Review

Review of the Durability of Polymer Electrolyte Membrane Fuel Cell in Long-Term Operation: Main Influencing Parameters and Testing Protocols

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Abstract: Durability is the most pressing issue preventing the efficient commercialization of polymer electrolyte membrane fuel cell (PEMFC) stationary and transportation applications. A big barrier to overcoming the durability limitations is gaining a better understanding of failure modes for user profiles. In addition, durability test protocols for determining the lifetime of PEMFCs are important factors in the development of the technology. These methods are designed to gather enough data about the cell/stack to understand its efficiency and durability without causing it to fail. They also provide some indication of the cell/stack’s age in terms of changes in performance over time. Based on a study of the literature, the fundamental factors influencing PEMFC long-term durability and the durability test protocols for both PEMFC stationary and transportation applications were discussed and outlined in depth in this review. This brief analysis should provide engineers and researchers with a fast overview as well as a useful toolbox for investigating PEMFC durability issues.

Keywords: fuel cells; PEM; degradation; durability; test protocol; hydrogen

1. Introduction

Clean, energy-efficient fuel cells are appealing power sources that can help solve major energy production and consumption issues. Polymer electrolyte membrane fuel cells (PEMFCs) have been considered as electric power sources for cars, as well as stationary and portable power sources, due to their high energy efficiency, ease of operation, and environmental friendliness [1,2]. Furthermore, as a promising power source, integrating PEMFCs into microgrids, which are a common structure in the smart grid framework, has been gaining traction around the world, encouraging the usage of hydrogen energy. Over the last few decades, significant change has been made. However, the most significant remaining problem impeding the effective commercialization of a wide range of PEMFC stationary and transportation energy applications is durability [3,4]. To ensure high efficiency and durability, fuel cells should meet lifetime goals and operate under a variety of operating conditions. This will help to validate fuel cells as a cost-effective alternative energy source [3,5]. The European Union (EU)-funded project STAYERS (Stationary PEM fuel cells with lifetimes beyond five years) addressed the cost-effective use of PEMFC power for stationary applications, noting that these cells must have a lifetime greater than 40,000 h or at least five years of continuous operation to compete with current power generation systems such as microturbines and allow for an acceptable return on investment.
to the end-user [1,6,7]. The Fuel Cell Technical Team (FCTT) Roadmap from 2017 outlined 
the targets and efficiency expectations for fuel cells in automotive applications [8]. The 
FCTT has set a durability target of 8000 hours (equivalent to 150,000 miles) with less than 
10% performance loss for the full range of automotive operating conditions from –40 °C 
to 40 °C, as well as to adapt to rapid changes in power demand needed in automotive 
applications [8]. This target for durability is comparable to the standards of existing in-
ternal combustion engines on the market. However, according to an FCTT analysis, the 
highest fuel cell vehicle lifetime is currently about 4100 h (2016) with 10% stack voltage 
degradation [8]. This estimate is close to the 5000 h goal for 2020, but well short of the 
8000 h target for 2025. To speed up widespread commercialization, more intensive R&D 
is required to resolve PEMFC durability or degradation. In addition, test protocols and 
methods must be used to determine the fuel cell lifespan.

The strategy for addressing durability entails defining the factors that influence the 
lifespan of a PEMFC and devising methods for minimizing negative effects [8]. Platinum-
deposition and sintering, carbon-support corrosion, membrane thinning, and 
bipolar plate corrosion are most factors that may shorten the lifespan of PEMFCs, according 
to researchers. Operating environments and conditions may also have an effect on efficiency 
and longevity. Impurities in the fuel and oxidant sources, subfreezing temperatures, fuel 
starvation, and transient operations during startup and shutdown are indeed critical 
operating modes that affect durability [2]. The influences of these factors were summarized 
by Xiao-Zi Yuan et al. [1]. Impurities such as CO, H2S, SOx, NOx, and others are known 
to cause kinetic changes because of contamination of both the anode and cathode catalyst 
sites; mass transfer effects as a result of structural change and permeability of the electrolyte 
layers and gas diffusion layers; and conductivity effects caused by rising membrane and 
imonomer resistance. Significant reductions in the cathode electrochemical surface area 
occurred as the PEMFC was launched at subfreezing temperatures. If the anode is turned 
off for a long time, the hydrogen will ultimately flow to the cathode, filling the anode 
flow channels with air. Because fuel cells must operate in a wide range of operating and 
cycling environments, including temperatures from below zero to above the boiling point 
of water, relative humidity from ambient to saturated, and half-cell potentials from 0 to 
>1.5 V, fuel cell failure is unavoidable [1]. Transitioning from low to high power can induce 
changes in operational factors, including temperature and relative humidity, putting the 
fuel cell system’s integrity at risk. Temperature and relative humidity have been shown 
to impact the rate of catalyst surface area loss owing to platinum particle growth. The 
imonomer swells as the water content in the membrane increases, resulting in tensile residual 
stresses during drying and contributing to membrane mechanical failures. The rate of 
catalyst surface area loss related to platinum particle accumulation has been found to 
be affected by temperature and relative humidity. In some cases, especially in transitory 
settings, the operation of a fuel cell can accelerate known decay modes or even activate 
totally new decay mechanisms [2]. Although the lifetime requirement for automotive 
applications is shorter than for stationary applications, operating conditions like as fast 
cycling, repetitive stops and starts, and freezing/thawing make the present technology’s 
longevity goal more difficult to accomplish [1,7]. Obtaining a deeper understanding of 
failure modes for user profiles is a major roadblock to overcoming the durability limitations. 
If appropriate, a better knowledge base will contribute to the advancement of mitigation 
and/or performance recovery strategies [7].

Different methods for assessing the durability of PEMFCs have been developed in 
the United States, the European Union, Japan, and China. These methods are designed to 
collect enough data about the cell/stack in order to understand its efficiency and durability 
before it fails. They also have a method for determining the cell/age, and stacks related 
to changes in cell/stack performance over time [9]. According to the US Department of 
Energy, a metric is used until the cell/stack either accumulates a certain number of cycles 
or fails to meet a performance criterion, such as a cumulative 20% decline in power or a 
10% loss of average cell voltage from initial voltage under cycling conditions [7,9,10].
An effective durability test can provide PEMFC lifetime data, which can be used to evaluate and analyze PEMFC lifetime degradation using lifetime degradation models based on the data. Durability tests can also serve as a solid foundation for testing the degradation behaviors of various components in PEMFCs. The numerous durability tests available vary greatly. Long-term durability tests are time-consuming and expensive, and they often produce lifetime results that are too late to monitor the advancement of fuel cell technology [4].

Methods to evaluate the durability of PEMFC cell/stacks for automotive applications have been suggested by institutions and organizations in India, South Korea, and Russia, and more remarkable in the United States, the European Union, Japan, and China [11]. In particular, the Dynamic Stress Test (DST) of the U.S. Department of Energy (DOE, USA) and the Fuel Cell and Hydrogen Energy Association (FCHEA), the Fuel Cell Technical Team (FCTT) of the U.S. Driving Research and Innovation for Vehicle Efficiency and Energy Sustainability (USDRIVE, USA) Fuel Cell Technical Team, the New European Driving Cycle (NEDC) of the United Nations Economic Commission for Europe (UNECE, EU), the Fuel Cell Testing and Standardization Network (FCTESTNEST) of the International Electrotechnical Commission (IEC, EU) and Fuel Cell Testing and Standardization Network (FC TestNet), the Giantleap of the Fuel Cell and Hydrogen Joint Undertaking, and the JC08 of the Japanese New Energy and Industrial Technology Development Organization (NEDO, Japan), as well as other protocols of Chinese universities, such as those at Tongji University, Tsinghua University, Wuhan University of Technology, and Dalian Institute of Chemical Physics, have been developed.

From multi-megawatt systems for large-scale power generation to small units (e.g., 1 kW) for backup power or micro-combined heat and power (CHP) systems, stationary fuel cells can be used in a wide variety of commercial, manufacturing, and residential applications [7]. Since the basic construction and materials of the PEMFC stack for stationary applications are similar to those used in vehicles, durability issues are similar [12]. PEMFCs in stationary applications, on the other hand, have much longer lifetimes than those in automobiles. Although the operating conditions of PEMFCs in static applications are normally milder than in vehicles, the 40,000 h lifetime goal cannot be easily tested in actual operation [6,12,13]. Testing protocols for stationary applications have received less development attention than testing protocols for vehicle applications.

The aim of this paper is to compile existing data on the key factors influencing PEMFC durability and the various durability test protocols. We hope to provide practitioners with a short overview as well as a useful toolkit for researching PEMFC durability issues. To begin, we will go through the fundamentals and how they affect long-term durability. After that, we will go through the durability testing protocols that have been developed for both vehicle and stationary applications. This summary is not intended to be an exhaustive list of protocols produced by universities and research organizations in America (US), the European Union (EU), Japan, and China, but rather a reference for reviewing and comparing them. The findings would make a significant contribution to the creation of brand-new PEMFC single cell/stack durability test protocols.

2. Main Parameters Influencing Long-Term Durability of Polymer Electrolyte Membrane Fuel Cells (PEMFCs)

This section will provide a quick overview of the main limiting factors that will inevitably have a significant effect on PEMFC long-term durability. First, the phenomenon of degradation and potential mechanisms of critical PEMFC components such as catalyst layers, membranes, gas diffusion layers, bipolar plates, and sealing gaskets will be thoroughly discussed. Following that, the significant impact of operational effects (contamination, subfreezing, and other contributing factors) will be properly addressed, accompanied by thermal management and operating conditions.
2.1. Degradation of Different Components of PEMFC

Degradation or corrosion of prominent features in PEMFCs, such as catalyst layers, membranes, gas diffusion layers, flow-field channels, and seals, are fundamental mechanisms that are detrimental to fuel cell lifespan and one of the most important challenges to PEMFC commercialization. The durability and degradation of key components vary greatly depending on the fuel cell’s current and operating conditions [5]. This article emphasizes the leading causes and mechanisms of the degradation of critical PEMFC components.

2.1.1. Catalyst Layer

Structure and Role

A catalyst layer (CL) is a porous structure made up of a high surface area carbon support that has been carefully filled with nanoscale platinum catalyst particles in an ionomer matrix [1,3]. The carbon support structure helps nanoparticles to maintain high dispersion (2–3 nm) while also providing CLs with a porous electronically conductive structure. This arrangement allows reactants and electrons to flow more easily to the Pt nanoparticles while also removing gases and water. The ionomer, which clusters sulfonated side chains, ensures that the hydrophilic regions are properly maintained in a generally hydrophobic substance, allowing reactants and protons to freely enter active catalyst sites within the CLs [14–17]. Because of their multiple phases, porosity, and electrochemical reactions, CLs are often the thinnest layers in the fuel cell, but they are often also the most complex.

Degradation Phenomena and Mechanisms

Platinum (Pt) or platinum alloy nanoparticles have been the most common catalysts for both the anode and cathode of PEMFCs [16]. The Pt nanoparticles are typically supported by a black powder carbon surface, which serves as the electrode’s mechanical foundation. Catalyst degradation and carbon support corrosion are the two most common phenomena that cause significant electrode performance loss [17]. These two extraordinary phenomena are linked but not always mutually exclusive [18]. Carbon corrosion is catalyzed by a Pt-based catalyst. As a result, the number of sites available for anchoring Pt particles on the carbon support surface is reduced significantly. Consequently, the catalyst’s redistribution and sintering is that the catalyst degrades, and the electrode’s structural integrity deteriorates [15,19].

The basic electrode structures in most modern PEMFC designs are virtually identical, and they are the same for both the anode and the cathode; however, the degradation impacts on both electrodes are not identical [17]. According to Borup R.L. et al. [20], the most substantial corrosion of catalyst is observed at the cathode. During constant voltage, constant current, and drive cycle tests, as well as offline cycling measurement conditions, anode phenomena are negligible.

Catalyst degradation refers to the loss of Pt particles on the carbon support surface, as well as changes in their structure and distribution [5,20]. Catalyst degradation is currently known to occur in many ways, including contamination-induced catalyst toxicity, mechanical deformation-induced cracks, catalyst wash out, carbon coarsening, and electrolyte degradation. Pt dissolution is the primary cause of catalyst corrosion among these factors [21]. A phenomenon called Ostwald ripening occurs when small Pt particles dissolve in an acidic operating atmosphere and then redeposit on the surface of larger particles. Pt migration occurs as a result of Pt dissolution. Pt species that are dissolved in water diffuse into the ionomer phase and precipitate in the membrane [1,18].

Dissolution of Pt is accelerated by voltage cycling, which gives rise to a varying load on the fuel cell; in essence, stop-and-go driving experiences of a fuel cell vehicle can typically generate high voltage load [22]. Wang et al. [23] reported that dissolved Pt concentration increased when potentials rose from 0.65 to 1.1 V and then decreased at
potentials above 1.1 V. In another publication, Pt dissolution took place as potentials that exceeded 0.9 V because electrochemical oxidation of Pt occurred, and Pt oxide instantly formed on the Pt surface [24].

b Carbon support corrosion

Another primary cause of CL degradation in PEM fuel cells is corrosion of the catalyst support, which results in obvious loss of active catalyst surface area and catalyst connection, as well as incrementally lowers the porosity and hydrophobicity of the CL, resulting in overall cell performance reduction [14]. Principally, electrochemical carbon corrosion may occur according to the following reaction [14,18,21]:

\[
\text{C} + 2\text{H}_2\text{O} \rightarrow \text{CO}_2 + 4\text{H}^+ + 4\text{e}^- , \quad E_o = 0.207 \text{ V}_{\text{RHE}} \text{ at } 25^\circ\text{C} \tag{1}
\]

Carbon oxidation can occur by heterogeneous water-gas reaction [21,22]:

\[
\text{C} + \text{H}_2\text{O} \rightarrow \text{CO} + 2\text{H}^+ + 2\text{e}^- , \quad E_o = 0.518 \text{ V}_{\text{RHE}} \text{ at } 25^\circ\text{C} \tag{2}
\]

where RHE denotes the reversible hydrogen electrode.

The corrosion rate of carbon in a normal PEMFC at typical operating temperatures (60–90°C) is reasonably slow at potentials lower than 0.9 V [22,25] or 1.1 V [1] due to the slow kinetics. The carbon content of CL will decrease over time due to the sluggish kinetics of the carbon oxidation reaction (COR). Furthermore, the regular presence of Pt at lower potentials may help to promote COR [2,18]. Carbon corrosion is normally insignificant during steady-state operation, but it becomes more problematic with frequent start-stop cycles [1,15,18]. When a fuel supply is cut off, the hydrogen in the anode side is replaced by air almost immediately, resulting in a potential drop at the anode and a gradual increase in the temporary potential at the cathode to 1.5 V. Carbon corrosion and Pt dissolution can be accelerated by the high potential at the cathode [3,15]. High potentials can cause severe damage to the CL in systems that require regular start-up and shutdown, such as automotive applications.

2.1.2. Membrane Degradation

Structure and Role

The proton exchange membrane (PEM) is the heart of the PEMFC, designed to transport hydrogen ions, i.e., protons, from the anode to the cathode without moving electrons. The PEM also serves as a separator reactant (hydrogen and oxygen gas) in the fuel cell’s anode and cathode. PEMs should have excellent proton conductivity and thermal and chemical stability, strong mechanical resistance, exceptional flexibility and low permeability to gases, and low water drag to achieve these particular functions [17,26]. According to [5,19,26], perfluorinated sulfonic acid (PFSA) membranes such as Nafion (DuPont TM), Aciplex, Flemion (Asahi TM), and Gore-Select (Gore TM) are the most popular and commercially available PEM materials in PEMFCs. Among these membranes, Nafion has been regarded as the industry standard for PEMFCs [27].

The membrane in Nafion is an ionomer, which means that the polytetrafluoroethylene (PTFE) polymer structure has been changed by sulfonation to give it an ionic character. Owing to the strong water attraction of the added sulfonic acid groups, the resulting polymer structure contains hydrophilic regions and is efficiently created within a typically hydrophobic material (−SO₃H) [27]. When Nafion is well hydrated, this ionomer structure provides sufficient proton conductivity, allowing protons to freely move inside the material. Up to a certain limiting value, the ionic conductivity of Nafion is proportional to the degree of hydration [17,27].

Degradation Phenomena and Mechanisms

Membrane durability has a profound effect on the operating lifetime of PEMFCs [28]. Membrane degradation results in hydrogen leaks and, eventually, cell failure [29]. The
membrane degradation rate of the anode side is higher than that of the anode side [30]. The membrane degrades first on the anode side, according to G. De Moor et al. After that, when the anode side has lost about 70% of its initial thickness, hydrogen crossover accelerates cathode degradation [31]. Mechanical and chemical degradation are the two most common interactive mechanisms that cause membrane failure [32].

Early failure of PEMs during fuel cell operation is often caused by mechanical membrane degradation. Crack, puncture, blister, and pinhole formation caused by the membrane, CLs, and membrane electrode assembly (MEA) fabrication, as well as variations in mechanical stress in the membrane, are the most common causes of failure [33,34]. The primary cause of mechanical failure in cracking and tearing of the membrane is alternating compressive/tensile stresses. The membrane usually appears to be in-plane during fuel cell activity, a location generally caused by the significant swelling that occurs with increasing RH and stretching after shrinkage in a low RH state [34]. Furthermore, non-uniform mechanical stresses in-plane can be caused by the interface between the MEA frame and the channels in the bipolar plate. This non-uniform stress distribution causes defects like pinholes, cracks, and delamination, which can spread widely across the membrane and eventually lead to reactant crossover. These mechanical stresses accelerate membrane failure by spreading defects and increasing chemical membrane degradation [35]. Gas crossover intensifies and accelerates membrane degradation when a possible break or perforation occurs in the membrane [31].

Membrane failure is also attributed to chemical attack [36,37]. As several papers have shown, chemical attacks cause severe membrane thinning and fluoride release [38–45]. The chemical degradation of the membrane is caused by the complex formation of peroxy- and hydroperoxy-radicals. PEM conductivity is reduced as a result of these radicals attacking the backbone and side chains directly (a process known as side chain unzipping) [46]. General Electric (GE) accurately describes the most widely accepted process for the typical formation of peroxy species in PEM, which involves the five steps mentioned below [33]:

Step 1: \( \text{H}_2 \rightarrow 2\text{H}^\bullet \) (via Pt catalyst)
Step 2: \( \text{H}^\bullet + \text{O}_2 \) (diffused through PEM) \( \rightarrow \text{HO}_2^\bullet \)
Step 3: \( \text{HO}_2^\bullet + \text{H} \rightarrow \text{H}_2\text{O}_2 \) (which can diffuse into PEM at locus near degraded front of PEM)
Step 4: \( \text{H}_2\text{O}_2 + \text{M}^{2+} \) (Fe\(^{2+}\), Cu\(^{2+}\), … found in MEAs…) \( \rightarrow \text{M}^{3+} + \text{OH}^\bullet + \text{OH}^- \)
Step 5: \( \text{OH}^\bullet + \text{H}_2\text{O}_2 \rightarrow \text{H}_2\text{O} + \text{HO}_2^\bullet \) (hydrogen peroxide radical attacks the PEM).

2.1.3. Gas Diffusion Layers (GDLs)
Structure and Role
Sandwiched between the CL and flow field, the gas diffusion layer (GDL) is a porous electrically and thermally conductive carbon-based layer. GDLs consist of two layers: a macro-porous substrate (MPS) made of carbon fibers and a microporous layer (MPL) made of carbon powder and a hydrophobic agent (PTFE) [1,47,48].

By controlling mass, heat, and electron transport, GDLs play a critical role in PEM fuel cell operation [47]. Furthermore, the GDLs are reliably strong enough to prevent MEA from stretching due to water absorption. GDLs, on the other hand, will reduce the efficiency and lifetime of other components by clogging pores and preventing gas diffusion from the flow channels to the catalyst sites [49,50].

Degradation Phenomena and Mechanisms
Several GDL degradation mechanisms have been proposed, including mechanical and physicochemical degradation. The first category includes physical and mechanical damage caused by clamping friction, freezing and thawing in automotive applications, and erosion caused by high-velocity gas flow [48,50,51]. Due to breakage and delamination of carbon layers under high pressure, these contributing factors cause GDL deformation and thickness changes, resulting in a significant reduction in the hydrophobicity of immersed GDL [52]. The primary cause of GDL degradation in the second group is carbon corrosion/oxidation.
caused by the formation of water droplets during such severe operating conditions, such as start-up, shutdown, and fuel starvation [48]. Despite the fact that carbon fibers in GDLs and carbon black particles in MPLs are more stable than carbon particles in CLs, carbon oxidation by water cannot be ignored as a determining factor in GDL hydrophobicity loss and porosity reduction, resulting in gas diffusion being hindered due to high gas transport resistance as a result of the increased water content in the GDLs and MPL, and the fuel cell’s efficiency being lowered [1,53].

2.1.4. Bipolar Plate
Structure and Role

The flow-field plate, also known as the bipolar plate (BPP), is a multifunctional feature in the PEMFC and stack [19]. It gathers and conducts current from the anode to the cathode, distributes reactant and product streams uniformly with a flow-field, and separates the reactants and liquid coolant. Furthermore, the BPPs serve as direct links, allowing multiple cells to be connected in series to form PEMFC stacks. BPPs are available in carbon (non-metallic) and metallic materials. These materials must meet a number of specific criteria, including good electrical conductivity, high thermal conductivity, high resistance to chemical attack and corrosion, and high mechanical stability, especially under compression; low gas permeability; and low density [17,19]. While specific materials like graphite and graphite composites have these properties, their shock and vibration resistance, hydrogen permeability, and manufacturability are inferior to metals like stainless steel, particularly in vehicular applications [19,54]. Many review papers associated with bipolar plates have been released in recent years [55–59].

Degradation Phenomena and Mechanisms

BPP degradation adversely impacts the performance and durability of PEMFCs. Christoph Hartnig et al. [60] reported how bipolar plate degradation powerfully affects cell performance. Two main aspects of BPP degradation effects in the cell can be observed in this comprehensive research: (1) an increase in ohmic cell resistance and (2) an increase in oxygen reduction overpotentials (kinetic losses) due to acid loss from the CL and a reduction in catalyst electrochemically active surface area. According to the report, using the wrong plate materials would cause MEA degradation to soar. In [5], there are three main degradation mechanisms, including:

1. Dissolution of plate materials leading to poisoning of membrane;
2. Formation of resistive surface layer resulting in high ohmic resistance;
3. Possible fracture and continuous deformation of BPPs due to mechanical stress caused by high compressive pressure when sealing the stack.

For this review, the degradation phenomenon for both carbon BPP and metallic BPP have been considered.

a Carbon bipolar plate degradation

In early PEMFC studies, graphite was the most common material for bipolar plates [61]. However, drawbacks such as low mechanical properties, high gas permeability, brittleness, and high cost restrict the suitability of graphite BPPs. Due to these limitations, researchers have been forced to create new BPP materials. To overcome these drawbacks of graphite, composite materials, which combine graphite powder with a polymer matrix, have been investigated to provide flexibility to the BPP while significantly enhancing its mechanical strength and chemical stability [62–65].

Carbon corrosion in graphite and graphite composite BPPs under PEMFC operating conditions cannot be ignored. Even though carbon is used in BPPs and other components including GDLs and CLs, research on corrosion analysis of carbon in BPPs is sparse. In an acidic atmosphere, the carbon corrosion process normally involves two steps: the formation of surface oxide and the evolution of CO₂ [66]. The electrochemical oxidation of carbon can be adequately expressed in Equations (1) and (2), as described in Section 2.1.1. Corrosion
is not a significant issue in regular PEMFC operation; however, under severe conditions such as start-stop or fuel starvation, high potential at the electrodes can promote carbon corrosion [67,68].

b Metallic bipolar plate degradation

Metallic BBPs have high heat and electricity conductivity, mechanical stability, good manufacturability and gas impermeability, and easy machining of flow field channels compared to carbon BBPs. Metallic BBPs have these distinct advantages that allow for substantial thickness reduction [62,65]. However, in the aggressively corrosive atmosphere of high temperatures of 60–80 °C and low pH of 2–4 within PEMFCs, these materials are vulnerable to corrosion [69,70]. As a result, a thin oxide layer forms on the PEMFC surface, increasing the contact resistance between BBP and GDL. As a result, the internal electrical resistance of the fuel cell increases, resulting in a decrease in fuel cell efficiency [18,55]. Furthermore, metal cations (Fe$^{3+}$, Cr$^{3+}$, Ni$^{2+}$, etc.) formed during BBL corrosion inside the PEMFC may contaminate the membrane and CLs. Metal cation pollution has been found to severely reduce fuel cell capacity and durability [61,71]. As a result, an additional coating technique must be used to create a thin protective and electronic conducting layer on the exterior surface of BBPs [65].

Metallic BBPs will corrode at both the anode and the cathode, according to [61]. Because of the deterioration of the protective metal oxide layer, undesirable hydride formation and metal dissolution can occur instantly at the anode. This issue could increase the risk of membrane contamination and have a negative impact on the catalyst layer’s operation. In the presence of water, the corrosion rate of BBPs at the cathode will significantly increase.

2.1.5. Sealing Gasket

Structure and Role

Between the BBPs and the MEA, sealing gaskets are strategically positioned in PEMFCs. The main purpose of gaskets is to keep reactant gases and coolants from leaking. Gaskets often act as electrical insulators between components, allowing for proper stack height and variability control [67]. Furthermore, the seals control the compression of stack components, especially the prevention of GDL over-compression [17]. The optimal clamping force of a PEMFC is a function of the mechanical properties of the gasket materials, according to I. Gatto et al. [72].

PEMFCs require a sealing structure that carefully holds the reactant gases within their respective regions for long-term operation [73]. PEMFC sealing systems are usually divided into four categories: (1) PEM direct sealing, (2) PEM-wrapped frame sealing, (3) MEA wrapped frame packaging, and (4) rigid protective frame sealing. These sealing systems were intensively reviewed in [74].

When gaskets are exposed to a highly corrosive fuel cell environment, chemical stability and corrosion resistance are important. Since the fuel cell components are clamped together with adequate assembly force to prevent overboard leakage of the reactants, and the internal pressure within the cell during prolonged service is relatively high, they must have excellent mechanical strength and compressibility. Because of their excellent stress behavior, good flexibility, high chemical resistance, and good processability, PEMFC seals are generally made of elastomers, which are usually amorphous polymers composed of carbon, oxygen, and silicon [75]. Elastomers also have exceptional physical properties, such as transparency, permeability, insulation, and the ability to withstand massive strain [76]. Fluorine caoutchouc (FKM), ethylene-propylene-diene monomer (EPDM), and silicone are the most popular commercially available gasket materials used in PEMFCs [19]. The material behavior of these materials is intensively reviewed in [73].

Degradation Phenomena and Mechanisms

The long-term reliability and durability of the gasket are crucial to overall fuel cell efficiency and durability. If any of the gaskets degrades naturally, the reactants can leak or mix during operation, affecting fuel cell efficiency and posing safety concerns [77,78].
Moreover, seal failure can result in a loss of force retention, and compression and plate electrical shorting [19].

Loss of elastic property and seal functionality, as well as leakage of seal components that could contaminate the MEA, are typical degradation phenomena of sealing gaskets [67,78]. Attila Husar et al. [79] carefully analyzed gasket failure in a seven-cell PEMFC stack. It was indicated that the increase in stack temperature and pressure fluctuations in the anode and cathode, which resulted in a shift in cell voltage, were two leading indicators of the apparent inability. The leak of elevated temperature produced by the reaction of hydrogen and oxygen, which ignited when it met the Pt catalyst on the membrane, began the gasket failure. It was intensified by the loosening of the stack due to thermal cycling.

Despite the fact that there are numerous peer-reviewed publications on the degradation of elastomeric gasket materials in various environments, only a few reports on degradation and its mechanisms have been published, with the majority of them focusing on silicone rubber in the PEMFC setting. Jinzhu Tan et al. [77,78,80,81] investigated the degradation of commercial silicone rubber in the PEMFC environment. Their direct findings strongly suggest that chemical degradation induces an initial change in gasket surface chemistry. De-crosslinking and chain scission in the backbones of samples occur as a result of this change. Furthermore, stress that occurs in the samples accelerates silicone rubber degradation [82]. In [83], it was found that as the temperature rose, the nature of the rubber deteriorated increasingly, suggesting molecular degradation of silicone rubber. Simultaneously, in the presence of water, the oxygen content on the silicone surface increased, causing side-chain groups to eventually oxidize. Following that, as the amount of time spent in the fuel cell environment increased, the rubber macromolecules became increasingly shorter. PEMFC mechanical strength deteriorated over time and sealing performance deteriorated.

In summary, Table 1 details the primary failure modes for PEM fuel cell components such as the membrane, catalyst/catalyst substrate, GDL, bipolar plate, and sealing material [19,84].

| Component                  | Failure Modes                                    | Causes                                                                 |
|----------------------------|--------------------------------------------------|-----------------------------------------------------------------------|
| Membrane                   | Mechanical degradation                            | Mechanical stress due to non-uniform pressing and swelling Penetration of the catalyst and seal material traces and foreign particles |
|                            | Chemical degradation                              | Radical attack, Contaminations                                        |
|                            | Conductivity loss                                 | Ionic contaminations                                                  |
|                            | Activation losses                                 | Sintering or de-alloying of electrocatalyst                            |
|                            | Conductivity loss                                 | Corrosion of catalyst support                                         |
| Catalyst/catalyst layer CLs| Loss of reformate tolerance                      | Dissolution of alloying elements and contamination                    |
|                            | Decrease in mass transport rate of reactants      | Mechanical stress                                                     |
|                            | Decrease in water management ability              | Changing in hydrophobicity of materials                               |
| GDL                        | Decrease in mass transport                        | Mechanical stress, compression, Degradation of backing material       |
|                            | Decrease in water management ability              | Mechanical stress, Change in hydrophobicity of materials              |
|                            | Thinning                                          | Corrosion                                                             |
Table 1. Cont.

| Component       | Failure Modes          | Causes                        |
|-----------------|------------------------|------------------------------|
| Bipolar plate   | Conductivity loss      | Corrosion, oxidation         |
|                 | Fracture/deformation   | Mechanical stress            |
| Sealing gasket  | Mechanical failure and brittleness | Deformation, compression, and chemical reaction |

2.2. Operational Effects on PEMFC Durability

Fuel cell durability is considered to be affected by operating conditions. Several operational effects on fuel cell durability are discussed in this section. Impurities, start-up from subfreezing temperatures are the main issues. Potential cycling, fuel starvation, and start/stop cycling are among the additional operating conditions.

2.2.1. Impurity Effects

Contamination can have a negative impact on the efficiency and life of PEMFCs. Both the hydrogen fuel source and the air intake contain contaminants. The most common impurities and their sources are mentioned in Table 2. As shown in Table 2, the hydrogen fuel stream contains CO, H$_2$S, NH$_3$, and hydrocarbons as a result of the reforming of hydrocarbon fuel (crude oil, natural gas, and methanol). Carbon combustion and the atmosphere produce impurities in the air inlet, such as SO$_x$, NO$_x$, and volatile matter. Corrosion of fuel cell components and de-icers may also contain impurities. Different sections of the cell, such as the anode, the electrolyte membrane, and the cathode, are affected by these impurities.

Table 2. Origin of common fuel and air impurities [2] (Reprinted with permission from Borup et al. [2]. Copyright (2007) American Chemical Society).

| Sources                  | Potential Impurities                                      |
|--------------------------|----------------------------------------------------------|
| Crude oil                | CO, NH$_3$, H$_2$S, HCN, hydrocarbons                   |
| Natural gas              | CO, NH$_3$, H$_2$S, HCN, hydrocarbons                   |
| Methanol                 | CO, odorants, alcohols                                   |
| Biomass                  | Cations, aldehydes, alcohols, formic acid, NH$_3$, H$_2$S, HCN |
| Water electrolysis       | Anions, cations                                          |
| Fuel combustion pollution| SO$_x$, NO$_x$, hydrocarbons, soot, and particulates     |
| Ambient air, farming     | NH$_3$                                                   |
| Natural sources          | Ocean salts, dust                                        |
| Deicers                  | NaCl, CaCl$_2$                                           |
| Fuel cell corrosion products | Cations, anions                                   |

CO is one of the most challenging pollutants because it is difficult to separate and measure [85]. By forming a strong CO–Pt bond and chemisorbing on the metal surface, CO decreases the active surface area by occupying the Pt anode site. The chemisorbed CO prevents hydrogen from adsorbing into active Pt sites for hydrogen electro-oxidation (HOR), causing performance losses [2]. Due to CO crossover from anode to cathode via pin-holes in the membrane, CO impurities in the anode may also have a major impact on cathode output [86].

NH$_3$ is present both in hydrogen fuel and at the air intake. Fuel cell performance is reduced by as little as 13 ppm of NH$_3$ in H$_2$ fuel [76]. Depending on the NH$_3$ concentration and exposure period, the poison effect of NH$_3$ can be both reversible and irreversible [85]. Short-term (1 h) exposure has reversible effects, whereas long-term exposure has irre-
versible effects [2]. By replacing $H^+$ ions with $NH^4+$ produced by the reaction of $NH_3$ with ionomeric $H^+$, $NH_3$ impurities indirectly poison the catalyst, resulting in protonic conductivity loss of the anode CL and membrane [2,85,86]. As the $NH_3$ diffuses deeper, the negative effect begins at the anode CL, the first area exposed, and progresses to the membrane [2].

Another common PEMFC contaminant is hydrogen sulfide ($H_2S$). $H_2S$ has major degrading effects in the fuel cell, mostly due to Pt catalyst poisoning [86]. Small amounts of hydrogen sulfide ($H_2S$) can degrade fuel cell efficiency significantly. One experiment found that one ppm of $H_2S$ in $H_2$ resulted in measurable performance loss after only four hours of exposure, and nearly complete failure of service after 21 h. As sulfur adsorbs onto Pt, creating a strong $S$–Pt bond, $H_2S$ poisons the cell. This chemisorption, like CO, prevents sulfur species from adsorbing hydrogen into active Pt sites [2]. In addition, the presence of $H_2S$ can cause a decrease in water at the anode, lowering membrane conductivity and PEMFC performance. Additionally, $H_2S$ causes rapid voltage drops, lowering fuel cell capacity [85].

In the fuel reforming process, hydrocarbons are normally present, with methanol being the most common. Methanol in the fuel stream, on the other hand, has no poisoning effect on fuel cells. On the anode, exposure to other hydrocarbons such as benzene and toluene did not result in significant fuel cell degradation [2,85].

Popular atmospheric contaminants resulting from fossil fuel combustion, such as SOx and NOx, are exposed on the cathode side. By occupying the Pt surface and lowering the pH value in MEA, SO$_2$ will poison the fuel cell, resulting in free acid in the fuel cell and a possible drop in performance [86]. SOx poisons the Pt active sites in the cathode in the same way as $H_2S$ poisons the anode. The irreversible performance loss is caused by strong chemisorption of SO$_2$ on the Pt catalyst surface [2].

The majority of NOx in the air is made up of NO$_2$ and NO, with NO$_2$ accounting for 80% and NO accounting for the majority of the remaining 20% [87]. This has a negative impact on the efficiency of fuel cells. The presence of NOx in the cathode’s oxidant flow has been shown to cause the fuel cell’s performance to accelerate at first, then drop and stabilize [2,85]. NO$_2$ causes less serious performance loss than SO$_2$ [87]. At higher concentrations (>1 ppm), the damaging effects of NO$_2$ and NO on fuel cells were close, according to one experiment. Even so, at a concentration of one ppm, NO$_2$ only caused a minor current decrease, while NO had a major impact. It has been noted that nitric oxide has a direct effect on the catalyst and is absorbed primarily in NO [85]. The poisoning effect of NO$_2$ was also discovered to not be a catalyst surface poisoning species. NO$_2$, like $NH_3$ impurities, reacts with ionomeric $H^+$ to form $NH^4+$, which lowers protonic activity [2,86].

Salts like NaCl and CaCl$_2$ can contaminate the cathode air supply (mostly from deicers on roads during the winter or from aerosols in the air near marine environments). The presence of sodium chloride (NaCl) and calcium chloride (CaCl$_2$) reduces the efficiency of fuel cells and reduces the longevity of cell components. At high current densities, high salt concentrations reduce GDL’s hydrophobicity, increase liquid water retention, and thus reduce oxygen transport to the electrocatalyst. Because of the exchange of $H^+$ by cations at the CL and membrane, cations (Na$^+$, Ca$^{2+}$) cause decreases in proton conductivity [2]. Water flux and proton conductivity within the membrane are specifically affected by $H^+$ displacement with external cationic ions, leading to membrane degradation [86]. Anion $Cl^-$ lowers the electrochemical catalyst surface area (ECSA) of the cathode. The decrease in ECSA was attributed to both the site-blocking effect of directly adsorbed $Cl^-$ on the Pt surface and the electrochemical formation of chloride complexes, which increased Pt dissolution [88]. $Cl^-$ has also been shown to have an effect on GDL products, affecting water and gas transport [2].

### 2.2.2. Subfreezing Effects

The ability to start up and operate from subfreezing temperatures of $-20$ $^\circ$C to 50% rated power in as little as 30 s, with less than 5 MJ of energy consumption, is a
mandatory requirement for PEMFC stacks and systems, according to the DOE, especially in automotive applications [12]. One problem affecting PEMFC longevity is exposure to freezing temperatures after long-term shut-down [5]. Water formed at the cathode via electrochemical reaction and water-filled pores in MEA structures can freeze and lead to ice formation in subfreezing conditions. Due to the phase transition to ice, there is a 9% volume expansion, which can reduce efficiency by destroying cell components, blocking the gas passage, and covering the catalyst, resulting in cold start failure and even physical breakdown [89,90]. Ice formation can also lead to increased CL porosity and subsequent membrane delamination, resulting in a substantial decrease in cathode ECSA and an increase in membrane resistance [2]. Furthermore, repeated freeze-thaw cycles will delaminate CLs from both the membrane and the GDL, resulting in a loss of thermal and electrical contact [5]. After operating at sub-zero temperatures, the membrane surface can become rough and broken, and pinhole formation has been observed [91]. This failure causes gas crossover and an uncontrolled hydrogen-oxygen reaction, destroying the membrane and shortening the cell’s life [5,89].

At sub-zero temperatures, retained water in the PEMFC can prevent cold startup and cause irreversible PEMFC degradation. The bipolar plate and follow-field channels can be blocked with ice when the fuel cell is worked at subfreezing temperatures, resulting in startup failure and electrode degradation due to localized fuel starvation [89].

2.2.3. Other Operating Conditions Effects

One of the most common causes of fuel cell failure is starvation, which is described as fuel cell activity at sub-stoichiometric reactant conditions [92]. Reactant starvation, according to [93], includes both local and overall starvation. Insufficient reactant gas supply in partial areas of the catalyst layer causes local starvation. External current and voltage performance degradation is not readily apparent. A lack of reactant gas supply at the cell or stack level causes overall starvation. Outward current and voltage performance have also deteriorated. The fuel cell’s durability does not suffer as a result of this degradation. Starvation, on the other hand, causes cell reversal in the anode and cathode, with hydrogen being released in the cathode and oxygen being produced in the anode. Because of the high anode potential, this reversal results in a negative potential between the anode and the cathode [5]. Carbon corrosion of GDLs and subsequent electrocatalyst corrosion may be caused by gas starvation in the cathode or anode [5,93]. Gas lack eventually causes CL damage due to coalescence or catalyst loss on the electrode [92]. According to Zunyan Hu et al. [86], who studied cell interaction phenomena in a multi-cell stack with one cell suffering from fuel starvation, cell interaction is the main factor resulting in shorter lifespan for multi-cell stacks than for single cells. Current convergence caused by fuel starvation in one cell may cause local current convergence in normal cells nearby. Aside from that, the ECSA of the defective cell was reduced by 28%.

The startup and shutdown cycles of a fuel cell may have a significant impact on its durability [1]. All hydrogen will gradually cross over from the anode to the cathode during a prolonged shutdown, resulting in the anode flow channels being filled with air. In this situation, fuel cell startup can generate a transient state where fuel is present at the anode inlet, but the anode outlet is still fuel-starved. Localized fuel lack will cause the cathode local potential to exceed 1.8 V, resulting in significant fuel cell efficiency and durability degradation [1,2]. The effect of 1800 start-up and shut-down cycles on fuel cell efficiency and MEA degradation was demonstrated by R. Lin et al. [94]. The ECSA decreased from 479.6 cm$^2$·mg$^{-1}$ (fresh MEA) to 335.4 cm$^2$·mg$^{-1}$ after 1800 cycles, with an average decrease rate of 0.08 cm$^2$·mg$^{-1}$·cycle$^{-1}$; the ohmic resistance increased by 35%; and the charge transfer resistance of the cathode increased by around 90% relative to the fresh MEA. Increased charge transfer resistance of the oxygen reduction reaction can result from a decrease in ECSA. The cathode catalyst layer was also seriously harmed, with a 60 percent reduction in thickness and extreme carbon corrosion.
The voltage change caused by rapid changes in load required to match variable power demands in automotive applications is another aspect of fuel cell operations that is likely to affect cell integrity [2]. The current density of the fuel cell changes frequently during the load changing operation. The cell potential would also shift, ranging from 0.6 to 1.0 V in most cases. As the anode approaches the reversible hydrogen potential, the cathode experiences potential swings. The electrode material properties will change as the cathode potential changes, most notably the degree of oxide coverage of both platinum and carbon, which will prevent platinum dissolution at higher potentials [2]. Load cycling causes fuel cell failure due to water control issues, in addition to potential change. The flow rate of the gas in the flow channel is decreased when the load is suddenly reduced; as a result, the water content of the gas decreases, causing bulk water to be retained within the cell, resulting in flooding. Otherwise, the flow channel gas flow rate will increase during a sudden load increase, entraining much emitted water out of the fuel cell and causing membrane dehydration [93]. The effects of flooding and dehydration on fuel cell lifetime are addressed in Section 2.3.

2.3. Water Management

Water management is one of the most important problems that has a major impact on PEMFC efficiency and long-term durability [2,95]. Maintaining adequate humidification in a PEMFC is the ideal solution for limiting degradation [96]. While insufficient water operation can result in dehydration, too much water speeds up the flooding process, causing fuel cell degradation [97].

2.3.1. Dehydration

Since water is produced as a consistent product at the cathode, the catalyst layer and membrane on this side are kept hydrated. During dry operation, anode components appear to dehydrate, resulting in immediate and long-term fuel cell degradation [98]. With insufficient humidification to the cell, insufficient water compensation from the cathode through back-diffusion, and high electro-osmotic drag, the reactant gas supply can cause membrane dehydration [5]. When a fuel cell operates at a high temperature (typically above 60 °C), the evaporation speed may be faster than the speed of back-diffusion of water to the anode, resulting in cell drying. Additionally, the high current density is a complex operating condition in which the effect of water back-diffusion is weaker than that of electro-osmotic drag. Therefore, it is challenging to keep the anode side membrane wet. If the membrane is dehydrated, the proton conductivity will decrease in proportion with the water content, and then the ohmic losses will increase, leading to a considerable drop in voltage potential and power loss [99–101]. Furthermore, dehydration condition may lead the membrane to shrink, which hinders the back diffusion or induces irreversible degradation processes such as delamination and formation of pinholes. The membrane can become brittle and crack, resulting in reactant gas crossover [102,103].

2.3.2. Flooding

The so-called flooding phenomenon appears when too much water accumulates inside a fuel cell. Although this phenomenon is more likely to occur at the cathode side because of continuous water formation from the oxidation-reduction reaction, it is also observed at the membrane and flow channel cathode side.

At the cathode side of the membrane, water accumulates from the products generated by the oxygen reduction reaction (ORR), from the water migrating with protons across membranes by electro-osmotic drag transport, and from over-humidified reactant gas feeding [5,104]. An increase of the current density leads to more water formation by oxygen reduction reaction and dominant electro-osmotic drag through the membrane compared to back diffusion. On the other hand, the removal of water can occur via evaporation at high operating temperatures, back diffusion to the anode in low current density conditions (less than 0.3 A/cm²) [105], water-vapor diffusion, and capillary transport through GDL to flow
channels. Cathode flooding can be caused by imbalances between the water accumulation and removal process, accelerating mass transport losses. The pores of GDL are also blocked, and thus oxygen transport becomes limited in the PEMFC (gas starvation) [12]. Adverse effects of prolonged flooding on fuel cell performance, such as substantial increases in gas pressure drop, cell voltage, and potential losses, as well as fuel cell durability issues such as corrosion and contamination of components, were shown in many studies [106–109].

The flooding phenomenon at the anode was observed when fuel cell operates at low current densities. Back diffusion prevails over electro-osmotic drag effect at low-temperature operation due to restricted evaporation of accumulated water and the supply of over-humidified hydrogen. This phenomenon also causes fuel starvation, carbon substrate oxidation, and oxygen evolution [103]. Mansu Kim et al. [110] confirmed the profound degradation of cell voltage and current density every 250 h during a 1600 h experiment under anode flooding conditions. The anode flooding was found to affect the fuel cell performance and durability.

Flow channel flooding is another barrier in the design and operation of PEMFCs. Redundant accumulated water may form water columns or water bands inside flow channels and block gas flow; thus, pressure drop is increased [111]. An excessive pressure drop increases the power consumption and reduces the fuel cell system performance and durability [112]. Furthermore, if the fuel cell is exposed to sub-freezing temperatures after the operation, the water may freeze and harm component internal structures. In order to remove and limit the retention of bulk water and big droplets in flow field channels, a water-clearing fuel cell is necessary for flow field designs. A purge is frequently conducted before freezing to eliminate liquid water [90].

2.4. Thermal Management

Apart from water formation, heat is another product of fuel cell operation, which significantly affects PEMFCs because cell temperature influences virtually all parameters such as mass diffusion coefficient, maximum theoretical voltage, electrochemical activity, ionic conductivity, etc. [106–109]. Improper thermal management under sub-zero and elevated temperature operation can lead to performance and lifetime degradation.

Because water is either generated within the fuel cell or often provided as external humidification, it poses a significant concern for low-temperature operation. As mentioned in Section 2.2.2, during start-up and operation at sub-zero temperatures over long times, ice generated inside the stack can block reactant channels and pores, causing voltage drop or even shutting down the electrochemical reaction [5,113,114]. Fuel cells under freeze-thaw cycles also show reversible and irreversible degradation and structural damage [115]. The repetition of ice formation and melting into water can delaminate the MEA, the backing layers, and the gas diffusion layers. The mechanical damage may lead to gas crossover and then an uncontrolled reaction between hydrogen and oxygen, deteriorating PEMFC performance and life. Also, lower temperature operation results in a decrease in the power output of PEMFCs. This does not mean that the high-temperature process always yields high performance and prolongs fuel cell life, despite its advantages of good electrochemical kinetics, increased tolerance of contaminants, enhancement of water management and cooling, and CO-poisoning reduction [5].

Fuel cells working at high temperature (above 100 °C) also show some limitations, such as increases of material degradation rates and associated contaminant levels and decreases of water content in membrane and ionomer [5,102]. Therefore, the ionic conductivity should be reduced, and holes or hot spots may appear, leading to reduction of longtime performance and structural damage [5]. Mohammad A. Barique et al. [116] reported that Nafion NR212 membrane’s conductivity increases continuously up to 80 °C, but remains stable at temperatures from 80 to 100 °C.

The amount of heat generated by PEMFCs should be detected first, and then removed from the fuel cell by using a suitable cooling technology to keep the cells at the required temperature. Mohamed H.S. Bargal et al. [117] reviewed various methods for cooling
PEMFC, particularly in automobile applications. Edge cooling, cooling using separate airflow, air cooling, cooling using the cathode air supply, phase change cooling, and liquid cooling are some of the heat management strategies that can be used.

3. Cell/Stack Durability Test Protocols

The durability testing protocols developed or used for single cells or stacks will be summarized in this section for both vehicle and stationary applications.

3.1. Durability Testing Protocols for Transport Applications

Fuel cells for transportation applications, especially for automotive propulsion, may have to operate under different operating and cyclical conditions, including temperatures from sub-zero to above the boiling point of water, relative humidity from ambient to saturation, half-cell voltages from 0 to >1.5 volts. Additionally, the operation conditions during the transient and cyclic modes can change relatively quickly, ranging from high to low voltages, temperatures, humidity, and currents to meet the rapid changes in power demand, some of them severe special materials and the stability of the whole fuel cell system [7,8,118]. Institutions and organizations in India, South Korea, and Russia, and especially in the U.S, EU, Japan, and China, have made significant efforts to propose methods to quantify the durability of the PEMFC cell/stack for automobile applications [11]. Sections 3.1.1–3.1.9 will discuss these test protocols.

3.1.1. The Dynamic Stress Test Protocol (DST)

The DST protocol was developed by the DOE and FCHEA for testing the long-term durability of PEMFC cell/stacks for vehicular applications and comparing cell and stack performance with that specified by the U.S. DOE [2,9,10]. This protocol involves stepping through a series of different current draws typical of vehicle loads based on real power demand from a car during city driving in a typical American urban area [4]. When average voltages are 0.88, 0.80, 0.75, 0.65, and 0.60 V, an initial polarization curve is used to determine current densities, according to the protocol’s guidelines. These current densities are designated C88, C80, C75, C65, and C60, respectively [10]. Figure 1 depicts the stack load period used for fuel cell testing based on the initial stack polarization curve and Table 3 shows the current density/time profile. The cycle includes open circuit voltage (OCV), low current, medium current, and high current (16 steps in total). The first stage (15 s) of each 360-s loop is OCV, as shown in Figure 1. Fuel and oxidant compositions, as well as flow rates, will be calibrated at each current level to represent realistic system conditions over a driving cycle. Except for periodic diagnostic testing, the periods would be continuous. Cell resistance and hydrogen crossover, shorting resistance, and a constant-stoichiometry polarization curve would be among the diagnostics.

Since several issues (such as fuel cell activity during freeze/thaw cycles) are not addressed, this protocol is not intended to be comprehensive [118]. As a result, the effects of this protocol may or may not correctly estimate the lifetime of a fuel cell under real-world driving conditions [2]. Therefore, additional tests to correlate these results to actual lifetime are needed, including actual driving, start/stop, and freeze/thaw cycles.

Since it is design-specific, this protocol is better suited to vehicles traveling on American roads, where speeds are higher and hill climbing is more common [119,120]. As a result, other countries should change the protocol to measure the durability of vehicular PEMFCs based on their own road characteristics [4].
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### Figure 1.
The Dynamic Stress Test (DST) duty cycle used for fuel cell testing [2] (Reprinted with permission from Borup et al. [2]. Copyright (2007) American Chemical Society).

### Table 3.
Current density vs. time for cycle profile (Reprinted with permission from Borup et al. [2]. Copyright (2007) American Chemical Society).

| Step | Duration (s) | Cxx | Step | Duration (s) | Cxx |
|------|--------------|-----|------|--------------|-----|
| 1    | 15           | OCV | 9    | 20           | C75 |
| 2    | 25           | C80 | 10   | 15           | C88 |
| 3    | 20           | C75 | 11   | 35           | C80 |
| 4    | 15           | C88 | 12   | 20           | C60 |
| 5    | 24           | C80 | 13   | 35           | C65 |
| 6    | 20           | C75 | 14   | 8            | C88 |
| 7    | 15           | C88 | 15   | 35           | C75 |
| 8    | 25           | C80 | 16   | 40           | C88 |

### 3.1.2. Fuel Cell Technical Team (FCTT)

The FCTT protocol was developed by the Fuel Cell Technical Team of USDRIVE. This drive-cycle protocol targets test system, stack, and MEA lifetime under simulated automotive drive cycle conditions. The FCTT, like the DST, incorporates typical operating conditions for fuel cell vehicles. However, it does exclude situations such as unmitigated start/stop, freeze/thaw, fuel starvation, and system fault conditions [8].

The FCTT proposes two similar-structured cycles for wet and dry conditions [8,9,120]. The quantities of humidification used and the highest current density applied vary. In comparison to the dry cycle, which uses 25% relative humidity and 0.1 A/cm² maximum current density, the wet cycle uses 92% relative humidity and a higher maximum current density, 1.2 A/cm². The membrane hydration in the dry protocol is dependent on the PEMFC’s self-humidification due to the lower relative humidity. FCTT’s duty cycle profile is depicted in Figure 2. The PEMFC stack, as shown in Figure 2, works on a 60 s cycle, alternating between 30 s of idling and 30 s of full power. In the wet process, two load steps of 0.02 A/cm² and 1.2 A/cm² are fixed, while gas dew point temperatures are set at 83 °C. The load is set at 0.02 A/cm² and 0.1 A/cm² in the dry cycle, and the gas dew point temperatures are set at 53 °C. The stack temperature in these two procedures is 80 °C, and the gas stoichiometry remains constant throughout.
3.1.2. Fuel Cell Technical Team (FCTT)

PEMFCs based on their own road characteristics [4]. In cities, such as low engine power and a top speed of 50 km/h [119]. The NEDC has an alternating characteristic of the idling and full power conditions, as seen in Figure 2, illustrates this protocol’s key distinction from others. It emphasizes the impact of idling and full power on the durability of PEMFC stacks. As a result, this protocol is better suited to testing the durability of city buses with low speeds and frequent stops, as well as investigating the impact of idling and full power on full-power PEMFC cars and hybrid PEMFC cars in daily use [4].

3.1.3. The New European Driving Cycle (NEDC)

The United Nations Economic Commission for Europe (UNECE) developed the NEDC, which is used to measure emissions and fuel efficiency in passenger cars in European cities. It uses load cycles to simulate real-world driving conditions in the lab [4,9]. By repeatedly exposing the cell to the same load cycle, the load cycle is used to measure fuel cell durability over a long period of time. The fuel cell cycle, as shown in Figure 3, is based on the NEDC [84]. The NEDC cycle, which involves acceleration, deceleration, and constant speed, is commonly used for light-duty vehicles. It consists of four 195 s repetitions of a low-speed urban phase, each followed by a 400 s simulation of a motorway (highway) driving period. The total time of the NEDC is 1180 s, which equates to a theoretical distance of about 11 km driven in about 20 min. The NEDC mode includes zero-velocity of the vehicle, which means the power propulsion system is in idle mode, as shown in Figure 3 (left-side). As a result, the cell’s operating demand is not zero. As a result, as shown in Figure 3 (right-side), the revised electric load demand schedule should include the idle mode [121]. The y-axis changes from velocity to percent load, and the percent load of zero velocity moves to 5% load. Since rated power specifications differ from vehicle to vehicle, the percent load is determined by the NEDC mode’s load profile. The 100% current load value used in the Fuel Cell Dynamic Load Cycle (FC-DLC) is defined as the average current density of the ascending and descending polarization curve measurements to yield a cell voltage of 0.65 V [84].

Although this protocol was designed for single-cell testing for automobile applications, it can be used to test the durability of PEMFC stacks because it includes all of the typical operating conditions found in PEMFC vehicle daily driving, such as start-up/shutdown, idling, partial power running, full power running, and dynamic load running [4,121]. Furthermore, it simulates the driving characteristics of both urban and extra-urban areas. As a result, this protocol is appropriate for evaluating the durability of PEMFC and PEMFC hybrid vehicles under a variety of road conditions. However, this protocol includes a set of assumptions about how a fuel cell car’s driver will operate the vehicle in crowded European cities, such as low engine power and a top speed of 50 km/h [119]. The NEDC has an inherent flaw in that it is based on a theoretical driving scheme and includes a number of tolerances and flexibilities that do not accurately reflect real-world driving conditions, such as slow accelerations and slow speeds [122,123]. As a result, PEMFC cell/stack durability and lifetime are unrealistic and impractical. Hence, a new test protocol that is more realistic to real-world driving conditions is required.
3.1.4. Worldwide Harmonized Light Vehicle Test Procedure (WLTP)

The European Union developed the Worldwide Harmonized Light Vehicle Test Procedure (WLTP) to replace the previous NEDC test and officially implemented all new car registrations throughout Europe from September 2018 [124]. The goal of WLTP was to provide a more reliable and precise test basis for automobiles. The WLTP was developed using real-world driving data from five different regions: the European Union plus Switzerland, the United States, India, Korea, and Japan, which covers a wide range of driving conditions, from city traffic to highways, to provide more accurate CO₂ and fuel consumption figures that better represent a car’s on-road efficiency [125,126].

Since the WLTP was designed to harmonize test procedures around the world, it involved the development of a new cycle called the World Harmonized Light-Duty Vehicle Test Cycle (WLTC), which represents common driving characteristics around the world. Based on a vehicle’s power-to-mass (PMR) ratio and its maximum speed, three separate driving cycles were set up to reflect three vehicle classes [127].

The cycles are divided into four phases: low-speed, medium-speed, high-speed, and an extra-high-speed phase that is common on European highways. Figure 4 depicts the speed pattern for a Class 3 vehicle, which is likely to account for the majority of the global light-duty vehicle market. Class 3 vehicles include hybrid and electric automobiles [128]. The cycle’s duration was set at 1800 s.

In comparison to the NEDC, the WLTP test is much more dynamic. It contains many braking and accelerating cycles, including four complex cycles that cover a wide range of speeds from low to extra-high. These WLTC dynamic cycles accurately simulate the regular driving behavior of light-duty vehicles under real-world conditions. The differences between NEDC and WLTP are shown in Table 4 [87,90].

The WLTC’s characteristics demonstrate that it is ideal for measuring the reliability of daily PEMFC and PEMFC hybrid cars, as well as other electric vehicles. Although the WLTP has been tested in a number of laboratories around the world, these assessments focused primarily on CO₂ emissions, fuel consumption, and other traditional aspects of vehicle efficiency. Since fewer studies have been conducted with this protocol for the PEMFC cell/stack around the world, its rationality and validity must be verified.
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![Figure 4. NEDC and World Harmonized Light-Duty Vehicle Test Cycle (WLTC) class 3 driving profiles over time [129] (Reproduced with permission from D. Tsokolis et al. [129], Applied Energy; published by Elsevier Ltd., 2016).](image-url)

**Table 4.** Main differences between NEDC and Worldwide Harmonized Light Vehicle Test Procedure (WLTP) [122,124,125,129–131].

| Parameter         | NEDC                  | WLTP (Class 3)          | Difference | Note                             |
|-------------------|-----------------------|-------------------------|------------|----------------------------------|
| Starting temperature | Cold                  | Cold                    |            |                                  |
| Test cycle        | Single test cycle     | Dynamic cycle more      |            |                                  |
| Cycle time        | 1180 s                | 1800 s                  | +35%       |                                  |
| Idle time         | 23.7%                 | 13%                     | −45%       | Less effect of engine start/stop |
| Cruise time       | 38%                   | 4%                      | −90%       | Less stationary operation        |
| Acceleration      | 20%                   | 44%                     | +117%      | More acceleration                |
| Deceleration      | 14%                   | 40%                     | +179%      | More deceleration                |
| Cycle distance    | 11 km                 | 23.25 km                |            |                                  |
| Driving phases    | Two phases, 66% urban and 34% non-urban driving | Four dynamic phases with more emphasis on high performance, 52% urban and 48% non-urban driving | Two more |                                  |
| Average speed     | 34 km/h               | 46.5 km/h               | +53%       |                                  |
| Maximum speed     | 120 km/h              | 131 km/h                | +4%        |                                  |

3.1.5. The Fuel Cell Testing and Standardization Network (FCTESTNEST)

The FCTESTNET of the Fuel Cell Testing and Standardization Network (FC TestNet) was developed based on cycling conditions with current density following a dynamic profile versus time to determine the evolution of the voltage of a PEMFC single cell
operating in specified conditions during a durability test [132]. This protocol is specifically designed for transportation applications since the load profile as defined is an adaptation of the type of power profiles defined for different vehicles, including power levels varying with dynamic variation. Two similar durability protocols were proposed based on different assumptions, a smooth power increase followed by power-off (Figure 5) and on-off cycling (Figure 6) [133].

![Figure 5](image.jpg)

**Figure 5.** The dynamic test profile proposed by FCTESTNET [133] (Reproduced with permission from Ira Bloom et al. [133], ESC Transactions; published by The Electrochemical Society, 2013).

![Figure 6](image.jpg)

**Figure 6.** Duty cycle from International Electrotechnical Commission (IEC) and Fuel Cell Testing and Standardization Network (FC TestNet) [120] (Reproduced with permission from Ira Bloom et al. [119], Journal of Power Sources; published by Elsevier B.V, 2013).

As can be seen in Figure 5, the whole duration of one cycle of this protocol is 60 s, including the alternative operation of low-power stationary phase (40 s at \( i \)@ 20% \( P_{\text{max}} \) A/cm\(^2\)) and high-power dynamic phase (20 s with an increase of \( i \) from \( I_{\text{load min}} \) @ 20% \( P_{\text{max}} \) to \( I_{\text{load max}} \) @ 100% \( P_{\text{max}} \) A/cm\(^2\)). \( P_{\text{max}} \) is defined here as the maximum power to be used during regular operation. The dynamic cycling phase is conducted after an initial polarization curve and followed by a final polarization curve. The comparison of these two polarization curves will be used to quantify the performance loss of the cell for the entire range of current density, allowing us to analyze the causes of performance degradation.

The other protocol was proposed by a collaboration of the IEC and FC TestNet, as shown in Figure 6 [119]. One cycle of this protocol lasts 3600 s, comprising the 1800 s alternative operation, full power condition, and 1800 s partial power condition. As
illustrated in Figure 6, this protocol is similar to FCTT. Both protocols merely switch between two types of normal operational conditions, ignoring all other scenarios. The distinction is that the specific operating conditions addressed in this protocol are full power and partial power. In addition, this protocol has greater cycle durations than the FCTT. This protocol primarily studies the durability of a PEMFC stack that switches between two operating modes (full power and partial power), making it appropriate for vehicles like city buses [4].

3.1.6. The Giantleap Protocol

Giantleap from the Fuel Cell and Hydrogen Joint Undertaking (FCH) was focused on using passenger-car fuel cell stacks and automotive balance of plant components in intercity buses. This protocol was proposed based on a dynamic load cycle for endurance testing and can be carried out in both single cells as well as in stacks. A detailed description of the Giantleap protocol is given in [134].

The cycle involves a two-hour cycle that begins with a short duration at OCV and a very gradual ramp-up rate \((0.04 \text{ A cm}^{-2} \text{s}^{-1})\) to achieve high-efficiency operation. A 650 s length at a high-efficiency process is followed by an extra-long length at a steady state nominal operation (2500 s). Gas flow rates should be changed at least 3 s before ramping up the load and 3 s after ramping down the load to preserve a strategic operation from gas starvation. Furthermore, continuous current for long periods of time can cause some degradation (reversible degradation). As a result, the protocol includes some dynamics, moving back to high-efficiency operation for 650 s and then back to high-power operation for another 2500 s. The stack (or the cell in single-cell testing) power is ramped down at the same rate as the ramp-up rate after the steady-state cycles. The stack is placed into idling mode for 420 s after a brief time of 10 s at OCV.

For the fuel cell system to be used in the Giantleap project, the Giantleap protocol was specifically proposed, with the operating conditions shown in Table 5. It is strongly advised that the working conditions be tailored to the particular fuel cell technology and application of the manufacturer.

Table 5. Operating parameters of Giantleap.

| Parameters                              | Unit | Ref of NEDC  | Giantleap Conditions at Low Current Density | Giantleap Conditions at High Current Density |
|-----------------------------------------|------|--------------|--------------------------------------------|--------------------------------------------|
| Nominal cell operating temperature     | °C   | 80           | 73                                         | 75                                         |
| Fuel gas inlet temperature             | °C   | 85           | 73                                         | 75                                         |
| Fuel gas inlet humidity                | % RH | 50% (DPT 64 °C @ 80 °C) | 45                                         | 45                                         |
| Fuel gas inlet pressure (absolute)     | kPa  | 250          | 140                                        | 190                                        |
| Fuel stoichiometry                     |      | 1.3          | 1.5                                        | 2.5                                        |
| Oxidation gas inlet temperature        | °C   | 85           | 73                                         | 75                                         |
| Oxidation gas inlet humidity           | % RH | 30 (DPT 53 °C @ 80 °C) | 75                                         | 75                                         |
| Oxidation gas inlet pressure (absolute)| kPa  | 230          | 130                                        | 180                                        |
| Air stoichiometry                      |      | 1.5          | 1.7                                        | 1.7                                        |
| Minimum current density for stoichiometry operation | A/cm² | 0.2          | 0.2                                        | 0.2                                        |
3.1.7. The Japanese JC08 Cycle

The JC08 protocol of Japan was first introduced by Japanese emission regulation authorities in 2005 for light vehicles (<3500 kg GVW) [135]; it was implemented officially in stages from October 2007 [136] and then formally proposed in October 2015, replacing the previous 11 and 10–15 protocols [4]. This protocol was initially used for emission measurement and fuel economy determination for internal combustion engine vehicles (e.g., gasoline and diesel vehicles) [135]. However, it has always been used to test PEMFC stack durability in hybrid cars [128,137]. It represents driving in congested traffic in Japanese cities, including idling periods and frequently alternating acceleration and deceleration.

For fuel efficiency measurement under the new fuel efficiency standards, the JC08 mode is used as a driving mode, consistent with the exhaust emission measurement method, and evaluates fuel performance more accurately [92]. Cold-start driving before the engine has warmed up, as well as hot-start driving after the engine has warmed up, are used to mimic real driving as closely as possible. A cold start and a warm start are used in the JC08 measurement.

As shown in Figures 7 and 8, the entire test in 10–15 mode takes 660 s, while the JC08 takes twice as long, at 1200 s. The total test distance in JC08 mode is 8.171 km, with an average test speed of 24.4 km per hour (including idling time) [135,136]. The rising and falling edges of the curve appear more frequently in the JC08 (total 26 times) than in the 10–15 mode, indicating that the JC08 frequently simulates the dynamic loading state. This indicates that the protocol closely resembles real-world driving conditions on city streets. In addition, the time for driving at a constant speed has been reduced, simulating the situation of constantly getting on and off the accelerator. Additionally, before being completely baked, a vehicle’s average speed is held at 50 km/h, simulating waiting for traffic lights in city driving. Furthermore, slight oscillations between 700 and 1000 s can be used as emergency braking. Finally, after hitting a top speed of 82 km/h, the car encountered a brief moment of vibration, simulating the condition of driving on non-urban highways [4,136]. JC08 is somewhat close to real driving conditions due to all of the urban and extra-urban driving scenarios used in the protocol. As a result, many automakers have used it in recent years to measure the reliability of their own PEMFC and hybrid vehicles [4].

![Figure 7. Evolution of speed versus time for the 10–15 protocol (Reproduced with permission from Huicui et al. [4], Applied Energy; published by Elsevier Ltd., 2018).](image-url)
3.1.8. Test Protocols in China

PEMFC durability test protocols were developed by leading universities and institutes in research PEMFC technology in China, including Tsinghua University, Tongji University, the Dalian Institute of Chemical Physics, and Wuhan University of Technology [138]. These protocols were intensively summarized in [4].

Tongji University’s test protocol was based on the NEDC and took into account real-world operating conditions. The protocol, like the NEDC, encompasses a wide range of real-world driving scenarios, including startup-shutdown, idling, partial power, full power, overload, and dynamic load running. The protocol lasts 1200 s, as shown in Figure 9. In this protocol, the operating conditions are mostly the same as in the NEDC. The fuel gas and oxidation gas inlets, on the other hand, are fully humidified [4].

Tsinghua University proposed a durability test protocol based on a typical Chinese bus driving cycle, which is divided into two phases as shown in Figure 10. Idling, dynamic load, and partial power running are all part of stage I; full power and overload running are part of phase II. Four phases I and one phase II make up a single driving cycle, with a high proportion of idling and full power conditions. The total length of this protocol is 7235 s, with 38.9% idling and 28.2% full power, similar to city bus driving characteristics. It demonstrates that this protocol can be used to evaluate the PEMFC city bus [4].
The Wuhan University of Technology’s test protocol is shown in Figure 11. This protocol was proposed to reveal the dominant impacts on PEMFC durability based on normal operating conditions in the actual driving process. The total length of this protocol is 25,800 s (430 min), which includes OCV, idling, dynamic load with RH cycling, full power, and overload. This protocol includes a large time portion for the dynamic load condition to represent the primary operating condition of PEMFCs in real-world road conditions. Aside from that, this load period includes RH cycling. Combining the two requirements helps the model to represent lifetime shift circumstances quicker, making it more suitable for daily-used PEMFC car durability and hybrid PEMFC car durability on city roads [4].

Centered on the Fuel Cell Engine (FCE) testing mode, the Dalian Institute of Chemical Physics developed a protocol for testing PEMFC stability under significantly severe operating conditions, as shown in Figure 12. The total length of this protocol is 5700 s (95 min), which includes idling, partial power, full power, and overload conditions. This protocol simulates the theoretical PEMFC stack operation (startup–incremental loading–full power running–overload running–shutdown) without the use of a dynamic load. The 20 min of idling operation is a demanding test for the fuel cell. The idling situation is a severe test of the PEMFC’s adaptability due to the relatively low stack working temperature and relatively high temperature at the air inlet [4].
Figure 11. Wuhan University of Technology test protocol [4] (Reproduced with permission from Huicui et al. [4], Applied Energy; published by Elsevier Ltd., 2018).

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Figure 12. Dalian Institute of Chemical Physics test protocol [4] (Reproduced with permission from Huicui et al. [4], Applied Energy; published by Elsevier Ltd., 2018).

Most of the standard operating conditions of actual driving actions are covered by the above test protocols, including startup-shutdown, idling, partial power running, full power running, overload running, and dynamic load running. The only difference, as shown in Table 6, is the length of each operation condition in the cycle [4].

Table 6. Comparison of PEMFC durability test protocol of universities and institutes in China [4] (P—rated power, T—whole duration) (Reproduced with permission from Huicui et al. [4], Applied Energy; published by Elsevier Ltd., 2018).

| Operating Condition | Tongji University | Tsinghua University | Wuhan University of Technology | Dalian Institute of Chemical Physics |
|---------------------|------------------|---------------------|-------------------------------|--------------------------------------|
| OCV                 | 7.3% of T        |                     | 4.6% of T                     |                                      |
| Idling              |                  | 38.9% of T          |                               |                                      |
| Partial power       |                  |                     |                               |                                      |
| 80% P is 6.2% of T  |                  | 20% P is 3.5% of T  | 40% P is 8.4% of T            |                                      |
| 70% P is 8.4% of T  |                  | 40% P is 3.3% of T  | 60% P is 26.3% of T           |                                      |
| 90% P is 0.8% of T  |                  | 60% P is 3.3% of T  | 80% P is 10.5% of T           |                                      |
| Full power          |                  |                     | 34.9% of T                    | 31.6% of T                           |
| Overload            | 5.1% of T        | 1.8% of T           | 2.3% of T                     | 2.1% of T                            |
| Dynamic load        |                  |                     |                               |                                      |
| OCV to 15% P, cycling four times, 8.7% of T | Idling to full power, cycling 40 times, 37.2% of T | Idling to overload, cycling five times, 4.7% of T | OCV to idling, cycling ten times, 9.3% of T |
| OCV to 32% P, cycling four times, 17.4% of T | Idling to full power, cycling 20 times, 17.7% of T |                               |                                      |
| OCV to 50% P to 32% P, cycling four times, 23.6% of T |                               |                               |                                      |

3.1.9. Summary

The key feature of fuel cells in vehicular applications is load cycling. A fuel cell’s ability to operate successfully is determined not only by its efficiency but also by its durability. Endurance testing is used to assess fuel cell durability, which involves applying a repeated load profile to the cell and calculating performance degradation in terms of cell voltage decrease as a function of operating hours. To determine the cell/stack degradation rate, protocols based on a dynamic load cycle for endurance testing were proposed. An example of a PEMFC stack long-term durability experiment is illustrated in Figure 13.
Table 6. Comparison of PEMFC durability test protocol of universities and institutes in China [4] (P—rated power, T—whole duration) (Reproduced with permission from Huicui et al. [4], Applied Energy; published by Elsevier Ltd., 2018).

| Operating Condition          | Tongji University | Tsinghua University | Wuhan University of Technology | Dalian Institute of Chemical Physics |
|------------------------------|-------------------|---------------------|---------------------------------|-------------------------------------|
| OCV                          | 7.3% of T         | 4.6% of T           |                                 |                                     |
| Idling                       | 38.9% of T        | 21.1% of T          |                                 |                                     |
| Partial power 80%P           | 6.2% of T         | 70%P is 8.4% of T   | 90%P is 0.8% of T               | 20%P is 3.5% of T                  |
| 60%P is 3.3% of T            |                   | 40%P is 8.4% of T   | 60%P is 26.3% of T              |                                     |
| 80%P is 3.3% of T            |                   |                     | 80%P is 10.5% of T              |                                     |
| Full power                  | 22.5% of T        | 28.2% of T          | 34.9% of T                      | 31.6% of T                         |
| Overload                     | 5.1% of T         | 1.8% of T           | 2.3% of T                       | 2.1% of T                          |
| Dynamic load OCV to 15%P, cycling four times | 8.7% of T |                                   |                                 |                                     |
| OCV to 32%P, cycling four times |                   |                     |                                 |                                     |
| OCV to 50%P to 32%P, cycling four times | 23.6% of T |                   |                                 |                                     |
| Idling to full power, cycling 20 times | 17.7% of T |                                   |                                 |                                     |
| Idling to full power, cycling 40 times | 37.2% of T |                                   |                                 |                                     |
| Idling to overload, cycling five times | 4.7% of T |                                   |                                 |                                     |
| OCV to idling, cycling ten times | 9.3% of T |                                   |                                 |                                     |

3.1.9. Summary

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Figure 13. An example of a PEMFC stack long-term durability experiment.

According to a review of protocols from various organizations, institutions, and institutes around the world, a systematic and useful durability test protocol for PEMFC cells, stacks, and systems for vehicles should represent real-world driving conditions. Startup/shutdown, idling operation, partial power operation, full power operation, overload operation, and dynamic load operation are examples of these situations. According to tests and various driving conditions, the time portion of each condition in the entire cycle can be changed. To more comprehensively simulate the driving situation of PEMFC vehicles on actual roads, the dynamic load condition for light-duty vehicles should be planned and arranged with a large portion of the total length. A protocol, on the other hand, consists of a significant portion of idling and steady service, making it ideal for city buses [4]. The applicable vehicle types of PEMFC durability test protocols are summarized in Table 7.

Table 7. Summary of applicable vehicle types of PEMFC durability test protocols.

| Protocol                  | Vehicle Types |
|---------------------------|---------------|
| Dynamic Stress Test (DST) | ✓             |
| FCTT                      | ✓             |
| NEDC                      | ✓             |
| WLTP                      | ✓             |
| FCTESTNET                 | ✓             |
| Giantleap                 |               |
| JC08                      | ✓             |
| Tongji University         | ✓             |
| Tsinghua University       |               |
| Wuhan University of Technology | ✓   |
| Dalian Institute of Chemical Physics | ✓   |

3.2. Test Protocols for Stationary Applications

The effects of long-term operation on cell/stack performance in terms of voltage decay rate as a function of operating hours are investigated in aging or durability tests for PEMFCs. In general, there are two types of durability tests for stationary applications: steady-state durability tests and load cycling durability tests.
3.2.1. Steady-State Durability Tests

For stationary applications, the majority of long-term lifetime experiments are carried out under steady-state conditions (constant voltage or constant current) for long periods of time, often up to several thousand hours [1,2]. In terms of durability, the DOE developed steady-state protocols under constant current density and compared them to DOE targets. The cell will be run at a current density that produces 25% maximum power under stable operating conditions such as temperature, pressure, and stoichiometry. Except for in situ periodic diagnostic testing (every 200–250 h), the testing will be continuous [119]. Diagnostic testing aims to characterize variations in fundamental properties such as polarization curves, membrane resistance, hydrogen crossover, and electrochemical surface area as a function of time [1].

A test module of FCTESTNET was proposed for a long-term durability steady test under constant current density for a single PEMFC. The aim of the module was to figure out how a PEFC single cell’s voltage changed over time during a long-term durability test, which was mostly done in steady operation at a fixed current density but also included recurrent polarization curves to qualify the cell’s output at set intervals during the test. The polarization curves will be run at fixed intervals of $t_{\text{max}}/10$, where $t_{\text{max}}$ is the maximum period of the test as specified by the particular objective of the test module. Depending on the operating conditions and the program, $t_{\text{max}}$ can be set anywhere between 500 and 10,000 h. The final polarization curve will be compared to the initial one to qualify the cell’s performance loss over the entire range of current density and to investigate the causes of performance degradation, if any. Unless there is another ending criterion, such as specifying the test stop if the cell voltage falls below a pre-defined minimum voltage, the long-term durability test length would be $t_{\text{max}}$. If the components under examination are to be used for further testing, a minimum voltage of 0.5 V is recommended, and 0.3 V is recommended for research purposes, such as qualifying the lifetime and researching degradation processes to prevent irreversible harm to cell components [139].

In recent years, the STACKTEST project (European Development of PEM Fuel Cell Stack Reference Test Procedures for Industry) identified and validated the industrially applicable generic test modules [140,141]. A constant load durability test module [120] was introduced for stationary applications, with specific test input parameters shown in Table 8. Under continuous operation load, a model was used to assess the irreversible voltage decay rate induced by specific operating parameters. Throughout the test, a specific set of operating conditions was maintained, and the stack voltage and individual cell voltages were reported as functions of time. Aside from that, for MEA integrity reasons, the minimum single cell voltage limit should be set no lower than 0.3 V to avoid stack harm.

Table 8. Recommended nominal reference operating conditions for stationary application in STACKTEST project.

| Parameter                     | Stationary Application | Combined Heat and Power (CHP) | Backup |
|-------------------------------|------------------------|-------------------------------|--------|
| Stack temperature (Coolant inlet) | 70 °C                  | 65 °C                         |
| Reactant inlet temperature    | 75 °C                  | 70 °C                         |
| Fuel stoichiometry            | 1.2                    | 1.25                          |
| Oxidation stoichiometry       | 2.0                    | 2.0                           |
| Fuel relative humidity        | 80%                    | 40%                           |
| Dew point temperature fuel    | 69.5 °C                | 45.5 °C                       |
| Oxidant relative humidity     | 80%                    | 40%                           |
| Dew point temperature oxidation| 65 °C                  | 45.5 °C                       |
| Fuel outlet pressure          | Ambient                | 120 kPa abs                   |
| Oxidant outlet pressure       | Ambient                | Ambient                       |
3.2.2. Load Cycling Durability Tests

As opposed to keeping the load constant, load cycling will put a strain on the catalyst, membrane, and carbon support materials. Typically, load cycling testing is done by cycling the stack current and recording the voltage responses. The duty cycle for stationary systems can vary depending on the applications (residential CHP, commercial power, standalone, or grid-connected). Cycling is supposed to be much less severe than transportation applications. With fewer on-off cycles and less time spent at OCV, start-up and transient operation would be easier [119].

While FCTESTNET’s dynamic load cycling was created specifically for transportation applications, it can also be used to assess the generic performance of PEMFCs [132]. The dynamic load cycling consists of a low power stationary step at $I = 0.2@20\% P_{\text{max}}$, followed by a ramp from $I_{\text{load min}}@20\% P_{\text{max}}$ to $I_{\text{load max}}@100\% P_{\text{max}}$. The total time of the test, $t_{\text{max}}$, is set between 500 and 10,000 h, and the interval is interrupted every $t_{\text{max}}/10$ h to record a polarization curve [120, 132]. The duration of the long-term durability test is determined in the same way as the steady test module.

The load-cycling durability test module was also added to the STACKTEST project with the goal of determining the effect of dynamic load changes on the irreversible performance loss of a fuel cell stack [142–144]. It is possible to use it to build a load profile. Figure 14 depicts an example load cycling profile for CHP systems. Extended profile times of up to 24 h, as well as long phase times, are characteristics of load profiles for stationary applications. The load profile shown is made up of nine stages, each with its own dwell time and stack load value. Since reformate-fuel gas is commonly used in CHP systems, the fuel gas composition should be simulated. If no other options are available, the composition can be set to: 30% CO$_2$, 5% CH$_4$, and 10 ppm of CO are added to hydrogen. It may use an extra moderate air bleed of 0.8% [144].

![Figure 14. Variation of the stack load for the CHP load profile.](image)

4. Conclusions

One of the most significant barriers to fuel cell commercialization is durability, as discussed in this review article. We present some of the most important factors influencing the long-term durability of PEMFCs in this study. Degradation of PEMFC materials, operating conditions, water control, and thermal management are among these factors. We discuss PEMFC durability test protocols for transportation applications developed by leading organizations, institutes, and universities around the world, such as the DOE, FCHEA, USDRIVE, UNECE, FC TestNet, IEC, and Japanese NEDO and Chinese universities including Tongji University, Tsinghua University, Wuhan University of Technology, and...
the Dalian Institute of Chemical Physics. These protocols were created to mimic real-world driving situations. However, they are not meant to be comprehensive for all driving situations encountered in the real world, such as long periods of idling due to traffic jams, etc. Researchers must make adjustments to adapt current operating conditions to test objectives. In addition, the protocols for reliability testing PEMFC stationary applications are briefly discussed. We hope that the results of this research will help to speed up the commercialization of PEMFCs.

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