**Electrochemical synthesis of hydrogen peroxide from water and oxygen**

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# Abstract

H2O2 is important in large-scale industrial processes and smaller on-site activities. The present industrial route to H2O2 involves hydrogenation of an anthraquinone and O2 oxidation of the resulting dihydroanthraquinone - a costly method and one that is impractical for routine on-site use. Electrosynthesis of H2O2 is cost-effective and applicable on both large and small scales. This Review describes methods to design and assess electrode materials for H2O2 electrosynthesis. H2O2 can be prepared by oxidizing H2O at efficient anodic catalysts such as those based on BiVO4. Alternatively, H2O2 forms by partially reducing O2 at cathodes featuring either noble metal alloys or doped carbon. In addition to the catalyst materials used, one must also consider the form and geometry of the electrodes and the type of reactor in order to strike a balance between properties such as mass transport and electroactive area, both of which substantially affect both the selectivity and rate of reaction. Research into catalyst materials and reactor designs is arguably quite mature, such that the future of H2O2 electrosynthesis will instead depend on the design of complete and efficient electrosynthesis systems, in which the complementary properties of the catalysts and the reactor lead to optimal selectivity and overall yield.

# Introduction

H2O2 is a strong oxidant with many uses both in a domestic setting as well as in industrial bleaching,1 waste water treatment,2-5 chemical synthesis6-10 and fuel cell technologies.11, 12 These applications see H2O2 produced on a scale of approximately 2.2 million tonnes per year.1 a value that can be expected to increase given that H2O2 is a more environmentally friendly oxidant than Cl-based oxidants such as HOCl. At present, 95% of H2O2 is produced through the anthraquinone process,13 which begins with the hydrogenation of a 2-alkyl-9,10-anthraquinone in an organic solvent over a Pd catalyst.14 In solution, the resulting dihydroanthraquinone then undergoes rapid oxidation by O2 to give H2O2 and regenerate the anthraquinone. The H2O2 is removed by solvent extraction, leaving anthraquinone in the organic phase to be recycled.15 This batch process typically affords large volumes of highly concentrated aqueous H2O2, which poses safety risks in terms of stability, storage and transport.16 Aside from the specialty aerospace industry,17 few sectors require H2O2 in a concentrated form, such that the risks associated with its preparation are unjustified. Moreover, the regeneration of the anthraquinone is not 100% efficient, but participates in a number of side reactions. The dihydroanthraquinone can be further hydrogenated to give more saturated products including tetra- or octahydroanthraquinone, and carbonyl groups can undergo hydrogenolytic cleavage to give 2ethylanthrone, which also participates in further reactions to give 2-ethyleneanthracene.18 Highly oxidizing conditions can also oxidize the dihydroanthraquinone to give the corresponding epoxide.19 The anthraquinone therefore requires constant replenishment to maintain a satisfactory rate of H2O2 production.13 With these shortcomings in mind, there is a push to develop a more energy-efficient and resource-efficient route that is amenable to on-site production of H2O2 in dilute working concentrations.20

A popular alternative to the anthraquinone process is the direct synthesis of H2O2 from H2 and O2. This partial hydrogenation is normally performed at low temperatures by passing gas mixtures over a noble metal catalyst such as Pd.21 These catalysts also mediate competing side reactions, including the combustion of H2 to H2O, the hydrogenation of H2O2 to H2O and the decomposition of H2O2 to H2O and ½O2.22 Perhaps the most substantial barrier to the development of this synthetic route is the need to work with mixtures of H2 and O2, which are explosive over a wide composition range (4–94 mol%H2).23 This danger means that the reactants are diluted with a relatively inert carrier gas such as CO2. However, this dilution and the inherent thermodynamic favourability of the side reactions reduce the practically achievable overall yields of the direct synthesis.

H2O2 is produced in nature through a range of enzymatic processes. Oxidase enzymes such as glucose oxidase,24 ᴅ-amino acid oxidase25 and cholesterol oxidase,26 produce H2O2 when presented with their native substrates. The limited turnover frequencies of these systems has limited their utility for H2O2 synthesis, although these enzymes are used in biosensing applications, in which quantifying an enzyme substrate can be performed indirectly by measuring how much H2O2 it affords.27, 28 Other possible applications include food sanitation, in which H2O2 generated by glucose oxidase could be used to protect food against microorganisms.29

Electrochemistry offers an economical and environmentally friendly route to H2O2 either from H2O or O2. Indeed, H2O2 can be produced at either an anode or cathode surface and can accumulate in useful concentrations over continued electrolysis. The first published example of H2O2 electrosynthesis came from Traube in 1887, who produced H2O2 from O2 at a Hg-Au electrode.30 At the time, this method was less efficient than chemical syntheses of H2O2, which included O2 oxidation of hydrazobenzenes,31, 32 hydrolysis of persulfates33 (which, in turn, are prepared by electrolysis of bisulfates in H2SO4) or direct synthesis.34 The first H2O2 synthesis plant was built by Dupont in 1953 and used the anthraquinone process.33 Although chemical syntheses have dominated industrial H2O2 production, since Traube’s report electrosyntheses have appeared continually in the literature, including methods with improved yields35 and anodic electrosynthesis.36-38 Indeed, understanding the kinetics of the redox reactions involving O2, H2O2 and H2O39-41 enabled rational designs of electrodes with high current densities35, 42 and/or selectivities41, 43 have resulted in the development of increasingly efficient catalysts. The major developments in this field are summarised in Figure 1.

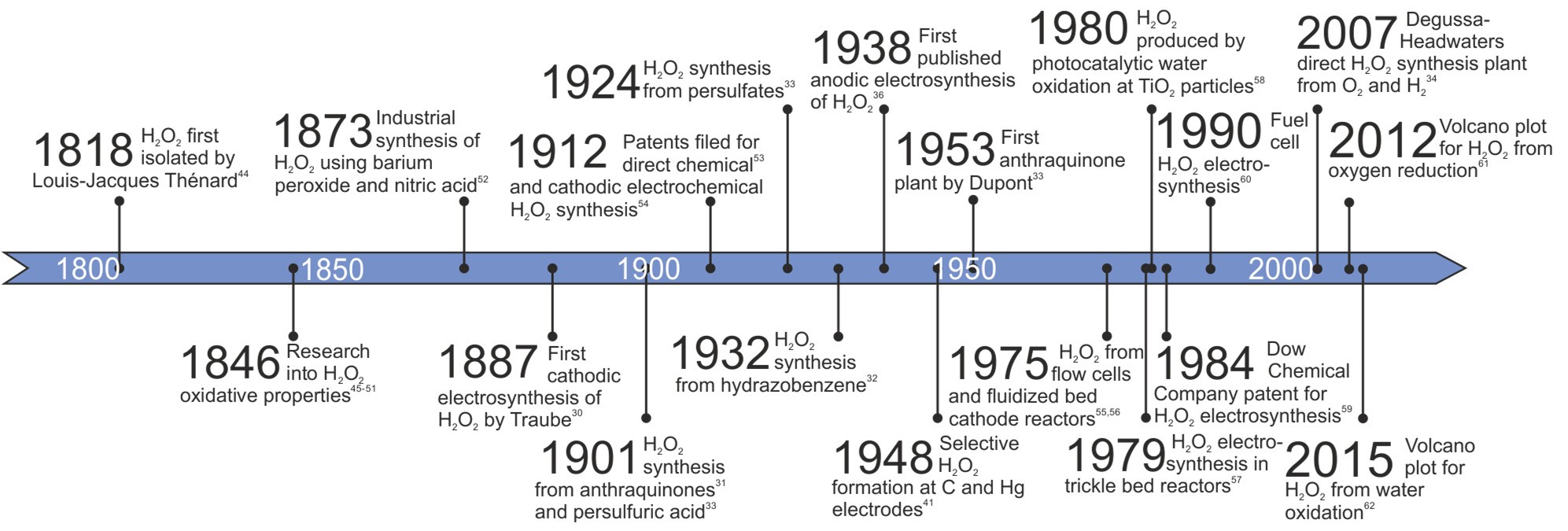


Figure 1 | **The major developments in the chemical and electrochemical synthesis of H2O2.**

Nowadays, laboratory H2O2 electrosyntheses are typically carried out in reactor volumes of hundreds of milliliters. Scaling these experiments up to industrially relevant volumes whilst preserving the same principles is likely possible given the number of reports that describe pilot plants operating on an intermediate scale.63-65 Alternatively, it is possible to produce H2O2 in situ for immediate consumption in oxidation reactions, such as the epoxidation of alkenes,66, 67 or the synthesis of peracetic acid68 or benzamide.69 Such in situ methods have distinct advantages over other synthetic routes, and in the case of epoxidation, for example, the local concentrations of H2O2 at the electrode are high enough to drive the reaction without the need for H2O2 to be present in high concentrations throughout the reactor.

As we mentioned above, the development of H2O2 electrosynthesis catalysts is complicated because many common electrode materials favour competing reactions. Thus, the 4e− reduction of O2 (the oxygen reduction reaction, ORR) or 4e− oxidation of 2H2O (the oxygen evolution reaction, OER) is undesirable because we require 2e− reduction of O2 or 2e− oxidation of 2H2O to get H2O2. The electrosynthesis of H2O2 is a relatively unusual process because it involves reversible redox reactions of starting materials, intermediates and products. This makes it more challenging than processes that focus on one reaction to give a stable product.

A lot of research on electrocatalytic O2 reduction and H2O oxidation is motivated by renewable energy applications, for which complete 4e− reduction of O2 is desirable because it affords the maximum power output. Thus, electrodes that produce H2O2 are considered flawed for fuel cell applications but are desirable in that they afford an industrially useful chemical in a green synthesis, particularly in the context of paired electrochemical processes. Most experimental research focuses on catalyst development, but the combination of experiment and theory enables more rational studies geared towards H2O2 formation whilst suppressing its redox reactions that afford either H2O or O2. A number of efficient electrosynthesis cell designs already exist for other reactions, but the key requirement particular to our reactions of interest is the separation of anode and cathode to attenuate the decomposition of H2O2 after its electrosynthesis.

This Review is an overview of the recently disclosed electrocatalysts that are selective for H2O2 as the end product, as well as the reactor designs that can best exploit these materials to generate the maximum possible H2O2 output. We describe the key advances required in H2O2 electrosynthesis to see it progress from laboratory-scale operations to larger industrial-scale applications.

# Thermodynamics of H2O2 production

The 2e− oxidation of 2H2O can proceed electrochemically to give H2O2:70

2H2O ⇌ H2O2 + 2H+ + 2*e*− *E*0 = 1.760 V *vs.* SHE (1)

Complications arise because the desired product H2O2 can undergo further oxidation:

H2O2 ⇌ O2 + 2H+ + 2*e*− *E*0 = 0.670 V *vs.* SHE (2)

The standard potential for the formation of H2O2 is more positive than that for the further oxidation of H2O2 to O2, so any H2O2 that forms may be easily oxidized, lowering the yield. Many electrochemical cells are optimized for the OER, either via H2O2 or by direct 4e− oxidation of 2H2O:

2H2O ⇌ O2 + 4H+ + 4*e*−  *E*0 = 1.229 V *vs.* SHE (3)

Thermodynamics also complicate the selective cathodic electrosynthesis of H2O2 because the reduction of O2 has a more negative standard potential than does H2O2, meaning that further reduction of H2O2 is favoured, lowering the yield of H2O2. Another deleterious reaction is the spontaneous disproportionation of H2O2 to give H2O and ½O2.71

2H2O2 ⇌ O2 + 2H2O (4)

In addition to the 2e− reductions and oxidations we have described, a number of 1e− redox reactions involving oxygenic species have been proposed to be operative:72-74

O2 + *e*− **→** O2**•**− *E*0 = 0.330 V *vs*. SHE (5a)

|  |  |  |
| --- | --- | --- |
| O2 + 3H+ + 3*e*− **→** HO• + H2O | *E*0 = 0.803 V *vs*. SHE | (5b) |
| H2O2 + *e*− **→** HO• + OH− | *E*0 = 0.104 V *vs.* SHE | (5c) |

The oxygenic radicals can contribute to the formation of H2O2, either by further electron transfers or through solution reactions following the superoxide dismutation:75

|  |  |  |
| --- | --- | --- |
| O2**•**− + H+ ⇌ HO2**•** |  | (6a) |
| O2**•**−+ HO2• **→** HO2− + O2 |  | (6b) |
| HO2• + *e*− ⇌ HO2− | *E*0 = -0.744 V *vs*. SHE | (6c) |
| HO2− + H+ ⇌ H2O2 |  | (6d) |

H2O2 may also undergo homolysis to HO• radicals,76 which are strongly oxidizing.72

H2O2 **→** 2OH**•** (7a)

HO• + H+ + *e*− ⇌ H2O *E*0 = 2.330 V *vs*. SHE (7b)

Systems for the preferential formation of H2O2 should favour initial redox to give H2O2, whilst also allowing for rapid diffusion of H2O2 away from the electrode before it can undergo (deleterious) further oxidation or reduction.

# Energy and bonding

We have described the reversible redox reactions between H2O, H2O2 and O2, but we stress that the interplay of these reactions is highly surface dependent. For O2 reduction, the rate determining step is as yet unknown even at relatively simple materials like Pt.77 Experimentally, Tafel analysis indicates that it is the first electron transfer that is rate determining.78 This does not reveal whether the electron transfer is coupled to O2 protonation or dissociative adsorption,79 although it has been proposed that the electron transfer precedes dissociation because the activation energy for the dissociation of adsorbed OOH is much smaller than for O2.80 Computational studies have also proposed other rate determining steps such as breaking the O−O bond81 or the reduction of PtO82 due to the large calculated activation energies for these reaction steps. This presents the possibility of different rate determining steps for different crystal faces, due to their vaied adsorption energies towards ORR intermediates.83 A consistent observation is that interaction between O2 and the electrode surface is important in predicting the efficiency of a catalyst for H2O2 production, hence the sizeable impact of crystal face and pH on the reaction rate.84 Most metal surfaces have a strong affinity for O2, such that the (electro)adsorption of O2 precedes e− transfer.79 One exception to this is Au, which first performs 1e− reduction on O2 through an outer-sphere mechanism.85 Regardless of which process occurs first, if we consider the complete 4e− reduction at a metal M there will be multiple oxygenic intermediates, each of which will have different M–O interactions.

There are a number of different ways in which O2 can adsorb onto a metal surface. O2 can bind sideon to a single metal centre in an M(η2-O2) arrangement referred to as the Griffith model.86 This model, by analogy with the Dewar–Chatt–Duncanson model for olefin binding, involves donation from the O2 2π orbital to the empty M-centred d*z*2 orbital, with synergistic backbonding from partially filled d*xy* or d*yz* M-centred orbitals into the 2π\* orbital of O2.87 Alternatively, O2 can bridge two metal sites in a M(μ,η2-O2)M motif called the bridge model .88 The strong σ interactions lead to strong π backbonding that weakens the O–O bond and promotes dissociative adsorption. In both these binding modes both O atoms are bound to a metal site, which hinders any H2O2 intermediates from leaving the electrode before they can undergo reduction as part of the complete 4e− ORR.80, 89 In contrast, the M(η1-O2) binding mode, known as the Pauling model features donation from an O-centred sp2 hybrid orbital into the M-centered dz2 orbital. This alone cannot lead to O2 dissociation and thus favours 2e− redox with H2O2 as the end product.87

In addition to the orientation of binding, the strength of the interactions also influences the products of the redox reactions. Stronger M–O bonds arise when M donates substantial electron density to the adsorbed O2 molecule. In such a case, the activation energies of the O-centred reductions that sequentially convert O2 to O2•−, H2O2 and then OH• are low. However, then the activation enthalpy associated with the final step — reduction of adsorbed OH• — becomes high89 and the electrode surface can become poisoned by the highly stable M–OH groups. Thus, in this strong bonding case, catalytic turnover can be slow and will favour H2O instead of H2O2 as the final product.

A number of studies have described tuning the binding energy between oxygenic species and metal surfaces (Δ*G*O) in order to rationally design catalysts to be selective for the ORR, OER or H2O2 synthesis.61, 90-94 As we have seen for the ORR, the highest catalytic activity occurs when Δ*G*O is an intermediate value — binding is strong enough to favour the initial adsorption, but weak enough to release OH• to prevent surface poisoning and complete the catalytic cycle. It is often thought that the optimal ORR catalyst can be arrived at by finding the optimal Δ*G*O value, which is sensitive to the identity of metal90, 94/alloy91-93 and facet.61, 84 This approach has led to the design of new ORR catalysts based on a number of Pt and Pd alloys,95-104 and OER perovskite catalysts such as Ba0.5Sr0.5Co0.8Fe0.2O3–δ.105, 106

When targeting the selective synthesis of H2O2 from either O2 or H2O we want a cathode or anode that binds H2O2 weakly, such that the product can dissociate from the surface instead of undergoing further 2e− reduction or oxidation as part of a complete ORR or OER. The binding energies of the superoxo intermediate (Δ*G*HOO\* for M + OOH → MOOH)107, 108 and of the hydroxo intermediate (Δ*G*HO\* for M + OH → MOH)61, 109-111 have been identified as useful descriptors for catalysts to favour H2O2 formation from O2 or H2O, respectively (FIG. 2). Alloying Hg with metals commonly used for the ORR has a large effect on Δ*G*HOO\*, giving a volcano-type relationship between Δ*G*HOO\* and the resulting activity for H2O2 formation, with Pd-Hg alloys outperforming other sampled alloys.107 This example also allows us to illustrate that Δ*G*HOO\* scales linearly with Δ*G*HO\*, such that the values can be used interchangeably.109 The scaling relationship between these descriptors can be detrimental because it can be difficult to change one without changing the other.112 For the ORR, catalyst turnover rate could be increased by reducing Δ*G*HO\* to prevent surface poisoning by the hydroxo intermediate, but this would need to be done without reducing Δ*G*HOO\*, otherwise the superoxo intermediate could be released before the ORR reaches completion. Fundamentally, the difference between the free energies of HO\* and HOO\* should be 2.46 eV (twice the equilibrium potential for the ORR) but the scaling constrains the difference to ≈ 3.3 eV, which gives limits the maximum efficiency of any catalyst towards the ORR.113 Countering the scaling between Δ*G*HOO\* and Δ*G*HO\* has already been the subject of works to develop new ORR catalysts, by modifying catalyst covalence114 or using a tandem bi-functional catalyst where one component is optimised towards Δ*G*HO\* and the other towards Δ*G*HOO\*.115

We have mentioned the large body of ORR and OER research, and noted a recent push to also design catalysts for the 2e− reduction of O2 to H2O2. There has also been attention paid to the rational design of catalysts for the 2e− oxidation of 2H2O to H2O2. As with the ORR, the bulk of the OER literature has focused on the complete 4e− pathway, in this case giving O2.116-128 Catalysts now also exist for the selective anodic synthesis of H2O2. These catalysts are often transition metal oxides, and include materials that are free of noble metals. Density functional theory (DFT) calculations on the adsorption of OER reaction intermediates revealed that Δ*G*HO\* and Δ*G*O\* are useful descriptors to predict the suitability of a material for anodic H2O2 electrosynthesis.62, 129, 130

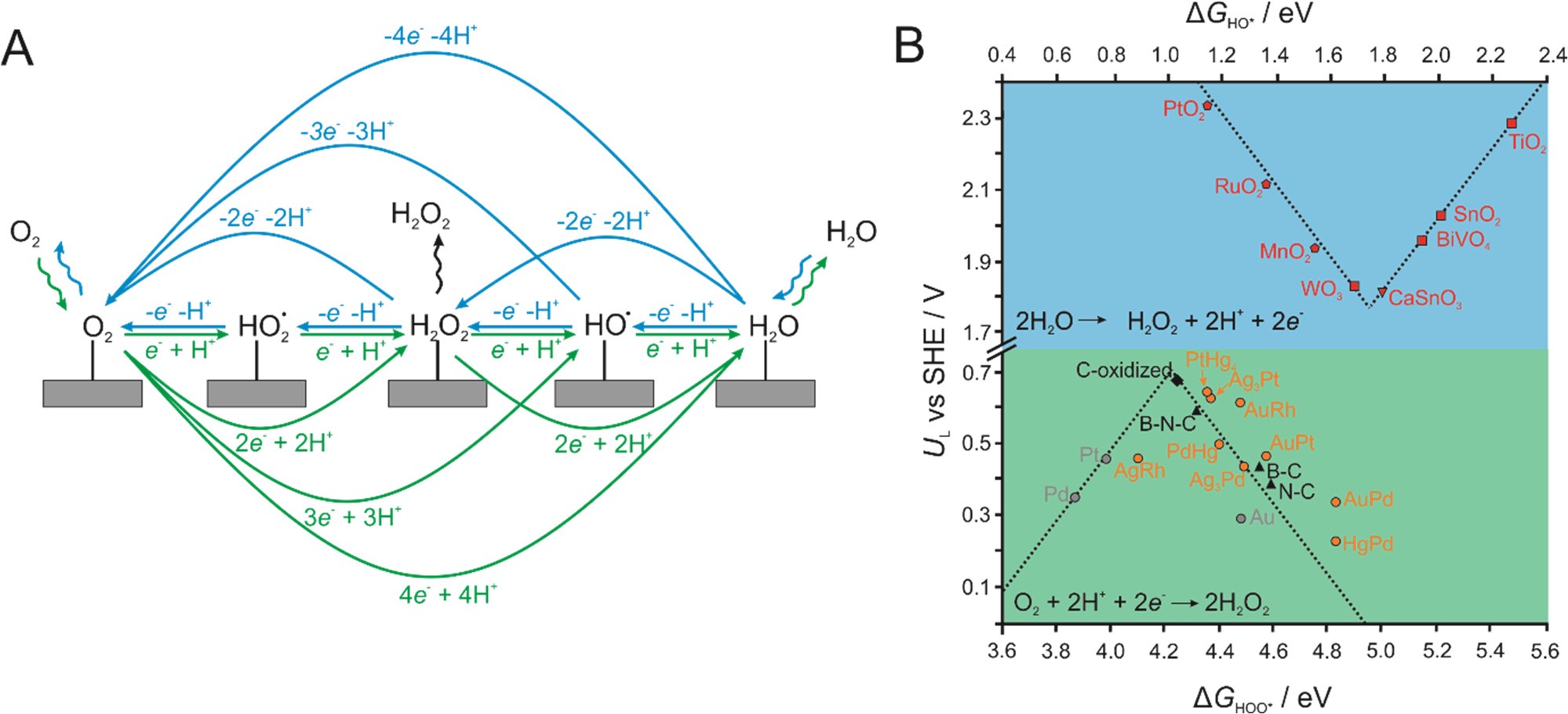


Figure 2. | **Interactions between intermediates and the catalyst surface strongly affect rates of H2O2 electrosynthesis. a |** The possible reduction pathways from O2 to H2O (green) and oxidation pathways from H2O to O2 (blue). The grey arrows denote mass transport of species between the bulk solution and the catalyst. Stoichiometric H2O is omitted for clarity. **b** | A plot of theoretical limiting potential (*U*L) against Gibbs free energies of binding \*OOH (Δ*G*HOO\*) and \*OH (Δ*G*HO\*) for H2O2 electrosynthesis. *U*L is the least positive (anodic) or negative (cathodic) potential at which both electron transfers are downhill in free energy (the potential where the free energy surface becomes completely exergonic). The green region represents O2 reduction and the blue region H2O oxidation. Catalysts are categorized as pure metals (grey), metal alloys (yellow), carbon-based materials (black) and metal oxides (red). The dashed lines represent the theoretical Sabatier volcanoes,61 the peaks of which corresponds to parameters of theoretical optimum catalysts. Part **b** was drawn from data in REFs 129 (■), 109 (●), 131 (▲) 108 (◆) 132 (▼) and 62 (⬟). For many materials, the exact *U*L depends on the surface site (step, edge, basal plane etc.) so the most favourable *U*L is shown in each case.

# Materials

The first published example of H2O2 electrosynthesis involved using a Hg-Au electrode to reduce O2 and afford only a few milligrams of H2O2.30 A major advancement came with instead using highly porous activated carbon electrodes, which greatly increased the electrode surface area to increase the overall rate of production.35 These porous carbon cathodes formed the substrate for subsequent electrocatalysts for many years.133, 134 As the field advanced, the electrode materials became more complex, moving towards doped carbons, noble metal alloys and metals oxides. The development of computational methods to predict the activity of materials towards H2O2 electrosynthesis drove further improvements in catalyst materials, as experimental papers worked towards the peak of the Sabatier volcano plot predicted from DFT calculations.107, 129

For all materials, there is an inverse relationship between the current density and the selectivity towards H2O2. A number of materials offer excellent selectivity towards H2O2 when working at small overpotentials and low total rates but then become poorly selective when working at higher total rates. Even the simplest of materials can have excellent selectivity for H2O2 formation provided the reaction is driven at low overpotentials. However, catalysts for real applications will need to maintain their selectivity even when working at current densities in the order of hundreds of milliamperes per square centimetre. When comparing materials, it is vital to consider the overpotential at which a given selectivity for H2O2 is reported, and one should determine the selectivity of a material over a range of overpotentials. Some of the leading materials for H2O2 electrosynthesis are given in Table 1, along with the conditions used to measure their selectivities. We now describe some of these materials and highlight mechanistic origins of their high selectivities.

Table 1 | **Electrode materials for the electrosynthesis of H2O2.** The methods and apparatus used to assess these methods include a gas diffusion electrode (GDE), rotating ring disc electrode (RRDE), chronoamperometry (CA), linear sweep voltammetry (LSV) and photovoltaic cell (PVC). Where multiple compositions of doped or alloyed materials were studied, the best recorded value is shown.

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Material** | **Faradaic**  **Efficiency**  **/ %** | ***n*** | **Anodic / Cathodic** | **Method** | **Solution** | **pH** | ***E* *vs.***  **RHE / V** | ***j* / mA**  **cm-2** | **Ref** |
| **Carbon** | | |  |  |  |  |  |  |  |
| Carbon nanotubes | 90a | 1.8 | Cathodic | RRDE | 0.1 M KOH | 13b | - | - | 135 |
| Mesoporous carbon | 95 | 2.1a | Cathodic | GDE | 0.5 M NaOH | 13.7b | 0.43 | -150 | 136 |
| **Doped carbon** | | |  |  |  |  |  |  |  |
| N-C  (mesoporous) | 95 | 2.1 | Cathodic | RRDE | 0.1 M H2SO4 | 0.3 | 0.10 | -1.25c | 137 |
| Boron doped diamond | 53 | 2.9a | Cathodic | CA | 0.5 M H2SO4 | 1b | -1.74c | - | 138 |
| **Carbon macrocycles** | | |  |  |  |  |  |  |  |
| Co(II) phthalocyanine | 81.5 | 2.3 | Cathodic | GDE | 0.1 M K2SO4 | 2b | 0.23 | -0.02c | 139 |
| Fe(II) phthalocyanine | 78.2 | 2.4 | Cathodic | GDE | 0.1 M K2SO4 | 2b | -0.18 | -0.01c | 140 |
| **Single atom catalyst** | | |  |  |  |  |  |  |  |
| Pt/S doped carbon | 96 | 2.1 | Cathodic | RRDE | 0.1 M HClO4 | 1.3b | 0.1c | -0.05c | 141 |
| **Metal Alloy Nanoparticles / Carbon** | | |  |  |  |  |  |  |  |
| Au0.92Pd0.08/carbo n black | 95 | 2.1a | Cathodic | RRDE | 0.1 M HClO4 | 1b | 0.0 | -0.8c | 142 |
| Sn6Ni/carbon black | 88 | 2.2 | Cathodic | RRDE | 1 M NaOH | 14b | 0.7d | -0.34d | 143 |
| **Metal Oxide Nanoparticles / Carbon** | | |  |  |  |  |  |  |  |
| CeO2/carbon black | 95 | 2.1a | Cathodic | RRDE | 1 M NaOH | 14b | 0.84 | -0.15c | 144 |
| WO3/Vulcan carbon | 84 | 2.3 | Cathodic | GDE | 0.1 M K2SO4 | 0.3b | -1.08 | - | 145 |
| **Metal oxides** | | |  |  |  |  |  |  |  |
| BiVO4 | 95 | 2.1a | Anodic | LSV | 1 M NaHCO3 | 8.3 | 2.8c | 15c | 129 |
| CaSnO3 | 76 | 2.5a | Anodic | CA | 2 M KHCO3 | 8.3 | 3.2 | 34c | 132 |
| **Mixed metal oxides** | | |  |  |  |  |  |  |  |
| BiVO4/WO3/  Al2O3 | 80 | 2.4a | Photooxidative | PVC | 2 M KHCO3 | 7.9 | - | 1 | 146 |
| IrO2/Ta2O5 | 79.3 | 2.4a | Cathodic | CA | 0.1 M  Na2SO4 | 7 | - | 0.001 | 147 |

a Value calculated assuming efficiency = (2 – *n*/2) × 100%, where *n* is the number of electrons.148 b pH was not explicitly reported but has been approximated here based on the electrolyte used. c Demonstrative potentials and current densities were read off steady state voltammograms. d Demonstrative potentials and current densities determined from a Koutecký–Levich plot.

## Pure metals

It has become common to analyze materials for their H2O2 electrosynthesis activity because applications in fuel cells demand ORR catalysts that produce a minimum of H2O2 even when working at low overpotentials. In practice, any H2O2 that forms can be detected either at the ring of a rotating ring–disc electrode (RRDE),149-153 by scanning electrochemical microscopy (SECM)154-158 or inferred from a decrease in the apparent number of electrons transferred (*n*) per O2 substrate from the ideal ORR value of 4.91, 159-164 Thus, ORR (or OER) catalysts that effect incomplete reduction or oxidation are less useful for fuel cells but may be of great use for H2O2 electrosynthesis, in which failed ORR or OER catalysts may be suitable for H2O2 production. The selectivity towards H2O2 formation at pure metals depends strongly on the applied potential. In the case of the ORR, SECM shows that Hg, Au, Ag, Cu, Pt and Pd all generate some H2O2 at low overpotentials. However, only Hg maintains high selectivity for H2O2 at all potentials studied (FIG. 3).154

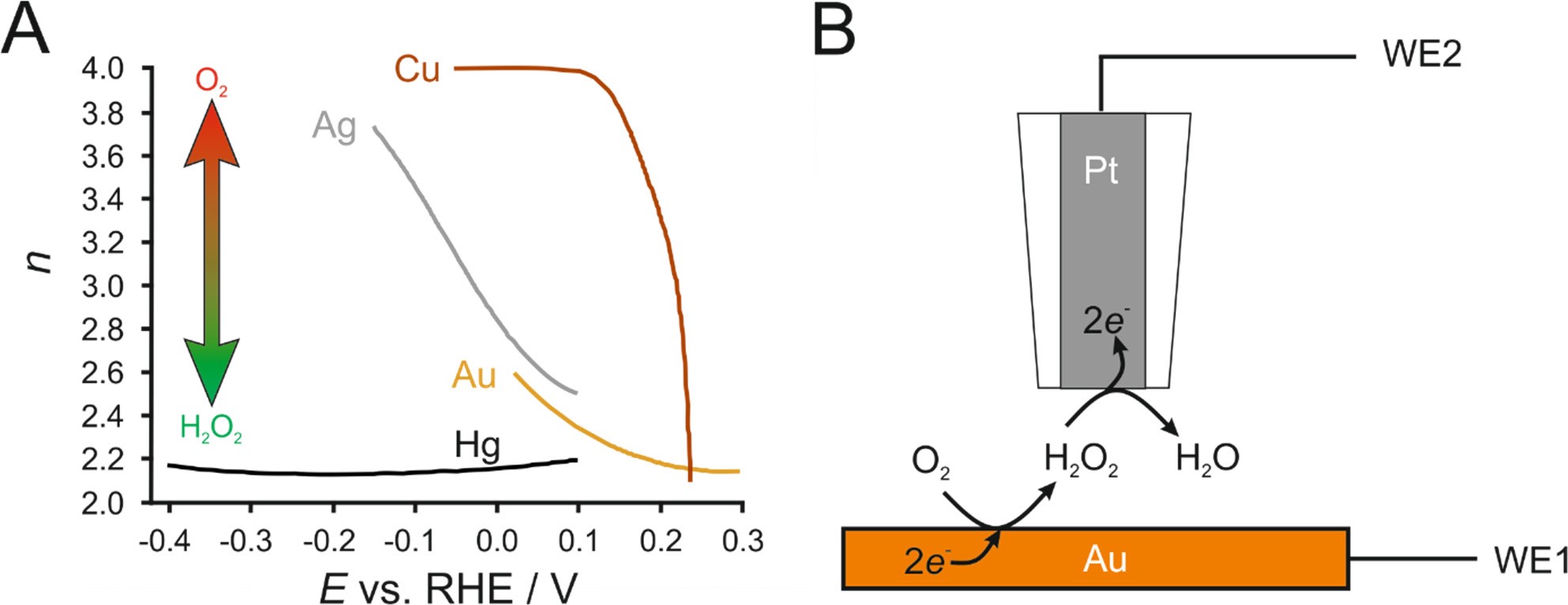


Figure 3. | **O2 reduction to H2O2 at metal electrodes. a |** The number of electrons transferred (*n*) to O2 at Hg, Au, Ag and Cu as a function of applied potential, with data recorded in H2SO4 (0.5 M aqueous) electrolyte. **b** | The electrode setup used to generate H2O2 and quantify its formation. Working electrode 1 (WE1) is 2 mm in diameter and is biased at potential *E* to generate H2O2. Located 50 μm away from WE1 is working electrode 2 (WE2), a 25 μm diameter microelectrode biased sufficiently positively to oxidize H2O2, thereby detecting it in the form of an anodic current. The ratio of currents at WE1 and WE2 are used to calculate *n*. Part **a** was drawn from data in REF 154.

Although it is encouraging that all materials can produce H2O2 when poised at a certain overpotential, the requirement of a low overpotential can result in inherently low current densities. Alternatively, a given material can be tuned to favour H2O2 production by increasing the rate of mass transport to/from the electrode surface, in which case H2O2 diffuses rapidly from the electrode before it can undergo further reactions. Increasing the rate of mass transport has been achieved using microelectrodes159, 160 rotating disc electrodes,159, 163 nanoparticles164, 165 and flow cells,166 by varying physical parameters such as electrode radius, catalyst loading, particle shape/size, rotation rate or flow rate. The effect of using small nanoparticles is not just a physical one — smaller particles favour the end-on O2 binding mode that can selectively afford H2O2. Indeed, the size of Pt clusters on the inert support indium tin oxide (ITO) is inversely proportional to the selectivity with which they electrogenerate H2O2 from O2.167 Although the increase in H2O2 production under conditions of high mass transport is substantial, the materials do not give 100% H2O2 selectivity (*n* = 2), so most works focus on more practical solutions such as alloys and composites.

## Metal alloys

Some of the most successful materials for H2O2 electrosynthesis by oxygen reduction are based on metal alloys, such as Pd-Au,16, 168 Pt-Pd,169 W-Au,170 or Pt-Hg.107, 109 As we described above, these catalysts are often designed through calculations that predict the optimal strength of O2 adsorption.90, 129 Computational studies have correctly predicted the performance of evenly dispersed107, 109 and core-shell alloy96, 170, 171 catalysts. The two metals that are alloyed do not both have to be individually active for H2O2 electrosynthesis. Indeed, alloying an active metal with another metal that is relatively inactive at the potentials applied can give discrete reactive sites surrounded by a relatively inert material.172 For example, in PdAu alloys it is the Pd sites that perform initial 2e− transfer to O2; Au binds O2 only weakly and cannot alone cleave the O–O bond, but gives too slow a reaction rate to be practically useful without the alloyed Pd.173, 174 The reactivities of alloy catalysts are dependent on the spacing between the more active metal atoms, such that finely tuning the ratio of inactive to active metal affords a volcano-type relationship between H2O2 selectivity and composition (FIG. 4).

Aside from PdAu, computational studies have shown that Pt or Pd, when alloyed with a wide range of metals, afford catalysts that are very selective towards H2O2, with PtSn2 approaching the peak of the theoretical activity volcano.109 Unfortunately, the dissolution potential of many of these metals is more negative than the reduction potential for oxygen, and so operating conditions would lead to their dissolution out of the alloy, leaving behind a Pt or Pd shell. Taking stability into account leaves PtHg4 as the optimal candidate. The same concept of active sites on a relatively inert substrate has been employed in the development of single atom catalysts,175 which feature single atoms of an active metal such as Pt isolated on (or in) a relatively inactive support such as S-doped carbon141 or TiN.176 The Pt1 active sites favour end-on O2 binding over other binding modes, such that the selectivity for H2O2 generation can reach 95%.

One must never lose sight that optimizing selectivity typically has a detrimental effect on reaction rates. In the case of Pd*x*Au1−*x*, lower values of *x* will favour H2O2 selectivity but afford lower current densities. The ideal electrode material will represent a compromise — diluting the active material enhances selectivity but lowers the total rate of reaction. This present discussion of catalytic activity assumes that the system maintains its integrity under the operating conditions. Some materials are only kinetically stable, and PdAu nanoparticles, when subjected to oxidizing or reducing potentials, can undergo partial phase separation to afford a Au-rich or Pd-rich shell, respectively. These new materials have different selectivities towards H2O2,177 highlighting the need to not only characterize activity but also the robustness of catalysts under operating conditions.

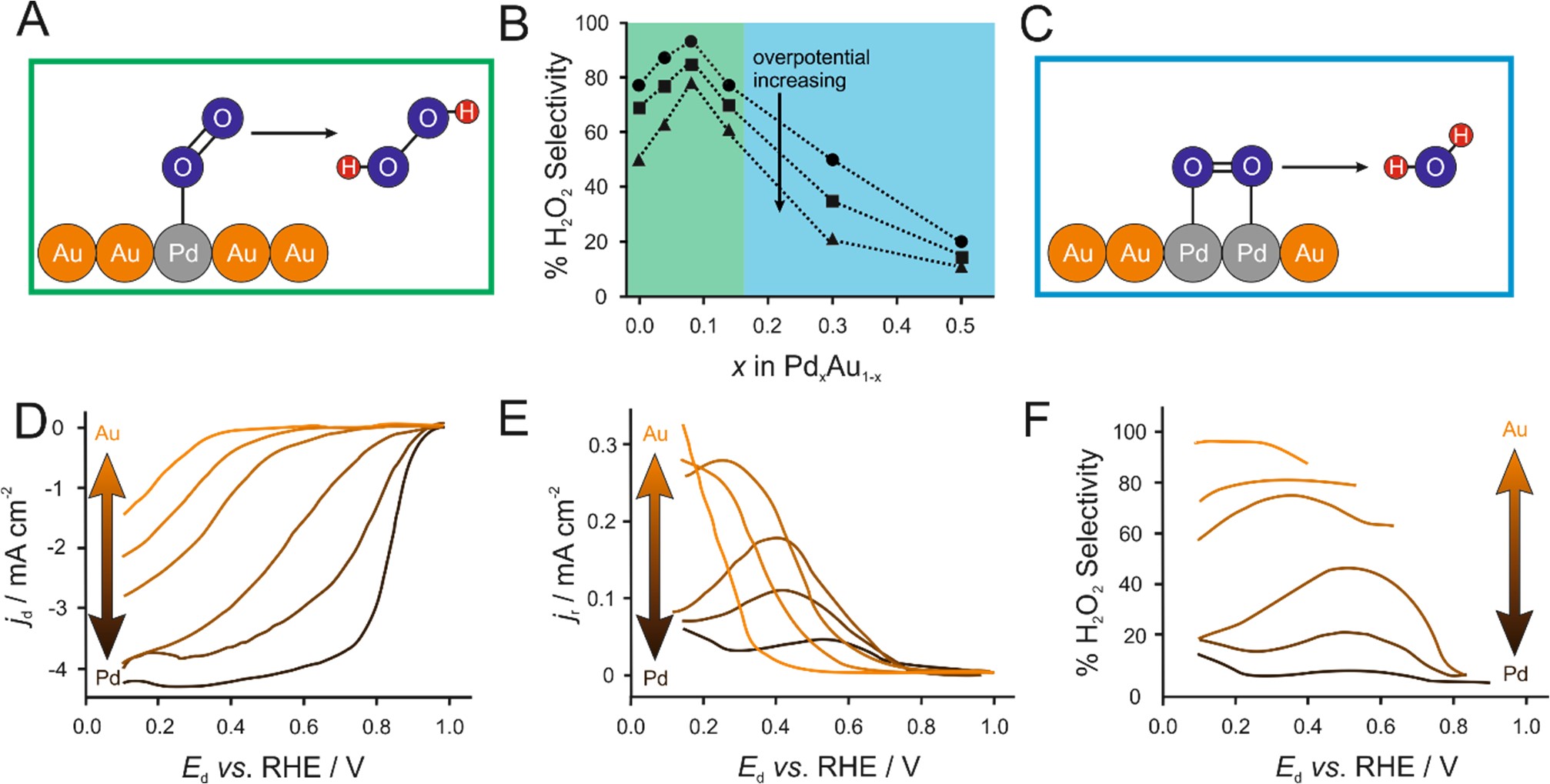


Figure 4 | **Selectivity in O2 reduction to H2O2 as a function of Pd content in Pd*x*Au1−*x*. a** | The Pd(η1-O2) end-on binding mode is known as the Pauling model and is favoured when the Pd content is low. **b** | The selectivity towards H2O2 at 0 (circles), −0.1 (squares) and −0.2 V vs. saturated calomel electrode (triangles) depends on the alloy composition. The green region represents the low Pd region, in which H2O2 formation is favoured. The blue region represents the high Pd region, in which H2O formation is favoured. **d–f** | Pd*x*Au1−*x* alloys can also be studied by immobilizing them on the disc electrode of a rotating ring–disc electrodes. **d** | The total current density at the electrode disc (*j*d) for O2 reduction as a function of disc potential *E*d. **e** | The observed current density at the ring (*j*r) corresponding to H2O2 detection. **f** | The corresponding selectivity for H2O2 for the given catalyst. In general, pure Pd exhibits the greatest current density but pure Au shows the best selectivity, highlighting how alloys give access to a compromise between the two. Part **b** was drawn from data in REF 142, parts **d**–**f** were drawn from data in REF 168.

## Carbon materials

Metal alloys have promising catalytic activities, but tend to include precious metals that are expensive and in limited availability. This has motivated the development of carbon materials as cheap and abundant electrocatalysts. The starting point for these materials can be one of a number of carbon allotropes, including graphite,178-181 graphene,182-184 C nanotubes,135, 185, 186 as well as porous137, 148, 187 and amorphous188-190 C. Porous C materials can have high surface areas and enable efficient mass transport, but can also favour H2O2 by offering more defect sites than the uniform materials.148 According to DFT calculations, certain defect configurations afford a material at pinnacle of the volcano plot for 2e− O2 reduction (FIG. 2), highlighting their role as active sites. In general, bulk coordinatively saturated C sites are less reactive than these defect sites. However, the optimum structure is not obvious because although a material with smaller pores has a higher density of defects, it also can lead to lower H2O2 yields because any H2O2 cannot easily diffuse from the electrode before undergoing further redox.191

In addition to modifying the structure of carbon electrode materials, one can also improve the performance of a H2O2 electrosynthesis catalyst by tuning the composition of the surface or bulk. For example, the current density of a H2O2-generating carbon material can be improved by oxidizing the surface anodically,192, 193 or thermally179 to provide a more active surface decorated with O and/or OH groups. In the case of carbon black, selectivity towards H2O2 formation can be substantially improved by annealing it to afford a more hydrophobic surface that favours the diffusion of O2 to the electrode and helps to maintain the characteristic three-phase boundary that is responsible for fast reaction rates at gas diffusion electrodes.194 As we noted, oxidized carbon materials bearing several –CO2H and –C–O–C– sites are predicted to have optimal activity for H2O2 formation.108

The activities of doped carbon materials are more dependent on the chemical effect of the dopant rather than the effects imparted on microstructure or defects.184, 195 A wide range of dopants have been studied, including B,131, 196 N,197-199 P,182, 184 S,183, 200 F201 and a number of transition metals.202206 Both mesoporous carbon and graphite, when doped with N atoms, can exhibit high selectivities towards H2O2.207, 208 For example, a N-C framework containing triazine and viologen groups has many redox-active sites that can accept electrons and transfer them to O2 to selectively give H2O2, even over an 8 h electrolysis.198 An exception to this has been observed with carbon nanotubes, where N-doping causes the apparent number of transferred electrons to increase (from *n* = 1.8 to 3.9).135

Most transition metal dopants are not suitable for H2O2 electrosynthesis, and the selectivity of metaldoped carbon materials for H2O2 can decrease with increasing dopant concentration.203, 205 Likewise, carbons with lower levels of N and Fe–N dopants have greater selectivities,209 a result that is intuitive given that higher concentrations of active sites favour 4e– ORR. One can obtain selective catalysts by immobilizing [M(porphyrinato)] or [M(phthalocyaninato)] complexes on a carbon support139, 140, 210 but these materials undergo degradation in the presence of H2O2, making them unsuitable for long-

term use.205, 211

## Metal oxides

Metal oxides can produce H2O2 through either cathodic or anodic electrosynthesis (FIG. 5a,b). Ru and Ir oxides mediate the anodic reaction at the lowest reported overpotentials,117, 212, 213 but are too expensive for large-scale applications214 and tend to favour O2 formation.215-218 Most promising metal oxides include SnO2,62 MnO*x*,214, 219 WO3-BiVO4220-222 and TiO2.223-227 Much of the literature dedicated to H2O2 synthesis from these materials is related to the overall photochemical reaction:228

O2 + 2H2O + *hν* **→** 2H2O2 (8)

The process involves the initial absorption of a photon to trigger charge separation: a hole in the valence band (hvb*+*) and an electron in the conduction band (ecb−). These enable H2O2 generation both by H2O oxidation and O2 reduction.225

MO*x* + *hν* **→** hvb+ + ecb− (9a) 2H2O + 2hvb+ **→** H2O2 + 2H+  (9b)

O2 + 2ecb− + 2H+ **→** H2O2 (9c)

Although the requirements for photocatalysts and electrochemical catalysts are expected to differ, exploiting both approaches simultaneously may afford a photoelectrode that forms H2O2 very efficiently.129 Conflicting views exist regarding whether it is the reduction of O2229, 230 or the oxidation of H2O58 that is the predominant source of H2O2. Experiments in degassed solutions suggest that the former reaction is more prevalent 225, 231

Present research into metal oxide catalysts for H2O2 electrosynthesis mostly focuses on refining the catalysts by changing their metal content. This can afford mixed-metal catalysts such as the ternary oxide CaSnO3, which shows good selectivity for H2O2 and was stable over a 12 h electrolysis.132. Alternatively, catalysts can be tuned for H2O2 formation by stacking thin layers of different metal oxides. For example, modifying the surface of a WO3-BiVO4 electrode with other metal oxides increases selectivity towards H2O2 production,228 with a combination of WO3, BiVO4 and Al2O3 being the optimal photoanode (FIG. 5c,d). This increase in selectivity during anodic H2O2 production is interesting given that previous works indicated that the purely photochemical reaction proceeds by a reduction step. Taken together, the mechanisms of the photocatalytic and electrocatalytic reactions are likely different.

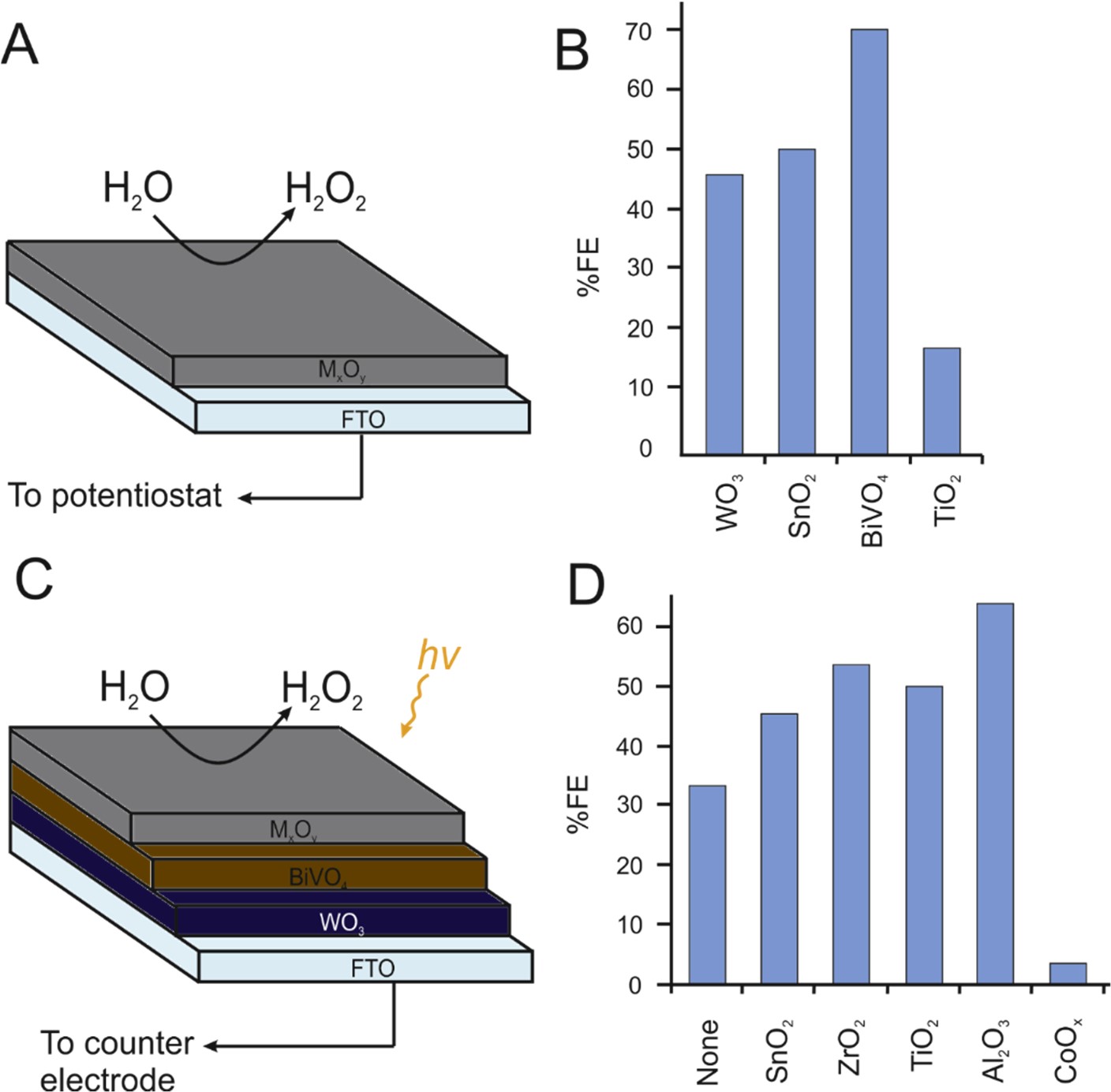


Figure 5. | **Metal-oxide-based electrodes and their performance in electrocatalytic and photoelectrocatalytic H2O2 production. a** | Schematic structure of a metal oxide electrode deposited on FTO glass and connected to a potentiostat. **b** | Each oxide has a different current efficiency, defined as the fraction of the total current density passed (*j*total) used to oxidize H2O to H2O2 (*j*H2O2). Electrodes were used in the dark and biased at 2.3 V vs. RHE (WO3), 3.1 V vs. RHE (BiVO4 and SnO2) and 3.3 V vs RHE (TiO2). **c** | Schematic structure of WO3-BiVO4 deposited on FTO glass and coated with a metal oxide catalyst M*x*O*y*. The electrode is not potentially biased but is illuminated as part of a photovoltaic cell. **d** | The current efficiencies of illuminated photoelectrodes with different catalysts M*x*O*y*. Overall, high current efficiencies are observed for both electrochemically-driven and photochemically-driven reactions. Part **b** was drawn from data in REF 129. Part **d** was drawn from data in REF 146. FTO, fluorine-doped tin oxide; RHE, reversible hydrogen electrode.

Metal oxide electrodes are more commonly used for anodic H2O2 electrosynthesis than they are for cathodic electrosynthesis, although a few suitable materials have been identified. These O2 reduction catalysts typically take the form of metal oxide nanoparticles dispersed on C supports. A number of oxides have shown promise, including WO3,145 Co*x*O*y*,232 CeO2,233-235 Ta2O5,147, 236 Nb2O5237 and V*x*O*y*.238 The usefulness of these species for cathodic H2O2 electrosynthesis is limited because they are intrinsically efficient materials for the 4e– ORR. Thus, their selectivity towards H2O2 can only come from physical engineering, such as loading a surface only sparsely with these oxides to lower the likelihood that H2O2 is reduced to O2.145, 235

Although metal-oxide-based catalysts can have useful activities, their electrical conductivities are lower than metals. Indeed, many of the best performing metal oxide catalysts are semiconductors, and the potential drop across this layer will affect the potential at the electrode surface. Additionally, in the case of metal oxide nanoparticles on a support such as C, the conductivity of the metal oxide will affect the electrical contact between the particles and the support and electrode, which can introduce additional potential drops and detract from the performance of the electrode.

# Electrolytes and additives

The number of electrons *n* accepted by each O2 molecule decreases with increasing pH, to the point where basic solutions favour *n* = 2 and the formation of H2O2. Increasing [OH−] can poison an electrode surface, and the lower density of active sites favours end-on adsorption and precludes the deleterious reduction of H2O2.239, 240 Unfortunately, OH− can also catalyze the disproportionation of H2O2 (equation 4).241, 242 Thus, one must make a compromise with pH — if it is too low then *n* increases towards 4, and if it is too high H2O2 undergoes decomposition.

With the above discussion in mind, it would make sense to deliberately partially poison an electrode surface with something other than OH−. For example, Pt electrodes for O2 reduction can be modified by exposing them to solutions containing Br-,243 I-,244 or Cl-,245 and even traces of these ligands can affect the surface.245-247 Similarly, Pt can be poisoned by adding S or Se atoms.248 Alternatively, Pt nanoparticles can be coated with a layer of porous amorphous carbon, which restricts substrate access and thereby favours end-on binding of O2 and formation of H2O2 as the end product.249 Partially poisoning an electrode will lower the current densities it can operate at because the electroactive area is lowered. This has motivated the development of alternative additives that do not detract from current density but can improve selectivity by affecting the kinetics at the electrode surface and slow down the reduction of H2O2. One option is to add halide salts of N(alkyl)4+, which not only affect the electrical double layer but also adsorb onto the electrode and raise the local pH, favouring localized H2O2 production.250 The selectivity towards H2O2 was the same in N(alkyl)4Br and N(alkyl)4I, so the cation effect was outweighing contributions from the halides. Only low concentrations of N(alkyl)4+ are needed, such that one can favour H2O2 without substantially increasing the bulk pH and catalyzing its decomposition.

H2O2 has a number of decomposition pathways, both through heterogeneous redox at an electrode (Equations 1 and 2) or through homogeneous reactions in bulk solution (Equations 4 and 7a). Heterogeneous decomposition can be slowed down by minimizing contact between H2O2 and the counter electrode by placing the latter in a separate cell compartment.214, 251 Homogeneous decomposition is harder to prevent because the process can occur by homolysis or by reaction with radicals or OH−. H2O2 decomposition can also be catalyzed by trace metal ions such as Fe2+, which participates in the Fenton reaction (equation 10):252-254

Fe2+ + H2O2 ⟶ Fe3+ + HO• + OH−  (10)

One can avoid the Fenton reaction by trapping aqueous metal ions as catalytically inactive complexes, including those of chelating ligands such as 2-(*N*-anilino)ethanol or aminophosphonates.254-257 The homolysis of H2O2 affords radicals that can proceed to react with H2O2:258

HO2• + H2O2 ⟶ HO• + H2O + O2  (11a)

HO• + H2O2 ⟶ HO2• + H2O (11b)

It is possible to prevent these deleterious cascade reactions by trapping HO• and HO2• with radical scavengers such as N-oxides or peracids.259, 260 and to stop their formation in the first place by shielding the cell from the UV light that would otherwise facilitate H2O2 homolysis.261 A distinct approach to avoiding H2O2 decomposition is to convert the product into a peroxosolvate salt, which can precipitate from solution and be stored as a stable solid and used as an on-demand oxidant or hydrolyzed back to H2O2.262-264 As well as avoiding its decomposition, sequestering H2O2 shifts reaction equilibria towards H2O2 as the end product. This requires separation of the peroxosolvate to isolate H2O2, although peroxosolvates have similar oxidizing properties to H2O2 and might be used directly in some applications.

# Electrodes, cells and their architecture

Complementary to research on developing catalyst materials with the ideal intrinsic properties is the equally important pursuit of engineering the catalyst and other cell components into an efficient device. The ideal electrode structure should have a sufficiently large number of adsorption sites, whilst also allowing H2O2 to rapidly desorb before it can undergo further reactions. The desorption of H2O2 is not only favoured when metal–H2O2 interactions are weak but also under conditions of fast mass transport. High surface areas and fast mass transport give rise to high current densities, which are particularly desirable when scaling up electrochemical cells from the laboratory scale to the industrial scale (FIG. 6).

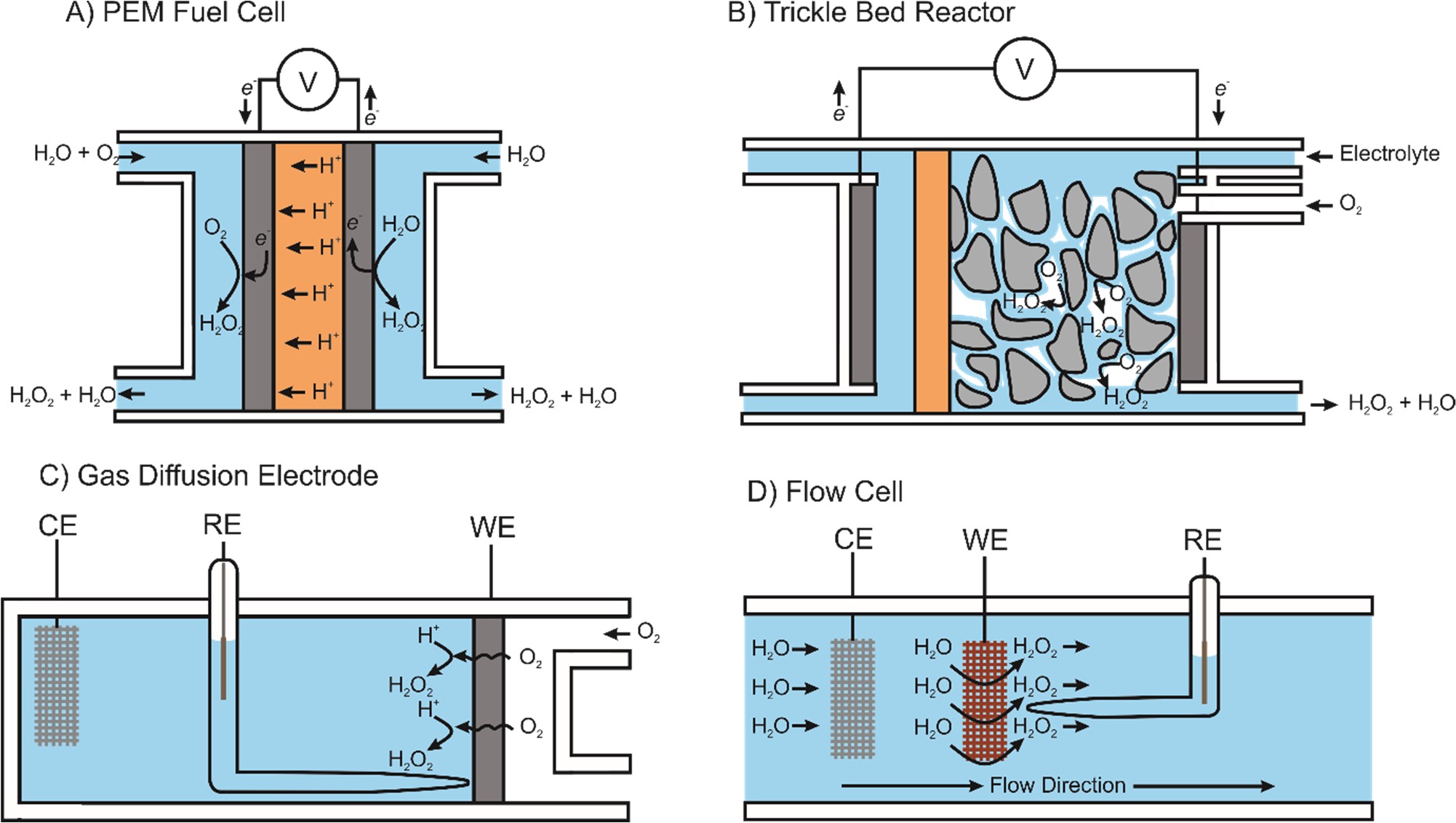


Figure 6. . | **Possible cell configurations for the electrosynthesis of H2O2**. Blue regions denote liquid electrolyte, white regions denote gas flow, and orange regions represent conductive membranes. **a** | In a gas diffusion electrode, O2 flows over the back of a porous hydrophobic electrode before being reduced at the solid–liquid interface. **b** | O2 dissolved in an electrolyte can flow through or by a working electrode and be converted into H2O2. Electrodes such as foams or meshes have high surface areas and can give high current densities. Both the gas diffusion cell and flow cell can also be operated with a membrane dividing the solution into two compartments, with the CE in a separate compartment to the RE and WE. **c** | In a trickle bed reactor, the electrolyte and O2 flow into the cathode chamber, which is packed with the catalyst. The electrolyte forms a thin film over the catalyst bed, so O2 can dissolve and be rapidly reduced to H2O2. The H2O2 solution is then collected at the bottom of the reactor. The anodic chamber is often a flow chamber over a porous electrode to balance the charge. **d** | O2 can be reduced at the cathode or H2O oxidized at the anode to produce H2O2 in a PEM fuel cell. The two electrodes are separated by the PEM and flowing solution over the electrodes gives access to greater current densities. CE, counter electrode; RE, reference electrode; WE, working electrode; PEM, proton exchange membrane.

A number of different electrode and cell designs tackle the same requirements in different ways. A popular choice for H2O2 electrosynthesis from O2 is the gas diffusion electrode (GDE, FIG. 6a),194, 265 which comprises a porous layer with a hydrophobic component, such as carbon paper impregnated with a hydrophobic polymer. The electrode acts as a membrane between the O2-containing gas and the liquid electrolyte. The catalyst is deposited onto the GDE at the solid–liquid interface, so that when O2 flows through the membrane it can be reduced as soon as it dissolves. Much of the research in this area has focused on the nature of the deposited catalyst, with metal nanoparticles being the most common choice. Care must be taken not to load the GDE with too much catalyst because this can physically hinder gas flow and lower the current density.194 GDEs have a number of advantages over normal electrodes (those optimized for solutions) because mass transport in the gas phase is typically faster that it is in solution, and one does not need to consider the solubility limits of O2 in the electrolyte.

We have described how a GDE can be suitable for the cathodic electrosynthesis of H2O2 because it can readily process the gaseous reactant O2. When one instead considers anodic H2O2 electrosynthesis, it is still possible to have a similar effect by using a liquid flow cell (FIG. 6b). This can collect the product in a reservoir or circulate an intermediate to allow it to undergo further reactions within the same volume of solution to give a high concentration of the desired product. A number of distinct reaction environments are established when the solution flow is changed using arrangements like such as flow-by systems over parallel plate electrodes or flow-through systems using mesh or channel electrodes266 These flow cells are particularly useful to assess a new material during its early development because they have well-defined mass transport regimes based on flow rate and geometry, such that setting up a reaction is relatively straightforward and is amenable to validation using finite element analysis.

Other reactor designs have been developed to increase overall currents by maximizing the active electrode area instead of mass transport. In the laboratory, this can be realized by using highly porous electrodes such as carbon felts or foams to offer large surface areas at which electrochemical reactions can occur.4, 267 When scaled up, this concept takes the form of a trickle bed reactor, in which the electrode is a fixed bed of catalyst particles in a large chamber (FIG. 6c).57, 268 The electrolyte is added into the top and then trickles down by gravity such that the effluent can be collected at the bottom. This combination of the large surface area and slow flow rate can afford concentrated solutions of

H2O2269-273

Another reactor option is the proton exchange membrane (PEM, FIG. 6d) fuel cell.60, 188, 210 In such a device, the anode and cathode are in two compartments separated by a H+-conducting polymer membrane. These cells are effective for H2O2 electrosynthesis because the separation of the two electrodes means that H2O2 formed at the anode cannot be degraded at the cathode or vice versa. Electrochemical synthesis within a fuel cell presents the possibility of simultaneously harvesting electrical energy together with a useful product, which has clear energetic benefits, compared to standard synthesis that only consumes energy.274

If one separates the compartments in an electrochemical cell it is possible to simultaneously produce useful products at the anode and cathode and not have the products interact with each other. Perhaps the best known example of such a divergent paired electrochemical process is the chloralkali process, which generates Cl2 and NaOH by oxidizing and reducing brine.275 Many more examples of divergent processes exist, including the conversion of 4-(*t*-butyl)toluene to phthalide,276 L-cystine to ʟ-cysteine and ʟ-cysteic acid,277 dienes and CO2 to diols and diacids278 and O2 and H2O to H2O2 and oxidants such as O3279 or HO•.280 An alternative approach is a convergent paired electrochemical process, in which the same product is generated in both compartments. This has been applied to the electrosynthesis of hydroquinone281, 282 and glyoxylic acid.283 Of course, concomitant cathodic and anodic H2O2 production from O2 and H2O is also possible, and encouragingly, a complete electrolysis cell operating at Faradaic efficiencies exceeding 90% has been patented.284 This convergent paired process appears to be the most feasible approach for H2O2 electrosynthesis on an industrial scale.

Diverse reactor designs are used both in the academic literature and in industry. Numerous patents have been filed for reactor designs for H2O2 electrosynthesis. These are mostly based on the cathodic production of H2O2 from O2 at a gas diffusion cathode in a PEM fuel cell, with the goal being to achieve give high mass transport rates and minimize the problems regarding the low solubility of O2 in aqueous electrolytes.285-287 By refining the design and orientation of the gas and liquid phase compartments, these have been incorporated into stackable, modular units, where the gas and liquid phases flow across multiple sequentially stacked gas diffusion electrodes, which greatly increases the overall output capacity.288 Similar bipolar electrolyzers to those used in the chloralkali process have been used to generate useful products at both the cathode and anode, such as H2O2 and HOCl,289 or H2O2 at both the anode and cathode.290

# Future research needs

A key figure of merit that requires improvement continues to be the selectivity towards H2O2 over H2O or O2. The selectivity of catalysts is often reported at low overpotentials in the kinetically controlled regime, and these catalysts can give rise to low overall yields for high current density electrosynthesis. There is plenty of scope for the improvement of catalyst materials through changing substrate composition, architecture, surface treatment or experimental design. A number of materials have come close to the theoretical maximum (CaSnO3, WO3 and BiVO4 at the anode129, 130, 132, PtHg and PdHg at the cathode107, 109), so these would be good candidates from which to further develop materials. The environmental concerns associated with Hg-based materials, and the costs and unsustainability related to using pure noble metals, mean that Pt or Au alloys109, 142 or N-doped carbon materials137, 197 are among the more practical candidates for cathode materials.

The redox activity of H2O2 makes 100% selectivity impossible. The presence of H2O2 adsorbed to electrode surfaces at large overpotentials means that some degree of further redox seems inevitable for both the anodic or cathodic mechanism. A more realistic aim would be to operate at a high selectivity (≈ 95% for example) while also stabilizing any H2O2 that forms and/or maximizing the current density. We reiterate that much of the available literature has described studies that focus on selectivity, with current density being relatively neglected in fundamental studies. A major challenge is to increase the current density of a H2O2 electrosynthesis cell without lowering selectivity, because, in general, a more active catalyst is also more likely to mediate the decomposition of H2O2 to O2 and H2O.

Once we can perform 2e− redox to favour H2O2 in preference to the 4e− ORR or OER, it will then be necessary to shift focus towards a desired working concentration of H2O2 within the experimental setup. This concentration will vary depending on application; solution phase alkene oxidation syntheses have been reported using between 1.5 and 4.5 equivalents of peroxide to alkene, where the exact ratio depends on the alkene and catalyst used.291 This might be achieved by operating at high selectivity and low current density, or higher current density and lower selectivity. In this regard, the electrosynthesis of H2O2 has a substantial advantage over other synthesis methods because the sideproducts of the reaction are simply O2 and H2O. Thus, obtaining a high yield even at reduced selectivity is not a problem because it will not affect the purity of the resulting H2O2 solution.

Aside from catalytic activity, the practical value of any electrode material also depends on its longterm stability. One should repeatedly cycle the potential and also conduct controlled potential electrolysis over a long time. Following these, it is important to once again characterize the electrode morphology using microscopy techniques, and to look for any changes in the rate of H2O2 formation by complementary electrochemical or spectroscopic techniques. Repeated use of an electrode even at moderate overpotentials can cause the metal and metal oxide catalysts to degrade. Indeed, H2O2 itself can break down a number of potential catalyst materials. For example, although promising in terms of activity, metalloporphyrin-based species have revealed themselves to be insufficiently robust, even in relatively short-term experiments139, 140 Other active materials may also prove unsuitable when subjected to more rigorous electrosynthesis trials conducted over hundreds of hours.

This Review has described different H2O2 electrosynthesis strategies from both cathodic and anodic directions. On the face of it, there appears to be no clear preference between these options, except perhaps that because the anodic route uses H2O as the starting material it is easier to realize practically because it does not require the same level of solution aeration as the cathodic route. The distinction between the two directions is more about what happens at the counter electrode, something that is often not considered during electrochemistry, so long as a sufficient current can be passed and no disruptive side-products are generated. However, a growing number of researchers are now designing paired electrochemical processes for the production of useful materials at both the anode and the cathode.281, 292 Thus, the choice of anodic or cathodic H2O2 electrosynthesis is dependent on the additional product that is desired; anodic H2O2 electrosynthesis also provides H2,220, 222 whereas cathodic H2O2 electrosynthesis can generate O2 or O3.279 Likewise, the oxidation of 2H2O or H2O2 can be coupled to reduction of CO2 to HCO2H or C2H4. Organic products might later be combined with H2O2 to give other useful products.293, 294 As mentioned in the preceding section, one can also design a cell purely for electrochemical synthesis of H2O2, which forms simultaneously at the anode

and the cathode.228, 290

The emphasis of H2O2 electrosynthesis research should be guided by the application for which the H2O2 product is intended. The various applications of aqueous H2O2 will each require a certain H2O2 concentration and may only tolerate a certain pH range. If we consider a plausible end goal of online H2O2 electrosynthesis, in which electrosynthesized H2O2 is immediately consumed en route to a value-added product, it is important that H2O2 can be synthesized at a comparable rate and at a concentration in a solvent/solute mixture appropriate for its final use. For example, if the second reaction is alkane oxidation, a big challenge is to produce H2O2 in sufficiently high concentrations to drive the effect the oxidation. Thus, instead of using a separate reactor for the second step it may be advantageous to conduct a one-pot synthesis, in which H2O2 is formed in the presence of the organic to be oxidized. This would favour alkane oxidation because H2O2 would be abundant at the electrode surface (it would be more difficult to achieve the necessary concentrations in a bulk phase to be delivered to a second reactor). H2O2 can also be used for H2O treatment because its catalytic decomposition by Fe cations effects the oxidation of organic impurities to less harmful products. This process is best performed at low pH in order to favour radical reactions and prevent precipitation of the catalytic Fe cations as hydroxides.295, 296 If H2O2 is to be used in situ for the Fenton reaction, the candidate catalysts are limited to the small fraction of catalysts that are known to be efficient in acidic media. Otherwise, a workup strategy must be found, involving an acidification step while the solution is moved from the first reactor to the second.

Long-term goals for H2O2 electrosynthesis are likely to focus on scalability — moving from benchtop experiments to syntheses on industrially relevant scales. Many of the necessary reaction cells and architectures already exist for other reactions and could be easily adapted to H2O2 electrosynthesis. Up-scaling would most likely require increased stability of catalyst materials in concentrated H2O2. Alternatively, modifications could be made to the industrial rig, such as incorporating a flow system to remove H2O2 from the electrode to give added protection. Of course, the efficiency with which electrochemistry can afford useful dilute solutions of H2O2 means that we do not necessarily have the same high yield requirements as present industrial methods.

When considering the advancement of H2O2 electrosynthesis in the industry, it is unfortunate that an electrochemical engineering approach has not been described in most studies published. These studies neglect to target a certain reaction environment, as can be done by considering fluid flow, mass transport and current distribution. Only in this way can scale-up and industrial integration of electrochemical systems be possible. Advances in the achievable H2O2 production rates are coming through the use of porous 3D electrodes decorated with catalysts such as coated metal or carbon mesh, felt, foam or fibres.297-299 Optimizing the channel depth, width and orientation within specifically designed flow fields can maximize the exposure of electrodes to the reactant — either H2O through liquid flow over the anode or O2 by gas flow over the cathode. Fundamental research into new catalyst materials for H2O2 electrosynthesis will find more relevance if it includes expanded analysis in which the entire reaction environment is considered in order to make materials ready for wide applications.

# Conclusions

A diverse range of electrode materials is available for the electrochemical synthesis of H2O2. For cathodic electrosynthesis, carbon-based electrodes perform well because of their poor efficiency for the 4e− ORR. The efficiency with which these catalysts produce H2O2 can be further improved by adding dopants or metal nanoparticles. For the anodic route, metal oxides such as CaSnO3, BiVO4 and WO3 perform well. Many of these materials were originally developed as photocatalysts but can still give impressive yields without illumination. Accordingly, there is much scope for further improvement of these materials to enhance the H2O2 yield by focusing on selectivity and/or current density. The choice of cathodic or anodic synthesis becomes more important when considering the reaction occurring at the counter electrode. Thus, H2O2 and another product can be formed in paired electrochemical processes or H2O2 can form at both the anode and cathode as part of dual electrosynthesis. The majority of papers published describe only short-term studies on model electrodes in laboratory electrolytes, so it is important to establish whether electrodes can exhibit long-term performance. Thus, one must consider their volumetric area, activity, resistance to degradation and their suitability to practical scale-up and industrial processing.

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