

ELECTROLYZERS *WORKING PRINCIPLES AND CHARACTERIZATION*

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Hydrogen can be produced by both renewable and non-renewable sources of energy. Nowadays, the most predominant methods for producing hydrogen still rely on non-renewable energy sources (> 90%). One method by which greenhouse gases emission due to hydrogen production can be minimized is by using solar or some other form of renewable energy source as the primary energy requirement for the hydrogen production.

HYDROGEN PRODUCTION – THE EU PLAN

European Commission 2020

HYDROGEN PRODUCTION – THE EU PLAN – HYDROGEN VALLEYS

A hydrogen valley is a regional initiative that aims to develop a hydrogen ecosystem, including hydrogen **production, storage, distribution, and utilization**, within a **defined geographical area**. The idea behind a hydrogen valley is to create a localized cluster of companies, research institutions, and public authorities that work together to promote the use of hydrogen as a clean energy vector.

Weichenhain U. *et al,* (2021) Insights, Luxembourg: Publications Office of the EU

IIVERSITÀ EGLI STUDI

North Adriatic Hydrogen Valley

HYDROGEN PRODUCTION WITHIN THE HYDROGEN VALUE CHAIN

https://www.idtechex.com/es/researchreport/hydrogen-economy-2023-2033 production-storage-distribution-andapplications/946

GREEN HYDROGEN PRODUCTION METHODS

Table 2.1 Classification of green hydrogen production methods

GREEN HYDROGEN PRODUCTION METHODS

Electrolysis is a technique that uses direct electric current to drive an otherwise non-spontaneous chemical reaction.

It is already used for different industrial purposes:

- In metallurgy, for extracting metals from ores, for metal purification, for metal deposition, for metal manufacturing, etching or polishing.
- For production of chemical compounds, such as NaOH, NaClO₃, KClO₃, or perfluorinated organic compounds, in example.
- For hydrogen production, starting from H_2O .

WATER ELECTROLYSIS

PEMWE CELL / CONSTITUENTS

Feng Q., et al., Journal of Power Sources, 366, 2017, 33-55

PEMWE CELL / CL

Kim *et al. Nat Commun* 4, 2473 (2013).

The CL has to be electronically conductive, chemically stable, its morphology has to promote gas diffusion. There have been three progressive improvements in MEA architecture:

- 1. CL on GDL (ease of manufacturing, reduced stability of catalyst nanoparticles)
- 2. Catalyst-coated-membrane-MEA (precious metal utilization is optimized and proton conductive resistance is reduced) – standard
- 3. Long-distance-ordered-structure-support for hosting the catalyst

PEMWE CELL / CL

The GDL/PTL should be:

- flat to make GDL contact CL closely, leading to uniform dispersion of the current on the electrode
- porous to provide enough mass transfer channels
- mechanically stable and made of certain elastic to meet the requirements of seal and assembly
- of low resistance for reducing current loss;
- anti-oxidative to enhance the durability of the MEA

Common materials used: Carbon paper or Titania-based microporous layers

The electrolyte should be:

- chemically and mechanically stable in the operating environment of the electrolyser
- good ionic conductor
- poor electron conductor
- avoid gas crossover

PROTON EXCHANGE MEMBRANE WATER ELECTROLYZER (PEMWE)

- PEM is usually constituted by Nafion, which is able to support high proton conductivities
- Catalysts are still expensive: Pt/C (HER), IrO₂ (OER)
- Cathode (HER): $4H^+ + 4e^- \rightarrow 2H_2$
- Anode (OER): $2H_2O \to O_2(g) + 4H^+ + 4e^-$
- Total: $2H_2O \rightarrow O_2 + 2H_2$

https://en.wikipedia.org/wiki/Polymer_elect rolyte_membrane_electrolysis

PROTON EXCHANGE MEMBRANE WATER ELECTROLYZER (PEMWE)

- They operate at low T (50-80 $^{\circ}$ C) and P (20-50 bar).
- They operate at high current densities: $1 2$ A/cm² with a hydrogen rate (per stack, around 400 Nm³ /h)
- Specific energy consumption is around $4 6.5$ kWh/Nm³
- Efficiency in commercial stacks is around 70% with a system efficiency around 50%.
- Stack lifetime can reach up to 100000 hours with performance losses around 0.5- 2.5%/year
- Investment cost: 1500-3800 \$/kW, with annual maintenance cost about 4% of the initial investment https://en.wikipedia.org/wiki/Polymer_elect

ALKALINE WATER ELECTROLYZER (AWE)

- A OH⁻-conducting diaphragm is needed for separating the anode and cathode.
- Electrolyte: KOH solution; keeps hydrogen and oxygen separated to prevent their recombination.
- Ni and Fe are used as catalysts for cathode and anode, respectively.
- Cathode (HER): $4H_2O(l) + 4e^- \rightarrow 2H_2(g) + 4OH^-(aq)$
- Anode (OER): $4OH^{-}(aq) \rightarrow O_{2}(g) + 2H_{2}O + 4e^{-}$
- Total: $2H_2O \rightarrow O_2 + 2H_2$

Chowdhury, American Journal of Renewable and Sustainable Energy Vol. 4, No. 3, 2018, pp. 40-46

ALKALINE WATER ELECTROLYZER (AWE)

- They operate at low T (60-90 $^{\circ}$ C) and P (10-30 bar).
- They operate at low current densities: $0.2 0.6$ A/cm² with a high produced hydrogen rate (per stack, around 1400 Nm³/h)
- Specific energy consumption is around $4 6$ kWh/Nm³
- Efficiency in commercial stacks is around 70% with a system efficiency around 50%.
- The electrolyte is corrosive at operating temperatures: this strongly limits their lifetime
- Stack lifetime can reach up to 120000 hours with performance losses around 0.25-1.5%/year
- Investment cost: 850-1500 \$/kW, with annual maintenance cost about 2% of the initial investment

Chowdhury, American Journal of Renewable and Sustainable Energy Vol. 4, No. 3, 2018, pp. 40-46

SOLID OXIDE ELECTROLYZER CELL (SOEC)

Bessarabov, Millet, 2018. PEM water electrolysis. Academic Press.

- Solid electrolyte is composed by thin film ceramics $(e.g., Y₂O₃)$ doped ZrO₂)
- Ni or Ni-doped ceramic materials are used as electrodes

• Catholic:
$$
2H_2O + 4e^- \rightarrow 2H_2 + 2O^{-2}
$$

- Anode: 2 $-2 \rightarrow O_2 + 4e^-$
- Total: $2H_2O \to O_2 + 2H_2$

SOLID OXIDE ELECTROLYZER CELL (SOEC)

Bessarabov, Millet, 2018. PEM water electrolysis. Academic Press.

- They operate at high temperatures ($700 1000$ °C) and low pressures (1-15 bar).
- Applied current density ranges between 0.3 an 1.0 A/ $cm²$ with resulting low rate of produced hydrogen (per stack, around 10 Nm³/h)
- Specific energy consumption is around $3 4$ kWh/Nm³
- Efficiency in commercial stacks is around 100% with a system efficiency around 80%.
- Thermal and mechanical stresses strongly jeopardize the lifetime of the device.
- Stack lifetime can reach up to 20000 hours with performance losses around 3-50%/year
- Investment cost: <2200 \$/kW

WATER ELECTROLYZERS / COMPARISON

PEMWE / OPERATING POINT

Feng Q., et al., Journal of Power Sources, 366, 2017, 33-55

The operating point of an electrolyzer is defined by the values do voltage and current applied. Nonetheless, being the water splitting reaction not spontaneous, applied voltage must satisfy some boundaries.

By considering that the water splitting reaction is characterised by a variation of Gibbs free energy (the amount of energy involved in a reaction) equal to ΔG = 237.22 kJ/mol, from the relation among the variation of Gibbs free energy and the so-called standard potential, E^0 :

the potential needed to start the reaction is:

$$
E^0 = \frac{\Delta G}{nF} = 1.23V
$$

However, from a thermodynamic point of view, the water splitting reaction at room temperature and pressure can be expressed as :

$$
H_2O + \Delta H \rightarrow 1/2O_2 + H_2
$$

Where ΔH can be expressed, by means of the second law of thermodynamics as:

 $\Delta H = \Delta G + T \Delta S$

Here, ΔG is the Gibbs free energy of the reaction and it defines the minimum energy required to induce for water splitting, while ΔS describes the variation of entropy due to temperature changes.

For liquid water, ΔS =163.15 J/(mol K), ΔH=285.840 J/(mol K), leading to the definition of the socalled thermo-neutral voltage:

$$
E_{TN} = \frac{\Delta H}{nF} = \frac{\Delta G + T\Delta S}{nF} = 1.48V
$$

The voltage efficiency of operation of an electrolyzer, is thus defined by the ratio among:

$$
\varepsilon_V = \frac{V_{TN}}{V}
$$

In addition to this, in order to determine the amount of hydrogen produced by an electrolyser, the Faraday's law can be used:

$$
\dot{n} = \frac{dn}{dt} = \frac{I}{zF}
$$

From the Faraday's law the faradaic efficiency can be defined as:

$$
\varepsilon_F = \frac{\dot{n}zF}{I}
$$

And the efficiency of the electrolyser can be calculated by:

$$
\eta=\varepsilon_V\varepsilon_F
$$

ELECTROLYZERS / OPERATING CONDITIONS

Generally the applied voltage has to be higher than the voltage defined form theoretic calculation:

$$
V = V_{TN} - V_{ACT} - V_{OHM} - V_{TRN}
$$

Where V_r are different types of voltage losses:

 V_{ACT} : losses due to the activation overpotential

 V_{OHM} : losses due to ohmic overpotential

 V_{TRN} : losses due to mass transport

PEMWE / ACTIVATION LOSSES

Activation overpotential

• Related to the additional energy which has to be provided at the system in order to start the HER and the OER

 $sinh^{-1}$

 -5

- Directly affected by the temperature, catalyst material, utilization, and loading
- The can be modelled by means of a simplified version Butler-Volmer equation

$$
V_{ACT} = \frac{RT}{\alpha} \operatorname{arcsinh}\left(\frac{i}{i_0}\right)
$$

Where α is the charge transfer coefficient and i₀ is the exchanged current density

Ohmic losses

Related to electronic and ionic transport; it is due to the resistance to the flow of electrons through the current collectors and separator plates, as well as the conduction of protons through the membrane. They can be described by means of the Ohm's law:

$$
V_{OHM} = R_{TOT}I + \frac{\delta_m}{A\sigma_m}I
$$

Where $\delta_{\rm m}$ is the membrane thickness and $\sigma_{\rm m}$ its proton conductivity

Mass transport losses

They are related to matter flow through a porous media and can be related to different phenomena:

- flow restriction to the catalyst sites
- gas bubbles formation from the reaction products
- surplus of reaction products at the catalyst sites blocking the reactant diffusion.

Diffusion is described by the Fick's law: $J=-D_{eff}\left(\frac{\partial c_i}{\partial x}\right)$, where J is the diffusion flux, D is the diffusion

coefficient and C the specimen concentration. Mass transport losses are related to the variation of specimen concentration at the membrane-electrode-interface (C_{mei}) by means of the Nernst equation:

$$
V_{TRN} = \frac{RT}{nF} \ln \left(\frac{C_{i,mei}}{C_{i,mei,0}} \right)
$$

ELECTROLYZERS / OPERATING CONDITIONS

PEMWE CELLS

PEMWE STACKS

While designing a PEMWE stack, homogeneous water and current distribution and appropriate compression in the cell and stack must be obtained. The stack must be compressed with an appropriate pressure to make sure the cell sealed and to avoid gas leak, or MEA crack resulted by lower or higher torque, as well as Ohmic losses due contact resistance decreases the stack efficiency

PEMWE SYSTEM

https://www.ise.fraunhofer.de/en/business-areas/hydrogen-technologies-andelectrical-energy-storage/electrolysis-and-power-to-gas.html

BALANCE OF PLANT

Composition: $H₂O$ subsystem H₂ subsystem Cooling subsystem Control subsystem

BALANCE OF PLANT / WATER SUBSYSTEM

BALANCE OF PLANT / HYDROGEN SUBSYSTEM

It takes advantage by the pressure difference in water contained in the form of moisture to dry the hydrogen. Once a high humidity gradient is reached in the high pressure separator, the wet (*dirty*) hydrogen flows into the low pressure separator, while the dry (*clean*) hydrogen continues to the dryers' stage

WATER ELECTROLYZERS / INDUSTRIAL PERSPECTIVES

UNIVERSITÀ Dipartimento di **DEGLI STUDI** Ingegneria **DITRIESTE** e Architettura

WATER ELECTROLYZERS / INDUSTRIAL PERSPECTIVES

WATER ELECTROLYZERS / BOTTLENECKS

PEMWE CELLS / POSSIBLE IMPROVEMENTS – MATERIAL SIDE

- **Improve the catalytic activity** for HER and OER by reducing catalyst load (in example by introducing binary, ternary or quaternary alloys with an advanced material design).
- **Improve the electrochemical active surface area**, catalyst utilization, **and stability** against corrosion. In the case of using supported catalysts, development of highly conductive supports that can sustain the corrosion environment and still provide high nanoparticle dispersion and homogeneity.
- Improve the **proton transport** across the catalytic layer, decrease gas crossover across the electrolyte and nanoparticle hindering, diminish the electronic resistance provided by the ionomer, improve the water transport across the triple-phase-boundary.

PEMWE CELLS / POSSIBLE IMPROVEMENTS (2)

- Develop low cost current collectors with tuned porous structure, high corrosion resistance, low ohmic resistance, and optimized mass transport
- Titanium-made separator plates could be replaced by using lower cost materials (e.g.: copper, graphite, stainless steel) coated with high electron transport and high corrosion resistance materials.
- Model the multi-phase transport of species through the current collectors and separator plates. This could prove to be very beneficial in the design of current collectors, especially for larger scale electrolysers.
- Develop a predictive model for the exchange current density for various catalysts. Although this task is very demanding, if successfully accomplished, modelling would prove to be a much more useful design tool capable of aiding in the design of all the individual components.

Books/Book Chapters

- Electrochemical Water Electrolysis, Fundamentals and Technologies, 2020 Taylor & Francis Group, 78-1-138-32932-4, chapter 1 (Fundamentals of water electrolysis - Xiaoxia Yan, Rida Javed, Yanmei Gong, Daixin Ye, Hongbin Zhao)
- Erik Wolf, Electrochemical Energy Storage for Renewable Sources and Grid Balancing, Chapter 9 Large-Scale Hydrogen Energy Storage, doi.org/10.1016/B978-0-444-62616-5.00009-7
- Solar Based Hydrogen Production Systems, Dincer, Joshi, 2013, Springer, 978-1-4614-7430-2

Optional

- Carmo, Stolten *et al.*, A comprehensive review on PEM water electrolysis, International Journal of Hydrogen Energy, 38, 12, 2013, 4901-4934, <http://dx.doi.org/10.1016/j.ijhydene.2013.01.151>
- Mancera, Calderón *et al.*, An Optimized Balance of Plant for a Medium-Size PEM Electrolyzer: Design, Control and Physical Implementation, Electronics 9, 5, (2020), 871, <https://doi.org/10.3390/electronics9050871>
- Kumar, Himabindu, 2019, Hydrogen production by PEM water electrolysis A review, Materials Science for Energy Technologies, 2, 3, (2019), 442-454, <https://doi.org/10.1016/j.mset.2019.03.002>

