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Sustainable seawater electrolysis: evaluating environmental impacts and technological development opportunities

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As the global energy landscape shifts towards sustainable development, seawater electrolysis is increasingly recognised as a promising technology for green hydrogen production. This approach utilises the abundant availability of seawater, presenting a more environmentally friendly alternative to conventional hydrogen generation techniques. Despite its promise, seawater electrolysis faces significant challenges in energy efficiency, catalyst performance, and the management of undesirable by-products. This review provides a comprehensive analysis of recent advancements in seawater electrolysis, with particular emphasis on its environmental impacts and the technological opportunities it presents. The challenges of seawater electrolysis are discussed, with issues related to electrodes, electrolytes and membranes being highlighted. Furthermore, recent solutions and technological developments aimed at addressing these challenges are outlined. Finally, the current state of seawater electrolysis is summarised, and its prospects are discussed, highlighting promising research directions that may enhance its viability in supporting the worldwide shift towards clean energy solutions.

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1. Introduction

The transition towards a sustainable energy economy is heavily dependent on the integration of renewable energy sources. However, the intermittent nature of resources such as wind and solar presents a major challenge to ensuring a reliable and



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continuous energy supply. To overcome this issue and meet the ever-increasing energy demands of society, efficient and scalable energy storage systems are required.¹ Electrochemical water splitting, specifically water electrolysis, offers a promising pathway for converting excess electricity produced by renewable energy into chemical energy in the form of hydrogen, which can be stored and utilised during periods of low electricity generation. Hydrogen production *via* water electrolysis serves as efficient energy storage while also being adaptable for use in diverse sectors, including households, transportation, and industry for the synthesis of valuable chemicals. Moreover, hydrogen holds the potential to play a critical role in combating climate change and achieving carbon neutrality, as its generation, consumption, and regeneration create a closed-loop cycle with minimal environmental impact. However, the associated costs, including investment, maintenance, and transportation, impose significant economic burdens and affect the viability and widespread adoption of hydrogen as a sustainable energy carrier.²

Renewable hydrogen, generated *via* water electrolysis driven by renewable energy, will establish novel interconnections between the energy and water sectors,³ and securing a sustainable water supply is fundamental to its success in transitioning the energy sector to meet global net-zero targets. Global demand for renewable hydrogen is expected to grow significantly over the coming decades. Utilising freshwater for electrolysis reduces the treatment requirements to reach the high quality needed by conventional electrolyzers; however, less than 1% of the water on earth exists as non-frozen freshwater.⁴ In countries with inland freshwater resources (*e.g.*, surface water and groundwater), such as Australia, hydrogen production through water electrolysis is feasible. However, the uneven distribution of water resources across the land presents a challenge for hydrogen production in inland regions.⁵ Additionally, many regions experiencing rapid growth in renewable energy are located along extensive coastlines, where the development of hydrogen production facilities may be challenged by limited access to freshwater resources.⁶

Freshwater availability is undergoing significant changes worldwide, with a general declining trend observed between 2001 and 2020, particularly in the southern hemisphere.^{7,8}



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Among the key drivers of this variability, precipitation has been identified as the dominant factor.⁸ Climate change is expected to intensify the global hydrological cycle, altering the spatial and temporal distribution of water resources and further constraining freshwater supplies.⁹ Projections indicate that river flows in regions such as South America and Australia will continue to decline under ongoing warming trends.⁸ In parallel, the past five decades have seen a dramatic rise in global groundwater extraction, largely driven by population growth, which heightens the risk of aquifer depletion and long-term water insecurity.¹⁰ The combination of multiple stressors on water availability such as population growth, industry, and climate change are highly challenging for water resource management. In this context, seawater electrolysis can potentially mitigate pressures on freshwater resources by utilising abundant ocean water, positioning it as a promising option for hydrogen production in regions with limited freshwater availability.

Commercially available water electrolysis systems commonly rely on freshwater resources for their water feedstocks. However, global freshwater systems increasingly threatened by the growing human population, rising water consumption, industrial development pressures, and anthropogenic climate change.¹¹ Recognising the need for alternative water sources, seawater as a limitless resource can be a viable approach to hydrogen production. Seawater electrolysis is still in its infancy compared to pure water electrolysis due to limitations raised by seawater characteristics and the complexity of electrochemical reactions. As a result, multiple methods, such as purification, desalination, and deionisation, are required to treat seawater. There are two approaches for seawater electrolysis (Fig. 1): direct seawater electrolysis and conventional electrolysis coupled with desalination. Each presenting several compelling arguments in its favour. Conventional electrolysis integrates external mature desalination technologies and pre-treatment plants with commercial water-electrolyzers. On the other hand, direct seawater electrolysis involves treating seawater inside the electrolyzers or consuming seawater with no pre-treatment.

Some studies suggest that conventional electrolysis is more practical because the cost of purifying seawater upfront is much lower than that of electrolysis. The initial stage of seawater treatment involves the removal of microorganisms and sediments through straightforward filtration methods.^{12–14} However, further research has shown that the water quality purified by seawater desalination technologies may not be adequate for direct use, and the soluble ion species remain within the seawater, entering the electrolysis system, which can harm the electrolyzers over time.¹⁵ Thus, it becomes imperative to consider the presence of these ion species during seawater electrolysis. Additionally, the chemical compounds and contaminants present in seawater progressively diminish the efficiency of the electrolyser.¹⁵ Therefore, additional purification steps are required to ensure high purity water feeds, significantly increasing the overall cost.^{13,14,16} Consequently, especially in large-scale offshore hydrogen production, bypassing the pre-treatment systems through direct seawater



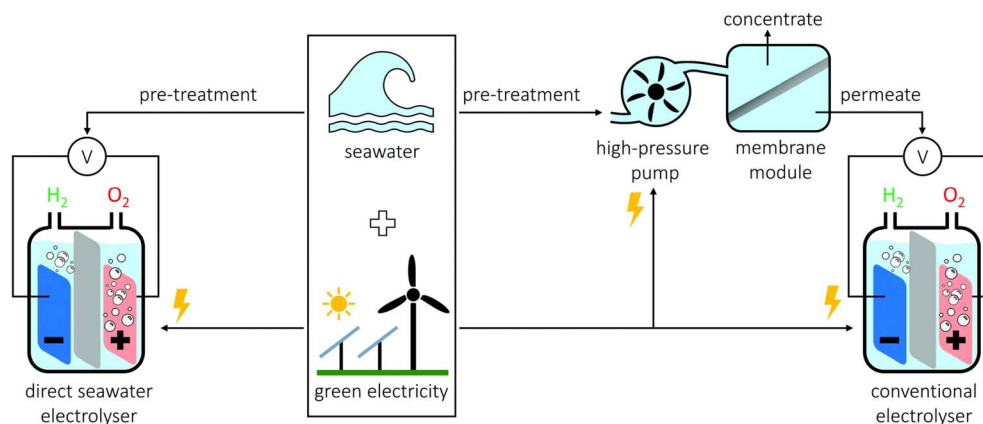


Fig. 1 Schematic of direct seawater electrolysis and conventional electrolysis coupled with desalination.⁴ Reproduced from ref. 4 with permission from the Royal Society of Chemistry, copyright 2021.

electrolysis offers tremendous potential to enhance economic viability regarding system design, reduced capital costs, minimised maintenance, and smaller physical footprints.

In summary, seawater electrolysis stands as a promising solution at the intersection of sustainable energy and water resource management. As global efforts intensify to address environmental challenges, further research, development, and implementation of seawater electrolysis technologies become imperative for a resilient and environmentally conscious future. This review mainly explores environmental impact, technical challenges and the state-of-the-art in direct seawater electrolysis in depth from various perspectives.

2. Environmental impact of seawater electrolysis

Physical and ecological features of the environment can be impacted during the construction, operation, and decommissioning of desalination and electrolysis plants through various causal pathways. The intensity of impacts depends on the degree of disturbance and the ecosystem's inherent sensitivity, varying based on the unique characteristics of the local habitat and communities.¹⁷ A high-level causal pathway for two drivers: conventional seawater electrolysis (combined desalination–electrolysis) and direct seawater electrolysis, has been developed for this review (Fig. 2).

The planned activities triggered by seawater electrolysis include several essential developments including electricity generation, earthworks/construction, desalination, electrolysis, and waste generation. These can lead to stressors, which are negative impacts on the environment resulting from activities. Stressors, in turn, cause a change in processes, which are natural mechanisms occurring in the environment, and these can affect endpoints, the marine and terrestrial ecosystems. In this review, potential impacts on the marine environment have been investigated.

2.1 Vegetation removal

Direct effects on coastal and marine systems may arise from habitat loss caused by vegetation clearance and seafloor disruption associated with constructing and operating seawater intake and outfall pipelines. In addition to marine ecosystems, terrestrial habitats are also affected, as vegetation is often cleared to accommodate desalination plants, electrolyser systems, pipeline corridors, and related infrastructure. These activities contribute to habitat fragmentation and loss, which can in turn lead to biodiversity decline.¹⁸ While the severity of these impacts varies depending on local ecological conditions, the degradation of natural habitats has been widely recognised as a key driver of biodiversity loss.¹⁸ The extent of such environmental impacts is closely linked to the physical footprint of the infrastructure involved.¹⁷ For example, the construction of the Alkimos seawater desalination plant in Western Australia is estimated to have caused the loss of approximately 8.3 hectares of reef, seagrass, and microalgal habitats,¹⁹ with largely irreversible effects on benthic communities.¹⁷ In this context, direct seawater electrolysis offers potential environmental advantages. By eliminating the need for large-scale desalination plants, it can substantially reduce land and marine area requirements, thereby minimising vegetation clearance and habitat disruption. This reduced physical footprint may help mitigate ecosystem fragmentation and the associated biodiversity impacts, particularly in sensitive coastal zones.

2.2 Atmospheric emissions

Emissions to the atmosphere from seawater desalination and electrolysis can include greenhouse gases (GHG), particularly CO₂, as well as chlorine (Cl₂) or oxygen (O₂), depending on the type of electrolysis applied.²⁰ Among these, CO₂ emissions are largely determined by the electricity source used to power various systems, including seawater intake and pre-treatment, reverse osmosis (RO), product water distribution, electrolyzers, and other balance-of-plant components.^{21,22} Electricity for these systems can come from onshore/offshore wind,





Fig. 2 Causal network of hydrogen production pathways for direct seawater electrolysis and conventional seawater electrolysis.

photovoltaic (PV) solar, or the local electricity grid, each carrying different emission intensities.²³ While seawater reverse osmosis (SWRO) is less energy-intensive than conventional water electrolysis (Table 1), direct seawater electrolysis offers a potential benefit in that it does not require additional energy input compared to current methods such as alkaline and PEM electrolysis.²⁴

Renewable energy such as wind or PV solar generates no CO₂ emissions compared to the Australian electricity grid mix in 2021–22, in which fossil fuels contributed 71% resulting in a total emission of 0.66 kg CO₂-eq. kWh. A comparison of different electricity mixes for seawater desalination²² found that emissions during the operation phase were 692 CO₂-eq. if using the Australian electricity mix and 26 CO₂-eq. if using 100% renewable energy over a 26-year period. Emissions for water electrolysis under two scenarios: COP26 (mostly solar, nuclear, and hydro energy) and EU27 (mostly coal, natural gas, nuclear, wind, and hydro) electricity mixes,³³ were estimated at 2.329 kg CO₂-eq. kg H₂ (COP26) and 22.964 kg CO₂-eq. kg H₂ (EU27) respectively. A separate study indicated CO₂-eq. emissions from grid-connected water electrolysis could be between 1.5–41.1 kg CO₂-eq. kg H₂.²³ The global warming potential of electrolysis powered by electricity from wind was estimated at 0.85 kg CO₂-eq. kg H₂ for alkaline electrolyser and 0.95 kg CO₂-eq. kg H₂ for PEM electrolyser.³⁴

The release of CO₂ into the atmosphere, if unmanaged, leads to the enhancement of the greenhouse effect and hastens climate warming. As more CO₂ dissolves in the ocean, seawater

pH decreases (becomes more acidic) due to the formation of carbonic acid and the release of hydrogen ions (H⁺).³⁵ Certain marine taxonomic groups are more vulnerable to acidification than others, as increased energy is required to form and sustain the exoskeletons of phytoplankton, crustaceans, mollusks, and corals under low pH conditions.³⁵ Under future acidification scenarios, the growth and survival of approximately half of benthic species could be reduced. Deoxygenation and hypoxia are also closely linked to ocean acidification. The effects of global warming, however, may differ between ocean basins due to differences in circulation patterns.

2.3 Seawater extraction

Stoichiometric calculations indicate that the production of 1 kg of hydrogen through electrolysis requires a minimum of 8.9 litres of water;³⁰ however, additional volumes are needed to meet the high purity requirement for the electrolyser, cooling of the system, and to account for system inefficiencies.³⁶ The recovery of water using reverse osmosis for desalination is around 40–50%³⁷ and electro-deionisation is around 90%.³⁸ This means that the required seawater intake volume to produce the 1 kg of hydrogen could be more than double the stoichiometric requirement. In addition, volumes for electrolyser cooling have been estimated at 88.1 L per kg H₂ for alkaline and PEM electrolysis, 152 L per kg H₂ for PEM electrolysis using evaporative cooling, and 645 L per kg H₂ for SOEC electrolysis.³⁰

Table 1 Energy consumption of seawater desalination, electrolysis, and seawater electrolysis methods

Process	Functional unit	Energy consumption	Reference
SWRO	kWh per m ³ desalinated water	2–7	4, 6, 14, 21, 23 and 25–28
SWRO	kWh per m ³ water	3.5–5	29
Alkaline	kWh per kg H ₂	47–66	30 and 31
PEM	kWh per kg H ₂	47–63	24, 27, 28, 30 and 31
SOEC	kWh per kg H ₂	42.3	30
Seawater electrolysis	kWh per kg H ₂	50–56.2	31 and 32
Electrolysis	kWh per kg H ₂	40–60	29



Impacts on the marine environment associated with seawater intake depend on the nature of the local environment at the intake area, the intake capacity, type, and structure.²⁵ Seawater is extracted from the ocean using either surface (open) intake or subsurface intake structures.²⁶ Surface intakes can be classified into three categories: offshore submerged intakes, nearshore submerged intakes, or nearshore surface intakes. In contrast, subsurface intakes can be categorised as onshore, utilising vertical wells, or offshore, employing wells or seabed infiltration galleries.²⁶

The extraction of large quantities of seawater may disrupt natural circulation of water, especially in regions with weak currents and waves.¹⁷ The method of extraction can result in the entrainment, entrapment, and/or impingement of marine organisms, thus reducing species population numbers which may impact on ecosystem productivity.^{17,26}

Entrainment occurs when small organisms with limited or no swimming capabilities are drawn into intake pipes, thereby being removed from the natural environment.¹⁷ Open seawater intake usually results in the entrainment and loss of fish eggs and larvae and benthic invertebrate species, algae and seagrass spores, phytoplankton and zooplankton, and other smaller marine organisms.¹⁷ The effects of entrainment can be substantial depending on local conditions. Impingement involves the physical confinement of marine organisms against intake screens due to high-flow conditions and the associated hydraulic forces. This typically results in injury, weakening, or even mortality, leading to a decrease in the local abundance and diversity of marine life.²⁵ Meanwhile, entrapment refers to the process where organisms are captured within the intake system without the ability to return to their natural environment, which may affect the overall diversity of the local marine ecosystem.²⁵

2.4 Waste and by-product disposal

The main waste and by-products from seawater electrolysis are (i) brines, (ii) chemicals added (additives) during pre-treatment of seawater and cleaning of RO membranes and (iii) inert solid waste including RO membranes.

2.4.1 Brines. Brine is the hypersaline byproduct generated from seawater electrolysis or desalination processes. Typically, seawater reverse osmosis (SWRO) systems achieve a recovery ratio between 40% and 50%,^{37,38} meaning that 50–60% of the original feedwater is concentrated into brine. In direct seawater electrolysis, the removal of water similarly leads to an increased concentration of solutes, producing brine with elevated salt levels. Beyond the high salt content, brine effluent may also contain chemical contaminants resulting from electrochemical reactions, such as alkalis or hypochlorites formed during seawater electrolysis.²⁰ This mixture of concentrated salts and potentially harmful chemicals complicates brine management.

Disposal of brine effluent represents a significant challenge due to both its environmental impact and the high costs involved.³⁸ While direct ocean discharge remains the most common practice,³⁹ alternative onshore disposal methods, including deep well injection or evaporation ponds, are hindered by factors such as substantial capital investment,

limited land availability, and stringent regulatory controls, as well as the inherent risks of soil and groundwater contamination. Offshore disposal *via* diffusers or coastal release attempts to mitigate some environmental risks, yet the long-term ecological consequences remain a concern. Emerging strategies emphasize transforming brine from a waste product into a resource, notably through ‘zero liquid discharge’ (ZLD) approaches.⁴⁰ These focus on recovering valuable salts and metals, offering economic incentives that could offset disposal costs. Additionally, innovative applications such as fish farming and halophyte cultivation leverage the unique chemical composition of brine to create sustainable bioproduct systems.³⁸ Despite these promising avenues, comprehensive assessments of environmental trade-offs and economic feasibility are crucial before widespread implementation can be achieved. Thus, managing brine not only poses a technical challenge but also an opportunity to rethink waste valorisation within seawater electrolysis frameworks.

The total dissolved solid (TDS) concentration of seawater is generally between 30 000–40 000 mg L⁻¹.³⁷ Seawater desalination plants typically produce brine which is 1.5 to 2 times more concentrated than ambient seawater (60 000–80 000 mg per L TDS).^{37,41} When brine is discharged into the ocean, it creates a high-density hypersaline plume that usually follows the gradient of the seabed.⁴² The dispersion and dilution of the brine plume are shaped by the discharge method (coastal or submarine outfall), the hydrodynamic characteristics of the environment (*e.g.*, currents, waves, and tides), and the bathymetry of the seabed, which makes the dispersal and impact extent location-specific.^{42,43} Variability in the flow can result in long-term chronic exposure at the plume centre as well as episodic short-term exposure of marine organisms to the waste brine at the perimeters of the plume.⁴⁰ Based on the aforementioned factors, a typical coastal desalination plant and its associated brine disposal system are illustrated in Fig. 3.⁴⁴

Marine organisms exist in an osmotic balance with their environment, and an increase in the concentration of salts can cause osmotic stress on a wide range of marine organisms.^{40,45,46} Osmotic imbalances from salinity increases may lead to irreversible cell dehydration (‘lethal osmotic shock’), reduced turgor pressure, and even death.^{14,47,48} However, different species show varying levels of sensitivity to increased salinity.⁴³ The scale and severity of impacts depends on factors such as species’ vulnerability and conservation status, the prevailing environmental conditions (*e.g.*, oceanography, climate, and bathymetry), and the chemical composition and disposal mechanism of the brine discharge.⁴⁹ Effects tend to worsen where water circulation is limited and for sessile organisms such as seagrass and benthic fauna. Seagrass, for instance, is known to be particularly sensitive to changes in salinity and may serve as a bioindicator for habitat degradation.⁵⁰ Numerous studies have investigated the effect of brine on seagrass species such as *Cymodocea nodosa*, *Posidonia oceanica*, and *Posidonia australis*.^{14,50} Laboratory experiments have demonstrated that salinity levels exceeding 39.1 g L⁻¹ reduce the vitality of *P. oceanica*, evidenced by decreased leaf growth, necrotic spots, and premature leaf senescence. The species *C. nodosa* showed





Fig. 3 Schematic illustrating a coastal desalination plant and its brine disposal system.⁴⁴ Reproduced from ref. 44 with permission from Elsevier, copyright 2018.

negative effects from increased salinity in both laboratory and *in situ* conditions. *P. australis* exhibited a broader tolerance range for salinity, from 27 to 60 g L⁻¹.¹⁴ Consequently, extensive studies advise that brine discharge should avoid seagrass ecosystems altogether or remain below a critical salinity threshold of 37.7 g L⁻¹.⁵⁰ Similarly, the abundance of Echinoderms has been used as an early warning of osmotic stress near the Javea facility in Spain.⁵⁰ Another study presents a seven-year evaluation of the impact of discharge from the Sydney Desalination Plant on fish communities near rocky reefs, showing a significant rise in fish assemblages at the outlet in comparison to reference locations.⁵¹

The discharge of the highly saline SWRO brine can also alter the abiotic marine environment by increasing the ambient ocean temperature, decreasing the dissolved oxygen content, altering the pH, and/or causing eutrophication.⁴⁷ Membrane-based desalination plants produce brine at a temperature close to that of the intake seawater, with reported temperature differences ranging from 1 to 2.5 °C.¹⁴ In the vicinity of the outlet of Australia's first desalination plant in Cockburn Sound, Western Australia, a notable reduction in dissolved oxygen (DO) levels was observed.⁵² Reduced dissolved oxygen availability may lead to changes in photosynthesis, metabolic, and growth rates.²⁵ Thermal stress and deoxygenation of receiving waters can occur in the case of thermal distillation. Rejected brine from saline electrolysis, containing precipitated Mg(OH)₂ and Ca(OH)₂, has a pH of >14 and, therefore a strong alkalinising effect on the receiving marine environment.²⁰

2.4.2 Additives. The hypersaline brine discharged into the ocean often contains residual chemicals introduced during the pre-treatment of seawater and cleaning of reverse osmosis (RO) membranes, which are essential steps to minimise scaling and fouling and to optimise water recovery.^{37,51} These chemicals typically include a variety of acids, coagulants such as FeCl₃ and Fe₂(SO₄)₃, flocculants like cationic polymers, disinfectants, sodium *meta*-bisulphite (Na₂S₂O₅), caustic soda, anti-scaling agents (including polyphosphates, phosphonates, and polycarbonic acids), as well as biocides.⁵³ The frequency of cleaning

is dependent on the quality of the seawater, efficiency of the pre-treatment process, and the anti-scaling agents used and can be as often as weekly for shock treatment where chemicals are applied periodically.⁴⁹ In the context of seawater electrolysis, pre-treatment often involves the use of hydroxide or phosphate buffers to selectively remove problematic ions.⁶ Additionally, periodic acid washing is necessary to control scaling caused by magnesium (Mg²⁺), calcium (Ca²⁺), sulphate (SO₄²⁻), and other impurities inherent in seawater.²⁰ While these chemical treatments are crucial for maintaining operational efficiency and prolonging equipment lifespan, their introduction into the brine effluent raises environmental concerns, particularly regarding the toxicity and accumulation of chemical additives in marine ecosystems. This underscores the need for developing more sustainable and less chemically intensive pre-treatment strategies to balance operational demands with ecological protection.

Sodium metabisulphite, used as a disinfectant in membrane-based desalination methods, can react with dissolved oxygen releasing sulphur dioxide (SO₂). This results in acidification and de-oxygenation (hypoxia) of the effluent.⁴⁹ Such conditions negatively impact the growth and survival of benthic marine organisms and may lead to changes in the composition and spatial arrangement of benthic habitats and communities. Anti-scaling agents containing organic phosphorus can enrich the effluent with nutrients stimulating plant growth. Coagulants are known to enhance water turbidity and reduce light penetration and/or lead to the burial of sessile benthic organisms.⁵⁴ Thermal desalination systems also employ anti-foaming agents and higher concentration of anti-biofouling additives, compared to membrane-based systems, which may pose risks to the life cycles of marine species.⁵⁵ Desalination, especially through thermal systems that experience higher corrosion rates, can lead to the release of heavy metals into effluent streams, contingent on the specific metallic alloys and their characteristics within the system.⁵⁶

In seawater electrolysis, chlorine Cl₂ (g) (0.23 kg per kg H₂) is produced at the anode as a by-product and released to the



atmosphere if no special measures are undertaken.²⁰ Chlorine is over twice as dense as air and tends to accumulate near its release point unless dispersed by air currents. Additionally, chlorine gas is highly soluble in water and exhibits significant reactivity.⁵⁷ Residual chlorine at low concentrations can be detrimental to aquatic life when the environment is heavily chlorinated. Research has demonstrated that marine invertebrates, like shrimps, hermit crabs, and amphipods, show particular sensitivity to chlorine exposure.⁵⁸ Chlorine undergoes hydrolysis to produce hydrochloric acid (HCl) and hypochlorous acid (HOCl), the latter being a powerful oxidising agent with sanitising properties. Chlorine also combines with organic compounds in seawater, leading to the creation of chlorinated and halogenated organic by-products, some of which could be harmful to aquatic life or even carcinogenic.⁵⁹

It's noteworthy that alkaline seawater electrolysis generates alkaline brine containing calcium and magnesium that can react with dissolved CO₂ in the ocean, providing a pathway for ocean CO₂ mineralisation. This approach not only reduces some brine waste but also aids in removing carbon dioxide from the ocean, which has positive environmental implications.

2.4.3 Inert solid waste. The periodic replacement of RO membranes contributes significantly to solid waste generation.²⁶ On average, approximately 100 RO modules are required to produce 1000 m³ of desalinated water, which each module having a typical lifespan of 5–8 years.⁶⁰ At the end of their service life, disposal options for these modules include landfill (waste burial), incineration, syngas combustion, electric arc furnace processing, material recycling, reuse as ultrafiltration modules, or direct reuse.⁶⁰ In practice, disposal to municipal landfills remains the most adopted method in Australia. While the polymeric composition of RO membranes renders them non-toxic, the environmental impacts associated with their disposal primarily stem from land use and the carbon footprint of transportation.⁶⁰ In comparison, electrolyser systems also have finite lifespans, though their solid waste profiles differ. Electrolyser stack lifetimes vary by technology—less than 20 000 hours for proton exchange membrane (PEM), less than 40 000 hours for solid oxide electrolyser cells (SOEC), and up to 90 000 hours for alkaline electrolysers—while overall system lifespans range from 10–20 years for PEM to 20–30 years for alkaline types.⁶¹ However, unlike RO modules, electrolysers do not require frequent large-scale physical replacements of polymer-based units.

From a life-cycle waste perspective, direct seawater electrolysis may offer advantages by avoiding the substantial volume of waste associated with RO membrane disposal. This distinction highlights another potential environmental benefit of direct electrolysis pathways, particularly in long-term sustainability assessments.

3. Challenges of seawater electrolysis

Seawater electrolysis presents a significant challenge due to the complex chemical composition of seawater. Seawater contains both inorganic and organic compounds, including dissolved salt ions (Cl⁻, Na⁺, K⁺, Mg²⁺, Ca²⁺), sediments containing heavy

metals (Sr²⁺, Cd²⁺, Pb²⁺, Cu²⁺ and Fe^{2+/3+}), sulphates (SO₄²⁻), carbonates (CO₃²⁻ and HCO₃⁻) in deep seawater, CO₂ in surface seawater, dissolved F⁻, N⁻ and B⁻ anions,^{62,63} as well as micro-organism species including bacteria, fungi and microalgae.⁶⁴ Moreover, the characteristics of seawater, such as composition, salinity, and pH, vary depending on the depth, geographical locations, and seasonal fluctuations.^{62,63} These property fluctuations necessitate adapted approaches for seawater electrolysis, as the electrolytic parameters must be constantly adjusted to accommodate these variations. The complex relationship of these factors underscores the complexity of seawater electrolysis, making it imperative for researchers and engineers to develop innovative solutions to harness the potential of this abundant resource for sustainable energy production.⁶⁵ Herein, the key challenges of direct seawater electrolysis are presented.

3.1 Conductivity

Although seawater contains various ionic compounds, its salinity and ionic conductivity are low, which results in low electrical conductivity for seawater. This low electrical conductivity decreases the electrolyser efficiency by increasing the cell voltage input and energy consumption.^{66,67}

3.2 pH

While the pH of seawater changes depending on the geographical locations and seasonal changes, another seawater splitting challenge is the variation of local pH at the cathode and anode during electrolysis. By increasing the potential, the local pH increases at the cathode during HER and reduces at the anode during OER. Generally, the bulk pH of seawater is close to neutral, about 8.⁶² However, the local pH variation at electrodes can be between 5 to 9 pH units, which causes electrode degradation and reduces their stability and efficiency. Although the carbonates in seawater can act as buffers during the electrolysis, their performance is not sufficient to prevent fluctuations in the local pH at electrodes.⁶⁸ The local pH increases at the cathode results in precipitation of metal hydroxides such as Mg(OH)₂ (at pH 10.7–11) and Ca(OH)₂ as well as Ca(CO₃)₃ on the cathode surface and block the cathode active sites,^{69,70} consequently reducing the HER efficiency of the cathode. Moreover, at the specific potentials, the soluble metal cations, including Cu, Pb and Cd, will electrodeposit on the cathode surface and block active sites for HER. These metal depositions not only compete with HER but also reduce the efficiency and durability of the cathode.

3.3 Chloride

The presence of electrochemical active Cl⁻ anions in seawater is another key challenge in seawater electrolysis. Chloride electro-oxidation chemistry in aqueous solutions is a complex mechanism that tends to compete with oxygen evolution reaction (OER) and is influenced by several variables, including pH, applied potentials, and temperature. At a temperature of 25 °C and a fixed total 0.5 M concentration of chlorine species, which is representative of seawater conditions, a Pourbaix diagram was constructed to depict the prevailing reactions under





Fig. 4 The Pourbaix diagram displays (a) chloride chemistry and (b) the oxygen evolution reaction (OER) in an aqueous saline electrolyte. The red line corresponds to the acidic oxidation of chloride to Cl_2 gas, while the green line represents the thermodynamic equilibrium between H_2O and O_2 . The black and purple lines indicate the onset of chloride oxidation to HOCl or ClO^- .^{71,72} Reproduced from ref. 71 with permission from Wiley-VCH, copyright 2016 and from ref. 72 with permission from Elsevier, copyright 2022.

different pH conditions (Fig. 4).⁷¹ When the pH drops below 3.0, under acidic conditions, the free chlorine evolution reaction (CLER) becomes the dominant chloride oxidation process. As the pH increases to the range of 3 to 7.5, hypochlorous acid (HOCl) formation becomes the major reaction, while hypochlorite (ClO^-) forms at pH values exceeding 7.5, corresponding to the $\text{p}K_a$ of hypochlorous acid.⁷¹ The presence of chlorine dissolved in water and the disproportionation of hypochlorite ions under higher temperatures add further complexity to the chemistry of chlorine species.

Thermodynamically, the chloride oxidation reactions are less favourable than the OER. The potential difference between these reactions and the OER grows with increasing pH, eventually reaching a maximum value near 480 mV when hypochlorite formation starts, at which point it stabilises.⁷¹ This implies that under alkaline conditions, by maximising the potential difference between the OER and the CLER, the formation of toxic chlorine compounds can be effectively suppressed, which is known as the “alkaline design criterion” in saline water electrolysis. Furthermore, the chloride reactions involve two electron transfers, whereas the OER involves four electrons. This discrepancy in the number of electrons participating in the OER reaction mechanisms contributes to the commonly observed higher overpotential for the OER compared to chloride oxidation, making the OER kinetically unfavourable. Consequently, the development of highly selective anode catalysts is crucial to prevent the evolution of chlorine gas during seawater electrolysis.^{73–75}

Moreover, the presence of Cl^- anions, along with the production of harmful Cl_2 gas and hypochlorite, degrades catalysts and electrode materials while also corroding other components of the electrolyser through the formation of metal chloride and hydroxide compounds. This results in reduced efficiency and stability of the electrolysis process, ultimately shortening the lifespan of the electrolyser.

3.4 Organic and biological impurities

Organic and biological impurities in seawater significantly impact the efficiency and longevity of water electrolysers.

During electrolysis, they can lead to various issues such as the production of undesired compounds, increased gas impurity, reduced performance, membrane fouling, and even the normal operation of electrolysers.¹⁵ Organic compounds, for instance, might not contribute to conductivity but can disrupt the electrolysis process. Biological impurities introduce complexities in electrochemical water splitting and can cause corrosion and affect the overall functionality of electrolysers. Therefore, considering the impact of these impurities, it is essential to design electrolyser components for the efficient functioning of seawater electrolysis, ensuring consistent and sustainable hydrogen production.

4. Potential solutions and recent advancements

In addressing the challenges faced in direct seawater electrolysis, researchers have adopted a multitude of strategies aimed at overcoming these obstacles. A pivotal focus of these efforts has been the examination and modification of key components of conventional water electrolysis systems for their applicability in seawater electrolysis. The primary components under study are electrodes, electrolytes, and membranes, each of which plays a crucial role in the electrolytic process. In addition to these fundamental components, researchers have explored innovative solutions and developed hybrid systems that combine different electrolysis technologies to optimise the efficiency and stability of seawater electrolysis. Fig. 5 represents a summary of these key concepts and innovations. The following sections examine the progress made in addressing the issues associated with seawater electrolysis.

4.1 Electrodes

Researchers have investigated the enhancement of electrode materials and designs to improve their catalytic activity and stability in seawater conditions. Noble metal-based electrodes have been a common choice due to their high catalytic activity and corrosion resistivity, but efforts have been made to replace them with transition metal-based counterparts, which are more





Fig. 5 The key concepts in seawater electrolysis research and development include three main areas: electrode design, electrolyte improvement, and membrane innovation.

cost-effective.⁷⁶ Approaches such as improving electrical conductivity, increasing the number of active sites, reducing the reaction energy barrier, and developing protective layers have been investigated to enable efficient electrolysis in the challenging seawater environment.⁶⁵ Some representative methods of electrode design are highlighted in Fig. 5.

4.1.1 Enhancing electrical conductivity. The low conductivity of seawater affects the electrocatalytic activity. Various studies have been conducted to improve the conductivity to enhance the electron transfer and facilitate the kinetic reactions.^{77,78} In this regard, both electrocatalysts and substrates were studied.

One of the investigated approaches to enhance the electronic properties of electrocatalysts is applying conductive supports/substrates. The structure, conductivity and wettability of substrates can significantly affect the electrocatalytic performance of catalysts. Studies showed that substrates with 3D structures (e.g. Ni/Co/Cu foams and carbon cloth/papers) have demonstrated better performance compared to flat surfaces, such as glassy carbon electrodes. It was shown that *in situ* synthesis of catalysts on substrates with 3D structures facilitates the electron and mass transfer, increase electrochemical active surface area, and provides better gas diffusion. It is noteworthy to say that the dispersion of catalysts on the substrate can affect its electrochemical performance. Uniform and mono dispersion of catalytic materials on substrates can increase the active sites, surface area and electrochemical performance while it results in less materials being used. For instance, nano-dispersion of noble metals on carbon-based substrates has been introduced as an efficient electrode for seawater electrolysis. These electrodes provide a large surface area and exposure to active sites due to the presence of noble metals. Furthermore, the carbon-

based substrates protect the noble metals from corrosion in seawater electrolysis.^{79,80} This type of catalyst not only enhances the conductivity and performance of OER/HER but also provides good durability in seawater due to its excellent mechanical stability and structural diversity. Moreover, other carbon-based materials and metal-based materials have been studied as suitable conductive supports for HER catalysts to enhance electron transfer.^{66,81–85} It should be noted that conductive supports do not alter the intrinsic catalytic activity but rather improve the electron transfer and increase catalytic surface area.⁸⁶

Another approach is to develop highly active electrocatalysts to enhance the electrical conductivity and increase seawater electrolysis performance. Researchers implement different methods to design electrocatalysts with excellent activity. One of the studied methods to enhance electrical conductivity involves the strategic design of heterostructures composed of two or more active components, which enable rapid charge transfer due to their optimised interfacial structure.^{87–89} Heteroatom doping of catalytic materials serves as an alternative way to adjust the electronic structure and enhance conductivity.⁹⁰ Furthermore, rational defect engineering is another way to optimise electrical conductivity.⁹¹

Overall, the high electrical conductivity is crucial for accelerating electron, facilitating gas production, and improving electrocatalytic activity. Over the last decades, many electrocatalysts have been developed with high faradaic efficiency for use in pure water electrolyzers that can be investigated for seawater electrolysis.

4.1.2 Improving activity and selectivity. Increasing the accessible active site on the catalyst surface can improve the electrocatalytic performance for seawater electrolysis while reducing the impact of impurities (e.g. bacteria), insoluble materials (e.g., Ca(OH)₂ and Mg(OH)₂), and complex ionic environment on the active sites of the catalyst.⁹² One of the ordinary methods to increase the active sites is to develop catalysts with large surface areas to enhance electrolyte diffusion increase the catalytic process.^{93–96} Another approach is to develop a catalyst with more active sites that expose higher OER selectivity compared to CLER.^{97–102} Moreover, conventional methods of defects engineering and more novel methods, such as *in situ* chemical conversion routes, can be applied to develop electrocatalysts that exhibit highly active sites and significantly improve the electrocatalytic activity in seawater electrolysis.^{75,103}

Another parameter that can improve the electrochemical performance is improving the reaction energy barriers. For both HER and OER, the electrocatalytic performance is determined by the free energy of adsorption and desorption of hydrogen and oxygen intermediates on the catalyst surface. The reaction energy barriers of the catalysts can be optimised by improving the electronic structure of the catalyst's surface. Researchers have studied various methods to reduce the energy barriers, such as interface and defect engineering phase transition and atom doping.^{88,90,99,104–108}

The creation of active sites is a crucial factor in developing highly efficient electrocatalysts, which can be achieved through two main approaches: (a) increasing the number of active sites



and (b) enhancing their intrinsic activity to optimise reaction energy barriers. While most research in this field focuses on designing electrocatalysts with a greater number of active sites, the effects of impurities or hybrid materials on their deactivation have received little attention. Therefore, identifying effective strategies or pre-treatment methods to mitigate the impact of seawater impurities on electrocatalytic active sites is essential.

4.1.3 Designing protective layer. In addition to high electrical conductivity, catalytic activity, and selectivity, electrocatalyst designed for seawater electrolysis should also demonstrate long-term stability and high resistance to Cl_2 , ClO^- , and other ions present in seawater.^{109,110} Tailoring preferential adsorption of oxygen-containing intermediates,¹¹¹ creating perm-selective layers,^{82,112,113} and altering electrode architectures^{114–116} are common techniques for inhibiting chloride corrosion.^{102,117} These approaches often necessitate complex structural engineering, with just a few succeeds to date in meeting the long-term stability criterion.^{82,96,118,119} Some of the introduced protecting layers on the catalyst surface include metal oxide layers (e.g., MnO_x ,¹¹² SiO_x ,¹¹³ CeO_x ¹²⁰ and TiO_2 ⁸⁸), cationic selective polyanion (e.g., NiFe/NiS_x ,¹²¹ $\text{Ni}_m\text{Co}_n\text{S}_x$ ¹²² and MoS_2 ⁸⁸), small organic molecules (SOMs)¹²³ and carbon coating (e.g., graphene oxide⁸²) that can provide superior resistance to chloride corrosion and increase the stability of electrocatalysts.¹²⁴ Moreover, researchers have employed structural buffer engineering to develop a catalyst that exhibits high stability and selectivity for OER. The Cl^- within the lattice structure of these catalysts acts as a structural buffer, and its gradual leaching during the OER process creates vacancies that allow seawater Cl^- to enter, preventing catalyst deactivation.¹²⁵ Recently, the intercalation of anions, sulphates, phosphate, carbonate, and selenide ions, within the structure of catalysts has been drawing attention due to their high chloride rippling ability.^{126,127}

4.1.4 Creating alkaline design criterion. Although seawater contains carbonates, they are not strong buffered enough to avoid changes in local pH during the electrolysis. On the other hand, in seawater electrolysis, CIER kinetically is more favourable than OER. However, seawater electrolysis in alkaline conditions maximise the potential difference between OER and CIER. Therefore, an alkaline design criterion has been introduced to provide overpotentials of catalysts lower than ~ 480 mV at the desired current density. In general, achieving the desired current densities below a 480 mV overpotential is challenging under alkaline conditions. However, studies have demonstrated that electrocatalysts, such as transition metal-based oxides, oxyhydroxides, and layered double hydroxides, can deliver current densities near the required levels for commercial-scale electrolyzers at potentials under 480 mV.^{71,128–134}

4.2 Electrolytes

The performance, hydrogen purity and longevity of commercial water electrolysis systems are widely recognised to be influenced by the quality of the water feed. Impurities in the electrolyte feed can detrimentally impact electrolyser efficiency and

the overall quality of produced hydrogen, as well as contribute to device degradation over time. Two different sources of impurities in water electrolysis processes are identified: exogenous and endogenous.¹⁵ Exogenous impurities are those introduced by the feed water, including ionic and organic impurities as well as dissolved gases (e.g. nitrogen, oxygen, argon and carbon dioxide). Endogenous impurities are generated *in situ* by degradation processes within the electrolyser and total plant itself. Impurities may also be introduced by the electrolyser due to leaching from its components and contamination during manufacturing and maintenance. Like freshwater electrolysis, seawater electrolysis struggles with both types of impurities. However, the complex composition of seawater makes the scenario more complicated. This section focuses specifically on exogenous impurities in the electrolyte, and some representative methods are listed, as shown in Fig. 5.

4.2.1 Applying additives. As mentioned earlier, the conductivity of seawater is low, which results in significant potential losses. On the other hand, the presence of diverse contaminants in natural seawater, primarily consisting of solid particles, bacteria, and organic matter, along with various ions such as Ca^{2+} , Mg^{2+} , Sr^{2+} , Cl^- , Br^- , and others, poses significant challenges to the effective and sustainable production of hydrogen through seawater electrolysis. The electrodes and membranes are integral components of the electrolyser, but they are also sensitive to contamination, corrosion, or poisoning caused by contaminants. Due to the low concentration of H^+/OH^- in natural seawater, higher current densities result in increased cathode precipitation, characterised by a milky appearance due to a decrease in pH. Additionally, anode corrosion is intensified by the presence of halide ions and other related species. The direct electrolysis of natural seawater is considered an optimal approach. However, due to the close-to-neutral pH and the abundance of halide ions compared to hydroxide ions, the selectivity of oxygen would be constrained. Moreover, during the electrolysis, the change in local pH at electrodes affects the thermodynamics and hydrogen and oxygen half-reactions at the cathode and anode, respectively (Fig. 6). Few studies covered the improvement of electrolyte conductivity. However, understanding the electrolyte conductivity is also crucial for improving electron transport and enhancing electrocatalytic performance.

In order to overcome the low conductivity of seawater and prevent local pH gradient in seawater electrolysis, a possible effective strategy can be applying additives to the electrolyte. Several research have used buffered seawater to get improved results. For instance, 1 M KOH + 1 M phosphate (KPi) + 1 M borate (KBi)¹³⁶ or NaH_2PO_4 + Na_2HPO_4 ¹³⁷ were added to natural seawater to control the pH during seawater electrolysis. While the utilisation of buffering seawater electrolytes may result in reduced pH gradients across the electrolytic cell and improved catalyst performance, the process of adding salts to seawater and then filtering them presents two minor challenges. Firstly, the process of direct seawater splitting has been modified to an indirect approach involving pre-treatment of seawater to reduce the concentration of ions. Secondly, regular addition of salts is necessary to maintain a steady reaction. These concerns are





Fig. 6 Schematic representations of (a) the creation of an alkaline microenvironment on anode with Lewis acid modification, promoting OER and suppressing chlorine chemistry, and (b) the formation of alkaline microenvironment on cathode with Lewis acid modification, aiding HER and preventing precipitation. EDL = electrical double layer, LA = Lewis acid site, E = external electric field.¹³⁵ Reproduced from ref. 135 with permission from Springer Nature, copyright 2023.

more intensified in the case of alkaline seawater, as the process of alkalisating seawater necessitates the use of KOH prior to electrolysis. Although several methods involve the use of more affordable salts such as Na_2CO_3 to eliminate $\text{Ca}^{2+}/\text{Mg}^{2+}$ cations, followed by the addition of NaOH/KOH to induce alkalinity, these pre-treatment steps effectively convert the process from direct electrolysis of natural seawater to indirect electrolysis of artificially alkalisated seawater.¹³⁸

4.2.2 Alkaline/acidic electrolyte. Current research efforts aim to identify materials capable of achieving high current densities at overpotential below 480 mV for catalytic reactions in alkaline seawater with a pH greater than 7.5, known as the “alkaline criterion”.¹³⁹ The rationale behind this strategy is that alkaline media can effectively limit chlorine electrochemistry at certain potentials due to the presence of preference for excessive OH^- ions around the anode, which thermodynamically promotes OER and inhibits ClOR.¹⁴⁰ Although the development of catalysts with the desired properties is feasible and has shown significant advancements, as mentioned earlier, the introduction of artificial alkylation to seawater introduces additional processes and energy costs that need to be considered in commercial settings. Moreover, it is important to consider that while the commercial electrolyzers of water commonly consume an electrolyte containing around 30% KOH, the electrocatalysts in alkaline seawater may experience faster degradation if the solution becomes too alkaline (e.g. in electrolysis of seawater + 6 M KOH).¹³⁷ Furthermore, in alkaline conditions, organic impurities are oxidised at the anode, generating various products such as formates and acetates that affect catalyst activity and electrode corrosion.¹⁵ Further research is crucial to comprehensively explore these impurity impacts.

An alternative approach is the process of electrolysis in acidic seawater that offers certain advantages, such as reduced precipitation of cations on electrodes and enhanced efficiency in HER. However, this method also introduces challenges, including unlimited chloride oxidation, acid etching and intensive corrosion. Currently, there is few research investigating acidic seawater electrolysis.^{79,135} Although direct electrolysis in acidic electrolytes offers a simpler process, there are

now more severe criteria for both electrodes and membranes. In a PEM electrolyser system, cationic impurities, substitute protons within the membrane, diminishing ionic conductivity and hindering electrochemical reactions. Organics often adsorb on the catalyst, impeding reactions. For example, those with $-\text{OH}$ and $-\text{CN}$ functional groups cause metal leaching, and some, like methanol, oxidise at the anode, but gaseous products can poison the cathode.¹⁵ It is evident that the cost-effectiveness of direct electrolysis compared to conventional electrolysis, using an acid electrolyte, is possible only when both electrode and membrane exhibit sufficient durability. Therefore, more comprehensive research is needed to evaluate the impact of these impurities on direct seawater electrolysis under acidic conditions.

4.2.3 Asymmetric electrolyte design. Although the traditional electrolyser systems (Fig. 7) enable to improve the OER selectivity *versus* ClOR, a significant cell voltage is still necessary, and the achieved current density is limited (i.e. $<200 \text{ mA cm}^{-2}$).¹⁴¹ The process will ultimately result in the occurrence of chloride electro-oxidation chemistry at the anode, leading to corrosion over extended periods of electrolysis and a substantial increase in energy consumption. Moreover, in these systems, not only the hydrogen production efficiency is low, but also a lot of additional chemicals/additives are consumed. Therefore, designing unconventional seawater electrolysis systems has significant potential for the cost-effective generation of hydrogen.

In this regard, another approach is developing electrolyzers with asymmetric electrolyte feed instead of using a symmetric electrolyte feed that removes the limitation of dependency on additives. A novel electrolyser system was tested with asymmetric electrolyte feed using synthetic seawater (0.5 M NaCl) as electrolyte for the cathode compartment and KOH solution (0.5 M KOH) for the anode compartment (Fig. 8).¹⁴² It was reported that this electrolysis system reduces the cost of electrolyte by removing the need for additives to the electrolyte, alleviates anode corrosion by improving OER selectivity and operates at high current density, triggering ClOR, consequently increasing the efficiency.





Fig. 7 Flowsheets illustrating the (a) suggested process for chlorine-free seawater electrolysis to produce H₂ and O₂, and (b) laboratory setup used for continuous electrolysis experiments.¹⁴¹ Reproduced from ref. 141 with permission from Elsevier, copyright 2018.

4.2.4 Organic molecules in electrolyte. Another strategy is replacing OER with the conversion of organic molecules that oxidised at the anode to produce high value-added compounds, such as CO₂, or N₂.^{143–150} The electrolyte used in this method involves the substitution of seawater solution with a reductive chemical saltwater solution, such as hydrazine, sulphion and urea. An energy-efficient hybrid seawater electrolyser was developed where 1 M KOH seawater solution was employed as the catholyte, and a 1 M KOH + 0.5 M hydrazine solution was utilised as the anolyte⁸¹ (Fig. 9). At the cathode, seawater splitting results in the separation of H₂ and OH⁻ ions. Subsequently, the OH⁻ ions are directed towards the anode, where they

facilitate the oxidation of hydrazine into chemically unreactive N₂ gas. The hybrid seawater electrolyser demonstrates an exceptionally low cell voltage range of 0.7–1 V while operating at a current density of 500 mA cm⁻². In this system, the H₂ production rate can reach 9.2 mol h⁻¹ g⁻¹ (of catalyst) at a current density of 500 mA cm⁻² for 140 hours, and the energy consumption is reduced by 30–50% when compared with commercial alkaline electrolyzers.

In another study, a combination of water electrolysis and sulphion degradation was reported (Fig. 10).¹⁵¹ In comparison to OER, the cell voltage for sulphion oxidation is decreased by a range of 1.3 to 1.4 V. This reduction effectively mitigates the



Fig. 8 Schemes for anion exchange membrane (AEM) electrolyzers utilising separate electrolyte feeds with varying compositions: (a) symmetric alkaline feeds, (b) asymmetric alkaline catholyte, (c) asymmetric KOH catholyte and NaCl anolyte, (d) symmetric KOH + NaCl feed, (e) asymmetric NaCl catholyte, (f) asymmetric NaCl catholyte and KOH anolyte.¹⁴² Reproduced from ref. 142 with permission from the Royal Society of Chemistry, copyright 2020.

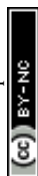




Fig. 9 (a) The Pourbaix diagram for OER, HER, HzOR, and ClOR in synthetic seawater containing 0.5 M Cl^- at pH values ranging from 7 to 14. (b) Schematic illustration of hybrid seawater electrolysis design (HSE) (top) vs. alkaline seawater electrolysis design (ASE) (bottom).⁸¹ Reproduced from ref. 81 with permission from Springer Nature, copyright 2021.

issues of corrosion and competition of ClER in the process of seawater electrolysis. The hydrogen production rate in this study was reported as $5.34 \text{ mol h}^{-1} \text{ g}^{-1}$ (of catalyst) at a current density of 300 mA cm^{-2} , and the energy consumption exhibits a reduction of 60% in comparison to the alkaline water electrolyzers. Since urea oxidation is thermodynamically more favourable than OER, urea sources, such as wastewater streams containing high concentrations of urea, can be potential electrolyte feed for hydrogen production from seawater.^{152,153}

The performance of Ru/P-NiMoO₄ catalyst with Ni foam support was evaluated as a bifunctional electrocatalyst for HER in alkaline seawater electrolyte and urea oxidation reaction (Fig. 11).¹⁵⁴ This system required 1.73 V cell voltage at a current density of 500 mA cm^{-2} for overall urea splitting, and H₂ production was stable for 145 h operating at a current density of 100 mA cm^{-2} .

4.3 Membrane

As discussed earlier, during water electrolysis, fluctuations in local pH levels can result in an acidic environment at the anode

and an alkaline environment at the cathode, which directly impacts the electrode potential and reaction rate. At the cathode, in alkaline conditions, insoluble deposits like Mg(OH)₂, Ca(OH)₂, and MgO are more likely to form on the electrode surface. Notably, these blockages are not limited to the cathode but also affect the anode, as well as the porous diaphragm or membrane. In PEM electrolyzers, metal ions may occupy the proton exchange sites of the membrane, increasing the resistance to charge transfer and slowing down the reaction. In a similar manner, Cl⁻ ions can hinder the ion exchange efficiency within the anion exchange membrane of AEM electrolyzers. Moreover, solid impurities and microbial contaminations, including bacteria and microbes, lead to precipitation and adhesion on the electrode and membrane surfaces. These factors further impede the transfer of charge and diffusion of water during the electrolysis process and affect the efficiency of seawater electrolysis. Consequently, in designing electrolyzers for direct seawater splitting, it is crucial to develop membranes that effectively prevent impurities from reaching the electrodes and also minimise membrane fouling caused by these



Fig. 10 (a) Pourbaix diagram of SOR, HER, OER, and ClOR under alkaline conditions. Below pH 13, the polysulfides in the aqueous solution are not stable. (b) Schematic illustration of hydrogen production at cell voltages below 1.0 V by coupling seawater reduction with SOR on Co-based electrocatalysts.¹⁵¹ Reproduced from ref. 151 with permission from Wiley-VCH, copyright 2021.





Fig. 11 Schematic illustration of renewable-powered electrolyser using seawater and industrial urea sewage as the feeds.¹⁵⁴ Reproduced from ref. 154 with permission from Elsevier, copyright 2023.

impurities. Some innovative works on membrane technology are listed as shown in Fig. 5.

4.3.1 Reverse osmosis membrane. Cost-effective reverse osmosis (RO) membranes were evaluated for seawater electrolysis as a substitute for the expensive proton exchange membranes (Fig. 12).¹⁵⁵ This approach presented a viable means of substantially decreasing the cost of water electrolysis membranes. The research findings indicate that the RO membrane has a high degree of selectivity towards protons in cation salts, particularly when employed in high concentration electrolytes. In comparison to ion exchange membranes, the cost of reverse osmosis (RO) membranes presents a compelling motivation for investigating their potential use in water electrolysis systems.

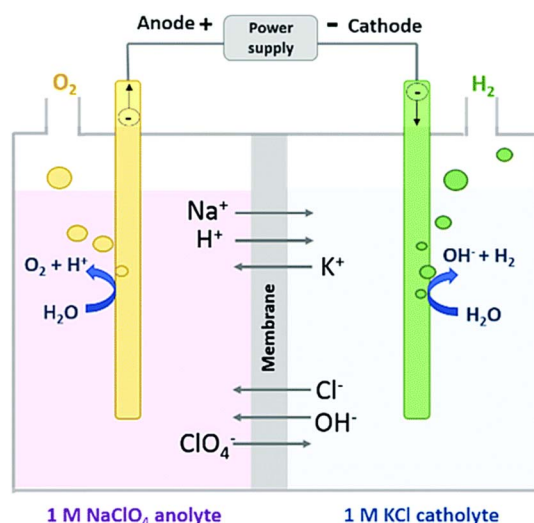
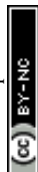


Fig. 12 Schematic illustrating ion transfer under a constant current, with 1 M NaClO_4 for the anolyte and 1 M KCl for the catholyte instead of NaCl to demonstrate cation transport under varying conditions.¹⁵⁵ Reproduced from ref. 155 with permission from the Royal Society of Chemistry, copyright 2020.

4.3.2 Forward osmosis membrane. Another study introduced a novel method of direct seawater electrolysis by integrating a forward osmosis (FO) membrane with water splitting (Fig. 13).¹⁵⁶ This process maintains a balance between water inflow and outflow rates, enabling continuous extraction of water from impure sources *via* the FO membrane. However, FO and RO membranes lack perfect selectivity, allowing chloride ions (Cl^-) to pass and undergo oxidation at the anode. This oxidation can lead to the formation of harmful byproducts like chlorine gas (Cl_2) and chlorate/perchlorate ions. To address this, supplementary procedures are needed to eliminate these byproducts from seawater electrolysis. Consequently, further adjustments were made by incorporating a cost-effective semi-permeable membrane between the electrodes, effectively lowering expenses and limiting chloride ion oxidation.¹⁵⁶ Further studies showed that the use of transition metal-based components allows for continuous water electrolysis over 5 days with a relatively low overpotential of 125 mV and no decrease in efficiency.



Fig. 13 Schematics illustrating (a) a forward-osmosis electrolyser that addresses the challenge associated with precipitation of $\text{Ca}(\text{OH})_2/\text{Mg}(\text{OH})_2$, and (b) the issue of operating an unprotected cathode and $\text{Ca}(\text{OH})_2/\text{Mg}(\text{OH})_2$ precipitation challenge in a conventional cell design for seawater electrolysis.¹⁵⁶ Reproduced from ref. 156 with permission from American Chemical Society, copyright 2022.



4.3.3 Waterproof breathable membrane. Recently, a novel membrane-based seawater electrolysis technology has been developed, eliminating the need for pretreatment processes like desalination.³² A commercial alkaline electrolyser integrated with a self-dampening electrolyte (SDE) and a hydrophobic porous polytetrafluoroethylene (PTFE) membrane. This setup enabled continuous H₂ production from seawater for over 72 hours at 400 mA cm⁻², with a scaled-up version generating 386 L of H₂ per hour operating stably for over 3200 hours at 250 mA cm⁻². The technology relies on the difference in water vapour pressure between seawater and SDE to promote seawater evaporation and water vapour diffusion. The vapour is then re-liquefied after adsorption on SDE, establishing a thermodynamic equilibrium. This innovative approach ensures a reliable and steady water supply for freshwater electrolysis *via* a “liquid-gas-liquid” mechanism. Featuring low energy requirements, high efficiency, and exceptional stability, this direct seawater electrolysis technology has the potential to make significant strides in the field, rendering pretreatment processes unnecessary and fostering the advancement of direct seawater electrolysis. While both PEM and AEM are utilised in certain aspects of electrolyser technology, electrochemical data primarily stem from a porous diaphragm. This self-driven purification technology faces two main challenges. Firstly, the use of a porous diaphragm often results in decreased hydrogen purity at the cathode and substantial crossover to the anode, which diminishes overall hydrogen production efficiency. Secondly, it is essential to ensure that seawater purification speeds align with electrolysis rates, particularly in long-term, high-current production scenarios. Achieving a balance between seawater purification rate and electrolysis rate forms the foundation for the continuous operation of this technology, necessitating advancements in purification rate enhancement and control over the electrolysis rate.

4.3.4 Bipolar membrane. The bipolar membrane (BPM) is an advanced ion-exchange membrane that comprises two layers: an anion exchange layer (AEL), which permits the selective movement of anions, and a cation exchange layer (CEL), which allows the selective passage of cations.^{157,158} BPM has been widely utilised in several applications, including water electrolysis, acid–base synthesis, electrodialysis, and CO₂ reduction.^{159–161} When comparing monopolar membranes (*e.g.* PEMWE) to bipolar membrane technology (BPM), it becomes evident that BPM possesses numerous notable benefits (Fig. 14). The distinct ionic selectivity of AEL and CEL enables the BPM to effectively link two separate reactions, while simultaneously reducing the interference of opposing ions in a single reaction.^{153,162} Additionally, the reduction in electrolyte ion crossover enhances the ability to collect and separate products and reactants. Since 2022, several studies have documented the integration of BPM for seawater electrolysis.^{163–165} In recent research, scientists explored the application of BPM in the development of an asymmetric bipolar membrane water electrolyser (BPMWE) aimed at seawater electrolysis (Fig. 14).¹⁶³ A unique design was employed, incorporating a catalyst layer placed between a cathode with an attached CEL and an anode

with an attached AEL, with deionised water and a 0.5 M NaCl solution supplied to the anode and cathode, respectively. The role of the CEL layer was pivotal in preventing the crossover transfer of cathodic Cl⁻, thereby avoiding interference with anodic OER and preserving the overall efficiency of the electrolyser. By utilising BPM, saline water was successfully purified by minimising the transmission of unwanted ion crossovers. Later, a comparative study with a conventional PEM electrolyser revealed that BPM electrolyser, when fed with impure water, exhibited enhanced stability and avoided competitive ClER. However, when real seawater was introduced to both cathode and anode of BPMWE, the system's stability decreased significantly due to the transport of Cl⁻ ions across AEL, leading to corrosive OCl⁻ species formation and reduced efficiency. This limitation highlighted the challenges of utilising BPM in full seawater electrolysis. Moreover, BPMWE faced barriers such as higher operational voltage requirements (above 3 V), a substantial increase in internal resistance caused by the application of two membranes and a high mass loading of precious metals catalysts (6.25 mg cm⁻²), which is not aligned with targeted values for utilisation of noble metal materials in PEM electrolysers by 2026. Despite its innovative approach, BPMWE, as it stands, requires further optimisation to meet the efficiency, stability, and cost targets necessary for practical application in seawater electrolysis.

Based on a comprehensive evaluation encompassing current density, faradaic efficiency, operational stability, energy demand, and overall cost, self-driven purification technology emerges as the most promising option. This approach demonstrates superior electrochemical characteristics, combined with relatively low operational costs and satisfactory durability. Although the performance of forward osmosis is not as high, it still shows considerable potential as an alternative method. On the other hand, both the vapour-fed system and the bipolar membrane approach are hindered by limited current output and stability issues, alongside elevated energy requirements and high implementation costs, which collectively constrain their feasibility for practical deployment.

5. Summary and outlook

In this review, the environmental impact of seawater electrolysis, the technical barriers posed by the complexity of seawater composition, and various strategies to address the challenges of direct seawater electrolysis are extensively discussed. These strategies encompassed diverse approaches, from exploring innovative electrodes to adjusting the electrolyte composition and membrane designs, aiming to achieve efficient, cost-effective, and sustainable seawater electrolysis. Some recent breakthroughs in the development of novel seawater electrolysis systems were also highlighted, shedding light on their potential but also acknowledging their existing limitations. Building upon these discussions, the potential future research directions are identified below, which could significantly shape the field of direct seawater electrolysis (Fig. 15).

(1) Although efforts have been carried out to design electrodes for direct seawater electrolysis by multi-scale design, the



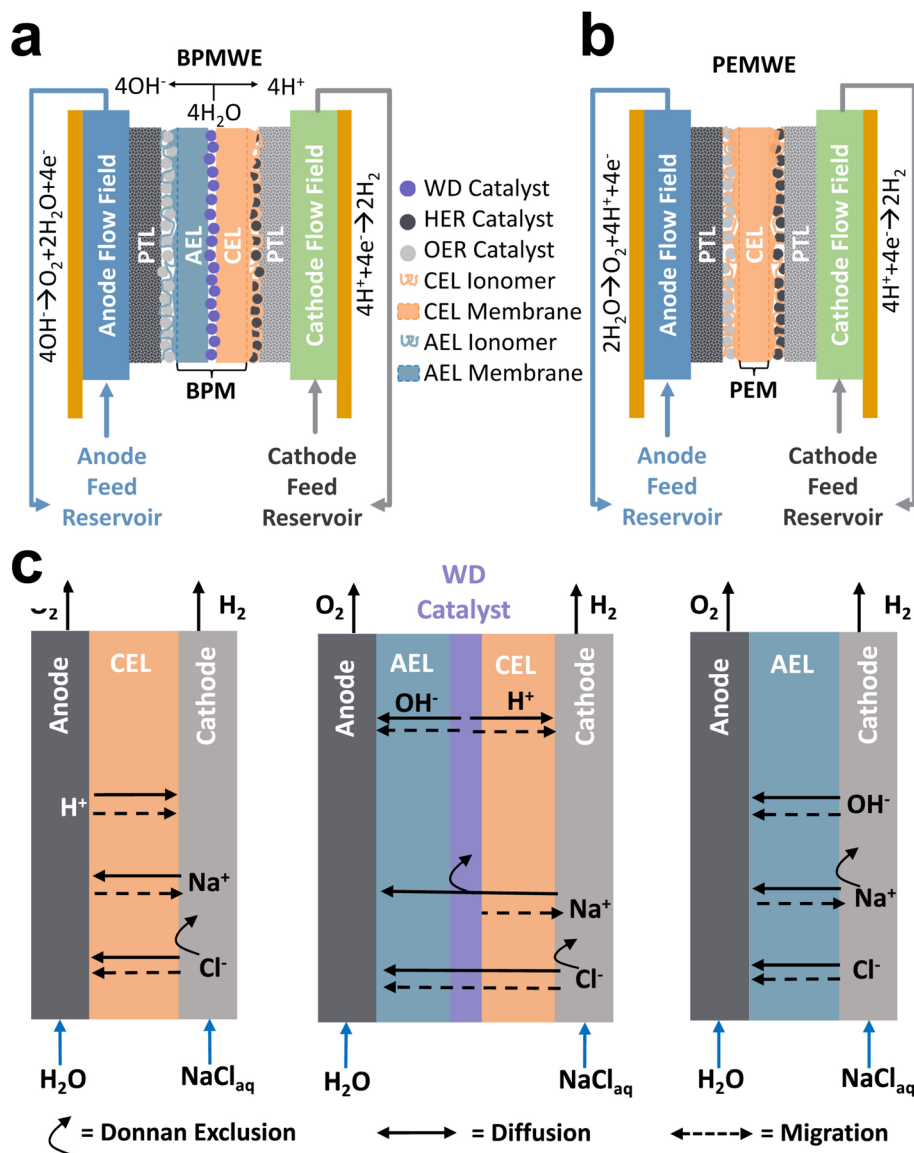


Fig. 14 Schematics showing the placement of ion-exchange membranes and catalyst/ionomer-coated porous transport layers (PTL), along with the feed circulation used for (a) zero-gap BPMWE and (b) zero-gap PEMWE electrolyser design. (c) Schematic illustrating the primary ion-transport mechanisms: Donnan exclusion (curved arrows), diffusion (solid arrows), migration effects (dashed arrows), and that govern the ion transport across ion-exchange membranes for PEMWE (left), BPMWE (middle), and AEMWE (right) electrolyser systems.¹⁶³ Reproduced from ref. 163 with permission from Elsevier, copyright 2023.

performance and stability of developed electrodes are still far from the industrial target values. Therefore, it is recommended to extend research on industrialising electrodes using multiple strategies.

(2) Efforts should focus on manipulating electrocatalysts' electronic structure for optimal adsorption energy and improving selectivity, particularly for OER. In this regard, both morphology and electronic engineering have demonstrated their efficacy in enhancing catalytic performance. Exploring specialised hydrophilic nanostructure catalysts, especially porous or multilayered ones, that provide abundant active sites and prevent bubble formations, enhancing overall efficiency and stability.

(3) Achieving high stability for electrodes requires enhancing chloride corrosion resistance. Recommended further investigations in this area include: (a) exploring suitable anti-corrosion coatings, considering diverse pH conditions without compromising catalytic performance, is crucial for industrial seawater splitting. For example, developing core-shell structures by integrating a catalytically active core with specific protective shells (manganese oxides, transition metal phosphides, or graphene) that enhance corrosion resistance and stability; (b) evaluating the performance of high-entropy materials with stable structure resilient to corrosion, for seawater electrolysis.



Future research directions



Fig. 15 Future research directions of direct seawater electrolysis development.

(4) Limited research has been conducted on electrode poisoning by insoluble and organic impurities in seawater. For example, the precipitation of magnesium/calcium hydroxides during electrolysis clogs catalyst channels and covers active sites, severely impairing catalytic performance and reducing catalyst lifespan. Exploring the impact of these impurities on electrode performance, preventing strategies, and electrode regenerating techniques are crucial for direct seawater electrolysis.

(5) Since the suitable choice of substrate composition and structure enhances the mass transfer, catalytic activity, and corrosion resistance, further research on substrates/supports for electrocatalysts is suggested. Particularly, extending research on developing self-supported electrocatalysts with controllability of morphology and structure is highly recommended.

(6) Transition metal-based electrocatalysts hold promise as replacements for precious metal-based catalysts, but their large-scale production remains a significant concern. Current synthesis methods involve high-temperature hydrothermal and solvothermal processes, leading to low yield, energy inefficiency, and scaling issues to meet industrial demand. There's a crucial need for cost effective, sustainable, and simplified production techniques to enable the broad application of these electrocatalysts in seawater splitting.

(7) Owing to the intricate chemical makeup of seawater, the electrochemical processes occurring during its electrolysis are considerably more complex than those in freshwater systems. As such, comprehensive investigations into the underlying reaction mechanisms are crucial for advancing knowledge of electrode structural dynamics, active site behaviour, electronic characteristics, and electrocatalytic pathways in seawater electrolysis. In this regard, combining electrochemical and *in situ* characterisation techniques with computational chemistry methods (including density functional theory calculations

(DFT) and Molecular Dynamics (MD) simulations) would provide a clear understanding of electrochemical reactions and structural alterations during seawater electrolysis.

(8) Despite significant investment in the development of seawater-based electrocatalysts, the actual hydrogen yields, power costs, and electrolyser testing for realistic applications have been disregarded, which are necessary to the commercialisation of extended hydrogen generation by seawater electrolysis. As a result, increasing hydrogen production experiments, appropriate power consumption studies, and the design of an efficient electrolytic cell were critical.

(9) The impact of organic and biological/microbial impurities is underexplored. Therefore, further comprehensive research is essential to evaluate the influence of these impurities on electrocatalysts, membranes, electrochemical reactions over a range of pHs and overall electrolyser performance.

(10) Future research could target the creation of membranes that possess high robustness in harsh seawater electrolytes and high selectivity for specific ions, ensuring that impurities do not hinder the electrolysis process. Additionally, the development of self-cleaning or fouling-resistant membranes could contribute to prolonged system operation without significant efficiency losses.

(11) The efficiency of seawater electrolysis is not solely dependent on individual components but also on the holistic system design. Research in this area could explore innovative reactor designs and flow patterns that enhance mass transfer and minimise energy losses, making the technology more scalable and adaptable to various applications.

(12) Innovative and efficient integration strategies are essential for direct seawater electrolysis. Technologies allowing simultaneous self-purification and electrolysis are desirable, such as solid oxide electrolysis cells (SOEC) dissociating water into hydrogen and oxygen at high temperatures, coupling interfacial-solar stream and water electrolysis or integrating



flow-electrode capacitive deionisation devices with electrolyzers.

(13) Further research into the environmental impacts of direct seawater electrolysis is critical as it could provide a more sustainable pathway for hydrogen production. To advance knowledge in this area, suggesting research directions include: investigate marine ecosystem impacts of brine discharge, develop mitigation strategies for chlorine emissions, assess the impact of alkaline brine on ocean acidification, investigate “zero liquid discharge” (ZLD) strategies and methods for resource recovery from brine, particularly valuable salts and minerals like magnesium or lithium, assess cumulative ecological impact of electrolysis plants along coastlines (particularly in regions with high biodiversity or sensitive ecosystems), develop predictive models and monitoring systems that track the long-term ecological impacts of direct seawater electrolysis and inform adaptive management.

(14) Comprehensive life cycle assessment (LCA) is essential to quantify and compare the environmental impacts of direct seawater electrolysis *versus* conventional electrolysis (using desalinated water) across their full life cycles.

Data availability

This article is a review and does not involve the creation or analysis of new data. All data supporting this work are derived from publicly available sources, as cited within the text.

Conflicts of interest

There are no conflicts to declare.

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