oil. A further study by Li et al. revealed the catalytic activity of zeolite-Y supported AuPd catalysts for the oxidation of cyclohexane, with only cyclohexanol observed as the desired product, however this is likely due to the relatively short reaction times investigated.

**Conclusion and Future Perspectives.**

The direct synthesis of H₂O₂ from H₂ and O₂ offers potential significant environmental and economic benefits over the current means of H₂O₂ production, i.e. the anthraquinone process. However, despite extensive research for over 100 years the direct synthesis of H₂O₂ from H₂ and O₂ is still not industrially viable, although research into this challenging reaction has gained momentum in the past decade.

Although Pd based catalysts have received the greatest attention perhaps one of the biggest breakthroughs in the field came from the finding that incorporating Au into supported Pd catalysts can dramatically enhance catalytic selectivity, without the need for acid or halide additives. Further investigations have since shown that the incorporation of several other secondary metals, including Pt and Ni into supported Pd catalysts can greatly enhance catalytic selectivity and activity. While the use of high surface and acidic supports have been shown to inhibit the agglomeration of metal nanoparticles and enhance H₂O₂ selectivity, respectively.

Numerous computational studies have led to a greater understanding of the active sites responsible for the direct synthesis of H₂O₂, its degradation and the competitive formation of H₂O as well as aiding experimentalists in the design of new catalysts. However, further study is required to improve catalyst activity for use in a semi-continuous / continuous regime, where contact time between H₂, O₂ and the catalyst is significantly shorter than that utilised in batch reactors. This is likely to be required if the direct synthesis route is to be utilised on an industrial scale. In addition further research is required to gain a better understanding of catalyst lifetime and how catalytic structure and morphology changes on-stream.

Selective oxidation using H₂O₂ generated in situ from H₂ and O₂ offers an attractive alternative to current processes, removing the need for the storage of large quantities of H₂O₂ on site prior to use as well as allowing for the use of milder reaction conditions, in particular high temperatures associated with auto-oxidation reactions, with obvious environmental and economic benefits. Furthermore the lack of stabilising agents, such as acids can promote catalyst and reactor lifetime, which are concerns that arise when pre-formed H₂O₂ is utilised. Often oxidative catalytic systems have utilised a two-step process; the production of H₂O₂, usually over noble metals, followed by its activation and use in the oxidation of the target substrate. This has often required the diffusion of H₂O₂ from catalytic species responsible for its generation to a secondary, sometimes homogeneous component, with the Fenton reaction a well-studied example. However, often reaction conditions utilised are unfavourable towards H₂O₂, resulting in the conversion of H₂O₂ to H₂O. To mitigate this, future catalytic development should, where feasible, centre on single active sites responsible for both H₂O₂ production and activation. Where consolidation into one active site is not possible, research should focus on means to avoid diffusion limitations and the resulting degradation of H₂O₂. The higher activity observed for H₂O₂ generated in situ compared for both the selective oxidation of methane as well as the degradation of waste stream contaminants is particularly exciting and highlights the potential benefits that can be unlocked through the production of an in situ oxidant.

In conclusion the growing global demand for commodity and fine chemicals, coupled with the environmental need to replace chloride-based bleaching agents particularly in the treatment of waste streams, will lead to an increasing need for H₂O₂ over the coming decades.

It is suggested that as it has been demonstrated that H₂O₂ can be synthesised with 100 % selectivity based on H₂ focus should now be placed on increasing catalytic activity towards H₂O₂ production. At present the highest concentration of H₂O₂ reported is approximately 1 wt.% however, to be competitive with current means of H₂O₂ production significantly greater concentrations are required. It is hoped that with
growing interest from the scientific community and increased collaboration between theoreticians, experimentalists and chemical engineers this target will soon be met by a commercialised direct synthesis process.

References.
