Research on water status detection method of high-power fuel cell system based on water balance model

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# Abstract

With the gradually accelerating pace of global decarbonization, highly efficient and clean proton exchange membrane fuel cells(PEMFC) are considered one of the solutions for future energy. During the operation of the fuel cell, it is necessary to keep the internal proton exchange membrane in a good state of hydration, so an appropriate method of detecting the hydration state is essential. At present, the fuel cell system is rapidly developing towards high power, but the methods for detecting the hydration state of high-power fuel cell systems are still relatively lacking. Therefore, this paper studies the hydration state of high-power fuel cell systems and builds a condensation tail gas water collection device for calculating the water flow out of the fuel cell system. To verify the water balance model, a controlled variable experiment was conducted on a 100kW fuel cell system under different working temperatures, air metering ratios, and load currents. Finally, based on the experimental data, the change rate of the internal water content of the fuel cell system under different conditions was calculated. The results show that, under the same load current, as the working temperature and air metering ratio increase, the change rate of the internal water content of the fuel cell system gradually decreases. Therefore, at low power, it is necessary to maintain an appropriate working temperature, while at high power, maintaining an appropriate metering ratio is more important.

*Keywords:* PEMFC, water content, water balance, high power fuel cell system

## **1. Introduction**

Proton exchange membrane fuel cells (PEMFCs), henceforth denoted as fuel cells, facilitate the direct conversion of chemical energy present in hydrogen into electrical energy. These fuel cells are characterized by their high energy utilization rate, low operating temperature, and the production of water as the sole reaction byproduct. Their versatility allows them to function not only as a small-scale distributed power supply

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*Preprint submitted to TBD April 11, 2024*

but also as the energy source for high-power transportation systems, thus demonstrating their broad applicability [1] . Nevertheless, the reliability, stability, and durability of the fuel cell stack, as the central component of the fuel cell power system, have emerged as critical factors influencing the extensive commercialization of fuel cell products. During the operational phase of the fuel cell, it is imperative to maintain the internal proton exchange membrane (PEM) in an optimal state of hydration to ensure the proton conduction ability remains at its peak, thereby achieving efficient and stable output performance of the fuel cell. Hydration state failure is the most prevalent failure mode of fuel cells, accounting for approximately 52 of total failures, primarily manifested as drying and flooding. Short-term hydration state failure can induce fluctuations and reductions in the output performance of the fuel cell. If this condition persists, it may lead to irreversible attenuation.

Under dry failure, the water content inside the stack is too low to keep an sufficiently hydrated state, and the internal resistance increases and heat production increases since some region of the PEM are not fully hydrated. Long-term dryness may even cause mechanical damage to the PEM, resulting in irreversible damage[2]. Under flooding failure, the water content inside the stack is too high, and the liquid water gradually block the flow channel or reaction area,preventing the reaction gas from reaching the reaction site, causing a local ”starvation” phenomenon[3, 4]. In severe cases, other side reactions will occur in the fuel cell at this time, causing damage to the carbon carrier[4–

6].

In addition to the direct impact on output performance, the internal hydration state also indirectly affects environmental adaptability and durability[7, 8]. In low temperature conditions, especially in sub-zero environments, the free water inside the fuel cell will freeze, thereby blocking the reactant channels, and even causing structural damage to the catalyst layer and diffusion layer[9, 10]. In addition, it will also affect the aging characteristics of the catalyst[2, 11]. Flooding conditions will accelerate the loss of Pt surface area, while dry conditions will accelerate the corrosion of the carbon carrier [12, 13].

Currently, studies focusing on the characterization of the internal hydration state of fuel cells have yielded significant findings. Standard indicators utilized for ascertaining the hydration state of fuel cells encompass direct observation or computation of water content, electrical signals, pressure drop signals, and electrochemical signals [14].

The identification of a fuel cell’s hydration state fundamentally involves determining the internal water content, making this the most basic and accurate assessment. Direct observation of water content can be facilitated through the transparent cell method and the neutron imaging method. The transparent cell method modifies the traditional fuel cell structure, employing transparent end plates and flow field plates to directly observe the quantity of liquid water within the flow field[15]. Hussaini et al[14] designed a fuel cell with an effective area of 14cm², containing seven straight channels, and conducted a visual examination of the liquid water content in the fuel cell channels under typical automotive conditions. Yang et al[16] utilized a transparent fuel cell for experiments to observe the process of liquid water transfer, discovering that water droplets initially appear on the GDL surface and adhere to the GDL surface due to surface tension. The water droplets progressively enlarge until they make contact with the flow channel wall, which exhibits stronger hydrophilicity. If the droplets on the flow channel are sufficiently thick, they may obstruct the channel and disrupt the airflow. The neutron imaging method is predicated on the principle that rays will attenuate when traversing an object. Given that different materials exhibit varying attenuation characteristics for neutron beams, particularly as water has higher absorbency than other materials such as metals, the internal water content can be measured using the transmitted neutron dose rate. Pekula[17] and Trabold[18] employed neutron imaging technology and found that water accumulation is likely to occur where the flow channel turns, providing appropriate explanations for each. However, the high cost and complex operation of current neutron imaging equipment have limited its widespread adoption.

Consequently, researchers have considered utilizing the law of conservation of mass to compute the water content within the fuel cell. When the fuel cell is operating stably, the flow rate of residual water can be calculated by measuring the discharged water. Zhao et al[19] employed a single transparent fuel cell with an active area of 25cm to conduct tests under varying operating conditions. By comparing the calculated rate of change in water content within the fuel cell and the water distribution in the flow channel, the accuracy of the model was validated.

The output voltage of the fuel cell serves as the most direct indicator of judgment, and faults in the internal hydration state will directly influence the output voltage, causing it to decrease or fluctuate. Particularly for high-power fuel cell systems, the Cell Voltage Monitoring module (CVM) is often the simplest standard of judgment. In terms of the impact of hydration state faults on voltage, recognition standards based on electrical signals encompass polarization curves and voltage fluctuation signals.

Li et al.[20] showed that at low current densities, water flooding have little effect on the polarization curve. As the current density increases, the voltage drop gradually becomes apparent. The more severe the water flooding, the more severe the voltage decay, and the earlier occurrence of voltage decay. Legros et al[21] introduced dry and wet gases into a single cell, respectively, and found that the fluctuation of the output voltage increased with the current density during dry periods.

The pressure drop between the inlet and outlet of the fuel cell is mainly caused by the friction between the flowing medium and the internal flow channel of the cel[22]. The pressure drop between the two ends increases with the increase of liquid water content, so the pressure drop can well reflect the degree of water flooding in the stack. The indicators include: pressure drop, pressure drop residual, pressure drop frequency, two-phase flow Multiplier coefficient, etc., and the, diagnosis is carried out through the above indicators by qualitative analysis, statistical analysis, artificial intelligence analysis, etc.[23]

The detection methods based on electrochemical signals mainly include electrochemical impedance spectroscopy, current interruption method, and cyclic voltammetry, etc. Among them, electrochemical impedance spectroscopy is the most common hydration state detection method because it can distinguish different electrochemical processes and can be tested online [12, 13]. The figure 1 shows the difference in fuel cell impedance spectra under different air humidity conditions. As the air humidity decreases, the value of the high-frequency impedance segment (i.e., the intercept of the impedance spectrum with the real axis) gradually increases, the impedance spectrum moves to the right, and the radius of the low-frequency arc segment also gradually increases [14], which can be used as an indicator of the hydration state of the fuel cell.

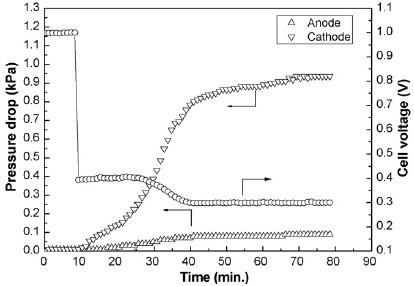


Figure 1: The change of cathode and anode voltage drop with time during flooding process

When analyzing the relationship between impedance spectra and the internal hydration state of fuel cells, equivalent circuit models are often introduced. The model generally includes resistors, capacitors, inductors, and Warburg elements[24]. The model is capable of quantitatively analyzing the electrochemical process. The key to this method is the establishment of equivalent models and parameter identification. Fouquet et al.[25] extended the standard capacitor to a constant phase element, proving that the three resistance values of the model are related to the relative humidity of the air supply. Zhang et al.[26] used a neural network method to complete parameter identification, and the predicted Nyquist plot almost completely coincides with the measured value, and by directly predicting the parameters of the equivalent circuit, they detailed the degree of influence of different processes on fuel cell performance. BMW proposed a method for online monitoring of impedance, which requires collecting five frequency points, divided into three groups to fit the Randles circuit, obtaining ohmic impedance, polarization resistance and double layer capacitance[25].

However, the above methods are mostly suitable for single cells or low-power fuel cells. Applyingthemtohigh-powerfuelcelldoesn’tgainanynoticeableimprovements[24, 26–29], since the assembly structure, flow field structure, and operating conditions of high-power fuel cell stacks are significantly different from those of single cells and low-power stacks [30].

Furthermore, research has indicated that smaller-scale fuel cell stacks exhibit reduced sensitivity to gas flow rates[31], which can also cause the difference between small and big stacks.

Therefore, the detection of the hydration state of high-power fuel cell systems is still in a relatively blank stage.

Additionally, the variation in the number of cells within a fuel cell stack can significantlyinfluencethepowergenerationperformance, leadingtotemperatureimbalances[28]. This, in turn, further exacerbates the uneven distribution of water present throughout the stack. Consequently, there exists a pronounced distinction between smaller and larger stacks regarding these phenomena. Researches indicates that there may exist notable differences in the water generation mechanisms between large-scale and small-scale polymer electrolyte membrane fuel cell (PEMFC) stacks[32]. Therefore, traditional approaches relying on electrochemical analysis and similar techniques may not be applicable to large-scale stacks. Due to the fact that large-scale fuel cell stacks frequently operate at lower power densities[33], this could potentially have an impact on the water generation mechanisms involved, further differentiating them from their smaller-scale counterparts.

Consequently, this study aims to investigate the hydration state of high-power fuel cell systems, construct, and validate a water balance model. Initially, the water balance model of the high-power fuel cell system is scrutinized, and a condensation-type exhaust water collection device is established to compute the water flow rate emanating from the fuel cell system. To authenticate the constructed water balance model, controlled variable experiments are executed under varying working temperatures, air metering ratios, and load currents. The experimental outcomes indicate that as the working temperature and air metering ratio escalate, the water flow rate emanating from the air side incrementally increases, and the water flow rate from the hydrogen side gradually diminishes. As the load current amplifies, the water flow rate emanating from both sides augments. Ultimately, predicated on the experimental data, the change rate of the internal water content of the fuel cell system under diverse conditions is calculated. The results reveal that under the same load current, as the working temperature and the air stoichiometric ratio augment, the change rate of the internal water content of the fuel cell system progressively decreases. Conversely, as the load current intensifies, the impact of the air stoichiometric ratio also incrementally escalates. Therefore, at low power, it is essential to maintain an appropriate working temperature, while at high power, upholding an appropriate metering ratio is of greater significance.

Based on the experiment results, we propose the following viewpoint: The water generation and water balance models for large-scale fuel cell stacks differ mechanistically from those of smaller-scale models and cannot be considered as linear scalings of the smaller models. The internal states of large-scale fuel cell stacks exceed the complexity of small-scale stacks due to their increased scale. Consequently, traditional approaches based on small-scale stacks may not be entirely applicable to large-scale stacks.

## **2. Experiment**

To verify the reliability of the water balance model for high-power fuel cell systems, it is necessary to record the water-containing state of the high-power fuel cell system when it reaches water balance under different operating conditions. Therefore, the experimental conditions should meet the following three points:

* The selected operating parameters should be within the normal operating range

Table 1: Water state experimental parameter table

|  |  |  |
| --- | --- | --- |
| **Load current(A) / currency density(***A*ꞏ*cm*−2**)** | **Working temperature(**◦*C***)** | **Air metering ratio** |
| 120/0.4 | 55,60,65,70,73 | 2,2.2,2.4,2.6,2.8 |
| 210/0.7 | 55,60,65,70,73 | 1.8,2,2.2,2.4 |
| 300/1.0 | 63,65,68,70,73 | 1.8,2,2.2,2.4 |

of the fuel cell system to avoid other faults in the fuel cell system other than water content faults (such as insufficient supply of reaction gas, air compressor surge, proportional valve failure, etc.)

* In order to study the phenomena when the fuel cell system has a water content fault, the designed experiment needs to reflect more obvious phenomena under dry and flooded conditions
* The steady-state operation time should be long enough to ensure that the water content inside the fuel cell is in dynamic balance during this period, avoiding the impact of dynamic characteristics on the experimental results.

### 2.1. Experiment Design

The literature indicates factors with greatest impact on water content inside fuel cell are air metering ratio, working temperature, and load current [21]. Given the temperature inside stack cannot be measured during operation, the inlet temperature of coolant is treated as working temperature in this research, and the working temperature in the following text refers to the inlet temperature of the coolant. According to the operating conditions recommended by the system side, a three-factor five-level water content experiment was designed. The parameters of the experiment are shown in Table 1.

Literatures[22] indicates that the time required for the fuel cell system to reach water balance is generally a few seconds to ten minutes. Therefore, each test point ran for 20 minutes, and it was assumed that the fuel cell was in the process of establishing water balance within the first 15 minutes, and no data was recorded for this process, and the data obtained in the last 5 minutes was recorded. During the test, if a single cell voltage is too low or other faults cause the fuel cell system to shut down, the data of this operating point will be removed.

### 2.2. Test platform introduction

The high-power fuel cell system used in this article is shown in Fig 2, manufactured by AT&M Environmental Engineering Technology Co. With rated net output power of 100kW, the stack is composed of 410 single cells. High-pressure dry hydrogen gas enters the anode of the reaction stack after being depressurized by the solenoid valve and proportional valve. The liquid water in the residual hydrogen is separated by the gas-water separation device and intermittently discharged through the drain valve.

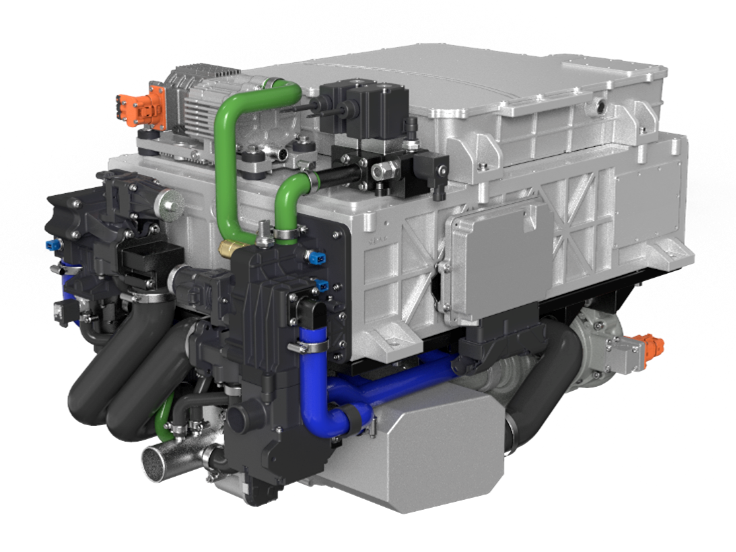
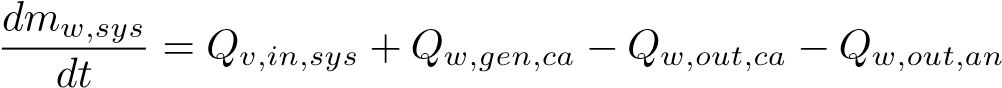


Figure 2: power system diagram

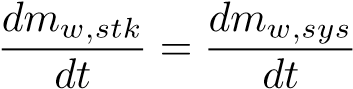
For the above fuel cell system, the water balance model can be represented as

 (1)

In formula (1), epresents the water

* *m*(*w,sys*) represents the water content inside the fuel cell system, in grams;
* *Q*(*v,in,sys*) represents the flow rate of water vapor entering the fuel cell system, in g/s;
* *Q*(*w,gen*) represents the water flow generated by the electrochemical reaction, in g/s;
* *Q*(*w,out,ca*) represents the water flow out of the system from the air subsystem, in g/s;
* *Q*(*w,out,an*) represents the water flow out of the system from the hydrogen subsystem, in g/s;

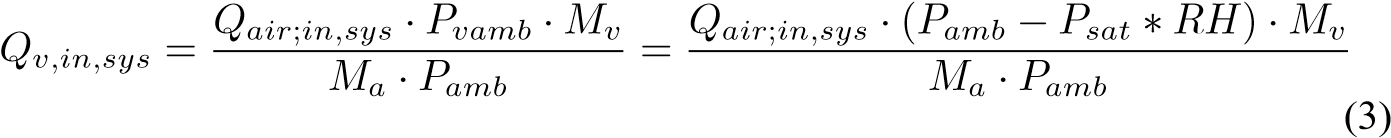
Taking the fuel cell system during steady-state operation as the research object, the change in the water content of the auxiliary system of the fuel cell system can be ignored, that is,

 (2)

Each part of the equation required for the water balance model is described below:

## Water flow into the system

Since the hydrogen entering the system is high-pressure hydrogen with a purity of 99.99%, water vapor mainly enters the system with the air. By recording the temperature, humidity, pressure of the environment during the reaction process, and the air flow entering the system, the water flow entering the system can be calculated.



In Formula(3),

* *Qair*;*in,sys* stands for air flow into the system, in g/s;

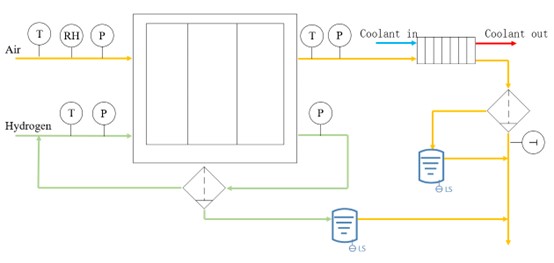
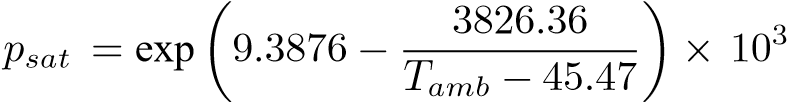


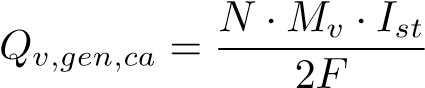
Figure 3: Schematic diagram of exhaust gas water content detection device

* *Pamb* stands for environmental pressure, in kPa;
* *Rv* represents the relative humidity of the environment;
* *Psat* represents the saturated vapor pressure of the environment, which can be calculated using the saturated water vapor pressure formula.

 (4)

* *Tamb* represents environmental temperature, K;
* *Mv* represents water relative molecule mass, in g/mol;
* *Ma* represents water relative molecule mass, in g/mol;

## Water flow by electrochemical reaction

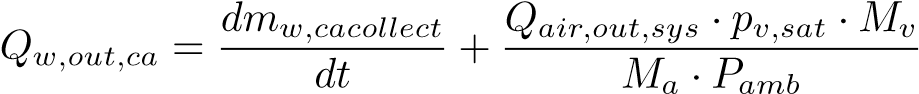
The water flow produced by electrochemical reaction can be presented as formula below:

(5)

* *N* represents the number of single cells in the fuel cell stack.
* *Ist* represents load current, in A.
* *F* stands for faraday constant, which is 96485C/mol.

## The water flow out of the system from the air side and the hydrogen side

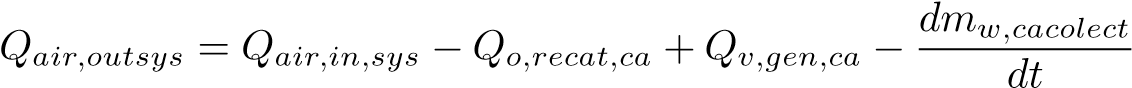
To measure the water flow out of the system from the air and hydrogen sides, this article has designed an exhaust water content detection device centered on condensation. The exhaust water content detection device on the air side is composed of a heat exchanger, a water separator, a temperature sensor, and a water collection device with a liquid level sensor. The basic principle is to condense the high-temperature supersaturated exhaust gas on the air side, and then separate the liquid water inside from the exhaust gas through the water separator, thus obtaining saturated low-temperature exhaust gas and liquid water. Collect the separated liquid water and collect the liquid level with a liquid level sensor. Therefore, the water flow out of the system from the air side is composed of two parts: the liquid water collected in the air side collection device and the water vapor carried in the saturated steam after water vapor separation, that is,

 (6)

In formula(6),

* *mw,cacollect* represents the mass of the liquid collected by the exhaust water collection device on the air side, in grams(g);
* *Qair,out,sys* representstheairflowoutofthesystem, ing(gram)/s(second), which can be calculated by the following formula.

The *Qair,out,sys* is represented as below:

 (7)

Since the hydrogen return circuit of the system is equipped with a water separator, the water content detection device on the hydrogen side mainly collects liquid water and a small amount of hydrogen, with less gaseous water, which does not significantly affect the experimental results. Therefore, the exhaust water content detection device on the hydrogen side only has a water collection device with a liquid level sensor, which is used to record the water flow discharged from the hydrogen side.

*dmw,ancollect*

*Qw,out,an* = (8) *dt*

*mw,ancollect* represents the mass of the liquid collected by the exhaust water collection device on the hydrogen side, in grams(g).

## Test methods

In sequence, turn on host computer, system controller, electronic load, auxiliary devices, and impedance testing system; Turn on the water pump, solenoid valve, proportional valve, back pressure valve, air compressor, drain valve, and hydrogen circulation pump in sequence, setting the anode inlet gas pressure 10kPa higher than the cathode inlet gas pressure (∆*p* = 20*kPa*).

Gradually load the system to 120A, adjust the speed of the air compressor and the opening angle of the back pressure valve, so that the air flow rate reaches the air metering coefficient of 2.8 corresponding to the target current, gradually raise the working temperature to 55 ± 1◦*C*, and start the temperature closed-loop control program to maintain this temperature; After the working temperature is stable at 55 ± 1◦*C*, start timing and run for a total of 15 minutes; Adjust the speed of the air compressor and the opening angle of the back pressure valve, so that the air metering ratio gradually decreases in the range of 2 2̃.8 with a step size of 0.2; Raise the working temperature to 60 ± 1◦*C*, 65 ± 1◦*C*, 70 ± 1◦*C* , 73 ± 1◦*C*, repeat the testing phase and record the experimental data.

Load up to 210A, 300A, repeat all recording steps, until all operating points water content detection experiments are completed.

Gradually reduce the load to 0A, purging circuit with nitrogen, turn off the machine, and power down in sequence.

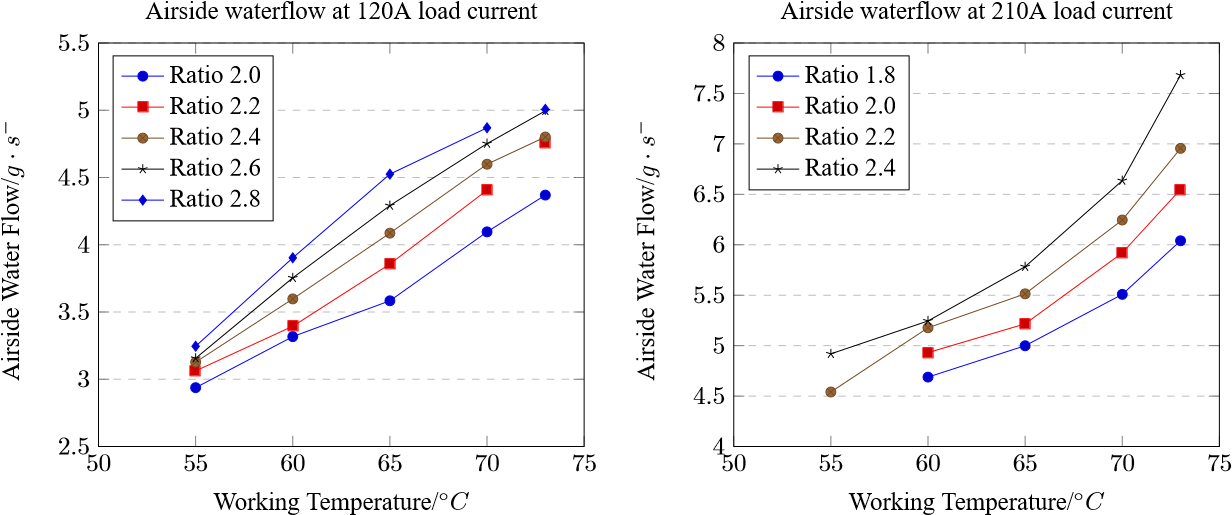
### **3. Results and discussion**

#### 3.1. Effect of operating temperature on water flow rate outflow from air side

Figure **??** shows the change in the water flow out of the air side at different working temperatures. As can be seen from the figure 5a, as the temperature rises, the water content in the air side exhaust gas generally shows an upward trend. The higher the temperature, the higher the saturation vapor pressure of the fuel cell. Under the same water content, the relative humidity will decrease at this time, which is equivalent to the generation of less liquid water inside the fuel cell, and the ability of the gas to carry water is improved. This change applies to both the air side and the hydrogen side, which is shown in figure 4b and figure 4c. Therefore, under the same other conditions, the water flow out of the fuel cell stack by the air increases with the increase in temperature, and more liquid water is collected by the air side exhaust gas collection device.

It is worth noting that when the load current is 120A, the impact of the working temperatureonthewaterflowoftheairsideisslightlydifferentunderdifferentmetering ratios. As the air metering ratio increases, the growth rate of the water flow on the air side gradually decreases. This is because at a small current density, the fuel cell produces less water due to the electrochemical reaction, and at the same time, the highspeed high-temperature reaction gas has a strong ability to carry water, and the water produced by the fuel cell has been completely carried out by the reaction gas. Therefore, even if the air metering ratio increases, the reaction gas cannot carry out more water. For the working conditions of low working temperature and large load current, the ability of the reaction gas to carry water is low or the fuel cell produces more water, so there is still some water vapor or liquid water remaining in the system. At this time, if the air flow is increased, the water flow out of the system will also increase accordingly.

Fig 5a shows that as the temperature rises, the water content in the hydrogen side exhaust gas generally shows a downward trend, and even at 120A/70◦*C*, the water content in the hydrogen side exhaust gas is 0, the same result appears in 210A(fig 5c) and 300A(fig 5c). There are mainly two reasons for this phenomenon. First, according to the principle of proton exchange membrane fuel cell reaction, the reaction product water is generated at the cathode (that is, the air side), and the fuel used by the fuel cell system is 99.99% pure hydrogen. Therefore, the water inside the hydrogen side is mainly the water that migrates from the air side to the hydrogen side under the action of concentration diffusion and pressure diffusion. Since the hydrogen-air pressure difference is basically consistent (20kpa) during the experiment, the influence of pressure on water migration can be ignored when comparing. Second, the exhaust gas collected on the anode side needs to first pass through the water separator built into the fuel cell system to separate the unreacted hydrogen from the liquid water, and then it can be



(a) (b)

60

62

64

66

68

70

72

74

7

7

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5

10

10

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5

WorkingTemperature/

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*C*

AirsideWaterflow/

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1

Airsidewaterflowat300Aloadcurrent

Ratio1.8

Ratio2.0

Ratio2.2

Ratio2.4

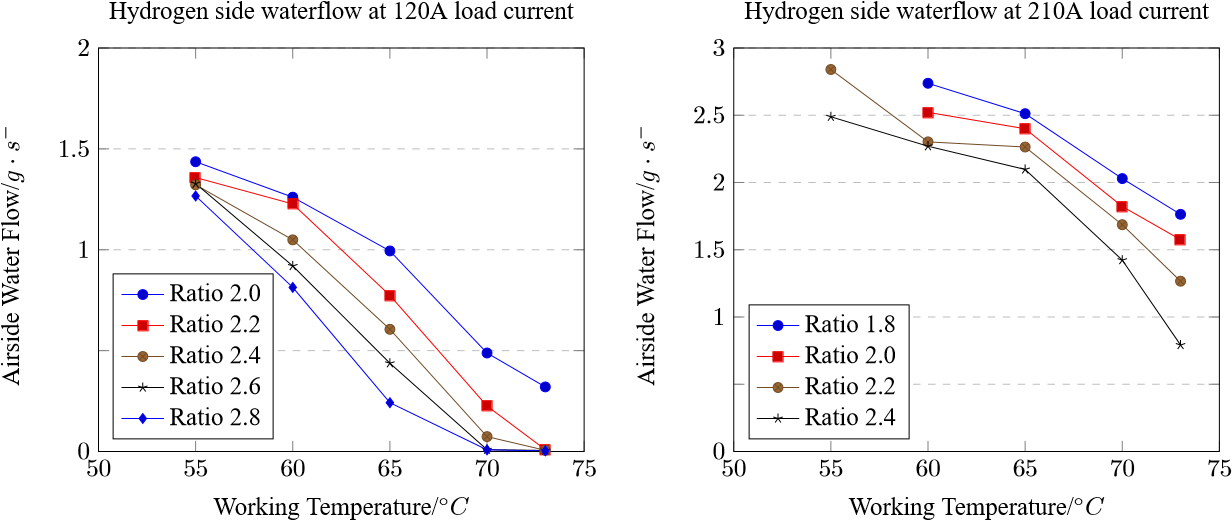
(c)

Figure 4: Comparison of water flow rate flowing out of the air side at different operating temperatures (a:120A; b:210A; c:300A; vertical water flow; horizontal as temperature; starting point from top to bottom is ordered by flow rate: 1.8/2.0/2.2/2.4)

collected by the exhaust water collection device on the hydrogen side. And at higher working temperatures, the saturation vapor pressure of the gas on the hydrogen side rises, and the content of liquid water decreases relatively. These two factors cause the working temperature to rise and the water content flowing out of the hydrogen side to decrease.

#### 3.2. Influence of air metering ratio on the water flow out of the cathode and anode

Compare the water flow out of the air side and the hydrogen side at 120A, 210A, 300A load currents, different coolant inlet temperatures, and different air metering ratios, as shown in Fig 6a. Similar to the impact of working temperature on the water flow out of the air side, under the same load current, the higher the air metering ratio, the more water is carried out of the stack by the unreacted air, which also applies to load current of 210A(fig 6b) and 300A(fig 6c). It should be noted that this growth is not unlimited. After reaching a certain air metering ratio, the growth of the water content in the air side exhaust gas slows down, especially when the load current is small and the working temperature is high. This is because at this time, the water produced by the electrochemical reaction of the fuel cell has been completely carried out of the

0*.*5

0*.*5

(a) (b)

60

62

64

66

68

70

72

74

2

2

*.*

5

3

3

*.*

5

4

4

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5

WorkingTemperature/

◦

*C*

AirsideWaterFlow/

*g*

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*s*

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1

Hydrogensidewaterflowat300Aloadcurrent

Ratio1.8

Ratio2.0

Ratio2.2

Ratio2.4

(c)

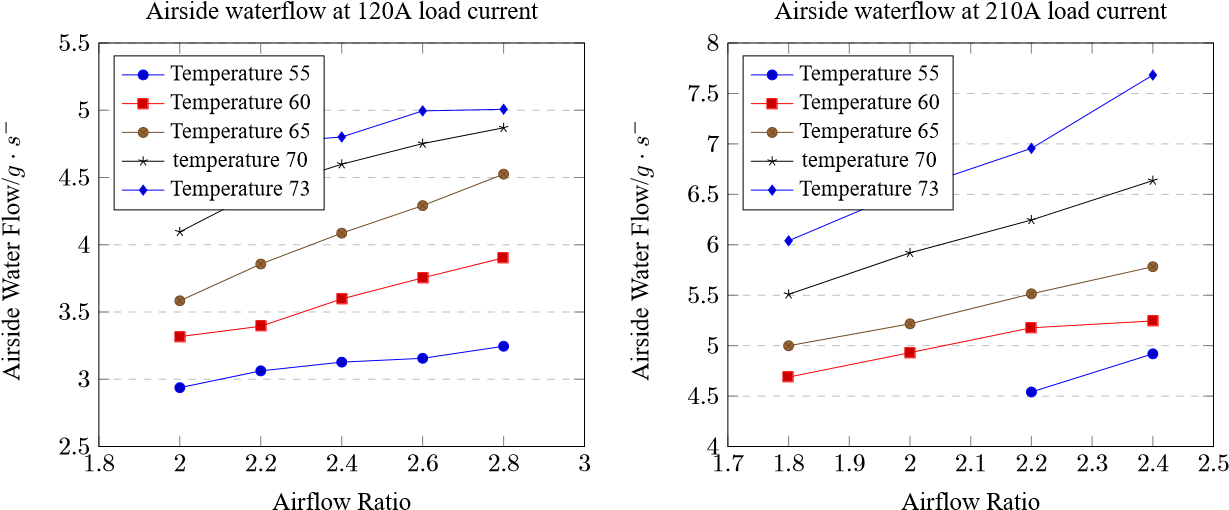
Figure 5: Comparison of water flow rate flowing out of the air side at different operating temperatures (a:120A; b:210A; c:300A; vertical water flow; horizontal as temperature; starting point from top to bottom is ordered by flow rate: 1.8/2.0/2.2/2.4)

fuel cell by high-temperature high-speed air. At this time, even if the air metering ratio continues to increase, the water content of the exhaust gas will not continue to increase.

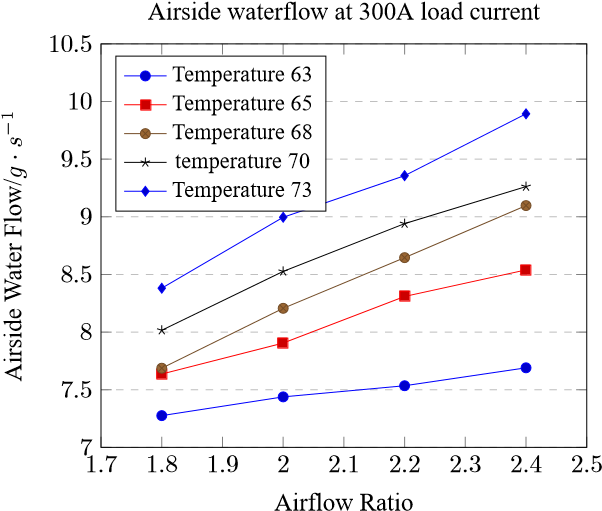
For the hydrogen side, similarly, as the air metering ratio increases, the water flow out of the hydrogen side basically continues to decrease. After the above analysis, as the air metering ratio increases, the water stored in the cathode will be taken out of the stack by the high-speed airflow, and the water flow diffused to the hydrogen side through concentration difference becomes less, so relatively speaking, the water collected on the hydrogen side will become less. However, the degree of decline is different. Under low temperature (120A/60◦*C*, 210A/60◦*C*, and 300A/63◦*C*), they all show a relatively gentle decline. As the temperature rises, the water flow out of the hydrogen side is also gradually affected by the air metering ratio.

#### 3.3. Effects of operating temperature and air metering ratio on internal water content of fuel cell system

According to the working principle of the fuel cell, during the reaction process, water mainly comes from the electrochemical reaction and the water vapor that enters



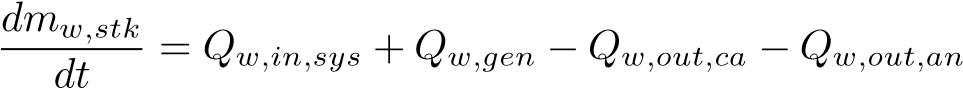
(a) (b)



(c)

Figure6: Comparison of waterflow rate flowing out ofthe air side underdifferentair metering ratios (a:120A; b:210A; c:300A; vertical as water flow; horizontal as air flow rate; starting point from top to bottom is ordered by temperature: 63/65/68/70/73℃)

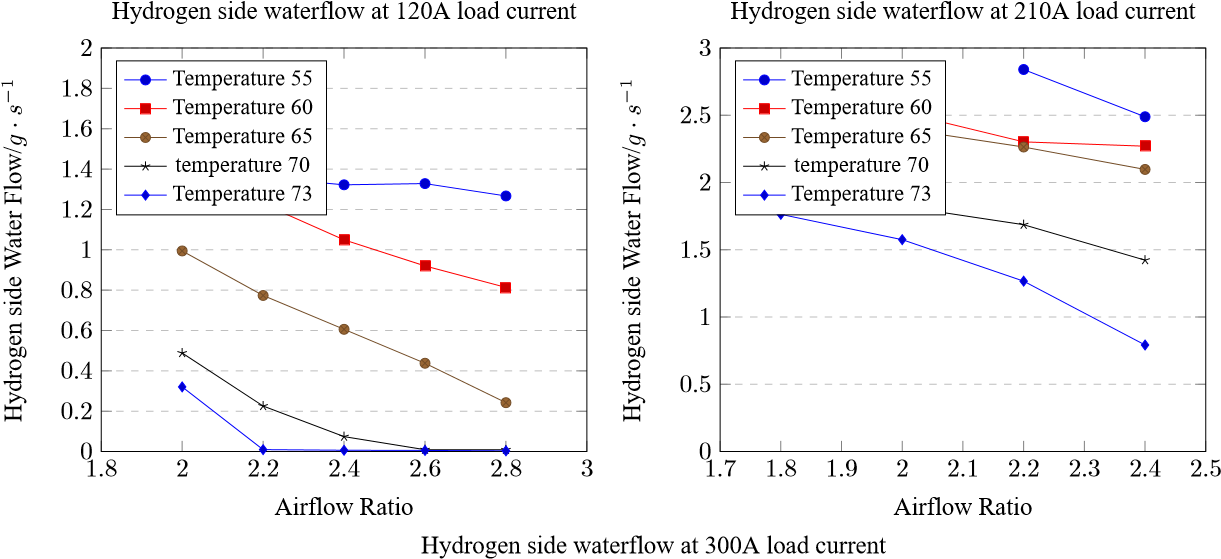
the fuel cell with the reaction gas. Then, a part of the water is discharged from the fuel cell with the exhaust gas after the reaction, and another part of the water stays inside the fuel cell. In this article, the change rate of the water remaining in the fuel cell will be used as the research object to evaluate the water content status inside the fuel cell.

 (9)

In formula 9, *m*(*w,stk*) is the water content inside the fuel cell, in g, *Q*(*w,in,sys*) is the water flow rate entering the fuel cell system, in (g/s), *Q*(*w,gen*) represents the water flow rate generated by the electrochemical reaction, in (g/s), *Q*(*w,out,ca*) represents the water flow rate flowing out of the system from the cathode, in (g/s), *Q*(*w,out,an*) represents the water flow rate flowing out of the system from the anode, in (g/s).

Although during the reaction process, under the effects of concentration diffusion and electro-osmotic drag, water at the cathode will migrate to the anode. However, the research object of the water balance method is the entire fuel cell, so the internal water migration process will not have an impact.

Using the method described above, the rate of change of the water content inside



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7

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8

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9

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AirflowRatio

HydrogensideWaterFlow/

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*s*

−

1

Temperature63

Temperature65

Temperature68

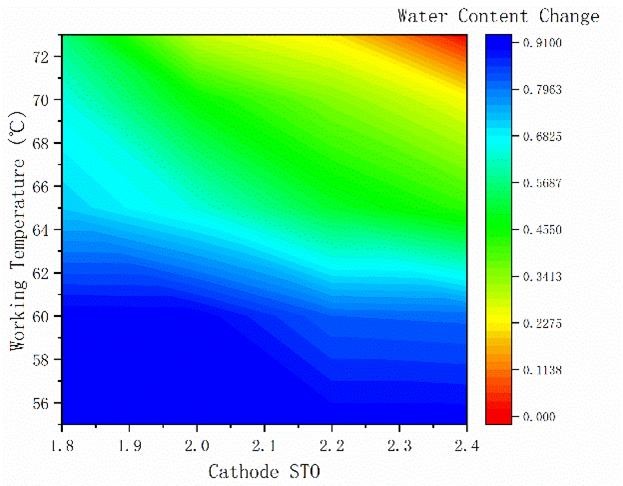
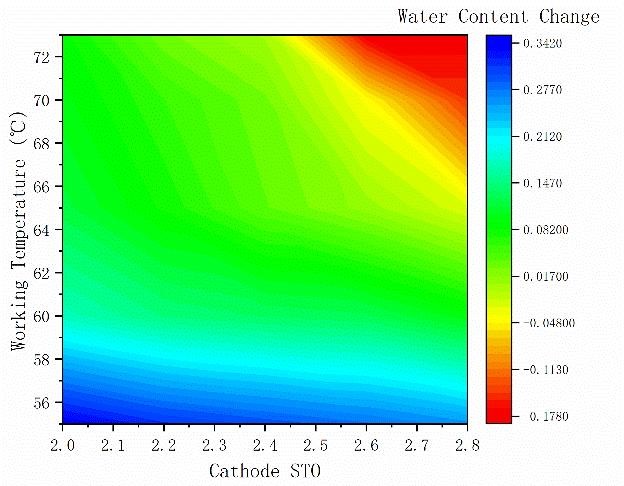
temperature70

Temperature73

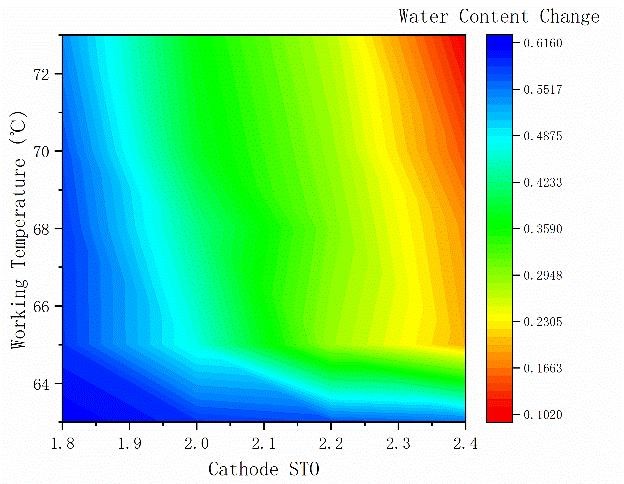
Figure7: Comparison of waterflow rate flowing out ofthe air side underdifferentair metering ratios (a:120A; b:210A; c:300A; vertical as water flow; horizontal as air flow rate; starting point from top to bottom is ordered by temperature: 63/65/68/70/73℃)

the fuel cell system under different load currents, working temperatures, and air metering ratios can be calculated. In the process of making the graph, to maintain the integrity of the graph. We use (Load current/Temperature/Coefficient) as the format of test environment description. The data of (210A/55◦*C*/1.8 ) and (210A/55◦*C*/2.0) are replaced with (210A/55◦*C*/2.2) test’s data. The obtained results are compared as shown in Fig 8a, Fig 8b and Fig 8c, where positive values indicate an increase in water content (represented in blue), and negative values indicate a decrease in water content (represented in red). For the convenience of subsequent discussions, the blue part is defined as flooding, the red part is defined as drying, and the green part is defined as the normal state.

As can be seen from Fig 8, under the same load current, as the working temperature increases and the air metering ratio increases, the rate of change of the water content inside the fuel cell system gradually decreases. When it is 120A/73◦*C*/2.8, 210A/73◦*C*/2.4, 300A/73◦*C*/2.4, the rate of change of the water content inside the fuel cell system reaches the minimum value under this load current. However, the degree of influence of the working temperature and the air metering ratio on the rate of change of the water content inside the fuel cell system is slightly different. When the load current is 120A and 210A, too low working temperature is more likely to cause flooding than too low air metering ratio. For example, at 120A, when the working temperature is



(a) (b)



(c)

Figure 8: Comparison chart of internal water content of fuel cell system under different conditions (a:120A; b:210A; c:300A)

55◦*C*, even if the air metering ratio reaches 2.8, the fuel cell system is still in a relatively flooded state; and when the working temperature is 73◦*C*, even if the air metering ratio is only 2.0, the fuel cell system is in a normal state. In a dry state, the degree of influence of working temperature and air metering ratio is not much different. However, when the load current is 310A, the trend is the opposite. Too low working temperature and too low air metering ratio will cause flooding, and too high air metering ratio is more likely to cause drying than too high working temperature.

Furthermore, this paper uses the method of linear regression to analyze the impact of working temperature and air metering ratio on the water content inside the fuel cell system. Before this, the data needs to be standardized. This paper uses the Z-score normalization method[34, 35], which is

*x*∗ = *x* − *µσ*

Table 2 shows the result of analysis: Similar to the phenomenon described above, the impact of the working temperature is more significant when load current is on 120A and 210A; while the impact of the air metering ratio is more significant when load current is 300A. Reason for this phenomenon may be the air metering ratio related to the load current. Under different load currents, the air metering ratio changes the same magnitude, but the change in air flow is different. Since the fuel cell system needs to rely on high-speed high-temperature gas to carry the water out, the higher the load current, the more significant the impact of the air metering ratio.

Table 2: Regression analysis under different load current

|  |  |  |
| --- | --- | --- |
| **Load current(A)** | **Temperature ratio regression coefficient** | **Air metering ratio regression coefficient** |
| 120 | -0.8469 | -0.4145 |
| 210 | -0.9341 | -0.4347 |
| 300 | -0.4637 | -0.7618 |

Table 3: Statistical table of the change rate of internal water content of different load currents

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Load current(A)** | **maximum** | **minimum** | **range** | **average** | **variance** |
| 120 | 0.342 | -0.177 | 0.519 | 0.086 | 0.019 |
| 210 | 0.909 | 0.009 | 0.900 | 0.562 | 0.067 |
| 300 | 0.615 | 0.103 | 0.512 | 0.402 | 0.028 |

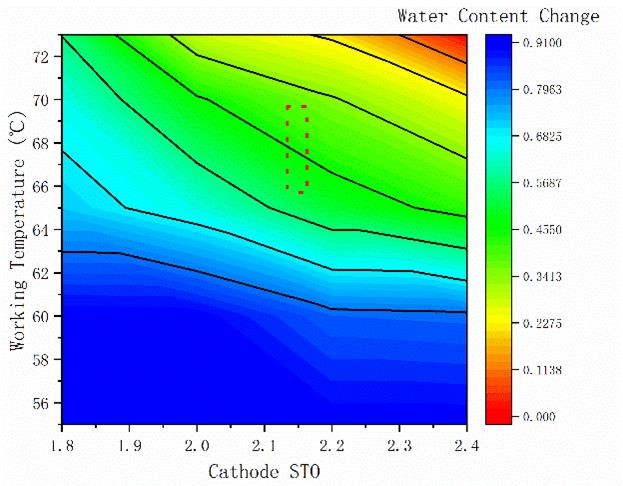
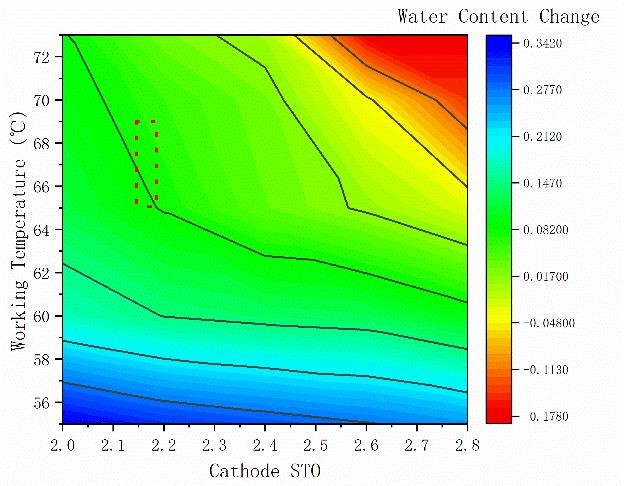
Therefore, to maintain the long-term stable operation of the fuel cell system, appropriate operating conditions need to be selected. The following principles should be considered when choosing:

* The fuel cell system should be in a normal state under this condition;
* In the actual operation process, there are certain fluctuations in the working temperature and air metering ratio. The fluctuation range of the working temperature is about ±2◦*C*, and the fluctuation range of the air metering ratio is about±0*.*1, so the selected conditions should make the points within the fluctuation range in a normal state;
* A larger air metering ratio and a lower working temperature will cause additional consumption of the auxiliary system, so try to choose working conditions with a low air metering ratio and a high working temperature.

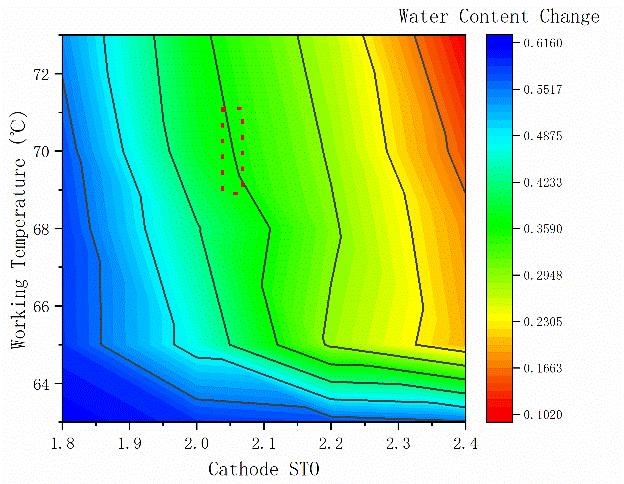
As shown in Fig 9, 120A/67◦*C*/2.15, 210A/68◦*C*/2.15, 300A/70◦*C*/2.05 are selected as the appropriate operating conditions under each load current. Next, comparing the rate of change of the water content inside the fuel cell system under different load currents.

From Table 3, it can be seen that as the load current increases, the rate of change of the water content inside the fuel cell system increases to some extent, but the increase is not large. This may be because the higher the load current of the fuel cell, the more water flow is generated, so the water accumulated inside correspondingly increases. In addition, itwas foundthat whentheloadcurrent is210A,therangeis largercompared to 120A and 300A. Correspondingly, the fuel cell system cannot operate stably under the conditions of 210A/55◦*C*/1.6 and 210A/55◦*C*/1.8. This further reflects the connection between the rate of change of the internal water content of the fuel cell system and the water content fault.

However, the above calculation method can only calculate the absolute value of the water content inside the fuel cell system. The output performance of the fuel cell in



(a) (b)



(c)

Figure9: Selectionof fuelcellOSoperating environment(Workingenvironmentsurroundedby dots, a:120A; b:210A; c:300A)

actual operation is related to the working temperature, intake pressure, and air metering ratio. To find the most suitable working conditions under each load current, it is necessary to establish a fuel cell stack model, and study the correlation between the difference between the actual output and theoretical output of the fuel cell stack under different conditions and the absolute value of the water content inside the fuel cell system.

### **4. Conclusion**

Different conditions and the absolute value of the water content inside the fuel cell system.

This study investigates the water content state of the high-power fuel cell system and constructs a water balance model. Initially, to authenticate this model, a tail gas water content detection apparatus centered on condensation was designed and established, based on the original high-power fuel cell system. Subsequently, control variable experiments involving three factors, namely, working temperature, air metering ratio, and load current, were executed to measure the water flow rate emanating from the air side and hydrogen side of the fuel cell system during steady-state operation. The experimental outcomes are as follows:

1. Under larger air metering ratios, elevated coolant inlet temperatures, and higher load currents, the water content in the exhaust gas on the air side is higher; conversely, under smaller air metering ratios, lower coolant inlet temperatures, and higher load currents, the water content in the exhaust gas on the hydrogen side is higher.
2. Under the same load current, as the working temperature and air metering ratio escalate, the rate of change of the water content inside the fuel cell system progressively decreases. However, the degree of influence of working temperature and air metering ratio on the rate of change of the water content inside the fuel cell system slightly varies. When the load current is 120A and 210A, the influence of working temperature is more pronounced; whereas, when the load current is 300A, the influence of the air metering ratio is more evident.
3. With the amplification of load current, the rate of change of the water content inside the fuel cell system somewhat increases, but the increase is not substantial. Simultaneously, the range of the rate of change of the internal water content at 210A is greater than that at 120A and 300A, and the corresponding fuel cell system cannot operate stably under the conditions of 210A/55◦*C*/1.6 and 210A/55◦*C*/1.8. This further substantiates the correlation between the rate of change of the internal water content of the fuel cell system and the water content fault.

In conclusion, the water balance model can effectively predict the water content inside the fuel cell system. The experimental results provide a reference for the water content detection of high-power fuel cell systems. The water content status of the fuel cell system can be detected by monitoring the tail gas water volume. The results of this study can provide a theoretical basis for the water content detection of high-power fuel cell systems and can be used as a reference for the operation and maintenance of high-power fuel cell systems.

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