

Calibration and Validation of a PEM Fuel Cell Hybrid Powertrain Model for Energy Management System Design

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




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Article

Calibration and Validation of a PEM Fuel Cell Hybrid Powertrain Model for Energy Management System Design

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Abstract

This paper presents a calibrated and dynamically responsive simulation framework for hybrid energy systems that integrate Proton Exchange Membrane Fuel Cells (PEMFCs) and batteries, targeting applications in light commercial vehicles (LCVs). The aim is to support the design and assessment of energy management strategies (EMS) under realistic operating conditions. A publicly available PEMFC model is used as the starting point. To improve its representativeness, calibration is performed using experimental polarization curve data, enhancing the accuracy of the stack voltage model, and the air compressor model—critical for maintaining stable fuel cell operation—is adjusted to reflect measured transient responses, ensuring realistic system behavior under varying load demands. Quantitatively, the calibration results are strong: the R^2 values of both the fuel cell polarization curve and the overall system efficiency are around 0.99, indicating excellent agreement with experimental data. The calibrated model is embedded within a complete hybrid vehicle powertrain simulation, incorporating longitudinal dynamics and control strategies for power distribution between the battery and fuel cells. Simulations conducted under WLTP driving cycles confirm the model's ability to replicate key behaviors of PEMFC-battery hybrid systems, particularly with respect to dynamic energy flow and system response. In conclusion, this work provides a reliable and high-fidelity simulation environment based on empirical calibration of key subsystems, which is well suited for the development and evaluation of advanced EMS algorithms.

Keywords: proton exchange membrane fuel cell; battery hybrid powertrain; energy management strategy; fuel cell calibration; light commercial vehicle



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1. Introduction

As global energy shortages and environmental concerns become increasingly pressing, fuel cell technology is gaining significant attention as a potential solution. A fuel cell is an electrochemical device that directly converts the chemical energy of hydrogen and oxygen into electrical energy through electrode reactions [1]. The main byproduct of this reaction is water, resulting in zero local emissions, which makes fuel cells a truly pollution-free energy source at the point of use [2]. Additionally, fuel cells are not constrained by the Carnot cycle, allowing for high energy conversion efficiencies that can reach between 60% and 80% [3].

To fully realize the potential of PEMFCs in automotive applications, accurate modeling and simulation are essential [4]. In practical scenarios, especially during the design and

testing of vehicle powertrains, it is neither efficient nor feasible to rely solely on experimental trials [5]. Such testing not only consumes considerable time and resources but also poses a risk of degradation to the fuel cell system [6]. A reliable simulation model enables efficient exploration of system behavior under various operating conditions, supporting the development of control strategies [7], energy management systems [8], and component integration [9].

Despite the growing body of literature on PEMFC modeling, two major limitations persist. First, many existing models are highly detailed [10], focusing extensively on the electrochemical processes [11,12] and auxiliary systems [13]. While informative, these models often require numerous physical parameters that are not always available in vehicle-level research. In such cases, engineers may have access only to basic datasets, such as polarization curves or rated performance points. Moreover, from a powertrain design perspective, the focus is generally on macroscopic input–output relationships—voltage, current, and power—rather than internal cell-level dynamics. Second, other models in the literature are oversimplified [14], with empirical or control-oriented formulations [15] that lack transparency and generalizability [16]. These approaches are often difficult to calibrate or adapt, even when basic reference data are available, thus limiting their practical utility in system-level applications [17].

In this study, a dynamic model of a complete proton exchange membrane fuel cell (PEMFC) system is developed to support energy management simulations. The model by Pukrushpan et al. [18], which includes a detailed cathode air supply loop from the compressor to the outlet manifold, is adopted as the baseline due to its comprehensive dynamic behavior. Building upon this foundation, the proposed model retains key transient characteristics while enhancing computational efficiency and facilitating integration into larger vehicle simulation frameworks, which includes a complete cathode air control loop, from the air compressor to the outlet manifold. The new model preserves dynamic features while maintaining computational efficiency and ease of integration.

To meet the performance requirements of vehicle propulsion and portable fuel cell applications, a fuel cell stack is commonly coupled with a battery via a DC/DC converter to form a hybrid power system. This configuration allows the system to leverage the strengths of both power sources, with the battery providing supplementary power during peak demands and enabling energy recovery through regenerative braking. Defining the performance parameters of the fuel cell and identifying potential limitations are critical steps in designing effective fuel cell hybrid power applications [19]. When data for battery and vehicle dynamics models are limited, a comprehensive methodology for modeling and simulating BEVs can be utilized to guide engineers and researchers in the early stages of EVs design [20–22]. This approach provides a robust and efficient framework for developing hybrid power systems and ensuring optimal performance [23]. A well-calibrated model enhances the accuracy of dynamic energy flow simulation, providing a more reliable representation of energy distribution over time [24]. Thus, this research lays crucial groundwork for advancing the understanding and optimization of hybrid energy management in fuel cell electric vehicles.

2. Fuel Cell and Battery Hybrid Vehicle Modeling

This chapter presents the modeling approach adopted for a hybrid electric vehicle powered by a fuel cell and a battery system. The primary objective is to build a simulation platform capable of supporting the development and evaluation of energy management strategies, with a particular focus on system efficiency and vehicle range. To this end, this chapter begins with a brief description of the fuel cell model, including essential calibration to ensure realistic performance representation. Following this, a simplified

modeling method is introduced for the battery pack and electric drive, suitable for early-stage simulations where rapid evaluation is prioritized over detailed physical fidelity. Finally, the vehicle’s longitudinal dynamics are described along with the implementation of a basic longitudinal controller. This structure ensures that all powertrain components are integrated within a consistent framework, enabling the effective assessment of energy flows and system behavior under various operating conditions.

2.1. Fuel Cell System

There are various approaches to modeling fuel cells. One common method is to represent the fuel flow as a function of power and efficiency, where efficiency is treated as a variable dependent on power output [14]. Alternatively, methods like response surface modeling can capture input–output relations but are impractical without sufficient experimental data [16]. Although this approach is simple, it lacks dynamic behavior, as it assumes fixed voltage and current values for each power level, while the model proposed in this paper is capable of simulating smooth transitions in voltage and current in response to changes in power demand, effectively avoiding abrupt discontinuities. Moreover, detailed models focusing on mass transfer or gas channel behavior in each layer of the fuel cell stack are computationally intensive and therefore unsuitable for powertrain energy management analysis, as such models require lots of physical parameters of FC stack and long simulation times to estimate total driving range [11,12].

A fuel cell system typically comprises four subsystems: hydrogen circuit, air circuit, coolant circuit, and humidifier circuit. Since most fuel cells use deionized water as the coolant, the coolant and humidifier subsystems are often integrated into a single water circuit. This chapter outlines the modeling principles of the key components involved in the hybrid fuel cell system, drawing upon the detailed framework proposed by Pukrushpan et al. [18]. While their model provides a well-established foundation, certain subsystems—particularly those affecting energy flow and control dynamics—require refinement to better support the objectives of this study.

Emphasis is placed on the stack voltage model and the air compressor model, which significantly influence system performance and energy management strategies. These components have been modified to improve numerical stability, capture transient behaviors more accurately, and better align with the hybrid architecture used in this study. For a comprehensive description of the complete PEMFC system, readers are referred to the original work by Pukrushpan et al. [18].

2.1.1. Stack Voltage Model

Stack voltage is calculated as the product of the number of cells by their individual voltage. The combined effect of thermodynamics, kinetics and ohmic resistance determines the output voltage of the cell v_{fc} as defined by [1]:

$$v_{fc} = E - v_{act} - v_{ohm} - v_{conc} \tag{1}$$

where E is the open circuit voltage, v_{act} is the activation loss, v_{ohm} is the ohmic loss, and v_{conc} is the concentration loss. The Nernst equation describes the theoretical open-circuit voltage of a fuel cell [7]:

$$E = -\frac{\Delta G_{react}}{2F} + \frac{R_{gas} \cdot T}{2F} \cdot \ln\left(\frac{p_{H_2} \cdot p_{O_2}^{0.5}}{p_{H_2O}}\right) \tag{2}$$

where ΔG_{react} is the Gibbs free energy, F is Faraday’s Constant, R_{gas} is the gas constant, T is expressed in Kelvin and p_{H_2} , p_{O_2} and p_{H_2O} are expressed in atm. In real applications, the

OCV is lower than the theoretical Nernst voltage due to additional losses such as hydrogen crossover and internal short circuits [25]. However, these effects will not be considered in this work.

As indicated by Equation (2), the operating temperature of the fuel cell stack has a significant impact on its performance. But due to the absence of a thermal management loop in the model and the lack of sufficient temperature-related experimental data for validation, the fuel cell temperature is assumed to remain constant at 80 °C throughout this study.

Activation loss arises from the energy required to drive electrochemical reactions at the electrodes, involving electron transfer and bond breaking/forming. While both electrodes contribute, the loss is primarily dominated by the slower oxygen reduction reaction at the cathode, as the hydrogen oxidation at the anode proceeds relatively quickly. The relation between the activation voltage drops and the current density is described by the Tafel equation [17]:

$$v_{act} = b \ln \left(\frac{i}{i_0} \right) \tag{3}$$

where i is the current density, i_0 is the exchange current density, and b is the Tafel slope. Though the Tafel equation was originally concluded from experimental data, it also has a theoretical foundation, that for a hydrogen fuel cell with two electrons, the constant b is given by:

$$b = \frac{RT}{2\alpha F} \tag{4}$$

where the constant α is called the charge transfer coefficient and is the proportion of the electrical energy applied that is harnessed in changing the rate of an electrochemical reaction. For the hydrogen electrode, its value is about 0.5 for a great variety of electrode materials. At the oxygen electrode, its value is between about 0.1 to 0.5 in most cases [1].

Meanwhile, the exchange current density i_0 is crucial in controlling the performance of the fuel cell electrode. The i_0 at cathode is much smaller than that at anode. Since the smaller the i_0 , the greater the voltage drop, the voltage drop at the anode is negligible compared to that of the cathode [1,2].

Ohmic loss is due to the resistance of the polymer membrane to the transfer of protons and the resistance of the electrodes and the collector plate to the transfer of electrons. The voltage drop that corresponds to the ohmic loss is proportional to the current density:

$$v_{ohm} = i \cdot \frac{t_m}{\sigma_m} \tag{5}$$

where t_m is the thickness of the membrane and the membrane conductivity, σ_m is a function of membrane water content, λ_m and fuel cell temperature [26]:

$$\sigma_m = b_1 \cdot \exp \left(b_2 \left(\frac{1}{303} - \frac{1}{T_{fc}} \right) \right) \tag{6}$$

$$\begin{cases} b_1 = 0.005139\lambda_m - 0.00326 \\ b_2 = 350 \end{cases}$$

Since membrane conductivity depends on water content and temperature—both fixed by the model—the only adjustable parameter to tune ohmic voltage loss is membrane thickness, t_m . Industry-optimized Nafion membranes typically range from 20 to 50 μm in thickness.

Concentration losses, also known as mass transport losses, arise from limitations in the diffusion of reactants to the catalyst sites as the electrochemical reaction proceeds. These

losses become increasingly significant at high current densities, where the depletion of reactant concentration near the electrode surface leads to a steep voltage drop. While classical models describe this phenomenon using logarithmic relations derived from the Nernst equation, the empirical expression presented in Equation (7) offers a better fit for the nonlinear voltage behavior observed at high current densities [27]:

$$v_{conc} = i \left(c_2 \frac{i}{i_{lim}} \right)^{c_3} \tag{7}$$

where c_2 , c_3 and i_{lim} are constants that depend on the temperature and the reactant partial pressure and can be determined empirically. The parameter i_{lim} is the current density that causes precipitous voltage drop. This formulation is particularly useful in dynamic system simulations, where accurately capturing performance degradation underload is critical.

2.1.2. Calibration of Stack Voltage Model

The tunable parameters in the stack voltage model are b , i_0 , t_m and i_{lim} , which are calibrated in the MATLAB/Simulink R2024b environment using the Parameter Estimator toolbox and Levenberg–Marquardt algorithm to minimize the cost function of sum square error between simulated and reference data [28,29].

Initial values are taken from the literature or based on information from datasheets; for example, platinum is considered the most common material as a catalyst. Table 1 illustrates the impact of each parameter, offering a general sense of their relative scales.

Table 1. Tunable parameters and tuning strategies.

Parameter	Range	Influence *
b [V]	0.03–0.15 [2]	Positive to Activation loss
i_0 [mA/cm ²]	around 0.5 [1]	Negative to Activation loss
t_m [μm]	20–50 [30]	Positive to Ohmic loss
i_{lim} [A/cm ²]	2.0–3.0 [31]	Negative to Concentration loss

* Positive influence refers that increasing the parameter will increase the voltage drop correspondingly, vice versa.

2.1.3. Compressor Model

The only dynamic state in the model is the compressor speed, ω_{cp} . The inputs to the model include inlet air pressure, $p_{cp,in}$, its temperature, $T_{cp,in}$, voltage command to compressor motor, v_{cm} , and downstream pressure, which is the supply manifold pressure, $p_{cp,out} = p_{sm}$. The inlet air is typically atmospheric, and its pressure and temperature are assumed to be fixed at $p_{atm} = 1$ atm and $T_{atm} = 25$ °C, respectively. The motor command is one of the inputs to the fuel cell system. The downstream pressure is determined by the supply manifold model.

The compressor model was well designed [7,18] with a lumped rotational model to represent the dynamic behavior of the compressor:

$$J_{cp} \frac{d\omega_{cp}}{dt} = (\tau_{cm} - \tau_{cp}) \tag{8}$$

where J_{cp} is the combined inertia of the compressor and the motor, ω_{cp} is the compressor speed, τ_{cm} is the compressor motor torque input, τ_{cp} is the torque required to drive the compressor. The compressor motor torque is calculated using a static motor equation:

$$\tau_{cm} = \eta_{cm} \frac{k_t}{R_{cm}} (v_{cm} - k_v \omega_{cp}) \tag{9}$$

where k_t , R_{cm} and k_v are motor constants and η_{cm} is the motor mechanical efficiency.

A lookup table of the compressor efficiency, η_{cp} , is used to find the efficiency of the compressor from the mass flow rate and pressure ratio across the compressor. The maximum efficiency of the compressor is 80%. The torque required to drive the compressor is calculated using the thermodynamic equation [32]:

$$\tau_{cp} = \frac{C_p}{\omega_{cp}} \frac{T_{atm}}{\eta_{cp}} \left[\left(\frac{p_{sm}}{p_{atm}} \right)^{\frac{\gamma-1}{\gamma}} - 1 \right] W_{cp} \tag{10}$$

where τ_{cp} is the torque needed to drive the compressor, C_p is the specific heat capacity of air = 1004 J/(kg·K), and γ is the ratio of the specific heats of air = 1.4.

A static compressor map is used to determine the air flow rate through the compressor, W_{cp} by the Jensen and Kristensen nonlinear curve fitting method [32]. The compressor model used here is for an Allied Signal compressor [33].

2.1.4. Calibration of Compressor Model

Compressor power can be calculated by Equation (10). Notably, the pressure ratio in the model increases along with the air mass flow rate, making it infeasible to calibrate both the compressor power and the mass flow rate simultaneously.

To reflect variations in the inlet condition of the compressor, which are the inlet flow pressure and temperature, the “corrected” values of mass flow rate, W_{cr} and compressor speed, N_{cr} are used in the compressor map. Using the Jensen and Kristensen method, the mass flow rate is considered as a function of pressure ratio, $\frac{p_{sm}}{p_{atm}}$ and motor speed, N_{cm} , with a group of parameters, a_i , b_i , and c_i , which are regression coefficients obtained by curve fitting of the compressor data [7], while the compressor data is calculated by the limited experimental data based on the following order:

1. Use the reference load I_{st} to obtain the nominal mass flow rate of the compressor W_{cp} ;
2. Input W_{cp} in the FC model to collect the corresponding pressure ratio;
3. Calculate compressor power P_{cp} with W_{cp} and pressure ratio, assuming $P_{cp} = P_{cm}$;
4. Obtain the compressor motor parameters from a map incorporating P_{cm} vs. N_{cm} ;
5. Create a new compressor map with pressure ratio, N_{cm} and W_{cp} .

After having the new compressor data, a_i , b_i , and c_i is obtained from the ‘Parameter Estimator’ in Simulink and listed in Table 2.

Table 2. Compressor map regression coefficients.

Parameter	Value	Parameter	Value
a_4	-8.110×10^{-7}	b_0	2.556
a_3	5.588×10^{-6}	c_5	-1.298×10^{-8}
a_2	3.508×10^{-6}	c_4	-1.298×10^{-8}
a_1	2.676×10^{-7}	c_3	0.215
a_0	4.821×10^{-4}	c_2	0.799
b_2	8.889×10^{-4}	c_1	81.190
b_1	0.111	c_0	-1.449

After updating the compressor model, the responses in terms of air flow rate and pressure were successfully corrected. However, a persistent deviation remained in the power consumption output, which could not be resolved due to the lack of more detailed system information. To address this issue, a correction function was applied between the actual power consumption and the reference power. This adjustment allowed the final output of the net fuel cell power to align with the expected values, ensuring consistency in system-level simulation results.

In the implemented fuel cell (FC) model, the control input is the current, whereas the energy management system (EMS) determines the power request as its output. To bridge this interface, the current input to the fuel cell at a given time step t , denoted as $i_{FC}(t)$, is computed based on the power demand $P_{req}(t)$ and the fuel cell voltage $v_{FC}(t - 1)$ from the previous time step [34].

$$i_{FC}(t) = \frac{P_{req}(t)}{v_{FC}(t - 1)} \tag{11}$$

This formulation is based on the assumption that the voltage variation of the fuel cell is relatively slow compared to the simulation time step, thereby justifying the use of the previous voltage value in the calculation.

2.2. Battery Pack

Various modeling approaches exist for battery packs, including electrochemical models [35,36], thermal–electrochemical-coupled models [37], and equivalent circuit models [38]. Electrochemical models, while offering high accuracy and detailed insight into internal reactions, typically require extensive parameter identification and computational resources, making them less practical in early-stage studies with limited data availability. In this work, an equivalent circuit model is adopted due to its balance between simplicity and effectiveness, which also enables rapid development and integration into the vehicle powertrain simulation, making it well suited for preliminary analysis and energy management strategy evaluation.

The battery pack consists of battery cells in parallel and in series, and the power of the battery pack is:

$$P_{batt,pack} = P_{batt,cell} \cdot N_p \cdot N_s \tag{12}$$

where the $P_{batt,pack}$ is the power of the battery pack and the $P_{batt,cell}$ is the power of a single battery cell. N_p is the number of cells in parallel and N_s is the number of cells in series. The battery current of a single cell is determined by an internal resistance model:

$$v_{batt,cell} = v_{nom} - i_{batt,cell} \cdot R_{int} \tag{13}$$

$$P_{batt,cell} = v_{batt,cell} \cdot i_{batt,cell} \tag{14}$$

where the battery open circuit voltage v_{nom} and the internal resistance R_{int} are static functions of the battery state of charge and temperature. The voltage and current of the battery pack is:

$$\begin{cases} v_{batt,pack} = v_{batt,cell} \cdot N_s \\ i_{batt,pack} = i_{batt,cell} \cdot N_p \end{cases} \tag{15}$$

where the $v_{batt,pack}$ and $i_{batt,pack}$ is the voltage and current of the battery pack, respectively. The SOC is based on Coulomb counting [39] and calculated from the battery current and the capacity of the battery, k_{batt} and the battery charging efficiency, η_{batt} :

$$\dot{SoC} = \begin{cases} k_{batt} \cdot i_{batt,pack} , i_{batt,pack} \geq 0 \\ -\eta_{batt} \cdot k_{batt} \cdot i_{batt,pack} , i_{batt,pack} < 0 \end{cases} \tag{16}$$

The pack dis- and charging current limit are also from a lookup table, which is a function of SoC and Temperature. In this model, the temperature is constant, which equals to the ambient temperature of 25 °C. The calculation of the battery’s State of Charge (SoC) is determined by taking the initial SoC value SoC_0 and subtracting the cumulative energy consumption over time.

$$SoC(t) = SoC_{t_0} - \int_{t_0}^t \dot{SoC} \cdot d\tau \tag{17}$$

2.3. Electric Drive

Electric motor modeling methods can generally be categorized into dynamic and static approaches. Dynamic modeling is based on electromagnetic principles, typically employing mathematical representations such as the d-q axis model to capture the motor’s transient behavior under various operating conditions [40,41]. In contrast, static modeling simplifies internal dynamics and commonly uses an efficiency map to describe the motor’s performance as a function of speed and torque [42,43]. This approach is well suited for system-level energy flow analysis and vehicle-level simulations. Each method involves a trade-off between computational complexity and modeling accuracy, and the choice depends on the specific application context.

From the efficiency map implemented in the model, η_{em} is a function of torque input and rotational speed. The overall efficiency is the product of the motor efficiency, η_{motor} , and the inverter efficiency, η_{inv} . The rotational speed of electric drive is calculated by:

$$\omega_{ed} = \frac{V}{R_e} \cdot \tau_{trm} \tag{18}$$

where V is vehicle longitudinal speed, R_e is the wheel radius and τ_{trm} is the transmission ratio. And the torque input is calculated by:

$$Trq_{eq} = P_{eq} \cdot \omega_{ed} \tag{19}$$

$$P_{eq} = P_{DC} - P_{aux} \tag{20}$$

where ω_{ed} is the rotational speed of electric, P_{eq} is the equivalent power input, which equals the DC power input, P_{DC} minus the power consumed by the auxiliary components, P_{aux} . Depending on the specific driving conditions, the electric drive performs as the generator or motor [43], and the torque output is:

$$\begin{cases} Trq_{act} = Trq_{eq} \cdot \eta_{em} , & Trq_{eq} \geq 0 \\ Trq_{act} = Trq_{eq} / \eta_{em} , & Trq_{eq} < 0 \end{cases} \tag{21}$$

where Trq_{act} is the actual torque output of the electric drive block to VDS for the calculation. η_{em} is the total efficiency, considering both the efficiency of motor and inverter.

2.4. Vehicle Dynamic System

Vehicle dynamic systems form the foundation for understanding and optimizing automotive performance, safety, and control. These systems describe how a vehicle responds to driver inputs and external forces in the longitudinal, lateral, and vertical directions. Accurate dynamic modeling enables the development of advanced control strategies for stability, ride comfort, and energy efficiency, especially important in modern applications such as electric vehicles and autonomous driving. From the classical single-track model to more complex multi-body simulations, vehicle dynamics encompasses suspension behavior, tire–road interaction, and drivetrain response.

Vehicle motion is opposed by several resistive forces that the powertrain must overcome, including aerodynamic drag, rolling resistance, grade force, and transmission losses. Considering a combination of aerodynamic drag and rolling resistance, the longitudinal resistive force is a quadratic function of velocity [27]:

$$F_{xr} = A + B \cdot V + C \cdot V^2 \tag{22}$$

where the coefficients A , B , and C are determined experimentally by fitting the data to reflect real-world conditions. This quadratic relationship highlights how resistance components scale with speed, significantly impacting vehicle dynamics and energy consumption. The grade force is the component of the resistance to motion of a vehicle that depends directly on the slope of the road, where α is the road slope:

$$R_{grad} = mg \sin \alpha \tag{23}$$

To correctly calculate the acceleration of the vehicle, the inertial contribution of the rotating mass also needs to be considered. Using the energy method, an expression for the “equivalent mass” can be written based on the relationship between rotational speed and vehicle speed [21]:

$$m_e = m + \frac{J_w}{R_e^2} + \frac{J_t}{R_e^2 \tau_f^2} + \frac{J_m}{R_e^2 \tau_f^2 \tau_g^2} \tag{24}$$

where J_w , J_t and J_m are the moments of inertia of wheel, transmission and motor, respectively. R_e is the effective wheel radius and τ_f and τ_g are the final and gearbox transmission ratios.

The net torque passes through the drivetrain and is converted into linear force by dividing it by the wheel radius. This force governs the vehicle’s acceleration, calculated by dividing the force by the vehicle’s equivalent mass. This relationship is expressed as:

$$a_v = \frac{F_{rem}}{m_e} = \frac{Trq_{rem}}{m_e \cdot r_w} \tag{25}$$

where F_{rem} and Trq_{rem} , are the remaining force and torque applied on the wheel, the r_w is the radius of wheels, m_e is the equivalent mass of the vehicle.

The input torque to the tire system is determined by a torque saturator, combining motor torque (scaled by the transmission ratio) and friction braking torque [44,45]. A torque limit ensures safe operation within physical and tire–road interaction constraints. Considering a rear-wheel-drive configuration, the torque limit under specific acceleration and slope conditions is [46]:

$$Trq_{lim} = (a_v \cdot h_{CG} \cdot M/b + Mg \cdot a/l \cdot \cos \alpha) \cdot \mu_x \cdot r_w \tag{26}$$

where h_{CG} is the height of the center of gravity, M is the vehicle mass, a , b and l represent wheelbase parameters, g is gravitational acceleration, α is the road slope angle, μ_x is the road-tire friction coefficient, and r_w is the wheel radius.

2.5. Longitudinal Vehicle Controller

Various types of longitudinal controllers have been proposed to regulate vehicle speed and ensure safe following distances, which include Model Predictive Control [47,48], fuzzy logic-based controllers [49], and adaptive cruise control systems using PID or hybrid approaches [50]. Since this study does not focus on the development of longitudinal control strategies, a simple and effective Proportional–Integral (PI) controller is employed to manage vehicle speed.

In powertrain simulations, the longitudinal vehicle controller acts as a virtual driver, controlling speed and acceleration to match a target cycle like WLTP. It compares the reference and actual speeds to compute the required acceleration or deceleration, which is then converted into a torque demand by the motor torque reference generator. An anti-saturation PI controller ensures stable control, with the output acceleration signal defined as [51]:

$$cmd(t) = V_{ref}(t) \cdot \frac{K_{ff}}{v_{nom}} + e_v(t) \cdot \frac{K_p}{v_{nom}} + \int_0^t \left(e_v(\tau) \cdot \frac{K_i}{v_{nom}} + e_{cmd}(\tau) \cdot K_{aw} \right) \cdot d\tau + \alpha(t) \cdot K_g \quad (27)$$

where V_{ref} is the reference vehicle speed, cmd is the controller command, of which the subscripts *accel*, *decel* stand for acceleration and deceleration. α is the road grade. e_v is the error of vehicle speed and e_{cmd} is the error of controller command. K_{ff} is the coefficient of forward velocity, K_p is the coefficient of proportional gain, K_i is the coefficient of integral gain, K_{aw} is the coefficient of anti-windup and K_g is the coefficient of feedforward grade angle.

During light braking, the motor handles deceleration to recover energy. For rapid deceleration, mechanical brakes assist by adding torque. To enhance efficiency and reduce brake wear, the control strategy favors motor braking whenever possible.

The motor torque reference value Trq_{ref} is obtained by the saturator. It is calculated by the acceleration or deceleration command,

$$Trq_{ref} = cmd_{accel} \cdot Trq_{peak} + cmd_{decel} \cdot Trq_{const} \cdot R_{lim} \quad (28)$$

where Trq_{peak} is the maximum output torque for acceleration and Trq_{const} is the continuous output torque for deceleration. The regeneration coefficient limit factor R_{lim} used in the case studied in the next chapter is 0.5.

2.6. Energy Management Strategy

EMS strategies can be classified into rule-based, optimization-based, and learning-based methods. Rule-based approaches rely on heuristic if-then rules derived from engineering intuition and are easy to implement but often suboptimal [52]. Optimization-based methods [53] aim to minimize a cost function over a prediction horizon. More recently, learning-based strategies, such as reinforcement learning [54], have emerged, enabling real-time adaptive control under uncertain conditions. Each method involves trade-offs among computational complexity, adaptability, and optimality. In this study, to focus on model validation, a rule-based approach, Baseline A, is adopted for its simplicity and practicality.

Baseline A is designed around two critical metrics: the State of Charge (SoC) and the H₂ Inventory (SoH₂). This framework categorizes the vehicle’s power demands into three distinct operational modes, each tailored to optimize the use of energy sources while ensuring efficient performance under varying conditions:

- **Battery Depleting Mode:** To prevent overcharging in a battery system, when the SoC level is higher than 85%, the system relies primarily on the battery to meet power demands. This strategy aims to prevent overcharging the battery:

When $P_{req} < P_{batt,max}$

$$\begin{cases} P_{batt} = P_{req} \\ P_{fc} = 0 \end{cases} \quad (29)$$

When $P_{req} > P_{batt,max}$

$$\begin{cases} P_{batt} = P_{batt,max} \\ P_{fc} = P_{req} - P_{batt,max} \end{cases} \quad (30)$$

where P_{batt} is the power split of battery system and P_{fc} is the power split of fuel cell system. $P_{batt,max}$ is the maximum power possibly delivered by the battery system.

- **Battery Sustaining Mode:** When the SoC falls below 80%, without risking overcharge, Baseline A enters battery sustaining mode. The 80–85% window is implemented to avoid oscillations during mode transitions. In this phase, the fuel cell output is

adjusted in real time to meet power demands, while additional energy is used to recharge the battery, maintaining SoC and improving overall efficiency.

$$\begin{cases} E_{DC}(t-1) = \int_0^{t-1} P_{DC}(\tau) \cdot d\tau \\ E_{fc}(t-1) = \int_0^{t-1} P_{fc}(\tau) \cdot d\tau \end{cases} \quad (31)$$

$$\begin{cases} P_{fc}(t) = K_p \cdot (E_{DC}(t-1) - E_{fc}(t-1)) + K_i \cdot \int_0^t (E_{DC}(\tau-1) - E_{fc}(\tau-1)) \cdot d\tau \\ P_{batt}(t) = P_{req}(t) - P_{fc}(t) \end{cases} \quad (32)$$

where E_{DC} is the energy consumed by the DC power system, and E_{fc} is the energy generated by the fuel cell block.

- Pure Electric Mode: As hydrogen depletes, the fuel cell shuts down and the vehicle enters pure electric mode, relying solely on the battery for propulsion.

$$\begin{cases} P_{batt} = P_{req} \\ P_{fc} = 0 \end{cases} \quad (33)$$

This transition is key to maintaining vehicle operation as hydrogen depletes, enhancing flexibility and reliability. Baseline A structures the energy management process to optimize efficiency and performance, enabling intelligent adaptation to varying conditions and improving sustainability in fuel cell electric vehicles.

3. Simulation Results

This chapter presents the results of two simulation stages. First, the performance of the calibrated fuel cell model is shown, demonstrating a high level of agreement with the reference experimental data. Second, the behavior of the complete powertrain model, equipped with a simple energy management algorithm, is illustrated over a WLTC driving cycle. The purpose here is not to evaluate the effectiveness of the energy management strategy, but rather to show that the model can reasonably represent the behavior of each subsystem.

3.1. Calibration Results

In the simulation, the reference data comes from an 80-kW FC that comprises 440 individual cells, each with an active area of 270 cm². This reference data has been obtained through empirical testing of a fuel cell

There are two polarization curves in Figure 1, the blue curve represents the data obtained from the reference values and the red curve corresponds to the results from the calibrated model, between which it demonstrates a strong agreement. Quantitatively, the model demonstrates a high level of accuracy with an R² value of 0.99. Additionally, the Relative Root Squared Mean Error (RRSME), which is normalized by the actual data values to ensure it is dimensionless and scale-independent, is calculated to be 0.013.

$$RRSME = \sqrt{\frac{1}{n} \sum_{i=1}^n (y_i - \hat{y}_i)^2} / \sqrt{\frac{1}{n} \sum_{i=1}^n y_i^2} \quad (34)$$

where y_i is actual value, \hat{y}_i is predicted value and n is the number of data points. The closer the RRSME value is to zero, the better the prediction is.

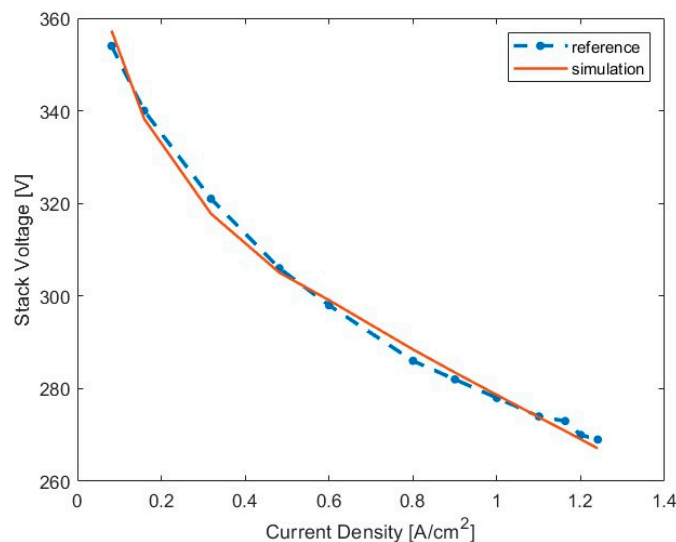


Figure 1. Comparison of polarization curves.

Slight overestimation at low current densities is likely due to the limited accuracy of the Tafel model in this region. At high current densities, the model exhibits a notable decline, primarily resulting from the sharp increase in concentration losses beyond 1 A/cm². The concentration loss model is difficult to calibrate due to its dependence on product-specific empirical formulas and parameters, which are often unknown.

Minor differences are present in the polarization curves but are barely visible in Figure 2, and the power–current calibration remains accurate at all current densities, which is critical for power-based drive cycle simulations.

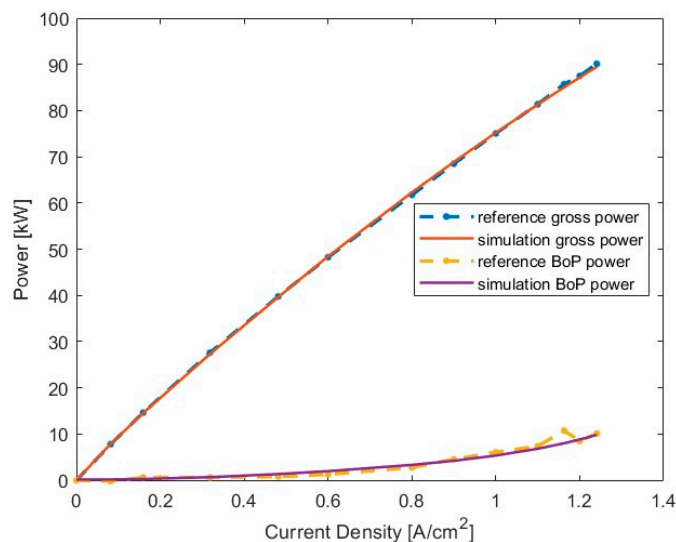


Figure 2. Comparison of power outputs.

However, Figure 2 also compares the auxiliary power output, showing that the reference data exhibits instabilities above 1 A/cm², likely due to real-world factors such as temperature variations and flow rate changes, highlighting the challenges of simulating high current density behavior.

The comparison between the reference data and the actual model efficiency shown in Figure 3 is primarily used to evaluate the correlation and trends between the two sets of data. Quantitatively, the RRSME is 0.012 and the R² is 0.989. Due to the inherent instability in fuel cell operation, if only the main operating range between 10 kW and 70 kW is considered,

it brings a better consistency. This analysis is critical to verifying the performance of the model and ensuring that it accurately reflects the operating characteristics of the fuel cell system.

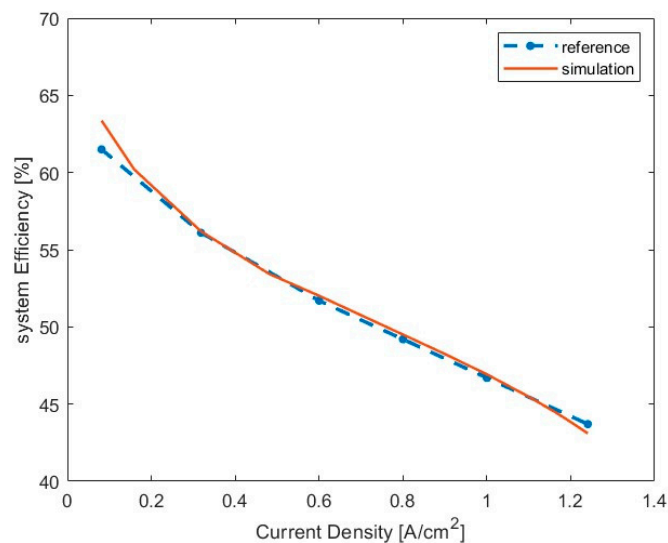


Figure 3. Fuel cell total efficiency comparison.

3.2. Drive Cycle Simulation

In the simulation, the WLTC was chosen due to its widespread acceptance across the automotive industry and its alignment with regulatory standards. Among the drive cycle tests, different initial SoC are chosen, while the initial SoH₂ remains the same, as it does not affect the fuel cell performance.

3.2.1. Battery Depleting Mode

As it is possible to see in Figure 4, the SoC decreases from 0.9 to approximately 0.75 over the test cycle, indicating the main operation in battery-depleting mode. During the initial phase, mild driving conditions allow the battery to meet power demand alone, keeping the fuel cell inactive and the H₂ tank level constant. However, when the SoC is less than 0.8, EMS switches from depleting mode to sustaining mode, which will be discussed in the next chapter.

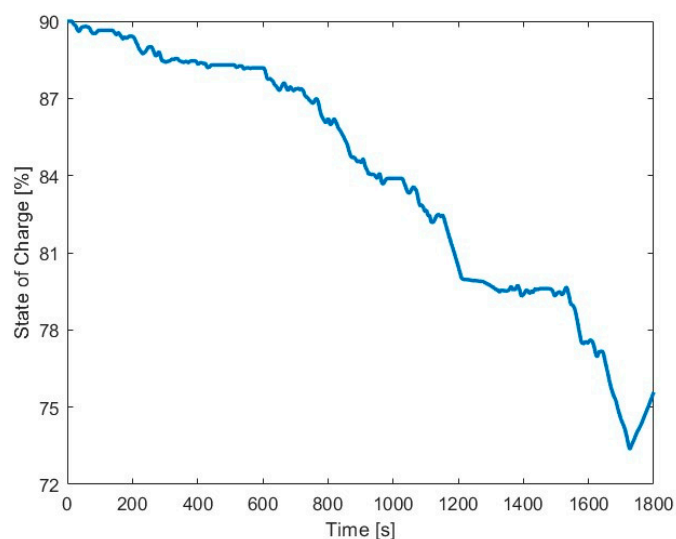


Figure 4. SoC variation in battery depleting mode.

Figure 5 demonstrates that the battery pack is sufficiently capable of meeting the power demand throughout the entire driving cycle, and the fuel cell system remains deactivated when the EMS operates in battery-depleting mode. The H₂ inventory is a number from 0 to 1 indicating the fraction of remaining hydrogen in the tank.

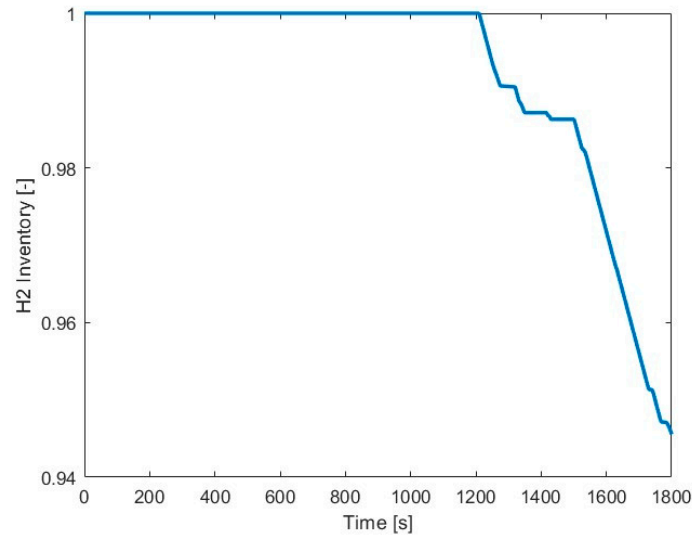


Figure 5. H₂ inventory variation in battery depleting mode.

The diagram in Figure 6 compares the reference speed with the actual vehicle speed to evaluate its ability to respond to varying driving conditions. It is observed that during the battery depleting mode, the actual vehicle speed closely aligns with reference speed. This behavior highlights the battery’s capability to effectively meet dynamic power demands. The high responsiveness of the battery ensures that the vehicle maintains the desired performance without significant deviations from reference speed.

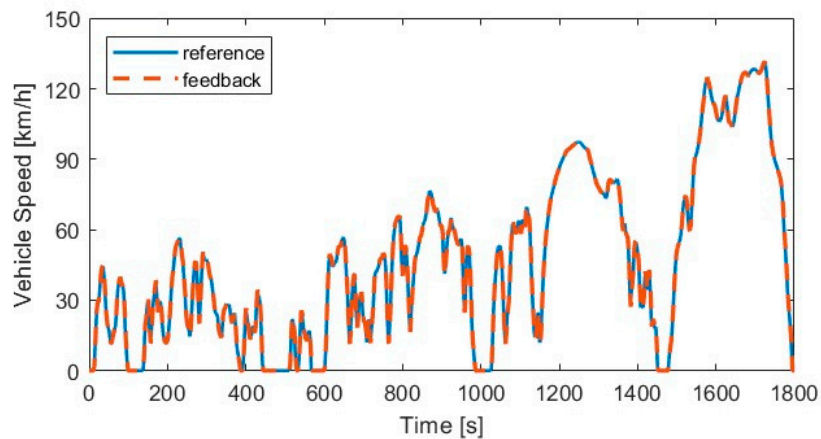


Figure 6. Vehicle speed comparison in battery depleting mode.

3.2.2. Battery Sustaining Model

Figure 7 illustrates the variation in SoC during a WLTP cycle for three different initial SoC values: 0.6, 0.4, and 0.2. To facilitate a direct comparison, Figure 7 presents the change in SoC rather than the absolute SoC values, allowing for a clearer evaluation of how battery depletion evolves from different starting points. The results indicate that the EMS, when operating in battery-sustaining mode, aims to keep the SoC relatively stable over the course of the driving cycle.

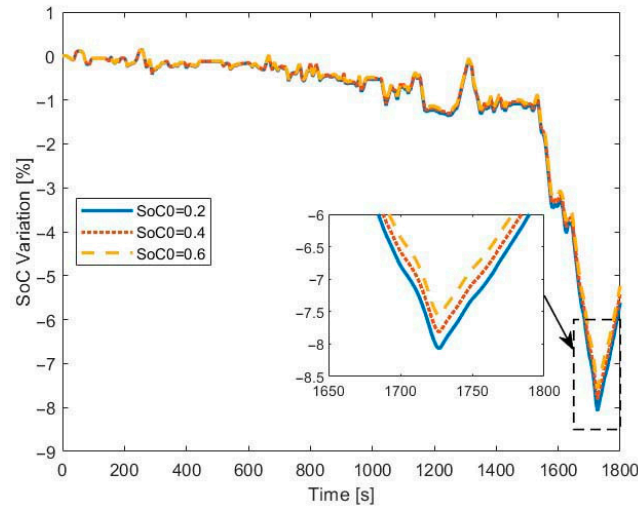


Figure 7. SoC variation with different initial values.

Although this stabilizing behavior is less obvious when the battery operates at low power outputs, a more evident trend emerges during high power demands, which confirms that the simulation model effectively captures the relationship between battery energy levels and performance. As the initial SoC decreases, the battery experiences a drop in nominal voltage, leading to reduced efficiency and increased depletion rates [55].

As the power demand increases, particularly during periods of high-power requirements, the SoC begins to exhibit a noticeable decline. This drop indicates a greater reliance on the battery to meet the immediate surge in power needs. The temporary reduction in SoC during these moments underscores the battery’s role as the primary energy source when rapid power delivery is required, while the increase in the SoC at the end shows the ability of FC to recharge the battery pack.

Meanwhile, Figure 8 suggests a continuous consumption of hydrogen throughout the cycle, indicating that the fuel cell remains consistently engaged. The strong overlapping in the curves shows that hydrogen consumption remains the same value under different initial SoCs with this EMS. In this mode, the fuel cell serves as the primary power source, while the SoC is maintained at a relatively high level. As the SoC decreases, the battery’s nominal voltage slightly drops, though this effect is not obvious; keeping the battery at a higher SoC helps optimize its performance and contributes to the overall efficiency of the powertrain.

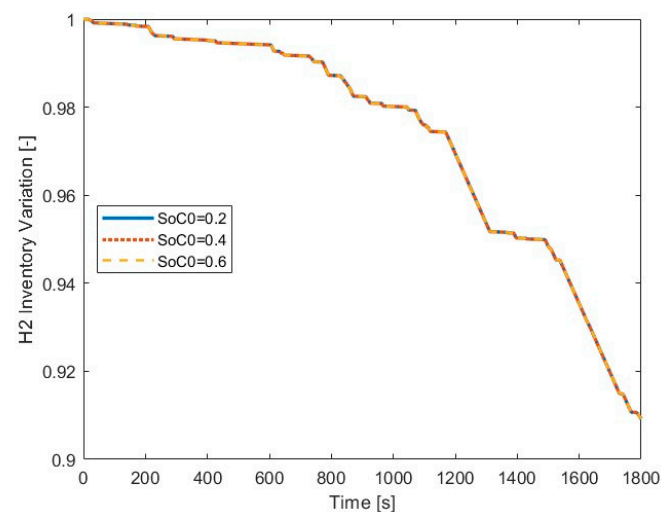


Figure 8. SoH₂ variation in a single drive cycle.

Figure 9 shows a perfect consistency between reference speed and the feedback speed. According to the reference data, the battery pack has a large capacity and wide power output range, making it a highly responsive energy source capable of meeting a wide range of power demands. This characteristic provides significant flexibility for the energy management algorithm, allowing future strategies to focus more on optimizing the fuel cell's operating range, thereby enhancing the overall system efficiency.

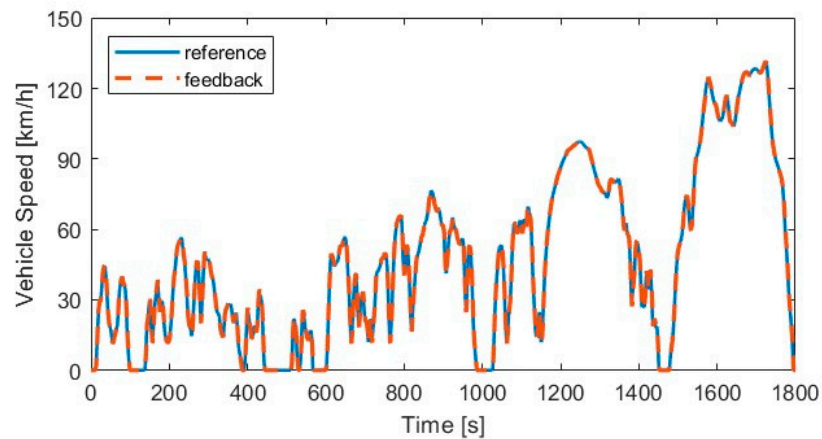


Figure 9. Vehicle speed comparison in battery-sustaining mode.

3.2.3. Repetitive Drive Cycles

A repetitive WLTC simulation, starting from a 90% SoC and a full hydrogen tank to a 0% SoC and an empty hydrogen tank, offers a comprehensive overview of the powertrain system and serves as a key method for evaluating EMS performance, particularly in terms of total driving distance.

Figures 10 and 11 provide a general idea of the entire behavior of the battery pack and the hydrogen tank. Following battery-sustaining mode, fuel cells become the primary energy source. As a result, battery discharge occurs more gradually, allowing the SoC to be maintained at a relatively high level. This operating condition enhances the efficiency of the battery system, as higher SoC levels are generally associated with improved energy conversion performance [38,55].

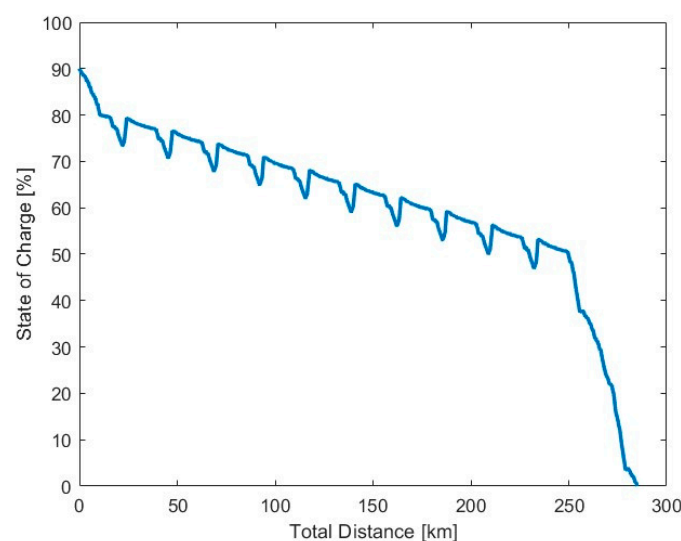


Figure 10. Full discharging profile.

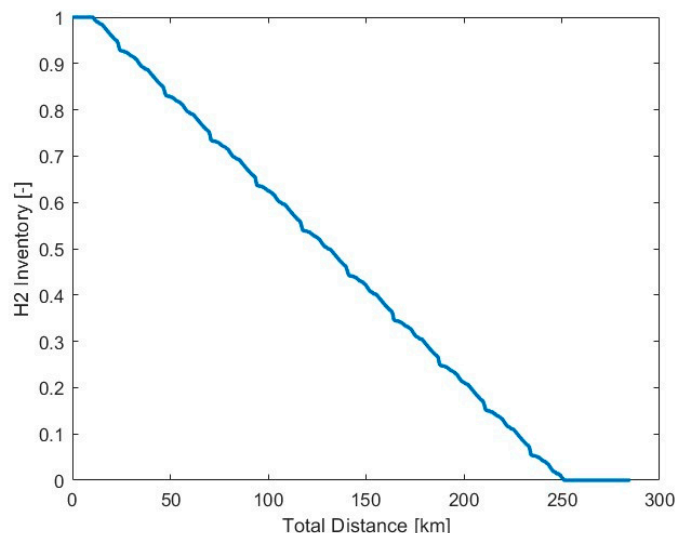


Figure 11. Hydrogen consumption profile.

Figure 12 illustrates the efficiency of the FC system. The simulation results demonstrate that the FC efficiency largely aligns with the reference curve. However, the presence of working points below the reference indicates transient operating phases that should be minimized.

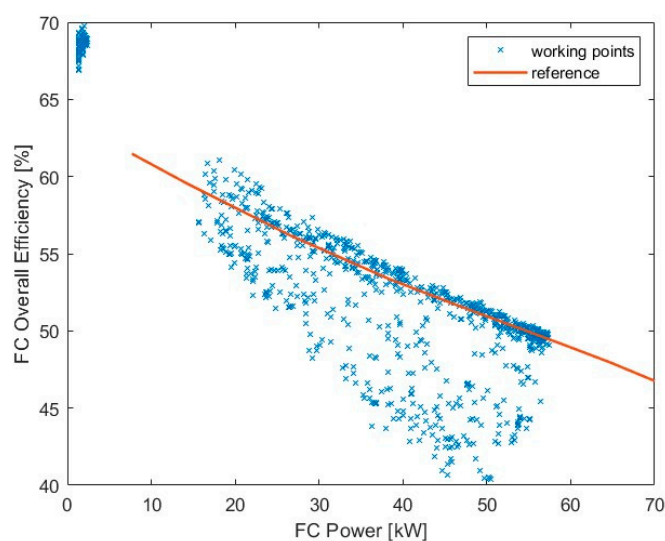


Figure 12. Real system efficiency vs. reference data.

While a degradation model has not yet been implemented in the current study, a qualitative analysis of the fuel cell power output profile has been conducted [56]. The results suggest that energy management strategies inducing less frequent power fluctuations are likely to be more beneficial for fuel cell longevity. Since the present model is primarily intended for the preliminary evaluation and comparison of energy management algorithms, degradation effects were not quantitatively modeled. However, given that the electrochemical model of the fuel cell is already well established, integrating a degradation model is both feasible and planned for future development.

Figure 13 presents the variation in driving modes throughout the entire simulation, during which a total distance of 284.9 km is covered. The powertrain operates in power split mode for over 80% of the time, highlighting the critical role of a well-designed EMS.

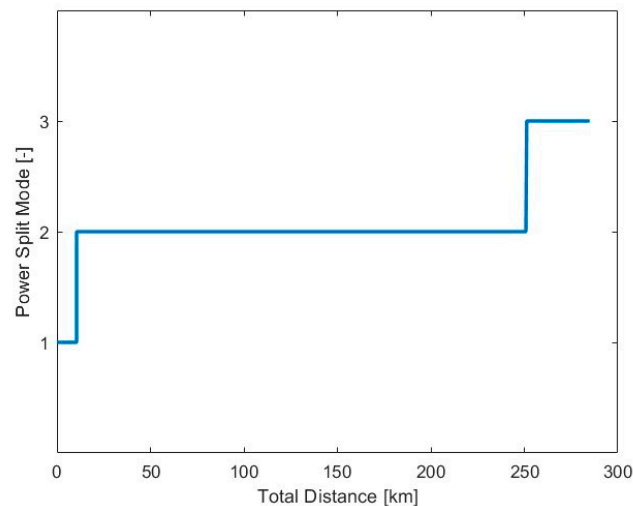


Figure 13. Power split mode variation.

The simulation results confirm that the developed model is both reasonable and reliable. This model can serve as a solid foundation for the design and validation of energy management strategies [48,53,54]. It enables the application of optimization-based algorithms to operate the fuel cell at higher-efficiency points, thereby extending the total driving range. Furthermore, the model's robustness also makes it suitable for integration with reinforcement learning approaches, providing a trustworthy environment for training and testing advanced EMS algorithms.

4. Conclusions

The calibration of the PEM fuel cell model presented in this paper has been demonstrated to be both effective and precise, despite limited available reference data. This calibration effectively reproduced the experimental polarization curve and the transient response characteristics, underscoring the robustness and accuracy of the developed model. Quantitatively, the calibration results are strong: the RRSME of both the fuel cell polarization curve and the overall system efficiency are approximately 0.01, while the R^2 values are around 0.99, indicating excellent agreement with experimental data.

The comprehensive powertrain simulation platform, integrating detailed models of the fuel cell stack, battery pack, electric motor, and vehicle longitudinal dynamics, was validated under realistic WLTP driving cycles. The energy management system presented in this study is intended as a representative example to demonstrate the feasibility and reliability of the proposed modeling approach. Its primary purpose is to support the simulation framework rather than to optimize control strategies. While the implemented algorithm is relatively simple, it provides a solid basis for future work involving the development and evaluation of more advanced energy management strategies. The simulation accurately captured critical dynamic behaviors, energy flows, and power distribution patterns inherent to hybrid PEMFC–battery systems. Simulation runtime is an important factor for real-time applications. For example, a full WLTC cycle—from 90% to 10% SoC and full to empty hydrogen tank—takes around 20 min. However, this duration can vary depending on the complexity of the energy management algorithm.

The energy management system (EMS) implemented in this study primarily serves as an illustrative example to demonstrate the model's capability for reliable analysis and to introduce the proposed modeling framework. While the EMS remains basic and fundamental, it provides a solid foundation for future development. The modeling approach enables systematic testing and benchmarking of various, potentially more advanced, EMS

algorithms. This platform therefore establishes valuable groundwork for future research aimed at improving and optimizing hybrid energy management in fuel cell electric vehicles.

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Data Availability Statement: Data is contained within the article.

Conflicts of Interest: The authors declare no conflicts of interest.

Abbreviations

The following abbreviations are used in this manuscript:

BEV	Battery Electric Vehicle
BoP	Balance of Plant
EMS	Energy Management Strategies
FC	Fuel Cell
FCEV	Fuel Cell Electric Vehicle
FCS	Fuel Cell Stack
ICE	Internal Combustion Engine
LCV	Light Commercial Vehicle
LHV	Lower Heating Value
PCU	Powertrain Control Unit
PEM	Proton Exchange Membrane
RRSME	Relative Root Square Mean Error
SoC	State of Charge
SoH ₂	State of Hydrogen
VDS	Vehicle Dynamic System
WLTP	Worldwide Harmonized Light vehicles Test Procedure

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