

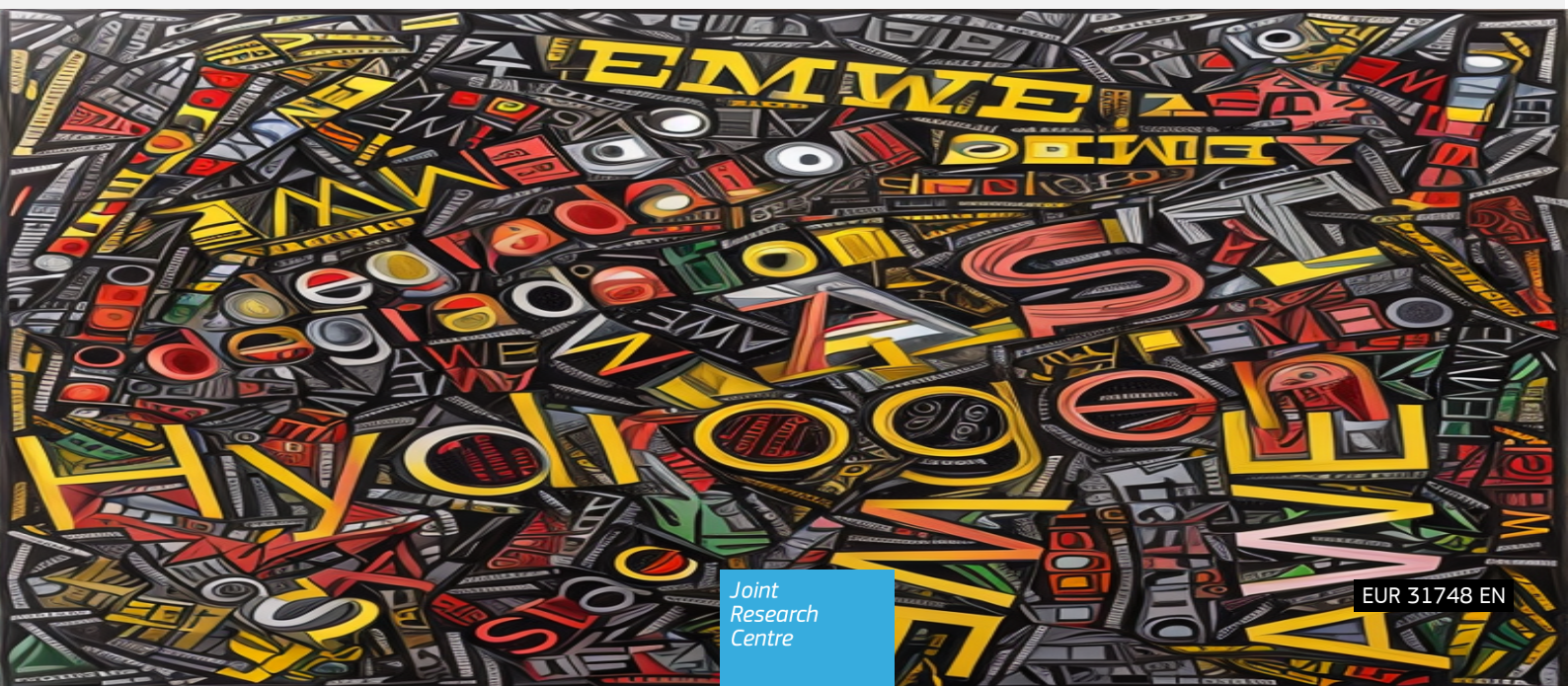


EU harmonised accelerated stress testing protocols for low-temperature water electrolyser

A proposal with testing guidance for assessing performance degradation in water electrolyser stacks

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Abstract

This document introduces proposed accelerated stress testing (AST) protocols for assessing the performance degradation of water electrolyser (WE) stacks. These stacks play a crucial role in generating clean hydrogen in bulk amounts through the electrolysis of water, primarily using electricity from renewable energy sources such as photovoltaic arrays and wind turbines.

By implementing these protocols, it becomes feasible to assess the performance degradation of various stacks systematically especially following a design of experiment approach. This allows for a thorough comparison of the three main low-temperature water electrolysis technologies: alkaline water electrolysis in an alkaline water electrolyser, anion exchange polymer membrane water electrolysis in an anion exchange polymer membrane water electrolyser, and proton exchange membrane or polymer electrolyte membrane water electrolysis in a proton exchange membrane or polymer electrolyte membrane water electrolyser.

It is important to note that this document does not delve into specific techniques for accelerating particular failure modes or enhancing different degradation phenomena at the component and sub-component levels within WE stacks. Instead, it offers broad guidelines for establishing AST procedures for stacks to ensure their reliable operation in water electrolyser systems utilising fluctuating renewable electricity.

These protocols are intended for use by both the research community and industry, serving purposes such as research and development (R&D), and stack prototype qualification, assessing R&D progress, setting priorities with cost targets, development milestones, and technological benchmarks, and making informed decisions regarding technology selection.

Foreword

This report was prepared under the framework contract between the Directorate-General JRC of the European Commission (EC) and the Clean Hydrogen Joint Undertaking (Clean H₂ JU), the successor to the Fuel Cells and Hydrogen second Joint Undertaking (FCH2JU)⁽¹⁾. The JRC contractual activities are stated in the strategic research and innovation agenda 2021-2027 of the Clean Hydrogen Partnership for Europe (SRIA)⁽²⁾. This report constitutes part two of the deliverable B.1 entitled "Report summarising the workshop findings on electrolyser lifetime degradation phenomena SoA and a preliminary proposal for setting up harmonised protocols for accelerated stress testing of low temperature electrolyzers" of the Rolling Plan 2023 contained in the Clean H₂ JU work programme 2023⁽³⁾. It is the result of a collaborative effort between partners from research and technology organisations (RTOs) in industry and academia participating to European Union (EU) funded R&D projects⁽⁴⁾ especially in power-to-hydrogen (P-to-H₂) applications involving low-temperature water electrolyser (LTWE) for demonstration and eventually, first industrial deployment (FID).



⁽¹⁾ According to Article 3(1)(c) of Council Regulation (EU) No 2021/2085 of 19/11/2021 (EU OJ L 427, 30.11.2021, p. 17), the Clean H₂ JU succeeds the FCH2JU as of 30 November 2021.

⁽²⁾ See p. 103 at https://www.clean-hydrogen.europa.eu/about-us/key-documents/strategic-research-and-innovation-agenda_en

⁽³⁾ See p. 163 at https://www.clean-hydrogen.europa.eu/about-us/key-documents/annual-work-programmes_en

⁽⁴⁾ For a list of projects, see online at https://www.clean-hydrogen.europa.eu/projects-repository_en. More comprehensive information is searchable at the Community Research and Development Information Service (CORDIS) under <https://cordis.europa.eu>.

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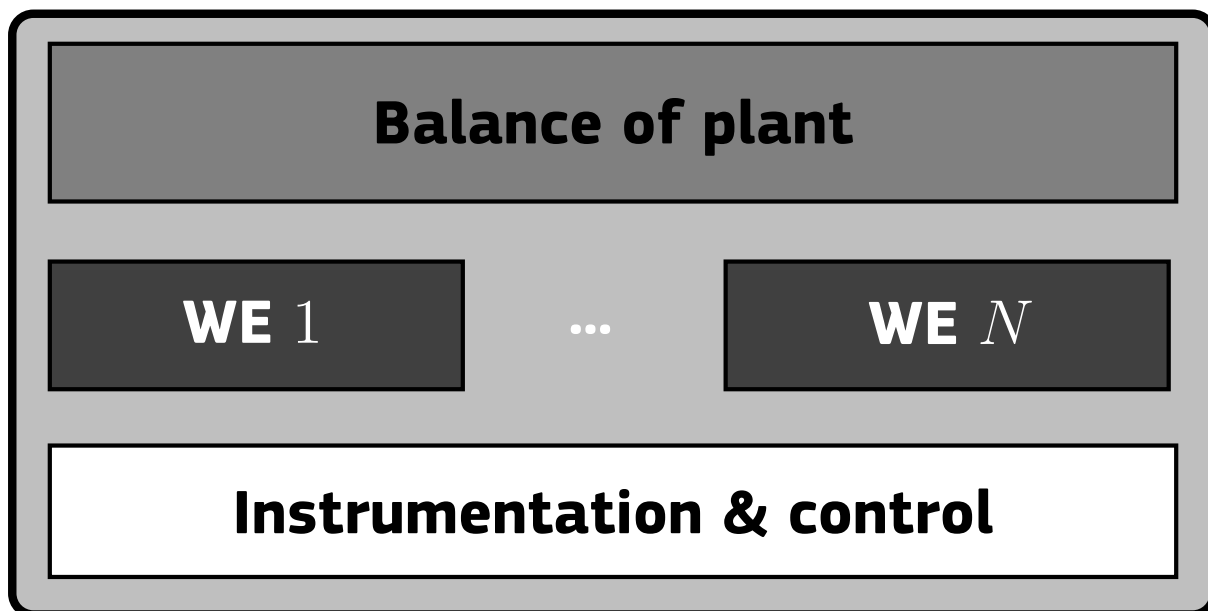
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1 Introduction

Water electrolyser (WE) stacks (**4.2.80**) used in water electrolyser systems (WE systems) (**4.2.81**) are at the core of generating clean hydrogen (H_2) in bulk amounts in addition to oxygen (O_2) and heat by the electrolysis (**4.2.36**) of water (H_2O) using fluctuating electricity particularly from sources of variable renewable energy (VRE). Most industrial WE systems (Figure 1) employing commercial WE stacks use low-temperature water electrolysis (LTWEL) technologies, namely alkaline water electrolysis (AEL) (**4.2.8**) in an alkaline water electrolyser (AWE) (**4.2.7**) and proton exchange membrane or polymer electrolyte membrane water electrolysis (PEMEL) (**4.2.64**) in a proton exchange membrane or polymer electrolyte membrane water electrolyser (PEMWE) (**4.2.63**) (Chatenet *et al.*, 2022, Shih *et al.*, 2022).

Figure 1: Schematic of a WE system comprising one or more WE stacks (WE 1 to WE N), common balance of plant (BoP) and instrumentation & control devices including safety sensors and software.



Source: JRC, 2023.

AWE have the advantage of being least reliant on the use of critical raw materials (CRM) (**4.2.22**). CRM are a serious concern for the European Union (EU) with regard to up-scaling and large-scale deployment of low-temperature water electrolyser (LTWE) technologies (Carrara *et al.*, 2023). PEMWE use CRM in catalysts (**4.2.16**), namely platinum (Pt) at the cathode (negative electrode) and platinum-group metals (PGM) (**4.2.61**) oxides such as iridium oxide (IrO_x) or iridium-ruthenium oxide ($IrO_x\text{-}RuO_y$) at the anode (positive electrode) to facilitate the water electrolysis (WEL) reactions in the electrodes of the proton exchange membrane or polymer electrolyte membrane water electrolysis cells (PEMECs), see equation (3.1.4).

PEMWE have the benefits of higher current density, small footprint and space requirements, non-toxic liquids and high hydrogen output pressure. They operate more flexible than other electrolyser making them suitable for delivering grid balancing services (Allidières *et al.*, 2019). Note, electricity grids (**4.2.30**) will exceedingly rely on balancing services in the future compared to the present situation with increasing use of diverse sources of VRE (solar, tidal, wave, wind, etc.) in the grid.

Today, WE stacks that use anion exchange polymer membrane water electrolysis (AEMEL) (**4.2.11**) in an anion exchange polymer membrane water electrolyser (AEMWE) (**4.2.10**) are less common. Potentially, AEMWE combine advantages of AEL and PEMEL (see Table 1) in a single device (Du *et al.*, 2022, Santoro *et al.*, 2022) when using de-mineralised liquid water (**4.2.25**) as in PEMWE whereas alkaline anion exchange polymer membrane water electrolyser (AAEMWE) uses dilute alkaline solution similar to AWE.

Commonly, manufacturer of WE systems specify the system boundaries while considering the BoP components (**4.2.13**) which form part of the system ⁽¹⁴⁾. Besides common hardware (piping, valves, actuators, sensors,

⁽¹⁴⁾ The immediate use of the hydrogen generated may require compression equipment (Sdanghi *et al.*, 2020, Durmuş *et al.*, 2021, Tahan, 2022, Marcuś *et al.*, 2022) as part of the BoP, especially in power-to-gas (P-to-G) applications and in industrial processes requiring high-pressure hydrogen. In applications of energy storage (ES) (**4.2.40**), including hydrogen-to-power (H_2 -to-P) with hydrogen stored as compressed gaseous hydrogen (CGH_2) (**4.2.19**) in vessels or large (seasonal) underground storage facilities, compression equipment may be part of the BoP of a particular WE system (Ausfelder *et al.*, 2017). In power-to-mobility (P-to-M) applications with hydrogen stored either as CGH_2 or as liquid hydrogen (LH_2) (**4.2.47**) in vessels, liquefaction equipment may be part of the BoP of a particular WE system in the latter case.

wiring/cabling, etc.), BoP usually consists of

- **electric power supply (4.2.31)** such as AC-to-DC (AC/DC) converter when grid-connected, or DC-to-DC (DC/DC) converter when directly coupled (off-grid) to one or another renewable energy source (RES) (4.2.67), for example, photovoltaic (PV) array (4.2.59) and/or wind turbine (4.2.83),
- **conditioning unit** including pumps, ion exchanger and heat exchanger for feeding de-mineralised water to PEMWE and AEMWE stack(s), alkaline solution (lye) to AWE, and diluted lye to AAEMWE stack(s) at the required inlet temperatures and
- **hydrogen purifier (4.2.43)** including liquid/gas separators, cooler(s), dryer(s), and de-oxidiser.

Where systems jointly use points of connection (PoCs) for electricity and/or fluid supply and for conveying exiting hydrogen and oxygen as part of a plant, the system boundary as the delineation between system interior and system exterior is to be defined by the manufacturer with the user's agreement.

Before their wider deployment in significantly large numbers and at scales ranging from a few hundred megawatts to several tens of gigawatt in capacity for use in ES and industrial applications across the EU and worldwide, stacks used in WE systems have to overcome a number of serious challenges (see section 3.4) by research and development (R&D) efforts and exceedingly, in real-world demonstrations, including first industrial deployment (FID) accompanied with capital investment in the said LTWE technologies (¹⁵).

⁽¹⁵⁾ For EU policy measures taken, see, for example, at https://energy.ec.europa.eu/topics/energy-systems-integration/hydrogen_en.

2 Objective and scope of this document

The objective of this document is to propose accelerated stress testing (AST) (4.2.2) protocols (4.2.3) for establishing the performance degradation (4.2.57) of WE stacks used for generating bulk amounts of hydrogen by LTWEL (AEL, AEMEL and PEMEL) at temperatures usually below 100 °C (373,15 K). Note, seawater electrolysis (Khan *et al.*, 2021), wastewater electrolysis (Cartaxo *et al.*, 2022), and bipolar polymer membrane water electrolysis (BPMEL) (Mayerhöfer *et al.*, 2020) are not considered⁽¹⁶⁾. This also applies to hybrid redox flow batteries (HRFBs), where, in addition to their use as ordinary redox flow batteries (RFBs), electrolysis to generate hydrogen is intended (Schmucker *et al.*, 2021). In addition, this document is primarily neither intended for AEL in an unipolar or monopolar AWE (LeRoy and Stuart, 1981), nor for membrane-less WE (Manzotti *et al.*, 2023).

WE stacks which can be rectangular, square, or circular in geometry, use electricity preferably from least dispatchable sources of VRE. A WE stack used in a WE system can be deployed in various applications where hydrogen is used as an energy carrier (4.2.38) (fuel or commodity) among others in ES such as P-to-G, P-to-M (road, rail, maritime) and power-to-X (P-to-X) including power-to-chemical (P-to-C), power-to-liquid (P-to-L), and power-to-fuel (P-to-F), as well as for direct use as feedstock or reducing agent in hydrogen-to-industry (H₂-to-I) processes. By applying the AST protocols (section 6) along with a test plan (4.2.78) to execute a test programme in a test campaign, the performance degradation of WE stacks are established under given test conditions (section 6.3), for example,

- to evaluate R&D progress made,
- to set research and innovation (R&I) priorities for development milestones and technological benchmarks to improve technology and assess impact on cost and
- to make well-informed business decisions regarding the selection of a particular WE stack technology.

The test methods suggested are mainly those contained in standards of the International Organization for Standardization (ISO) and the International Electrotechnical Commission (IEC)⁽¹⁷⁾. **Readers are advised to sufficiently familiarise with the referred standards, testing procedures and test methods described or cited therein.**

Furthermore, we also consider testing procedures previously developed as part of the EU water electrolysis harmonisation activities (Malkow *et al.*, 2018b, Malkow *et al.*, 2018a, Malkow and Pilenga, 2023a). Note, it is not intended to exclude any other suitable testing procedure or test method. The operation profiles (4.2.52) presented (section 6.8.2) serve as examples to establish the durability (4.2.28) of WE stacks by performing accelerated lifetime testing (ALT) (4.2.1) under reference test and operating conditions (section 6.4) as well as AST under stressing operating conditions (4.2.72) (section 6.5) especially during R&D and stack prototype qualification. They can be complemented by duty cycles, for example, to reflect realistic RES power generation profiles (section 6.8.2) for on-demand stack operation including the performance of services especially to balance variable loads of renewable energy (4.2.66) on the electricity grid known as balancing services⁽¹⁸⁾.

The use of RES-derived power profiles for stack testing distinguish these testing protocols from test protocols (4.2.74) for particular components of WE stacks developed in EU-funded research projects such as Enhanced performance and cost-effective materials for long-term operation of PEM water electrolyzers coupled to renewable power sources (ELECTROHYPEM), Hydrogen meeting FUTURE needs of low carbon manufacturing value chains (H2FUTURE), High Performance PEM Electrolyzer for Cost-effective Grid Balancing Applications (HPEM2GAS), Next Generation PEM Electrolyser under New Extremes (NEPTUNE), Novel modular stack design for high pressure PEM water electrolyzer technology with wide operation range and reduced cost (PRETZEL), Cost-effective PROton Exchange MEmbrane WaTer Electrolyser for Efficient and Sustainable Power-to-H₂ Technology (PROMETH2), Next Generation Alkaline Membrane Water Electrolyzers with Improved Components and Materials (NEWELY) and Anion Exchange Membrane Electrolysis for Renewable Hydrogen Production on a Wide-Scale (ANIONE) (Aricò *et al.*, 2013, Aricò *et al.*, 2016, Aricò *et al.*, 2018, Strataki, 2018, Stiber *et al.*, 2020, Fouda-Onana, 2020, Aricò *et al.*, 2020).

These protocols constitute testing guidance including mandatory requirements and agreed reference operating conditions for WE stacks to establish their performance degradation in a given power-to-hydrogen (P-to-H₂) application. They allow for sufficient flexibility when the test plan (section 6.6) of a scheduled test campaign is drawn up for a specific test programme addressing the use of the test item (4.2.76) in the target application. Thus, the test plan is to provide further details on

⁽¹⁶⁾ A bipolar polymer membrane water electrolyser (BPMWE) composed of bipolar polymer membrane water electrolysis cells (BPM-WECs) performs BPMEL without gas evolution at the AEM-PEM bipolar junction.

⁽¹⁷⁾ Standards, Technical Specifications (TSs) and Technical Reports (TRs) are not open access, but they can be purchased from ISO and IEC directly or from their constituting national committees (NCs).

⁽¹⁸⁾ Currently, Working Group (WG) 32 of ISO Technical Committee (TC) 197 is preparing the Approved Working Item (AWI) entitled "ISO 22734-2 Hydrogen generators using water electrolysis - Industrial, commercial, and residential applications - Part 2: Testing guidance for performing electricity grid service".

- test execution including
 - setting of test input parameters (TIPs) (**4.2.75**) with permissible variations,
 - test criteria for acceptance, failure and emergency stop, and
 - operation profiles (section 6.8)
- based on the stated purpose(s) and objective(s) of the tests and
- where necessary, provide more specific details on
 - test set-up (*e.g.*, sensor positions, stack compression, etc.) including specification and requirements of test equipments,
 - testing procedures including start-up and shut-down including emergency stop,
 - instrumentation, test and measurement methods (section 6.2), and
 - data acquisition (DAQ) (**4.2.24**) and post-processing of test results including an agreed set of test output parameters (TOPs) (**4.2.77**).

These protocols do not provide detailed examples on how to accelerate certain failure modes or enhance different degradation phenomena at component and sub-component level in WE stacks. Rather, they provide general guidelines for setting up accelerated stress testing of stacks for their reliable use in WE systems utilising fluctuating electricity from RES such as PV and wind turbine (WT) electric power. Note, the various key performance indicators (KPIs) suggested in section 6.10 are for assessing performance degradation of WE stacks exclusive of BoP components.

Importantly, the application of these AST protocols to WE stacks does not require the specification of the type and characteristics of the tested stack. Also users may selectively execute tests that are suitable for the objective and purpose of their test campaign from among those described herein.

3 Overview of low-temperature water electrolysis technologies

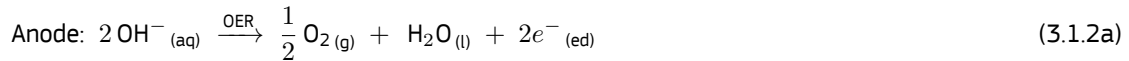
3.1 WEL electrode reactions

The generation of one mole of gaseous hydrogen, $\text{H}_2\text{(g)}$ (subscript (g) denotes gaseous phase), along with half a mole of gaseous oxygen, $\text{O}_2\text{(g)}$, by the electrolysis of one mole of liquid water, $\text{H}_2\text{O}(\text{l})$ (subscript (l) denotes liquid phase), as shown in the overall reaction

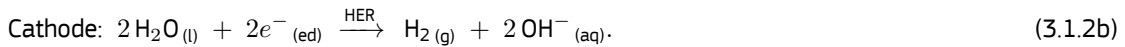


is performed in a water electrolyser. The three LTWEL technologies considered are

- **Alkaline water electrolysis:** Gaseous oxygen is formed by oxidising hydroxide ions (OH^-) in the aqueous phase (denoted by subscript (aq)) of the alkaline solution, typically 20–40 wt-% KOH (potassium hydroxide or lye), as electrolyte (4.2.37) at the anode or oxygen electrode in the oxygen evolution reaction (OER):

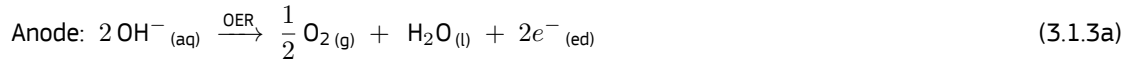


under an applied positive difference in potential (voltage) in excess of the open circuit potential (OCP) (U_{OCP}) sometimes called open circuit voltage (OCV) (U_{OCV})⁽¹⁹⁾ resulting from the supplied direct current (DC) (I_{dc}). Simultaneously, at the cathode or hydrogen electrode, gaseous hydrogen is formed by reducing liquid water in the hydrogen evolution reaction (HER):

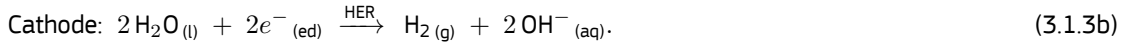


The electrons (e^-) are conducted via the electrodes (subscript (ed) denotes electrode) connected to an external circuit (DC power supply) entailing an ohmic resistance. The hydroxide ions diffuse along the potential-induced concentration gradient within the electrolyte of the alkaline water electrolysis cell (AEC) in the AWE stack from cathode to anode via a diaphragm.

- **Anion exchange polymer membrane water electrolysis:** Gaseous oxygen is formed by oxidising hydroxide ions at the anode in the OER:

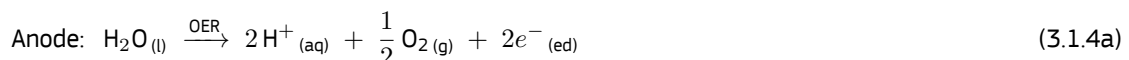


under an applied potential in excess of OCV. Gaseous hydrogen is formed simultaneously by reducing liquid water at the cathode in the HER:



Whereas electrons are conducted via the electrodes connected to an external circuit, hydrated hydroxide ions are conducted by the vehicular mechanism (standard diffusion) and the Grotthuss (proton hopping) mechanism (Dong *et al.*, 2018, Chen *et al.*, 2016) through the anion exchange polymer membrane (AEM) electrolyte of the anion exchange polymer membrane water electrolysis cell (AEMEC) in the AEMWE stack. In the case of an alkaline anion exchange polymer membrane electrolysis cell (AAEMEC), water is substituted by a dilute lye solution. Note that the identical electrode reactions (3.1.2) and (3.1.3) proceed in different media, namely alkaline solution in AWE according to reactions (3.1.2) and liquid water in AEMWE according to reactions (3.1.3).

- **Proton exchange membrane or polymer electrolyte membrane water electrolysis:** Gaseous oxygen is formed by oxidising water at the anode in the OER:



under an applied potential in excess of OCV. Gaseous hydrogen is formed simultaneously by reducing protons (H^+) at the cathode in the HER:



Whereas electrons are conducted via the electrodes connected to an external circuit, hydrated protons (H_3O^+) are conducted by the vehicular and Grotthuss mechanisms through the proton exchange membrane or polymer electrolyte membrane (PEM) electrolyte of the PEMEC in the PEMWE stack. That is, water is also yielded on the cathode due to electro-osmosis (4.2.32).

⁽¹⁹⁾ Note, OCV refers to the voltage across a complete electrochemical cell (4.2.33), while OCP refers to the potential measured between a half-cell and a suitable reference electrode (del Olmo *et al.*, 2021).

3.2 Materials, operating conditions and technology readiness levels

AWEs with technology readiness level (TRL) 9 at the MW scale are mature as they benefit from many decades of operational experience in the chlor-alkali electrolysis process, simplicity when compared to ion exchange membrane (IEM) (4.2.44) based WE, low cost, and suitability for larger scale projects. PEMWEs with PEMECs as constituting units having TRL 8 to 9 at the kW to MW scale are most mature among the IEM based WEs. The least mature, with TRL 6 at the kW scale, are AEMWEs using AEMECs as constituting units employing dilute alkaline solution. In the future, AEMWEs may be fed by pure water.

Most common in AWEs are Zirfon[®], a porous composite made of zirconia-polysulfone coated open mesh polyphenylene sulfide (PPS) polymer fabric, as porous separator membrane, nickel (Ni) or Ni/Ni Fe alloys (on steel core) as anode, and nickel or Ni alloy coated stainless steel as cathode. Current collectors (4.2.23) are made of nickel plates or Ni-coated steel. Typically, AWEs operate at temperatures between 60 °C to 90 °C (333,15 K to 363,15 K), current densities of between 0,2 A/cm² and 0,9 A/cm², and atmospheric pressure or pressures up to 30 bar (3 MPa) (Ehlers *et al.*, 2023; Brauns and Turek, 2020).

AEMWEs often use permeable fluorine-free hydrocarbon polymers as electrolyte membranes, non-PGM especially Co, Ni or Fe, their alloys and (mixed) oxides as anodes, and Ni and its alloys as cathodes, besides Pt. Typically, AEMWEs operate at temperatures between 40 °C to 80 °C (313,15 K to 353,15 K), current densities of between 0,5 A/cm² and 2 A/cm², and atmospheric pressure or at pressure from 8 bar to 35 bar (800 kPa to 3,5 MPa). The gas diffusion layer (GDL) (4.2.41) are made of carbon paper or cloth, titanium sheets, stainless steel felts, or Ni foam (Miller *et al.*, 2020; Du *et al.*, 2022). They provide for electronic conductivity between the catalyst layer (CL) (4.2.17) and the bipolar plates (biPs) (4.2.14) and remove gaseous products (hydrogen and oxygen).

Most often, PEMWEs use perfluoro sulfonic acid (PFSA) (4.2.55) as electrolyte membrane, PGM oxides such as IrO_x and IrO_x-RuO_y as anode catalysts, and PGM such as Pt as cathode catalyst. Typically, PEMWEs operate at temperatures between 50 °C to 90 °C (323,15 K to 363,15 K), current densities of 1 A/cm² to 4 A/cm², and atmospheric pressure, equal pressures (anode and cathode), or at differential pressures of up to 50 bar (5 MPa). Their biPs are made of titanium (Ti) or graphite (Kumar and Lim, 2023; Carmo *et al.*, 2013).

3.3 Stack operation modes

Under galvanostatic conditions, direct current provided to a WE stack results in a DC voltage (U_{dc}) across each cell. Adding the voltage of all series-connected water electrolysis cells (WECs) results in the stack voltage. Under potentiostatic conditions, a DC voltage applied to a WE stack results in a current flowing through the stack perpendicular to the active electrode area (A_{act}) (4.2.6) of all in-series WECs of the stack. DC electricity is in the form of electric energy (E_{el}):

$$E_{el} (\text{kWh}) = P_{el} (\text{kW}) \cdot t (\text{h}) \text{ where} \quad (3.3.1a)$$

P_{el} is electric power and t is the duration of applied electric power. Specifically, the electric power of a stack is DC power calculated as follows

$$P_{el,dc} (\text{kW}) = U_{dc} (\text{kV}) \cdot I_{dc} (\text{A}). \quad (3.3.1b)$$

The electric power density of a stack ($P_{el,d}$) is calculated as follows

$$P_{el,d} (\text{kW/cm}^2) = U_{dc} (\text{kV}) \cdot J (\text{A/cm}^2) \text{ where} \quad (3.3.1c)$$

$$J (\text{A/cm}^2) = \frac{I_{dc} (\text{A})}{N_{cell} \cdot A_{act} (\text{cm}^2)} \quad (3.3.1d)$$

is the current density of the stack and N_{cell} is the number of cells in the stack electrically connected in series. Depending on temperature (T), the three operation modes of a WE stack are

- *Endothermic operation:* The water temperature decreases from input to output of the stack with its voltage below the thermal-neutral voltage⁽²⁰⁾ but above the reversible potential⁽²¹⁾. Among the three modes of stack operation, this mode corresponds to the highest energy efficiency (η_e) (4.2.39) of the stack (section 6.7.13). However, it comes at the expense of a low hydrogen output rate (section 6.7.8). The heat required for the WEL reactions (3.1.2), (3.1.3), and (3.1.4) to proceed as desired stems, under presumed adiabatic conditions, from the supplied water rather than from Joule (ohmic) heating (4.2.45) due to an insufficient supply of electricity.

⁽²⁰⁾ At standard ambient pressure and standard ambient temperature (4.2.70) of liquid water (pH = 0), the thermal-neutral voltage (U_{tn}) is 1,481 V, while this voltage is 1,473 V at 80 °C (353,15 K). However, the thermal-neutral voltage decreases with increasing temperature and higher pH values.

⁽²¹⁾ At standard ambient pressure and standard ambient temperature of liquid water (pH = 0), the reversible potential (U_{rev}) is 1,229 V vs SHE, while this potential is 1,184 V vs SHE at 80 °C (353,15 K). However, the reversible potential decreases with increasing temperature and higher pH values, while it slightly increases with increasing pressure.

- **Isothermal (thermal-neutral) operation:** The water temperature is virtually the same at both the input and output of the WE stack. The stack voltage is basically the thermal-neutral voltage. The additional heat required to sustain the equilibrium of the WEL reactions (3.1.2), (3.1.3), and (3.1.4) usually stems from Joule heating due to the externally supplied electricity required to establish the reversible potential.
- **Exothermic operation:** The water temperature increases from the input to output of the WE stack with its voltage above the thermal-neutral voltage. As a result, heat is formed by Joule heating due to the supplied excess electricity. In this mode, the heat generated is more than that required to sustain the WEL reactions (3.1.2), (3.1.3), and (3.1.4). An advantage of this mode is that more supplied electricity means a higher hydrogen output. It comes at the expense of high overvoltages (overpotentials) (**4.2.53**) or voltage gains and an increase in performance degradation upon prolonged operation at high current densities ($> 1 \text{ A/cm}^2$). Hence, voltage limits not exceeding 3,0 V for AEC, 2,0 V for AEMEC and 2,5 V for PEMEC are common (Kumar and Lim, 2022) to prevent excessive stack degradation. At WE system level, any recovered heat boosts the overall energy efficiency of the system.

3.4 Advantages, disadvantages and challenges

Table 1 lists common advantages, disadvantages and main challenges of the three LTWE technologies at present.

Table 1: Common advantages, disadvantages and main challenges of three major LTWEL technologies (AWE, AEMWE and PEMWE)

	AWE	AEMWE	PEMWE
Advantages	relatively high energy efficiency ^(a) use of less expensive non-PGM catalysts materials ^(d)	use of less expensive non-PGM catalyst ^(b) low water impurity sensitivity relatively low CAPEX	high energy efficiency ^(c) fast response time including rapid start-up and shut-down ^(e) small footprint, compact and light-weight
	low current density, pressure and gas purity limited operational flexibility ^(g)	limited response time low ionic conductivity	use of expensive cell materials ^(f) sensitive to feed water quality with limited tolerance to (foreign ion) impurities ^(h) higher degradation rates and shorter lifetime safety risk due to hydrogen and oxygen crossover ⁽ⁱ⁾
Challenges	relatively high overpotentials for improving chemical and thermo-reduction and eventual replacement of PGM-free OER electro-catalysts in alkaline media	mechanical membrane stability and durability	reduction and eventual replacement of PGM oxide as catalysts ^(j)
	operating at higher temperatures (enabling higher efficiencies)		counteracting corrosion and low conductivity in passive layers on current collectors substitution of PFAS (4.2.54) by PEM with lower gas diffusivity ^(k)
	improvement of (micro-porous) membranes to reach higher ionic conductivities (enabling higher current densities) with better mechanical properties (enabling thinner membranes) and reduced gas crossover (enabling lower power operation without compromising safety) scalability of durable, low-power bipolar WE modules to larger scales at reduced manufacturing cost ^(l)		

Note: This table does not claim to present an exhaustive list of advantages and disadvantages.

(a) Typically, the energy efficiency of AWE stacks is 70-80 % (higher heating value (HHV)).

(b) For example, nickel or cobalt.

(c) Typically, the energy efficiency of PEMWE stacks is far greater than 70 % (HHV) considering their higher current densities.

(d) Often, Ni and Ni alloys are used.

(e) Relevant to ensure rapid responses to fluctuating power from variable RES.

(f) These include IEMs as electrolytes, Pt-based catalysts at the cathode and iridium oxides based catalysts at the anode, with high CAPEX.

(g) Note, unipolar WEs can operate anywhere from 0 to 100 % power and have a fast response time, including rapid start-up and shut-down.

- (^h) Typically, these impurities are dissolved minerals and other contaminants.
- (ⁱ) Crossover of hydrogen to the oxygen electrode and oxygen to the hydrogen electrode can lead to the formation of explosive atmospheres.
- (^j) Especially, IrO_x and IrO_x-RuO_y as OER catalysts in the anode and Pt as HER catalyst in the cathode.
- (^k) In PEMECs, the PEM electrolyte, ionomer, gaskets, and sealants frequently use PFAS, including PFSA polymers (fluoropolymers).
- (^l) This would require repeating BoP and power electronics as well as design changes to electrical substations.

Source: Joint Research Centre (JRC), 2023.

The most mature among these three technologies is the AWE technology. AWEs are initially prime candidates for reliable hydrogen generation in bulk amounts. In the midterm, they will likely be increasingly replaced or complemented by more versatile PEMWEs (²²). In the more distant future, less mature AEMWE technology could become the dominant LTWE technology. Often, AEMWEs are portrayed as beneficially uniting the advantages of AWEs and PEMWEs without necessarily sharing their drawbacks.

However, the two most demanding challenges encountered today by the three LTWE technologies are:

- Upon completion of the ongoing REACH restriction process (²³), the possibility of a future ban by the EU on the use of PFAS-containing materials (EC, 2020b) in products placed on the single market (²⁴) and
- Without realising high iridium (Ir) recycling rates, the scarcity of iridium on earth threatening the scale-up of PEMEL technology to the TW-scale globally (Clapp *et al.*, 2023, Kibsgaard and Chorkendorff, 2019, Kiemel *et al.*, 2021).

As a result, some WEC materials used in stacks will inevitably be different to those used presently once these challenges are met progressively in the future (²⁵). Ideally, this happens without changing too many KPI (**4.2.46**) targets. Also, the operating conditions of WE stacks and their mode(s) of operation may require adaptation since new WEC materials may have different properties and stack design could change. In addition, WEC materials and their morphologies integrated into WE stacks and interfaces within WE stacks may undergo iterative optimisation, including possible modifications in configuration and design suitable for high-throughput processing and mass manufacture of WE stacks.

Also, the generation and consumption patterns of electricity will change. Future electricity supply will be more variable as more and more RES installations of increasingly larger size are connected and their type and scale variety increases. Most probable, future smart grids (**4.2.69**) will exceedingly rely on autonomous distributed energy resources (DER) (**4.2.27**) managed by continuously improving artificial intelligence (AI)-based software. It includes automated on-demand deployment and disengaging of one or another WE system as part of a grid with fluctuating electricity.

Further, a WE system directly coupled to a large-scale RES installation as part of solar or wind energy farm will increasingly use AI-based supervisory control and data acquisition (SCADA) (**4.2.73**) software with predictive energy and weather forecasting and monitoring so that the supply of electricity to such a system or WE plant will not only be dependent on weather conditions and actual energy demand but also revenue considerations as regards the sale of renewable electricity versus clean hydrogen.

As a result, the mechanisms of material degradation in individual WECs (*e.g.*, iridium dissolution and redeposition, carbon catalyst support corrosion with agglomeration of platinum nano-particles, membrane thinning with fluoride release, and blocking of ion exchange sites by foreign cations in lye/liquid water feed) and thus of the performance degradation of WE stacks (*e.g.*, passivation by titanium oxide formation on Ti-based PTL and bIP) as currently known are likely to differ in their significance and extent. New degradation phenomena could emerge while degradation phenomena of little relevance today, could become more dominant.

Along with future material developments and stack optimisation, ongoing research into degradation phenomena of WEC materials and stack components and the mechanisms for their explanation will need to address the set-up of AST protocols for assessing the performance degradation of WE stacks in real-world applications dominated by fluctuating RES-derived electricity for the production of hydrogen. Developed AST protocols therefore need to consider the mentioned challenges and must provide for the necessary user flexibility.

⁽²²⁾ Note, unipolar AWEs can be as versatile as PEMWEs.

⁽²³⁾ See <https://echa.europa.eu/regulations/reach/restrictions/restriction-procedure>

⁽²⁴⁾ Due to risks (toxic and bioaccumulative effects) for human health, animals, and the environment, actions proposed by the European Commission aim at phasing out PFAS use in the EU unless proven essential for society (EC, 2020a). This is primarily owing to the strength of the carbon-fluorine (C-F) covalent bond in fluoropolymers, which makes them water-, oil- and grease-repellent and highly resistant to chemical and thermo-mechanical attacks. See also <https://www.eea.europa.eu/publications/emerging-chemical-risks-in-europe> and <https://echa.europa.eu/hot-topics/perfluoroalkyl-chemicals-pfas>. Note, PFAS-containing materials are used in several BoP components of WE systems, for example, in pumps, valves, cables, and electronics. In addition, fluoropolymers (PFAS membranes) are used in the electrolysis of potassium chloride (KCl) to create KOH, which is used in AEL technologies.

⁽²⁵⁾ The manufacture and use of alternative materials to PFAS in stacks and BoP components may pose other challenges.

4 Terminology

4.1 General

Terms and definitions used in this document are given below and in two JRC EUR reports (Tsotridis and Pilenga, 2018, Malkow *et al.*, 2021). In addition, ISO and IEC maintain terminological databases at the following websites:

- ISO Online browsing platform available at <https://www.iso.org/obp>.
- IEC Electropedia available at <http://www.electropedia.org>.

The verbal forms used have the following meaning:

- “shall” indicates a requirement,
- “should” indicates a recommendation,
- “may” indicates a permission and
- “can” indicates a possibility or a capability.

Reference to Système International d’Unités (SI) coherent (derived) units includes, as appropriate, metric prefixes. Following Clause 9.1 of ISO/IEC Directives, Part 2 (ISO and IEC, 2021), decimal fractions are denoted by comma. Alongside SI units, non-SI units may be used as customary. For example, we use degree Celsius ($^{\circ}\text{C}$) as unit of temperature (T) alongside Kelvin (K) and kilo Watt hours (kWh) as unit of energy (E) instead of kilo Joule (kJ).

4.2 Terms and definitions

4.2.1 accelerated lifetime testing (ALT)

destructive testing of a test item (4.2.76) by subjecting it to aggravated conditions (*e.g.*, current, pressure, temperature, voltage, etc.) in excess of nominal conditions of real-life use, in an attempt to reveal likely faults and modes of failure in a short amount of time by increasing their frequency of occurrence, magnitude, duration, or any combination thereof, and thereby to assess the reliability of the item, mainly for commercial purposes

Note 1 to entry: ALT may help to predict the remaining useful life and required maintenance intervals of the test item. It shall not alter the basic failure modes and mechanisms, or their relative prevalence.

4.2.2 accelerated stress testing (AST)

testing of a test item (4.2.76) by applying high levels of stress (*e.g.*, current density, pressure, temperature, voltage, etc.) in excess of those under normal conditions of use to shorten the test duration in an attempt to trigger the same performance degradation mechanism(s) as would presumably occur for a longer exposure of the test item when tested under normal conditions of use, mainly to advance the maturity of the test item

Note 1 to entry: AST is intentionally non-destructive and mainly for identifying potentially detrimental operating conditions and modes of operation as well as unsuitable designs and ineffective materials and components. It may also be performed to identify and to characterise performance degradation (4.2.57) and their mechanism(s) occurring in the test item. Design of experiment (DoE) (4.2.26), along with physics-based modelling and post-mortem characterisation of the test item, may help to gain insight into and understand the relationship between the applied stress and performance degradation and their mechanism(s).

4.2.3 accelerated stress testing protocol

protocol with guidance specific for accelerated stress testing (4.2.2)

Note 1 to entry: Typically, testing protocols include objective and scope of testing, definition of terms, description of test item(s) (4.2.76), reference test and operating conditions, list of test procedures and measurement methods to be applied and instructions on processing, presentation and documentation of test results. Specific test procedures and measurement methods may also be listed in individual test protocols (4.2.74).

4.2.4 acceleration factor

ratio between the life at the normal use stress level and the life at the accelerated stress level

[Source: ISO/TR 16194:2017, 3.2 (ISO, 2017b)]

4.2.5 activation polarisation

part of the electrode polarisation (**4.2.34**) arising from a charge-transfer step of the electrode reaction

[Source: IEV 482-03-05]

Note 1 to entry: The electrode reactions are given in equation (3.1.2) on page 12 for AWE (**4.2.7**), equation (3.1.3) on page 12 for AEMWE (**4.2.10**), and equation (3.1.4) on page 12 for PEMWE (**4.2.63**).

4.2.6 active electrode area (A_{act})

geometric area of the electrode perpendicular to the direction of the current flow

[Source: IEV 485-02-08]

Note 1 to entry: For calculating current density (see equation (3.3.1d) on page 13) and electric power density (see equation (3.3.1c) on page 13) of a water electrolyser stack (**4.2.80**), the active electrode area is to be multiplied by the number of cells in the stack electrically connected in series.

Note 2 to entry: Active electrode area is expressed in cm^2 .

4.2.7 alkaline water electrolyser (AWE)

water electrolyser using alkaline solution as electrolyte (**4.2.37**)

[Source: JRC EUR 30324 EN report, term 680 (Malkow *et al.*, 2021)]

Note 1 to entry: AWE configurations are unipolar (also known as monopolar) or bipolar.

4.2.8 alkaline water electrolysis (AEL)

electrolysis (**4.2.36**) that employs an alkaline solution as electrolyte (**4.2.37**)

[Source: JRC EUR 30324 EN report, term 678 (Malkow *et al.*, 2021)]

4.2.9 anion exchange polymer membrane (AEM)

polymer based membrane with an anion conductivity, which acts as an electrolyte (**4.2.37**) and a separator between anode and cathode

[Source: JRC EUR 30324 EN report, term 681 (Malkow *et al.*, 2021)]

4.2.10 anion exchange polymer membrane water electrolyser (AEMWE)

electrolyser (**4.2.35**) that employs a polymer with (hydroxide) ion exchange capability as the electrolyte (**4.2.37**)

[Source: JRC EUR 30324 EN report, term 684 (Malkow *et al.*, 2021)]

4.2.11 anion exchange polymer membrane water electrolysis (AEMEL)

electrolysis (**4.2.36**) that employs an anion exchange polymer membrane (**4.2.9**) as electrolyte (**4.2.37**)

[Source: JRC EUR 30324 EN report, term 682 (Malkow *et al.*, 2021)]

4.2.12 artificial intelligence (AI)

set of methods or automated entities that together build, optimize and apply a model so that the system can, for a given set of predefined tasks, compute predictions, recommendations, or decisions

[Source: ISO/TR 6026:2022, 3.3 (ISO, 2022b)]

4.2.13 balance of plant (BoP)

supporting and auxiliary components, associated subsystems and structures based on the source of electricity and site-specific requirements and integrated into a comprehensive water electrolyser system (**4.2.81**) necessary to generate hydrogen

4.2.14 bipolar plate (biP)

conductive plate separating individual cells in a water electrolyser stack (**4.2.80**), acting as current collector (**4.2.23**) and providing mechanical support for the electrodes

4.2.15 Bode plot

combined graphical representation of impedance modulus (absolute value) and phase angle (argument)

as functions of frequency on a logarithmic scale

Note 1 to entry: This plot is named after Hendrik Wade Bode (1905-1982).

4.2.16 catalyst

substance that accelerates an electrochemical reaction without being consumed itself

Note 1 to entry: The catalyst lowers the activation energy of the reaction, allowing for an increase in the reaction rate.

[Source: IEV 485-01-01]

4.2.17 catalyst layer (CL)

porous region adjacent to either side of the electrolyte (**4.2.37**), containing the electro-catalyst, typically with ionic and electronic conductivity

Note 1 to entry: The catalyst layer comprises the spatial region where the electrochemical reactions take place.

[Source: IEV 485-02-06]

4.2.18 cold start

start-up when the test item (**4.2.76**) is at ambient temperature with no power input

Note 1 to entry: The ambient temperature can vary by location, season and time of the day.

4.2.19 compressed gaseous hydrogen (CGH₂)

gaseous hydrogen, which has been compressed and stored for later use

4.2.20 compression factor (*f*_{compr})

positive multiplier of less than unity used to shorten the original duration of an operation profile (**4.2.52**)

Note 1 to entry: The compression factor is calculated as follows:

$$f_{\text{compr}} = \frac{t_{\text{compr}} (\text{h})}{t_{\text{origin}} (\text{h})}, \quad 0 \ll f_{\text{compr}} < 1, \quad (4.2.1)$$

where t_{compr} is the duration of the compressed operation profile and t_{origin} is the duration of the original operation profile.

4.2.21 concentration polarisation

part of the electrode polarisation (**4.2.34**) arising from concentration gradients of electrode reactants and products

[Source: IEV 482-03-08]

Note 1 to entry: Concentration polarisation is most relevant at high current densities. In water electrolysis cells, concentration polarisation can result in a non-proportional increase in voltage.

4.2.22 critical raw materials (CRM)

materials that, according to a defined classification methodology, are economically important and have a high-risk associated with their supply

[Source: ISO 14009:2020, 3.2.14 (ISO, 2020c)]

4.2.23 current collector

conductive material in a water electrolyser stack (**4.2.80**) that collects electrons from the anode side and conducts electrons to the cathode side

4.2.24 data acquisition (DAQ)

process of collecting and entering data

[Source: ISO 15143-1:2010, 3.1.4 (ISO, 2010a)]

4.2.25 de-mineralised water

water of the mineral matter or salts have been removed by de-ionisation

[Source: ISO 23321:2019, 3.1 (ISO, 2019b)]

Note 1 to entry: De-mineralised water shall also be free of pollutants such as organic matter and gases.

4.2.26 design of experiment (DoE)

systematic methodology for collecting information to guide improvement of any process

Note 1 to entry: Statistical models are developed to represent the process under analysis.

Note 2 to entry: Simulation tools and optimisation can be applied to test and confirm specific improvements.

[Source: ISO 13053-2:2011, 2.12 (ISO, 2011a)]

4.2.27 distributed energy resources (DER)

generators (with their auxiliaries, protection and connection equipment), including loads having a generating mode (such as electrical energy storage systems), connected to a low-voltage or a medium-voltage network

[Source: IEV 617-04-20]

4.2.28 durability

ability of a test item (4.2.76) to maintain its performance characteristics (4.2.56) as required, under specified conditions of use and maintenance

4.2.29 durability test

test intended to verify whether or to evaluate to what degree a test item (4.2.76) is able to maintain its performance characteristics (4.2.56) over a period of use

4.2.30 electricity grid

public electricity network

[Source: ISO 52000-1:2017, 3.4.8 (ISO, 2017a)]

4.2.31 electric power supply

provision of electric energy from a source

[Source: IEV 151-13-75]

4.2.32 electro-osmosis

flow of water induced by a direct electric current applied across a membrane separating two electrolytes

[Source: IEV 891-02-84]

Note 1 to entry: The membrane can also be a diaphragm.

4.2.33 electrochemical cell

composite system in which the supplied electric energy mainly produces chemical reactions or, conversely, in which the energy released by chemical reactions is mainly delivered by the system as electric energy

[Source: IEV 114-03-01]

Note 1 to entry: In the first case, an electrochemical cell is also known as an electrolytic cell.

4.2.34 electrode polarisation

accumulation or depletion of electric charges at an electrode, resulting in a difference between the electrode potential with current flow, and the potential without current flow or equilibrium electrode potential

[Source: IEV 114-02-15]

4.2.35 electrolyser

device that performs electrolysis (4.2.36)

[Source: ISO/TR 15916:2015, 3.33 (ISO, 2015)]

4.2.36 electrolysis

method of separating and neutralising ions by an electric current in an electrolytic cell

[Source: IEV 114-04-09]

4.2.37 electrolyte

liquid or solid substance containing mobile ions that render it ionically conductive

[Source: IEV 485-03-01]

Note 1 to entry: The electrolyte is the main distinctive feature of the different LTWEL technologies.

4.2.38 energy carrier

substance or phenomenon that can be used to produce mechanical work or heat or to operate chemical or physical processes

[Source: ISO 52000-1:2017, 3.4.9 (ISO, 2017a)]

4.2.39 energy efficiency (η_e)

ratio of useful energy output to the total energy input, including all parasitic and auxiliary energy needed to operate the system

Note 1 to entry: Energy efficiency is expressed in % on the basis either of lower heating value (LHV) or higher heating value (HHV), which should be stated.

4.2.40 energy storage (ES)

action or method used to accumulate, retain and release energy for later use in an energy using system

[Source: ISO/IEC 13273-1:2015, 3.1.5 (ISO and IEC, 2015a)]

4.2.41 gas diffusion layer (GDL)

porous substrate placed between the catalyst layer (4.2.17) and the bipolar plate (4.2.14) to serve as an electric contact and allow the access of reactants to the catalyst layer and the removal of reaction products

[Source: IEV 485-04-05]

Note 1 to entry: A GDL coated with catalyst material is called gas diffusion electrode (GDE).

4.2.42 hot start

start-up when the item test (4.2.76) is within its normal operating temperature range

4.2.43 hydrogen purifier

equipment to remove undesired constituents from the hydrogen

Note 1 to entry: Hydrogen purifiers can comprise purification vessels, dryers, filters and separators.

[Source: ISO 19880-1:2020, 3.41 (ISO, 2020d)]

4.2.44 ion exchange membrane (IEM)

polymer sheet that contain negatively or positively charged functional groups in its polymer matrix designed to conduct cations or anions while blocking opposite charged ions

[Source: ISO 20468-6:2021, 3.1.18 (ISO, 2021a)]

4.2.45 Joule heating (ohmic heating)

heating caused by an electric current through a resistive material

[Source: IEV 815-15-41]

Note 1 to entry: It is named after James Prescott Joule (1818-1889).

4.2.46 key performance indicator (KPI)

quantifiable level of achieving a critical objective

Note 1 to entry: The KPIs are derived directly from, or through an aggregation function of, physical measurements, data and/or other KPIs.

[Source: ISO 22400-1:2014, 2.1.5 (ISO, 2014)]

4.2.47 liquid hydrogen (LH₂)

hydrogen that has been liquefied, i.e. brought to a liquid state

[Source: ISO 13984:1999, 3.4 (ISO, 1999)]

4.2.48 machine learning (ML)

process using algorithms rather than procedural coding that enables learning from existing data in order to predict future outcomes

[Source: ISO/TR 22100-5:2021, 3.2 (ISO, 2021b)]

4.2.49 Nyquist plot

graphical representation of the real part of impedance versus the negative of the imaginary part of impedance in rectangular coordinates

Note 1 to entry: This plot is named after Harry Nyquist (1889-1976).

4.2.50 ohmic polarisation

polarisation caused by the resistance to the flow of ions in the electrolyte (**4.2.37**) and of electrons in the electrodes, bipolar plates (**4.2.14**), and current collectors (**4.2.23**)

Note 1 to entry: The term "ohmic" refers to the fact that the voltage drop follows Ohm's law proportional to the current with an ohmic resistance (called "internal resistance" of the cell) as the proportionality constant.

[Source: IEV 485-15-03]

Note 2 to entry: The ohmic resistance of the electrolyte encompasses the electronic resistance of the polymer membrane (PEM and AEM) and the diaphragm (AWE), depending on its type and thickness.

4.2.51 operating (working) point

point on a characteristic performance curve representing the values of variable quantities at which usual operation is expected and optimum efficiency is desired

Note 1 to entry: Characteristic performance curves of a water electrolyser stack (**4.2.80**) are direct current-DC voltage curves and energy efficiency/electrical efficiency-electric power curves.

4.2.52 operation profile

curve representing electric power, current, or voltage against time used to illustrate the variation in electric power, current or voltage during a given time interval

4.2.53 overvoltage (overpotential)

voltage difference between the measured electrode potential and the equilibrium potential

[Source: ISO 8044:2020, 7.1.30 (ISO, 2020a)]

Note 1 to entry: In WE stacks, overvoltage relates to a given current density under specified conditions.

4.2.54 perfluoroalkyl and polyfluoroalkyl substances (PFAS)

commonly used international abbreviation for organic compounds with replacement of most or all hydrogen atoms by fluorine in the aliphatic chain structure

Note 1 to entry: The term is used in the broader sense for per- and polyfluoroalkyl substances (PFAS), and per- and polyfluorinated compounds (PFC) as well.

[Source: ISO 21675:2019, 3.1 (ISO, 2019c)]

4.2.55 perfluoro sulfonic acid (PFSA)

chemical compounds of the formula $C_nF_{(2n+1)}SO_3H$ and thus belong to the families of perfluorinated and polyfluorinated alkyl compounds

Note 1 to entry: PFSA belongs to PFAS (**4.2.54**).

4.2.56 performance characteristics

characteristics defining the ability of a test item (**4.2.76**) to operate as intended under specified conditions of use and maintenance

4.2.57 performance degradation

process leading to a significant change in the performance of the test item (**4.2.76**), typically characterised by a change of properties, whether reversible or irreversible, or by decay affected by environmental and test conditions, proceeding over a period of time and comprising one or several steps that affect the test item (**4.2.76**) to operate as intended, under specified conditions of use

4.2.58 performance test

test intended to verify whether or to evaluate to what degree a test item (**4.2.76**) is able to accomplish its performance characteristics (**4.2.56**)

4.2.59 photovoltaic (PV) array

two or more photovoltaic modules at one location that together provide a photovoltaic solar energy system

[Source: ISO 6707-3:2022, 3.3.7 (ISO, 2022a)]

4.2.60 photovoltaic (PV) power

technology that turns sunlight directly into electricity

[Source: ISO/IEC TR 15067-3-8:2020, 3.19 (ISO, 2020e)]

4.2.61 platinum-group metals (PGM)

group of six noble, precious metallic elements (ruthenium, rhodium, palladium, osmium, iridium, and platinum) clustered together in the periodic table

Note 1 to entry: These transition metals are located in the d-block of the periodic table. They have similar physical and chemical properties.

4.2.62 proton exchange membrane or polymer electrolyte membrane (PEM)

polymer based membrane with cation (proton) conductivity, which acts as an electrolyte (**4.2.37**) and a separator between anode and cathode

Note 1 to entry: PEM is a cation exchange membrane exclusively in the acidic H^+ form.

[Source: JRC EUR 30324 EN report, term 695 (Malkow *et al.*, 2021)]

4.2.63 proton exchange membrane or polymer electrolyte membrane water electrolyser (PEMWE)

electrolyser that employs a polymer with (proton) ion exchange capability as the electrolyte (**4.2.37**)

[Source: IEC 62282-8-102:2019, 3.1.26 (IEC, 2019b)]

4.2.64 proton exchange membrane or polymer electrolyte membrane water electrolysis (PEMEL)

electrolysis (**4.2.36**) that employs a proton exchange membrane or polymer electrolyte membrane (**4.2.62**) as electrolyte (**4.2.37**)

[Source: JRC EUR 30324 EN report, term 696 (Malkow *et al.*, 2021)]

4.2.65 principle of superposition

principle that the time response to the sum of several input variables is the same as the sum of the time responses caused by the individual input variables

Note 1 to entry: The principle of superposition includes the special case, that at multiplication of an input variable by a constant factor the accompanying time response is multiplied by the same factor (often called “principle of amplification”).

[Source: IEV 351-45-01]

Note 2 to entry: In electrochemical impedance spectroscopy (EIS) measurements, the input variable is current (voltage) under galvanostatic (potentiostatic) conditions. Under these conditions, the time response is the resulting voltage (current).

Note 3 to entry: This principle is attributed to Ludwig Boltzmann (1844-1906) and John Hopkinson (1849-1898).

4.2.66 renewable energy

energy obtained from a renewable energy source (**4.2.67**)

[Source: ISO/IEC 13273-2:2015, 3.1.6 (ISO and IEC, 2015b)]

4.2.67 renewable energy source (RES)

energy source not depleted by extraction as it is naturally replenished at a rate faster than it is extracted

[Source: ISO/IEC 30134-3:2016, 3.1.4 (ISO, 2016a)]

4.2.68 reverse current

flow of induced current in a commissioned AWE (**4.2.7**) stack during OCV operation (zero supply current) given the electrical connections of a closed circuit made of the bipolar plates (**4.2.14**) with electronic conduction and ionic conduction of the electrolyte (lye) solution in the manifolds of the stack

Note 1 to entry: Reverse current is also driven by remaining product gas (O_2 and H_2) after electrolyser operation, which triggers a fuel cell (FC)- like operation mode generating such current.

4.2.69 smart grid

electric power system that utilises information exchange and control technologies, distributed computing and associated sensors and actuators for purposes such as:

- to integrate the behaviour and actions of the network users and other stakeholders,
- to efficiently deliver sustainable, economic and secure electricity supplies

[Source: IEV 617-04-13]

Note 1 to entry: Smart grids enable enhanced and automated monitoring and control of electricity generation, transmission and distribution for added availability, reliability, efficiency, and cost-effective operations.

4.2.70 standard ambient temperature and pressure (SATP) conditions

conditions of standard ambient pressure ($p^0 = 100$ kPa) and standard ambient temperature ($T^0 = 298,15$ K)

4.2.71 stray current

current which follows paths other than the intended paths

[Source: IEV 811-35-05]

Note 1 to entry: For operating AWE (**4.2.7**) (**4.2.7**) stacks, stray current is also known as shunt current. Shunt currents occur when the flow of current instead of passing through the ionically conducting diaphragm, it passes through the lye solution in the manifolds of the stack.

4.2.72 stressing operating conditions

operating conditions intentionally in excess of normal operating conditions exerted onto a test item (**4.2.76**), which are likely to cause performance degradation (**4.2.57**) of the item during its operation

4.2.73 supervisory control and data acquisition (SCADA)

process control system generally used to control dispersed assets using centralised data acquisition (**4.2.24**) and supervisory controls

[Source: ISO/IEC 27019:2017, 3.15 (ISO and IEC, 2017)]

Note 1 to entry: Systems operate using coded signals over communication channels with remote equipment to acquire information about the status for display or recording functions, and accessing process control set points as well as current and historical online data.

4.2.74 test protocol

list of the steps to be followed in the test

[Source: ISO/IEC/IEEE 26513:2017, 3.141 (ISO *et al.*, 2017)]

Note 1 to entry: Test protocols contain specific test procedures and measurement methods to address particular aspects of test item performance or durability, including at component and sub-component levels. Usually, test protocols follow an established scheme with comprehensive decision rules, including pass and fail criteria, while often employing purpose-oriented test cases or set(s) of test cases.

4.2.75 test input parameter (TIP)

parameter whose values can be set in order to define the test conditions of the test system, including the operating conditions of the test object

Note 1 to entry: TIPs have to be controllable and measurable. Values of TIPs are known before conducting the test. TIPs can be either static or variable. Static TIPs stay constant and variable TIPs are varied during the test.

[Source: IEC 62282-8-101, 3.1.33 (IEC, 2020c)]

4.2.76 test item

electrolyser (**4.2.35**) stack of type AWE (**4.2.7**), AEMWE (**4.2.10**) including AAEMWE, or PEMWE (**4.2.63**)

4.2.77 test output parameter (TOP)

parameter that indicates the response of the test system/test object as a result of variation of test input parameters (**4.2.75**)

[Source: IEC 62282-8-101, 3.1.34 (IEC, 2020c)]

Note 1 to entry: Values of TOPs are unknown before conducting the test and are measured during the test or calculated subsequently.

4.2.78 test plan

planning document detailing the principles, test methods, conditions, procedures and data quality required to carry out testing and to produce test data

[Source: ISO 14050:2020, 3.4.19 (ISO, 2020b)]

Note 1 to entry: The test plan outlines the objective(s), purpose(s), requirements, and strategy for testing, including the type of test(s), descriptions of test environments and conditions, including TIP (**4.2.75**) set-point values, responsibility for the testing, the equipment and instrumentation for use in the testing, as well as the process for performing (workflow of testing) and documenting the test(s) (recording, analysing, and reporting of test results), and for handling test failure(s).

4.2.79 uncertainty

parameter, associated with the result of a measurement, that characterises the dispersion of the values that could reasonably be attributed to the measurand

[Source: International Electrotechnical Vocabulary (IEV) 415-05-13]

4.2.80 water electrolyser (WE) stack

assembly of two or more electrochemical cells (**4.2.33**), separators, manifolds, and a supporting structure, using DC electricity to generate hydrogen and heat by the electrolysis (**4.2.36**) of liquid water

Note 1 to entry: Alkaline water electrolyser (**4.2.7**) stacks and alkaline anion exchange polymer membrane water electrolyser stacks employ an alkaline solution rather than liquid water.

4.2.81 water electrolyser (WE) system

assembly of interrelated components of a defined configuration with one or more water electrolyser stacks (**4.2.80**) at its core, which delivers hydrogen

Note 1 to entry: A water electrolyser system can include components such as power supply terminals, fluid connectors, compressors, storage vessels, piping, valves, pressure-relief devices, pumps, expansion joints, gauges, cabling, and control, monitoring, and safety subsystems, including communication interfaces.

Note 2 to entry: A water electrolyser system can refer to a site, a facility at a site, or an installation at a facility. A water electrolyser plant may contain multiple water electrolyser systems.

4.2.82 wind power

use of wind to provide mechanical power through wind turbines (**4.2.83**) to turn electric generators

[Source: ISO 6707-3:2022, 3.6.19 (ISO, 2022a)]

4.2.83 wind turbine (WT)

device that converts kinetic energy from the wind into electricity

[Source: ISO 6707-3:2022, 3.2.5 (ISO, 2022a)]

4.3 Abbreviations and acronyms used

A list of abbreviations and acronyms used in this report is appended, see page 72.

4.4 Symbols used

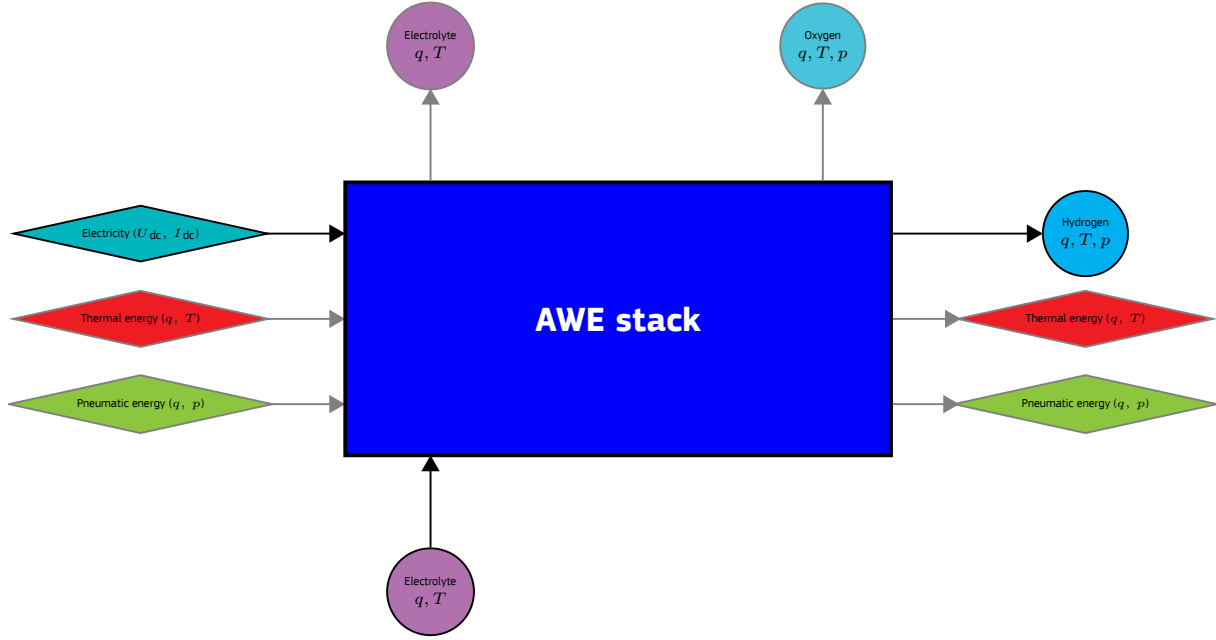
A list of symbols used in this report is appended, see page 79.

5 Description of test items

5.1 AWE stack

Figure 2 shows schematically the input and output streams of energy forms and substances of an AWE stack.

Figure 2: Schematic of the input and output streams (directional arrows) of energy forms (diamond shape) and substances (circular shape) of an AWE stack (rectangular shape); q and p represent flow rate and pressure, respectively. The thick line around the grey-shaded box denotes the stack boundary. The use of the grey colour indicates streams of secondary relevance in the context of accelerated stress testing.



Source: JRC, 2023.

At its PoCs (current/voltage terminals and fluid inlets), the input energy streams to an AWE stack are

- **Electricity** in the form of electric energy, see equation (3.3.1a), by supplying DC power, see equation (3.3.1b),
- **Thermal energy (E_{th})**, if any, in the form of heat/cold:

$$E_{th} \text{ (kWh)} = P_{th} \text{ (kW)} \cdot t \text{ (h)} \text{ where} \quad (5.1.1a)$$

P_{th} is thermal power given by equation (5.1.1b) and t is the duration of heat/cold supply.

$$P_{th} \text{ (kW)} = \sum_i q_m^i \text{ (kg/s)} \cdot c_p^i \text{ (kJ/(kg K))} \cdot (T^i \text{ (K)} - T^0 \text{ (K)}); \quad (5.1.1b)$$

q_m^i , c_p^i and T^i are mass flow rate, specific heat capacity at constant pressure and temperature of fluid i , respectively. On input, the heat transfer fluid i (input substance stream) is the aqueous electrolyte.

- **Pneumatic energy (E_{compr})** is only relevant for pressurised stacks:

$$E_{compr} \text{ (kWh)} = P_{compr} \text{ (kW)} \cdot t \text{ (h)} \text{ where} \quad (5.1.2a)$$

P_{compr} is the power of compression given by equation (5.1.2b) and t is the duration of pressurised stack operation.

$$P_{compr} \text{ (kW)} = \sum_j \left(\frac{\gamma^j}{\gamma^j - 1} \right) \frac{\bar{Z}^j \cdot R_g \text{ (kJ/(mol K))} \cdot T^0 \text{ (K)} \cdot q_n^j \text{ (mol/h)}}{3600 \text{ (s/h)}} \\ \left(\left(\frac{p^j \text{ (kPa)}}{p^0 \text{ (kPa)}} \right)^{\frac{\gamma^j - 1}{\gamma^j}} - 1 \right); \quad (5.1.2b)$$

R_g , \bar{Z}^j , q_n^j and p^j are universal gas constant, average compressibility factor, molar flow rate and pressure of fluid j , respectively. The isentropic expansion factor of fluid j is calculated as follows

$$\gamma^j = \frac{c_p^j (\text{kJ}/(\text{kg K}))}{c_v^j (\text{kJ}/(\text{kg K}))}; \quad (5.1.2c)$$

c_p^j and c_v^j are specific heat capacity at constant pressure and constant volume of fluid j , respectively. For a pressurised stack, the pneumatic fluids j (output substance streams) are hydrogen and oxygen.

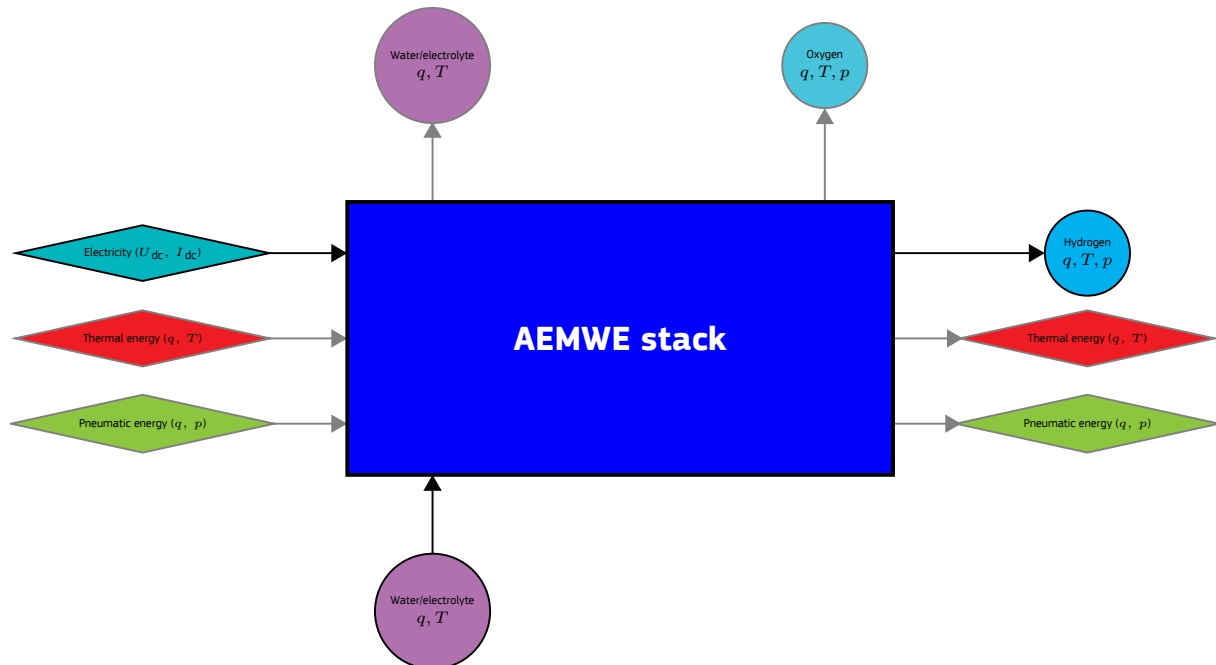
At fluid outlets, the output energy and substance streams from an AWE stack are

- Thermal energy carried by heat transfer fluids:
 - aqueous electrolyte,
 - hydrogen and
 - oxygen,
- Pneumatic energy carried by compressible fluids:
 - hydrogen and
 - oxygen,
- Hydrogen (mixed with electrolyte) at the cathode and
- Oxygen and water (mixed with electrolyte) at the anode.

5.2 AEMWE stack

Figure 3 shows schematically the input and output streams of energy forms and substances of an AEMWE stack.

Figure 3: Schematic of the input and output streams (directional arrows) of energy forms (diamond shape) and substances (circular shape) of an AEMWE stack (rectangular shape). The thick line around the grey-shaded box denotes the stack boundary. The use of the grey colour indicates streams of secondary relevance in the context of accelerated stress testing.



Source: JRC, 2023.

At its PoCs (current/voltage terminals and fluid inlets), the input energy streams to an AEMWE stack are

- **Electricity** in the form of electric energy, see equation (3.3.1a), by supplying DC power, see equation (3.3.1b),

- **Thermal energy**, see equation (5.1.1a), if any, in the form of heat/cold where on input the heat transfer fluid i (input substance stream) is liquid water for AEMECs and aqueous electrolyte for AAEMECs.
- **Pneumatic energy**, see equation (5.1.2a), which is only relevant for pressurised stacks where the pneumatic fluids j (output substance streams) are hydrogen and oxygen.

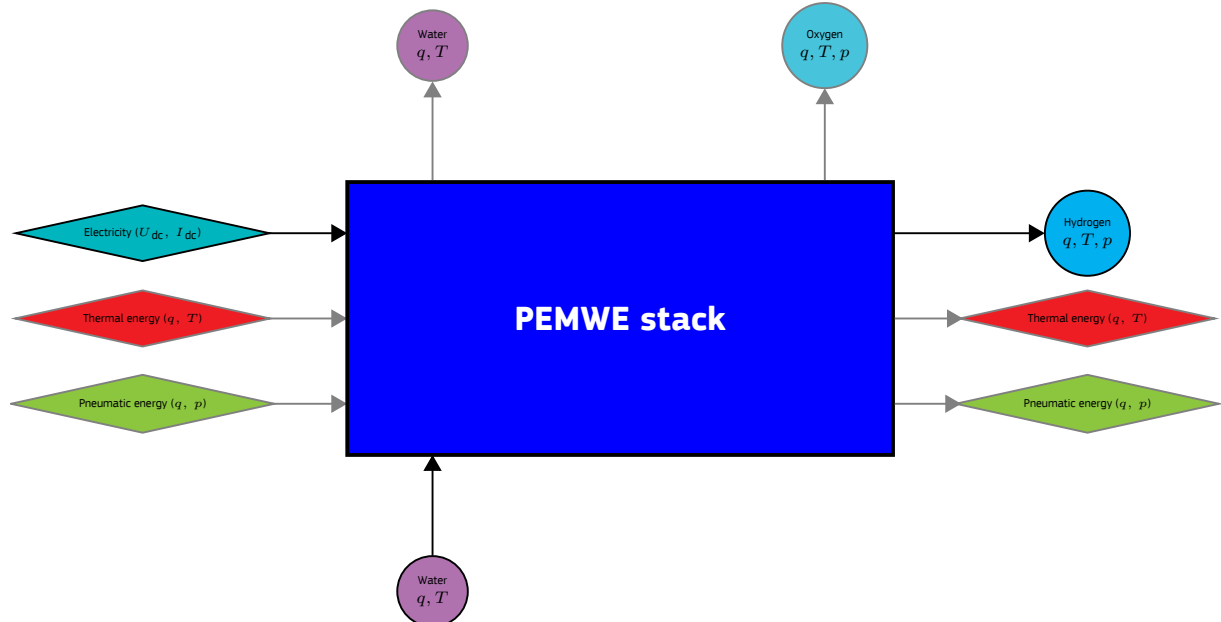
At fluid outlets, the output energy and substance streams from an AEMWE stack are

- Thermal energy carried by heat transfer fluids:
 - liquid water for AEMECs and aqueous electrolyte (lye) for AAEMECs,
 - hydrogen and
 - oxygen,
- Pneumatic energy carried by compressible fluids:
 - hydrogen and
 - oxygen,
- Hydrogen and water at the cathode and
- Oxygen and water (mixed with electrolyte) at the anode.

5.3 PEMWE stack

Figure 4 shows schematically the input and output streams of energy forms and substances of a PEMWE stack.

Figure 4: Schematic of the input and output streams (directional arrows) of energy forms (diamond shape) and substances (circular shape) of a PEMWE stack (rectangular shape). The thick line around the grey-shaded box denotes the stack boundary. The use of the grey colour indicates streams of secondary relevance in the context of accelerated stress testing.



Source: JRC, 2023.

At its PoCs (current/voltage terminals and fluid inlets), the input energy streams to a PEMWE stack are

- **Electricity** in the form of electric energy, see equation (3.3.1a), by supplying DC power, see equation (3.3.1b),
- **Thermal energy**, see equation (5.1.1a), if any, in the form of heat/cold where on input the heat transfer fluid i (input substance stream) is liquid water.
- **Pneumatic energy**, see equation (5.1.2a), which is only relevant for pressurised stacks where the pneumatic fluids j (output substance streams) are hydrogen and oxygen.

At fluid outlets, the output energy and substance streams from a PEMWE stack are

- Thermal energy carried by heat transfer fluids:
 - liquid water,
 - hydrogen and
 - oxygen,
- Pneumatic energy carried by compressible fluids:
 - hydrogen and
 - oxygen,
- Hydrogen and water (due to electro-osmosis, see section 3.1) at the cathode and
- Oxygen and water at the anode.

6 Proposal for AST protocols

6.1 General

A test campaign for assessing the performance degradation of WE stacks under specified test and operating conditions may have different objectives such as

- Determining qualitatively and/or quantitatively one or another identified degradation phenomenon including interactions (coupling) among different and possibly simultaneously occurring phenomena for a given set of WEC materials in a particular stack configuration of known design,
- Identifying or verifying hypothesised degradation mechanisms associated with one or more known phenomena occurring for a set of WEC materials in a given stack,
- Evaluating the performance and durability of improved or newly developed materials for WEC components deployed in a representative stack design and configuration,
- Checking on suitable stack designs and configurations, as well as
- Identifying potentially detrimental operating conditions and modes of operation as well as optimising operation modes of a stack in a given WE system deployed in a particular application.

As for any test campaign, an AST campaign on WE stacks follows a test program with a test plan (section 6.6) and schedule to conduct tests according to protocols describing precisely what type of tests, whether performance tests (**4.2.58**) or durability tests (**4.2.29**), should be carried out under which test and operating conditions (section 6.3) and how tests should be performed, in which order, when, and for how long or how often. It includes post-processing and presentation of the test results (section 7) as well as their reporting (Annex B).

What makes an AST campaign different from a test campaign testing the durability of WE stacks, is that the former test campaign aims to significantly shorten the time required for stack testing to save cost on R&D of WE stacks. Alternatively, it is to test a greater variety of WEC materials and stacks in the same amount of time using the available test hardware and equipment more effectively. It seeks an earlier demonstration including FID with the market entry of the developed WE stacks to eventually accelerate their commercialisation. For this reason, WE stacks are operated for a short time under operating conditions different from their normal operating conditions in an attempt to trigger similar performance degradation mechanism(s) as occur for longer stack exposures. It leads inevitably to more and higher transient operation of WE stacks primarily supplied by fluctuating RES electricity with a higher number of and more frequent changes in current or voltage (including start/stop (ON/OFF) operation) as would occur under real-life operating conditions.

In an AST campaign, available real-world operation profiles of RES-derived power (section 6.8.2) may be compressed in duration by a compression factor (f_{compr}) (**4.2.20**) and combined to obtain simulated duty cycles for dynamic operation of a WE stack undergoing accelerated stress testing. Additional stress is by setting other operating parameters especially stack operating temperature (T_{stack}) and differential (anode-to-cathode) pressure to their extreme values but not out of the range specified by the stack manufacturer to avoid dysfunction or destruction of the WE stack. Additional stress can also be applied to a WE stack when deliberately adding known contaminants in sufficient quantities to the lye solution in AWE and AAEMWE and to liquid water in AEMWE and PEMWE but not too high in amount and combination to risk dysfunction or destruction of the stack.

However, the operation of the WE stacks (section 5) shall follow applicable safety requirements (Annex A) and the manufacturer's instructions. WE stacks should not be subjected to test and operating conditions jeopardising the safety or risk the dysfunction or destruction of the stack. Then, the selection of the stressing operating conditions (section 6.5) and their values and ranges shall, in principle, be within the specification of the stack manufacturer. Also, the test equipment used shall be suitable to apply the stressing operating conditions to the WE stack properly and in a safe manner.

When an aspect of performance degradation of a WE stack is initially not or insufficiently known, an iterative and possibly step-wise approach may be applied where one condition and mode of operation is varied within a reasonable parameter range considering the limiting operating conditions specified by the stack manufacturer. Progressively, more than one such condition or operation mode may be changed to arrive at a set of suitable stressing operating conditions and modes of operation of a WE stack. The established conditions and operating mode are for use in subsequent AST campaigns during R&D including stack prototype qualification.

Note, in comparison to presently identified stressing operating conditions and modes of operation suggested in available test protocols (Aricò *et al.*, 2013, Aricò *et al.*, 2016, Aricò *et al.*, 2018, Strataki, 2018, Stiber *et al.*, 2020, Mennemann *et al.*, 2021, Aricò *et al.*, 2020, Fouada-Onana, 2020), the various challenges experienced by WE stacks (section 3.4) will likely entail seeking to re-establish such conditions and modes of operation when testing improved or newly developed materials in one or another WEC component of a WE stack. Also, the design and configuration of stacks may adapt to changes in the set-up and operation of WE system in response

to these challenges. An ever-increasing share of VRE in the electricity grid across Europe and ongoing market developments in exceedingly diverse P-to-H₂ applications worldwide may also lead to stack adaptations.

The testing of WE stacks under given test conditions (section 6.3) consists of executing, usually at their beginning-of-life (BoL), all or selected types of tests according to a defined test plan (section 6.6). BoL of a stack shall be the start of its first operation following complete conditioning according to manufacturer instructions. Performance tests (section 6.7) of a stack at BoL are followed by durability tests (section 6.9) conducted for a prescribed period of use of the stack (section 6.9.2) employing application-oriented operation profiles of power, current, or voltage (section 6.8.2) as appropriate for the intended use of the stack (section 6.9.3).

Optionally, performance tests can be executed intermittently at specified intervals to assess how the stack has maintained or altered its performance characteristics ⁽²⁶⁾. Performance tests are mandatory at the end of the test campaign to evaluate the final degree of stack degradation using suitable KPIs (section 6.10). Changes in the performance characteristics of stacks are usually also presented graphically versus the total test duration or the number of performed operation profiles or sequence(s) of operation profiles (section 7).

The test plan (section 6.6) may require intermittently performing safety checks (Annex A) on the stack. Testing shall not continue for stacks unsafe to operate. It shall also be verified that neither the test equipment used nor employed BoP components are a cause for leakage.

6.2 Measurement techniques

The test equipment, measuring instruments, and measurement methods shall conform to relevant standards (*e.g.*, IEC 61010-1:2010+AMD1:2016 CSV (IEC, 2016)), test methods and procedures or best testing practices. Instruments shall be calibrated following applicable standard(s), measurement method(s), or procedure(s) as recommended by the manufacturer of the stack to meet the targeted uncertainty **(4.2.79)** of the concerned test parameter, whether TIP **(4.2.75)** or TOP **(4.2.77)**. The measurement set-up shall be documented. Calibration records and certificates of the measuring instruments should be available. Guidance on how to carry out an uncertainty analysis of the test results is provided by the Guide to the Expression of Uncertainty in measurement (GUM) (JCGM, 2008, JCGM, 2009, JCGM, 2020).

6.3 Test conditions

The test conditions with permissible variations are

- environmental conditions of the immediate surrounding (ambient) of the item under test: air (composition, velocity, pressure, temperature, humidity), salinity, ultraviolet (UV) radiation and other (weather) conditions, see Clause 4.4 of IEC 60204-1:2016+AMD1:2021 CSV (IEC, 2021a).
- actual operating conditions and modes of WE stack operation: start-up (hot and cold), shut-down, emergency stop and quiescence (standby).

They shall be defined before testing by the purpose(s) and objective(s) of the test campaign and conform with the specification of the stack as provided by the manufacturer.

However, the TIPs used in the various performance and durability tests constitute the operating conditions of the stack. In the test plan (section 6.6), the individual set point values of these TIPs shall be listed per test to be performed. The test plan should also list and name the TOPs (test results) to be measured or calculated per each test.

Further, while the inlet flow rates, temperatures, and compositions of liquid feeds (lye and water) are TIPs set (operating conditions), the electrode gas pressures and outlet temperatures are TOPs needing regulation. Other sets of operating conditions to be set are stack power (P_{stack}), stack current (I_{stack}), or stack voltage (U_{stack}).

In the first case, stack power is the TIP set while stack current and DC voltage are measured TOPs. In the second case, stack current is the TIP set. The stack voltage is a TOP measured. In the third case, stack voltage is the TIP set. The stack current is the TOP measured. In both cases, the stack power is a derived TOP to be calculated.

6.4 Reference test conditions

Reference test conditions are agreed upon before testing to facilitate the comparison of test results ⁽²⁷⁾. That is, reference test conditions are defined in agreement with manufacturer specifications to assess the performance

⁽²⁶⁾ The operation regime applied to the stack during a performance test can affect its degradation rate.

⁽²⁷⁾ For the three WEL technologies (AEL, AEMEL, and PEMEL), the strategic research and innovation agenda 2021-2027 of the Clean Hydrogen Partnership for Europe (SRIA) states 2024 and 2030 KPI targets for current density and hydrogen pressure but not for stack operating temperature (lye/water output temperature); see online at https://www.clean-hydrogen.europa.eu/knowledge-management/sria-key-performance-indicators-kpis_en.

degradation of WE stacks upon durability tests (section 6.9) conducted under stressing operating conditions (section 6.5) as appropriate, using polarisation curve measurements (section 6.7.11) and EIS measurements (section 6.7.12). Table 2 provides exemplary reference operating conditions recommended for a typical WE stack of type AWE, AEMWE, and PEMWE. For a test campaign aiming at a direct comparison of the three LTWE technologies or at a comparison of one stack technology versus another stack technology, the reference values of current density and stack operating temperature may be chosen to be the same. The actual reference operating conditions used shall be mentioned in the test report (Annex B).

Table 2: Recommended reference operating conditions for typical WE stacks (AWE, AEMWE and PEMWE)

Description	Unit	Symbol	AWE	AEMWE	PEMWE
Current density ⁽¹⁾	A/cm ²	J	0,4 (\pm 2,5 %)	1,0 (\pm 1 %)	2,0 (\pm 0,5 %)
Stack operating temperature ⁽²⁾ K	K	T _{stack}	353,15 (\pm 2 K)	333,15 (\pm 2 K)	353,15 (\pm 2 K)
Hydrogen pressure ⁽³⁾	kPa(g)	p _{H₂}	100 (\pm 2 %) ⁽⁴⁾	100 (\pm 2 %)	100 (\pm 2 %) ⁽⁴⁾

Note: The test plan (section 6.6) may list other conditions, including ambient environmental conditions of the WE stack, other than standard ambient temperature and pressure (SATP) conditions (**4.2.70**); see, for example, Clause 5.2.3.1 of ISO 22734:2019 (ISO, 2019a). This is especially relevant for WE stacks designed to operate at high pressure. Such stacks may not be able to operate at nominal power whilst at atmospheric pressure. This may be due to a different gas/liquid ratio or heat rejection efficiency, for example.

⁽¹⁾ This value should correspond to the maximum current density at BoL.

⁽²⁾ Unless otherwise agreed or specified by the manufacturer, the sensor (*e.g.*, thermocouple) position to determine the stack operating temperature should be at the centre of the tubing where lye/water is expelled by the WE stack close to its outlet.

⁽³⁾ The gauge pressure of hydrogen is measured at the outlet of the WE stack, preferably upon removal of residual liquid (lye or water). Gauge pressure is chosen to allow comparison when testing is conducted at sites having different atmospheric pressure levels. Balanced pressure or differential pressure operation of the stack shall be reported.

⁽⁴⁾ The SRIA anticipates 3 MPa as the pressure of hydrogen for the AEL and PEMEL technologies, see footnote 27. This pressure level is not selected as not all test equipment may allow testing at such pressure.

Source: JRC, 2023.

A WE stack should first be subjected to testing at reference test conditions before proceeding to testing under other specified test conditions including stressing operating conditions (section 6.5). Particularly, this regards polarisation curve measurements (section 6.7.11) and EIS measurements (section 6.7.12) at beginning-of-test (BoT) and at end-of-test (EoT) when WE stacks should again be tested under these reference test conditions to assess their overall performance degradation in the test campaign.

6.5 Stressing operating conditions

Major stressing operating conditions for WE stacks are (Tsotridis and Pilenga, 2021)

- *High current density operation:* This concerns operation at current densities in excess of the recommended value (see Table 2), for example, at up to 200 % of the recommended value.
- *Dynamic electric operation:* This concerns variable electric power, direct current, or DC voltage (including frequent ON/OFF operation) (Sayed-Ahmed *et al.*, 2024) employing simulated operation profiles (Alia *et al.*, 2019, Allidières *et al.*, 2019, Abmann *et al.*, 2020, Tsotridis and Pilenga, 2021, Malkow and Pilenga, 2023b, Reissner *et al.*, 2020), or RES-derived "real-world" operation profiles (section 6.8.2). For AWE stacks, OFF operation (zero supply current or OCV conditions), the phenomenon of the flow of a reverse current (**4.2.68**) occurs which leads to materials oxidation at the negatively charged cathode and materials reduction at the positively charged anode on the respective biP side (Haleem *et al.*, 2021).
- *AC ripple:* This concerns current (or voltage) fluctuations due to non-optimum AC/DC conversion, which are inevitable in real converters, superimposed onto DC (or DC voltage) supplied to the stack (Parache *et al.*, 2022).
- *Stack operating temperature:* The stack operating temperature influences the thermal-neutral voltage of the stack and the overvoltages as well as temperature-activated reaction kinetics at the electrodes.
- *Pressure:* This concerns especially anode-to-cathode differential pressure of the stack. For example, differential and absolute pressure cycling can be performed within the range of 60 % and 100 % of the respective nominal pressure levels (as specified by the manufacturer) or beyond to test the mechanical integrity of the stack. For operational safety, the stack may intermittently be subject to leak testing to identify significant external and internal leaks.
- *Lye/water inlet flow rate:* Too low lye/water inlet flow rates may result in insufficient wetting, risking dry active area and a reduction in gas bubble removal which could result in hotspots and overvoltage increases. Too high lye/water inlet flow rates could accelerate catalyst loss due to dissolution/erosion effects.

- **Lye/water impurities:** This concerns deliberate additions of calcium, copper, iron, potassium, magnesium, sodium, nickel, zinc, chloride, and carbonate ions (dissolved CO₂) to mainly affect IEM capability and catalyst activity (Becker *et al.*, 2023). Also, organics stemming from BoP components or products of in-stack materials deterioration, which is introduced to the stack by re-circulated lye/water, may deliberately be added.

Considering the specification of the stack and the test equipment capabilities, the test plan (section 6.6) shall specify the values and ranges of these stressing operating conditions including the compression factor applied to the original operation profile. The actual choice of a compression factor shall be such that the test equipment used can readily apply the intended dynamic operation of the stack and the ramp rates resulting from the simulated duty cycle(s) do not exceed those specified by the stack manufacturer, if any. That is, the compression factor has a lower limit per stressing operating condition individually and when combining two or more of such conditions as well as for the applied operation profile (section 6.8). An incremental approach to successively decrease the compression factor in discrete steps is recommended unless known, for example, from previously conducted single-cell or short-stack testing. The lower limit of compression factor is established when for a subsequent step no significant change occurs for the specified KPI (section 6.10) within its tolerance range.

In addition to accelerated stress testing of WE stacks, accelerated lifetime testing of the same type of stack may be conducted employing the original operation profile (section 6.8.2) while applying the same stack operating temperature and differential pressure without deliberately adding lye/water impurities. Note, performing accelerated lifetime testing of stacks as part of a test campaign using a suitable DoE (**4.2.26**) methodology along with physics-based modelling will assist in

- Identifying ranges of aggravated test conditions (stressing operating conditions) representative of the targeted application considering contributions from the use of BoP components used in WE systems (*e.g.*, liquid pollutants and electric power quality) and duty cycle in real-world stack operation which do not significantly alter the interdependency or eminence of intrinsic degradation mechanisms,
- Deriving robust AST transfer functions to correlate accelerating factors (**4.2.4**) with the occurrence or amplification of performance degradation in stacks which may not necessarily be expressible as linear relationships and eventually,
- Quantifying reliable rates of performance degradation allows prediction of durability under real-world conditions that can contemporaneously or sequentially be combined for evaluating various relevant stack operation modes.

This is especially useful to lower the overall R&D cost of WE stacks as test durations could significantly be reduced.

6.6 Test plan

The test plan shall take into account

- the manufacturer's specifications and instructions (*e.g.*, maximum voltage, pressure, and anode-to-cathode differential pressure, stack operating temperature, range of heating/cooling rates, and electrode gas compositions, cut-off voltage, etc.),
- an activation/conditioning procedure for initial stack operation upon manufacture, refurbishment, or prolonged non-operation (Lickert *et al.*, 2023, Tomić *et al.*, 2023),
- the test conditions: reference test conditions (section 6.4) and stressing operating conditions (section 6.5),
- the measurement techniques and instrumentation (section 6.2),
- the test type (section 6.7 and section 6.9), sequence, frequency, duration, and operation profiles,
- the DAQ with the number and logging rates of the data points ⁽²⁸⁾,
- re-start procedure upon unintended test interruptions other than pre-mature stack failure (*i.e.* test equipment break down, emergency stops, power supply failure, etc.),
- one or more KPIs, whether measured or derived TOPs, as a result of performance tests, and
- one or more test stop criteria with decision rules to prevent unintended failure or destruction of the stack.

⁽²⁸⁾ Considering the duration of the individual tests and the expected standard uncertainty in the measurements, different numbers, ranges, and data logging rates may apply to performance tests (section 6.7) and durability tests (section 6.9).

One or more KPIs shall be defined to assess the performance and durability of the test item. For this purpose, TIPs and TOPs should be specified to obtain KPIs as functions of such parameters; for example,

- TIPs:

- input electric power ($P_{\text{el,in}}$) (section 6.7.1), input current (I_{in}) (section 6.7.2), or input voltage (U_{in}) (section 6.7.3) to the stack,
- input temperature of lye ($T_{\text{in}}^{\text{lye}}$) to AWE and AAEMWE stacks and
- input water temperature (T_{in}^{w}) to PEMWE and AEMWE stacks.

- TOPs:

- stack operating temperature (T_{stack}),
- output pressure of hydrogen (p_{H_2}) and
- temperature of hydrogen (T_{H_2}) generated by the stack.

Environmental conditions other than SATP conditions may also be considered; for example, conditions likely to be experienced at intended installation sites of WE systems.

Consistent with the test campaign purpose(s) and objective(s), the test plan should specify the test methods and measurement techniques employed where standards, testing procedures, and manufacturer's instructions provide different possibilities. The test plan may also list (micro-structural) characterisation methods, for example, to perform post-test analysis of stack materials to gain more insight into the obtained test results.

6.7 Performance tests

6.7.1 Input electric power

The input electric power to a WE stack shall be determined in accordance with Clause 5.2.1 of ISO 16110-2:2010 (ISO, 2010b).

6.7.2 Input direct current

For a specified DC voltage set, the input direct current to a WE stack shall be determined from electric power measurements (section 6.7.1). The input direct current is the measured input electric power divided by the specified DC voltage.

6.7.3 Input DC voltage

For a specified direct current set, the input DC voltage applied to a WE stack shall be determined from electric power measurements (section 6.7.1). The input DC voltage is the measured input electric power divided by the specified direct current.

6.7.4 Input thermal power

The input thermal power ($P_{\text{th,in}}$) to a WE stack conveyed by heat transfer fluid(s), if any, shall be determined in accordance with Clause 5.2.2 of ISO 16110-2:2010 (ISO, 2010b).

6.7.5 Input power of compression

The input power of compression ($P_{\text{compr,in}}$) to a WE stack conveyed by pneumatic fluid(s), if any, shall be determined in accordance with Clause 5.2.2 of ISO 16110-2:2010 (ISO, 2010b).

6.7.6 Response time and ramp energy

Especially for delivering grid balancing services by electrolyzers (Alladières *et al.*, 2019), the response time (t_{resp}) of a WE stack to a given positive or negative ramp rate of input electric power, input direct current or input DC voltage (section 6.6) should be determined in accordance with Clause 5.6.1 of IEC 62282-8-201:2020 (IEC, 2020d). Ramp rates shall be consistent with the manufacturer's instructions. In addition to response time, the ramp energy (E_{ramp}) should be determined in accordance with Clause 14.6.3.2 of IEC 62282-3-201:2017+AMD1:2022 CSV (IEC, 2022) where, by analogy, reference to FC is replaced by water electrolyser.

The response time and ramp energy shall be recorded separately for positive and negative ramps in the test report (Annex B). The test plan shall specify appropriate indices to be added to t_{resp} and E_{ramp} to distinguish

between different ramps, whether positive or negative. The state before ramping may include a cold start (**4.2.18**) and hot start (**4.2.42**).

The determination of response time and ramp energy should at least be at BoT and EoT. It is for assessing whether or not changes in the ability of a WE stack to respond as intended to variations in input electric power (input direct current or input DC voltage) occurred due to durability testing of the stack (section 6.9).

6.7.7 Measurements of fluid feeds

For AWE and AAEMWE stacks, the pH value and the electrical conductivity (σ_{el}) of the alkaline solution (KOH electrolyte) entering the stack should, by analogy to water, be determined in accordance with Clauses 7.1 and 7.2 of ISO 3696:1987 (ISO, 1987), respectively.

For AEMWE and PEMWE stacks, the pH value and the electrical conductivity of the feed water to the stack should be determined in accordance with ISO 10523:2008 (ISO, 2008) and ISO 7888:1985 (ISO, 1985), respectively.

The stack manufacturer may specify other means to measure the pH value and the electrical conductivity of lye solution and feed water.

6.7.8 Hydrogen output rate and quality

The product gas output rate, also known as the product gas molar flow rate ($q_{n,out}$) of the WE stack, should be determined in accordance with Clause 5.2.11.1 of ISO 22734:2019 (ISO, 2019a). From the product gas molar flow rate, the hydrogen output rate, also known as the molar flow rate of hydrogen (q_{n,H_2}), shall be calculated as follows

$$q_{n,H_2} \text{ (mol/h)} = x_{n,H_2} \text{ (mol/mol)} \cdot q_{n,out} \text{ (mol/h)}; \quad (6.7.1)$$

x_{n,H_2} is the molar concentration of hydrogen in the product gas to be determined by gas analysis in accordance with, for example, Clause 5.2.2.2 of ISO 16110-2:2010 (ISO, 2010b). From the molar flow rate of hydrogen, the volumetric flow rate of hydrogen (q_{V,H_2}) generated under SATP conditions is calculated as follows

$$q_{V,H_2} \text{ (m}^3/\text{h)} = q_{n,H_2} \text{ (mol/h)} \cdot V_{m,H_2} \text{ (m}^3/\text{mol)}; \quad (6.7.2)$$

$V_{m,H_2} \approx 24,79 \cdot 10^{-3} \text{ m}^3/\text{mol}$ is the molar volume of hydrogen under SATP conditions ⁽²⁹⁾. The hydrogen output quality of a WE stack other than the molar concentration of hydrogen in the product gas, in particular, humidity, is to be determined in accordance with, for example, Clause 5.2.11.2 of ISO 22734:2019 (ISO, 2019a).

Note, neglecting gas crossover and leakages in the stack as well as stray currents (**4.2.71**) in AWE stacks, the theoretical volumetric flow rate of hydrogen generated under SATP conditions can be calculated as follows

$$q_{V,H_2}^{\text{theo}} \text{ (m}^3/\text{h)} = \frac{\eta_F \%}{100 \%} \cdot \frac{V_{m,H_2} \text{ (m}^3/\text{mol)} \cdot N_{\text{cell}} \cdot I_{dc} \text{ (A)}}{z \cdot F \text{ (C/mol)}} \cdot 3600 \frac{\text{s}}{\text{h}}; \quad (6.7.3)$$

η_F is the Faradaic efficiency, $F = 96485,3321 \text{ C/mol}$ is Faraday constant and z is the number of electrons exchanged in the WEC reactions (3.1.2), (3.1.3) or (3.1.4). For the various modes of stack operation, the dependency of the volumetric flow rate of hydrogen on Faradaic efficiency and current may be established by preliminary testing.

The mass flow rate of hydrogen (q_{m,H_2}) generated by a WE stack related to SATP conditions is calculated as follows

$$q_{m,H_2} \text{ (kg/h)} = q_{n,H_2} \text{ (mol/h)} \cdot m_{H_2} \text{ (kg/mol)}; \quad (6.7.4)$$

$m_{H_2} \approx 2,02 \cdot 10^{-3} \text{ kg/mol}$ is the molar mass of hydrogen under SATP conditions ⁽³⁰⁾.

6.7.9 Oxygen output rate and concentration

The oxygen output rate, also known as the molar flow rate of oxygen (q_{n,O_2}) of the WE stack, should be determined in accordance with Clause 5.2.11.1 of ISO 22734:2019 (ISO, 2019a). The molar concentration of oxygen (x_{n,O_2}), is to be determined in accordance with, for example, Clause 5.2.11.2 of ISO 22734:2019 (ISO, 2019a).

⁽²⁹⁾ At a temperature of 273,15 K and a pressure of 101,3 kPa, the molar volume of hydrogen is $V_{m,H_2} \approx 22,41 \cdot 10^{-3} \text{ m}^3/\text{mol}$.

⁽³⁰⁾ The mass of deuterium and tritium as hydrogen isotopes are not considered due to their negligible natural abundance.

6.7.10 Water/lye quality measurements

For AEMWE and PEMWE stacks with IEMs and other materials containing fluorine (F), PFAS, and polycyclic aromatic hydrocarbons (PAH) such as membranes, coatings, and sealants, the quality of the water entering and exiting the stack should be determined regarding

- (a) fluoride concentration (c_F) in accordance with ISO 10359-1:1992 (ISO, 1992), ISO 10359-2:1994 (ISO, 1994), ISO 10304-1:2007 (ISO, 2007), ISO/TS 17951-1:2016 (ISO, 2016b), ISO/TS 17951-2:2016 (ISO, 2016b) or ISO/TS 15923-2:2017 (ISO, 2017c), as appropriate.
- (b) PFAS concentration (c_{PFAS}) in accordance with ISO 21675:2019 (ISO, 2019c) or any other suitable analysis technique ⁽³¹⁾, as feasible. Information available from cell/stack material suppliers could help determine potential species likely to occur as a result of degrading PFAS.
- (c) PAH concentration (c_{PAH}) in accordance with ISO 17993:2002 (ISO, 2002), ISO 7981-1:2005 (ISO, 2005a), ISO 7981-2:2005 (ISO, 2005b) or ISO 28540:2011 (ISO, 2011b), as appropriate.

The content of fluoride, PFAS, and PAH in the water on the stack inlet should, in principle, be subtracted from the respective content at the stack outlet unless in-loop water filtration materials are subject to analysis ⁽³²⁾. Upon subtracting any fluoride inlet content from that at the stack outlet, the occurrence of fluoride in the water exiting the stack as an end product of the decomposition of fluoropolymers is conclusive evidence for performance degradation of the stack by materials deterioration. Also, upon applying a similar subtraction procedure, the occurrence of PFAS and PAH in the exit water of a stack as fragments of partially decomposed polymers is likewise conclusive evidence for performance degradation of the stack by materials deterioration. The identified type of fragments along with their analysed quantities may assist in determining the pathways of material deterioration to gain useful insight into the performance degradation of WE stacks.

Especially for WE stacks tested using ion impurities in water/lye as stressing operating conditions (section 6.5), the quality of water/lye entering and exiting the stack shall be determined regarding

- calcium concentration (c_{Ca}) in accordance with ISO 6058:1984 (ISO, 1984a), ISO 7980:1986 (ISO, 1986a), ISO 14911:1998 (ISO, 1998) or ISO/TS 15923-2:2017 (ISO, 2017c), as appropriate,
- chloride concentration (c_{Cl}) in accordance with ISO 9297:1989 (ISO, 1989), ISO 10304-4:1997 (ISO, 1997), ISO 15682:2000 (ISO, 2000), ISO 10304-1:2007 (ISO, 2007), ISO 15923-1:2013 (ISO, 2013) or ISO 10304-4:2022 (ISO, 2022c), as appropriate,
- copper concentration (c_{Cu}) in accordance with ISO 8288:1986 (ISO, 1986b),
- iron concentration (c_{Fe}) ⁽³³⁾ in accordance with ISO 6332:1988 (ISO, 1988) or ISO/TS 15923-2:2017 (ISO, 2017c), as appropriate,
- potassium concentration (c_K) ⁽³⁴⁾ in accordance with ISO 9964-2:1993 (ISO, 1993b), ISO 9964-3:1993 (ISO, 1993c) or ISO 14911:1998 (ISO, 1998), as appropriate,
- magnesium concentration (c_{Mg}) in accordance with ISO/TS 15923-2:2017 (ISO, 2017c) or ISO 6059:1984 (ISO, 1984b), as appropriate,
- sodium concentration (c_{Na}) ⁽³⁵⁾ in accordance with ISO 9964-1:1993 (ISO, 1993a), ISO 9964-3:1993 (ISO, 1993c) or ISO 14911:1998 (ISO, 1998), as appropriate,
- nickel concentration (c_{Ni}) in accordance with ISO 8288:1986 (ISO, 1986b) and
- zinc concentration (c_{Zn}) in accordance with ISO 8288:1986 (ISO, 1986b).

The content of calcium (Ca), chloride (Cl⁻), copper (Cu), iron (Fe), potassium (K), magnesium (Mg), sodium (Na), nickel (Ni), and zinc (Zn) in the water on the inlet of the stack shall be subtracted from those at the stack outlet.

⁽³¹⁾ Note, the appendix to Annex E.4 of the Annex XV Restriction Report on the manufacture, placing on the market, and use of PFAS contains an overview of analytical methods for the analysis of PFAS in different matrices; see at <https://echa.europa.eu/restrictions-under-consideration/-/substance-rev/72301/term>.

⁽³²⁾ For sampling water and subsequent analysis as well as water filtration, materials free of fluoride, PFAS, and PAH shall be used.

⁽³³⁾ Notably, in AWE, low iron concentration due to leaching of Fe from BoP components into lye can cause significant changes in stack performance (Demnitz *et al.*, 2024).

⁽³⁴⁾ Potassium is not relevant for AWE stacks using 30 % KOH as a lye solution.

⁽³⁵⁾ Sodium is not relevant for AWE stacks using Na-containing lye solution.

6.7.11 Polarisation curve measurements

The measurement of the current-voltage (I_{dc} - U_{dc}) characteristics or polarisation curves of WE stacks shall be determined by applying the EU harmonised polarisation curve test method for LTWEL (Malkow *et al.*, 2018b). Polarisation curve measurements are commonly performed under reference test conditions (section 6.4) by sweeping from a start set point to an end set point or stepwise at a minimum of three different set points to be specified in the test plan (section 6.6). In deciding on the actual set points, the test plan shall take into account the possibility of gas crossover especially at low current densities not to jeopardise testing safety.

Polarisation curve measurements under reference test conditions may act as an operation phase for *in-situ* stack regeneration to recover reversible degradation (Tsotridis and Pilenga, 2021). The test plan (section 6.6) may also foresee another type of procedure for stack regeneration to be applied prior to measurements.

In case the cell voltage difference in galvanostatic polarisation measurements is negligible (*i.e.* less than 10 mV for all data points) when starting from (near) zero current to proceeding to maximum current (I_{max}) (ascending polarisation curve) compared to starting from maximum current and proceeding to (near) zero current (descending polarisation curve), subsequent measurements may be conducted unidirectionally with respect to changes in current while maintaining a steady stack operating temperature (Lettenmeier *et al.*, 2016).

The maximum current is the current at which the voltage of any one cell in the stack is for three consecutive samplings equal to the cut-off voltage ($U_{cut-off}$) defined before testing as part of the test plan (section 6.6) or following manufacturer recommendations. As a voltage limit, the cut-off voltage is to prevent excessive stack degradation. Thus, polarisation curve measurements are preferably conducted under galvanostatic conditions.

For AWE stacks, no OCV (zero supply current) shall be used as a set point in the polarisation curve measurement to prevent the flow of a reverse current in the stack leading to materials deterioration (see section 6.5).

From the I_{dc} - U_{dc} curve, the current-electric power (I_{dc} - P_{el}) curve is calculated, see equation (3.3.1b). The energy efficiency (η_e) given by equation (6.7.16) versus the electric power density given by equation (3.3.1c) and the electrical efficiency (η_{el}) given by equation (6.7.16) versus the electric power density is also plotted to assess the optimum operating (working) point (**4.2.51**) of the WE stack under the test conditions.

The distribution of the cell voltage is a measure of voltage homogeneity in the stack. The mean absolute error of average cell voltage ($MAE_{\overline{U}_{cell}}$) and standard deviation of average cell voltage ($SD_{\overline{U}_{cell}}$) are two complementary statistical indicators of cell voltage distribution calculated as follows

$$MAE_{\overline{U}_{cell}} (\text{mV}) = \frac{1}{N_{cell}} \sum_n |U_{cell,n} (\text{V}) - \overline{U}_{cell} (\text{V})| \cdot 10^3 \frac{\text{mV}}{\text{V}} \quad (6.7.5a)$$

$$SD_{\overline{U}_{cell}} (\text{mV}) = \sqrt{\frac{1}{N_{cell}-1} \sum_n (U_{cell,n} (\text{V}) - \overline{U}_{cell} (\text{V}))^2} \cdot 10^3 \frac{\text{mV}}{\text{V}} \quad (6.7.5b)$$

where

$$\overline{U}_{cell} (\text{V}) = \frac{1}{N_{cell}} \sum_n U_{cell,n} (\text{V}) \quad (6.7.5c)$$

is the average cell voltage of all series-connected WECs in the stack; $U_{cell,n}$ is the voltage of cell number n. A significant deviation of the average cell voltage times the number of cells from the measured stack voltage, $U_{stack} \not\approx N_{cell} \cdot \overline{U}_{cell}$, indicates ohmic resistances in the stack materials other than cell materials and contact resistances at the various in-stack interfaces are not negligible. Such deviation also depends on the in-stack positions of the terminals across which the individual cell voltages and the stack voltage are measured.

To gain more insight into the performance degradation by voltage increases of the WECs in a WE stack, a voltage breakdown analysis may optionally be performed on the measured polarisation curves for attributing voltage increases to individual WEC components (Flick *et al.*, 2015, Gerhardt *et al.*, 2021, Dizon *et al.*, 2022). This is accomplished by additive contributions of the various overvoltages to the OCV of the WEC given by equation (6.7.7a). Then, the voltage of the WEC is calculated as follows

$$U_{WEC} (\text{V}) = U_{OCV} (\text{V}) + U_{act} (\text{V}) + I_{dc} (\text{A}) \cdot R_{lf} (\Omega) + U_{conc} (\text{V}); \quad (6.7.6)$$

U_{act} is the activation polarisation voltage given by equation (6.7.7b), R_{lf} is the low-frequency resistance estimated as the slope of the polarisation curve ⁽³⁶⁾ and U_{conc} is the concentration polarisation voltage given by equation (6.7.7c). Specifically, the voltage contributions are

- Open circuit voltage or open circuit potential (U_{OCP}) which is different from the reversible voltage due to
 - gas crossover leading to mixed electrode potentials,
 - locally different catalyst surface concentrations and

⁽³⁶⁾ The low-frequency resistance can also be determined by EIS measurements (section 6.7.12), see equation (6.7.12).

- gas solubility at the water-ionomer-catalyst interface.
- Ohmic polarisation (**4.2.50**) represented by $I_{dc} \cdot R_{lf}$ where the low-frequency resistance or ohmic resistance of the WEC is due to
 - the electrodes including diffusion media,
 - the electrolyte, namely lye solution in AWE and AAEMWE and hydrated IEM in AEMWE and PEMWE, and
 - interfacial contact surfaces of the stack between WEC, biPs, and current collectors.
- Activation polarisation (**4.2.5**) due to the kinetics of
 - the OER (3.1.2a), (3.1.3a) or (3.1.4a) at the anode and
 - the HER (3.1.2b), (3.1.3b) or (3.1.4b) at the cathode.
- Concentration polarisation (**4.2.21**) due to mass transfer limitations in the electrodes of the stack including hindrance caused by gas bubble formation.

In the absence of gas crossover and other non-ideal conditions, the temperature-dependent open circuit voltage is calculated as follows (Bernt, 2019)

$$U_{OCV}(T) \text{ (V)} = 1,2291 \text{ V} - 0,0008456 \text{ (V/K)} (T \text{ (K)} - 298,15 \text{ K}) + \frac{R_g \text{ (J/(mol K))} \cdot T \text{ (K)}}{z \cdot F \text{ (C/mol)}} \log \left(\frac{p_{H_2} \text{ (kPa)}}{p^0 \text{ (kPa)}} \cdot \left(\frac{p_{O_2} \text{ (kPa)}}{p^0 \text{ (kPa)}} \right)^{0,5} \right); \quad (6.7.7a)$$

p_{H_2} and p_{O_2} are the partial pressure of hydrogen and partial pressure of oxygen, respectively. The activity of liquid water (a_{H_2O}) is taken as unity. The exponent of 0,5 stems from the fact that half a mole of oxygen is generated by electrolysis from one mole of liquid water, see reaction (3.1.1). The temperature-dependent activation and concentration polarisation voltages are calculated as follows (Hernández-Gómez *et al.*, 2020)

$$U_{act}(T) \text{ (V)} = \frac{R_g \text{ (J/(mol K))} \cdot T \text{ (K)}}{z \cdot F \text{ (C/mol)}} \log \left(\left(\frac{I_{0,a}(T) \text{ (A)}}{I_a \text{ (A)}} \right)^{\alpha_a} \cdot \left(\frac{I_{0,c}(T) \text{ (A)}}{I_c \text{ (A)}} \right)^{\alpha_c} \right) \text{ and} \quad (6.7.7b)$$

$$U_{conc}(T) \text{ (V)} = \frac{R_g \text{ (J/(mol K))} \cdot T \text{ (K)}}{z \cdot F \text{ (C/mol)}} \log \left(\left(\frac{c_{O_2} \text{ (mol)}}{c_{O_2}^0 \text{ (mol)}} \right) \cdot \left(\frac{c_{H_2} \text{ (mol)}}{c_{H_2}^0 \text{ (mol)}} \right) \right); \quad (6.7.7c)$$

$I_{0,a}$, I_a , $I_{0,c}$, I_c , α_a and α_c are anodic exchange current, anodic current, cathodic exchange current, cathodic current, anodic charge transfer coefficient, and cathodic charge transfer coefficient, respectively while c_{O_2} and c_{H_2} are the oxygen concentration and hydrogen concentration at the electrolyte-electrode interfaces, respectively and $c_{O_2}^0$ and $c_{H_2}^0$ are their respective equilibrium concentrations. Further information on voltage breakdown analysis is given elsewhere (Ma *et al.*, 2021, Falcão and Pinto, 2020, Gerhardt *et al.*, 2021).

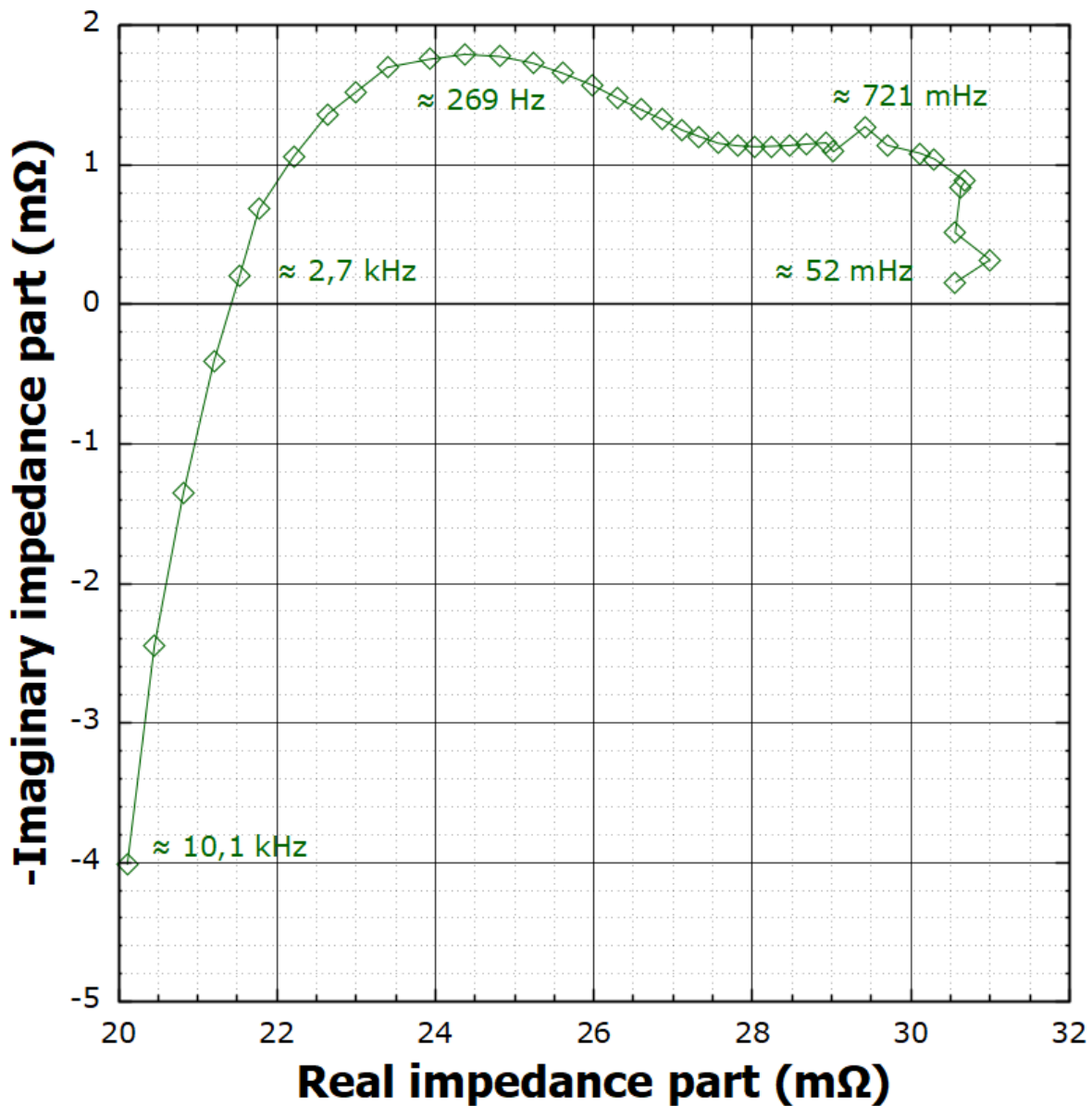
6.7.12 EIS measurements

The electrical impedance (Z) of individual cells in a WE stack as a function of perturbation frequency (f) ⁽³⁷⁾ may be determined optionally and where feasible. The EU harmonised test procedure on EIS for WECs (Malkow *et al.*, 2018a) should be applied. The measurements should be performed under the same test and operating conditions including the DC set point values as applied in the polarisation curve measurements (section 6.7.11). In case EIS measurements and polarisation curve measurements are conducted simultaneously, AC and DC contributions are contained in the current and voltage data. The test plan (section 6.6) shall specify the excitation type, magnitude, frequencies, and the number of repetitions.

In Nyquist plots (**4.2.49**), the (negative) imaginary part of the electrical impedance is plotted against the real part of the electrical impedance, $(-\Im Z)$ versus $\Re Z$, Figure 5. The resulting graph displays one or more semi-arcs whereby the number of distinguishable semi-arcs coincides with the number of relaxation times or time constants (τ) due to polarisation processes such as charge build-up at and transfer across the WEC

⁽³⁷⁾ In EIS measurements, the output under potentiostatic conditions is electrical impedance, $Z=|Z|^2 Y^* =|Z| e^{i\arg(Z)}$; $|\cdot|$ denotes modulus or gain, $\arg(\cdot)$ denotes phase or argument, superscript * denotes complex conjugation and i is the imaginary unit with property, $(\pm i)^2 = -1$. Under galvanostatic conditions, the output is electrical admittance, $Y=|Y|^2 Z^* =|Y| e^{i\arg(Y)}$.

Figure 5: Graphical representation of the real and imaginary parts of the electrical impedance of a water electrolysis cell in a Nyquist plot.⁽¹⁾



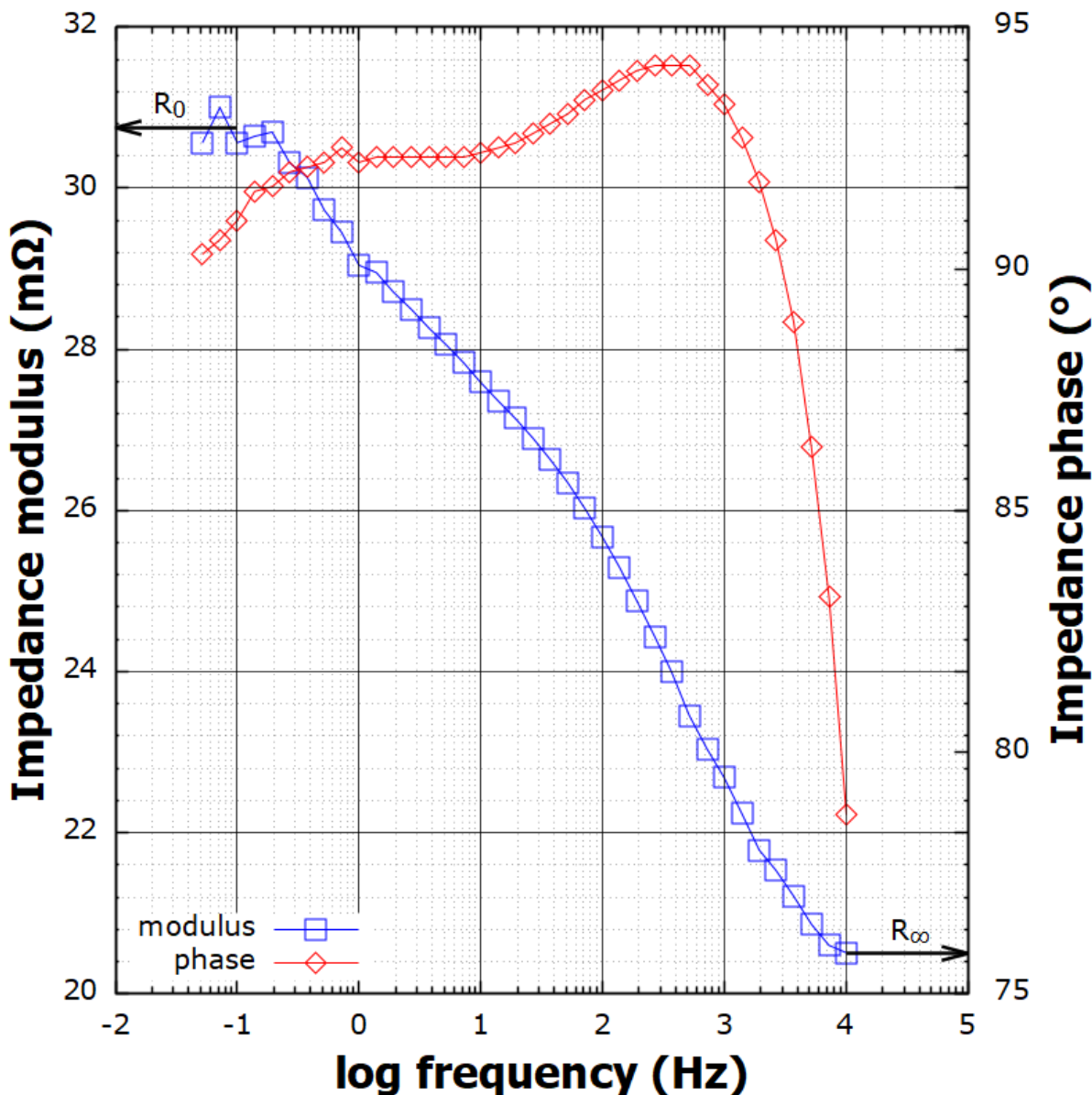
⁽¹⁾ Note that the high-frequency part of the spectrum is affected by inductance arising from the measurement set-up, where inductances are induced in the wiring used to connect the cell to the power supply and measurement device (leakage current), as well as converters used in the device instrumentation.

Source: JRC, 2023.

interfaces as well as species transport (convection, diffusion, migration, reaction, etc.) within the WEC⁽³⁸⁾. Often, semi-arcs overlap considerably so that not all time constants are identifiable. That is, not all polarisation processes may visually be detectable from Nyquist plots, see Figure 5.

In Bode plots (4.2.15), the modulus of electrical impedance⁽³⁹⁾ and its phase⁽⁴⁰⁾ are plotted against the frequency in logarithmic scale, $|Z|$, $\arg(Z)$ versus $\log f$, Figure 6. From a Bode plot, the values of the low-

Figure 6: Graphical representation of the electrical impedance modulus and phase of a water electrolysis cell in a Bode plot. The data are the same as in Figure 5.



Source: JRC, 2023.

⁽³⁸⁾ The combination of any two distinct passive (lumped) circuit elements such as a capacitor having electrical admittance, $Y=\omega C$, an inductor having electrical impedance, $Z=\omega L$, and a resistor having electrical impedance, $Z=R$, arranged in parallel or series, makes up a characteristic time constant; $\omega=2\pi f$ is angular frequency. It is attributable to the dissipation of electric energy (E_{el}) by resistors accounting for the resistivity in bulk materials and the resistance (R) of the various interfaces of a WEC or a WE stack due to electronic and ionic conduction, as well as energy storage by capacitors accounting for capacitance (C , negative reactance) at electrically charged interfaces (*i.e.* electrolyte-electrode intersections) and inductors accounting for inductance (L , positive reactance) in conductors (*i.e.* wires, converters, etc.). Non-ideal circuit elements include constant phase elements (CPEs) having electrical impedance, $Z=((\omega)^{\alpha} Q_C)^{-1}$ or $Z=(\omega)^{\alpha} Q_L$ ($0 < \alpha < 1$ where α is a power exponent, Q_C is a non-ideal capacitance and Q_L is a non-ideal inductance) to account for frequency dispersion (Alexander *et al.*, 2016, Córdoba-Torres, 2017, Kartci *et al.*, 2020, Fitzek *et al.*, 2022) and distributed circuit elements (Warburg, Gerischer, etc.) to account for species transport (Huang, 2018, Chowdhury and Kant, 2018, Boukamp, 2017). They too have individual (mean) time constants.

⁽³⁹⁾ Also, the real and imaginary parts of the electrical impedance may be displayed in Bode plots.

⁽⁴⁰⁾ Instead of phase or argument of electrical impedance, the tangent of the loss angle of electrical impedance may be plotted against the logarithm of frequency (or angular frequency), $\tan(\Im Z / \Re Z)$ versus $\log f$ (or $\log \omega$).

frequency resistance (R_{lf}) (see equation (6.7.12)) and the high-frequency resistance (R_{hf}) (see equation (6.7.9)) can often be read directly, which is not necessarily the case for a Nyquist plot, cf. Figure 6 and Figure 5.

In principle, the ohmic resistance is the infinite-frequency resistance (R_∞) that is the electrical impedance at high frequencies ($f \rightarrow \infty$) with vanishing reactance, $\Im Z [f \rightarrow \infty] = 0$,

$$\lim_{f \rightarrow \infty} \Re Z [f] (\Omega) = R_\infty (\Omega). \quad (6.7.8)$$

Practically, the ohmic resistance is the electrical impedance measured at the highest of the probed frequencies (f_{max}) where $\Im Z [f \rightarrow f_{max}] \rightarrow 0$,

$$\lim_{f \rightarrow f_{max}} \Re Z [f] (\Omega) \approx R_{hf} (\Omega) \quad (6.7.9)$$

to represent the resistance of the electrolyte. Unfortunately, the reactance at f_{max} is often not negligible, $\Im Z [f_{max}] > 0$, see Figure 5. The polarisation resistance is the difference between the zero-frequency resistance (R_0) and the infinite-frequency resistance,

$$R_{pol} (\Omega) = R_0 (\Omega) - R_\infty (\Omega). \quad (6.7.10)$$

The zero-frequency resistance is the electrical impedance at low frequencies ($f \rightarrow 0$) with vanishing reactance, $\Im Z [f \rightarrow 0] = 0$,

$$\lim_{f \rightarrow 0} \Re Z [f] (\Omega) = R_0 (\Omega). \quad (6.7.11)$$

Practically, the zero-frequency resistance is the electrical impedance measured at the lowest of the probed frequencies (f_{min}) where $\Im Z [f \rightarrow f_{min}] \rightarrow 0$,

$$\lim_{f \rightarrow f_{min}} \Re Z [f] (\Omega) \approx R_{lf} (\Omega). \quad (6.7.12)$$

Consequently, the polarisation resistance is approximated to the difference between the low-frequency resistance and the high-frequency resistance,

$$R_{pol} (\Omega) \approx R_{lf} (\Omega) - R_{hf} (\Omega). \quad (6.7.13)$$

The area-specific resistance is calculated as follows

$$R_{ASR} (\text{m}\Omega\cdot\text{cm}^2) = R_{lf} (\Omega) \cdot A_{act} (\text{cm}^2) \cdot 1000 \frac{\text{m}\Omega}{\Omega}. \quad (6.7.14)$$

Also, the estimated value of the slope of the polarisation curve (section 6.7.11) may be used in place of R_{lf} to calculate R_{ASR} according to equation (6.7.14).

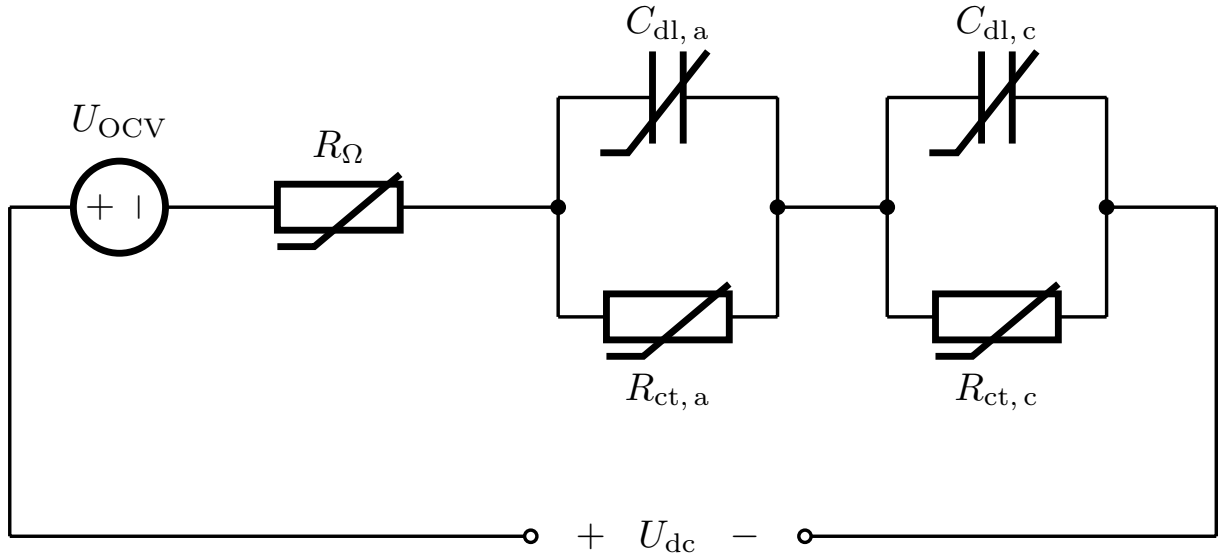
Guidance on EIS measurements (Siracusano *et al.*, 2018, Szekeres *et al.*, 2021) is provided by Clause 10.7.2.2 of IEC 62282-7-2:2014 (IEC, 2014b) and Clause 6.3.10 of IEC 62282-8-101:2020 (IEC, 2020c) while guidance on post-processing of EIS data is provided by Clause 7.6.3 of IEC 62282-8-101:2020 (IEC, 2020c). Note, useful software tools to perform such data post-processing are listed in term 403 on p. 66 (online version) of the recently published electrolysis terminology document (Malkow *et al.*, 2021).

Further, advanced EIS data analysis includes the method of distribution of uncorrelated relaxation times (DRT) (Plank *et al.*, 2024, Boukamp, 2020) for detecting better resolved time constants (relaxation times) without explicit assumptions except for the applicability of the principle of superposition (**4.2.65**). Primarily, DRT analysis provides for the total number and the values of relaxation times being the time constants of relaxation phenomena occurring in WECs due to heat and species transport (conduction, convection, diffusion, migration, reaction, etc.). Significant changes in these parameters upon prolonged in-stack exposure of a WEC under normal operating conditions or stressing operating conditions may occur due to degradation. Further, the number of time constants in the equivalent electric circuit (EEC) model should necessarily match the number of relaxation times unambiguously identified by DRT analysis. Also, the polarisation resistance is the area under the graph of the DRT (Malkow, 2019). Depending on the EIS data, the value of the DRT function can be real or complex (Malkow, 2021) (⁴¹).

Advantageously, the determined values of the time constants and the polarisation resistance may serve as initial start values for subsequent complex non-linear least squares (CNLS) fitting of the measured EIS data to a suitable EEC model simulating the observed WEC electrical impedances (Boukamp, 2004, Macdonald and Potter, 1987, Boukamp, 1986). This type of data post-processing, also known as parameter identification, may eventually identify WEC parameters as values of sought microscopic quantities (diffusion coefficients, permeabilities, reaction rates, etc.). A simplified EEC model of an electrolyser connected to a DC source is presented in Figure 7 (Chen *et al.*, 2022).

⁽⁴¹⁾ Unfortunately, software codes to determine complex-valued DRT are not available to date. Also, a theory of DRT functions with correlated relaxation times representative of interdependent (hierarchical, interacting, coupled, or cooperative), spatially distributed or delayed polarisation processes (resistive-capacitive or resistive-inductive) prevailing in electrochemical cells including WECs is yet missing.

Figure 7: Simplified EEC model of an electrolyser connected to a DC source; $C_{dl,a}$, $C_{dl,c}$, $R_{ct,a}$ and $R_{ct,c}$ are the temperature-dependent anodic double-layer capacitance, cathodic double-layer capacitance, anodic charge transfer resistance, and cathodic charge transfer resistance, respectively, and R_Ω is the temperature-dependent ohmic resistance.



Source: JRC, 2023.

The anodic double-layer capacitance, cathodic double-layer capacitance, anodic charge transfer resistance, and cathodic charge transfer resistance are calculated as follows

$$C_{dl,a}(T) \text{ (mF)} = \frac{\epsilon(T) \text{ (F/m)} \cdot A_{act} \text{ (cm}^2\text{)}}{d_{dl} \text{ (nm)}} \cdot 10^5 \text{ cm}^2/\text{nm/m} \cdot 10^{-3} \frac{\text{mF}}{\text{F}}, \quad (6.7.15a)$$

$$C_{dl,c}(T) \text{ (mF)} = \frac{\epsilon(T) \text{ (F/m)} \cdot A_{act} \text{ (cm}^2\text{)}}{d_{dl} \text{ (nm)}} \cdot 10^5 \text{ cm}^2/\text{nm/m} \cdot 10^{-3} \frac{\text{mF}}{\text{F}}, \quad (6.7.15b)$$

$$R_{ct,a}(I, T) \text{ (\Omega)} = \frac{R_g \text{ (J/(mol K))} \cdot T \text{ (K)}}{z \cdot F \text{ (C/mol)} \cdot I_{0,a}(T) \text{ (A)}} \text{ and} \quad (6.7.15c)$$

$$R_{ct,c}(I, T) \text{ (\Omega)} = \frac{R_g \text{ (J/(mol K))} \cdot T \text{ (K)}}{z \cdot F \text{ (C/mol)} \cdot I_{0,c}(T) \text{ (A)}}; \quad (6.7.15d)$$

ϵ is the absolute permittivity of the double-layer having length d_{dl} . Note that the charge transfer resistances depend on current.

6.7.13 Efficiency determination

For WE stacks, the energy efficiency based on HHV under SATP conditions of hydrogen ($\eta_{HHV,e}^0$), the energy efficiency based on LHV under SATP conditions of hydrogen ($\eta_{LHV,e}^0$), the electrical efficiency based on HHV under SATP conditions of hydrogen ($\eta_{HHV,el}^0$) and the electrical efficiency based on LHV under SATP conditions of hydrogen ($\eta_{LHV,el}^0$) shall be determined by applying the recently published energy performance testing procedure (Malkow and Pilenga, 2023a) under the same test and operating conditions as applied in the polarisation curve measurements. These efficiencies are calculated as follows

$$\eta_{HHV,e}^0 \text{ (\%)} = \frac{q_{n,H_2} \text{ (mol/h)} \cdot HHV_{H_2} \text{ (kWh/mol)}}{P_{el,dc} \text{ (kW)} + P_{th} \text{ (kW)} + P_{compr} \text{ (kW)}} \cdot 100 \%, \quad (6.7.16a)$$

$$\eta_{LHV,e}^0 \text{ (\%)} = \frac{q_{n,H_2} \text{ (mol/h)} \cdot LHV_{H_2} \text{ (kWh/mol)}}{P_{el,dc} \text{ (kW)} + P_{th} \text{ (kW)} + P_{compr} \text{ (kW)}} \cdot 100 \%, \quad (6.7.16b)$$

$$\eta_{HHV,el}^0 \text{ (\%)} = \frac{q_{n,H_2} \text{ (mol/h)} \cdot HHV_{H_2} \text{ (kWh/mol)}}{P_{el,dc} \text{ (kW)}} \cdot 100 \% \text{ and} \quad (6.7.16c)$$

$$\eta_{LHV,el}^0 \text{ (\%)} = \frac{q_{n,H_2} \text{ (mol/h)} \cdot LHV_{H_2} \text{ (kWh/mol)}}{P_{el,dc} \text{ (kW)}} \cdot 100 \% ; \quad (6.7.16d)$$

$\text{HHV}_{\text{H}_2} = 79,4 \text{ kWh/mol}$ and $\text{LHV}_{\text{H}_2} = 67,2 \text{ kWh/mol}$ are respectively the higher heating value and the lower heating value of hydrogen under SATP conditions (Tsotridis and Pilenga, 2018), q_{n,H_2} is given by equation (6.7.1), $P_{\text{el,dc}}$ is given by equation (3.3.1b), P_{th} is given by equation (5.1.1b) and P_{th} is given by equation (5.1.2b).

6.8 Operation profiles

6.8.1 General

Operation profiles, whether profiles of the input electric power, input direct current, or input DC voltage versus time (t), are intended to simulate, under given test conditions (section 6.3), stack operation for use in the application concerned. Apart from simulated operation profiles (Alia *et al.*, 2019, Allidières *et al.*, 2019, Aßmann *et al.*, 2020, Tsotridis and Pilenga, 2021, Malkow and Pilenga, 2023b, Reissner *et al.*, 2020), this includes real-world operation profiles (duty cycles) derived from RES-based power profiles typical for the intermittent supply of electricity to a WE system by various RES types, namely PV power (4.2.60) and wind power (4.2.82).

Herein, we use open data of the Belgian electricity grid with a 15-minute time resolution (Elia Transmission Belgium SA, 2023) by normalising the measured electric power (P_{el}) by the monitored peak electric power to derive operation profiles for PV electric power (Figure 8 and Figure 9) and WT electric power (Figure 10 and Figure 11).

For testing, the derived profiles are expressed in terms of input electric power or translated into input current using a typical value of rated voltage, or translated into input voltage using the rated current. The time interval of a profile (duty cycle) is usually a fixed period comprising the time required to carry out a given number of consecutive profiles of the same type or a sequence of profiles of different types. Thus, individual profiles (duty cycles) may constitute building blocks of a test sequence specified in the test plan (section 6.6).

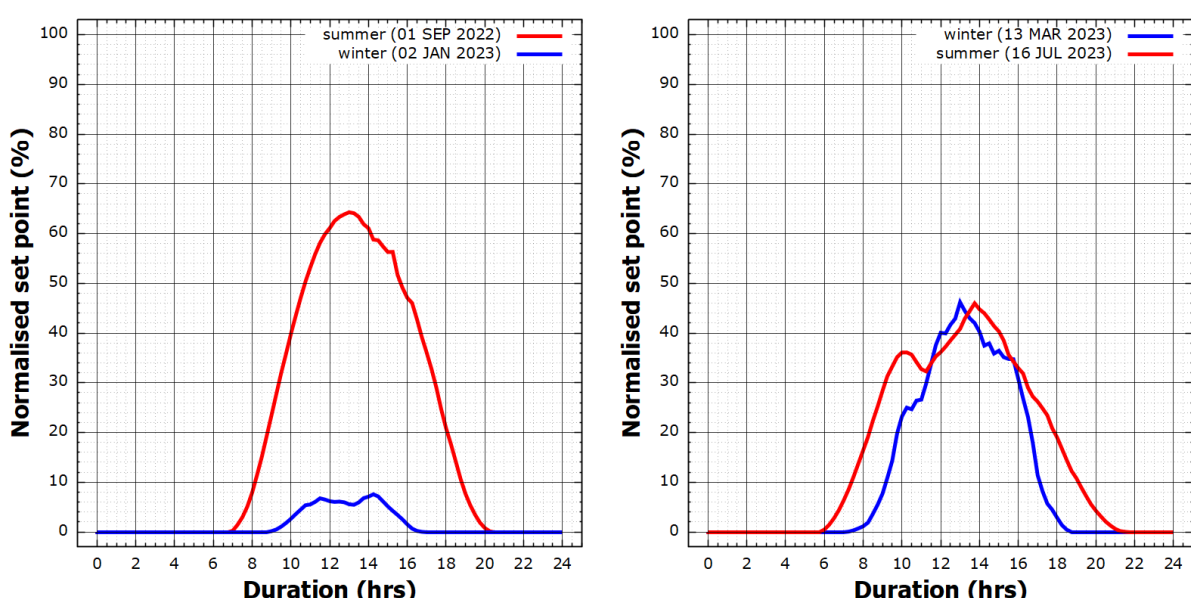
6.8.2 Graphical representation

Figure 8 shows graphical representations of daily operation profiles (normalised set point versus duration) derived from PV electric power for two randomly selected summer and winter weeks in Brussels (Malkow, 2023e). Figure 9 shows the graphical representation of an operation profile for one randomly selected year of PV electric power, which include the daily profiles of the four 7-day periods presented in Figure 8.

Figure 10 shows the graphical representation of daily operation profiles derived from onshore WT electric power in Flanders and offshore WT electric power in Belgium (Malkow, 2023e) for the same four 7-day periods as selected in Figure 8. These profiles may likewise be used as building blocks for a sequence of operation profiles to test a WE stack supplied by WT electric power.

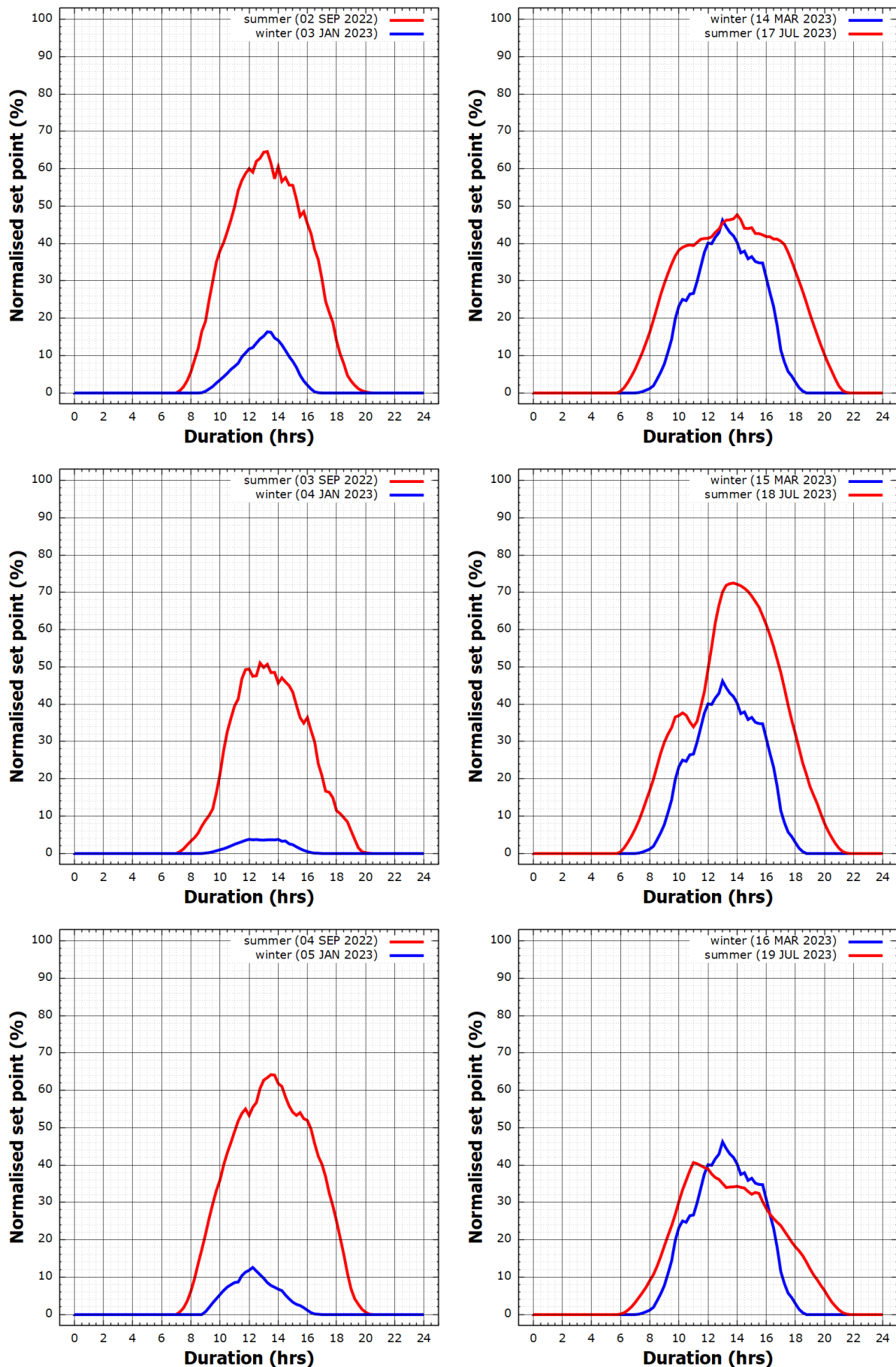
Figure 11 shows the graphical representation of an operation profile for one year of onshore and offshore WT electric power, which include the daily profiles of the four 7-day periods presented in Figure 10.

Figure 8: Daily operation profiles of PV electric power for WE stack testing (Malkow, 2023d).



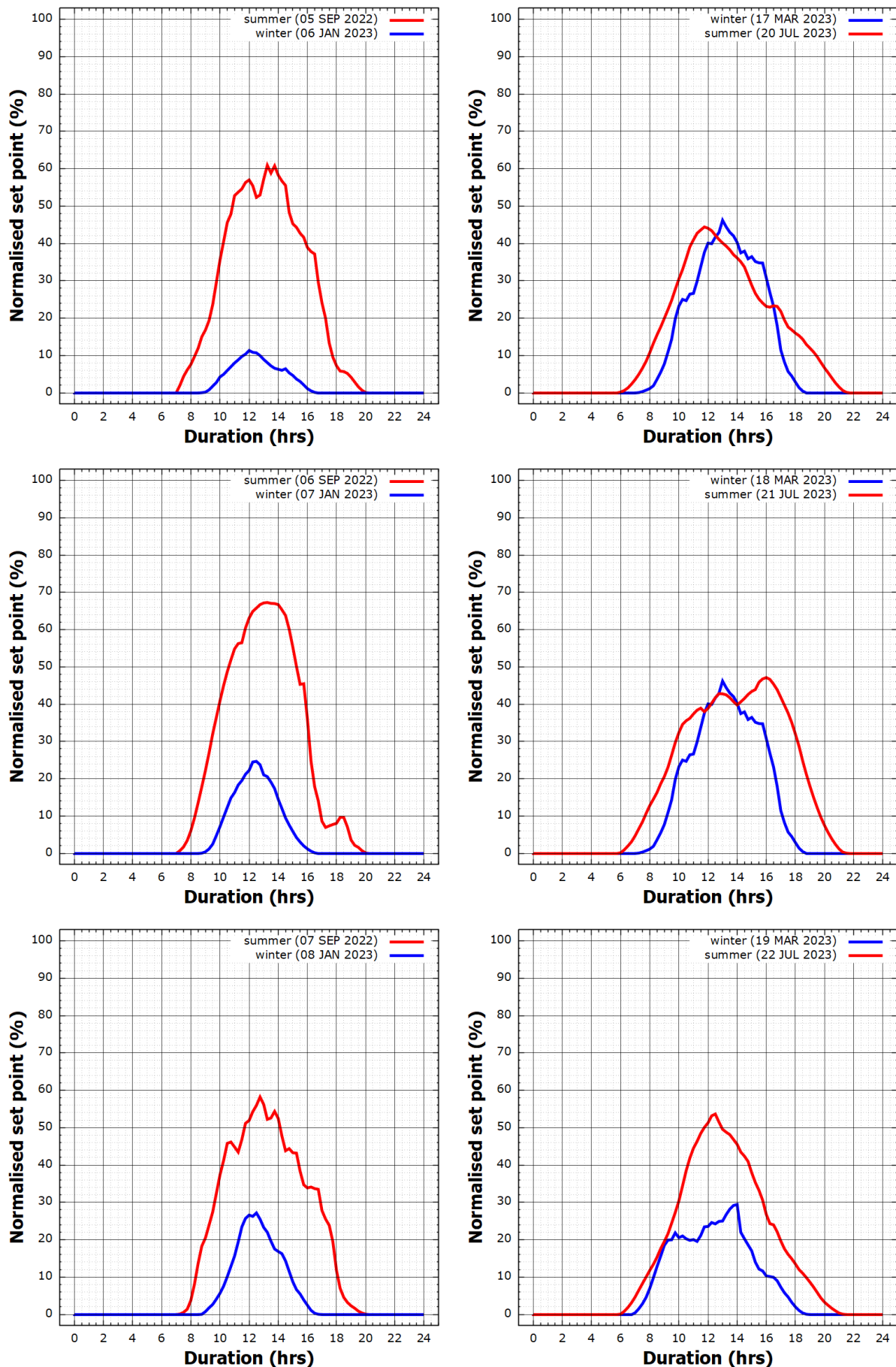
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Figure 8 – continued from previous page



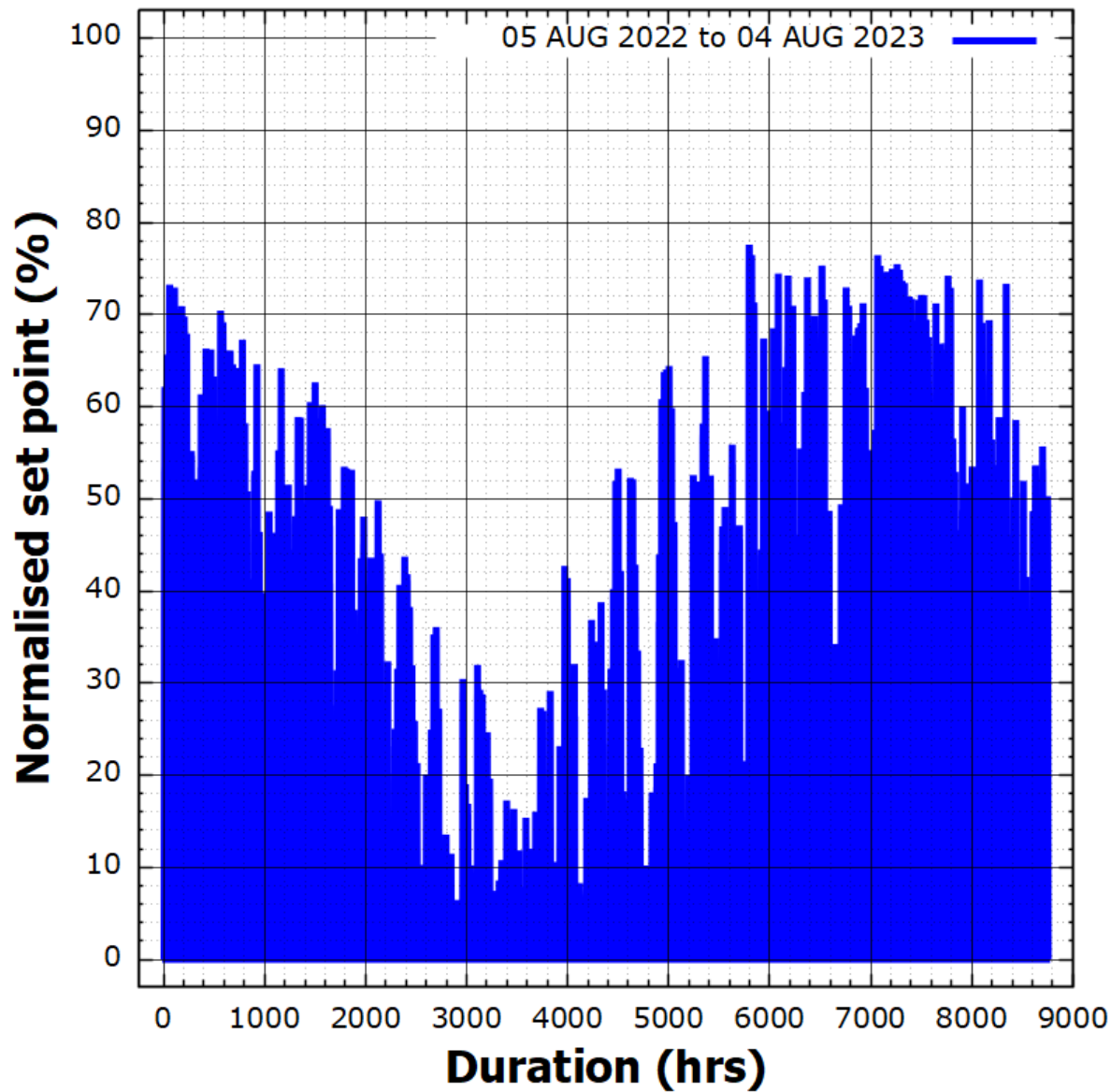
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Figure 8 – continued from previous page



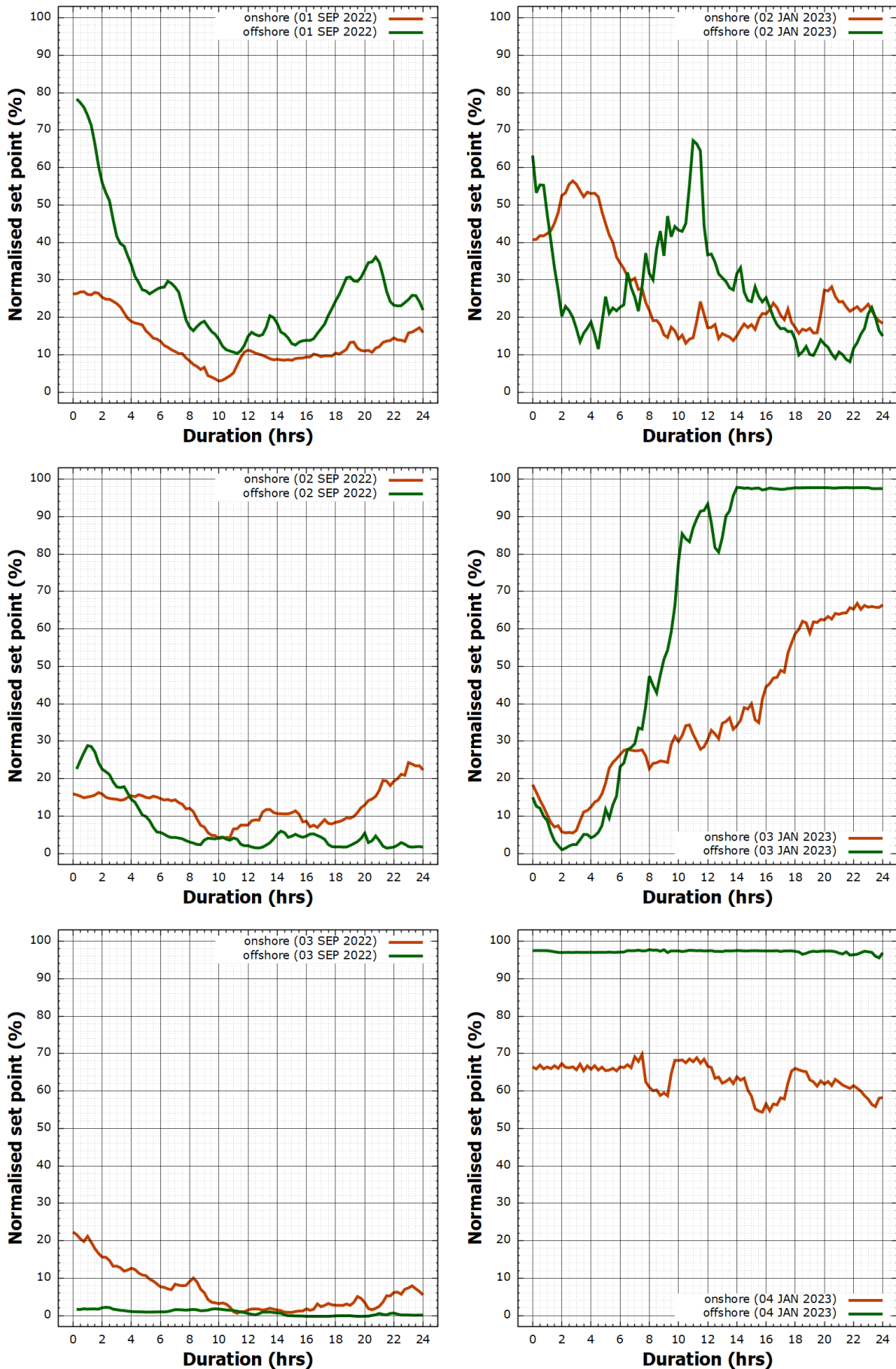
Source: JRC, 2023 (Malkow, 2023a).

Figure 9: Operation profile derived from one year of PV electric power for long-term WE stack testing (Malkow, 2023c). The year, with its start and end dates, is a random selection.



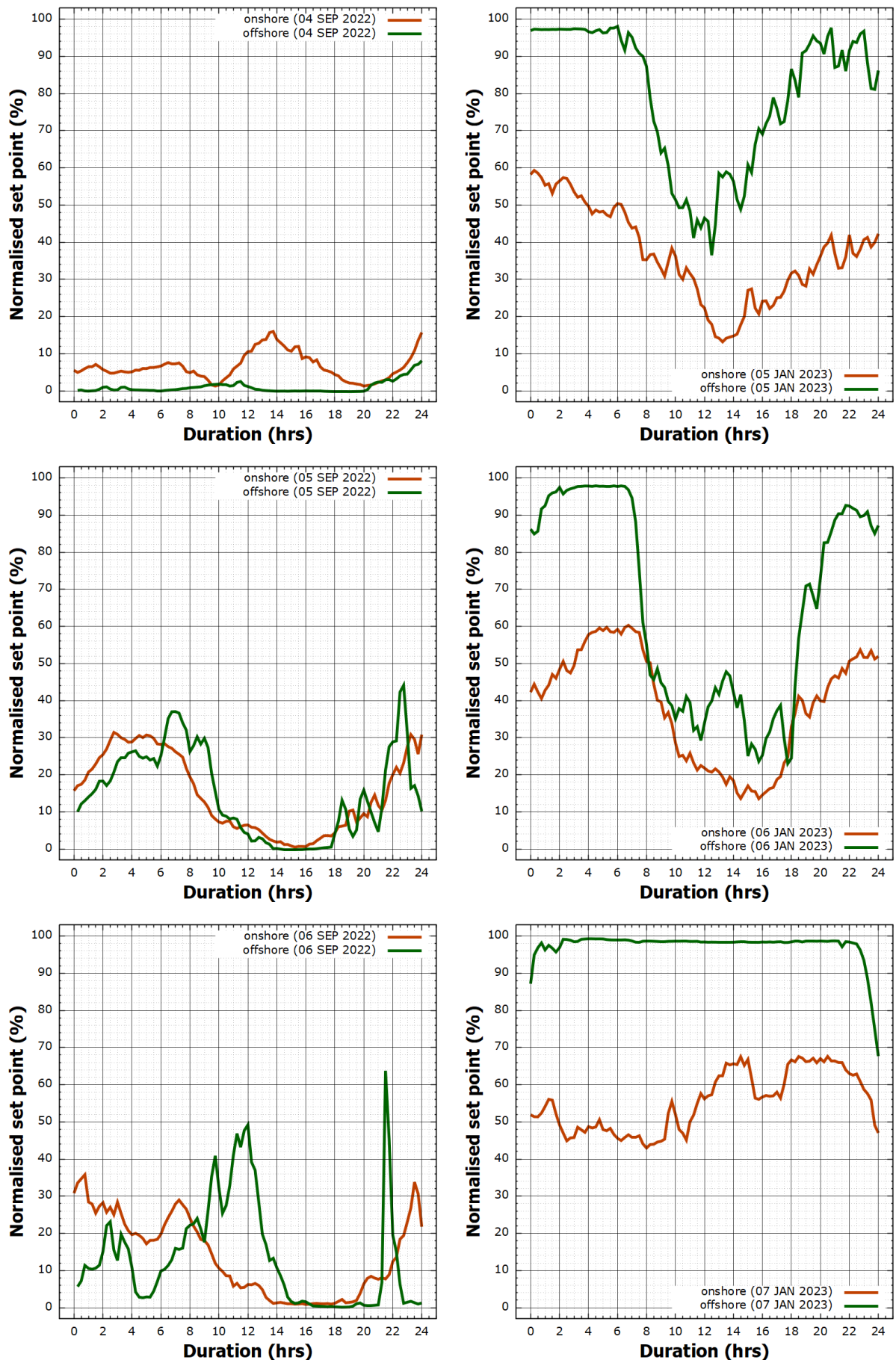
Source: JRC, 2023 (Malkow, 2023a).

Figure 10: Daily operation profiles of onshore/offshore WT electric power for WE stack testing (Malkow, 2023c).



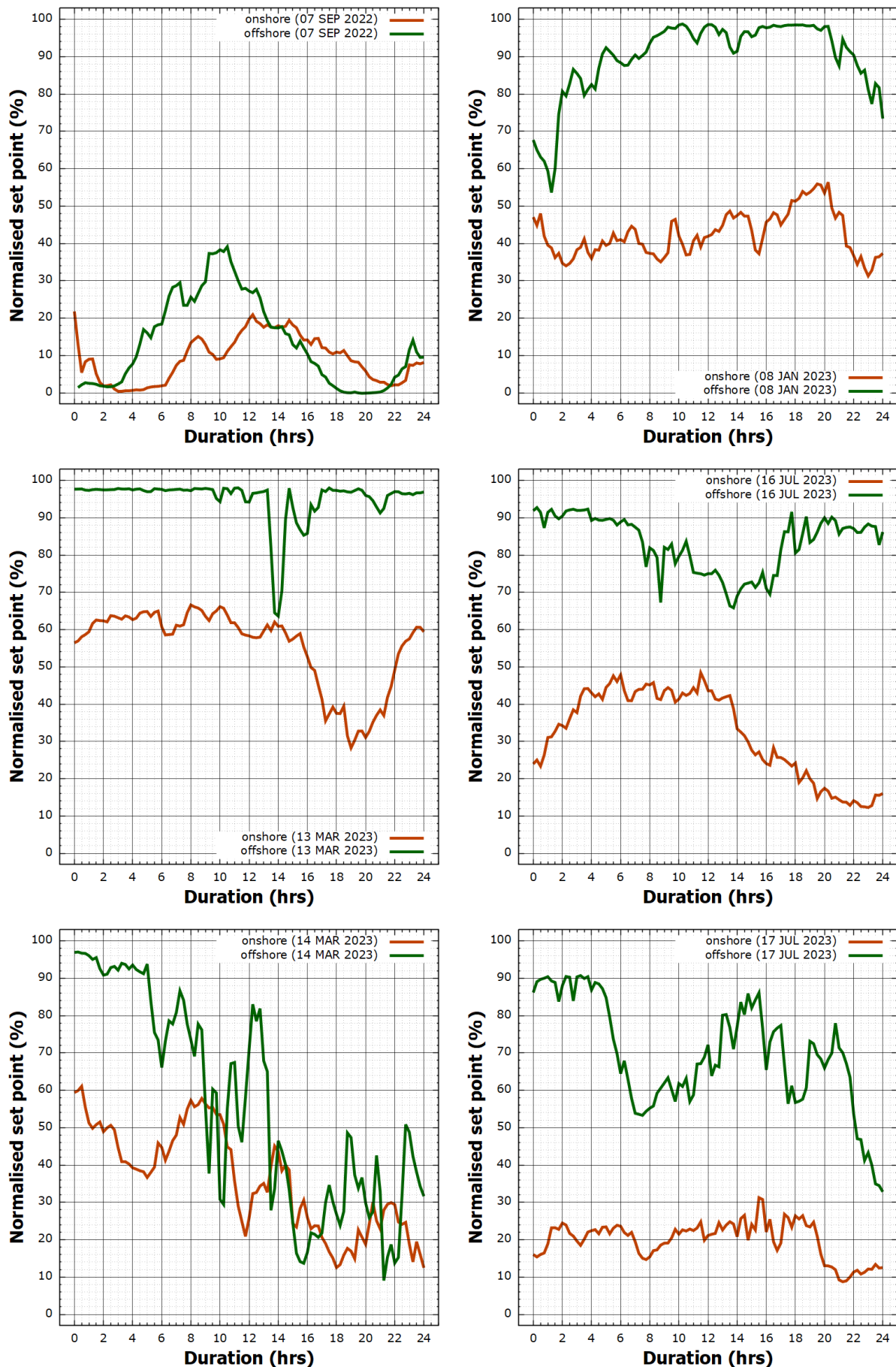
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Figure 10 – continued from previous page



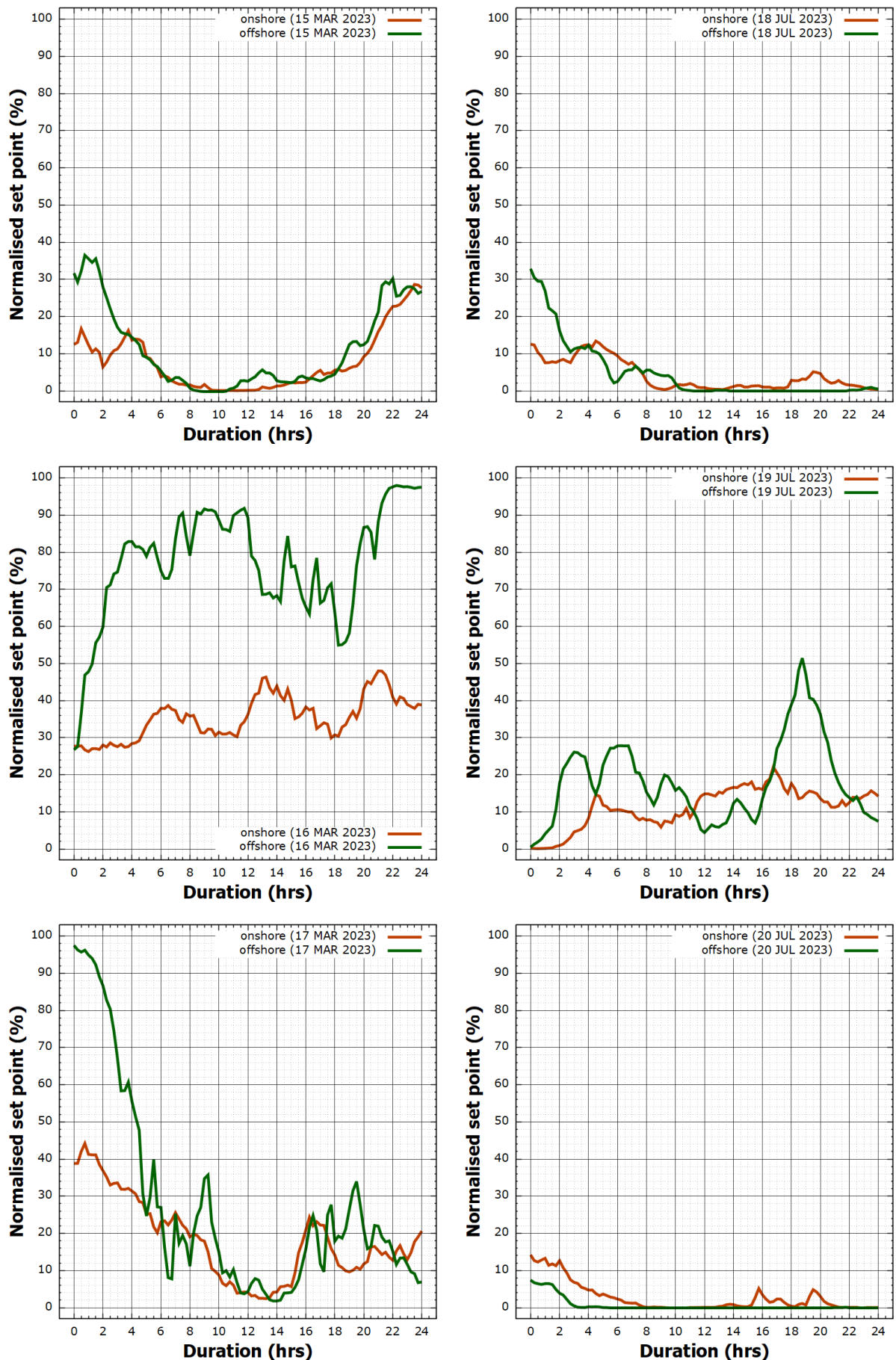
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Figure 10 – continued from previous page



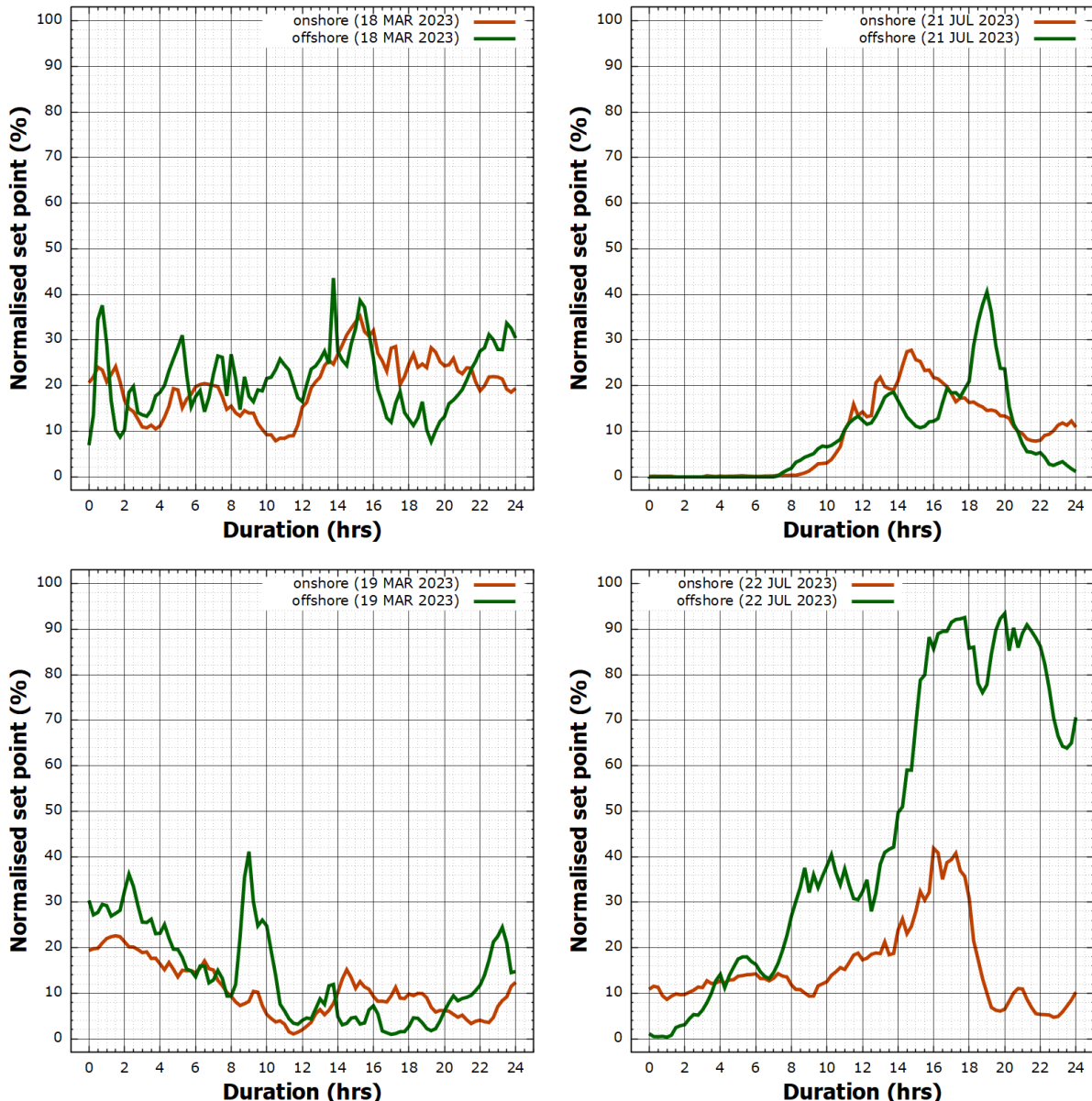
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Figure 10 – continued from previous page



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Figure 10 – continued from previous page



Source: JRC, 2023 (Malkow, 2023b).

In Figure 8 and Figure 11, the normalised set point is either the ratio of the specified input

- electric power to its nominal (rated) value ($P_{el, nom}$) specified by the manufacturer, namely

$$\text{Normalised electric power set point (\%)} = \frac{P_{el,in} (\text{kW})}{P_{el,nom} (\text{kW})} \cdot 100 \%,$$

- current to its nominal (rated) value (I_{nom}) specified by the manufacturer, namely

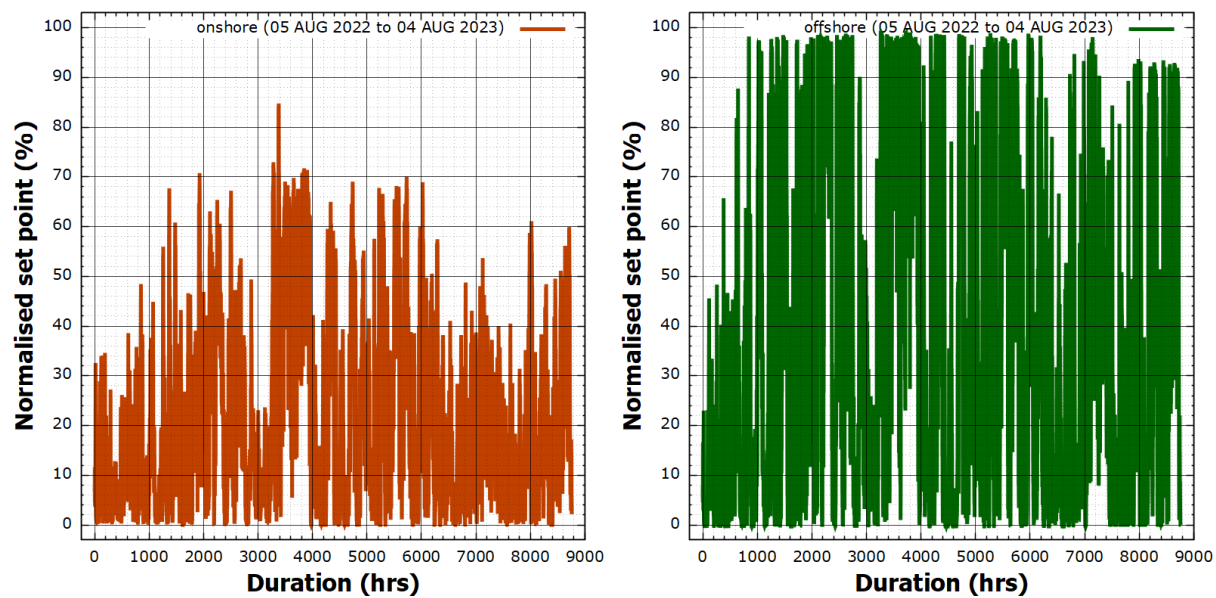
$$\text{Normalised current set point (\%)} = \frac{I_{in} (\text{A})}{I_{nom} (\text{A})} \cdot 100 \% \text{ or}$$

- voltage to its nominal (rated) value (U_{nom}) specified by the manufacturer, namely

$$\text{Normalised voltage set point (\%)} = \frac{U_{in} (\text{kV})}{U_{nom} (\text{kV})} \cdot 100 \%.$$

The operation profiles displayed in Figure 8 to Figure 9 should only be applied where the specification of the WE stack by the manufacturer allows operation at zero input electric power or zero input current. If not, the profiles should be adapted accordingly, for example, by adding a positive offset.

Figure 11: Operation profile derived from one year of onshore and offshore WT electric power for long-term WE stack testing. The start and end dates are the same as in Figure 9.



Source: JRC, 2023 (Malkow, 2023b).

In an AST campaign, the daily operation profiles displayed in Figure 8 and Figure 10 may be combined and used together with appropriate compression factors as building blocks for one or other sequences of operation profiles forming simulated duty cycles to test WE stacks supplied by fluctuating RES-derived electricity stemming from variable PV electric power and/or WT electric power.

Similarly, the yearly operation profiles displayed in Figure 9 and Figure 11 may also be combined using appropriate compression factors for accelerated stress testing of WE stacks. Without any compressed duration, these profiles could also serve in accelerated lifetime testing of WE stacks.

6.9 Durability tests

6.9.1 General

Durability tests (**4.2.29**) on WE stacks evaluate the ability of the stack to maintain its performance characteristics under specified test conditions for a given time interval ⁽⁴²⁾, either at constant 'steady-state' operation (section 6.9.2) or variable operation (section 6.9.3). Combinations of both modes of operation typical for a stack in a given application are possible. Preferably, the original duration of each interval comprises one thousand or more hours of stack operation.

In addition to performance tests at BoT and EoT, performance tests (section 6.7) may, as an option, be conducted intermittently at intervals ($k=1,2,\dots$) to determine one or more KPIs. Importantly, BoT should not be BoL for a stack. The stack should operate for a sufficiently long period recommended by the manufacturer with a minimum of 1000 hours of operation in order to overcome the phase of possible initial high degradation, which is common after operating the stack for the first time.

The inability of a stack to maintain its performance characteristics during durability testing in accordance with a specified stop criterion is considered that the stack has not passed the test.

6.9.2 Constant stack operation

Durability testing of a WE stack under constant current or constant voltage should be conducted in accordance with Clause 7.3 of IEC 62282-8-102:2019 (IEC, 2019b).

Constant stack operation is performed by two similar methods, namely

Method A) Constant current method: Setting the current to its specified value according to the test plan (section 6.6) and maintain it until the stack voltage stabilises to within ± 20 mV for every cell in the stack. If

⁽⁴²⁾ The interval may comprise a specified duration or the time required to complete a given number or sequence of operation profiles (duty cycles).

individual cell voltage measurements are not available, maintain it until the average cell voltage of the stack stabilises to within ± 20 mV. Conduct the test for its specified duration (section 6.6) while recording at least the current, stack voltage, and stack operating temperature.

Method B) Constant voltage method: Setting the stack voltage to its specified value according to the test plan (section 6.6) and maintain it until the current stabilises to within $\pm 2\%$. Conduct the test for its specified duration (section 6.6) while recording at least the current, stack voltage, and stack operating temperature.

6.9.3 Variable stack operation

Durability testing of a WE stack under variable power, current, or voltage using operation profiles (section 6.8) should be conducted in accordance with Clause 7.4 of IEC 62282-8-102:2019 (IEC, 2019b).

The test is carried out by setting the specified test conditions (section 6.3), including stressing operating conditions (section 6.5), to the stack and maintaining these conditions either for a specified duration or until a specified voltage is obtained, according to the test plan (section 6.6), while applying the operation profiles to the stack. Throughout the test, the current, stack voltage, and stack operating temperature shall be recorded as minimum requirements.

6.10 Determination of KPIs

For a given current density (J), stack operating temperature (T_{stack}) and pressure of hydrogen (p_{H_2}), the durability of a WE stack at an elapsed time interval t_k is assessed from the difference (deviation) of the stack voltage at that instant, $U(t_k)$, and the stack voltage at t_0 , $U(t_0)$, by calculating the total rate of change of voltage ($\Delta_{tot}^k U$), whether positive (degradation) or negative (improvement), as follows⁽⁴³⁾

$$\Delta_{tot}^k U \text{ } (\mu\text{V}/\text{h}) = \frac{U(t_k) \text{ (kV)} - U(t_0) \text{ (kV)}}{t_k \text{ (h)} - t_0 \text{ (h)}} \cdot f_{compr} \cdot 10^9 \frac{\mu\text{V}}{\text{kV}}; \quad (6.10.1)$$

f_{compr} is given by equation (4.2.1), and t_k is the time elapsed from BoT at t_0 until the time at the end of interval k, whether for constant stack operation (section 6.9.2) or variable stack operation (section 6.9.3). At both instants, t_0 and t_k , the stack voltages shall be determined from polarisation curve measurements (section 6.7.11) conducted under galvanostatic conditions. In case the stack voltage at t_k exceeds the cut-off voltage of the stack, the decision rules of the test plan shall be followed. The current density of the stack is usually the one when operating the stack under given conditions of temperature and pressure of hydrogen at BoT⁽⁴⁴⁾.

The relative rate of change of voltage ($\Delta_{rel}^k U$), corresponding to a minimum of one thousand hours of stack operation times the compression factor⁽⁴⁵⁾, whether positive (degradation) or negative (improvement), is calculated as follows (McPhail *et al.*, 2022)

$$\Delta_{rel}^k U \text{ } (\%) = \left(\frac{U(t_k) \text{ (kV)}}{U(t_0) \text{ (kV)}} - 1 \right) \cdot \frac{1000 \text{ (h)} \cdot f_{compr}}{t_k \text{ (h)}} \cdot 100 \%. \quad (6.10.2)$$

Multiplying the total rate of change of voltage by the direct current at which the stack voltages, $U(t_k)$ and $U(t_0)$, were determined in polarisation curve measurements, the total rate of change of electric power ($\Delta_{tot}^k P_{el}$) is calculated, whether positive (degradation) or negative (improvement), as follows

$$\Delta_{tot}^k P_{el} \text{ } (\text{mW}/\text{h}) = \frac{P_{el}(t_k) \text{ (kW)} - P_{el}(t_0) \text{ (kW)}}{t_k \text{ (h)} - t_0 \text{ (h)}} \cdot f_{compr} \cdot 10^6 \frac{\text{mW}}{\text{kW}} \quad (6.10.3)$$

where the electric power at t_k and t_0 , $P_{el}(t_k)$ and $P_{el}(t_0)$, are calculated as follows

$$P_{el}(t_k) \text{ (kW)} = U(t_k) \text{ (kV)} \cdot I_{dc}(t_k) \text{ (A)} \text{ and} \quad (6.10.4a)$$

$$P_{el}(t_0) \text{ (kW)} = U(t_0) \text{ (kV)} \cdot I_{dc}(t_0) \text{ (A)}. \quad (6.10.4b)$$

Here, $I_{dc}(t_k)$ and $I_{dc}(t_0)$ represent the direct current at t_k and t_0 , respectively. Notably, a positive total rate of change of electric power implies an increase in nominal DC power of the stack. In principle, the current applied during the polarisation curve measurements at t_k and t_0 should be the same, but measured current values

⁽⁴³⁾ The SRIA 2024 and 2030 targets are 0,11 % and 0,10 % of performance degradation per one thousand hours of AWE operation. These values are 0,9 % and 0,5 % for AEMEL and 0,15 % and 0,12 % for PEMEL (see footnote 27 and Table 2).

⁽⁴⁴⁾ In case the total rate of change of voltage is determined for more than one value of current density, stack operating temperature, or pressure of hydrogen, appropriate indices should be added to $\Delta_{tot}^k U$ and similarly to $\Delta_{rel}^k U$ given by equation (6.10.2), $\Delta_{tot}^k P_{el}$ given by equation (6.10.3), and $\Delta_{rel}^k P_{el}$ given by equation (6.10.5), $\Delta_{qV,H2}^k U$ and $\Delta_{qV,H2}^k U$ given by equation (6.10.6), as well as $\Delta_{qV,H2}^k E_{el}$ and $\Delta_{qV,H2}^k E_{el}$ given by equation (6.10.7).

⁽⁴⁵⁾ The compression factor is unity for accelerated lifetime testing.

may eventually deviate slightly from this assumption. Accordingly, the relative rate of change of electric power ($\Delta_{\text{rel}}^k P_{\text{el}}$) is calculated, whether positive (degradation) or negative (improvement), as follows

$$\Delta_{\text{rel}}^k P_{\text{el}} (\%) = \left(\frac{P_{\text{el}}(t_k) (\text{kW})}{P_{\text{el}}(t_0) (\text{kW})} - 1 \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 \%. \quad (6.10.5)$$

Note, the total rate of change of electric power and the relative rate of change of electric power are useful KPIs when comparing performance degradation of WE stacks determined under potentiostatic conditions as opposed to the recommended galvanostatic conditions.

Importantly, equation (6.10.1) and equation (6.10.2) determine the performance degradation of a WE stack only in terms of absolute and relative voltage deviation, respectively. They do not encompass accompanying changes in the hydrogen output rate, see equation (6.7.1), due to leakage and gas crossover, if any.

Then, the performance degradation of a WE stack should also be assessed from the difference in the stack voltage at t_k normalised by the volumetric flow rate of hydrogen, see equation (6.7.2), at that instant ($q_{V,H_2}(t_k)$) and the stack voltage at t_0 normalised by the volumetric flow rate of hydrogen at the latter instant ($q_{V,H_2}(t_k)$) being the total change of voltage per unit of hydrogen volumetric flow rate ($\Delta_{q_{V,H_2}}^k U$) as well as the difference in the stack voltage at t_k normalised by the mass flow rate of hydrogen, see equation (6.7.4), at that instant ($q_{m,H_2}(t_k)$) and the stack voltage at t_0 normalised by the mass flow rate of hydrogen at the latter instant ($q_{m,H_2}(t_0)$) being the total change of voltage per unit of hydrogen mass flow rate ($\Delta_{q_{m,H_2}}^k U$) calculated, whether positive (degradation) or negative (improvement), as follows (Suermann *et al.*, 2019)

$$\Delta_{q_{V,H_2}}^k U (\mu\text{V}/\text{m}^3/\text{h}) = \left(\frac{U(t_k) (\text{kV})}{q_{V,H_2}(t_k) (\text{m}^3/\text{h})} - \frac{U(t_0) (\text{kV})}{q_{V,H_2}(t_0) (\text{m}^3/\text{h})} \right) \cdot f_{\text{compr}} \cdot 10^9 \frac{\mu\text{V}}{\text{kV}} \text{ and} \quad (6.10.6a)$$

$$\Delta_{q_{m,H_2}}^k U (\mu\text{V}/\text{kg}_{H_2}/\text{h}) = \left(\frac{U(t_k) (\text{kV})}{q_{m,H_2}(t_k) (\text{kg}/\text{h})} - \frac{U(t_0) (\text{kV})}{q_{m,H_2}(t_0) (\text{kg}/\text{h})} \right) \cdot f_{\text{compr}} \cdot 10^9 \frac{\mu\text{V}}{\text{kV}}. \quad (6.10.6b)$$

As for the stack voltages used in equation (6.10.1), both types of hydrogen flow rates (section 6.7.8) are determined as average values simultaneously with their corresponding stack voltages during polarisation curve measurements. Then, the total change of electric energy per unit of volume of hydrogen ($\Delta_{q_{V,H_2}}^k E_{\text{el}}$) and the total change of electric energy per unit of mass of hydrogen ($\Delta_{q_{m,H_2}}^k E_{\text{el}}$) are calculated, whether positive (degradation) or negative (improvement), as follows

$$\Delta_{q_{V,H_2}}^k E_{\text{el}} (\text{mWh}/\text{m}^3_{H_2}) = \left(\frac{P_{\text{el}}(t_k) (\text{kW})}{q_{V,H_2}(t_k) (\text{m}^3/\text{h})} - \frac{P_{\text{el}}(t_0) (\text{kW})}{q_{V,H_2}(t_0) (\text{m}^3/\text{h})} \right) \cdot f_{\text{compr}} \cdot 10^6 \frac{\text{mW}}{\text{kW}} \text{ and} \quad (6.10.7a)$$

$$\Delta_{q_{m,H_2}}^k E_{\text{el}} (\text{mWh}/\text{kg}_{H_2}) = \left(\frac{P_{\text{el}}(t_k) (\text{kW})}{q_{m,H_2}(t_k) (\text{kg}/\text{h})} - \frac{P_{\text{el}}(t_0) (\text{kW})}{q_{m,H_2}(t_0) (\text{kg}/\text{h})} \right) \cdot f_{\text{compr}} \cdot 10^6 \frac{\text{mW}}{\text{kW}}. \quad (6.10.7b)$$

The uniformity of the performance degradation of a WE stack in terms of cell voltages may be assessed by the total rate of change of mean absolute error average cell voltage ($\Delta_{\text{tot}}^k \text{MAE}_{\overline{U}_{\text{cell}}}$), the relative rate of change of mean absolute error average cell voltage ($\Delta_{\text{rel}}^k \text{MAE}_{\overline{U}_{\text{cell}}}$), the total rate of change of standard deviation of average cell voltage ($\Delta_{\text{tot}}^k \text{SD}_{\overline{U}_{\text{cell}}}$), and the relative rate of change of standard deviation of average cell voltage ($\Delta_{\text{rel}}^k \text{SD}_{\overline{U}_{\text{cell}}}$), calculated as follows⁽⁴⁶⁾

$$\Delta_{\text{tot}}^k \text{MAE}_{\overline{U}_{\text{cell}}} (\mu\text{V}/\text{h}) = \frac{\text{MAE}_{\overline{U}_{\text{cell}}}(t_k) (\text{mV}) - \text{MAE}_{\overline{U}_{\text{cell}}}(t_0) (\text{mV})}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}} \cdot 10^3 \frac{\mu\text{V}}{\text{mV}}, \quad (6.10.8a)$$

$$\Delta_{\text{rel}}^k \text{MAE}_{\overline{U}_{\text{cell}}} (\%) = \left(\frac{\overline{U}_{\text{cell}}(t_0) (\text{V}) \cdot \text{MAE}_{\overline{U}_{\text{cell}}}(t_k) (\text{mV})}{\overline{U}_{\text{cell}}(t_k) (\text{V}) \cdot \text{MAE}_{\overline{U}_{\text{cell}}}(t_0) (\text{mV})} - 1 \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 \%, \quad (6.10.8b)$$

$$\Delta_{\text{tot}}^k \text{SD}_{\overline{U}_{\text{cell}}} (\mu\text{V}/\text{h}) = \frac{\text{SD}_{\overline{U}_{\text{cell}}}(t_k) (\text{mV}) - \text{SD}_{\overline{U}_{\text{cell}}}(t_0) (\text{mV})}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}} \cdot 10^3 \frac{\mu\text{V}}{\text{mV}} \text{ and} \quad (6.10.8c)$$

$$\Delta_{\text{rel}}^k \text{SD}_{\overline{U}_{\text{cell}}} (\%) = \left(\frac{\overline{U}_{\text{cell}}(t_0) (\text{V}) \cdot \text{SD}_{\overline{U}_{\text{cell}}}(t_k) (\text{mV})}{\overline{U}_{\text{cell}}(t_k) (\text{V}) \cdot \text{SD}_{\overline{U}_{\text{cell}}}(t_0) (\text{mV})} - 1 \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 \%; \quad (6.10.8d)$$

$\text{MAE}_{\overline{U}_{\text{cell}}}(t_k)$, $\text{SD}_{\overline{U}_{\text{cell}}}(t_k)$ and $\overline{U}_{\text{cell}}(t_k)$ are respectively the mean absolute error of average cell voltage, standard deviation of average cell voltage and average cell voltage at t_k , while $\text{MAE}_{\overline{U}_{\text{cell}}}(t_0)$, $\text{SD}_{\overline{U}_{\text{cell}}}(t_0)$ and $\overline{U}_{\text{cell}}(t_0)$ are

⁽⁴⁶⁾ In case the total rate of change of mean absolute error average cell voltage, the relative rate of change of mean absolute error average cell voltage, the total rate of change of standard deviation of average cell voltage, and the relative rate of change of standard deviation of average cell voltage are determined for more than one value of current density, stack operating temperature, or pressure of hydrogen, appropriate indices should be added to $\Delta_{\text{tot}}^k \text{MAE}_{\overline{U}_{\text{cell}}}$, $\Delta_{\text{rel}}^k \text{MAE}_{\overline{U}_{\text{cell}}}$, $\Delta_{\text{tot}}^k \text{SD}_{\overline{U}_{\text{cell}}}$, and $\Delta_{\text{rel}}^k \text{SD}_{\overline{U}_{\text{cell}}}$ given by equation (6.10.8).

the same quantities but at t_0 . They are calculated from the cell voltages determined during polarisation curve measurements.

Further, the performance degradation of a WE stack for a given current density, stack operating temperature, and pressure of hydrogen may be assessed by the total rate of change of area-specific resistance ($\Delta_{\text{tot}}^k R_{\text{ASR}}$) and the total rate of change of ohmic resistance ($\Delta_{\text{tot}}^k R_\Omega$), whether positive (degradation) or negative (improvement), calculated as follows⁽⁴⁷⁾

$$\Delta_{\text{tot}}^k R_{\text{ASR}} (\text{m}\Omega \cdot \text{cm}^2/\text{h}) = \frac{R_{\text{ASR}}(t_k) (\text{m}\Omega \cdot \text{cm}^2) - R_{\text{ASR}}(t_0) (\text{m}\Omega \cdot \text{cm}^2)}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}} \text{ and} \quad (6.10.9\text{a})$$

$$\Delta_{\text{tot}}^k R_\Omega (\text{m}\Omega/\text{h}) = \frac{R_\Omega(t_k) (\text{m}\Omega) - R_\Omega(t_0) (\text{m}\Omega)}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}}; \quad (6.10.9\text{b})$$

$R_{\text{ASR}}(t_k)$ and $R_{\text{ASR}}(t_0)$ are the area-specific resistances at respectively t_k and t_0 , while $R_\Omega(t_k)$ and $R_\Omega(t_0)$ are the ohmic resistances at these two instants. They are determined from EIS measurements (section 6.7.12) conducted under galvanostatic conditions.

The relative rate of change of area-specific resistance ($\Delta_{\text{rel}}^k R_{\text{ASR}}$) and the relative rate of change of ohmic resistance ($\Delta_{\text{rel}}^k R_\Omega$) corresponding to a minimum of one thousand hours of stack operation times the compression factor, whether positive (degradation) or negative (improvement), are calculated as follows

$$\Delta_{\text{rel}}^k R_{\text{ASR}} (\%) = \left(\frac{R_{\text{ASR}}(t_k) (\text{m}\Omega \cdot \text{cm}^2)}{R_{\text{ASR}}(t_0) (\text{m}\Omega \cdot \text{cm}^2)} - 1 \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 \% \text{ and} \quad (6.10.10\text{a})$$

$$\Delta_{\text{rel}}^k R_\Omega (\%) = \left(\frac{R_\Omega(t_k) (\text{m}\Omega)}{R_\Omega(t_0) (\text{m}\Omega)} - 1 \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 \%. \quad (6.10.10\text{b})$$

For a given input current (input voltage or input electric power) and stack operating temperature, the durability of a WE stack at an elapsed time interval t_k is assessed from the difference of the energy efficiency based on HHV under SATP conditions of hydrogen ($\eta_{\text{HHV},e}^0$), the energy efficiency based on LHV under SATP conditions of hydrogen ($\eta_{\text{LHV},e}^0$), the electrical efficiency based on HHV under SATP conditions of hydrogen ($\eta_{\text{HHV},el}^0$), and the electrical efficiency based on LHV under SATP conditions of hydrogen ($\eta_{\text{LHV},el}^0$) at t_k and t_0 by calculating the total rate of change of energy efficiency based on HHV under SATP conditions of hydrogen ($\Delta_{\text{tot}}^k \eta_{\text{HHV},e}^0$), the total rate of change of energy efficiency based on LHV under SATP conditions of hydrogen ($\Delta_{\text{tot}}^k \eta_{\text{LHV},e}^0$), the total rate of change of electrical efficiency based on HHV under SATP conditions of hydrogen ($\Delta_{\text{tot}}^k \eta_{\text{HHV},el}^0$), and the total rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen ($\Delta_{\text{tot}}^k \eta_{\text{LHV},el}^0$), whether negative (degradation) or positive (improvement), as follows⁽⁴⁸⁾

$$\Delta_{\text{tot}}^k \eta_{\text{HHV},e}^0 (\%/\text{h}) = \frac{\eta_{\text{HHV},e}^0(t_k) (\%) - \eta_{\text{HHV},e}^0(t_0) (\%)}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}}, \quad (6.10.11\text{a})$$

$$\Delta_{\text{tot}}^k \eta_{\text{LHV},e}^0 (\%/\text{h}) = \frac{\eta_{\text{LHV},e}^0(t_k) (\%) - \eta_{\text{LHV},e}^0(t_0) (\%)}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}}, \quad (6.10.11\text{b})$$

$$\Delta_{\text{tot}}^k \eta_{\text{HHV},el}^0 (\%/\text{h}) = \frac{\eta_{\text{HHV},el}^0(t_k) (\%) - \eta_{\text{HHV},el}^0(t_0) (\%)}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}} \text{ and} \quad (6.10.11\text{c})$$

$$\Delta_{\text{tot}}^k \eta_{\text{LHV},el}^0 (\%/\text{h}) = \frac{\eta_{\text{LHV},el}^0(t_k) (\%) - \eta_{\text{LHV},el}^0(t_0) (\%)}{t_k (\text{h}) - t_0 (\text{h})} \cdot f_{\text{compr}}; \quad (6.10.11\text{d})$$

$\eta_{\text{HHV},e}^0(t_k)$, $\eta_{\text{LHV},e}^0(t_k)$, $\eta_{\text{HHV},el}^0(t_k)$ and $\eta_{\text{LHV},el}^0(t_k)$ are respectively the energy efficiency based on HHV under SATP conditions of hydrogen, the energy efficiency based on LHV under SATP conditions of hydrogen, the electrical efficiency based on HHV under SATP conditions of hydrogen, and the electrical efficiency based on LHV under SATP conditions of hydrogen at t_k while $\eta_{\text{HHV},e}^0(t_0)$, $\eta_{\text{LHV},e}^0(t_0)$, $\eta_{\text{HHV},el}^0(t_0)$, and $\eta_{\text{LHV},el}^0(t_0)$ are these efficiencies at t_0 . At both instants, t_k and t_0 , these energy efficiencies are determined in accordance with section 6.7.13 during polarisation curve measurements. The relative rate of change of energy efficiency based on HHV under SATP conditions of hydrogen ($\Delta_{\text{rel}}^k \eta_{\text{HHV},e}^0$), the relative rate of change of energy efficiency based on LHV under SATP conditions of hydrogen ($\Delta_{\text{rel}}^k \eta_{\text{LHV},e}^0$), the relative rate of change of electrical efficiency based on HHV

⁽⁴⁷⁾ In case the total rate of change of area-specific resistance and the total rate of change of ohmic resistance are determined for more than one value of current density, stack operating temperature, or pressure of hydrogen, appropriate indices should be added to $\Delta_{\text{tot}}^k R_{\text{ASR}}$ and $\Delta_{\text{tot}}^k R_\Omega$ given by equation (6.10.9) and similarly to $\Delta_{\text{rel}}^k R_{\text{ASR}}$ and $\Delta_{\text{rel}}^k R_\Omega$ given by equation (6.10.10).

⁽⁴⁸⁾ In case the total rate of change of energy efficiency based on HHV under SATP conditions of hydrogen, the total rate of change of energy efficiency based on LHV under SATP conditions of hydrogen, the total rate of change of electrical efficiency based on HHV under SATP conditions of hydrogen, and the total rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen are determined for more than one value of input current (input voltage, input electric power) or current density, appropriate indices should be added to $\Delta_{\text{tot}}^k \eta_{\text{HHV},e}^0$, $\Delta_{\text{tot}}^k \eta_{\text{LHV},e}^0$, $\Delta_{\text{tot}}^k \eta_{\text{HHV},el}^0$, and $\Delta_{\text{tot}}^k \eta_{\text{LHV},el}^0$ given by equation (6.10.11) and similarly to $\Delta_{\text{rel}}^k \eta_{\text{HHV},e}^0$, $\Delta_{\text{rel}}^k \eta_{\text{LHV},e}^0$, $\Delta_{\text{rel}}^k \eta_{\text{HHV},el}^0$, and $\Delta_{\text{rel}}^k \eta_{\text{LHV},el}^0$ given by equation (6.10.12).

under SATP conditions of hydrogen ($\Delta_{\text{rel}}^k \eta_{\text{HHV}, \text{el}}^0$), and the relative rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen ($\Delta_{\text{rel}}^k \eta_{\text{LHV}, \text{el}}^0$) corresponding to a minimum of one thousand hours of stack operation times the compression factor are calculated, whether negative (degradation) or positive (improvement), as follows

$$\Delta_{\text{rel}}^k \eta_{\text{HHV}, \text{el}}^0 (\%) = \left(\frac{\eta_{\text{HHV}, \text{el}}^0(t_k) (\%) - 1}{\eta_{\text{HHV}, \text{el}}^0(t_0) (\%)} \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 (\%), \quad (6.10.12\text{a})$$

$$\Delta_{\text{rel}}^k \eta_{\text{LHV}, \text{el}}^0 (\%) = \left(\frac{\eta_{\text{LHV}, \text{el}}^0(t_k) (\%) - 1}{\eta_{\text{LHV}, \text{el}}^0(t_0) (\%)} \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 (\%), \quad (6.10.12\text{b})$$

$$\Delta_{\text{rel}}^k \eta_{\text{HHV}, \text{el}}^0 (\%) = \left(\frac{\eta_{\text{HHV}, \text{el}}^0(t_k) (\%) - 1}{\eta_{\text{HHV}, \text{el}}^0(t_0) (\%)} \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 (\%) \text{ and} \quad (6.10.12\text{c})$$

$$\Delta_{\text{rel}}^k \eta_{\text{LHV}, \text{el}}^0 (\%) = \left(\frac{\eta_{\text{LHV}, \text{el}}^0(t_k) (\%) - 1}{\eta_{\text{LHV}, \text{el}}^0(t_0) (\%)} \right) \cdot \frac{1000 (\text{h}) \cdot f_{\text{compr}}}{t_k (\text{h})} \cdot 100 (\%). \quad (6.10.12\text{d})$$

The input direct current (input DC voltage or input electric power) and the stack operating temperature are their rated values given by the stack manufacturer unless otherwise specified in the test plan.

7 Presentation of test results

Table 3 lists the TOPs as results of performance tests (section 6.7) and durability test (section 6.9).

Table 3: Test output parameter as test results

TOP (unit)	Description	Test method
<i>Performance tests</i>		
$P_{\text{el,in}}$ (kW)	input electric power	section 6.7.1
I_{in} (A)	input current	section 6.7.2
U_{in} (kV)	input voltage	section 6.7.3
P_{th} (kW)	thermal power	section 6.7.4
P_{compr} (kW)	power of compression	section 6.7.5
t_{resp} (s)	response time	section 6.7.6
E_{ramp} (kJ/s)	ramp energy	section 6.7.6
pH _{lye}	pH value of lye solution ⁽¹⁾	section 6.7.7
$\sigma_{\text{el,lye}}$ (mS)	electrical conductivity of lye solution ⁽¹⁾	section 6.7.7
pH _w	pH value of liquid water ⁽²⁾	section 6.7.7
$\sigma_{\text{el,w}}$ (mS)	electrical conductivity of liquid water ⁽²⁾	section 6.7.7
q_{n,H_2} (mol/h)	molar flow rate of hydrogen	section 6.7.8
x_{n,H_2} (mol/mol)	molar concentration of hydrogen	section 6.7.8
q_{v,H_2} (mol/h)	volumetric flow rate of hydrogen	section 6.7.8
q_{m,H_2} (mol/h)	mass flow rate of hydrogen	section 6.7.8
q_{n,O_2} (mol/h)	molar flow rate of oxygen	section 6.7.9
x_{n,O_2} (mol/mol)	molar concentration of oxygen	section 6.7.9
c_F ($\mu\text{g/l}$)	fluoride concentration ⁽³⁾	section 6.7.10
c_{PFAS} (ng/l)	PFAS concentration ⁽³⁾	section 6.7.10
c_{PAH} (ng/l)	PAH concentration ⁽²⁾	section 6.7.10
c_{Ca} ($\mu\text{g/l}$)	calcium concentration	section 6.7.10
c_{Cl} ($\mu\text{g/l}$)	chloride concentration	section 6.7.10
c_{Cu} ($\mu\text{g/l}$)	copper concentration	section 6.7.10
c_{Fe} ($\mu\text{g/l}$)	iron concentration	section 6.7.10
c_K ($\mu\text{g/l}$)	potassium concentration	section 6.7.10
c_{Mg} ($\mu\text{g/l}$)	magnesium concentration	section 6.7.10
c_{Na} ($\mu\text{g/l}$)	sodium concentration	section 6.7.10
c_{Ni} ($\mu\text{g/l}$)	section 6.7.10	
I_{stack} (A)	stack current	section 6.7.11
J_{stack} (A/cm ²)	stack current density	section 6.7.11
U_{stack} (kV)	stack voltage	section 6.7.11
$U_{\text{cell,n}}$ (V)	voltage of cell n	section 6.7.11
$\overline{U}_{\text{cell}}$ (V)	average cell voltage	section 6.7.11
MAE $\overline{U}_{\text{cell}}$ (mV)	mean absolute error of average cell voltage	section 6.7.11
SD $\overline{U}_{\text{cell}}$ (mV)	standard deviation of average cell voltage	section 6.7.11
R_{Ω} (Ω)	ohmic resistance	section 6.7.12
R_{ASR} (m $\Omega\cdot\text{cm}^2$)	area-specific resistance ⁽⁴⁾	section 6.7.12
$\eta_{\text{HHV,e}}^0$ (%)	energy efficiency based on HHV under SATP conditions	section 6.7.13
$\eta_{\text{LHV,e}}^0$ (%)	energy efficiency based on LHV under SATP conditions	section 6.7.13
$\eta_{\text{HHV,el}}^0$ (%)	electrical efficiency based on HHV under SATP conditions	section 6.7.13
$\eta_{\text{LHV,el}}^0$ (%)	electrical efficiency based on LHV under SATP conditions	section 6.7.13
<i>Durability tests</i>		
$\Delta_k^k U$ ($\mu\text{V}/\text{h}$)	total rate of change of voltage	section 6.10
$\Delta_{\text{rel}}^k U$ ($\mu\text{V}/\text{h}$)	relative rate of change of voltage	section 6.10
$\Delta_k^k P_{\text{el}}$ (mWh/h)	total rate of change of electric power	section 6.10
$\Delta_{\text{rel}}^k P_{\text{el}}$ (%)	relative rate of change of electric power	section 6.10
$\Delta_k^k q_{v,\text{H}_2} U$ ($\mu\text{V}/\text{m}^3\text{H}_2/\text{h}$)	total change of voltage per unit of hydrogen volumetric flow rate	section 6.10
$\Delta_k^k q_{m,\text{H}_2} U$ ($\mu\text{V}/\text{kgH}_2/\text{h}$)	total change of voltage per unit of hydrogen mass flow rate	section 6.10
$\Delta_k^k E_{\text{el}}$ (mWh/m ³)	total change of electric energy per unit of volume of hydrogen	section 6.10
$\Delta_k^k E_{\text{el}}$ (mWh/kgH ₂)	total change of electric energy per unit of mass of hydrogen	section 6.10

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Table 3 – continued from previous page

$\Delta_{\text{tot}}^k \text{MAE } \overline{U_{\text{cell}}} (\mu\text{V}/\text{h})$	total rate of change of mean absolute error average cell voltage	section 6.10
$\Delta_{\text{rel}}^k \text{MAE } \overline{U_{\text{cell}}} (\%)$	relative rate of change of mean absolute error average cell voltage	section 6.10
$\Delta_{\text{tot}}^k \text{SD } \overline{U_{\text{cell}}} (\mu\text{V}/\text{h})$	total rate of change of standard deviation of average cell voltage	section 6.10
$\Delta_{\text{rel}}^k \text{SD } \overline{U_{\text{cell}}} (\%)$	relative rate of change of standard deviation of average cell voltage	section 6.10
$\Delta_{\text{tot}}^k R_{\text{ASR}} (\text{m}\Omega \cdot \text{cm}^2/\text{h})$	total rate of change of area-specific resistance	section 6.10
$\Delta_{\text{tot}}^k R_{\Omega} (\text{m}\Omega/\text{h})$	total rate of change of ohmic resistance	section 6.10
$\Delta_{\text{rel}}^k R_{\text{ASR}} (\%)$	relative rate of change of area-specific resistance	section 6.10
$\Delta_{\text{rel}}^k R_{\Omega} (\%)$	relative rate of change of ohmic resistance	section 6.10
$\Delta_{\text{tot}}^k \eta_{\text{HHV,e}}^0 (\%/\text{h})$	total rate of change of energy efficiency based on HHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{rel}}^k \eta_{\text{HHV,e}}^0 (\%)$	relative rate of change of energy efficiency based on HHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{tot}}^k \eta_{\text{LHV,e}}^0 (\%/\text{h})$	total rate of change of energy efficiency based on LHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{rel}}^k \eta_{\text{LHV,e}}^0 (\%)$	relative rate of change of energy efficiency based on LHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{tot}}^k \eta_{\text{HHV,el}}^0 (\%/\text{h})$	total rate of change of electrical efficiency based on HHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{rel}}^k \eta_{\text{HHV,el}}^0 (\%)$	relative rate of change of electrical efficiency based on HHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{tot}}^k \eta_{\text{LHV,el}}^0 (\%/\text{h})$	total rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen	section 6.10
$\Delta_{\text{rel}}^k \eta_{\text{LHV,el}}^0 (\%)$	relative rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen	section 6.10

Note: According to the test plan (section 6.6), TOPs may be obtained as functions of TIPs or other TOPs as well as time (test duration), the number of operation profiles, or the sequence(s) of such profiles. By adding appropriate indices to the concerned TOP, TOPs of the same type are distinguished.

⁽¹⁾ Measured for AWE and AAEMWE stacks.

⁽²⁾ Measured for AEMWE and PEMWE stacks.

⁽³⁾ Measured for PEMWE stacks.

⁽⁴⁾ The method of R_{lf} estimation, whether polarisation curve measurements or EIS measurements, shall be stated.

Source: JRC, 2023.

The test results should, as appropriate, be reported along with their standard uncertainties (u) or combined standard uncertainties (u_c) in accordance with the GUM (JCGM, 2008, JCGM, 2009, JCGM, 2020).

In addition to tabulated test results, TOPs of durability tests should be presented graphically, for example, to show their evolution with time or the number and sequence(s) of operation profiles. Their standard uncertainties or combined standard uncertainties should constitute error bars for a specified level of confidence (JCGM, 2008).

8 Conclusions with final remarks

This report proposes testing protocols for accelerated stress testing of low-temperature WE stacks to assess their performance degradation when used in WE systems generating bulk amounts of clean hydrogen using fluctuating electricity from RESs such as PV and WT electric power. These protocols draw upon test methods outlined in ISO and IEC standards, as well as testing procedures developed as part of EU water electrolysis harmonisation activities.

These protocols enable a comprehensive comparison of WEL technologies within stacks. They also facilitate the comparison of performance degradation among different stacks of the same type but varying in design, configuration, and materials used in WECs. Designed for use by both the research community and industry in R&D and stack prototype qualification, these protocols offer inherent flexibility. Users can selectively perform performance tests, and in addition to the RES-based operation profiles provided, they can employ application-oriented duty cycles in durability tests.

Moreover, users have the freedom to substitute one test method for another during a specific campaign if deemed more suitable for the intended application of the WE stack. All tests must be conducted safely (Annex A), with due care taken to record all relevant test parameters, whether TIPs or TOPs, as required. Reported test results (Annex B) should be accompanied by their uncertainties.

While the application of various stress parameters and their combinations to WE stacks during an AST campaign may not guarantee the induction of previously identified degradation phenomena, the wealth of test data collected from accelerated stress testing can be subjected to advanced statistical analysis (IEC, 2010, IEC, 2011, IEC, 2017b) and physics-based modelling especially when a DoE approach was followed. This could reveal performance degradation patterns in WE stacks and establish correlations with applied stress conditions, including acceleration factors.

The utilisation of machine learning (ML) (**4.2.48**) algorithms (Mohamed *et al.*, 2022, Sayed-Ahmed *et al.*, 2024) and other artificial intelligence (AI) (**4.2.12**) techniques (Chavez-Ramirez *et al.*, 2011, Jha *et al.*, 2017, Bahr *et al.*, 2020) on open-access test results holds promise for developing transfer functions expressing the relationship between WE stack performance degradation, applied stress parameters, and test duration reduction. Ultimately, the goal of accelerated stress testing is to minimise R&D costs and expedite the maturation of WE stacks for cost-effective, long-term operation in generating clean hydrogen using renewable electricity.

Finally, it is emphasised that stacks operated in WE systems, along with BoP components under conditions simulating real-world situations, including RES-based intermittent electricity supply, are likely to exhibit different performance degradation with possible intermediary recovery compared to operations in stack-only tests. This is due, not the least, to transitions in system operational states and depends on the actual system design and configuration, including electrolyser modularity and BoP components' responsiveness and performance evolution (Nguyen *et al.*, 2024). Thus, a DoE approach for assessing the performance degradation of WE stacks through an AST campaign applying simulated duty cycles needs to account for these factors.

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List of abbreviations and acronyms

Abbreviation	Description
A/S	Aktieselskab
AAEMEC	alkaline anion exchange polymer membrane electrolysis cell
AAEMWE	alkaline anion exchange polymer membrane water electrolyser
AAS	atomic absorption spectrometry
AB	Aktiebolag
AC	alternating current
AC/DC	AC-to-DC
AEC	alkaline water electrolysis cell
AEL	alkaline water electrolysis
AEM	anion exchange polymer membrane
AEMEC	anion exchange polymer membrane water electrolysis cell
AEMEL	anion exchange polymer membrane water electrolysis
AEMWE	anion exchange polymer membrane water electrolyser
AG	Aktiengesellschaft
AI	artificial intelligence
ALT	accelerated lifetime testing
AMD	amendment
ANIONE	Anion Exchange Membrane Electrolysis for Renewable Hydrogen Production on a Wide-Scale
ANPR	automatic number plate recognition
ARMINES	Association pour la Recherche et le Développement des Méthodes et Processus Industriels
AS	Aktsiaselts
ASA	Allmennaksjeselskap
ASR	area-specific resistance
AST	accelerated stress testing
ATEX	Appareils destinés à être utilisés en atmosphères explosives
AWE	alkaline water electrolyser
AWI	Approved Working Item
biP	bipolar plate
BoL	beginning-of-life
BoP	balance of plant
BoT	beginning-of-test
BPM	bipolar polymer membrane
BPMEL	bipolar polymer membrane water electrolysis

Abbreviation	Description
BPMWE	bipolar polymer membrane water electrolyser
BPMWEC	bipolar polymer membrane water electrolysis cell
BV	besloten vennootschap
BVBA	Besloten Vennootschap met Beperkte Aansprakelijkheid
CAPEX	capital expenditure
CC BY 4.0	Creative Commons Attribution 4.0 International
CEA	Commissariat à l'énergie atomique et aux énergies alternatives
CERTH	Centre for Research & Technology, Hellas
CFA	continuous flow analysis
CGH ₂	compressed gaseous hydrogen
CH	Switzerland
CHFCA	Canadian Hydrogen and Fuel Cell Association
CL	catalyst layer
Clean H ₂ JU	Clean Hydrogen Joint Undertaking
CNLS	complex non-linear least squares
CNR	Consiglio Nazionale delle Ricerche
CNRS	Centre national de la recherche scientifique
CORDIS	Community Research and Development Information Service
CPE	constant phase element
CRM	critical raw materials
CSIC	Consejo Superior de Investigaciones Científicas
CSV	consolidated version
CUA	Customs Union Agreement
DAQ	data acquisition
DC	direct current
DC/DC	DC-to-DC
DER	distributed energy resources
DIS	draft international standard
DLR	Deutsches Zentrum für Luft- und Raumfahrt e. V.
DoE	design of experiment
doi	digital object identifier
DRT	distribution of uncorrelated relaxation times
EC	European Commission
ec	electrochemical cell
ECN	Stichting Energieonderzoek Centrum Nederland

Abbreviation	Description
EDTA	ethylenediaminetetraacetic acid
EEA	European Economic Area
EEC	equivalent electric circuit
EESS	electrical energy storage system
EIS	electrochemical impedance spectroscopy
ELECTROHYPEM	Enhanced performance and cost-effective materials for long-term operation of PEM water electrolyzers coupled to renewable power sources
EMC	electromagnetic compatibility
EN	English
ENDURE	alkaline electrolyzers with ENhanced DURability
ENSMP	École nationale supérieure des mines de Paris
EoT	end-of-test
ES	energy storage
EU	European Union
EUR	European Union Report
EIIIE	Επαρχία Περιορισμένης Ευθύνης
FBK	Fondazione Bruno Kessler
FC	fuel cell
FCH2JU	Fuel Cells and Hydrogen second Joint Undertaking
FES	flame emission spectrometry
FHa	Fundación para el Desarrollo de las Tecnologías del Hidrógeno en Aragón
FIA	flow injection analysis
FID	first industrial deployment
FZJ	Forschungszentrum Jülich GmbH
GC	gas chromatography
GDE	gas diffusion electrode
GDL	gas diffusion layer
GLP	good laboratory practice
GmbH	Gesellschaft mit beschränkter Haftung
GUM	Guide to the Expression of Uncertainty in measurement
H2FUTURE	Hydrogen meeting FUTURE needs of low carbon manufacturing value chains
H ₂ -to-I	hydrogen-to-industry
H ₂ -to-P	hydrogen-to-power
HER	hydrogen evolution reaction
HES	Home Electronic System

Abbreviation	Description
HHV	higher heating value
HPEM2GAS	High Performance PEM Electrolyzer for Cost-effective Grid Balancing Applications
HPLC	high-performance liquid chromatography
HPTLC	high-performance thin-layer liquid chromatography
HRFB	hybrid redox flow battery
IC	ion chromatography
ICT	Institut für Chemische Technologie
IEC	International Electrotechnical Commission
IEEE	Institute of Electrical and Electronics Engineers
IEM	ion exchange membrane
IEV	International Electrotechnical Vocabulary
IFAM	Fraunhofer-Institut für Fertigungstechnik und Angewandte Materialforschung
Inc.	Incorporation
ISBN	international standard book number
ISO	International Organization for Standardization
ISSN	international standard serial number
JCGM	Joint Committee for Guides in Metrology
JRC	Joint Research Centre
KG	Kommanditgesellschaft
KIST	Korea Institute of Science and Technology
KPI	key performance indicator
L	Luxembourg
LC	liquid chromatography
LH ₂	liquid hydrogen
LHV	lower heating value
LT	low-temperature
LTWE	low-temperature water electrolyser
LTWEL	low-temperature water electrolysis
LV	low-voltage
LVD	Low-Voltage Directive
ML	machine learning
MRA	Mutual Recognition Agreement
MS	mass spectrometry
MV	medium-voltage
NC	national committee

Abbreviation	Description
NEPTUNE	Next Generation PEM Electrolyser under New Extremes
NEWELY	Next Generation Alkaline Membrane Water Electrolysers with Improved Components and Materials
NV	naamloze vennootschap
OCP	open circuit potential
OCV	open circuit voltage
OER	oxygen evolution reaction
OHS	occupational health and safety
OJ	Official Journal
OÜ	Osaübung
P-to-C	power-to-chemical
P-to-F	power-to-fuel
P-to-G	power-to-gas
P-to-H ₂	power-to-hydrogen
P-to-L	power-to-liquid
P-to-M	power-to-mobility
P-to-X	power-to-X
PAH	polycyclic aromatic hydrocarbons
PDF	portable document format
PED	Pressure Equipment Directive
PEM	proton exchange membrane or polymer electrolyte membrane
PEMEC	proton exchange membrane or polymer electrolyte membrane water electrolysis cell
PEMEL	proton exchange membrane or polymer electrolyte membrane water electrolysis
PEMWE	proton exchange membrane or polymer electrolyte membrane water electrolyser
PFAS	perfluoroalkyl and polyfluoroalkyl substances
PFC	per- and polyfluorinated compounds
PFSA	perfluoro sulfonic acid
PGM	platinum-group metals
plc	public limited company
PoC	point of connection
PPS	polyphenylene sulfide
PRETZEL	Novel modular stack design for high pressure PEM water electrolyzer technology with wide operation range and reduced cost
PROMETH2	Cost-effective PROton Exchange MEmbrane WaTer Electrolyser for Efficient and Sustainable Power-to-H ₂ Technology
PTL	porous transport layer
PV	photovoltaic

Abbreviation	Description
R&D	research and development
R&I	research and innovation
REACH	registration, evaluation, authorisation and restriction of chemicals
REF	renewable energy factor
RES	renewable energy source
RFB	redox flow battery
RLV	redline version
RTO	research and technology organisation
RUL	remaining useful life
S.L.	Sociedad Limitada
SA	Société anonyme
SARL	Société à responsabilité limitée
SAS	Société par actions simplifiée
SATP	standard ambient temperature and pressure
SCADA	supervisory control and data acquisition
SHE	standard hydrogen electrode
SI	Système International d'Unités
SINTEF	Stiftelsen for industriell og teknisk forskning
SoA	state-of-the-art
SpA	Società per azioni
SRIA	strategic research and innovation agenda 2021-2027 of the Clean Hydrogen Partnership for Europe
TC	Technical Committee
TEU	Treaty on European Union
TFEU	Treaty on the Functioning of the European Union
TIP	test input parameter
TNO	Nederlandse Organisatie voor Toegepast Natuurwetenschappelijk Onderzoek
TOP	test output parameter
TR	Technical Report
TRL	technology readiness level
TS	Technical Specification
UG	Unternehmergeellschaft
UQTR	Université du Québec à Trois-Rivières
URL	uniform resource locator
UV	ultraviolet

Abbreviation	Description
VRE	variable renewable energy
VSCHT	Vysoká Škola chemicko-technologická v Praze
WE	water electrolyser
WE system	water electrolyser system
WEC	water electrolysis cell
WEL	water electrolysis
WG	Working Group
WT	wind turbine
ZSW	Zentrum für Sonnenenergie- und Wasserstoff-Forschung Baden-Württemberg

List of symbols

Symbol	Description
(aq)	subscript denoting aqueous phase
(ed)	subscript denoting electrode
(g)	subscript denoting gaseous phase
(l)	subscript denoting liquid phase
$ Y $	modulus of electrical admittance
$ Z $	modulus of electrical impedance
A_{act}	active electrode area
$a_{\text{H}_2\text{O}}$	activity of liquid water
α	power exponent
α_a	anodic charge transfer coefficient
α_c	cathodic charge transfer coefficient
$\arg(Y)$	argument of electrical admittance
$\arg(Z)$	argument of electrical impedance
C	capacitance
C	carbon
$c_{\text{H}_2}^0$	equilibrium hydrogen concentration
$c_{\text{O}_2}^0$	equilibrium oxygen concentration
Ca	calcium
c_{Ca}	calcium concentration
c_{Cl}	chloride concentration
c_{Cu}	copper concentration
$C_{\text{dl},a}$	anodic double-layer capacitance
$C_{\text{dl},c}$	cathodic double-layer capacitance
c_F	fluoride concentration
c_{PAH}	PAH concentration
c_{PFAS}	PFAS concentration
c_{Fe}	iron concentration
c_{H_2}	hydrogen concentration
c_K	potassium concentration
Cl^-	chloride
c_{Mg}	magnesium concentration
c_{Na}	sodium concentration
c_{Ni}	nickel concentration
Co	cobalt
CO_2	carbon dioxide
c_{O_2}	oxygen concentration
CO_3	carbonate
c_p	specific heat capacity at constant pressure
c_p^i	specific heat capacity at constant pressure of fluid i
c_p^j	specific heat capacity at constant pressure of fluid j
Cu	copper
c_v^j	specific heat capacity at constant volume of fluid j
c_{Zn}	zinc concentration
d_{dl}	double-layer length
$\Delta_{q_{m,\text{H}_2}}^k E_{\text{el}}$	total change of electric energy per unit of mass of hydrogen
$\Delta_{q_{m,\text{H}_2}}^k U$	total change of voltage per unit of hydrogen mass flow rate
$\Delta_{q_{v,\text{H}_2}}^k E_{\text{el}}$	total change of electric energy per unit of volume of hydrogen
$\Delta_{q_{v,\text{H}_2}}^k U$	total change of voltage per unit of hydrogen volumetric flow rate
$\Delta_{\text{rel}}^k \eta_{\text{HHV,e}}^0$	relative rate of change of energy efficiency based on HHV under SATP conditions of hydrogen
$\Delta_{\text{rel}}^k \eta_{\text{HHV,el}}^0$	relative rate of change of electrical efficiency based on HHV under SATP conditions of hydrogen
$\Delta_{\text{rel}}^k \eta_{\text{LHV,e}}^0$	relative rate of change of energy efficiency based on LHV under SATP conditions of hydrogen
$\Delta_{\text{rel}}^k \eta_{\text{LHV,el}}^0$	relative rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen
$\Delta_{\text{rel}}^k \text{MAE}_{\overline{U}_{\text{cell}}}$	relative rate of change of mean absolute error average cell voltage
$\Delta_{\text{rel}}^k P_{\text{el}}$	relative rate of change of electric power
$\Delta_{\text{rel}}^k R_{\text{ASR}}$	relative rate of change of area-specific resistance

Symbol	Description
$\Delta_{\text{rel}}^k R_\Omega$	relative rate of change of ohmic resistance
$\Delta_{\text{rel}} \text{SD } \overline{U}_{\text{cell}}$	relative rate of change of standard deviation of average cell voltage
$\Delta_{\text{rel}}^k U$	relative rate of change of voltage
$\Delta_{\text{tot}}^k \eta_{\text{HHV,e}}^0$	total rate of change of energy efficiency based on HHV under SATP conditions of hydrogen
$\Delta_{\text{tot}}^k \eta_{\text{HHV,el}}^0$	total rate of change of electrical efficiency based on HHV under SATP conditions of hydrogen
$\Delta_{\text{tot}}^k \eta_{\text{LHV,e}}^0$	total rate of change of energy efficiency based on LHV under SATP conditions of hydrogen
$\Delta_{\text{tot}}^k \eta_{\text{LHV,el}}^0$	total rate of change of electrical efficiency based on LHV under SATP conditions of hydrogen
$\Delta_{\text{tot}}^k \text{MAE } \overline{U}_{\text{cell}}$	total rate of change of mean absolute error average cell voltage
$\Delta_{\text{tot}}^k P_{\text{el}}$	total rate of change of electric power
$\Delta_{\text{tot}}^k R_{\text{ASR}}$	total rate of change of area-specific resistance
$\Delta_{\text{tot}}^k R_\Omega$	total rate of change of ohmic resistance
$\Delta_{\text{tot}}^k \text{SD } \overline{U}_{\text{cell}}$	total rate of change of standard deviation of average cell voltage
$\Delta_{\text{tot}}^k U$	total rate of change of voltage
E	energy
e^-	electron
E_{compr}	pneumatic energy
E_{el}	electric energy
ε	absolute permittivity
E_{ramp}	ramp energy
U_{rev}	voltage under reversible (equilibrium) conditions
η	efficiency
η_e	energy efficiency
$\eta_{\text{HHV,e}}^0$	energy efficiency based on HHV under SATP conditions of hydrogen
η_{el}	electrical efficiency
$\eta_{\text{HHV,el}}^0$	electrical efficiency based on HHV under SATP conditions of hydrogen
$\eta_{\text{LHV,e}}^0$	energy efficiency based on LHV under SATP conditions of hydrogen
$\eta_{\text{LHV,el}}^0$	electrical efficiency based on LHV under SATP conditions of hydrogen
η_F	Faradaic efficiency
E_{th}	thermal energy
F	Faraday constant
F	fluorine
f	frequency
F^-	fluoride
f_{compr}	compression factor
Fe	iron
f_{max}	maximum frequency
f_{min}	minimum frequency
γ_j	isentropic expansion factor of fluid j
H	hydrogen
H^+	proton
H_2	molecular hydrogen or dihydrogen
H_2O	steam
H_3O^+	hydrated proton
HHV_{H_2}	higher heating value of hydrogen
I	current
i	imaginary unit
$I_{0,a}$	anodic exchange current
$I_{0,c}$	cathodic exchange current
I_a	anodic current
I_c	cathodic current
I_{dc}	direct current
I_{in}	input current
I_{max}	maximum current
$\Im Z$	imaginary part of electrical impedance
I_{nom}	nominal (rated) current
Ir	iridium
IrO_x	iridium oxide
I_{stack}	stack current
J	current density

Symbol	Description
J_{stack}	stack current density
K	potassium
KCl	potassium chloride
KOH	potassium hydroxide
L	inductance
LHV_{H_2}	lower heating value of hydrogen
$\text{MAE}_{\overline{U}_{\text{cell}}}$	mean absolute error average cell voltage
Mg	magnesium
m_{H_2}	molar mass of hydrogen
Na	sodium
N_{cell}	number of cells
Ni	
O	oxygen
O_2	molecular oxygen or dioxygen
OH^-	hydroxide ion
ω	angular frequency
P	power
p	pressure
p^0	standard ambient pressure
P_{compr}	power of compression
$P_{\text{compr,in}}$	input power of compression
P_{el}	electric power
$P_{\text{el,d}}$	electric power density
$P_{\text{el,dc}}$	DC power
$P_{\text{el,in}}$	input electric power
$P_{\text{el,nom}}$	nominal (rated) electric power
p_{H_2}	pressure of hydrogen
p_{H_2}	partial pressure of hydrogen
p_{H_2}	pressure of hydrogen
pH_{lye}	pH value of lye solution
pH_w	pH value of liquid water
p_j	pressure of fluid j
p_{O_2}	partial pressure of oxygen
P_{stack}	stack power
Pt	platinum
P_{th}	thermal power
$P_{\text{th,in}}$	input thermal power
q	flow rate
Q_c	non-ideal capacitance
Q_L	non-ideal inductance
q_m	mass flow rate
q_{m,H_2}	mass flow rate of hydrogen
q_m^i	mass flow rate of fluid i
q_n	molar flow rate
q_{n,H_2}	molar flow rate of hydrogen
q_n^j	molar flow rate of fluid j
q_{n,O_2}	molar flow rate of oxygen
$q_{n,\text{out}}$	product gas molar flow rate
$q_{V,\text{H}_2}^{\text{theo}}$	theoretical volumetric flow rate of hydrogen
q_{V,H_2}	volumetric flow rate of hydrogen
R	resistance
R_0	zero-frequency resistance
R_{ASR}	area-specific resistance
R_{ct}	charge transfer resistance
$R_{\text{ct,a}}$	anodic charge transfer resistance
$R_{\text{ct,c}}$	cathodic charge transfer resistance
$\Re e Z$	real part of electrical impedance
R_g	universal gas constant
R_{hf}	high-frequency resistance

Symbol	Description
R_∞	infinite-frequency resistance
R_{lf}	low-frequency resistance
R_Ω	ohmic resistance
R_{pol}	polarisation resistance
RuO_y	ruthenium oxide
$\text{SD } \overline{U_{\text{cell}}}$	standard deviation of average cell voltage
T	temperature
t	time
T^0	standard ambient temperature
t_0	time at beginning-of-test
$\tan(\Im m Z / \Re e Z)$	tangent of the loss angle of electrical impedance
t_{compr}	duration of compressed profile of operation
T_{H_2}	temperature of hydrogen
T^i	temperature of fluid i
Ti	titanium
TiO_x	titanium oxide
t_k	time at interval k
$T_{\text{in}}^{\text{lye}}$	input temperature of lye
t_{origin}	duration of the original operation profile
t_{resp}	response time
T_{stack}	stack operating temperature
T_w	water temperature
T_{in}^w	input water temperature
U	voltage
u	standard uncertainty
U_{act}	activation polarisation voltage
u_c	combined standard uncertainty
U_{cell}	cell voltage
$\overline{U}_{\text{cell}}$	average cell voltage
$U_{\text{cell,n}}$	voltage of cell n
U_{conc}	concentration polarisation voltage
$U_{\text{cut-off}}$	cut-off voltage
U_{dc}	DC voltage
U_{in}	input voltage
U_{nom}	nominal (rated) voltage
U_{OCP}	open circuit potential
U_{OCV}	open circuit voltage
U_{rev}	reversible voltage
U_{stack}	stack voltage
U_{tn}	thermal-neutral voltage
U_{WEC}	water electrolysis cell voltage
V	volume
$V_{\text{m,H}_2}$	molar volume of hydrogen
X	reactance
$x_{\text{n,H}_2}$	molar concentration of hydrogen
$x_{\text{n,O}_2}$	molar concentration of oxygen
Y	electrical admittance
Y^*	complex conjugate of electrical admittance
Z	electrical impedance
z	number of electrons exchanged
Z^*	complex conjugate of electrical impedance
\bar{Z}	average compressibility factor
\bar{Z}^j	average compressibility factor of fluid j
Zn	zinc
σ_{el}	electrical conductivity
$\sigma_{\text{el,lye}}$	electrical conductivity of lye solution
$\sigma_{\text{el,w}}$	electrical conductivity of liquid water
τ	time constant

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Annex A Test safety

In LTWE stacks, hazards arises from various factors including

- the generation of hydrogen and oxygen gases,
- the use of alkaline solution,
- temperature,
- pressure and
- voltage.

Throughout the phases of installation, commissioning, operation, quiescence, maintenance, and decommissioning, ensuring the safety of personnel requires diligence and caution from all involved parties. Entities conducting testing and chemical analysis must adhere to the occupational health and safety (OHS) requirements outlined in ISO 45001:2018 (ISO, 2018) and follow good laboratory practice (GLP). Tests conducted on WE stacks should comply with applicable legislation, granted licenses, and issued permits to prevent harm or unacceptable risk to humans, property, and the environment.

Guidance on the safety of electrical equipment (IEC, 2021b, IEC, 2018) and alkaline ES devices (IEC, 1996) has been published by the IEC, while ISO published guidance on the safety of hydrogen systems (ISO, 2015) (49). These guidances should be observed when testing WE stacks (50). Standards related to FC safety issued by the IEC (IEC, 2019a, IEC, 2020b) may be applied by analogy. Additionally, IEC guidance on the classification of areas where explosive atmospheres can occur (IEC, 2014a, IEC, 2013, IEC, 2017a, IEC, 2020a), should be followed.

In the European Economic Area (EEA) (51), directives such as the Appareils destinés à être utilisés en atmosphères explosives (ATEX) Directives 2014/34/EU (EP and Council, 2014b) and 94/9/EC (EP and Council, 1994) apply, along with other EU legislation including the electromagnetic compatibility (EMC) Directive 2014/30/EU (EP and Council, 2014a), the Low-Voltage Directive (LVD) 2014/35/EU (EP and Council, 2014c), the general product safety Directive 2001/95/EC (EP and Council, 2001), the machinery Directive 2006/42/EC (EP and Council, 2006), and the Pressure Equipment Directive (PED) 2014/68/EU (EP and Council, 2014d) (52). In principle, test items that do not conform to EU legislation shall not be used within the EEA.

(49) WG 29 of TC 197 is currently reviewing ISO/TR 15916:2015.

(50) Note that ISO 22734:2019 Hydrogen generators using water electrolysis — Industrial, commercial, and residential applications (ISO, 2019a) specifies safety-related aspects of WE. Additionally, WG 34 of TC 197 issued recently ISO/DIS 22734-1:2024 Hydrogen generators using water electrolysis - Part 1: General requirements, test protocols and safety requirements (ISO, 2024), which contains clauses on safety among others, to replace ISO 22734:2019.

(51) The EEA comprises the EU territory as defined by Article 52 of the Treaty on European Union (TEU) and Article 355 of the Treaty on the Functioning of the European Union (TFEU), Iceland, Norway and Liechtenstein. It also applies to Switzerland under a Mutual Recognition Agreement (MRA) and Türkiye under a Customs Union Agreement (CUA) with the EU.

(52) The European Commission publishes guidance online for various directives and regulations, including:

- ATEX Directive (https://single-market-economy.ec.europa.eu/single-market/european-standards/harmonised-standards/equipment-explosive-atmospheres-atex_en),
- EMC Directive (https://single-market-economy.ec.europa.eu/sectors/electrical-and-electronic-engineering-industries-eei/electromagnetic-compatibility-emc-directive_en),
- LVD (https://single-market-economy.ec.europa.eu/sectors/electrical-and-electronic-engineering-industries-eei/low-voltage-directive-lvd_en),
- General product safety Directive (https://single-market-economy.ec.europa.eu/single-market/european-standards/harmonised-standards/general-product-safety_en),
- Machinery Directive (https://single-market-economy.ec.europa.eu/sectors/mechanical-engineering/machinery_en), and
- PED (https://single-market-economy.ec.europa.eu/sectors/pressure-equipment-and-gas-appliances/pressure-equipment-sector/pressure-equipment-directive_en).

Annex B Test report

B.1 General

The test report shall accurately, clearly, and objectively present all relevant information to demonstrate the purpose(s) and objective(s) of the test. As a minimum requirement, the test report shall contain a title page (section B.2) and a summary (section B.3) with the measured or calculated TIPs and TOPs at least as mean values along with their (combined) standard uncertainties, whether absolute, relative, or both. The test plan (section 6.6) forms part of the report. Calibration records or certificates of the measuring instruments shall be documented in the report and shall be available upon request.

B.2 Title page

The titlepage(s) shall present the following information:

- (a) Report identification, *i. e.* report number (optional),
- (b) Type of report (summary, detailed or full),
- (c) A reference to this document,
- (d) Author(s) of the report,
- (e) Entity issuing the report with name and address,
- (f) Date of the report,
- (g) Person(s) conducting the test when different from the reporting author(s),
- (h) Organisation conducting the test when different from report issuing entity,
- (i) Date and time per test run,
- (j) Location per test run when different from the address of the report issuing entity,
- (k) Descriptive name per test, and
- (l) Identification (model name, serial number, type and specification) of the WE stack tested (including manufacturer).

The titlepage(s) may be followed by a contents page before the summary report.

B.3 Summary

The summary shall include the following information:

- (i) test purpose(s) and objective(s),
- (ii) description of the test(s) with sufficient information on the test conduct and measurement set-up, including test methods, measurement techniques (section 6.2), and test conditions (section 6.3),
- (iii) all relevant test parameters, namely TIPs and TOPs with uncertainties (section 7), and
- (iv) conclusion(s) with remark(s) and/or observation(s) as appropriate. Unless a full test report is to be issued, where all test results shall also be presented graphically (section 7) and properly discussed, a brief discussion with graphical presentation of the main test results (section 7) supporting the conclusion(s) may be appended to the report.

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