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1 Introduction

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Development of a double-side ordered membrane electrode assembly based on titanium nitride nanoarrays

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The membrane electrode assembly (MEA) plays a crucial role in the functionality of proton exchange membrane fuel cells (PEMFCs). The channels present within the catalyst layer of MEAs exhibit a disordered configuration, which consequently give rise to low efficiency in mass transportation. In order to enhance the mass transfer performance and the corrosion resistance of the catalyst layer, this paper developed a double-side ordered MEA based on TiN nanorod arrays. We synthesized TiN nanorod arrays on the ITO surface by a seed-assisted hydrothermal reaction and nitriding treatment, and coated the catalyst uniformly on the TiN support by ultrasonic spraying. Then the double-side ordered MEA was fabricated by transfer printing, and achieved a peak power of 678.30 mW cm⁻² with a cathode platinum loading of 0.2 mg cm⁻² at 80 °C and anode saturated humidity. After 200 hours of accelerated stress test (AST) at 90 °C and 30/ 30% relative humidity, the peak performance only dropped by 4.8%. These results provide substantial evidence for the effectiveness of our developed double-side ordered MEA which can mitigate catalyst polarization corrosion. Thus, this study reveals the immense potential of the TiN nanorod array-based double-side ordered MEA in advancing the development of efficient and stable MEAs. **PAPER**
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Keywords: PEMFC; Preparation of MEA; Ordered MEA; TiN nanorod array; Catalyst carrier.

Proton exchange membrane fuel cells (PEMFCs) can be regarded as a clean and highly efficient energy conversion technology that employs renewable hydrogen as a fuel source, and solely generates water as a by-product.^{1,2} Fuel cells have been widely used in the spheres of renewable energy, including transportation, power generation, and portable devices.3,4 Nonetheless, the crucial challenges of hydrogen– oxygen fuel cells for large-scale commercial application are the high platinum load and insufficient durability of the MEA.⁵ The MEA comprises a proton exchange membrane (PEM) sandwiched between two catalyst layers (CLs) and two gas diffusion layers (GDLs).⁶ The conventional configuration of the MEA exhibits several limitations, such as hindered mass transportation, catalyst aggregation, and membrane degradation.^{$7-9$} Consequently, it is essential to investigate novel MEA structures that can enhance the performance and surmount the catalyst polarization corrosion problem. The

ordered structure of the MEA facilitates the regular arrangement of catalyst supports on the PEM surface, thereby providing well-defined channels for reactant gas diffusion and effective water management. $10-12$

The preparation of corrosion-resistant nanomaterials is important to promote the advancement of ordered catalyst layers. Diverse materials, such as organic compounds, $13-16$ carbon nanotubes, $17,18$ and metal compounds, $19,20$ have been extensively utilized. The nanostructured thin film electrode (NSTF) developed by 3M® is one of the most typical ordered nanostructured MEAs and has been commercialized.^{21,22} Yang et $al.^{23}$ proposed a new catalyst/electrode structure, named the coaxial nanowire electrode (CANE), which consists of vertically aligned nanowires, each of which has an ionconducting core wrapped in a nanoscale Pt film. Through accelerated stress tests, it was found that the CANE only had 2% and 5% performance losses in the catalyst support and catalyst tests, respectively. Cho et $al.^{24}$ fabricated a Lego-like multi-layer multi-scale nanocolumnar PEM by using multiple photolithography techniques, which has a lower membrane resistance and larger electrochemical active surface area than the traditional flat Nafion membrane, thereby improving the performance of the fuel cell, while ensuring the mechanical stability of the membrane. Considering the inherent poor

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electrical conductivity and corrosion resistance of polymer whiskers, carbon nanotube arrays as catalyst layer carriers have been used.²⁵ Du et al.²⁶ employed a gas diffusion layer as the substrate and utilized PECVD to grow vertically aligned carbon nanotube arrays as catalyst supports. In the single-cell test, the electrode composed of nitrogen-doped carbon nanotube-supported CL not only showcased significantly higher power density but also remarkable durability compared to that of the Pt/C electrode. To further enhance the corrosion resistance of catalyst supports under high potentials, researchers employed metal compounds as catalyst supports.²⁷ Jiang et al.²⁸ synthesized TiO₂ nanorods (NRs) by a hydrothermal method on carbon paper, and then modified their surfaces with carbon as a supporting material. The testing results showed that the NR electrode can generate a maximum power density of 343 mW $\rm cm^{-2},$ also this electrode demonstrated remarkable stability throughout the testing duration. Although metal compounds may possess generally low conductivity, they present great potential due to their wide range of sources, robust stability, ease of processing, and ability to enhance catalyst activity.²⁹ The high conductivity and excellent corrosion resistance of titanium nitride (TiN) make it an excellent material choice. These characteristics enable TiN to play an important role in the catalyst support, which helps to improve the performance and stability of the catalyst layer. According to these advantages, the strategy of constructing ordered TiN nanorod arrays as a catalyst carrier will be studied with the aim of enhancing the activity, stability and performance of the catalyst layer, and finally developing an efficient and stable MEA. **Excelse Constraint Chemistry & Materials**

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The cooperation between the gas diffusion layer, porous layer, catalyst layer, and PEM will greatly affect the performance of the PEMFC. Therefore, it is crucial to investigate the preparation process involved in manufacturing of the MEA. In early stages of MEA preparation, catalyst layers are typically fabricated on gas diffusion layers to form a gas diffusion electrode (GDE).³⁰ Although the GDE method offers a straightforward fabrication process, the resulting MEAs pose challenges in achieving high-performance fuel cells. Subsequently, in order to improve the poor combination of the catalyst layer and PEM, a method called catalyst-coated membranes (CCM) was developed.³¹ This approach directly applies the catalyst to the PEM, resulting in a strong bond between the CL and PEM, as well as higher catalyst utilization. However, it imposes higher requirements on the manufacturing process. $32,33$ Several methods have been proposed for the preparation of MEAs, including scraping, 34 transfer printing 35 and spraying. 36 Park et $al.^{37}$ demonstrated a scraping method via a pre-expanded PEM, and the results showed that the height and speed of the scraper can be adjusted according to the state of the paste, to realize the reproducibility of the scraping process. Lee et $al.^{38}$ fabricated a novel MEA with a cathode groove electrode by transfer printing, which consisted of catalyst ridges with high ion polymer content and empty grooves,

providing effective channels for H^+ and O_2 transport, respectively. Compared with the optimized flat baseline electrode, it provided up to 50% performance improvement. As for the spraying method, Sassin et $al.^{39}$ developed a highperformance MEA using ultrasonic spray deposition of Pt/C catalysts directly onto a PEM. The results showed that the spray-deposited MEAs exhibited high power density, low ohmic resistance, and improved mass transport properties compared to those prepared through conventional methods. However, most of the fabrication processes can only achieve single-sided ordering on the cathode side of the CCM, but the MEA can only obtain better cell performance when both half-reactions of the anode and cathode are at a high level. In this study, we utilized a combined ultrasonic spraying and transfer printing process to fabricate CCMs with ordered structured catalyst layers at both the cathode and anode, which can not only enhance the electrochemical performance of the catalyst, but also improve catalyst utilization.

In this study, we propose a cost-effective method for constructing a MEA with an ordered anode and cathode using a corrosion-resistant material. Initially, the TiN nanorod arrays were synthesized on an ITO substrate through a seed-assisted hydrothermal process followed by nitridation treatment. Subsequently, the catalyst was uniformly applied onto the ordered support surface by ultrasonic spraying, and an ordered MEA was fabricated through thermal transfer printing. The physical and electrochemical properties of the prepared double-side ordered MEA were investigated experimentally. The results demonstrated that the doubleside ordered MEA can achieve a peak power of 678.30 mW cm−² with a platinum loading of 0.20 mg cm−² . Further, even after AST for 200 h, the output performance of the MEA only degraded by 4.8%. Overall, the fabricated double-side ordered MEA not only offers well-defined pathways for mass transfer but also provides a large surface area and reactive sites.

2 Results and discussion

2.1 Modulation of TiN nanorod array microstructure

The TiN nanorod arrays were obtained through the nitridation reaction of TiO₂. In situ growth of TiO₂ on ITO substrates requires the preparation of a seed layer, which facilitates the growth of rod-shaped $TiO₂$ on the substrate. In this study, TiCl₄ was selected as the precursor solution, and the morphology control of the ordered catalyst layer support of TiN can be achieved by controlling the concentration of the $TiCl₄$ solution in the precursor solution and the content of TBT and hydrochloric acid during the hydrothermal reaction.

In the seed-assisted hydrothermal reaction, the array density of $TiO₂$ is positively correlated with the concentration of TiCl₄ solution. Fig. 1 illustrates the SEM images of TiO₂ nanorod arrays grown from precursor solutions at various $TiCl₄$ concentrations. As the concentration of $TiCl₄$ increases from 0.05 mol L^{-1} to 0.20 mol L^{-1} , the directional alignment of TiO₂ array structure on ITO improves, the array density

Fig. 1 The SEM images of the TiO₂ ordered array: (a) 0.05 mol L^{−1}, (b) 0.10 mol L^{−1}, (c) 0.15 mol L^{−1}, and (d) 0.20 mol L^{−1} concentration of TiCl₄ seed solution respectively.

increases, and the diameter of the grown nanoscale $TiO₂$ decreases. When the concentration of TiCl₄ is 0.05 mol L^{-1} , the $TiO₂$ nanorods are dispersed in a clustered form on the surface of ITO, as shown in Fig. 1a1. Due to insufficient of growth sites, the nanorods tend to aggregate during the growth procedure, resulting in the generated $TiO₂$ being unable to cover the ITO surface completely, as shown in Fig. 1a2. At a TiCl₄ concentration of 0.10 mol L^{-1} , TiO₂ nanorods with a diameter of approximately 200 nm are grown on ITO. The alignment of the array is generally consistent, and suitable spacing between nanorods provides sufficient space for subsequent catalyst loading and facilitates efficient multi-phase channels for electrochemical reactions, as shown in Fig. 1b. When the $TiCl₄$ concentration reaches 0.15 mol L^{-1} , the ITO substrate is completely covered by $TiO₂$ nanorods, and noticeable cracks are observed at low magnification. Although the nanorod diameter is consistent at this concentration, the density is too high to serve as a catalyst support, as shown in Fig. 1c. At a $TiCl₄$ concentration of 0.20 mol L^{-1} , the nanorod array becomes excessively dense,

and numerous cracks are visible at low magnification, caused by stress concentration leading to fragmentation during cooling or cleaning processes, as shown in Fig. 1d. The overly dense seed distribution results in significant adhesion between nanorods, preventing further experiments. Additionally, the formation of bulk titanium dioxide can be observed, which would waste catalyst material and block multi-phase reaction channels. Therefore, a TiCl₄ concentration of 0.10 mol L^{-1} is ultimately selected as the precursor solution for seed growth.

After determining the precursor solution concentration for seed formation, it is crucial to investigate the key parameters during the hydrothermal reaction. Next, the TiN nanorod array on the ITO surface after the nitridation reaction will be observed. The content of TBT in the hydrothermal reaction precursor greatly influences the morphology of the nanorod array, as shown in Fig. 2. When 0.5 mL of TBT solution was added to the precursor solution, due to the inadequate TBT content, the TiN array fails to cover the entire ITO surface, as illustrated in Fig. 2a1. Additionally, an insufficient nanorod

Fig. 2 The SEM images of the TiN ordered array: the TBT content of (a) 0.5 ml; (b) 1.0 ml; (c) 1.5 ml; (d) 2.0 ml.

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array was obtained, and only discrete TiN clusters are formed, as shown in Fig. 2a2. By increasing the TBT content to 1.0 mL, a satisfactory TiN nanorod array is achieved. Furthermore, the TiN nanorods prepared under this condition exhibit uniform thickness, clear directional alignment, appropriate spacing distribution, and remarkably smooth surface when observed at low magnification, as shown in Fig. 2b. When the TBT content is further increased to 1.5 mL, some bulk metal compounds start to appear on the ITO surface, as shown in Fig. 2c1, impeding water vapor transport and potentially leading to the rupture of the PEM. Meanwhile, the nanorod array becomes inconsistent in length and uneven in thickness, as illustrated in Fig. 2c2. At a TBT content of 2.0 mL, the prepared TiN completely loses its directional alignment. A large amount of bulk materials is distributed on the ITO surface, blocking the channels for material exchange, rendering it unsuitable as a catalyst support, as shown in Fig. 2d. It is evident that the TBT content in the hydrothermal reaction precursor solution is significantly correlated with the surface morphology of the TiN nanorod array. A higher TBT content initially enhances and then degrades the vertical alignment of TiN nanorods, along with more bulk metal compounds on the surface. Therefore, they are unfit as a catalyst support. So, we choose a TBT content of 1.0 mL to create an ordered catalyst support. Industrial Chemistry & Materials

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During the hydrothermal reaction, hydrochloric acid acts as an inhibitor of TBT hydrolysis, so the ratio of water to hydrochloric acid in the solution has a significant effect on the uniformity and thickness of the whiskers, and only a suitable ratio can yield catalyst layer supports that meet the requirements. When the ratio of water to acid is $1:2$, the resulting nanorods are extremely small and exhibit poor uniformity, accompanied by noticeable agglomeration. Irregular spherical particles are observed to be irregularly distributed on the ITO surface, as shown in Fig. 3a. On the other hand, when the ratio of water to acid is $1:1$, the whiskers exhibit a highly uniform and smooth surface distribution, with appropriate thickness and spacing, making it an ideal and ordered catalyst layer support, as illustrated in Fig. 3b. As the water content increases, the inhibitory effect of dilute hydrochloric acid on the hydrothermal reaction weakens. When the water-to-acid ratio is 2 : 1, a large number of irregular and large-scale aggregations appear on the ITO surface. Furthermore, at high magnification, the whiskers are observed to become very thick with minimal spacing, as shown in Fig. 3c. When the water-to-acid ratio is 3 : 1, the ITO surface no longer possesses whiskers, it forms a highly dense coating with large spherical aggregations instead, as shown in Fig. 3d1. In observation under high magnification, the whiskers at the aggregation sites are extremely short and exhibit uneven thickness, rendering them unsuitable as catalyst supports, as illustrated in Fig. 3d2. Based on the findings, the final choice was a water-to-acid ratio of $1:1$ to achieve a smooth surface morphology, moderate whisker thickness, and reasonable spacing.

2.2 Nanoarray element composition determination

EDS tests were conducted on the samples to determine the composition of the product after the nitridation reaction, and the results are presented in Fig. 4. The surface of the samples exhibited clear detection of three elements: titanium (Ti), nitrogen (N), and oxygen (O). Titanium was identified as the predominant element, accounting for 64.27% of the mass and 35.67% of the atomic composition. Nitrogen was the second most abundant element, comprising 21.02% of the mass and 39.89% of the atomic composition. Additionally, a residual amount of oxygen was observed, constituting 14.71% of the mass and 24.44% of the atomic composition. These findings provide compelling evidence for the successful conversion of whiskers into TiN through the nitridation reaction. However, the presence of oxygen suggests that the nitridation reaction may not have reached complete

Fig. 3 The SEM images of the TiN ordered array: the ratios of water to hydrochloric acid in the hydrothermal reaction solution were (a) 1: 2, (b) 1: 1, (c) 2 : 1, and (d) 3 : 1, respectively.

Fig. 4 The EDS layered images of (a) TiN ordered array, (b) Ti, (c) N, and (d) O, respectively.

conversion, resulting in a core–shell structure with an outer layer of TiN and an inner layer of TiO₂. Furthermore, the thinness of the nanorod array necessitates consideration of potential interference from the quartz glass substrate during EDS analysis. In conclusion, the EDS experiment confirms the synthesis of uniformly distributed TiN nanorod arrays, which hold promise as catalyst supports within the internal region of the MEA for PEMFCs.

2.3 Fabrication results of double-side ordered MEA

The photographs showcasing the manufacturing process of the ordered membrane electrode assembly (MEA) are presented in Fig. 5. Initially, the desired morphology, size, and spacing of a $TiO₂$ nanorod array were synthesized on a 3

Fig. 5 Fabrication of the double-side ordered MEA: (a) ITO after the hydrothermal reaction; (b) ITO after the nitriding reaction; (c) MEA prepared by hot pressing; (d) ordered-MEA after PET encapsulation. Fig. 6 Cross-sectional view of the catalyst layer in the MEA.

 \times 3 cm² indium tin oxide (ITO) substrate using a seedassisted hydrothermal reaction as shown in Fig. 5a. The resulting ITO surface exhibited a characteristic white color, attributed to the physical properties of $TiO₂$. Subsequently, the $TiO₂(Q)$ ITO sample was subjected to a nitridation process within a tubular furnace under an ammonia atmosphere. Consequently, the ITO surface transformed into a distinct black color, predominantly emanating from the TiN nanorod array's properties (Fig. 5b). In the process of ultrasonic spraying, isopropanol was used as the dispersant of the catalyst, and Nafion solution was used as the proton conductor. The volume ratio of isopropanol to water was 5, the mass ratio of Nafion solution to PtC was 0.7, and the PtC content in the slurry was 2 mg ml⁻¹. The four components were added to the vessel in the order of PtC, water, isopropanol, and Nafion solution, and then ultrasonically mixed for 1 h in an ice bath to form the ultrasonic spray coating slurry. At this stage, the platinum-based catalyst was effectively loaded onto the black surface using ultrasoundassisted spray coating to ensure their complete infiltration into the array's interior. In contrast to the GDE configuration, the catalyst does not get trapped in the micropores of the carbon paper during the coating and transfer process. Instead, it is evenly distributed on both sides of the PEM with minimal waste, leading to a high level of catalyst utilization efficiency in this approach. Due to the much higher diffusion of hydrogen gas at the anode than oxygen gas at the cathode, in this study, by controlling the ultrasonic spray time and nozzle speed, the platinum loading of the cathode was determined to be 0.2 mg cm^{-2} , while that of the anode was 0.1 mg cm−² . It should be noted that due to the weak bonding between TiN and ITO, and the susceptibility of thin TiN layers to fracture, the spray coating process required the adjustment of the load-carrying airflow to its minimum. Subsequently, two catalytically-loaded ITO substrates sandwiched a 5×5 cm² proton exchange membrane (PEM) during a thermal transfer press operation, resulting in the preparation of the catalyst-coated membrane (CCM). Carbon papers were then placed on both sides of the CCM to form a complete MEA (Fig. 5c). Finally, to mitigate moisture-induced **Fraction Context Commons Articles.** The company of the stationary and the stationary of the stationary and the stationary and the stationary of the stationary and the stationary of the stationary of the stationary of the

deformation, the prepared MEA underwent PET encapsulation before final testing. The resulting ordered MEA exhibited a total area of 8 \times 8 cm², with an effective active area of 3×3 cm², as shown in Fig. 5d.

We sectioned the CCM after transfer hot-pressing to investigate the morphology of the TiN nanorod array in the MEA by SEM, as depicted in Fig. 6. The cross-sectional image demonstrates that the TiN nanorod array was entirely transferred to the PEM and retained its original shape well. The CL had a thickness of 4.2 μm, and the TiN nanorod array displayed good orientation. Moreover, the upper and lower surfaces of the ordered CL were smooth, with a flatness variation of only 0.6 μm, which could not only ensure a snug fit with the PEM without causing puncture damage, but also diminish the dead zone between the CL and diffusion layer, enhancing the expulsion of water. Owing to the ultrasonic spraying before transfer, the catalyst in CL exhibited a denser feature near the PEM side, which was favorable to exploit the catalyst to a greater degree.

2.4 Electrochemical properties of double-side ordered MEA

The performance of the ordered MEA can be evaluated by the polarization curve and the power density curve, as shown in Fig. 7. The polarization curve reveals how the voltage changes with the current density, whereas the power density curve demonstrates how the output power changes with the current density. The fabricated MEA demonstrates an initial performance that achieves a peak power density of 678.30 mW cm−² . When the output voltage reaches 0.6 V, it can be considered as the working voltage of the MEA. Due to the larger reactive surface area and superior catalyst dispersion, the working current density remains stable at 971.62 mA cm−² , with an output power density of 583.08 mW cm−² . During the operation of fuel cells, high-power operation is inevitable, which involves high current density but lower voltage. In such cases, water flooding and electrode material corrosion are prone to occur. However, thanks to the efficient mass transfer channels provided by the TiN nanorod arrays, the ordered MEA fabricated showed a nearly linear

Voltage of MEA

Voltage after AST

Power density of MEA

Power density after AST

1000

Current Density (mA cm⁻²)

1500

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500

relationship between the increased current density and output voltage, without experiencing severe voltage decay. This observation indicates that the internal membrane electrode can still maintain a favorable oxygen reduction reaction (ORR) process and efficient removal of product water.

In this study, we employed the chemical accelerated stress test (AST) for MEAs provided by the U.S. Department of Energy (DOE) to investigate the durability of the ordered MEAs fabricated. The electrochemical performance of the MEAs was also evaluated. After AST operation, the performance of the membrane electrode can decline due to various factors such as chemical degradation of the proton exchange membrane, catalyst deactivation, and carbon support corrosion. As shown in Fig. 7, the performance of the membrane electrode in the low current density region showed a minimal decline, while a significant drop was observed in the high current density region. This discrepancy can be attributed to the dominant influence of activation polarization overpotential in the low current density region, whereas the high current density region is mainly affected by the ohmic polarization overpotential and concentration polarization overpotential. The ordered membrane electrode exhibited a 4.8% decrease in output performance at a working voltage of 0.6 V. However, as the current density increased, the decline in the membrane electrode performance became more pronounced, with a relative decrease of 7% in the MEA current density when the output voltage reached 0.3 V. Simultaneously, the peak power density decreased to 642.13 mW cm^{-2} , and there was a noticeable leftward shift in the peak power inflection point. Due to its exceptional chemical stability and excellent corrosion resistance and conductivity, TiN has been shown in multiple studies to exhibit outstanding corrosion resistance when structured in nanoscale arrays. $44-46$ It can operate stably in acidic, neutral, and alkaline electrolytes for extended periods without altering its morphology and structure. Therefore, we believe that even in the presence of an ordered carrier, prolonged operation at high current densities may lead to the loss of catalytic activity and dispersion, causing some loosely attached catalysts on TiN to aggregate, resulting in increased ohmic polarization and concentration polarization overpotentials. **Contractive State Article Contract Article Consumer Contract Consumer Consume**

> To comprehensively evaluate our double-side ordered MEA based on TiN nanorod arrays, we conducted extensive research on current typical PEMFC electrode structures and preparation methods. Table 1 showcases various MEA structures prepared using different methods. The MEAs with disordered structures typically necessitate higher platinum loading to attain optimal performance, as depicted in Table 1. This is attributed to their internal multiphase reaction channels being in a disordered state, resulting in generally lower platinum utilization. The MEAs prepared through ultrasonic spraying and transfer printing exhibit similar performance but display distinct characteristics. Ultrasonic spraying, owing to its superior catalyst dispersion,

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MEA structural Preparation process Pt load (mg cm−²) Peak power $(mW cm^{-2})$ Current density@0.6 V $(mA cm^{-2})$ Disordered Ultrasonic spraying 0.4 695 1100 40 Disordered Transfer printing 0.5 920 540 540 41
Cathode-ordered Ultrasonic spraving 0.2 538.33 819.57 42 Cathode-ordered Ultrasonic spraying 0.2 538.33 819.57 42
Cathode-ordered Active screen plasma pitriding 0.19 640 900 43 Cathode-ordered Active screen plasma nitriding 0.19 640 900 900 43
Doubleside-ordered Ultrasonic spraying & decal transfer 0.2 678.30 917.62 Th Ultrasonic spraying & decal transfer 0.2 678.30 917.62 This work **Paper**
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demonstrates exceptional working voltage output performance, while transfer printing exhibits higher peak power. Interestingly, ultrasonic spraying and active screen plasma nitriding processing show similar performance, but the former is obviously faster and cheaper in comparison. Although the cathode single-sided ordered MEA can achieve performance similar to disordered structures with lower catalyst content, there still exists a discernible gap. It can be seen from the comparison that the cathode ordering has a significant improvement on the performance of the MEA, which can achieve a similar performance to the conventional

MEA with only half of the platinum loading. This indicates that the ordered structure has a promoting effect on the ORR, which benefits from its provision of good mass transfer and water drainage channels. By comparing the bilateral and cathode unilateral ordered structures, it can be found that the performance of the membrane electrode has been further improved. This is because the fuel cell involves two halfreactions at the anode and cathode, and only when both sides are at a high level can a better cell performance be obtained. Therefore, it can be proved that the anode ordered structure can also provide more reaction sites for the hydrogen oxidation reaction.

In this study, we utilized a combination of ultrasonic spraying and transfer printing techniques to create a doubleside ordered MEA utilizing TiN nanorod arrays. This innovative approach resulted in output performance comparable to that of traditional methods but with only half the platinum loading. The fabrication of double-side ordered MEAs through ultrasonic spraying and transfer printing showcases a promising, efficient, stable, and low-platinum solution.

3 Conclusions

This paper presents a novel double-side ordered MEA for PEMFCs based on TiN nanorod arrays. The arrays were synthesized by a seed-assisted hydrothermal reaction and a subsequent nitriding treatment, and then transferred to both sides of the PEM by using ultrasonic spraying and transfer printing. The morphology and density of the TiN nanorod arrays were controlled by adjusting the concentration of $TiCl₄$ in the seed solution, the content of TBT and the ratio of water to hydrochloric acid in the hydrothermal reaction solution. The TiN nanorod arrays exhibited high electrical conductivity, corrosion resistance, and catalytic activity as) Ref.

catalyst supports for PEMFCs. The ordered structure of the catalyst layers provided well-defined channels for reactant gas diffusion and product water removal, as well as a large surface area for catalyst loading and reaction sites. The fabricated ordered-MEA showed high performance and durability under various operating conditions. The peak power density reached 678.30 mW cm−² , and the working current density remained stable at 971.62 mA cm−² at a working voltage of 0.6 V. After AST operation, the ordered-MEA exhibited a relative decrease of 4.8% in output performance at a working voltage.

This study provides a new approach for developing highperformance, low-platinum MEAs. In future work, we will further optimize the membrane electrode structure design and preparation process with the goal of high power density through a combination of MEA half-reaction simulation and experimental verification. Additionally, we will conduct research on high-power density single-cell packaged fuel cells.

4 Experimental section

4.1 Structure of double-side ordered MEA

The MEA is an essential component of proton exchange membrane fuel cells (PEMFCs) that can efficiently convert the chemical energy from hydrogen and oxygen into electricity. Here, we present a novel double-side ordered MEA based on TiN nanorod arrays. These nanorod arrays are prepared by seed-assisted hydrothermal reaction and nitriding treatment. Meanwhile, the catalyst loading is achieved through the ultrasonic spraying method. The structure of the double-side ordered MEA is illustrated in Fig. 8a, and it adopts a seven-in-one configuration, comprising the anode gas diffusion layer (AGDL), anode micro-porous layer, anode ordered catalyst layer (ACL), proton exchange membrane (PEM), ordered cathode catalyst layer (CCL), cathode micro-porous layer, and cathode gas diffusion layer (CGDL). Among these layers, the catalyst layer plays a crucial role in facilitating the major electrochemical reactions within the fuel cell. It requires materials that are corrosion-resistant, have low impedance, and exhibit efficient channels to ensure unobstructed mass transport and timely removal of product water. To fulfill these requirements, we develop ordered catalyst layers composed of TiN nanorod arrays that are directly constructed on the substrate using a hydrothermal reaction and then transferred to both sides of

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Fig. 8 (a) Structure of double-side ordered MEA structure, and working principle of (b) traditional MEA and (c) ordered MEA.

the PEM through the transfer printing process. The TiN arrays are oriented perpendicularly to the micro-porous layer and PEM. This ordered catalyst layer structure provides additional reaction sites for oxygen, protons and electrons, thus efficiently facilitating the removal of generated water. These features contribute to the enhanced promotion of oxygen reduction reaction (ORR).

4.2 Working principle

As for conventional MEAs, the catalyst particles are dispersed in a disordered manner on both sides of the PEM, leading to tortuous or even blocked transport channels, resulting in detrimental factors such as water flooding, polarization corrosion and catalyst agglomeration, as illustrated in Fig. 8b. The working principle of the ordered MEA is shown in Fig. 8c. Due to the small molecular weight of hydrogen, it can easily diffuse into the anode catalytic layer. Therefore, the structured anode catalytic layer can provide a large number of catalytic sites, promoting the decomposition of hydrogen into protons and electrons. When the protons are conducted to the cathode through Nafion and PEM, the electrons reach the cathode through the external circuit, and combine with oxygen in the reduction reaction to form water. The ordered nanorod TiN arrays offer an optimal environment for mass transfer and provide a large reaction surface area for the ORR, resolving the issues associated with the conventional MEA's disordered structure. Furthermore, thanks to TiN's corrosion resistance and low electrical resistance properties, the electrode's vulnerability to damage from high potentials and strong acids is effectively reduced. Consequently, the multi-phase mass transfer channels remain unobstructed. Moreover, the catalyst is loaded by ultrasonic spraying and decal transferring, so that the catalyst concentration increases from the GDL to PEM gradually, guaranteeing a high catalyst utilization.

The function and advantages of the TiN double-side ordered MEA are mainly reflected in its highly ordered channel structure, which can effectively promote mass transfer and water management, while maintaining high catalytic activity and stability. Firstly, the channel structure of the double-side ordered MEA can increase the permeability of hydrogen and air, thereby improving the utilization of reactants and the performance of the cell. Due to the low mass transfer resistance of the TiN nanorod array channel structure, the gas diffusion ability can be significantly improved. Secondly, due to the radial capillary force and low surface tension of the channel structure of the double-side ordered MEA, the water discharge speed can be significantly increased, thus avoiding water accumulation and blockage, and improving the stability and durability of the cell.

The proposed ordered-MEA offers several benefits, including: (1) enhanced mass transport through the implementation of structured anodes and cathodes, (2) increased durability attributed to the corrosion resistance and low electrical resistance properties of TiN, and (3) improved catalyst utilization resulting from the gradient distribution of catalyst concentration.

4.3 Fabrication process of double-side ordered MEA

The preparation process of the double-side ordered MEA can be illustrated as shown in Fig. 9; the green, yellow and red arrows represent the fabrication process of nanorod arrays, the loading and transfer process of catalysts, and the manufacturing process of the ordered-MEA, respectively. Hence, the fabrication process of the double-side ordered MEA can be mainly divided into three parts: preparation of an ordered carrier for the catalyst layer, ultrasonic spraying and decal transferring of the catalyst, and thermal pressing encapsulation of the membrane electrode. These correspond to the upper, middle, and lower sections of Fig. 9.

The key process in the fabrication of the double-side ordered MEA lies in the rational preparation of TiN nanorod arrays. The nanorod arrays on ITO were synthesized using a seed-assisted hydrothermal method. The seed-assisted

hydrothermal process is rather simple and does not require complex conditions to obtain products with complete crystalline shape, uniform particle size distribution and good dispersion. For the fabrication of TiN nanorod arrays, $TiCl₄$, tetrabutyl titanate (TBT), concentrated hydrochloric acid, ethanol and isopropanol were purchased from HUSHI® (Shanghai, China). All chemicals were of analytical grade and were used as received without further purification. A detailed description of the fabrication process will be provided, which consists of 3 procedures corresponding to the green arrows in Fig. 9. The specific steps are as follows:

Step 1-Synthesis of seed crystals: a 3 cm \times 3 cm ITO substrate was subjected to ultrasonic cleaning in a solution containing deionized water, ethanol, and acetone (in a volume ratio of $1:1:1$) for 20 min. Subsequently, the ITO substrate was heated to 120 °C and dried for another 20 min. To ensure that only the conducting layer remains exposed, the non-conductive surface of the cleaned ITO was carefully sealed with polytetrafluoroethylene (PTFE) tape. Then, the cleaned ITO glass substrate was immersed in a TiCl₄ chloride aqueous solution and heated at 70 °C for 15 minutes in a water bath. During this process, $TiCl₄$ underwent hydrolysis reaction and formed $TiO₂$ seeds on the surface of ITO. The corresponding chemical reaction can be described as

$$
TiCl_4 + 2H_2O \triangleq TiO_2 + 4HCl
$$
 (1)

The ITO glass was taken out from the solution and annealed at 400 °C for 30 min to remove the surface stress and the impurities involved on the surface.

Step 2-Seed assisted hydrothermal reaction: to prepare the hydrothermal reaction precursor, a TBT solution with a volume ranging from 0.5 to 2.0 ml was added to a mixture of 20 mL concentrated hydrochloric acid and 20 mL deionized water. The resulting solution was stirred for a period of 15 min. Subsequently, the mixture along with the ITO substrate coated with TiO₂ seeds was transferred into a 100 mL Teflonlined stainless-steel autoclave. During the transfer, it was crucial to position the conducting layer with seeds facing downward to prevent undesired $TiO₂$ deposition, which could lead to changes in whisker morphology or stomatal blockage. The hydrothermal treatment commenced with heating at a rate of 5 °C min−¹ and was maintained at a temperature of 170 °C for a duration of 9 hours. The ensuing chemical reactions occurring during this step are

$$
Ti(O-C4H9)4 + 2H2O = TiO2 + 4CH3(CH2)3OH
$$
 (2)

Finally, the autoclave was cooled to room temperature within a fume hood. The treated ITO was cleaned with deionized water and ethanol to remove excess ions, then dried by heating at 130 °C. To ensure the cleanliness and purity of the $TiO₂$ nanorod arrays, the cleaning step mentioned above was repeated a minimum of 5 times. This rigorous repetition was undertaken to guarantee the absence of any contaminants or impurities on the surface of the $TiO₂$ nanoarrays.

Step 3-Nitriding treatment: the last step involves nitridation of the TiO₂ nanorod arrays generated by the seed-assisted hydrothermal reaction to convert the product into TiN. The nitridation process was carried out by suspending the synthesized $TiO₂$ nanorod arrays in an alumina boat and placing it in a tube furnace. The temperature was raised to 450 °C under a nitrogen atmosphere, followed by switching the gas source to ammonia. The sample was then heated to 850 °C and kept at that temperature for 3 hours while maintaining an ammonia atmosphere. Subsequently, the gas source was switched back to nitrogen and the temperature was gradually cooled down to room temperature. The chemical reaction equation during this process is

$$
6TiO2 + 8NH3 \triangleq 6TiN + 12H2O + N2
$$
 (3)

The previously white surface of the ITO substrate turns black, indicating the coating of TiN onto the ITO surface.

4.4 Characterization tests

To characterize the performance of the fabricated doubleside ordered MEA, the following tests were conducted. The single fuel cell is evaluated at ambient pressure and the temperature was set to constant. The reactants can be fed into the fuel cell at a fixed flow rate to meet the reaction requirements.

4.4.1 Scanning electron microscopy (SEM). SEM was used to investigate the influence of different process parameters on the formation of TiN nanorod arrays. Through SEM images, we can clearly observe the shape, size, and arrangement of the TiN nanorods. Additionally, SEM is utilized to detect surface defects and potential contaminants. To perform data analysis on the SEM images, image processing software is employed to measure parameters such as the length, diameter, and spacing of the nanorods. By statistically comparing these parameters under different experimental conditions, we can assess the extent to which process parameters affect the morphology of the nanorods. This aids in validating our control methods and provides crucial guidance for further process optimization.

4.4.2 Energy dispersive spectrometry (EDS). EDS was primarily utilized to examine the successful conversion of the nanorod array into TiN following the nitridation process. To determine the specific composition of the nanorod array, the surface of ITO was examined before and after the nitridation reaction. During the EDS analysis, X-ray signals emitted from the sample were collected and processed to identify characteristic energy peaks associated with different elements present in the nanorod array. By comparing the obtained spectra with reference spectra of known elements, the elemental composition of the nanorod array was determined. Additionally, elemental mapping using EDS was employed to investigate the spatial distribution of elements within the nanorod array, providing crucial evidence for the desired chemical transformation. This comprehensive analysis utilizing EDS played a pivotal role in elucidating the conversion mechanism and confirming the successful synthesis of TiN within the nanorod array.

4.4.3 Polarization curve tests. The polarization curve is a critical parameter for evaluating the performance of the MEA. In this study, a single cell was utilized to collect the polarization curve data, aiming to comprehensively characterize the electrochemical performance of the MEA. Before starting the test, the MEA was activated using the method described by the US Fuel Cell Council. The MEA was first held at 0.6 V for 60 min, then cycled between 0.7 V and 0.5 V nine times, each for 20 min. Finally, the MEA was operated at a constant current of 200 mA cm−² for 12

hours. To ensure consistency, the gas temperatures at the anode and cathode were precisely controlled at 80 °C. To obtain the polarization curves, the current was measured at various applied voltages. Each voltage was maintained for a duration of 10 minutes, allowing enough time for the system to reach a steady state. Data collection was conducted at a sampling rate of 6 points per minute, ensuring an adequate number of data points for accurate analysis. The obtained polarization curve data were then subjected to detailed analysis. The current–voltage relationship was plotted, and key performance parameters such as the open circuit voltage, peak power density, and cell voltage at a specific current density were determined. This rigorous experimental procedure allowed for a comprehensive characterization of the electrochemical performance of the MEA. Industrial Chemistry & Materials
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4.4.4 Accelerated stress test (AST). The AST can simulate the performance degradation of fuel cells after prolonged operation, serving as a metric for the activity and stability of the catalyst layer as well as the degradation of the membrane electrode. This study adopts the AST testing method proposed by the U.S. DOE, with a temperature of 90 °C and a relative humidity of anode/cathode 30/30%. The hydrogen/air inlet is set at stoichiometric ratios of 10/10 at 0.2 A cm^{-2} equivalent flow, and the inlet gas pressures for the anode and cathode are 250/200 kPa, respectively. By comparing the performance before and after testing, we aim to validate the durability of the double-side ordered TiN-coated electrode.

Conflicts of interest

There are no conflicts to declare.

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