Models and Mechanisms of Oxygen Evolution Reaction on Electrocatalytic Surface

Sulay Saha, Koshal Kishor, Sri Sivakumar, and Raj Ganesh S. Pala

Abstract | A profound change has taken place in understanding surface electrochemistry during water-splitting reaction due to the accumulated knowledge over the past decades and supported by recent advances in spectroscopic techniques and high-performance quantum chemical simulations. The design of electrocatalysts has been improved due to better understanding of surface structures of electrocatalysts and their active sites. This review provides insights into both theoretical and experimental electrochemistry that are directed towards a better understanding of the rate-determining step of water splitting, i.e., oxygen evolution reaction (OER). The emphasis of this review is on the origin of the electrocatalytic activity of nanostructured catalysts toward the aforementioned reaction by correlating the electrode performance with their intrinsic electrochemical properties. Also, the design aspects of acidic- and alkaline-medium electrocatalysts is discussed from preliminary discussion on active site engineering to a more applied concern of achieving highly stable and active electrode fabrication. The design concerns while choosing a support for OER electrocatalyst has also been discussed. At the end, challenges in electrolyser designs and problems faced by the industry to commercialize the electrolyser in a cost-effective manner have been discussed.

1 Introduction
Generation of clean energy is one of the main issues of the 21st century because the use of the fossil fuels has problems associated with diminishing reserves and environmental compatibility. An alternative source of the energy carrier can be hydrogen, which has high energy density and can also generate clean discharge (water) on burning. Also, hydrogen is the most important chemical in chemical industry as petrochemical, fertilizer and semiconductor industry are significantly dependent on its supply. However, pure hydrogen as a gas is scarce in the earth’s environment and its demand is mainly met through steam-reforming of hydrocarbons, which generates CO and CO₂ as byproducts. Electrochemical water electrolysis is another process, which can generate hydrogen. Though this process does generate clean fuel, it is an energy-intensive limiting its industrial application.

Hydrogen is generated at the cathode and oxygen is generated at the anode during water electrolysis. The process involves oxidation (loss of electrons) at anode and reduction at cathode (protons accepting electrons, which come from the conducting wire) with the help of supporting electrolyte in aqueous medium. The generated hydrogen can be used as an industrial fuel, or for hydrogenation reaction as well as source of electricity when “burnt electrochemically” in a fuel cell. Generation of hydrogen through water electrolysis was preferred over steam-reforming until the flourishing of petrochemical industry with accompanying decrease in crude oil prices, which led to closure of many electrolysis plants. The upward trend in crude prices along with environmental concerns these petrochemical processes bring has led to relook of water electrolysis with increasing interest. The general efficiency of these electrolysis processes is around
−80% and the lab-scale efficiency is around −94% (See for definition of efficiency in equation 14). Since the whole electrolysis process is electrical energy-intensive due to inherent endothermic nature of the reaction, coupling the electrolyzer with a renewable energy source is being suggested to increase the viability of the electrolysis process. Among different energy sources that are coupled with water electrolyzers, steam\textsuperscript{11}, wind\textsuperscript{12}, solar\textsuperscript{13,14} and geo-thermal energy\textsuperscript{15} have been mainly explored. Also, instead of electrochemical hydrogen generation, photo-electrochemical hydrogen generation is also being tried and recently getting investigated in great detail.\textsuperscript{16,17} Table 1 summarizes the different types of electrolyzers that are used in the industry and their efficiency.

Ideal water splitting requires 1.23 V potential at standard thermodynamic conditions. However, the activation barrier associated with the chemical reaction and ohmic loss leads to splitting of water at much higher potential. The hydrogen evolution (HER) at the cathode is a two-electron process and OER at the anode a four-electron process. It has been observed that OER at the anode is the rate-determining step and requires higher overpotential when compared to HER. Thus, the performance and efficiency of the water electrolysis cells is intricately related to the performance of OER electrocatalyst at the anode. Until the 1970s, expensive OER catalysts, made up of late transitional metal oxides (e.g., Ru, Ir), were generally used in the electrolyzers. This led to increase in cost of the overall process, making the whole process economically less viable. With the introduction of alkaline water electrolyzers\textsuperscript{20} and invention of dimensionally stable anodes\textsuperscript{21} in acidic medium in the 1980s, the focus of electrocatalysis research has shifted to finding active electrocatalysts via earth-abundant and cheap material like Ni, Co, Fe have lower activity as well as suffer from corrosion. Thus, it is not only important to understand the reaction mechanism of electrochemical OER but also the associated dissolution reaction of anodes under electrochemical OER condition. Understanding of the reaction mechanism, identification of active and stable sites in electrocatalysts is greatly expected to help in the design of better electrocatalysts for the future.

This work summarizes the recent progress in the development of OER electrocatalysts in acidic and alkaline media. Since, both these electrocatalysts can be used in neutral medium and neutral medium electrolyzers show lower efficiency due to low ionic conductivity of water and do not have any limitations in electrocatalysts, they are not discussed in this review. Instead of focusing on catalyst preparation and its role in increasing the activity, we have paid attention to the nature of the active sites of electrocatalysts and their conditioning for designing superior electrocatalysts with better stability and activity. Fundamental aspects of the OER and its electrochemical aspects are also addressed particularly as well as practical requirements and benchmarking criteria while reporting OER electrocatalysts. Since the electrode comprises support also, we have discussed the design criteria for supports. However, we note that finding a better OER electrode is only one aspect of making an efficient electrolyzer and several additional aspects of water electrolysis must be studied, e.g., contact resistance, polymer exchange membranes.\textsuperscript{22}

1.1 Electrode kinetics and benchmarking of activity

1.1.1 Electrode kinetics: An understanding of the reaction kinetics is of utmost importance in the design of a better electrolyzer. The reactions at both the cathode and anode are redox reactions

<table>
<thead>
<tr>
<th>Serial no.</th>
<th>Type of electrolyzer</th>
<th>Type of energy used</th>
<th>Efficiency (%)</th>
<th>Operating temperature</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Alkaline</td>
<td>Electrical</td>
<td>80</td>
<td>60°C\textsuperscript{18}</td>
</tr>
<tr>
<td>2</td>
<td>Acidic</td>
<td>Electrical</td>
<td>85</td>
<td>80°C\textsuperscript{19}</td>
</tr>
<tr>
<td>3</td>
<td>Steam</td>
<td>Electrical and Heat</td>
<td>92</td>
<td>800–1000°C\textsuperscript{11}</td>
</tr>
<tr>
<td>5</td>
<td>Photo-voltaic</td>
<td>Solar</td>
<td>90</td>
<td>105°C\textsuperscript{14}</td>
</tr>
<tr>
<td>6</td>
<td>Wind</td>
<td>Wind</td>
<td>78%</td>
<td>80°C\textsuperscript{12}</td>
</tr>
<tr>
<td>7</td>
<td>Geothermal</td>
<td>Geothermal</td>
<td>80</td>
<td>900°C\textsuperscript{15}</td>
</tr>
<tr>
<td>8</td>
<td>Photo-electrochemical</td>
<td>Solar</td>
<td>13</td>
<td>65°C\textsuperscript{16}</td>
</tr>
</tbody>
</table>

* Pressurized water is used in high-temperature electrolyzers.
$O + ne \rightarrow R$ involving electrons. Nernst equation governs the thermodynamic potential for such a reaction. This equation correlates the electrode potential ($E$) with the concentration of the products and reactants:

$$E = E^o - \frac{RT}{nF} \ln \left( \frac{C^R_C}{C^O_C} \right)$$

(1)

Here, the term $E^o$ is the thermodynamic equilibrium potential of an overall reaction, $R$, the universal gas constant, $T$, the temperature, $n$, the number electrons associated with the reaction, $F$, the Faraday constant, and $(C^*/C_o^*)$, the concentration of oxidized/reduced species in the system. The thermodynamic equilibrium potential is the minimum potential bias that is required for any electrochemical reaction to calculate using the following equation:

$$E_{eq} = \frac{\Delta G}{nF}$$

(2)

During electrochemical reaction, redox reaction occurs at both anode and cathode. The current density obtained from a specific reaction at a particular electrode (e.g., anode) is given by,

$$j_f = nFk_f C_o$$

(3)

where $j_f$ and $k_f$ are the current density and reaction rate constant, respectively. Here, a standard rate constant $k^e$ and rate constant can be related with Arrhenious relationship by:

$$k_f = k^e \exp\left(-\alpha f \left(E - E^{eq} \right)\right)$$

(4)

where $f$ is equivalent to $F/RT$ and $\alpha$ is termed as transfer coefficient with value between 0 to 1, which can be found out by Marcus theory.\textsuperscript{[23-25]}

The electrochemical reactions do not occur at equilibrium thermodynamic potential and certain excess potential is required to overcome the energy barriers involved in the activated processes correlated with specific reaction intermediates and this excess potential is known as overvoltage or overpotential in electrochemist's parlance. This overpotential is written as:

$$\eta = E_{eq} - E$$

(5)

$$j = j_o \left[ \frac{C^*_R}{C^*_o} \exp(\alpha f \eta) - \frac{C^*_R}{C^*_o} \exp((1 - \alpha) f \eta) \right]$$

(6)

where $C^*_R$ and $C^*_o$ are the bulk concentration of the reduced and oxidized species and $j_o$ is known as the exchange current density of the reaction. The exchange current density is an important descriptor for electrochemical reaction and is indicative of the activation barrier of the reaction. If the electrochemical reaction is not diffusion limited, i.e., either the electrolyte bath is well stirred, or the reaction rate is slow, then the bulk and the surface concentration can be taken as approximately almost equal (i.e. $C^*_R = C^*_o$ and $C^*_o = C^*_o$). In such a situation, the above equation would reduce to an equation which is more famously known as the Butler-Volmer equation,

$$j = j_o \left[ \exp(-\alpha f \eta) - \exp((1 - \alpha) f \eta) \right]$$

(7)

According to the Butler-Volmer equation at small $j_o$, the reaction kinetics would be slow. The Butler-Volmer equation shows that current density at any overpotential is the summation of oxidation (forward) and reduction (backward) reaction occurring at electrode surface. At acertain overpotential the contribution from the backward reaction is almost zero and only forward reaction dominates and in that case, Butler-Volmer equation for OER is reduced to the following form,

$$j = j_o \left[ \exp(-\alpha f \eta) \right]$$

(8)

$$\log j = \log j_o - \alpha f \eta$$

(9)

The above equation simply states that the generated current due to electrochemical reaction is exponentially linked to the overpotential in the absence of diffusion resistance imposed on reactants and this reaction is better known as the Tafel equation. The above equation can be written in the following form,

$$\eta = a + b \log j$$

(10)

where $a$ and $b$ are the Tafel constants ($a = \frac{2.3}{\alpha f} \log j_o, b = -2.3 \frac{1}{\alpha f}$). Tafel equation is valid for all electrochemical reactions and is useful in the analysis of electrochemical energy storage devices (fuel cell, battery and electrolyzer, etc.).\textsuperscript{[26-28]}

1.1.2 Benchmarking parameters for electrocatalyst: The performance of an electrocatalyst is usually gauged from the other electrocatalysts by using two typical metrics, (i) apparent total electrode activity (e.g.,
current density expressed in mA/cm²\text{geometric} or overpotential required to attain certain current density), which is the most important factor for device-oriented electrocatalyst design, and (ii) the intrinsic activity of each catalytic site (e.g., turnover frequency (TOF) or specific current density expressed in mA/cm²\text{ECSA}) and this metric is more useful for fundamental studies on catalyst design. Among other electrochemical metrics, Tafel slope, exchange current density (\(J_0\)) or charge-transfer resistance (\(R_q\)) calculated using electrochemical impedance spectroscopy at a potential in Faradaic region are also used in comparing the activity of electrocatalyst and understanding the basic phenomena occurring at the electrode/electrolyte interface. Generally, the activity of OER electrocatalysts is benchmarked through metrics of (i) overpotential required to attain a current density of 10 mA/cm²\text{geometric} and (ii) current density at an overpotential of 0.35 V.\text{29,30} To attain higher efficiency, a lower Tafel slope and higher current densities are desirable. Preparation of three-dimensional porous electrocatalyst is an effective strategy to enhance OER activity. In these electrodes, the number of active sites would be significantly larger than that may arise from geometrical surface area though almost neither electronic properties nor surface structure of active sites has been altered. To compare the increased surface area from the apparent geometrical surfacearea, the following metric named Roughness Factor (RF) is introduced.

\[
\text{Roughness Factor (RF)} = \frac{\text{Electrochemical surface area (ECSA)}}{\text{Geometrical surface area (GSA)}}
\]

where RF can be as high as 500–4000 in a porous 3D electrode. Consequently, while assessing the activity of individual sites, it is imperative to normalize it by real surface area or the TOF is to be calculated. However, the key issue is the accurate measurement of the electrocatalytically active surface area (ECSA), which can reflect the total number of active sites on a given surface for a certain reaction. While for metals, either underpotential deposition (upd) hydrogen (H\text{upd}) adsorption voltammetry\text{31} or CO stripping voltammetry\text{32} is widely used, it is not useful for metal oxides, especially for OER. The surface area calculated using BET does not reflect the ECSA and sometimes differs by an order of magnitude.\text{33,34} For example, in ABO, perovskite material, either A or B would be electrocatalytically active for OER, but BET measurement would show both of them as active sites.\text{35} Therefore, it is advisable not to use BET surface area for measuring specific activity of OER. Other methods include measurement of double-layer capacitance (\(C_{dl}\)) through cyclic voltammetry or impedance spectroscopy at non-Faradaic region and then correlating it with ECSA\text{36} by using the following equation,

\[
\text{ECSA} = \frac{C_{dl} \cdot d}{\varepsilon_0 \varepsilon_r}
\]

where \(C_{dl}\), \(\varepsilon_0\), \(\varepsilon_r\) and d are the double-layer capacitance electrical permittivity, relative electrical permittivity and distance between two layers, respectively (d = 10 Å).\text{37} However, the measurement of the interfacial double-layer capacitance is methodologically difficult due to the fact that some semiconducting and capacitive properties as well as the double-layer frequency dispersion has contribution towards total capacitance, leading to questions on the reliability of the computed value.\text{30,38–40}

In porous 3D electrodes, porosity is another useful parameter but it needs to be measured. The porosity observed in liquid solution under electrochemical conditions would differ from that measured through BET, which are more accurate for solid-gas-phase heterogeneous catalytic conditions. Trasatti et al. suggest that porosity calculation through electrochemical means mirror more porous conditions in solid-liquid solution catalysis.\text{41} They suggest that the total charge (\(q_t\)) and outer charge (\(q_o\)) of metal oxide can be calculated by integration of voltammograms at different scan rates. The voltammetric charge corresponding to the total surface area (\(q_t\)) can be calculated by plotting the reciprocal of \(q_t\) against the square root of the potential scan rate and then by extrapolation of the linear plots at \(v = 0\). The values of outer charge (\(q_o\)) can be calculated from the extrapolation to \(v \rightarrow \infty\) in the plot of \(q\) vs \(v^{-0.5}\).

The electrochemical porosity is measured as

\[
\text{Electrochemical porosity} = \frac{q_t - q_o}{q_t}
\]

Apart from this, Tafel slope and exchange current density (\(J_0\)) are used for comparing the OER activities of the electrocatalysts. Though Tafel slope is widely used as performance metric of OER electrocatalyst, it cannot be used as a singular measure since electrocatalysts having the same Tafel slope and different \(J_0\) would show different OER activity. For calculation of \(J_0\), special care is required as the current axis is in logarithmic scale and small errors during extrapolation of Tafel slope or ohmic-drop correction may result in
large errors in $I_0$ values. Also, the experimentally obtained current densities are dependent on various measuring conditions such as (i) scan rate, (ii) electrolyte concentration, and (iii) distance between electrodes. One of the alternative ways of benchmarking condition is potentiostatic measurement in which steady-state potential is applied or galvanostatic measurement in which steady-state current density is maintained for a fixed time limit, say 1 h. The steady-state current density after a particular time gives a measure of activity while the reduction in current density after a fixed time is indicative of the stability of electrocatalyst. In this way, both activity and stability of different electrocatalysts can be measured.

One must note that the current density obtained in electrochemical OER conditions does not fully correspond to OER. It includes contribution from (i) dissolution current of electrocatalyst, (ii) pseudo-capacitive current, (iii) current arising from surface reconstruction the electrocatalyst, and (iv) other possible electrochemical reactions due to interaction with anions of supporting electrolyte, particularly in acidic conditions. Predominance of one of these above phenomena during OER condition would lead to wrong conclusion over electrocatalyst performance. Hence, it is better to calculate the Faradaic efficiency of OER through actual O$_2$ gas collection or measurement of O$_2$ concentration through in-situ spectroscopy. The Faradaic efficiency is given as

$$\text{Faradaic efficiency} = \frac{\text{Experimentally collected total volume of oxygen gas}}{\text{Theoretical amount of oxygen gas should evolve}} \times 100\%$$

(14)

Overall, depending on the electrocatalyst material, one should decide the benchmarking conditions they would choose.

2 Molecular Mechanisms in Oxygen Evolution Reaction

2.1 Mechanistic pathway of OER

It is essential to understand the mechanistic pathway and reaction barriers associated with OER for designing a better electrocatalyst. With the advent of density functional theory (DFT)-based simulations, it has become far easier to (i) discover new reaction mechanistic pathways for reactions, and (ii) reveal active sites of catalysts, which may help in the design of better catalysts. These active sites are generally under coordinated surface sites of the electrocatalyst, which interact with reactants to form different reaction intermediates. Using DFT methodology, the energy of the possible reaction intermediates is calculated and the lowest activation barrier pathway is identified as the reaction pathway. The highest energy barrier reaction step of the lowest energy pathway is identified as the activation barrier with the reaction step identified as the rate-determining step.

The mechanistic pathway of OER on catalyst surface is dependent on electrochemical potential and surface coverage. Previously, it was believed that OER follows Langmuir-Hinshelwood mechanistic pathway involving neighboring active sites. Bockris et al. suggest the following mechanistic pathway (Figure 1):

$$S + H_2O \rightarrow S - OH + H^+ + e^-$$
$$S + OH \rightarrow S - O + H^+ + e^-$$
$$2S - O \rightarrow 2S + O_2$$

where S is the active site. The existence of this reaction pathway for OER has been proved by isotope-labelling studies for RuO$_2$ electrocatalyst. With the advent of more powerful experimental spectroscopic and computational facilities, another parallel pathway involving OOH intermediate was proposed by Norskov et al.

The elementary reaction of this mechanistic pathway is as follows,

$$S + H_2O \rightarrow S - OH + H^+ + e^-$$
$$S - OH \rightarrow S - O + H^+ + e^-$$
$$S - O + H_2O \rightarrow S - O - OH + H^+ + e^-$$
$$S - O - OH \rightarrow S + O_2 + H^+ + e^-$$

where S is the active site. The schematic pathway of this mechanism for RuO$_2$ is given in Figure 2.

The preferred mechanistic pathway depends on the hydroxyl coverage over the active catalyst, which in turn is dependent on the electrochemical potential. At lower electrochemical potential, $-\text{OH}$ coverage is found to be more in comparison to $-\text{O}$ coverage. In this zone, $-\text{OOH}$ formation pathway is favored. With increase in electrochemical potential, $-\text{OH}$ coverage over the surface is reduced and interaction between two neighboring $-\text{O}$ adsorbates becomes possible thus making Langmuir-Hinshelwood mechanistic pathway plausible.

Due to the presence of these two mechanistic pathways, the rate-determining step for OER differs depending on the electrochemical potential applied. This is reflected in the switch of Tafel slopes commonly observed during OER. At lower overpotential, it is common to observe a Tafel slope $< 120$ mV/dec, while at higher overpotential a Tafel slope of $> 120$ mV/dec is observed.
Also, the mechanistic pathway differs depending on the oxide surface. As for example, RuO$_2$ (110) surface is found to show peroxide formation as a rate-determining step as it is far more easier to oxidize Ru-5c sites on (110) the surface compared to Ru-6c.$^{52}$ However, the rate-determining step of TiO$_2$ (110) surface is found to be different and is found to be the adsorption of oxygen adsorbates over Ti-5c sites as it is far more harder to oxidize Ti-5c sites on (110) surface compared to Ti-6c.$^{52}$ Also, in the same material, different surfaces behave differently depending on the electrochemical potential. This is exemplified in the case of Co$_3$O$_4$ through DFT studies by Santen et al.$^{46}$ At lower electrochemical potential, the most active surface is found to be (001) surface with the rate-determining step being the desorption of O$_2$ with the involvement of dual Co surface sites and Langmuir-Hinshelwood dominating the mechanistic pathway. But as the electrochemical potential is increased, the structure-sensitivity of other surfaces comes into play and (311) surface is found to be more active with the formation of –OOH as the rate-determining step.
of activation barrier due to the involvement of two Co (IV) sites instead of a single Co (V) site during OER leads to the formation of a bridged intermediate having lower potential energy (Figure 3). The delocalization of holes around dual sites is also responsible for the lowering of potential energy. Further increase in electrochemical potential leads to single Co-site of (110) surface as the most active one. Similar results are obtained in the case of RuO$_2$, where OER activity between single-crystal-cut (100) and (110) surfaces are compared, and (100) surfaces are more active in comparison to (110) surfaces. This type of structure sensitivity is due to interaction between neighboring sites and higher density of active metal sites per geometrical surface area.

It is being observed that surface metallic sites get oxidized to a higher oxidation state during OER (M$^{n}$ $\rightarrow$ M$^{n+m}$). If the chemical potential of surface oxides with higher oxidation state is lower than the chemical potential of bulk oxides with lower oxidation state in aqueous medium under electrochemical OER conditions, then the dissolution of electrocatalyst takes place, which seriously hampers the performance of electrolyzer in the long run. For example, RuO$_2$ undergoes dissolution to RuO$_4$ with loss of activity under a potential of E $>$ 1.4 V in aqueous medium.

RuO$_2$ (s) + 2H$_2$O (l) $\rightarrow$ RuO$_4$ (s) + (4H$^+$ + 4e$^-$)

Similar dissolution has been observed in the case of MnO$_2$ with generation of MnO$_4^-$ species in aqueous medium,

MnO$_2$ (s) + 2H$_2$O (l) $\rightarrow$ MnO$_4^-$ + 4H$^+$ + 3e$^-$

This dissolution reaction occurs in parallel with OER. In an ideal OER electrocatalyst, the dissolution reaction would be suppressed without affecting the OER kinetics. However, the intermediates of dissolution reaction of electrocatalyst and OER are identical in most of the cases, which makes increasing OER activity over dissolution reaction hard to achieve. Current research and developments on OER electrocatalyst are focused on enhancing OER activity, while suppressing the dissolution of electrocatalyst to electrolytes. The current method of suppressing electrocatalyst dissolution into electrolytes involves preparation of doped/alloyed oxide or solid solution involving stable oxides (e.g., mixing stable TiO$_2$ with active RuO$_2$). However, lack of understanding of dissolution mechanism of electrocatalysts has made development and design of stable electrocatalyst largely empirical in nature.

### 2.2 Origin of volcano plot and its utility in catalyst design

OER reaction involves 4-electron transfer with the formation of various reaction intermediates. Ideally each electron transfer would require the same amount of energy. However, variation in the requirement of energies due to energetics of reaction intermediates imposes activation barrier for the reaction. For designing a better catalyst, Sabatier principle is recalled, which states that the catalyst should bind the reaction intermediates not too weakly or nor too strongly so that adsorption and desorption kinetics becomes faster. In the case where the intermediates bind too weakly, they cannot activate the surface leading to lower activity. However, when they bind too strongly, all available surface sites are occupied leading to poisoning and desorption is difficult. Hence, moderate binding energies of intermediates on catalyst surface are a good compromise between these two extreme situations.

![Figure 3: Mechanism of water adsorption to (a) a terminal Oxo on a single-Co Site, and (b) a bridging Oxo on a dual-Co Site at different points along the reaction trajectory. PCET stands for proton-coupled electron transfer. The figure is adapted from American Chemical Society.](image)
To understand the reaction kinetics better, a number of ‘descriptors’ are used and they are plotted against the actual activity of electrocatalysts. These descriptors are derived from the binding energies of adsorbates on the electrocatalyst surface. The binding energy of –OH, –O and –OOH adsorbates on the electrocatalyst surface vary between 0 and 5 eV. Plotting of activity against descriptor results in generally “volcano”-shaped correlation (Figure 4) and further, such descriptors are used for predicting the activity of an unknown material.

Trasatti et al. suggest enthalpy of formation of oxides from a lower oxidation state to higher oxidation state as a descriptor \((M^{\text{Ox}} \rightarrow M^{m+\text{Oy}})\) and plotting this energy against the activity of electrocatalyst resulted in a “volcano plot” (Figure 4a).\(^{54-66}\) It is suggested that OER can be simply viewed as a transition between lower to higher oxidation state during adsorption of reactant water molecule and a transition between higher to lower oxidation state during desorption of \(O_2\) gas molecule. As a result, metal oxides that are hard to be further oxidized would not be a good catalyst since reaction intermediates are weakly adsorbed. In these cases, water dissociation over catalyst surface would be the rate-determining step (e.g., \(\text{TiO}_2\)). On the reverse, oxides that are easily oxidized further would not be catalytically active since intermediates are strongly adsorbed and desorption of \(O_2\) would be the rate-determining step (e.g. \(\text{OsO}_2\)). Later, Norskov et al. correlated OER reactivity with difference in the formation energy of oxide intermediate (–O) and hydroxyl intermediate (–OH) due to reaction of adsorbate-free surface and \(H_2O\) molecule (Figure 4b).\(^{48,52}\) This descriptor works well for various oxide surfaces such as rutile, pervoskite, spinel, rock salt, and bixbyite.\(^{48}\) Regardless of the quality of electrocatalyst material, the above descriptor imposes an overpotential limitation of 0.4 V for OER and this corresponds to the peak of the volcano.\(^{48}\) However, this imposed limitation is not sacrosanct and can be circumvented by changing the nature of active sites.\(^{67,68}\) Rossmeisl et al. showed through DFT simulation that OER on Ni- and Co-doped \(\text{RuO}_2\) electrocatalyst has proton donor–acceptor functionality on otherwise inactive under coordinated oxygen sites at the surfaces, which enhance the surface reactivity at under coordinated (cus) \(\text{Ru}(-5c)\) sites by reducing the formation energy of –O and –OOH reaction intermediates.\(^{67}\) This is also reflected experimentally as Ni-doped \(\text{RuO}_2\) is far more active than conventional \(\text{RuO}_2\) as predicted from volcano relationship.

While designing an electrocatalyst with doped oxide or alloy of an oxide, the target is to ascend towards the peak of the volcano. To achieve this, the general design principle is to combine materials from different legs of the volcano curve as a synergistic effect is expected. However, it has to be kept in mind that energetics of doping follows its own rule and dopant in doped oxide might not occupy the active sites due to thermodynamic conditions during synthesis. This is especially true for aliovalent and guest-host size-mismatched dopants, where surface segregation and undesirable defect formation is quite common.\(^{69}\)
Usually, the free energy and related quantities for intermediates formed in oxygen-involving reactions on various surfaces change are correlated. This phenomenon has been observed for different surfaces including metals and metal oxides and is known as a scaling relationship between the oxygen intermediates. Specifically, in the case of metal surfaces and transition metal oxides, the lines corresponding to –OOH and –OH are almost parallel to each other with the slope of one, as shown in Figure 5. The underlying principle for such behaviour is that the adsorbates, –OOH and –OH, form single bond with the catalyst surface through an oxygen atom via a single bond while –O forms double bond with the catalyst surface. This is reflected in the slope of formation energy of –O intermediate vs –OH intermediate which is nearly 2 (i.e. \( \Delta G_{\text{O}} / \Delta G_{\text{OH}} = 2 \)). Although surface-adsorbate scaling relationships provide qualitative arguments for tuning catalytic activity based on the bond strength between catalyst surface and adsorbate, they provide only a limited guideline for prediction of OER activity with increasing/decreasing coordination in lattice or change of elements, which can further help in the search for better catalyst. In this regard, the relationship between the surface adsorbate, binding strength and the electronic band structure has to be investigated for engineering better catalysts. The electronic structure of d-bands of transition metal surfaces was found to have a correlation with chemisorption energies for adsorbates on the surfaces and is useful for both explanatory and predictive tool in the catalysis. In metal oxides, metal 3d-band splits into two kinds of electronic states: with doubly degenerate \( e_g \) symmetry and triply degenerate \( t_{2g} \) symmetry. It is found that there exists a correlation between adsorption energies and the occupancy level of \( t_{2g} \) orbital. Since the \( e_g \) and \( t_{2g} \) orbitals are normally nonbonding or antibonding and the 2p orbitals of oxygen are the bonding orbitals in octahedral complexes, an increase in the number of outer electrons in transitional metal would correspond to a weakening of the binding strength while interacting with electrophilic adsorbates (–OH, –O, –OOH). Distortions to the octahedral symmetry of the metal cation coordination structure lead to changes in both the shape of the \( t_{2g} \)-band and adsorption energy which lead to changes in OER activity. Also, the binding strength of atomic oxygen adsorbate on surfaces of metal oxides can be correlated to lattice oxygen atoms’ p-band centre in a metal oxide. Also, the p-band centre position scales are found to vary linearly with oxygen vacancy formation energy and oxygen surface exchange kinetics. Though these scaling relations are helpful in understanding some of the features of catalyst design, it is important to recognize their limitations. While the predicted trend provides insight into understanding the difference in OER activity across different electocatalysts, these relations are applicable for surfaces that follow a similar reaction mechanism. Change in reaction mechanism may result in change in the scaling relations with breakdown of volcano relations.

### 2.3 Nature of electrocatalyst-electrolyte interface

An aspect worth considering but is often disregarded during computational catalyst design is the nature of the electrocatalyst–electrolyte interface during electrochemical OER conditions. Oxides having metallic electronic conductivity results in lower ohmic drop in the electrode bulk material. Among oxides, RuO\(_2\), IrO\(_2\), and LaNiO\(_3\) show quasi-metallic electronic conductivity. However, most of the other transition metal oxides used for OER are semiconducting in nature which...
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3.1 OER electrocatalyst for PEMWE

3.1.1 Monometallic oxides: The electrochemical splitting of water performed in acidic polymer electrolyte membrane water electrolyzers (PEMWEs) has several distinct advantages compared to that of alkaline systems as discussed in earlier sections. The main challenge here is to design a cost-effective, and active but stable electrocatalyst in acidic medium. Only a small number of metallic oxides are stable in the acidic medium (see Table 2), which restricts the manoeuvrability in designing superior catalysts. The experimentally observed overpotential sequence for OER in acidic solution is as follows: Ru < Ir < Pd < Rh < Pt < Au, while the stability order is Pt – Pd < Ir < Rh < Au – Ru.79,80

The above relation is true for sputtered thin film oxides and both their activity and stability can be tuned through engineering of electrocatalyst surfaces via different synthesis conditions. For example, RuO₂ (100) surface is more stable and active than RuO₂ (110).86 Therefore, both theoretical and experimental results for OER catalysts should be analysed with due care since the former study only considers ideal surfaces and the latter provides results strongly influenced by the catalyst preparation process and surface state. Also, crystallinity of prepared electrocatalyst also affects the electrocatalytic activity due to defect formation and surface stress. The activity ordering is found to be: amorphous RuO₂ > polycrystalline RuO₂ > single crystal RuO₂, while the stability is inverse of this order.81,82

The evolution of RuO₂ surface structure has been a subject of much interest.55,60,79,83,84 In-situ XANES study of Ru-electrode revealed that under 0 < E < 0.8 V RHE potential Ru-remains in metallic state, between 0.8 < E < 1.0 V RHE potential the oxidation state is between 3 and 4, under 1.0 < E < 1.3 V RHE rutile oxide is formed with oxidation state of 4 and above E > 1.4 V potential, the oxidation state is higher than 4.85 Prior to OER, it has oxidation state of 6 and with its onset its oxidation state increases to 8. RuO₂ undergoes dissolution mainly as RuO₄⁻. Though corrosion rate increases with onset of OER, small amount of dissolution is observed even below OER. As expected, the dissolution rate is expected to depend on the stability of the intermediates formed during OER and dependent on the crystallinity, defects and the nature of the exposed surface which is again dependent on the synthesis process.

Besides RuO₂, another very active OER electrocatalyst in acidic medium is IrO₂. Though it is expensive and its electrocatalytic activity towards OER is inferior to RuO₂, its stability in high anodic potentials in acidic environment makes it attractive.79,86 As in RuO₂, both the activity and stability of IrO₂ can be tuned through controlling the exposed facet in the surface, crystallinity and defect formation, which are again dependent on the...
Ir-metal also undergoes change in surface oxidation state similar to Ru but the potentials are higher than that of Ru. Even at high potential of 1.95 V RHE, sputtered IrO₂ catalyst shows no corrosion.

3.1.2 Doped and alloyed oxides: The idea of synthesising mixed oxide electrocatalysts is to introduce synergistic effect of both components and climb uphill in the volcano curve. To enhance the stability of RuO₂, the general strategy is to mix it with stable oxides even if they are passive oxides such as IrO₂, CeO₂, SnO₂, TiO₂. Like RuO₂, these oxides possess rutile structures (e.g., IrO₂, CeO₂, SnO₂, TiO₂) but have high overpotential for the OER as well as lower electrical conductivity, which makes them a poor choice if they are used as stand-alone OER electrocatalysts. However, when these oxides are mixed with RuO₂, the current density is higher than separate phases of individual oxides and the solid solution has greater stability. On detailed surface study after prolonged galvanostatic treatment of RuO₂–IrO₂ catalyst, the surface contains almost no Ru, but still its activity is higher. This is attributed to the pure Ir surface skeleton arising due to surface Ru-dissolution that now contains higher number of steps and kinks which might be having greater activity than planar surface and higher density of active sites per geometrical surface area. The DFT simulation study of OER in defective surfaces of electrocatalyst has not yet been investigated.

Similar increase in activity is also observed in RuO₂–TiO₂ electrocatalyst, which is generally known as dimensionally stable anode. Naslund et al. suggest that the increase in activity due electron transfer from Ru to Ti site, which activates otherwise benign Ti atoms. Indeed, the increased activity is achievable with Ru concentration in the doped oxide as low as 30%. DFT simulation suggests that activity of Ru surface atom on TiO₂ (110) surface does not get affected due to change in the neighboring environment. However, the activation barrier for neighboring Ti-atoms gets reduced due to distortion of surface in the presence of bigger Ru-atom. Yet, the activation barrier for OER for Ti surface dopants in RuO₂ almost remains unaffected. Hence, it is more beneficial to dope Ru on TiO₂ matrix. One other way of activating these electrocatalysts further is to introduce another dopant, which may activate the TiO₂ electrocatalyst. Lower-valent cations such as Ni, Co and Zn have been found to activate both RuO₂ and TiO₂ individually for OER even beyond volcano limitations as found through both experimental and simulation. Additionally, these aliovalent dopants improve charge transfer behavior through formation of oxygen vacancy in bulk electrocatalyst, which result in improved conductivity in addition to lowering activation barrier for OER through strain and electronic effects. These non-Noble dopants are unstable in acidic medium and their dissolution would generate porous electrocatalyst as well as higher number of kinks, steps, and defects on the surface (see Figure 7). Indeed Pala et al. showed that the OER activity of Zn-doped RuO₂–TiO₂ electrocatalyst increases with electrochemical cycling and time. This is also accompanied by an increase in ECSA and electrochemical porosity. Additionally, this is achieved without compromising stability and also reducing the cost of electrocatalyst. This can be a model for generating porous, active and stable electrocatalyst for acidic conditions. However, the intrinsic specific activity of generated active sites of this type of electrocatalyst is yet to be investigated.

Search for activated cheap electrocatalyst resulted in further research on activating TiO₂ electrocatalyst, which is stable in acidic medium and good for photo-electrocatalysis but performs poorly as an electrocatalyst due to its n-type semiconductor nature and high OER

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**Table 2:** Stability of the mono-elementary oxides in acidic medium under electrochemical OER conditions.
The yellow color denotes formation of unstable oxide and green that of stable oxides. Blue color denotes the presence of two types of oxides under different electrochemical potential.
A TiO$_2$–MnO$_2$ electrocatalyst was found to show good stability and activity in acidic medium. Manganese dioxide is stable only up to 1.6 V RHE and beyond this it starts to undergo dissolution. The dissolution front in doped-

MnO$_2$ stops once it encounters Ti-surface atoms and the stability is retained for long duration. Also, Ti prefers to be under coordinated sites in MnO$_2$, hence additional synthetic treatment is not required for making the electrocatalyst stable.

**Table 3:** Comparison of electrochemical OER activity of different electrocatalysts in acidic medium.

<table>
<thead>
<tr>
<th>Electrodes</th>
<th>Overpotential needed to reach 10 mA/cm$^2$ (mV) or * Activity at an overpotential of 350 mV (mA/cm$^2$)</th>
<th>Tafel slope (mV/dec)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crystalline RuO$_2$</td>
<td>450 mV (*7)</td>
<td>52</td>
<td>61</td>
</tr>
<tr>
<td>RuO$_2$ (110) thin film</td>
<td>#0.65</td>
<td>56</td>
<td>56</td>
</tr>
<tr>
<td>RuO$_2$ (100) thin film</td>
<td>#0.18</td>
<td>54</td>
<td>56</td>
</tr>
<tr>
<td>Amorphous RuO$_2$</td>
<td>220 mV (*42)</td>
<td>33</td>
<td>82</td>
</tr>
<tr>
<td>IrO$_2$</td>
<td>450 mV (*4.6)</td>
<td>58</td>
<td>55,105</td>
</tr>
<tr>
<td>IrO$_2$ (110) thin film</td>
<td>#0.0005</td>
<td>55</td>
<td>56</td>
</tr>
<tr>
<td>IrO$_2$ (100) thin film</td>
<td>#0.003</td>
<td>61</td>
<td>56</td>
</tr>
<tr>
<td>Ru$<em>{0.5}$Ti$</em>{0.5}$O$_2$</td>
<td>*37</td>
<td>56</td>
<td>61</td>
</tr>
<tr>
<td>Ru$<em>{0.5}$Zn$</em>{0.5}$Ti$_{0.5}$O$_2$</td>
<td>*55</td>
<td>33</td>
<td>61</td>
</tr>
<tr>
<td>Ru$<em>{0.5}$Ir$</em>{0.5}$ (RuO$_2$–IrO$_2$)</td>
<td>—</td>
<td>58</td>
<td>55</td>
</tr>
<tr>
<td>IrO$_2$@RuO$_2$</td>
<td>250 mV</td>
<td>58</td>
<td>106</td>
</tr>
<tr>
<td>Ru$<em>{0.5}$Ir$</em>{0.5}$Ce$<em>{0.5}$O$</em>{2.5}$</td>
<td>—</td>
<td>46</td>
<td>95</td>
</tr>
<tr>
<td>Mn$<em>{0.78}$Ti$</em>{0.24}$O$_2$</td>
<td>^4</td>
<td>113</td>
<td>104</td>
</tr>
</tbody>
</table>

^ Activity at an overpotential of 600 mV (mA/cm$^2$).
# Activity at an overpotential of 300 mV (mA/cm$^2$).
3.2 OER electrocatalyst for alkaline electrolysis

Though there are several challenges with alkaline water electrolyzer, lower cost coupled with the fact that there is greater material choice makes it a worthy option for extensive exploration. Unlike acidic medium, oxides of non-Noble elements (e.g., Ni, Co, Mn) are stable in alkaline medium over a higher range of anodic potential. The trend of activity among electrochemically prepared oxyhydroxide is Ni > Co > Fe\(^{107}\) and monolayer thin-film oxyhydroxides also have similar activity order with Ni > Co > Fe > Mn.\(^{108}\) However, computational studies indicate a different activity order Co > Ni > Mn.\(^{48}\) Depending upon anodic potential, these metals undergo transformation from hydroxide to oxyhydroxide to oxide forms. In addition to their surface OER activity, the electronic properties and conductivity also widely vary due to the formation of different phases. This influences the overall OER activity and the activity trends among non-Noble oxides are found to be potential dependent.\(^{109}\)

Cobalt-based electrocatalyst: Cobalt oxide is an important OER catalyst in alkaline medium and the literature has exclusive review on its behavior.\(^{110–112}\) Co-electrocatalyst forms different forms of water oxidation catalysts such as CoO\(_x\), CoOOH and Co(OH)\(_x\). During OER, Co undergoes surface oxidation from Co(II) \(\rightarrow\) Co(III) \(\rightarrow\) Co(IV).\(^{112,113}\) Also, surface reconstruction takes place depending upon electrochemical potential and electrochemical activity being proportional to the surface density of Co (III) sites. DFT studies indicate that CoOOH (1014) surface is the most stable below 1.13 V RHE, while CoOOH (0112) surface is predominantly above this potential.\(^{114}\) In spinel oxide of CoO\(_x\), Co (II) site remains inert, while Co (III) site is responsible for OER activity. This was proved by the fact that blocking Co (II) site by inert Zn atom does not hamper OER activity but blocking Co (III) site by inert Al atom significantly reduces it.\(^{115}\)

Also, the OER activity is facet dependent and modulated synthesis procedure has to be employed to attain high active facets.\(^{46,116–119}\) Another option is amorphization of the catalysis through (i) low temperature heating, or (ii) controlled electrochemical reduction.\(^{120,121}\) Also, metal-organic framework decomposition-derived CoO\(_x\) electrocatalysts show high activity due to their porous 3D configuration.\(^{122–124}\)

Introduction of dopant in cobalt-based electrocatalyst can substantially increase its OER activity. DFT simulation studies predict that doping of Ni and V in Co-sites of (1014) and (0112) surfaces would lower the activation barrier of OER.\(^{114}\) It has also been demonstrated experimentally that NiCoO\(_x\) is five times better in terms of specific activity compared to CoO\(_x\).\(^{13}\)

Fe incorporation also increases the intrinsic activity of CoOOH \(-100\)-fold, with peak activity for 40–60\% Fe.\(^{125}\) The Co films have Tafel slopes of \(-62\) \(\text{mV/dec}\) and Fe dopant between 0.33 < \(x\) < 0.79 leads to Tafel slopes of 26–39 \(\text{mV/dec}\). The stability will decrease with increase in Fe content. Incorporation of Fe dopant does not increase conductivity in CoOOH and after a certain Fe percentage, conductivity will even decrease. Fe dopant shifts the anodic peak of Co(II) \(\rightarrow\) Co(III) compared to what occurs in Co-oxide/oxyhydroxide and makes oxidation of Co (II) more difficult. Fe provides the active sites instead of CoOOH and the energetics of OER for surface Fe sites in CoOOH is far more different than FeOOH. Among other dopants, Cu\(^{126}\), Zn\(^{127,128}\), Cr, Mn\(^{129}\) increase the OER activity of co-oxides or their oxyhydroxide. However, the increase in activity does not appear significant in comparison to either Fe or Ni dopants.

Nickel-based electrocatalyst: Nickel oxide is among the other important electrocatalysts due to its high activity though its stability is an issue. Ni has several oxide, oxyhydroxide (\(\beta\)-NiOOH, \(\gamma\)-NiOOH) and hydroxide polymorphs \([\alpha\text{-Ni(OH)}_2, \beta\text{-Ni(OH)}_2]\).\(^{107,130}\) Less crystalline \(\alpha\text{-Ni(OH)}_2\) polymorph gradually changes into a more crystalline \(\beta\text{-Ni(OH)}_2\) phase during potential cycling in strong alkaline medium. It has also been suggested that the \(\beta\text{-Ni(OH)}_2\) phase is the most active for OER since its oxidation to the \(\beta\text{-NiOOH}\) phase represents the optimal condition for the OER to occur.\(^{107,131}\) However, during the initial electrochemical cycling, \(\alpha\text{-Ni(OH)}_2\) was found to be a better electrocatalyst than \(\beta\text{-Ni(OH)}_2\).\(^{132}\) Similarly, ordered \(\beta\text{-NiOOH}\) phase is generally formed during electrochemical cycling though disordered \(\gamma\text{-NiOOH}\) phase is more active for OER. The onset of OER is accompanied by Ni (II) \(\rightarrow\) Ni (III) \(\rightarrow\) Ni (IV) transformation.\(^{133–135}\)

To improve electrocatalytic performance of Ni-based electrocatalyst, doped oxides are being made. Among the first-row transition metals, doping of Fe in NiO\(_x\) results in significant increase in its activity.\(^{136–138}\) Even small amount of Fe doping results in significant increase in activity.\(^{134}\) Incorporation of Fe dopant result leads to almost 30-fold enhancement of conductivity. Additionally, Fe incorporation leads to changes in electronic structure of NiOOH. Electrochemical studies show an anodic shift of the Ni redox waves with increasing Fe content in NiFeOOH indicating...
lowering of Ni-oxidation states\textsuperscript{139-142} contrary to claims of some researchers that Fe doping and associated partial charge transfer and formation of Ni (IV) are easier in NiOOH structure.\textsuperscript{108,134,140} Computational studies with the support of experimental studies suggest that the Fe-O bond distance in Fe-doped NiOOH is unusually short, which is responsible for lowering of overall OER intermediates leading to the conclusion that Fe is the actual active site in NiFeOOH.\textsuperscript{142,143} As like other electrocatalysts, morphology modulation towards higher aspect ratio nanostructures, generation of 3D microstructure of electrocatalyst and amorphization of Fe-doped NiOOH leads to higher apparent electrocatalytic activity. The addition of other elements (e.g., Ce, Al, Co) may have additional effects, but does not appear to dramatically enhance OER activity.\textsuperscript{138}

**Iron-based electrocatalyst:** Due to the abundance of Fe in earth’s crust and nontoxicity, Fe-based electrocatalysts are appealing. The Fe-based oxide, oxyhydroxide, and hydroxides are less active, unstable and have low bulk conductivity.\textsuperscript{65,107,108,146,147} Fundamental studies on pure Fe-based electrocatalyst are severely constrained by the following factors: (i) Pure Fe-oxides and hydroxides are prone to corrosion in alkaline medium, and the relationship between activity and dissolution is difficult to deconvolute,\textsuperscript{30,33,109} (ii) Fe does not exhibit clear reversible redox waves in the experimentally accessible electrochemical window making it difficult to calculate ECSA through integrating redox waves, and (iii) the low electrical conductivity of bulk Fe-oxide/hydroxides leads to further lowering of activity.\textsuperscript{148} Stabilization by using other dopants leads to increase in activity, however, poor conductivity is always a problem. However, Fe(III) is indeed the best electrocatalyst for OER.\textsuperscript{109,148} It is better to dope Fe in other oxides/hydroxides or oxyhydroxide than doping other transition metals in Fe-based matrices.

**Manganese-based electrocatalyst:** Mn forms oxides with different oxidation states depending upon the electrochemical potential.\textsuperscript{149} At lower oxidation potential (below 0.5 V RHE), Mn$_3$O$_4$ was observed and Mn$_2$O$_3$ formed above 0.85 V RHE. Near OER potentials, Mn(III) $\rightarrow$ Mn(IV) transformation occurs and MnO$_2$ was formed above $\sim$1.4 V RHE potential. Above 1.7 V RHE, dissolution sets in with the formation of MnO$_4^-$ species with solution colour changing to violet. The presence of di-m-oxo-bridged Mn ions in the layered structure results in a pronounced redox and charge capacity behaviour but a relatively large Tafel slope by developing a layered and 3D cross-linked MnOx catalyst structure.\textsuperscript{150} In contrast, the 3D cross-linked structures with both mono- and di-m-oxo-bridged Mn ions present lower intrinsic OER activity but a smaller Tafel slope.

### 3.3 Support design for electrocatalyst

An ideal support should be: (i) stable under reaction conditions, (ii) provide low charge-
transfer resistance, (iii) lower charge density around metal atoms of metal oxide electrocatalyst through electrocatalyst-support interactions, which in turn lowers the activation barrier for OER, and (iv) provide mechanical stability to the electrocatalysts, which are often in the form of nanostructures or thin films. It is generally agreed that among all the support materials Au performs best for thin film (< 5 monolayer) electrocatalysts since it is (i) the most electronegative metal and can increase the oxidation state of metal atom (e.g., Co, Mn, Ni) population on the surface by acting as an electron sink,\textsuperscript{112,120,134–136} (ii) it has high electrical conductivity, and (iii) is stable in OER conditions. However, it is prohibitively expensive, which restricts its utility in electrolyser. Also, taking account of 3D electrocatalyst, which may have a thickness of greater than >10 monolayer, the electronegativity effect of Au gets reduced.\textsuperscript{112} This leads to search for alternative cost-effective supports, which can influence the electrocatalyst via (i) electronegativity that can cause changes in electronic structure, (ii) electrical conductivity that can reduce the charge-transfer resistance in the bulk, and (3) support faceting. The exposure of highly active facets due to support faceting is extremely case-specific in nature and is dependent on crystal-growth mechanism of electrocatalyst on the surface. During conformal thick film growth, the effect of support faceting is sometimes lost due to reorientation of stacking of planes and generation of polycrystalline films. However, activity of OER for thin film electrocatalyst is significantly affected due to support-faceting. This has been demonstrated by Switzer \textit{et al.} by epitaxial growth of Co$_3$O$_4$ on Au and stainless steel support.\textsuperscript{117} The morphology and XRD of the grown thin films have been found to be different for the same amount of electrocatalyst deposited.

To increase the OER activity, three-dimensional porous supports are also used. Highly porous carbon paper or zeolite support enables achieving high electrochemical surface area (ECSA).\textsuperscript{157,158} However, low electrical conductivity of these support increases the charge-transfer resistance of electron transport thereby reducing the overall Faradaic efficiency of the OER support.\textsuperscript{155,163} Furthermore, they have stability-related problems in higher potential regimes of OER. It is generally observed that OER rate increases at higher electrochemical potential, which would increase oxygen diffusion in the electrocatalyst and ultimately oxidizing the carbon support itself. To alleviate electrical conductivity-related problems, some researchers suggest using porous metallic framework such as Ni or Cu foam as a supporting material.\textsuperscript{159}

Other approaches to increasing efficiency include making a metallic interlayer within the porous carbon paper itself, thereby providing an alternate low resistance path of electron transport. A recent study by Pala \textit{et al.} showed that Cu would be the best metal among non-Noble transition metals for making interlayer in porous carbon support for increasing OER activity.\textsuperscript{151} It is found that OER activity of the catalyst varies almost linearly with conductivity of the mellic interlayer (see Figure 8). When OER activity is plotted with the conductivity of metallic interlayer, two classes of behavior of metal supports are discernible: (i)

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure8.png}
\caption{Specific activity (activity/ECSA) at an overpotential of 0.4 V (vs Ag/AgCl) vs conductivity of Co$_3$O$_4$ on metallic interlayer plot of different EAM-modified carbon supports.\textsuperscript{151}}
\end{figure}
metals behaving like electron-sink give higher activity, and (ii) metals behaving like electron donor give lower activity. Supra- or sublinear behavior of the electrocatalyst was observed depending upon whether the metal is electron-sink or electron-donor. Correlations between OER activity with conductivity is especially important in the case of thick film three-dimensional electrocatalysts, while in the case of thin films, electronic interaction between support-electrocatalysts is more important as mentioned earlier.

The supports are broadly classified as: (i) porous three-dimensional (3D) supports, which can provide high surface area (e.g., graphene, Ni foam, silica, zeolite), and the porous structure of carbon or zeolite support enables achieving high electrochemical surface area (ECSA) for electrocatalysts resulting in higher activity though they have low electrical conductivity of carbon paper in comparison to metal support, which increases resistance for electron transport, thereby reducing overall efficiency.

It is imperative to have strong interfacial bonding between electrocatalysts and support for stabilizing the electrode. However, lattice mismatch at interfaces make its formation difficult, which in turn reduces the stability of the electrode. One way to reduce the strain at electrode-support interface is to link electrocatalyst with support through conducting covalent linkage, e.g., click linkage of azide-alkyne. Click-linked electrocatalysts and support show same degree of electrocatalytic activity in comparison to unclicked electrocatalysts due to conducting click-linkage with a higher degree of stability.

4 Electrolysers for Oxygen Evolution Reaction
4.1 Electrolyser design
An electrolyser comprises the following components: (i) cathode, (ii) anode, (iii) current collector, and (iv) ion-exchange membrane. Cathode, anode and ion-exchange of membrane are in conjugation known as gas-diffusion layer (GDL). The gas-diffusion layer and current collector are connected to make the whole membrane electrode assembly (MEA). Separator is used in between cathode and anode for both kinds of cells to avoid the mixing of oxygen and hydrogen produced during water electrolysis. Asbestos was commonly used as a separator but now it has been replaced with oxide ceramic materials or polysulfone. Generally, the gap between the electrodes is kept minimal for reducing the ohmic loss occurring in the electrolyte of the water electrolysis cell. To reduce the gaps, compressive pressures are applied to make a compact electrolysis setup and lower contact resistance. Typically, high compressive pressure between gas diffusion layer (GDL) and current collector (CC) is applied through a nut-bolt assembly. The applied pressure reduces ohmic resistances offered by different components and interfaces of the cell (e.g., electrode-CC plate/bipolar plate interface), which are broadly termed as contact-resistance. However, the application of pressure affects the mesoporosity of the electrodes and reduces its expected electrocatalytic activity. This makes the design of pressurized electrolyzer more complicated and efforts are being made to rationalize some of the manufacturing techniques.

4.1.1 Alkaline electrolysers: Alkaline electrolysers were first made available in 1975. Earth-abundant electrocatalysts can be used in these types of electrolysers. They generally work at a temperature of 60–90°C. The electrode is typically synthesized by the nickel metal/metal oxide electrodeposited on metal plates. Alkaline electrolysis cells are available in two main categories, a filter press type and a tank type, which contain monopolar electrodes fitted in its tank. The filter press-type water electrolyzer can run at 32 bar. Newly developed alkaline water electrolyzer shows an overall efficiency of 80% and a current...
efficiency 98% with 4–5 kWh electrical energy consumption. The purity of produced hydrogen and oxygen is higher than 99%.

**4.1.2 Acidic electrolyzers:** Acidic electrolyzers were developed by the General Electric Company back in 1950. The liquid electrolyte was exchanged by the solid polymer electrolyte membranes like Nafion, which is a cation-conductive polymer membrane. An acidic water electrolyzer contains a stack of numerous MEA (membrane electrode assembly), which is separated by current collector diffusion layers that permit the produced gases and reactant water. The gas diffusion layer promotes optimal concentration of water and produced gases. The solid cation exchange membrane prevents the recombination of produced gases, oxygen and hydrogen. Acidic electrolyzers perform better in comparison to alkaline water electrolyzers with several advantages like achieving current density of 2 A/cm² at a comparatively lower power consumption. The pressurized acidic electrolyzers can work under higher pressure with pressures up to 150 bar.

**5 Conclusion and Future Challenges**

Water electrolysis to hydrogen and oxygen is one alternative way to balance fluctuating electricity generated through renewable energy sources such as solar, wind and tidal energy. To encourage clean energy generation and storage, both increase in efficiency and reduction in the cost of water electrolyzers have to be accomplished. For this, the key challenge is to develop low-cost electrocatalysts, which can show low overpotential as well as low Tafel slope for both HER and OER. Though electrocatalysts for PEM electrolyzers show excellent performance for water splitting, they are generally made of Noble late transition elements, which increase the cost.

Various strategies have been considered for enhancing the performance of electrocatalysts either through increasing (i) specific activity, or (ii) active site density per geometrical surface area (GSA). Synthesis of mesoporous three-dimensional networks to increase surface area and active sites or exposing higher percentage of active facets through various morphologies has been explored. Employing multiple synthesis techniques, e.g., dealloying has been explored to increase the number of active sites in electrocatalysts. Even then, non-Noble electrocatalysts generally show lower activity and stability in comparison to Noble metal counterparts.

This leads to investigation of newer strategies to increase specific activities such as, (i) doping or alloying to get an active site distribution, which has Noble-metal-like properties or (ii) stabilization of non-native polymorphs of the material. In some cases, it has been observed that small changes to bond lengths/angles result in orders of magnitude differences in activity. Doping leads to strain formation as well as change in d-band structure of host metal oxides, resulting in change in OER activity. Aliovalent doping leads to vacancy formation and it may produce additional effects like (i) bulk vacancies that increase the conductivity of the materials significantly leading to overall enhancement of OER activity due to reduced charge-transfer resistance, or (ii) surface vacancies leading to distorted metal octahedra configuration and to changes in energy barriers of reaction intermediates. The above reasons are thought to be valid for enhanced OER activity of Zn-doped RuO₂ and TiO₂. Enhancing specific activity through polymorph engineering is still among less-investigated techniques. Kitchin et al. showed that coulombite IrO₂ would show 0.15 eV less activation barrier than rutile IrO₂. The change in polymorphic structure results in huge change in d-band structure and considerable change in adsorption energies of intermediates. Also, it increases the possibility of stabilization of surfaces, which are structure-sensitive towards OER and thus can enhance the reaction rate considerably. One of the effective ways to stabilize these non-native polymorphs having low surface energy is through tuning surface to bulk ratio. Also, certain supports preferentially stabilize specific non-native polymorphs in low-dimensional regime during epitaxial growth due to formation of coherent interfaces having favorable interfacial energy. However, the stabilization of these metastable and non-native polymorphs still remains a challenge.

Although some progress has been made in obtaining low-cost electrocatalysts, which are both active and stable, some serious issues have not been resolved in relation to large-scale applications such as (i) large-scale synthesis, (ii) pressure-mediated loss of porosity in electrolysers, or (iii) inherent conductivity-related issues. Electrocatalysts should show excellent electrochemical stability, and have to be mechanically durable for many years so that the maintenance of electrolyzer does not create a problem. Furthermore, coating the electrocatalysts on the support should create a proper conductive interface to facilitate charge-transfer. This has been a problem for especially integrating solution-synthesized electrocatalysts.
onto support for large-area electrodes. In addition, electrodes having carbon supports (CNTs, graphene) and some three-dimensional microstructures (core–shell nanostructures) with excellent electrocatalytic performances often show unsatisfactory mechanical stability. Therefore, the process of scaling up of electrocatalysts from lab-scale to industrial scale has to be addressed.

Furthermore, the electrochemical corrosion of the electrodes has to be addressed. Even non-precious pure metallic catalysts and many alloys suffer acidic corrosion for HER in acid solution. Corrosion problem is more severe for OER. At high anodic potentials for thin film electrodes, both the electrocatalyst and support show higher oxidation states. Furthermore, some of the metal oxide supports may show excellent stability but their conductivity is a big issue. Thus, finding electrochemically corrosion resistant and conductive OER electrocatalysts as well as their support is an urgent issue.

To be viable in the industrial front, the water electrolyser work at generally high current densities, such as 1–2 A/cm². The Department of Energy (DOE) of the United States has set a target for water electrolysis to operate at 2 A cm⁻² with a cell potential of 1.5 V vs RHE by 2017. However, the lab-scale benchmarking conditions for OER electrocatalysts are 10–100 mA/cm². This makes them viable only for low-current devices or where hydrogen demand is less. On the other hand, it should be noted that high-current conditions require different electroanalytical techniques beyond cyclic voltammetry or Tafel measurement, since they involve mass-transport limitations. Moreover, during the measurements, the generated oxygen could diffuse into the cathode chamber where it can be reduced to water on reaction with hydrogen. These phenomena would be significant when dealing with high current densities. In addition, the low current density achieved is also due to the high ohmic resistance offered by the liquid electrolyte and membrane. This supports the fact that there are additional components in addition to better electrocatalyst synthesis, which have to be looked into to increase the performance of electrolyser.

Better design of electrolyser would only be possible with increased understanding of several phenomena that occur during water electrolysis, e.g., (i) OER pathway under different electrochemical potential, (ii) charge transfer between electrocatalysts and support and electrode-electrolyte interface, (iii) corrosion reaction pathway and corrosion fronts during OER, (iv) electronic nature of active sites, and (v) diffusion of oxygen through electrolyte during electrolysis. Understanding of all these phenomena at basic atomistic level would help us to design better electrocatalysts, which would enable efficient cost-effective electrolyser.

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