Electrochemical Compression Technologies for High-Pressure Hydrogen: Current Status, Challenges and Perspective

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Abstract

Hydrogen is an ideal energy carrier in future applications due to clean byproducts and high efficiency. However, many challenges remain in the application of hydrogen, including hydrogen production, delivery, storage and conversion. In terms of hydrogen storage, two compression modes (mechanical and non-mechanical compressors) are generally used to increase volume density in which mechanical compressors with several classifications including reciprocating piston compressors, hydrogen diaphragm compressors and ionic liquid compressors produce significant noise and vibration and are expensive and inefficient. Alternatively, non-mechanical compressors are faced with issues involving large-volume requirements, slow reaction kinetics and the need for special thermal control systems, all of which limit large-scale development. As a result, modular, safe, inexpensive and efficient methods for hydrogen storage are urgently needed. And because electrochemical hydrogen compressors (EHCs) are modular, highly efficient and possess hydrogen purification functions with no moving parts, they are becoming increasingly prominent. Based on all of this and for the first time, this review will provide an overview of various hydrogen compression technologies and discuss corresponding structures, principles, advantages and limitations. This review will also comprehensively present the recent progress and existing issues of EHCs and future hydrogen compression techniques as well as corresponding containment membranes, catalysts, gas diffusion layers and flow fields. Furthermore, engineering perspectives are discussed to further enhance the performance of EHCs in terms of the thermal management, water management and the testing protocol of EHC stacks. Overall, the deeper understanding of potential relationships between performance and component design in EHCs as presented in this review can guide the future development of anticipated EHCs.

Keywords Hydrogen storage · Hydrogen energy · Non-mechanical compressor · Electrochemical hydrogen compressor · Purification function
1 Introduction

Although energy is a significant vector in modern society, the unrestrained use of fossil fuels in the past few centuries has led to energy shortages along with global warming, air pollution and other environmental concerns. To address these issues, the development of novel renewable energy technologies with abundant resources, wide distributions, renewability and environmental friendliness has become the best choice to reduce fossil fuel consumption. And although current research has been focused on new energy technologies based on wind energy, tide energy and solar energy [1], the large-scale application of these new energy technologies is hindered by intrinsic unpredictability and intermittency [2]. Therefore, the search for novel energy carriers that can enable transfer between renewable resources and end-use customers is vital. Here, hydrogen as a completely pollution-free, carbon–neutral energy carrier is extremely promising in the addressing of these issues.

Hydrogen as a carbon-free fuel produces only water in related processes in which a carbon-free cycle is formed through the generation of water and hydrogen with clean and abundant energy. Hydrogen as an energy vector can also play an important role in society in which at the consumer or end-customer level, approximately three-quarters of primary energy is used as fuel and one-quarter is used as electricity [3]. Based on this, primary energy sources must be transformed into energy carriers for consumers. Here, hydrogen is a versatile primary energy source that can be converted into other forms of energy through five different approaches, including flame combustion, direct steam production, catalyst combustion [4], chemical reaction (hydrating) [5] and electrochemical conversion (fuel cells) [6]. In addition, based on the conversion efficiency of hydrogen into thermal, mechanical and electrical forms, it is more efficient to convert hydrogen into desired energy forms than other fuels [7]. Moreover, hydrogen also possesses an attractively high energy density between 120 MJ kg⁻¹ (the lower heating value, LHV) and 142 MJ kg⁻¹ (the higher heating value, HHV) [8] and has shown great promise in many fields. In terms of industrial application, hydrogen is primarily used as a reactant in fertilizer production and as a refinement material for metal processing [9]. Moreover, hydrogen is being increasingly used as a fuel in hydrogen refueling stations (HRSs) and proton exchange membrane fuel cells (PEMFCs) [10].

However, the promotion of hydrogen still faces a series of challenges, including production, storage and transportation. For hydrogen production, most industrial hydrogen is currently produced from fossil fuels through partial oxidation, autothermal oxidation, steam reforming and gasification [11] in which hydrogen from steam reforming suffers from serious issues (i.e., large numbers of impurities such as carbon monoxide) that need to be resolved before use. Hydrogen storage technologies are also limited by several issues, including low volume energy density, low efficiency compression and high-cost pressure vessels. And due to low-volume densities, hydrogen storage also requires high-pressure vessels or liquefaction under low temperatures in which due to bulky storage tanks, most hydrogen storage systems can only be used in stationary applications. As for hydrogen transport, industrial standards in the large-scale use of hydrogen require airtight shipping containers, the transport and storage of which can lead to further cost increases in hydrogen systems [12].

One key technology for the advancement of hydrogen technology application is hydrogen storage in which ideal storage technologies require: (1) reversible storage, (2) high efficiency, (3) high gravimetric and volumetric energy densities, (4) enhanced safety and (5) cost-effectiveness. Here, common technologies used for increasing the volumetric energy density of hydrogen involve compression and liquefaction. For liquefaction, the critical liquefaction process (temperature of −252.882 °C and pressure of 1.298 MPa) is an energy-intensive process that requires the use of cryogenic refrigeration to compress gaseous hydrogen into liquid hydrogen [13]. Alternative to the harsh requirements of liquefaction, direct compression is a more economical and convenient process in which associated hydrogen compressors are important for the development of hydrogen energy industry. Here, two main methods exist for the direct compression of hydrogen which are mechanical and non-mechanical methods. And although the use of mechanical compressors is the most conventional method, it is accompanied by several disadvantages. First, mechanical compressors possess lower efficiency in terms of the adiabatic compression process as compared with isothermal compression under the same compression ratios. Second, the moving parts of mechanical compressors can easily be damaged due to harsh operating conditions (huge pressure and low temperature) [14] and last, mechanical compressors are noisy due to vulnerable moving parts in which the US department of energy (DOE) reported that hydrogen compressors were the second most problematic component in HRSs to account for 18% of total incidents and a model designed by the national renewable energy laboratory (NREL) showed that compression accounted for the largest percentage of HRS operating costs (Fig. 1a) [15].

In terms of non-mechanical compressors, the absence of moving parts with the reduction in noise is an obvious advantage that can not only reduce the possibility of compressor failure to deliver cost savings but also improve the safety of devices. Currently, non-mechanical hydrogen compressors include cryogenic compressors, adsorption compressors, metal hydride compressors and electrochemical hydrogen compressors (EHCs) and with the exception of modular EHCs, all require large-scale sites. And because
EHCs can allow for the electrochemical compression of hydrogen with lower electric power consumption, reduced thermal loss and higher efficiency as compared with other non-mechanical hydrogen compressors as well as simple device designs, they have attracted significant attention in the last decade. In addition, the low-pressure requirements at the initial input of EHCs for the storage of hydrogen can reduce safety risks and EHCs can allow for the simultaneous compression and purification of gaseous hydrogen, making EHCs one of the most promising choices for hydrogen compression in the future.

More importantly, the electricity consumed by EHCs during operation can be supplied from surplus solar, wind or biomass energy and low-pressure hydrogen can be...
supplied from hydrogen reforming, electro-water splitting or hydrogen storage. Furthermore, the outlet high-pressure and pure hydrogen from EHCs can be directly input into various applications including industry, storage and automobile applications. And as a result of these advantages, EHCs can also be applied in anodes because a recirculating hydrogen system can reduce the frequency of purging operations and voltage pulsations to result in higher efficiency [16, 17] in which Barbir et al. [18] in 2006 evaluated the use of EHCs for the recirculation of hydrogen in a 10-cell EHC and achieved satisfactory and stable results. More recently, many projects based on EHCs have been developed for freezer compressors [19], infrared detectors in aerospace applications [20], gas-cleaning installations [21], etc. and other studies have explored the industrial-scale separation of hydrogen isotopes based on EHCs (Fig. 1b) [22].

Based on all of this and to guide the further development of EHCs, this review will first briefly introduce the operational principles and characteristics of all hydrogen compressors followed by the emphasis of EHC principles and their significant potential for real-world application. Subsequently, this review will analyze the basics of EHC operating procedures and levels of performance along with technologies and design features potentially achievable. More specifically, this review will discuss the current components of EHCs with a focus on technological limitations and current performances and outline existing methods for performance improvements.

2 Hydrogen Compression Technologies

In general, hydrogen compressors can be divided into mechanical and non-mechanical compressors [23], and in this section, both compressors along with their advantages and disadvantages are introduced with a focus on EHCs.

2.1 Mechanical Hydrogen Compressor

Mechanical compressors are the most widely used compressors to compress low-pressure hydrogen into high-pressure tanks by using mechanical energy in which with the lowering of confined volume, gaseous hydrogen can be squeezed into smaller chambers and result in higher gas pressures. In the following sections, general mechanical compressors including reciprocating piston compressors, hydrogen diaphragm compressors and ionic liquid compressors will be briefly introduced.

2.1.1 Reciprocating Piston Compressor

One example of mechanical compressors is the reciprocating piston compressor which is equipped with two separate automatic valves for inlet and outlet hydrogen, and a piston-cylinder system constitutes a single-stage reciprocating compressor (Fig. 2a). Here, a connecting rod connects a piston to a crankshaft and the piston transforms the rotary motion of a moving unit into the approximate linear motion of the piston. This movement is referred to as reciprocating motion [24] in which specifically, the piston moves forth and back in a cylinder to compress hydrogen gas with its head. In most applications, this sequence of movements is driven by a pump that is connected to two cylinders in which the compression step follows an expansion step, which comprises the working principle of reciprocating piston compressors. Overall, reciprocating compression is a mature technology that can be used to compress almost all gases and high-pressure hydrogen can be obtained by using reciprocating compressors with multi-stage configurations. Because of this, reciprocating compressors can provide flexibility during compression. Despite this, reciprocating compressors also possess several limitations that hinder application in hydrogen compression. As one disadvantage, the durability of compressor components is affected by the use of lubricants and can lead to risks of explosion, meaning oil-free compressors are preferred [25]. The presence of moving parts can also lead to increased costs due to manufacturing complexities and difficulties as well as increased maintenance frequencies during operation. Moreover, moving parts also generate heat and decrease the efficiency of overall systems [26]. Furthermore, the reciprocating process of pistons can cause pressure fluctuations that can lead to vibrations, noise and even disastrous explosions [27]. Reciprocating compressors are also not efficient at high flow rates despite high flow rates being potentially achievable [28] depending on the number of cycles and the dimension of cylinders in which in addition to the weight of minimum mechanical compressors being normally heavy (200–400 kg), increases to cylinder dimensions can further lead to bulkier systems that can in turn increase internal forces. And in order to reduce the influence of mechanical stress, lower speeds are preferred, meaning that high-compression speeds can only be achieved in the small scale, which restricts allowable flow rates.

As for industrial application, reciprocating piston compressors may be a good solution for large-scale mechanical compressors used in HRSs in which many projects can attain high-compression levels up to 70 MPa and 85 MPa through multi-stage compression. In addition, outlet pressures of 100 MPa and flow capacities of 300 N m$^3$ h$^{-1}$ have been attained in a Hitachi infrastructure system [29].

2.1.2 Diaphragm Compressor

Diaphragm compressors follow a similar working principle to reciprocating piston compressors in which with pistons pulling downwards, hydraulic oil can flow back to the
cylinder. Here, the resulting differential pressure can cause the downward elastic deformation of a diaphragm to increase the volume of a corresponding chamber and automatically open a gas inlet valve to suck gas in. Alternatively, as a con-rod pushes the piston upward, the piston will further push the hydraulic oil to cause the upward elastic deformation of the diaphragm (Fig. 2b) and decrease the volume of the gas chamber to result in hydrogen compression. And as gas pressures reach a certain level, a gas discharge valve will open automatically [30]. Based on this mechanism, diaphragm compressors possess lower-power consumption, high throughput and cooling requirements and therefore are considered to be effective for hydrogen compression [31]. In addition, diaphragm compressors also possess the potential to achieve high levels of volumetric efficiency, which further demonstrates its energy-saving advantages [32]. Despite this, the durability of diaphragm compressors is a critical issue in which reciprocating parts can easily be broken by mechanical stress during operation. And similar to reciprocating piston compressors, diaphragm compressors also possess relatively complex operating systems that are more suitable for applications requiring low flow rates in limited volume compression chambers [33]. Moreover, diaphragm failure can also occur as caused by radial stress related to diaphragm deflection [34] and the geometric design of cavity spaces. And because high flow rates can invalidate diaphragms, corresponding concavities and grooves need to be properly designed to ensure proper flow distribution. As a result, new designs for diaphragm plates are based on numerical simulation models [35, 36].
As for current diaphragm compressor applications, a German company named Hofer that is well-known for its hydrogen diaphragm compressor has equipped several HRS systems with diaphragm compressor to compress gases at ~ 390–581 N m⁻³ h⁻¹ with discharge pressures up to 28.1 MPa [37]. In addition, PDC machines have developed products to operate at discharge pressures ranging from 48.2 to 103.4 MPa with production rates ranging from 5 to 2500 kg daily [38, 39]. The Arizona Public Service company also built a three-stage diaphragm compressor that is capable of compressing hydrogen up to 41 MPa in a hydrogen and natural gas fueling station in downtown Phoenix [40].

### 2.1.3 Ionic Liquid Compressor

Ionic liquid compressors are also based on the same operating principles as reciprocating compressors. In the designed structure, solid pistons are replaced by ionic liquids in which ionic liquids are directly pumped into the compression chamber/chambers to compress hydrogen and send it back to the hydraulic system afterward [41]. Here, researchers have shown that the replacement of hydraulic oil with appropriate ionic liquids in a compressor can contribute to remarkable improvements in volumetric efficiency of approximately 10%–30% [42]. In addition, Kermani et al. [43] found that two imidazolium-based ionic liquids, [EMIM][Tf₂N] and [EMIM][CF₃SO₃], with significantly lower viscosity and corrosion current densities for AISI 316L can demonstrate desirable performances in ionic liquid hydrogen compressors. Moreover, Linde, a German international company, was able to develop a ionic liquid compressor with only eight moving units for hydrogen applications [44].

The fewer moving parts of ionic liquid compressors can further lead to the reduction in mechanical loss and the improvement in overall efficiency for hydrogen compression. And due to the unique properties of ionic liquids, especially their insolubility with gases, many complex designs involving bearings and sealings have been removed to significantly reduce overall system complexities. In addition, the reduction in mechanical parts can also significantly decrease operational expenses and extend the mechanical life in which the service life of ionic liquid compressors is almost ten times longer than that of regular reciprocating compressors. Furthermore, ionic liquid compressors can compress hydrogen gas to 90 MPa in five steps and the removal of lubricating oils allows for extremely pure hydrogen. Moreover, the ionic liquids used in these compressors display good coolant properties and lubricating performances. Overall, ionic liquid compressors are designed especially for increased hydrogen compression efficiency with ionic liquids possessing good chemical and thermal stability, enhanced fire retardance, high ionic conductivity, high polarity, negligible volatility and moderate viscosity [45]. Ionic liquids are also non-toxic and possess low compressibility, low volatility and good lubricating performances, especially for high-pressure applications [46, 47].

Despite these desirable attributes, ionic liquid compressors are not perfect and possess obvious drawbacks, such as the risk of corrosion, the possibility of leakage and the complexity of system design. In addition, ionic liquids can leave compression chambers through discharge tubes with small amounts of hydrogen, resulting in the need for additional cleaning processes for outlet hydrogen [48].

In terms of real-world application, the Linda Group has developed an ionic liquid compressor that can achieve compression pressures up to 45–90 MPa with a net capacity of 8–30 kg h⁻¹ [44]. Nasrin has also designed a hydraulic to pneumatic transformer to integrate pneumatic and hydraulic systems for an ionic liquid compressor that can build high pressures above 70 MPa [49].

### 2.2 Non-mechanical Hydrogen Compressor

Despite the widespread adoption of mechanical compressors, many problems exist. Based on this, other types of compressors with non-mechanical principles have been developed, including cryogenic compressors, adsorption compressors, metal hydride compressors and EHCs. In this section, the general principles and features of these non-mechanical hydrogen compressors are introduced.

#### 2.2.1 Cryogenic Compressor

In general, several modular elements make up a cryogenic compressor, including a container for low-pressure liquid hydrogen storage, a cryogenic compression container and a cryogenic pump in which liquid hydrogen is fed into a cryogenic pump through a vacuum-insulated tube and stabilized to a desired pressure through a cryogenic compressor (Fig. 2c) [50]. Vaporizers are also placed downstream of cryogenic pumps to ensure the acquisition of high-pressure hydrogen [51]. Here, cryogenic compressors can attain higher (more than two times) volumetric efficiency than mechanical compressors [52] and compressed hydrogen possesses high hydrogen densities, meaning cryogenic compressors possess high gravimetric and volumetric capacities and are suitable for large-scale hydrogen storage and compression.

Despite these advantages, hydrogen cryogenic compressors also possess an obvious drawback involving the energy cost that is a critical barrier hindering future application in which of total input energy, only 30% is stored based on the lowered heating value of hydrogen. The storage of liquefied hydrogen is also an issue hindering application due to the
difficulty in ensuring long-term vacuum stability for about 10 years. And although the high-temperature baking of metal surfaces can increase the degree of vacuum stability in internal pressure vessels made of composite materials [53], it poses great challenges in the rational design of materials and the investigation of compressor fabrication methods.

As for industrial applications, a cryogenic compressor manufactured by the Linda Group company used in hydrogen refueling stations can reach pressures up to 35–90 MPa with high throughput (100 kg h^{-1}, enough to fill an automobile in 5 min). In addition, the benefits of cryogenic container technologies have been comprehensively evaluated by establishing a method to evaluate the filling density in any initial thermodynamic state of containers [50] and Kunze et al. [54] introduced a cryogenic compressor that can compress hydrogen up to 30 MPa. Despite this, many problems exist in the development of cryogenic compressors that need to be conquered before industrial application [55, 56].

### 2.2.2 Adsorption Compressor

In terms of adsorption compressors, corresponding structures can be seen as a thermodynamic engine by using heat exchange as the driving force of compression (Fig. 2d) in which low-pressure hydrogen is injected into high-adsorption potential porous materials in a compressor, and physical hydrogen adsorption is desorbed at a certain volume after reaching high pressures under specific temperature and pressure conditions [57]. And depending on the physical nature of porous materials, the flow of compressed hydrogen is obtained through multiple cycles of adsorption/desorption.

Adsorption compressors have been widely used as hydrogen compressors due to several advantages. Not only do adsorption compressors require less pressure to store hydrogen and enhance safety, but these compressors also do not possess any moving parts and therefore do not generate noise or experience mechanical damage. In addition, more hydrogen can be obtained through the use of porous materials per unit volume as compared with mechanical compressors [58] in which zeolites, carbonaceous materials (i.e., activated carbons, carbon nanotubes or fullerenes) and metal–organic frameworks (MOFs) are usually used as porous materials in adsorption compressors [59, 60]. Here, the interaction between hydrogen and solid bed surfaces through weak Van der Waals forces can drive the formation of monolayered hydrogen molecules on absorbent surfaces. Moreover, required adsorbent materials in adsorption compressors are readily available and the driven forces of industrial waste heat can be used to lower system costs.

Adsorption compressors possess weaknesses as well, especially due to issues concerning heat and mass transfer in adsorbent beds, leading to the need for cooling systems and increased complexity in overall systems [61]. In addition, because the cycling of adsorption and desorption in adsorption compressors is intermittent in nature, adsorbent beds experience thermal gradient differences that can affect operating efficiency [62]. Moreover, the need for high vacuum conditions also creates technical engineering problems which if not evaluated properly, adsorption heat would increase temperatures and reduce adsorption capacity and system performance [63]. Furthermore, because the heat energy of hydrogen molecules is directly proportional to the temperature of the system [64], the interactions between adsorbent surfaces and gas molecules can increase with cooling systems. However, the adsorption of hydrogen usually occurs at reaction temperatures (~77 K) that are too low.

Despite these issues, adsorption compressors have proven to be an appropriate choice for industrial hydrogen compression. For example, various prototypes [65, 66] have demonstrated high-efficiency compression capabilities in which through the use of activated carbon as an adsorbent material, hydrogen with a pressure of 10 MPa can be obtained. Richard et al. [67] also used Maxsorb MSC-30TM-activated carbon as an adsorbent material to successfully compress hydrogen from 0.25 to 35 MPa. Based on these results, new technologies for adsorption compressors should be further developed.

### 2.2.3 Metal Hydride Compressor

As for metal hydride compressors (MHCs), they are made up of a slender central artery to distribute hydrogen inside a reactor and an annular space between the artery and a tank wall for metal hydride placement. In principle, low-pressure hydrogen can enter the metal hydride tank through the central artery and diffuse into the metal hydride bed to allow for the exothermic reaction of hydrogen absorption in which hydrogen compression can occur due to the continuous cooling and heating of the metal hydride as controlled by heat transfer [68]. Here, the selection of well-suited metal hydrides with BCC, AB_{5} and AB_{2} structures is vital [69] in which Ni-based AB_{3} hydrides have been found to be particularly promising due to their low costs and acceptable performances, especially at moderate temperatures.

In terms of advantages, MHCs possess straightforward designs with no moving parts and can operate with no noise and are not energy intensive. In addition, MHCs are also more compact and can more easily integrate into existing hydrogen refueling infrastructures [70]. Furthermore, MHC systems can be powered by using industrial waste heat rather than electricity in which high-pressure hydrogen can be obtained in situ from water through connections with electrolytic cell outlets to recover heat loss from electrolysis [71].
Unfortunately, the efficiency of MHCs is generally less than 25% at 423 K [69] and strictly depends on the compression rate and the amount of heat provided to the system due to several types of energy loss, including heat transfer as well as heat for hydrogen desorption and cooling. Furthermore, the use of MHCs results in large volumes, slow refueling times and low gravimetric densities and therefore is unsuitable for small mobile power stations. Moreover, the durability of MHCs is still currently unsatisfactory.

As for practical application, many MHCs have been developed with outlet pressures up to 35–70 MPa [72, 73]. For example, Pickering et al. [74] reported promising hydrogen absorption capacities in vanadium-based BCC solid solution alloys with high absorption/desorption kinetics at ambient temperature in which the addition of small amounts of niobium and manganese to Ti–V based alloys can result in a pressure of 65 MPa at moderate temperatures.

### 2.2.4 Electrochemical Hydrogen Compressor

Electrochemical hydrogen compressors (EHCs) are devices that use the electrochemical principle to compress low-pressure hydrogen into high-pressure hydrogen in which the application of voltage can lead to the generation of localized pressure difference due to hydrogen oxidation at anodes and hydrogen reduction at cathodes. Here, protons and electrons produced through hydrogen oxidation are transported to the cathode side through a proton exchange membrane (PEM) (for protons) and an external path (for electrons) to recombine to form new hydrogen molecules (Fig. 2e) in which electric power is converted to chemical potential in high-pressure hydrogen gas through the electrochemical process. As a result, EHC systems are analogous to proton exchange membrane fuel cells (PEMFCs) and contain PEMs, catalyst layers (CLs), gas diffusion layers (GDLs), flow field plates and end plates.

Overall, ideal compression should be isothermal in nature and operate without heat generation as shown in Eq. (1) as dictated through thermodynamic principles [the energy needed for the adiabatic process in mechanical compressors is expressed in Eq. (2) for comparison]:

\[
W = RT\ln\left(\frac{P_c}{P_a}\right)
\]

\[
W = \frac{\gamma}{\gamma - 1} \left[\frac{(\gamma - 1)}{\gamma}\right] \frac{P_c}{\gamma P_a}
\]

in which \(W\) represents power required for compression, \(R\) is the gas constant of 8.314 J mol\(^{-1}\) K\(^{-1}\), \(\gamma\) is the ratio of specific heat (for H\(_2\), \(\gamma\) is 1.4), \(P_c\) and \(P_a\) represent hydrogen pressure at high-pressure and low-pressure sides, respectively, and \(T\) represents the temperature of hydrogen on the low-pressure side. Based on these equations, it can be seen that the theoretical efficiency of isothermal compression is almost two times higher than that of adiabatic compression with a compression ratio of 300 (Fig. 2f). And based on the fact that the EHC process is isothermal in nature and through the increased profitability of hydrogen produced from EHCs (current hydrogen prices are around 11€ kg\(^{-1}\) due to the price reduction in membrane-electrode-assemblies (MEA) [75], EHCs are expected to replace conventional mechanical compressors. Like PEMFCs, EHCs also possess other attractive advantages, including the potential for higher efficiency, lower or zero emissions, increased simplicity and reduced costs. As another significant advantage, the use of EHCs can allow for the extraction of hydrogen from gaseous mixtures in which although current global hydrogen consumption is estimated at 50 Mt y\(^{-1}\), only ~4% is generated from electrolysis with the majority being produced from fossil fuels (~96%) along with small amounts from other sources such as medical and aerospace applications [76]. Recently, EHCs have also been applied to purify hydrogen similar to traditional pressure swing adsorption (PSA) and dense metal membrane techniques [77] in which Nordio et al. [78] compared EHC systems with PSA systems and reported that EHCs were more worthwhile in small-scale operations with elevated outlet hydrogen pressures. Moreover, EHCs have also been applied to coordinate MHCs in the effective reduction in operating costs [79–81].

Investigations into EHCs from different research institutes and companies have also revealed potential for future practical application. For example, ANALYTIC POWER conducted a two-stage approach to study EHCs as supported by the US DOE in 2005 due to the positive shifting in national policies on EHCs, and Fuel Cell Energy together with Sustainable Innovations were able to achieve a hydrogen pressure of 88.25 MPa in a single-stage mode and 41.37 MPa in a two-stage mode along with hydrogen recovery efficiency of over 98% by using EHCs. Other companies such as Proton Energy, Nuvera and H2Pump have also demonstrated hydrogen purification together with compressions up to 1.03 MPa by using EHCs along with pressures of 2.28 MPa (differential) by using polybenzimidazoles (PBI) as a membrane in 2012. In addition, Giner Inc. developed an EHC that can provide pressures of 87.5 MPa in a single-stage mode with a voltage of 0.159 V (each cell) and an inlet pressure of 10 MPa and also reported that maximum pressures up to 140 MPa can be achieved [82]. Grigoriev et al. [83] also developed an EHC that allowed for the compression of lower pressure into high-output pressure up to 13 MPa with...
a flow rate of 0.01 N m⁻³. Moreover, an EHC developed by HyET Hydrogen in the DON QUICHOTE project and the PHAEDRUS project achieved a compression up to 100 MPa [84] and in Japan, an EHC with an active surface area of 10 cm² was designed that demonstrated high pressures above 70 MPa under 1.2 A cm⁻² through burst testing [85].

By comparing commonly used hydrogen compressors mentioned above, the advantages and disadvantages of different compressors are summarized in Table 1. Overall, EHCs are prominent technology for hydrogen compression based on their advantages.

3 Electrochemical Hydrogen Compressor

3.1 Principle of EHCs

For EHCs using PEMs, low-pressure hydrogen is usually converted into high-pressure hydrogen through direct current. Here compression can be divided into three steps as driven by an imposed cell voltage similar to the cathodic reaction of water splitting involving the anodic oxidation of molecular hydrogen to form protons, the migration of protons through the PEM and the cathodic reduction in protons to molecular hydrogen. As for electrons, they are transferred through an external circuit to cathode CLs to allow for the simultaneous mass transport and purification of hydrogen. Overall, the EHC process involves low-pressure hydrogen \( P_a \) being fed into the anode side in which hydrogen oxidation reactions \( (\text{HORs}) \) occur to split hydrogen into protons and electrons whereas hydrogen evolution reactions \( (\text{HERs}) \) occur at the cathode side to reform hydrogen. The corresponding electrochemical reactions are as follows:

Anode: \( \text{H}_2 (P_a) \rightarrow 2\text{H}^+ + 2e^- \) (3)

Cathode: \( 2\text{H}^+ + 2e^- \rightarrow \text{H}_2 (P_c) \) (4)

Overall: \( \text{H}_2 (P_a) \rightarrow \text{H}_2 (P_c) \) (5)

in which \( P_c \) and \( P_a \) are the hydrogen pressures at the high-pressure cathode side and the low-pressure anode side, respectively.

The Volmer–Heyrovsky–Tafel mechanism can also be used to describe hydrogen electrode reactions [86] in which adsorbed hydrogen \( (\text{H}_{ad}) \) as a reaction intermediate species in hydrogen electrode reactions can be expressed by using three elementary reactions including the Volmer reaction, the Heyrovsky reaction and the Tafel reaction and the overall reaction can occur through either the Volmer–Heyrovsky route or the Volmer–Tafel route. All of these reaction routes can be represented as follows:

Volmer: \( \text{H}_2 \xrightarrow{v_f} \text{H}^+ + e^- \) (6)

Heyrovsky: \( \text{H}_2 \xrightarrow{v_f} \text{H}_ad + \text{H}^+ + e^- \) (7)

Tafel: \( \text{H}_2 \xrightarrow{v_f} \text{H}_ad + \text{H}_ad \) (8)

As for detailed mechanisms, they can be analyzed by using electrochemical impedance spectroscopy (EIS) with the spectra of the cathode displaying charge transfer rate-limiting features whereas the spectra of the anode exhibiting mass transfer rate-limiting features. Cathodic HER is dominated by the Volmer–Heyrovsky route and anodic HOR is dominated by the Volmer–Tafel route in which Chen and Kucernak [87] in their study proved that HOR follows the Volmer–Tafel reaction by using conventional ex situ characterization methods. Due to the slow reaction rate imposed by the Volmer–Heyrovsky route; however, cathodic HER requires high non-ohmic overpotentials. Despite this, HER kinetics can also increase by increasing cathode pressures, suggesting that the increased coverage of adsorbed hydrogen on catalyst surfaces can result in the reduction in non-ohmic overpotential for HER.

In EHCs, protons flow through a PEM and electrons flow through an external circuit to a cathode to eventually recombine into hydrogen molecules, thus resulting in the conversion of electrical energy into compression energy. Here, the reaction rate of hydrogen through PEMs can be calculated by using Faraday’s law:

\[
J_{\text{H}_2,\text{Theoretical}} = \frac{dn}{dt} = \frac{I}{2F}
\]

in which \( n \) is the inlet hydrogen flow in mol, \( F \) is Faraday’s constant \((9.648 \times 10^4 \text{ C mol}^{-1})\) and \( I \) is the current in A. In addition, the potentials of anodes and cathodes can be defined by using the Nernst equation as shown in Eqs. (10) and (11) and the relationship between the ratio of outlet gas and external circuit voltage can be calculated using Eq. (12):

\[
V_a = E_0 - \frac{RT}{2F} \ln \left( \frac{P_a}{P_0} \right)
\]

\[
V_c = E_0 - \frac{RT}{2F} \ln \left( \frac{P_c}{P_0} \right)
\]

\[
V_{\text{Nernst}} = \frac{RT}{2F} \ln \frac{P_c}{P_a}
\]

in which \( E_0 \) is the cell potential at standard conditions (considered 0 in the case of electrochemical compression), \( R \) is the universal gas constant \((8.3144 \text{ J mol}^{-1} \text{ K}^{-1})\) and \( T \)
Table 1 Different characteristics of common hydrogen compressors

| Category                        | Reciprocating piston compressors | Diaphragm compressors | Ionic liquid compressors | Cryogenic compressors | Adsorption compressors | Metal hydride compressors | Electrochemical hydrogen compressors |
|---------------------------------|----------------------------------|-----------------------|--------------------------|-----------------------|------------------------|--------------------------|--------------------------------------|
| Outlet pressure (MPa)           | 100                              | 100                   | 35–90                    | 35                    | 70                     | 70–100                   | 70–100                              |
| Typical capacity (N m$^{-3}$ h$^{-1}$) | 300                              | 51.7                  | 376–753                  | > 1000                | 72.6                   | 2                        | 0.93–1                               |
| Advantages                      | Mature technology                | Mature technology     | Fewer moving parts       | High hydrogen density | No moving parts         | No moving parts           | No moving parts                     |
|                                 | Flexibility in size and capacity | High throughput       | (compared to the first two) | High volumetric efficiency  | Simple design          | Simple design            | Simple design                      |
|                                 |                                  | Low cooling requirements | Long lifetimes           | High efficiency       | No noise               | No noise                  | No noise                            |
|                                 |                                  | Ideal for pure gases  | Low-energy consumption   | High gravimetric and volumetric capacities | Thermal energy driven | Thermal energy driven | Low energy cost                    |
| Disadvantages                   | Moving parts                     | Moving parts          | Possibility of leaking   | Low temperatures       | Limited heat transfer  | Slow compression time     | No moving parts                     |
|                                 | Contamination and danger due to lubrication oils (if used) | Low durability (diaphragm failure) | System complexity       | Low durability         | Additional cooling      | Limited heat transfer       | Simple design                       |
|                                 | System complexity                | Low flow rates        |                          | High vacuum quality    | system required        | Low efficiency            | No noise                            |
|                                 | Vibration                       | System complexity     |                          | High gravimetric and volumetric capacities |            | Low durability       | Hydrogen back-diffusion            |
|                                 | Noise                            |                        |                          | Low reaction temperature |                        |                          | Designed water management          |
|                                 | High heat loss                   |                        |                          |                       |                        |                          | High-strength accessory design required |
|                                 | Low efficiency                   |                        |                          |                       |                        |                          |                                     |
|                                 | High-cost                        |                        |                          |                       |                        |                          |                                     |
represents the temperature of hydrogen at the anode side. And although the theoretical voltage for hydrogen compression is only related to pressure difference and operating temperature based on the Nernst equation, the voltage of practical EHCs can be affected by several factors, including catalyst activity and PEM ion conductivity in which for voltage loss caused by catalysts and membranes, they can be classified into three causes, including activation overpotential-related to electrocatalysts, ohmic overpotential related to proton transport through PEMs and mass-transfer overpotential. Rochilz et al. [75] were also able to develop a zero-dimension and stable EHC model to calculate overall voltages by considering ohmic overpotential and activation overpotential and reported that different overpotentials can be separated based on anodes and cathodes. As for current density \( j \) applied to EHC cells, Eq. (13) can be used to express cell voltage \( V_{cell} \):

\[
V_{cell}(j) = V_{\text{Nernst}} + V_{\text{AnodeOhmic}}(j) + V_{\text{AnodeNonohmic}}(j) + V_{\text{CathodeOhmic}}(j) + V_{\text{CathodeNonohmic}}(j)
\]

Furthermore, the efficiency of EHCs can be classified into voltage efficiency and current efficiency in which voltage efficiency is defined by dividing theoretical voltage by cell voltage as expressed by Eq. (14), whereas current efficiency is defined by dividing the real amount of compressed hydrogen by the theoretical value as expressed by Eq. (15). Moreover, the overall efficiency of EHCs can be determined by both voltage and current efficiency as expressed by Eq. (16).

\[
\eta_V = \frac{V_{\text{Nernst}}}{V_{\text{cell}}}
\]

\[
\eta_I = \frac{J_{H_2,\text{Real}}}{J_{H_2,\text{Theoretical}}}
\]

\[
\eta_{\text{EHC}} = \eta_I \times \eta_V
\]

### 3.2 Operating Temperature of EHCs

The operating temperature of EHCs is highly dependent on the performance of membranes and in general, EHCs can be divided into low-temperature EHCs with low-temperature PEMs as membranes (e.g., Nafion), high-temperature EHCs with high-temperature PEMs as membranes (e.g., acid-doped PBI) and solid oxide EHCs with perovskite-type oxides as membranes [e.g., Ba(Zr_{0.30}Ce_{0.54}Y_{0.15}Cu_{0.01})O_{3−δ}].

#### 3.2.1 Low-Temperature EHCs

In terms of low-temperature EHCs, because the ion conductivity of perfluorosulfonic acid (PFSA) is highly dependent on water uptake, EHCs employing PFSAs such as Nafion as membranes can only operate at temperatures lower than 100 °C with operating temperatures normally ranging from 50 to 90 °C if considering catalyst activity. In recent years however, sulfonated hydrocarbon membranes such as SPEEK have also been employed as membranes due to low costs and high conductivities [88, 89]. And in general, single-stage low-temperature EHCs are more suitable for small-scale compressions.

#### 3.2.2 High-Temperature EHCs

As for high-temperature EHCs, phosphoric acid-doped polymers (e.g., the PBI/H₃PO₄ complex) are usually used to conduct protons. Various acids have also been studied for doping with PBI membranes, including HCl, HNO₃, H₂SO₄ and HClO₄, to allow for the conduction of protons with doped acids instead of hydrated protons and result in EHCs with these acid-doped polymers being considered as promising candidates for high-temperature EHCs at temperatures above 100 °C. These high-temperature EHCs are also promising due to associated cost reductions and reliability improvements through the elimination of issues in low-temperature EHCs in terms of water management, heat rejection, catalyst tolerance and reaction kinetics [90]. Furthermore, researchers have reported that the dipping of PBI in aqueous phosphoric acid to form an acid-base complex can result in enhanced proton conductivity, lowered gas permeability and a nearly zero water drag coefficient at high temperatures even in an anhydrous state. Moreover, the elevated operating temperatures of high-temperature EHCs can provide excellent thermal stability, a nearly zero water drag coefficient and improved impurity tolerance (especially carbon monoxide) for Pt catalysts as compared with low-temperature EHCs and the notorious flooding issue in the electrodes of low-temperature EHCs can be prevented due to the gaseous form of water at high temperatures.

#### 3.2.3 Solid Oxide EHCs

Perovskite-type oxides such as Ba(Zr_{0.30}Ce_{0.54}Y_{0.15}Cu_{0.01})O_{3−δ} can conduct protons at 600–900 °C. Based on this, solid oxide EHCs that are similar to solid oxide fuel cells and solid oxide water electrolysis systems have been proposed in recent years [91, 92]. Here, protons in solid oxide EHCs move through a lattice through a hopping process that is generated by water hydration reactions [93] and as compared with other fuel cell technologies, the main benefit of solid oxide EHCs is the existence of a solid electrolyte...
that can avoid the need for corrosive environments. Another great advantage of solid oxide EHCs is fuel flexibility in which a variety of hydrocarbon fuels can be used. Moreover, the production of hydrocarbon fuels such as ammonia can occur during hydrogen compression, thus allowing for the reduction of total costs for hydrogen compression [94, 95]. Furthermore, solid oxide EHCs are also suitable as large stationary compressors.

4 Components of Electrochemical Hydrogen Compressors

4.1 Membranes

PEMs are critical components in EHCs and need to possess high proton conductivities, good mechanical properties and low gas crossover rates [84] in which PEM proton conductivities can affect the efficiency of EHCs and good mechanical strength is needed to ensure the integrity of membranes under high-pressure. Here, HyET hydrogen and NREL have committed great efforts to the defect characterization of EHC membranes and the design of databases [96] in which to develop durable PEMs, the exploration of degradation mechanisms, especially the effects of mechanical stress in EHCs, is vital.

As for hydrogen back-diffusion, it occurs due to the existence of partial pressure differences between anodes and cathodes and is affected by many factors including the water content, temperature, membrane thickness and pressure [97]. In terms of the water content, the proton conductivity of PFSA membranes is strongly dependent on the water content and temperature in which water is needed to guarantee good proton conductivity in PEM hydrophilic phases because protons can move into hydrated sections through the dissociation of sulfonic acid bonds. Here, a low water content can cause PEM dehydration due to increasing operating temperatures and result in decreased proton conductivity whereas well-hydrated membranes can better accommodate increasing temperatures to enhance proton conductivity in which low conductivities can limit the access of protons to catalyst surfaces, reduce the number of reactive active sites in CLs and increase activation polarization [98]. And in cases of a very low water content, severe drying can occur that will lead to irreversible degradation (e.g., delamination, pinholes) [99] to result in significantly increased ohmic resistances. Alternatively, excess water can block catalyst sites and cause water flooding to result in poor performance. The water content in EHCs can also affect the swelling ratio of PEMs in which as the water content increases, the swelling ratio of PEMs also increases and can cause increased hydrogen back-diffusion [100].

System temperatures have a remarkable effect on transportation, including water diffusion, permeability and proton conductivity, and the influence of temperature on hydrogen crossover can be determined based on the hydrogen permeability coefficient that is closely related to temperature and can be expressed as the Arrhenius form [101]:

\[
\ln \psi_{H_2}^{PEM} = \ln \psi_0^{PEM} + \left( \frac{E_{H_2}^{PEM}}{R} \right) \frac{1}{T}
\]

in which \( \psi_{H_2}^{PEM} \) represents the hydrogen permeability coefficient, \( \psi_0^{PEM} \) is the maximum permeability coefficient (e.g., at infinite temperature), \( E_{H_2}^{PEM} \) is the activation energy for hydrogen crossover, \( R \) is the gas constant and \( T \) is the temperature in Kelvin. Based on this equation, it can be seen that hydrogen permeability coefficients increase with increasing temperature. Here, researchers reported that temperature decreases from 80 to 21 °C can lead to hydrogen gas permeability ratio decreases of 3 and 5 orders of magnitude under cathode pressures of 70 and 30 MPa, respectively [102]. Truc et al. [103] also proposed a numerical model for hydrogen crossover and the dependence of permeability on the membrane water content and temperature and reported that the reduction in EHC operating temperature can slow down hydrogen back-diffusion, particularly in the case of high-pressure differentials and thin or commercial membrane materials with greater porosity. As a result, optimum membrane operating temperatures for EHC applications can only be achieved through a compromise between high proton conductivity and low hydrogen back-diffusion to minimize overall energy requirements for compression [104].

As for the impact of pressure on hydrogen back-diffusion, there is a universal agreement that hydrogen back-diffusion rates increase with increasing outlet pressures at any temperatures and RH (the water content of membrane) and that increased hydrogen back-diffusion rates will lead to performance degradation and efficiency reduction [105]. Here, gas pressure applied on each side of EHC membranes is balanced with the solubility coefficient \( (H_1) \) (mol m\(^{-3}\) Pa\(^{-1}\)) of the facing side to form a concentration gradient in the corresponding PEM, which can cause gas permeation from one side of the membrane to the other in which the gas permeation rate \( (N_i) \) of species \( i \) through a membrane can be expressed as:

\[
N_i = D_i \frac{H_i^{p_1} \rho_i^{p_1} - H_i^{p_0} \rho_i^{p_0}}{\delta}
\]

In addition, the definition of \( k_i \), which is the gas molar permeability coefficient (mol m\(^{-1}\)s\(^{-1}\)Pa\(^{-1}\)), can be expressed as:
\[ k_i = D_i H_i \]  

(19)
in which \( D_i \) is the effective diffusion coefficient in \( \text{cm}^2 \text{s}^{-1} \), \( p_i \) is the partial pressure of gas (Pa), and \( \delta \) is the thickness of the membrane.

And as a key parameter in the above equations, the thickness of membranes can significantly affect performance in which for thick PEMs, water distribution becomes uneven, thus leading to uneven current distributions and current hot-spots as well as membrane swelling [106]. The thickness of PEMs can also cause ohmic loss, which can further lead to power loss. Therefore, although thick PEMs are usually used in EHCs due to their high mechanical strength under pressure, optimal PEMs in EHCs need to be simultaneously thin and robust.

Overall, all of the above controllable factors are crucial in the determination of hydrogen back-diffusion and the performance of EHCs. And in order to focus on electrochemical purification, EHC cells should operate with a zero total pressure gradient across the PEM. As for one-step EHCs, total pressure gradients can vary significantly between different experiments and because several cells are usually included in a single stack, even if overall compression ratios are specified, it is difficult to know the pressure difference on either side of PEMs for individual cells within the stack. Nevertheless, 5 MPa is commonly accepted as the large pressure gradient for low-temperature commercial PEMs whereas in laboratory settings, higher-pressure gradients can be achieved with higher compressions (a positive effect) due to the use of reinforced or high-temperature membranes. However, such large gradients can also cause increased gas permeation and back-diffusion (a negative effect). In summary, the proton conductivity, the mechanical property and the back-diffusion rate are three critical properties for PEMs in EHCs.

As for the membrane design of EHCs, it is based on predefined operating conditions (specifications), including inlet and outlet pressure, lifetimes, duty cycles, operating expense and capital expenditure, in which the total active area of membranes needs to be first defined followed by the consideration of operating current density, the compression ratio and optimal temperature for selected materials [84]. And based on a 3D graph of the properties of a proprietary low-temperature membrane visualizing energy requirement as a function of current density and the compression ratio (Fig. 3a), a positive correlation between energy requirement and current density exists based on Ohm’s law in which all energy is converted into heat at zero compression ratio (i.e., \( P_{in} = P_{out} = 10 \text{ bar} \)). At elevated cathode pressures, energy requirements also show a vertical tangent with low-current densities due to the compensation of the Faradaic compression rate by back-diffusion. And because these correlations are also appropriate for membrane thickness and materials, specific energy requirements can be optimized by adjusting membrane thickness or materials accordingly. Alternatively, high-temperature membranes suffer less hydrogen back-diffusion as compared with low-temperature membranes (Fig. 3b). In conclusion, the operation of lower current densities requires lower energy and corresponding membranes are more suitable for higher compression ratios. Based on these characteristics, several types of polymer membranes have been used in EHCs, including perfluorosulfonic acids (PFSA) (e.g., Nafion), ion-solvating polymers (ISPs) (e.g., polybenzimidazole), hydrocarbon polymers (e.g., sulfonated-polyether-ether-ketone) and proton-conducting oxides.

### 4.1.1 Perfluorosulfonic Acids

Currently, PFSA combined with hydrophobic backbones and hydrophilic side chains are the most widely used membranes in EHCs due to good chemical stability, high proton conductivities and excellent physical properties [107, 108]. Catalano et al. [109] were also able to formulate a theoretical framework with non-equilibrium thermodynamics to describe electro-kinetic effects in gas–liquid two-phased membranes based on Nafion 117. Despite this, the large-scale application of PFSA polymers in EHCs is limited by extensive hydrogen back-diffusion, management of the water content, low mechanical strength and high costs and up to now, the reduction of back-diffusion rates under high-pressure differences in EHCs remains difficult [104, 110]. Here, two main methods can be used to suppress hydrogen back-diffusion involving the use of thick membranes and the doping of materials to decrease the water content. For example, Analytic Power Corp. used Nafion 117 rather than Nafion 112 as a membrane in their EHC [111] because Nafion 112 with a thinner membrane led to larger hydrogen back-diffusion during high-pressure power consumption per unit hydrogen as compared with Nafion 117 due to higher resistivity. Stobel et al. [112] also studied the effects of PEM thickness on back-diffusion and found that PEM thickness was a crucial design parameter to balance back-diffusion and proton resistance and that increasing temperatures can lead to higher hydrogen back-diffusion rates. Alternatively, Sdanghi et al. [113] proposed that membrane thickness affected the water content, which in turn can influence the overall performance of EHCs. Hence, the influence of membrane thickness on back-diffusion needs to be comprehensively investigated. In a further study, Grigoriev et al. [83] found that zirconyl phosphate (ZP) particles can replace un-bonded water molecules in PEMs and this replacement is beneficial for the modification of membrane hydration levels to allow for the reduction in hydrogen back-diffusion (Fig. 3c). Despite this, these researchers also found that internal resistances increased as a result of the modification, which led to lowered compression efficiency in the corresponding EHC.
The addition of other inorganic compounds has also been explored to increase water uptake within polymer matrices, including SiO₂ [114], TiO₂ [115] and Al₂Si₂O₅(OH)₄·nH₂O [116]. Furthermore, researchers have reported that the use of low water content hydrogen as a feedstock can reduce the power input requirements for water evaporation and increase overall hydrogen compression efficiency [83]. Giner ELX Inc. also developed a modified PFSA membrane with a back-diffusion loss of 27% at 35 MPa in which as compared with an unmodified PFSA membrane, the back-diffusion rate in the modified membrane was reduced by more than 50% whereas the energy consumption of an EHC using the modified PFSA membrane was 2.0 kWh kg⁻¹ at 0.5 A cm⁻² under 35 MPa [82]. Moreover, McDonald et al. [117] reported that the mixing of a quaternary monomer can affect the polymerization of the monomer to exhibit increased strength and reduce permeability to water and gas as well as proton conductivity and acidity in which by modifying certain interior regions of the membrane, the membrane was divided into multiple sealed segments. Similarly, HyET hydrogen also mentioned membranes with high differential pressure characteristics in a patent filed in 2010 [118].

In general, the high-pressure conditions of EHCs can not only cause serious hydrogen back-diffusion issues but also
accelerate membrane degradation due to high mechanical stress to accelerate the chemical degradation of PFSA membranes in which resulting defects can be observed through optical imaging to identify in situ defect formation \cite{96} and from corresponding EHC performances. In addition, the release rate of fluoride in Nafion 117 at various applied pressures also possessed a positive correlation with compression pressure \cite{119} in which the deformation accumulation of membranes can lead to polymer decomposition and lowered activation energy in decomposition reactions \cite{119, 120}. Based on this, Greenway Inc. was able to develop a Nafion 117 membrane that can endure temperatures of 130–150 °C and up to 10 MPa for both EHCs and MHCs \cite{121}. And by considering the fact that the boiling point of water is 150 °C at 0.5 MPa, PFSA can potentially operate well at temperatures beyond 100 °C with stable proton conductivity \cite{122}.

More attention also needs to be paid to the separation of hydrogen from various gas mixtures in PFSA-based EHCs \cite{83, 123}. Here, Gardner et al. \cite{124} demonstrated that the issue of CO contamination can be resolved to improve hydrogen separation efficiency by periodically pulsing voltage to oxidize absorbed CO. Casati et al. \cite{125} also found that hydrogen recovery increased and performance coefficients (defined as the ratio of hydrogen produced to hydrogen consumed) decreased with differences in applied potential. Based on these results, optimal energy efficiency for separation should be provided and its functional dependence on process parameters in separating H₂ from H₂/N₂ mixtures should be identified. Here, Abdulla et al. \cite{108} developed single-stage and multi-stage gas recovery prediction models to study the separation of mixed gases, including CO₂, water vapor and hydrogen, and found that the multi-stage recovery model can achieve 90% energy efficiency and 98% hydrogen recovery. Using these models, these researchers were also able to confirm that membrane resistances and mass transfer coefficients of GDLs at the anode were two critical parameters of EHCs.

4.1.2 Hydrocarbon Membranes

Despite widespread usage, Nafion possesses inherent drawbacks such as high costs, limited operation temperatures and gas crossover \cite{126}. Alternatively, polyaromatic hydrocarbon membranes have emerged as popular candidates for EHCs, especially PEEK, due to low permeabilities (owing to narrower channels), high thermal and mechanical stability and low prices. In general, hydrocarbon membranes are synthesized through a simple sulfonation reaction involving the grafting of a sulfonic acid group into a polyether-ketone chain in which the resulting proton conductivity depends on the degree of sulfonation (DS). However, these hydrocarbon membranes usually possess poor proton conductivity due to the poor compatibility of imidazole ionic liquid-based composite membranes with Pt catalysts. In addition, continuous operations using these membranes often result in losses of ionic liquid. To resolve these issues, great efforts have been devoted to the investigation of improvement strategies for the various properties of hydrocarbon membranes (e.g., mechanical property, dimensional stability, proton conductivity) through methods including blending with other polymers \cite{127}, cross-linking \cite{128} and the addition of inorganic fillers \cite{129}.

The physicochemical characterizations of these modified membranes and corresponding EHC performances have also been widely investigated. For example, Rico-Zavala et al. \cite{130} studied membranes based on SPEEK prepared through the impregnation of Halloysite nanotubes with phosphotungstic acid (PWA) and Halloysite nanotubes (HNTs) and found that the incorporation of fillers can reduce water uptake and swelling (area and volume) through the effective modification of SPEEK membranes in which as compared with benchmark [S70] membranes (3.873 × 10⁻⁸ mol bar⁻¹ s⁻¹ cm⁻²), the modified [{S70/HNT₁₅} and [S70/(PWA/HNT₃₀₁₅)] membranes showed lower back-diffusion rates of 7.296 × 10⁻¹⁰ and 9.103 × 10⁻¹⁰ mol bar⁻¹ s⁻¹ cm⁻², respectively. These researchers also reported that proton conductivities were increased by 42% and 88% for membranes impregnated with HNTs and (PWA/HNT₃₀₁₅), respectively, and concluded that the presence of HNTs can improve the mechanical strength of composite membranes whereas the presence of PWA mainly gave rise to high proton conductivity, thus demonstrating that modified nano-composite membranes can present low-energy consumption at high pressure \cite{88, 130}.

Moreover, a “P29 series” hydrocarbon membrane developed by NEDO in Japan showed higher tensile stresses, elastic rates and better performances in EHCs than membranes developed in 2018 by Toray Industries (Fig. 3d) \cite{85} and Giner ELX Inc. developed multiple hydrocarbon membranes such as BPSH and biphenyl series membranes (BP-ArF₄, BP-ArSA, BP-SA) for EHCs to report that their biphenyl series membranes can display higher proton conductivities at lower ion exchange capacities with less swelling in water \cite{131}. In addition, Giner ELX Inc. also reported that their BP-ArF₄ membrane demonstrated a much reduced back-diffusion loss of 7% at 35 MPa as compared with a BPSH membrane (20%) and that an EHC using BP-ArF₄ and BPSH membranes exhibited the lowest cell voltage of 0.100 V cell⁻¹ at a current operating density of 1000 mA cm⁻² \cite{132}.

Many other types of membranes have also been explored of use in the specific work environment of EHCs. For example, to overcome the problem of water management, Yang et al. \cite{133} developed a water-free PEM that can operate in temperatures ranging from 120 to 180 °C in which at a low compression ratio of 1.5, a voltage of 0.073 V at 2 mA cm⁻² was obtained as well as a proton transport number in the
range of 0.17–0.20. Sustainable Innovations also conducted a cutting-edge study of phosphoric acid-doped aromatic polystyrene sulfonic acid (PA-APE) for EHCs with promising results [134] and Wu et al. [89] synthesized a semi-interpenetrating network (sIPN) containing a SPEEK membrane and a cross-linked polystyrene sulfonic acid. In this study, these researchers reported that the use of their SPEEK/CrPSSA membrane resulted in the rapid increase in current to the limited current, which indicated high mass-transfer resistance and that the energy efficiency of the corresponding hydrogen compressor was ~30%.

The study of durability has also been conducted based on realistic drive cycles. For example, Lipp et al. [135] completed a 10000 h durability test compressing hydrogen from 0.2 to 20.4 MPa and set a US compression record of 81.6 MPa with energy consumption of ~20 kWh kg⁻¹ in which a possible correlation existed between observed degradation behaviors and local mechanical stress experienced in EHCs.

4.1.3 Protonic Ceramic Membranes

Proton-conducting oxide membranes for EHCs can typically operate at high temperatures of 800–900 °C and therefore have gained attention in EHC processes focused on the purging of impurities in hydrogen. In addition, these materials can potentially be used as electrolytes for the separation of hydrogen from reformed natural gas despite low conductivities. For example, Catalano et al. [109] investigated Ba(Zr0.30Ce0.54Y0.15Cu0.01)O3−δ (LBYb-91) as a protonic ceramic membrane in which Y and Zr were added as dopants to improve chemical stability against steam and CO₂ whereas Cu was added as a dopant to enhance sintering ability and reported that the corresponding EHC demonstrated excellent performances at a current density of 2 A cm⁻² at 973 K. Sakai et al. [91] also found that La₀.₉Ba₀.₁YbO₃₋δ (LBYb-91) possessed high chemical stability against CO₂ contamination based on post-mortem XRD and TGA analysis and reported that the total cell voltage of an LBYb-91 EHC can reach ~0.7 V at 800 °C with an applied current density of 40 mA cm⁻². Despite these performances however, harsh working temperatures remain a serious issue that limits large-scale research and promotion as high-temperature EHCs. Regardless, the study of relationships between physical properties and nanostructures in protonic ceramic membranes such as proton mobility and gas permeability for EHC applications remains popular in the development of high-temperature EHCs. However, membranes designed for operation with pressure differences across membranes that are significantly affected by spacing, domain segregation, crystallization and the presence of water and electric fields were not further investigated. Overall, the meticulous design of protonic ceramic membranes with superior membrane support can significantly increase compression capability in which the fabrication of multilayered dense thin membranes with porous and robust support layers to reduce thickness is a viable strategy. In addition, the use of robust membrane supports can improve corresponding mechanical and thermal stability in membranes.

However, due to the lack of data concerning the membrane creep and long-term performance in the literature, the summarization of correlations between mechanical property and operational property in protonic ceramic membranes is difficult. As a result, significant research needs to be conducted on EHC protonic ceramic membranes in the future. In addition, due to intricate interactions among microscopic structures, membrane composition, reorganizations upon swelling and effective material properties (strain, stress, modulus) along with statistical fluctuations in pore ensembles, the comprehensive understanding of protonic ceramic membrane characteristics in EHCs also remains difficult.

4.1.4 Ion-Solvating Polymers

Due to PFSAs exhibiting large hydrogen back-diffusion rates as well as the release of corrosive gas after degradation, alternative membranes have been investigated in the last decade. Here, PBI is a high-temperature membrane with an operating temperature between 160 and 220 °C and therefore can demonstrate high-temperature operational capabilities. And because PBI membranes are generally doped with phosphoric acid to form an ion-solvating polymer (ISP), advantageous properties can be attained, including low back-diffusion [136], zero electron osmotic draw [137], high tolerance against impurities and facile water management in which the doped acid in PBI membranes is used to conduct protons. Because of this, the conductivity of corresponding membranes is directly dependent on the level of acid doping. And as result of this unique merit, acid-doped PBI membranes can operate at high temperatures (160–220 °C). However, because doped acids interact with PBI backbones through weak Van Der Waals forces, the progressive release of doped acids during EHC operation can also occur, which will cause gradual increases in internal resistance and significantly reduce the compression efficiency of EHCs.

More significantly, the higher operating temperatures achieved in the use of PBI membranes can enhance the tolerance of impurities such as CO and CO₂ [138–140] to reduce the inhibition of electrochemical activity for both anodic and cathodic reactions. For example, although industrial hydrogen is inevitably mixed with impurity gases, including CO or CO₂, the concentration of CO can be successfully reduced by more than 150 folds from 1906 to 12 ppm and the concentration of CO₂ can be successfully reduced by 32 folds and 62 folds from 11.9% to 0.37% and from 11.9% to 0.19%, at 0.4 and 0.8 A cm⁻², respectively, through the use
of corresponding EHCs [139]. Moreover, Thomassen et al. [141] found that the performance of a PBI membrane-based EHC supplied with a reformate mix containing 100 ppm CO was nearly identical to a cell running on 40% hydrogen in nitrogen, demonstrating that the CO content did not affect catalyst activity at low concentrations and that reformate species only had a diluting effect. These researchers also reported, however, that cells fed with reformate gas containing 1.36% CO can experience larger increases in cell voltage at high current densities despite increasing hydrogen concentrations, indicating that enhanced CO concentration in inlet gas can decrease EHC performance. Moreover, these researchers also found that the PBI membrane displayed excellent abilities to purify H2–CO2 mixtures in which a 21% CO2 content in a H2–CO2 mixture was reduced by over 98% at different current densities (Table 2). Strikingly, Robeson et al. also reported that both PFSA and PBI membranes can show enhanced performances in the separation of mixed gases at 230 °C, which is far beyond upper operating temperature limits. Sustainable innovations LLC also developed a phosphoric acid-doped polybibenzimidazole (PA-PBI) membrane for in-depth testing in which the resulting PA-PBI membrane was subjected to various testing conditions including the variable inlet CO2 content, the inlet relative humidity and the reaction temperature with Nafion 117 as a benchmark (Fig. 4) [134]. Based on the results however, these researchers concluded that due to the release of acid during long-running operation, their PA-PBI membrane was not a good choice for EHCs. And aside from the acid leakage of membranes, the control of the polyphosphoric acid content is also critical in PBI membranes in which effective catalyst areas can sharply decline due to the adsorption of concentrated phosphoric acid [142]. Furthermore, Pingitore et al. [143] synthesized a poly(2,2ʹ-(1,4-phenylene)5,5ʹ-bibenzimidazole) meta/para-PBI random copolymer through a poly (phosphoric acid) Table 2 Concentrations of CO2 in the cathode outlet and relative reduction in CO2 concentration in the separated hydrogen of a PBI membrane-based EHC [141]. Sources from the US Department of Energy

| Current density (A cm⁻²) | Cathode outlet CO₂ concentration (%) | Reduction in CO₂ concentration (%) |
|--------------------------|-------------------------------------|-------------------------------------|
| 0.5                      | 0.4 ± 0.05                          | 98                                  |
| 1.0                      | 0.21 ± 0.09                         | 99                                  |
| 1.5                      | 0.11 ± 0.04                         | 99.5                                |

Fig. 4 Parametric testing of an EHC with flow field modifications. Reprinted with permission from Ref. [134]. Sources from the US Department of Energy
process in which the compressive creep compliance was less than $2 \times 10^{-6} \text{ Pa}^{-1}$ and reported that this membrane exhibited a high proton conductivity of 150 mS cm$^{-1}$ at typical operating temperatures of 160–200 °C and an exceptionally low-voltage decay of 0.67 μV h$^{-1}$ at 160 °C for more than 2 years.

Researchers have also reported that gases mixed with water vapor can accelerate the leakage of doped acid in PBI membranes. For example, Bueltea et al. [142] reported a cell voltage increase from 15 to 450 mV with a supply gas dew point decrease from 60 to 20 °C in 250 h in which the power requirement of the un-humidified supply gas was 30 times higher than that of the humidified gas. Power consumption has also been found to be highly sensitive to water vapor pressure in gas stream supplies, which also depends on the concentration of phosphoric acid-based PBI membranes. However, Yang et al. [144] found that the existence of water vapor in feed gas streams did not significantly affect the permeability of H$_2$ and CO$_2$ or the selectivity of H$_2$/CO$_2$ due to the super thermal stability of their ZIF-8/PBI nano-composite material in hot steam, especially in 30% and 60% ZIF-8 composites. Perry et al. [139] also achieved long-term durability for nearly 4000 h under non-humidified and humidified conditions using their PBI membrane and attributed the lower-power requirement of their humidified operation to lower losses in the CL. Moreover, Buelte et al. [142] reported that anode side gases of PBI-based EHCs still require humidification with water vapor pressures greater than 55 mmHg (a dew point of 40 °C) to allow for efficient operations.

In terms of application, H2Pump LLC in 2012 was able to develop a third-generation PBI-based EHC with an area of 50 cm$^2$ that operated for more than 1000 h at a differential pressure of 2.275 MPa whereas a single-cell operated for 4000 h at the same pressure [145]. Greenway Inc. also applied an MHC and an EHC as smart technologies in a new hybrid solid-state hydrogen compressor system in which four constituent membranes [Nafion 212, PBI, an Advent membrane (a Pyridine-based aromatic polyether electrolyte) and a Fumatech membrane] were chosen as the membrane for the MEA of an EHC (Table 3) and found that although Nafion 212 in the EHC system possessed a slightly lower cell voltage than that of the PBI membrane, it did not provide any heat recovery opportunities for the MHC [131] (Fig. 5).

Overall, low operation temperatures can mitigate back-diffusion, particularly in situations (with back-diffusion of hydrogen gas) encountered with high-pressure differentials, thin thicknesses and high porosity in corresponding membranes. And in cases in which the hydrogen source is pure as a result of electrochemical water splitting, the mechanical properties of membranes are more important factors than the contamination of impurity gases in the use of high-temperature membranes. And based on overall design, the optimal balance of these characteristics is different for different applications. Furthermore, current membranes as reported by researchers have not been designed for operation under different pressures and therefore, membrane supports may be useful in the improvement of anti-compression ability. However, the absence of data on the creep and long-term properties of high-pressure membranes makes it difficult to establish accurate relationships between mechanical properties and membrane lifespans. As for the prevention of mechanical failure in ISP membranes, membrane clamping systems should be carefully designed to avoid uneven mechanical stress distribution and membrane reinforcements or modifications in polymer chemistry are also needed to reduce the creep characteristics of PFSA membranes. Here, membrane reinforcement can be achieved through the incorporation of inert, non-conducting reinforcement materials [58] and the combination of self-humidifying designs.

### Table 3 Membrane properties of an EHC system developed by Greenway Inc. Reprinted with permission from Ref. [121]. Sources from the US Department of Energy

| Metric                        | Nafion® | PBI Film® | Advent® | Fumatech® |
|-------------------------------|---------|-----------|---------|-----------|
| Operating $T$ (°C)            | 60–150  | 120–180   | 120–200 | 160–180   |
| Membrane thickness ($\mu$m)  | N212 = 50 | 70       | N/A     | 53        |
|                               | N211 = 25 |         |         |           |
|                               | N115 = 125 |        |         |           |
| Membrane area (cm$^2$)        | 500     | 500       | N/A     | 500       |
| Range of current density (A cm$^{-2}$) | 1–4     | 1–4       | 1–4     | 1–4       |
| Membrane conductivity (Ω cm$^{-1}$) | 0.5–0.11 | 0.1      | 0.08    | 0.1       |
| Electrolyte                   | PFSA    | PA        | PA      | PA        |
| Water crossover (kg H$_2$O kg H$_2$·h$^{-1}$) | 28.5    | Negligible | Negligible | Negligible |

### 4.2 Electrocatalysts and Catalyst Layers

Catalysts play an essential role in EHCs to decrease the activation energy of reactions and promote the electrochemical kinetics of both HOR and HER. Therefore, the category, the dosage, the structural design and the durability of catalysts for EHCs need to be further studied. And because gases, electrons and protons all participate in the electrochemical reactions of EHCs on catalyst surfaces in which all species can access, it is necessary to improve the microstructure of CLs to obtain...
high efficiency 3D contact points and avoid the low utilization of Pt catalysts. The loading of precious metal Pt in CLs remains a costly issue that severely hinders the popularization of PEM-based devices in which in the initially proposed concept of an EHC by Sedlak et al. [146] in 1981, a Pt-based catalyst with a loading of 4 mg cm\(^{-2}\) was required. However, additional research has significantly lowered this loading to 0.2–0.8 mg cm\(^{-2}\) [18, 107, 147]. To further reduce Pt loading, researchers have synthesized Pt nanocatalysts using atomic layer deposition [148] or nanostructured thin membranes with a loading of 0.1 mg cm\(^{-2}\) [149]. As a result of these findings, research has shifted away from the thickness and cost of electrodes to the uniform distribution of catalyst particulates. Furthermore, the study of catalyst loading limitations is also important in which the reduction in Pt loading from 1.1 to 0.2 mg cm\(^{-2}\) in both anodic and cathodic electrodes can lead to significant decreases in anode potential but no obvious changes at the cathode, indicating that HOR can significantly influence EHC performance [138]. Moreover, relatively few studies have been conducted in the optimization of Pt catalyst ratios in anodes and cathodes as well as the exploration of catalyst durability under high pressure. Researchers [147] have also detected that the equipotential line in thick membranes decreases, operating conditions: 160 °C, 0.2 A cm\(^{-2}\) and with pure hydrogen anode supply gas. Reprinted with permission from Ref. [142]; copyright of The Electrochemical Society. (d) \( \text{H}_2/\text{CO}_2 \) mixed gas permeation results of a ZIF-8/PBI nano-composite membrane (solid lines—\( \text{H}_2 \) and \( \text{CO}_2 \) permeability; dotted lines—\( \text{H}_2/\text{CO}_2 \) selectivity) [144]; copyright 2013 Elsevier B.V

In terms of carbon supports for EHC catalysts, the corrosion of carbon supports is not considered because the voltage required to drive reduction and oxidation reactions is by definition zero (except for a small overvoltage). However, characteristics such as high surface area, ability to maximize triple-phase boundaries through porous structures, good metal catalyst support interactions, high electrical conductivity and good water management need to be considered in the selection of carbon supports. Based on this, nano-carbon supports such as nanotubes, fibers and aerogels are promising replacements for traditional carbon supports for electrocatalysts and can be effectively used to reduce Pt loading. The alloying
of Pt catalysts with non-Pt group metals such as ruthenium, rhodium, iridium and osmium is also a promising strategy to decrease costs. And although the oxygen reduction reaction (ORR) demands high Pt catalyst loadings, there are still many reports for HER or HOR catalysts. Therefore, the holistic analysis of the catalytic activities of a single catalyst based on all reactions in a hydrogen electrode is important.

To further ensure the low loading of Pt, nanostructured CLs for MEAs in industrial production also need to be studied. For example, HyET hydrogen and NREL performed the X-ray fluorescence mapping and electrode structure characterization of HyET-produced electrodes for quality evaluation [96] and found that high-pressure environments can lead to the obvious destruction of CLs. Scanning electron microscopy and transmission electron microscopy also showed the degradation of CLs and the increased agglomeration of Pt nanoparticles at higher pressures. Therefore, the structural design of CLs plays a decisive role in durability [152]. Mass transport can also be affected by damages to catalyst structures under high-gradient pressures; however, Kim et al. [153] suggested that special pressure-resistant designs such as an inverse-opal-structured electrode (with a porous structure that can withstand high pressures and improve mass transfer and water management in CLs) can alleviate this issue. As for uneven reaction rates in anodes and cathodes, Taylor et al. [154] proposed that the optimization of Pt distribution and gradient loading in CLs is an effective solution. Here, these researchers showed that electrodes with a gradient design (Fig. 6b) were beneficial for the conduction of protons and the transport of different species such as gases or liquid water in which corresponding results revealed that the proper gradient distribution of Pt can effectively improve EHC performance, especially in areas of high current density, and significantly enhance mass transfer inside electrodes. Debe et al. [155] also reported that NSTF with an ordered microstructure can realize the separation and the order of electrons, protons and mass transfer channels to enable the improvement in catalyst utilization rates in electrodes and reduce the amount of Pt required. Moreover, carbon nanotube arrays as ordered catalyst carriers can also display high mechanical strength and conductivity; however, the corresponding stability is inferior water removal capability at low temperature [156]. Furthermore, metal oxides such as TiO2, WO3 and NbO2 can reportedly replace carbon carriers in which the design principle of the interface between the CL and the membrane is to increase adhesion and therefore improve cell durability. Other studies

![Fig. 6](https://example.com/fig6.png)

(a) Performance of catalysts developed by Giner ELX Inc. Reprinted with permission from Ref. [82]. Sources from the US Department of Energy. (b) Schematic of GDEs based on CLs with uniform and graded Nafion content: GDE-I (30 wt% Nafion), GDE-II (20/30/40 wt% Nafion) and GDE-III (40/30/20 wt% Nafion). Reprinted with permission from Ref. [190]; copyright of The Electrochemical Society. (c) Electrochemical yield versus space–time for two types of gas distribution plates (cell voltage=100 and 150 mV, \(T=298\) K). Reprinted with permission from Ref. [125]; copyright 2008 Elsevier B.V. (d) Baker diagram movement. Reprinted with permission from Ref. [111]; sources from the US Department of Energy. (e) Graph showing exemplary IV curves of identical MEAs testing hydrogen pumping performances in an EHC compression cell setup and in a PEMFC measurement setup. Reprinted with permission from Ref. [84]; copyright of CRC Press.
have also focused on CL designs [157] based on the shape of flow fields [158, 159]. More significantly, Giner ELX Inc. was able to enhance catalytic performances by threefolds at 1000 mA cm\(^{-2}\) (Fig. 6a) [82]. As for the design of nanostructures on CLs, ordered and gradient designs can be used in anodes to increase the three-phase interface of reactions to further optimize anodic water management and durability whereas proper high-pressure structural designs in high-pressure cathodes can reduce CL damage. In addition, researchers have suggested that the substitution of metal oxides for carbon can enhance adhesion between CLs and membranes [160, 161]; however, metal oxides can also offer high electrochemical stability and high surface areas, and mechanical stability and durability. To achieve distribution uniformity, EHCs also require a high degree of rigor in the spraying process in which plasma sputtering and electrodeposition methods are promising in the development of highly efficient catalyst systems that can deliver good performance and stability with long-term durability.

In actual application, hydrogen usually comes from coal gasification and methane reforming. Therefore, hydrogen is always mixed with small amounts of CO and CO\(_2\) that can poison catalysts in EHCs and decrease energy conversion efficiency in which small amounts of CO can be adsorbed onto Pt catalyst surfaces and compete with hydrogen adsorption to reduce the electroactive area of Pt. Here, Ru can be added to Pt catalysts to improve CO tolerance [124]. Based on this, Tokarev et al. [162] developed a CO adsorption model using the density functional theory to study CO tolerance capacities in which obtained experimental results showed that the adsorption energy of Pt\(_x\)Ru\(_2\) alloy was higher than that of PtRu\(_2\). These researchers also found that CO was readily adsorbed at the edge of Pt and that the (110) surface (with a small coordination number) of Pt atoms adsorbed CO stronger than the (111) surface. They were also able to use Ru to purify hydrogen from a mixture of methane and hydrogen. In another study, Kim et al. [163] prepared an Ir/C electrocatalyst to remove CO\(_2\) and compared its performance to that of traditional Pt/C catalysts. Here, these researchers reported that the exchange current density of the Ir/C catalyst treated at 300 °C (0.059 mA cm\(^{-2}\)) was larger than that of commercial Pt/C (0.043 mA cm\(^{-2}\)) and that through CO\(_2\) stripping analysis, the performance of the Ir/C catalyst was found to be unaffected by CO\(_2\) as demonstrated by the disappearance of the electrochemical desorption peak. To further avoid catalyst poisoning, MEMPHYS reported that catalysts can be renewed by using O\(_2\) or O\(_3\) in which O\(_2\) is a better choice due to its quick speed. However, the addition of O\(_2\) or O\(_3\) requires a second-stage EHC stack system to obtain pure hydrogen and maintain hydrogen output flows constant [21]. Overall, the addition of Ru and Ir to form Pt alloys can effectively inhibit CO adsorption and intake strategies should consider the addition of small amounts of O\(_3\) and O\(_2\) to oxidize CO in the consideration of the cost and complexity of catalyst preparation. In general, HER kinetics is slower than HOR kinetics and more attention needs to be paid to the enhancement of HER [123] in which HER has received great attention as a half reaction of electrolytic water due to its slow kinetics [164, 165]. However, challenges remain in the development of nanocatalysts to improve HER performance and current research on HER is mainly focused on Pt-based electrocatalysts, single-atom Pt catalysts, Pt-based alloys and non-Pt catalysts. However, more attention needs to be paid to the exploration of low-Pt or even non-Pt-based electrocatalytic materials to achieve low-cost, highly active and sustainable electrocatalytic materials for HER [166–168].

Although Pt-based catalysts are currently the main catalysts for HER, they also possess inherent problems mainly involving high prices and poor stability. Here, several strategies have been proposed to address these issues, including the downsizing of Pt to single-atom catalysts (SACs) [169] and the formation of alloys [170], all of which can significantly enhance catalytic performance by modifying catalyst surface hydrogen adsorption behaviors. Despite this, SACs possess relatively low-density active sites as well as an undesirable tendency toward aggregation. In addition, the preparation of noble metal-based alloys is a complex process in which many of the proposed bimetallic structures tend to change during reactions [171]. Therefore, the design of alloy catalyst atomic arrangements by using powerful characterization methods such as in situ spectroscopy is necessary. Furthermore, poor interfacial design strategies can lead to metal segregation and diffusion during long-term operation, meaning that the durability of Pt-based catalysts in EHCs is an issue that requires further research. Based on general physical and chemical properties, various earth-abundant and non-precious transition-metal-based compounds [layered transition-metal dichalcogenides (MoS\(_2\) and WS\(_2\)), transition-metal phosphides, sulfides, nitrides, etc.] have also been widely studied as promising electrocatalysts that are expected to replace noble metals in water electrolysis applications. And with inspiration from the structures and compositions of nitrogenase and hydrogenase, researchers have also exploited a series of metal sulfides as efficient electrocatalysts for HER [172], which is a profound achievement in the field of non-noble metal HER electrocatalysts.

As for MoS\(_2\) and WS\(_2\)-based materials, several designs involving nanoscale catalysts have been proposed to enhance catalytic activity based on instructional studies on active sites and semi-conductive natures in which as compared with original bulk materials, nanostructured [173, 174] and porous structured [175] materials were found to usually possess larger specific surface areas with higher densities of active sites. To further improve electrical contact in active sites, researchers have also proposed several methods such as doping with
suitable heteroatoms into base material lattices and the coupling of base materials with conductive species such as carbon nanotubes and graphene. Different from carbides and nitrides with relatively simple crystal structures (e.g., face-centered cubic, hexagonal close-packed or simple hexagonal), the crystal structures of phosphides are based on trigonal prisms due to the large radius of phosphorus atoms (0.109 nm) in which metal phosphides tend to form more isotropic crystal structures rather than the layered structures observed in metal sulfides [176]. And as a possible result of this structural difference, metal phosphides possess greater numbers of coordinatively unsaturated surface atoms than metal sulfides [177–179].

Many other materials have also shown great potential in HER catalysis, such as CoP [180], FeP [181], MoP [182, 183], MoN [184], WC [185] and MoSe2 [186], and overall, the design of catalysts has attracted special attention in the field of HER in the attempt to decrease or even replace rare noble metal electrocatalysts.

4.3 Flow Field and Gas Diffusion Layer

The flow field and the GDL of EHCs can affect the distributions of mass transportation and temperature and are crucial components in EHCs.

4.3.1 Flow Field

The main role of flow fields is the accommodation of mass distribution and transportation in porous electrodes with minimal pressure drops in which careful designs are required for flow fields at EHC anodes. Alternatively, porous electrodes can produce high-pressure hydrogen gas directly in EHC cathodes without the existence of a flow field.

Currently, most existing flow fields used in EHCs are similar to those used in PEMFCs and include single serpentine [125, 140, 187], pin-type (grid-type) [20] and parallel channel types. Here, Casati et al. [125] suggested that between single serpentine and parallel channel flow fields, the parallel channel flow fields showed slightly higher hydrogen recovery rates but low values of space–time, indicating the influence of different shapes in flow fields on performance under different pressures (Fig. 6c). In addition, Shimpalee et al. [188] reported that both the channel-land (rib) dimension and the shape of flow fields can have significant influences on the flow rate, the water removal, the pressure drop, the mechanical stability and the temperature distribution in which through the hydrogen void fraction in cathodes and the sensitivity study of the pressure drop, corresponding flow field channels can be determined. Bloomberg et al. [111] also used the variables involved in determining the location on a baker diagram [7, T_{cell}, T_{amb}, the pressure, amps per square foot (ASF), the void fraction and the channel area] to construct a spreadsheet to determine movement on the baker diagram as a function of individual parameters (Fig. 6d). Furthermore, Analytic Power Corp. reported that the main flow patterns found in their experiments were on the border of the bubble flow and the plug flow, which led to a decrease in the void fraction from 7.620 to 3.810 mm in which the channel depth was set to 0.127 mm and a higher current density of 350 A cm−2 as compared with 300 A cm−2 was selected based on expected performance improvements from their proposed design. Here, these researchers settled channels wider than lands with a ratio of 2 to 1 and reported that this ratio of land-to-channel can decrease pressure drops across the flow field and improve the water removal from the GDL to result in more contact area between reactants and the GDL as well as a reduction in total voltage loss [189]. Interestingly, Bessarabov et al. [84] reported that in two duplicate datasets obtained from the same membrane and catalyst, an expected linear trend was observed in a PEMFC whereas an EHC showed greater limitations above a certain current density followed by a leveling off, which suggested that the limitation was caused by the environment of the MEA (Fig. 6e).

4.3.2 Gas Diffusion Layer

GDLs are porous structures that play critical roles in EHCs in which common examples include carbon paper (e.g., Sigracet, Freudenberg, Toray) made by pressing carbon fibers and carbon cloth (e.g., ELAT) made by weaving carbon fibers and porous metal. GDLs should possess high electric conductivity, great mechanical strength, appropriate porosity and good chemical stability for high-performance EHCs. One function of GDLs is to support CLs and accommodate the diffusion of reactants, water, heat and electrons in which for the support of CLs, pressure differences between anodes and cathodes can lead to distortions on MEAs and flexible GDLs can play an important role in contacting CLs to avoid damage. GDLs can also facilitate diffusion across the channels of bipolar plates and CL membranes based on surface area and porosity in which increased surface areas of GDLs can increase the current between catalyst sites in MEAs and current collectors. As for the diffusion of water, GDLs can also handle the moisture control of EHCs in which GDLs can not only remove water outside CLs from the anode to prevent chamber flooding, but also maintain the wetting of CL surfaces to improve conductivity throughout the membrane. Moreover, GDLs also play an important role in the control of heat transfer during EHC operations. And based on all of this, the rational design of the configuration and coating of GDLs is critical to EHC performance.

Because of the obvious pressure difference between anodes and cathodes in EHCs, single and multi-layer porous metal materials [191, 192] as well as matched double GDLs (porous metal and carbon cloth) with carbon cloth [193] can be used as GDLs to support membranes on the anode...
side whereas for the cathode side, highly elastic materials can be used to provide shock absorption and supplement membrane deformation to inhibit the increase of contact resistance between GDLs and CLs.

As examples, a graded GDL with a porosity gradient possessing fewer numbers and larger sizes of holes as well as softer materials and rougher surfaces from the inside to the outside was used for the anode side by Panasonic Ip Management Ltd. (Fig. 7a) and Analytic Power Corp. considered a sintered titanium cell support on the anode side of an EHC to increase the percentage of holes in which the resulting active area exposed to reactant hydrogen was ~25% because of the porous support of the metal electrode [111]. These researchers also investigated porous metal supports that can be manufactured to be less than 0.127 mm thick and with up to 80% open ratios (Table 4) but reported that suitable corrosion-resistant materials were not found. As for double GDLs, porous metals can play the role of flow fluid and carbon cloth can be used to remove direct mechanical damage between porous metal and membranes. In terms of materials used in GDLs, sintered Ti is generally expensive and special designs with anisotropic pore sizes are even more costly. In addition, titanium can be oxidized by water and formed TiO2 layers are not conductive. Analytic Power Corp. also found that TiN coating layers used to protect Ti plates can be oxidized in EHC cell environments. Alternatively, researchers from both LANL and Giner Inc. [111] have suggested that optimal materials for electrode support screens in EHCs should be either molybdenum or niobium expanded metals. Toray Industries also investigated the effects of porous metal characteristics in EHCs including different thicknesses, porosity and shapes (Table 5) [85], and although many researchers have mentioned the use of porous metals in EHCs in their research, no detailed discussions have been provided.

Other than the development of GDL materials, the application of effective surface coatings is another important method to improve water management capabilities. For example, Lee et al. [193] used a low-cost fabrication method to prepare a hydrophobic coating consisting of Nafion and O-CNTs for porous GDLs to overcome anode flooding and reported that as compared with an EHC using an untreated GDL that could not recover from flooding at ambient temperature, an EHC using the GDL modified by O-CNTs and Nafion recovered completely in ~100 s. The use of high-elastic materials on the cathode side can compensate for the deformation of membranes and thereby reduce contact resistance between CLs and GDLs and provide shock absorption in which the degradation of EHC cell performance and damage to GDL microstructure are mainly caused by mechanical degradation [195].

Compression pressures can also strongly influence the properties of current collectors, CLs and bipolar plate interfaces, thereby affecting overall EHC performance in which optimal compression ratios can allow for maximized EHC performances. And because gas permeability decreases nonlinearly with GDL compressed thickness, differences in permeability can experience orders of magnitude in variation between compressed (250 μm) and uncompressed (380 μm) GDL sections. In addition, mass, heat and charge transport phenomena in non-uniformly compressed GDLs can considerably differ from those in uncompressed GDLs [196]. Here, an FEM model was adopted in many studies to obtain the properties and real shapes of elastically deformed GDLs as a function of clamping pressure and a species transport model was also applied to deformed GDLs to predict species distribution and cell performance [197, 198], thus allowing for greater insights into the effects of GDL compression on interfacial contact resistance, species transport and EHC performance.

In terms of GDL materials, common materials used in commercial GDLs contain carbon paper, carbon cloth or carbon felt. Here, comparisons of resistance and structural stability at high pressures revealed that increased load cycles can cause the gradual consumption of resistance and the cracking of fibers in carbon paper and carbon cloth whereas the stability of carbon felt performed much better in terms of electrical resistance and microstructure due to its tortuous and thick fibers (Fig. 7b, c). Moreover, the influence of GDL compression on carbon paper and carbon cloth was found to be relatively large under high current densities [194, 199]. Based on this, Giner ELX LLC proposed a mixture of chopped carbon fibers, milled carbon fibers and PVDF powders that was able to maintain above 90% original thickness under a pressure of 13.789 MPa [200]. Despite this, most studies concerning GDLs have been based on the clamping force of fuel cells and few studies have focused on the effects of high-pressure gas on GDL compression.

Overall, future research should focus on the exploration of new GDL materials such as Al/Mg/Si alloys and polymer-coated materials. In addition, the investigation of expanded metal screen electrode supports and metal-felt-based GDLs should be addressed in which metal felts can be effectively coated with imide or siloxane polymers. However, metal felts are designed for high differential pressures and corresponding membranes or GDLs may be pressed into channels under high-pressure situations. Therefore, the maximum pressure on EHC cathode sides is restricted.

5 System of EHCs

5.1 Thermal Management

An EHC is a fundamentally isothermal compressor in which temperature remains in principle constant during
Fig. 7  (a) A graded GDL with a porosity gradient developed by Panasonic Ip Management Ltd.; (b) cyclic and steady load effects on cell assembly with Tenax cloth: cyclic and steady load effects on resistance. Reprinted with permission from Ref. [194]; copyright 2018 Elsevier B.V. (c) Cyclic and steady load effects on cell assembly with Freudenberg carbon felt: cyclic and steady load effects on resistance. Reprinted with permission from Ref. [194]; copyright 2018 Elsevier B.V.
compression; however, Corgnalea et al. [80] reported that EHC stacks can potentially generate 4.75 kWh kg\(^{-1}\) of heat at 2.5 A cm\(^{-2}\). Based on this, proper thermal control is necessary to maintain constant performance and avoid temperature-induced variation in the cells of EHC stacks in which air convection cooling and liquid cooling are two main strategies. In addition, if cooling requirements are low at low current densities, convective air-cooling can also be used. As for single cells and short stacks, metal endplates can also act as radiators to dissipate excess heat from cells. Here, because of the existence of a parabolic temperature distribution in EHC stacks with the hottest part in the middle in which heat is transported to adjacent cells, the external heating of end plates can be used to increase external temperatures before operation to allow the temperature of central cells to be the lowest. And although external heating cannot correct the distribution of parabolic temperature during operation, it can affect the temperature offset of the overall stack profile and therefore increase middle cell temperatures as well. Imprecise designs can further lead to extreme temperature differences that can reach ten degrees between the hottest middle cell and the coolest external cell. However, as the number of cells increases, heat loss through the endplates remains almost the same whereas the demand for heat rejection increases proportionately. Sustainable Innovations also studied thermal management solutions for EHC stacks [201] and based on the assumption that heat was uniformly generated over the active area, a convection coefficient was estimated for free convection in a standard multi-cell stack in which the convective surface area of their stack was nearly 20 cm\(^2\) with the exposed ends of the cells used for cooling. As a result, these researchers reported that the protruding cells in their stack increased the effective convective surface area by nearly 17 times (Fig. 8a). Moreover, this company constructed thermal models with both thick and thin membranes for EHC stacks and found that temperatures can vary greatly with or without fins at different current densities (Fig. 8b). Aside from natural convection air-cooling, the circulation of cooling water is another common method to deal with thermal issues as well as provide humidification to cells. Concentration gradients can also be created through the addition of solutes (e.g., sugar) to cathode water to drive water formed in the anode compartment to the cathode. Despite these solutions, corrosion remains an issue that has not been resolved [111]. In addition, although proton conductivity, water management and back-diffusion are all affected by the temperature of EHC systems, no comprehensive mathematical models have been provided in literature focusing on the simultaneous simulation of these aspects in EHCs. Here, a model to predict the required level of thermal control in EHC systems is greatly beneficial to the avoidance of unstable and detrimental off-spec operating conditions.

### 5.2 Water Management

Water management is a key factor in the performance of EHCs because of the water content dependency of membrane conductivity, particularly for low-temperature EHCs. In addition, excess water can cover reaction sites on electrodes and hinder reactant flows through flooding to also affect EHC performance. Overall, water transport in EHC membranes occurs through three mechanisms, including electro-osmotic drag, concentration gradient induced diffusion and pressure gradient induced convection.

In terms of PEMFCs, modeling results have shown that performances can be improved by increasing differential pressure between cathodes and anodes to result in the increase of water back-diffusion across membranes [202]. In addition, several water transport models have been built based on PEMFCs [203, 204]. However, the scale of applied high-pressure differentials in PEMFC models may generate different predicted values from EHC physical models. And although Kusoglu et al. [205] discussed the internal balance between chemical and mechanical forces in the

### Table 4

| Dexmet product code | 1.3NB3-020F | 5NB5-031F | 10NB10-050F |
|---------------------|-------------|-----------|-------------|
| Base metal thickness (mm) | 0.033 | 0.127 | 0.254 |
| Material | Niobium | Niobium | Niobium |
| Strand width (mm) | 0.076 | 0.127 | 0.254 |
| LWD (mm) | 0.508 | 0.787 | 1.270 |
| Open area (%) | 50 | 50 | 31 |

### Table 5

| Test number | Thickness (mm) | Porosity (%) | Shape | Laser microscope image (μm) |
|-------------|----------------|--------------|-------|---------------------------|
| A           | 1.3            | 31.9         | Particle (size: 100 μm) | 25 |
| B           | 1.1            | 49.1         | Particle (size: 100 μm) | 50 |
| C           | 0.58           | 67.2         | Fibrous (size: 20 μm)   | 20 |
| D           | 0.01–1         | 10–20        | Etch holes               | No data |
determination of the water content and the $d$-spacing of water domains in PFSA membranes and showed that direct hydrostatic pressure confinement can reduce the water content by using small-angle X-ray scattering (SAXS), more experiments are needed to verify that decreases in the water content become less significant at 25 MPa. Casati et al. [125] also showed that the better control of the membrane hydration degree can be achieved by adopting variational operating conditions instead of galvanostatic ones and Onda et al. [187] found that both hydrogen flow relative humidity and current density distribution decreased along the channel direction during EHC operation because of unbalanced contributions from two main water transport mechanisms across the membrane (i.e., back-diffusion and electro-osmotic drag). This uneven distribution of transport mechanisms can also lead to localized membrane dehydration, which can increase local resistances. In a further study, Sdanghi et al. [206] developed a pseudo-2D model to prove that the stability of current density distribution strictly depended on the local water content of membranes but that contrary to PEMFC cathodes, no constant resupply of water was produced at the cathode side in EHCs in which the amount of water at the EHC cathode from electro-osmotic drag may not be enough to humidify the membrane, especially at low current densities and high temperatures. Therefore, EHCs need to be improved based on other water management strategies (internal humidification) due to the inlet of dry hydrogen aside from the strategy of humidifying inlet hydrogen.

Researchers have also reported that the use of internal humidifiers in cathode end plates is an improved design [207] as compared with humidification achieved through bubblers and that internal humidifiers can be used at both anode and cathode sides [83]. Unfortunately, water convection as caused by pressure gradients is usually ignored in these studies with more attention being paid on concentration gradients [147]. Analytic Power LLC also developed a hydraulic cathode that circulated water through the compressor cathode to cool the cell and humidify the membrane, thus accommodating the use of dry hydrogen in anodes and avoiding the fundamental problem of anode flooding, which is a principal challenge in prior designs. In this study, the diffusion of water was also accelerated through the addition
of a substance to create a concentration gradient in the water [111].

Aside from the humidification of EHC devices, Giner Inc. reported the development of a membrane called WaMM in which the composite membrane combined an ionomer with an electrical conductor composed primarily of carbon nanotubes that enabled improved water retention capacities in which water can be fed to one side of the WaMM and permeate through the membrane to the adjacent PEM, thus allowing water in the PEM to be self-replenishing due to the creation of a concentration gradient between the WaMM and the PEM. As a result, the developed WaMM with a high water flux (more than 0.10 g min⁻¹) can display 1.0 S cm⁻¹ of through-plane conductivity. And as assembled in a bipolar stack configuration, the WaMM experienced insignificant cell voltage losses of less than 6 mV cell⁻¹ at a current density of 1000 mA cm⁻² [133].

5.3 Models of EHC

Models of EHCs can allow for the analysis of influences from design parameters including electrode thickness, porosity and catalytic activity on cell performance. For example, because transport and electrochemical phenomena occur perpendicularly to the MEA direction in EHCs and significant variations in water and hydrogen concentration can occur as binary mixtures that flow along gas channels, 1D models cannot adequately describe the transport phenomena. Therefore, Nordio et al. [78] developed a 1D + 1D model of EHCs based on MATLAB by taking into account the changes of total pressure, total flow and hydrogen concentration in anodes and cathodes in which through the Butler–Volmer equation, simulations with different hydrogen concentrations for mixtures at different voltages can be performed. Here, a diffusive membrane model approach along with steady-state operations and gas phases was chosen in this model to treat the main system and therefore enabled the model to predict the polarization curve and purity of outlet water. Water content, pressure and back-diffusion effects on EHCs are also important research topics in which experiments at higher pressures and temperatures are needed to verify the accuracy of corresponding models. Here, Sdanghi et al. [208] developed a pseudo-2D model that was validated through comparison with obtained experimental data that can take into account both overall mass and energy balance in EHCs as well as heterogeneity in relative humidity along gas distribution channels. And by using this model, these researchers proved that the stability of current density distribution strictly depended on the local water content of membranes and was able to provide good predictions for both current density distribution and relative humidity at anode outlets under different current densities. These researchers also found based on this model that unstable operating conditions can occur if low relative humidity hydrogen was fed to EHCs due to water consumption at the cathode side. Sdanghi et al. [209] also investigated the uneven contribution of electro-osmosis flow and back-diffusion of water across membranes along with a series of parameters including humidity, temperature, membrane thickness, outlet pressure and the stoichiometric ratio in order to optimize overall efficiency.

Bampaou [210] also developed a zero-dimensional steady-state EHC model (temperature—up to 343 K; pressure—15 MPa) to calculate overall voltage by taking into account the Nernst voltage, ohmic loss and activation overpotential using commercial software (AspenPlus™). These researchers also used this model to evaluate required power and cooling loads based on water management and were able to quantify the extent of water flooding issues and liquid water that should be removed. In a further study, Rahbari et al. [211] combined working conditions and used molecular simulations and thermodynamic modeling to study the phase coexistence of H₂O–H₂ systems based on temperatures ranging from 283 to 423 K and pressures from 1 to 100 MPa. A thermodynamic model was also proposed with results that agreed with previous experimental data [112, 140]. Toghyan et al. [16] also compared a compressor with an ejector and observed that the performance of the ejector recirculation system was similar to that of the EHC recirculation system and that EHC recirculation system efficiency was higher than that of a mechanical compressor at low current densities (<0.1 A cm⁻²). These researchers also reported that at intermediate and high current densities, the performance of the ejector was better than that of the EHC because the operation of PEM fuel cells at higher current densities can lead to increases in parasitic power. Furthermore, this study focused on anode recycling systems and evaluated the effects of general parameters on PEMFCs, including operating temperature, pressure, relative humidity, the number of cells and active area of PEMFC stacks with EHCs.

5.4 Performance Testing of EHC Stack

The durability and voltage consistency of individual cells are key considerations in the design of EHCs in which the endurance of EHC cells is considered to be a critical parameter in life-cycle costs. Here, Fuel Cell Energy Inc. has devoted significant attention to the durability testing of EHCs [201] in which a developed single-cell system showed significant enhancements to endurance with operational time increasing from 6000 to 10,000 h at an elevated current density of 750 mA cm⁻² from 2012 to 2013. Endurance testing further shifted in 2014 and 2015 from hardware with 81 cm² (a 30-cell stack with endurance testing for more than 1600 h at 31.03 MPa) to 185 cm² with one cell operating for more than 22000 h with a hydrogen recovery rate above 95%. To
further demonstrate the robustness of their EHC, stress testing of similar single-cell stacks was also simulated, which underwent over 1000 pressure cycles with minimal losses in cell performance. And due to the efforts by Fuel Cell Energy Inc., the durability of their EHC increased by several orders of magnitude from 2007 to 2015 (Fig. 8c). Recently, Toray Industries also verified that their products can show durability above 5000 h at constant output pressures between 40 and 45 MPa without damage to cell components (Fig. 8d) in which in pressure variation tests between 30 and 40 MPa, no damage to MEAs or to cell components was found after 400 cycles (Fig. 8e). Despite this, Toray Industries also observed a worse phenomenon occurring in multi-cell EHC stacks in which in a 5-cell stack for example, the 5th cell was consistently almost twice the voltage of other cells. In addition, voltages were not evenly distributed among the cells in the 5-cell stack EHC, which ranged from 50 to 700 mV (Fig. 8f). And although the report did not mention the harm of this voltage inconsistency, the lifespan of individual cells may be threatened, which will subsequently affect the entire stack in a multi-cell EHC [85]. Furthermore, through the continuous modification of one design and improvements to fabrication and system controls, researchers were able to obtain a uniform cell-to-cell potential in a 30-cell stack across every cell in which within the 30-cell stack, variations were over 70% lower than previous stack designs after 4 years of improvements, suggesting that individual cells need to be designed carefully. In 2006, Barbir et al. [18] also evaluated performances using an EHC for the recirculation of hydrogen in a 10-cell stack and achieved satisfactory and stable results.

The current parameters and application of EHCs are shown in Table 6.

6 Conclusions, Challenges and Prospects

The issue of low hydrogen volume density in storage needs to be resolved for the future large-scale application of hydrogen as a widespread renewable fuel to replace traditional energy sources in which hydrogen compression is a direct approach. Here, commonly used hydrogen compressors can be divided into mechanical compressors and non-mechanical compressors in which mechanical compressors include widely used reciprocating and diaphragm compressors as well as novel ionic liquid compressors whereas non-mechanical hydrogen compressors include cryogenic compressors, adsorption compressors, metal hydride compressors and electrochemical hydrogen compressors. Based on this, this review has presented the advantages and disadvantages of these different types of compressors. In terms of EHC systems, this review has provided a comprehensive overview with discussions on key components and system management strategies. Here, EHC membranes require high proton conductivity, good mechanical integrity (i.e., durability, strength) and low hydrogen back-diffusion and major membranes explored include PFSAs, hydrocarbon membranes, protonic ceramic membranes and ion-solvating polymers, all of which have been described in detail in terms of both the current research status and existing challenges. As for CLs in EHCs, structural designs can reduce catalyst usage and potential failure in high-pressure situations whereas the reasonable design of three-phase interfaces can improve CL activities as well as mass transfer. And in terms of actual catalysts in EHCs, various metal catalysts can be used to purify hydrogen mixtures to reduce Pt inactivation and similar to the research direction in fuel cells and water electrolysis systems, the development of low-Pt and non-Pt catalysts remains the main goal in the future. With regard to flow fields and GDLs, the distribution of water and gas in flow channels and GDLs is a critical issue that is connected to geometric design, support material property and coating material in which the further exploration of high-pressure differentials in channels and GDL designs remains lacking. In addition, heat generation from cell resistance and excess driving voltage in large-scale EHCs also need to be reduced through air-cooling or liquid cooling methods to achieve thermal management in which models to predict the required level of thermal control to evaluate unstable or even detrimental off-spec operating conditions are necessary. Water management is also a critical focus in EHCs in which anode humidification alone is insufficient and requires water generation at cathodes as well at low current densities and high temperature. Here, main approaches include gas humidification, water admission in both electrodes and wetting coatings. Moreover, because the working principles of EHCs are different from that of PEMFCs, high-pressure differences can lead to different predicted values in physical models, which will require further exploration through experimental testing. In terms of the overall performance of EHCs, durability and voltage consistency are vital determinants to evaluate the proper design of EHC stacks in which the design of active area, pressure, temperature and other factors can affect the results of performance tests.

Overall, the main challenges of EHCs can be summarized as follows.

1. The design of membranes that can determine the performance of EHCs in which back-diffusion and mechanical strength are two critical parameters remains lacking. To resolve this, operation conditions and policies (the temperature, the outlet pressure, the membrane water content and the inlet flow) need to be considered and optimal combinations need to be elucidated. Here, a simple method to decrease hydrogen back-diffusion and enhance mechanical strength is to increase membrane thickness. As for uneven pressure distributions in membranes, membrane packaging designs should be
Table 6: Performances and characteristics in the application of various EHCs

| Author and Ref. (year) | Membrane electrode assembly (MEA) | Operating conditions | Performance in EHC |
|------------------------|-----------------------------------|----------------------|-------------------|
|                        | Membrane (thickness) | Electrodes (active area) | Proton conductivity (S/cm) | Back-diffusion | Mechanical strength | GDL | Temp (°C) | Gas composition | $P_{in}$ (MPa) | Flow (Nm³/h) | Applied voltage | Performance in EHC |
|                        | | | | | | | | | | | | | |
| Rohland et al. (1998) | ETEK MEA Nafion 117 | Pt, 2 mg/cm² (100 cm²) | No data | 24 °C: 0.08 ml/bar·min cm⁻²; 70 °C: 0.27 ml/bar·min cm⁻² | No data | Metal fiber matrix | 24, 70 | H₂: 0.1 | 4.3 | 0.04–5.7 | 100 mA/cm², 200 mA/cm², 350 mA/cm² | Efficiency loss of 5%, at 350 mA/cm², 1600 MPa at 24 °C, 46 MPa at 70 °C |
| Strobel et al. (2002) | WL GORE MEA 6000 (40 μm); ETEK MEA Nafion 105 (120 μm); ZSW MEA Nafion 115 (120 μm); ETEK MEA Nafion 117 (200 μm) | Pt, A/C: 167 μg/cm² (25 cm²) | No data | (1) 1.85 × 10⁻⁷ cm²/s; (2) 2.44 × 10⁻⁷ cm²/s; (3) 2.56 × 10⁻⁷ cm²/s; (4) 2.10 × 10⁻⁷ cm²/s | No data | No data | 24, 70 | H₂, H₂–N₂–CO₂ gas; 0.1–0.3 | 1.85 × 10⁻⁷ cm²/s; (2) 2.44 × 10⁻⁷ cm²/s; (3) 2.56 × 10⁻⁷ cm²/s; (4) 2.10 × 10⁻⁷ cm²/s | 200 mbar/cm² | 0.008–0.11 | 300 mV/cm² | 0.25V | 1 cell |
| Lee et al. (2004) | Nafion 115 (127 μm) | Pt, 0.4 mg/cm² (5 cm²) | No data | No data | No data | No data | 30, 50, 70 | H₂–N₂–CO₂ gas; 0.1 | No data | 0.008–0.11 | 300 mV/cm² | 0.25V | 1 cell |
| Analytic Power Corp (2006) | | | | | | | | | | | | | |
| Ibeh et al. (2007) | | | | | | | | | | | | | |
| Perry et al. (2008) | Nafion117 (279 μm, 381 μm, 431 μm) | Pt, 0.5 mg/cm² (25 cm²) | No data | No data | No data | No data | 20–70 | H₂–CO₂–CO gas; 0.1 | No data | 0.009, 0.018, 0.036 mL/s gas mixture of H₂ and methane | 190 mV/cm², 340 mV/cm² | No data |
| Gardner et al. (2007) | Nafion 15 | Pt, 1 mg/cm², Ru, 0.5 mg/cm², Pt, 1 mg/cm² (25 cm²) | No data | No data | No data | No data | 20 | H₂–CO₂–CO gas; 0.1 | 0.0195 | 0.00375, 0.00495 | 240 mV/cm² | Over 80% with 80% H₂ extraction |
| Bai et al. (2007) | | | | | | | | | | | | | |
| Onda et al. (2009) | | | | | | | | | | | | | |
| Nguyen et al. (2011) | Nafion117 SPE 40 wt% Pt on a Vulcan XC-72 carbon carrier of large (250 m²/g) (25 cm²) | Pt, 0.1 mg/cm² (50 cm²) | No data | No data | No data | No data | 20–70 | H₂–N₂–CO₂ gas; 0.1 | 0.00450 | 0.00564 | 200, 400, 800 mV/cm² | The content of CO and CO₂ decreased obviously |
| Grigoriev et al. (2011) | Nafion 117 (183 μm) | Pt/C, 0.4–0.8 mg/cm² (25 cm²) | No data | No data | No data | No data | 23–70 | H₂, H₂–N₂–CO₂ gas; 0.1 | 0.013 | 0.00228, 0.00228 | 0–50 or 500 mA/cm² | No data |
| Yang et al. (2011) | Membrane based on polyp-2-vinyldipyridinium dihydrogen phosphate (625 μm) | Pt, 8 mg/cm² (5 cm²) | 0.01 at 140 °C, The proton transport number 0.17–0.20 | No data | No data | No data | 120–180 | No data | 2.5 | No data | No data | No data |
Table 6 (continued)

| Author and Ref. (year) | Membrane electrode assembly (MEA) | Operating conditions | Performance in EHC |
|------------------------|----------------------------------|----------------------|-------------------|
|                         | Membrane (thickness) | Electrodes (active area) | Proton conductivity (S/cm) | Back-diffusion | Mechanical strength | GDL | Temp (°C) | Gas composition and $P_{in}$ (MPa) | $P_{out}$ (MPa) | Flow (Nm³/h) | Applied voltage or applied current density |
| Buelte et al. (2011) [142] | PBI | Brok Pt, 1 mg/cm² | No data | No data | No data | No data | 160 | H₂–CO₂ gas | No data | 1.2 X the requirement for the hydrogen pumping current | 200 mA/cm² | No data |
| Sakai et al. (2013) [91] | La₀.9Ba₀.1YbO₃₋ₓ, LBYb-91 (500 μm) | Porous Palladium | 0.0042 | No data | No data | No data | 800 | H₂ | No data | No data | 40 mA/cm² | High chemical stability towards CO₂ and H₂O |
| Kim et al. (2013) [138] | GORE MS5 series (25 μm), PBPA membrane (30 μm) | Pt/C, 1.1–0.2 mg/cm², 0.4–2.2 mg/cm², (25 cm², 10.24 cm²) | No data | No data | No data | No data | 160 | Humidified H₂–CO₂ gas | No data | 0.01308 | 800 mA/cm² | 80 mV at 800 mA/cm² |
| Wu et al. (2014) [215] | SPEEK/C/PSSA sIPN (100 μm) | Pt/C, Pt, 0.5 mg/cm², (1.9 cm²) | 0.08–0.1 | No data | No data | No data | 80 | Humidified 80% H₂, 20% CO₂ gas | No data | 0.00096 | No data | Energy efficiency of 30% |
| Lipp et al. (fuel cell energy) (2014) [135] | No data | No data | No data | No data | No data | No data | No data | H₂, 0.1–14 | 83 | 0.06 | No data | No data |
| Moton et al. (2014) [216] | Nafion® (127 μm) | Pt, 0.1 mg/cm², 200cm², 400cm² | No data | No data | No data | No data | 70 | H₂, 0.35 | 7–100 | 116–467 | 400 mA/cm², 1000 mA/cm² | No data |
| PHAF-DRUS project (2014) [217] | No data | No data | No data | No data | No data | No data | 20.80 | No data | 100 | 0.93 | No data | No data |
| DON QUI-CHOTE project (2014) [217] | No data | No data | No data | No data | No data | No data | 20.80 | No data | 40 | 28 | No data | No data |
| Huang et al. (2016) [218] | SPPESK (70 μm) | Pt/C, Pt, 0.5 mg/cm², (1.9 cm²) | 0.0338 | No data | No data | No data | 40 | Dry 25% H₂, 75% CO₂ gas | No data | 0.003 | 56.7 mA/cm² | > 9999% H₂, Energy efficiency: 35% at 60% H₂ |
| Hao et al. (2016) [147] | Nafion®117 membrane | Pt/C, Pt, 0.2 mg/cm², (5.3 cm²) | No data | No data | No data | No data | 30, 60 | H₂, 0.1 | 2 | 0.00132 and 0.00240 | 100–500 mA/cm² | 500 mA/cm² > 95% |
| HyEt Group (2017) [219] | No data | No data | No data | No data | No data | No data | No data | H₂, 0.1 | 70–100 | 1 | No data | No data |
| Author and Ref. (year) | Membrane electrode assembly (MEA) | Operating conditions | Performance in EHC |
|------------------------|----------------------------------|----------------------|---------------------|
| **Author and Ref. (year)** | **Membrane (thickness)** | **Electrodes (active area)** | **Proton conductivity (S/cm)** | **Back-diffusion** | **Mechanical strength** | **GDL** | **Temp (°C)** | **Gas composition and \( P_{in} (\text{MPa}) \)** | **Flow (Nm\(^3\)/h)** | **P_{out} (MPa)** | **Applied voltage or applied current density** |
| Choi et al. (2017) [92] | Ba\((Zr_{0.30}Ce_{0.54}Y_{0.15}Cu_{0.01})O_{3−}\delta\) (85 μm) | N-doped LaVO\(_3\) and Cu and Y-doped Ba \((\text{Ce, Zr})O_3\) (both 0.080 mm) | No data | No data | No data | No data | 600, 700 | Humidified 50% \( H_2 \) gas balanced with He; 10 sccm | No data | No data | 1000 mA/cm\(^2\) Faraday efficiency of 84%; Applied voltage: (1) 1.07 V, (2) 0.51 V |
| Greenway Energy (2017) [144] | Nafion N212 (50 μm), N211 (25 μm) N115 (125 μm); PBI Film (70 μm); Advent; Fumatech (53 μm) | No data (500 cm\(^2\)) | 0.5–0.1, 0.1, 0.08, 0.1 | No data | No data | No data | No data | No data | No data | No data | No data | 1000–4000 mA/cm\(^2\) Evaluation in area and volume swelling, water uptake, proton conductivity |
| Rico-Zavala et al. (2018) [130] | (1) SPEEK, (2) SPEEK/HNT\(_x\), (3) SPEEK (PWA/HNT\(_x\)) | PVC, Pt, 0.5 mg/cm\(^2\), (25 cm\(^2\)) | (100% relative humidity): (1) 0.01158, (2) 0.00967, (3) 0.02037 | No different pressure: (1) \( 2.25 \times 10^{-6} \text{mol}^{-1} \text{cm}^{-2}\) (2) \( 5.0 \times 10^{-6} \text{mol}^{-1} \text{cm}^{-2}\) (3) \( 1.0 \times 10^{-6} \text{mol}^{-1} \text{cm}^{-2}\) | 29 MPa | Sigracet SGL24-RC | 30, 60, 80 | 20–80 | \( H_2 \) \( 80% \) \( N_2 \) \( 20% \) gas; \( H_2 \) \( 50% \) \( N_2 \) gas mixture, 0.105–10 \( 20 \) \( H_2 \): 0.0156, \( N_2 \): 0.0188, \( H_2 \): 0.0188, \( N_2 \): 0.0188 | No data | Compression of the purified \( H_2 \); up to 20 MPa |
| MEMPHYS (2019) [21] | No data | No data | No data | No data | No data | No data | 20–80 | 87.5 | 5.5617 | 1000 mA/cm\(^2\) At 35MPa ≤ 0.250 V, ≥ 1000 mA/cm\(^2\) |
| Giner ELX (2019) [219] | PFSA, BP-ArF4 | No data (50–300 cm\(^2\)) | 0.106 | No data | No data | Scaling 140 MPa | No data | No data | \( H_2 \): 0.1–10 | 87.5 | 5.5617 | 1000 mA/cm\(^2\) The power consumption: 40% less than that of the mechanical compressor (0.29 kW/Nm\(^3\)) at 500 mA/cm\(^2\) |
| Toray Industries (2019) [85] | Nafion NRE212, Toray20 μm, 50 μm | Pt, AC: 0.3 mg/cm\(^2\); 0.7 mg/cm\(^2\); \( 100\text{mm}^2\) | 80°C, 80%RH | Nafion 0.11; Toray 0.22 | 0.5 MPa different pressure 80°C, 90%RH, Nafion 5.5 \( 10^{-9} \text{cm}^2/\text{s}\) cmHg, Toray 5.7 \( 10^{-9} \text{cm}^2/\text{s}\) cmHg | 35 MPa | 30–80 | \( H_2 \) | 45–82 | 0.005, 0.55 | No data |
| Pingitore et al. (2019) [220] | meta/para-PBI, meta-PBI (104 μm, 39 μm) | No data >0.150 | No data | The compressive creep compliance \( <2 \times 10^{-4} \text{Pa}^{-1} \) | No data | No data | 160–200 | Humidified \( H_2 \) and reformate test gas | No data | No data | 200–1000 mA/cm\(^2\) No data |
| Sdanghi et al. (2019) [206] | Nafion XL, Nafion 117 (30 μm, 175 mm) | No data (30 cm\(^2\)) | No data | 2.53 \( 10^{-7} \) | No data | 60 | Sigracet (240 mm), a gold-coated porous titanium (PTL) pore size: 5 mm and thickness: 1 mm | Humidified \( H_2 \) (0.1 MPa) | 3.2 | 0.0026–0.0103 | 0–1000 mA/cm\(^2\) 14–56% |
emphasized in which membrane segment designs are promising. As for high-pressure conditions in which membranes are sandwiched in the middle, the edges of membranes are subjected to stronger mechanical forces as compared with the middle and can break. Here, clamping systems for MEAs need to be carefully designed to avoid uneven mechanical compression based on force analysis. And equipped with locally strengthened segment-designed membranes, the reliability of membranes can be ensured with high-pressure difference loading. In practice, multi-stage compression is also promising for modularity, easiness in expansibility and so on.

2. The design of CLs in EHCs requires further attention in which well-designed CLs can reduce reaction times, improve overall efficiency and lower startup costs. In terms of costs, the development of non-Pt catalysts is also important in which the tailoring of catalyst morphology and the control of accessible active sites are required to achieve comparable activities to Pt catalysts. In addition, the design of high-pressure CLs is also important in which CLs need to maintain high specific surface areas and stable structures under high pressure in EHCs because catalyst particles can easily agglomerate under high pressures and CLs can drop off or crack. And although current research into pressure-resistant nanostructures for either PEMFCs or PEMWEs is rare, it should become a promising direction for EHCs. Moreover, the reduction of noble metal loading and the exploration of appropriate coating methods on CLs should be given more attention and the research of non-Pt catalysts is imperative in which overall costs can be substantially reduced. Furthermore, the performance and durability of EHC catalysts will require further attention.

3. The search for water management strategies in EHCs is vital. Unlike fuel cells and water electrolysis systems, EHCs do not involve water as a reactant or a product. Therefore, effective measurements to add water and maintain water in EHCs to achieve balance are required. And although many methods have been proposed to add water to EHCs, the easiest is to periodically add humidified hydrogen from the anode side. Water conduction models are also required to allow for the better understanding of EHCs in many conditions, especially for high-pressure conditions.

In general, the main barriers of EHCs for practical application and commercialization involve technical issues (low robustness, reliability and durability) and component costs. And although EHCs possess great potential and have attracted increasing attention, the determination of device durability, reliability and robustness remains difficult through the large amount of available experimental data. However, practical experimental testing remains the most effective approach to achieve incremental progress in the resolution of these issues. And due to the rapid development of fuel cells, corresponding technologies can potentially be applied to EHCs as well and in the foreseeable future, the issues of materials, chemistries and water management in EHCs will gradually be resolved. As for the commercialization of EHCs, it will depend on the research and development of overall hydrogen systems based on a future hydrogen economy that includes public acceptance and low overall costs. Ultimately, EHC technologies with superior, maintenance-free life cycles will become a compelling choice over existing solutions in the future.

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