RSC Advances



View Article Online

View Journal | View Issue

PAPER

Check for updates

Cite this: RSC Adv., 2022, 12, 4145

Received 13th January 2022 Accepted 27th January 2022

DOI: 10.1039/d2ra00246a

rsc.li/rsc-advances

1 Intorduction

A µDMFC is a portable and removable power source while it has high energy conversion efficiency, simple structure, easy operating conditions, low pollution, and is economical.1-3 It relies on the conversion of chemical energy of redox reactions into electrical power. In this process, the function of the current collector is to distribute the reaction gases and collect the incoming electrons. Therefore, the current collector material should have high electrical conductivity, corrosion resistance, hydrophobicity, permeability, etc. 4-6 The different materials and field structures have a significant impact on the performance of the µDMFC. Researchers did a lot on the materials and flow field structures of this. In 2000, Makkus et al.7 prepared a stainless steel collector plate, and the experimental results showed that contamination occurs between the stainless steel collector and the membrane electrode assembly (MEA) on the anode side. In 2010, Zhang et al.8 designed an experiment using titanium-coated stainless steel as a collector plate, and the TiN coating could reduce corrosion and contact resistance. In 2018, Li et al.9 used stainless steel to prepare a current collector for fuel cells with improved discharge stability. In 2021, Choi et al.¹⁰ proposed a Gas Diffusion Layer (GDL) of FT as an anode and

Performance study of μ DMFC with foamed metal cathode current collector

Fan Zhang, 跑 a Yanhui Zhang, 跑 a Chuan Luo, a Dacheng Zhang ab and Zhengang Zhao 跑 *ab

Micro Direct Methanol Fuel Cells (μ DMFCs) often have application in moveable power due to their green and portable nature. In a μ DMFC's structure, a current collector of the μ DMFC needs to have high corrosion resistance such that the μ DMFC can work for a long time in a redox reaction and respond to variable environmental conditions. To this end, four cathode current collectors were prepared. The materials selected were foam stainless steel (FSS) and foam titanium (FT), with fields of hole type and grid type. The performance of μ DMFC with different cathode collector types was investigated by I–V–P polarization curves, Electrochemical Impedance Spectroscopy (EIS), and discharge test. The experimental results show that the maximum power density of the hole-type FSS cathode current collector μ DMFC (HFSS- μ DMFC) is 49.53 mW cm⁻² at 70 °C in the methanol solution of 1 mol L⁻¹, which is 70.15% higher than that of the hole-type FT cathode current collector μ DMFC (HFT- μ DMFC). The maximum power density of the grid-type FS cathode current collector μ DMFC (GFSS- μ DMFC) is 22.60 mW cm⁻², which is 11.99% higher than that of the grid-type FT cathode current collector μ DMFC (GFT- μ DMFC). The performance of the HFSS- μ DMFC is optimal in the methanol solution of 1 mol L⁻¹.

> improved the performance of the cells in accelerated corrosion tests. Stainless steel and titanium are thus highly represented in the preparation of collector plates. To further enhance the performance of fuel cells, many scholars have used foam metals to prepare current collectors. In 2003, Kumar et al.¹¹ used foam metals to prepare bipolar plates for fuel cells and investigated the effect of foam metal permeability on fuel cell performance. In 2018, Park et al.12 used graphene foam metals to prepare current collectors and investigated their effect on mass transport of reactants and products. The experimental results showed that graphene foam metal collectors exhibited lower mass-transport resistance than conventional collectors. In 2018, Shin et al.13 embedded foam metals in collectors, which significantly enhanced the maximum power of fuel cells. In 2021, Sun et al.14 used foam metals to prepare collectors and demonstrated that foam metals have great potential for battery applications. Also, scholars have studied the flow field extensively. In 2010, Spernjak et al.15 investigated the effect of different flow fields on the cell performance. The experimental results showed that the serpentine flow field had a more stable output, and the parallel and staggered flow fields exhibited a higher water content and lower differential pressure. To improve the performance of µDMFC, the researchers tried many fields in collectors. In 2012, Friess et al.16 designed a radial flow field structure, and the cell with this structure outperformed the cell with parallel flow field structure under the same experimental conditions. In 2013, Guo et al.¹⁷ prepared an optimized needle flow field and improved the performance of the cells by

^aFaculty of Information Engineering and Automation, Kunming University of Science and Technology, Kunming 650500, China. E-mail: zhaozhengang@kust.edu.cn ^bYunnan Key Laboratory of Green Energy, Electric Power Measurement Digitalization, Control and Protection, Kunming, China

this design. In 2019, Wilberforce *et al.*¹⁸ investigated the performance of the cells with different flow field structures. The type of flow field is related to the mass transfer and current density of the cell. Thus, it is important to choose a suitable flow field.^{19,20}

In summary, stainless steel and titanium have good potential for current collector fabrication. Also, foam metals can effectively increase the oxygen transport channels and the reaction area in the cathode collector. And the wettability of the collector material directly affects the performance of μ DMFC. Therefore, the material selections are hydrophilic FT and hydrophobic FSS for preparing the cathode collector. In this study, four μ DMFC with different cathode collectors (HFSS- μ DMFC, GFSS- μ DMFC, HFT- μ DMFC, and GFT- μ DMFC) were assembled. The performance of the μ DMFC was evaluated by I-V-P polarization curves, EIS, and discharge test. Finally, the influence of material, wettability, and flow field on cathode gasliquid flow were analyzed to find a suitable cathode current collector.

2 Materials and methods

2.1 µDMFC structure

 μ DMFC consists of Cathode Current Collector (CCC), Anode Current Collector (ACC), Membrane Electrode Assembly (MEA), Teflon Gaskets (TG), Cathode Extremity Plate (CEP) and Anode Extremity Plate (AEP), and Liquid Storage Cavity (LSC).²¹ The chemical reaction equation within μ DMFC is as follows,

total chemical reaction:

$$CH_3OH + \frac{3}{2}O_2 \rightarrow CO_2 + 2H_2O \tag{1}$$

anode reaction:

$$CH_3OH + H_2O \rightarrow CO_2 + 6H^+ + 6e^-$$
 (2)

cathodic reaction:

$$\frac{3}{2}O_2 + 6H^+ + 6e^- \to 3H_2O$$
 (3)

As shown in Fig. 1, to observe the internal reaction of μ DMFC during operation, the CEP, AEP, and the LSC are prepared from acrylic plates. Its open window frame is 1.44 cm². The effective liquid storage area of LSC is 2.5 cm³ and the effective area of the

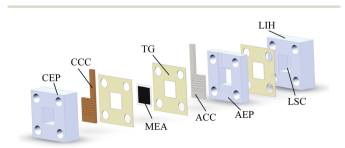


Fig. 1 Schematic diagram of µDMFC.

MEA is 1 cm². The TG is placed between the LSC and the AEP to prevent methanol liquid leakage from the reservoir chamber. The TG is placed between the current collector and the MEA to make close contact and prevent the methanol liquid on the anode side leak to the cathode. The AEP and CEP press the current collector closely to the MEA to reduce its contact resistance. A 3 mm diameter Liquid Injection Hole (LIH) is located above the LSC for liquid injection and CO₂ venting.²²

For the MEA preparation, the proton exchange membrane of choice is Nafion 117 (DuPont, Wilmington, DE, USA). The anode catalyst is 50% Pt and 25% Ru, and the cathode catalyst was 60% Pt. The catalyst loadings for the anode and cathode are 4 mg cm⁻² and 2 mg cm⁻², respectively. Finally, the carbon paper with catalyst is hot-pressed together with the proton exchange membrane to obtain the MEA. For the CCC preparation, the selected materials are FSS and FT to prepare the cathode collector.23 The foam metal is sintered under vacuum conditions using the powder metallurgy method. Their surface form is shown in Fig. 2. It can be seen that the foam metal has a three-dimensional porous structure, which can provide a path to solutions for problems in materials and flow fields.24-27 Further, the sintered foam metal is cleaned and placed on the laser cutting platform (Model 6060L to 1000W) to cut according to the design. Finally, different types of foam metal CCC are prepared.

The opening ratio of the cathode collector is calculated as follows:

$$\beta = \frac{A_{\rm fd}}{A_{\rm eff}} \tag{4}$$

where $A_{\rm fd}$ is flow field area and $A_{\rm eff}$ is effective area of current collector in contact with MEA. The opening ratio of the cathode collector was 38.5%,^{28–30} the flow field was hole and grid type, as shown in Fig. 3. The ACC remains unchanged. It is stainless steel and hole-type flow field, with the specifications of the CCC. The contact angle of the FSS was 96.525°, and the FT was 70.751°, as shown in Fig. 4. The FT has a contact angle of less than 90° while the FSS is more than 90°. Therefore they are hydrophilic and hydrophobic, respectively.³¹

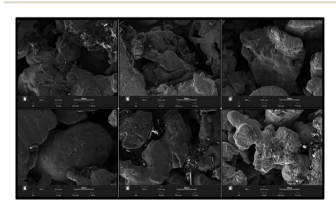


Fig. 2 Surface characterisation of the foam metal.

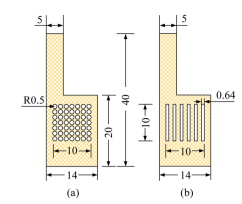


Fig. 3 Specification and flow field structure of cathode collector, unit: mm. (a) Hole type (b) grid type.

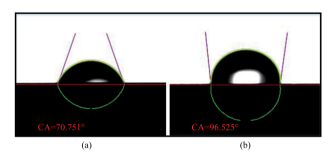


Fig. 4 Contact angle test. (a) Foam titanium; (b) foam stainless steel.

2.2 Preparation and experiment of µDMFC

For the µDMFC preparation:²²

(1) Milled the LSC, AEP, and CEP. Then, cut the TG according to the endplate shape. Cut the foam metal plate according to the cathode current collector and the flow field size. Finally, polish the surface of the current collector smoothly.

(2) Put the processed parts of μ DMFC into the ultrasonic cleaning machine and ultrasonically clean them with CH₃OH, C₂H₅OH, and deionized water in turn for 15 min to remove the oil on the surface. Then, put them into the oven at 100 °C to dry.

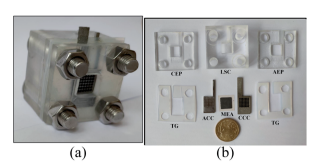
(3) Place a TG between the LSC and the AEP, between the MEA and the endplate to prevent fuel leakage and provide support protection for the MEA.

(4) Assembled the μ DMFC and fixed it with screws to obtain μ DMFC, as shown in Fig. 5.

The four different types of μ DMFC were HFT- μ DMFC, HFSS- μ DMFC, GFT- μ DMFC, and GFSS- μ DMFC. They use the same components. A hydraulic press is used to control the pressure of the battery assembly.

The test setup in this study consists of a DC electronic load (Model IT8511A+), an electrochemical workstation (Model CHI660E), and a temperature-controlled oven, as shown in Fig. 6. The test temperature is 70 $^{\circ}$ C.

During the experiment, the μ DMFC was placed in a oven. The electronic load was attached to the cathode and anode of μ DMFC. The three electrodes of the electrochemical workstation were respectively to the cathode and anode of μ DMFC



View Article Online

RSC Advances

Fig. 5 Physical diagram of the battery. (a) Overall diagram (b) split diagram.

according to the actual test needs. The DC electronic load was used to discharge test the μ DMFC and record the voltagecurrent data. The electrochemical workstation was used to test the EIS and record the related data.

3 Results and discussion

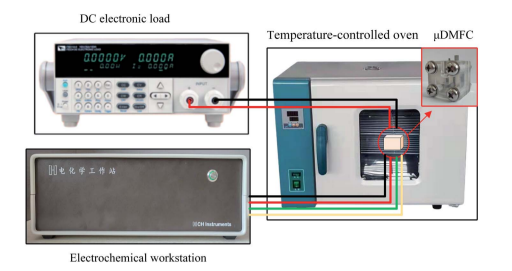
3.1 Analysis of polarization characteristics

To investigate the effect of four different types of cathode current collectors on the performance of μ DMFC. The power density tests of μ DMFC with various concentrations of methanol solutions (0.5–5 mol L⁻¹) were carried out at 70 °C. Their polarization curves are as in Fig. 7.

The power density, current density, and operating voltage of the FSS μ DMFC are higher than those of the FT μ DMFC at methanol concentrations of 0.5 mol L⁻¹ to 5 mol L⁻¹. The electrochemical and ohmic polarization of μ DMFC for four cathode current collectors is essentially the same at low concentrations of methanol solution. The concentration difference polarization of μ DMFC with FSS cathode collectors is superior to that of μ DMFC with FT cathode collectors. Differences in polarization properties appear with the increasing concentration of methanol solution. As the concentration increases, the power density of μ DMFC increases and then decreases.

The difference in the concentration polarization of μ DMFC for the four cathode collectors at 0.5–1 mol L⁻¹ methanol solution appeared. This is because of an increase in the reaction rate at the high potential region, resulting in insufficient oxygen supply. The electrochemical polarization, ohmic polarization, and concentrated polarization of μ DMFC for four cathodic current collectors at 2–5 mol L⁻¹ methanol solution concentrations are different. At the low potential region, the electrochemical polarization of FSS μ DMFC is less. It indicates that the molecular activation performance of FSS μ DMFC is better than that of FT μ DMFC. The concentration polarization of 3–5 mol L⁻¹ was significantly smaller than that of the grid-type flow field μ DMFC.

The power densities of μ DMFC with different types of cathode current collectors at various methanol concentration solutions are shown in Table 1 and Fig. 8. The maximum power density of HFSS- μ DMFC is 49.53 mW cm⁻² at 70 °C in 1 mol L⁻¹





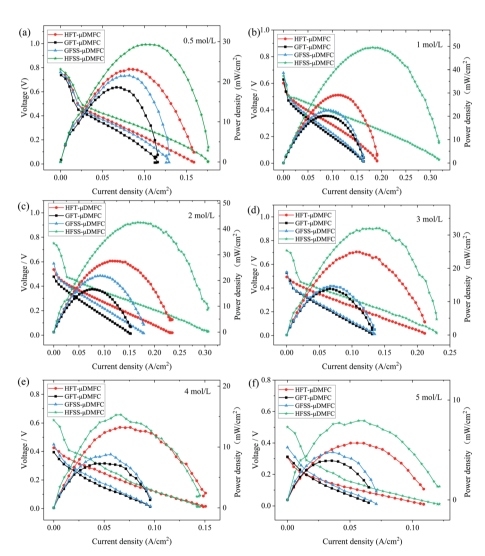


Fig. 7 Polarization curves of μ DMFC at different methanol solution concentrations. (a) 0.5 mol L⁻¹ (b) 1 mol L⁻¹ (c) 2 mol L⁻¹ (d) 3 mol L⁻¹ (e) 4 mol L⁻¹ (f) 5 mol L⁻¹.

Table 1 Maximum power density values for different current collectors at 70 $^\circ\text{C}$

Concentration $(mol L^{-1})$	$ m HFT-\mu DMFC$ $(m mW~cm^{-2})$	$ m GFT-\mu DMFC$ $ m (mW~cm^{-2})$	$ m HFSS-\mu DMFC \ (mW~cm^{-2})$	GFSS- μ DMFC (mW cm ⁻²)
0.5	23.11	18.61	29.26	21.55
1	29.11	20.18	49.53	22.60
2	27.58	16.63	42.56	21.89
3	24.89	13.69	32.01	14.66
4	13.23	7.29	15.29	8.78
5	5.70	3.93	7.94	4.78

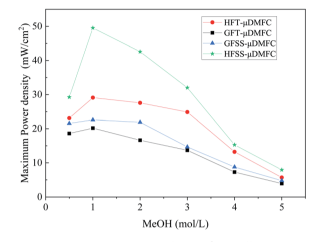


Fig. 8 Maximum power density of μ DMFC at different methanol solution concentrations.

methanol solution concentration, which is 70.15% higher than that of HFT- μ DMFC and 119.16% higher than that of GFSS- μ DMFC. The maximum power density of GFSS- μ DMFC is 22.60 mW cm⁻², which is 11.99% higher than that of GFT- μ DMFC. HFT- μ DMFC achieves a maximum power density of 29.11 mW cm⁻², which is 44.25% higher than that of GFT- μ DMFC. It implies that, under the same flow field structure, the performance of the FSS μ DMFC is better than that of the FT μ DMFC. Under the same collector plate material, the performance of the hole-type flow field μ DMFC is better than that of the grid-type flow field μ DMFC. In addition, compared to previously published work, the μ DMFC in this study has some enhancements in terms of maximum power density, as shown in Table 2.

The power density of the HFSS- μ DMFC at the methanol solution concentrations of 0.5–5 mol L⁻¹ is the highest, and the GFT- μ DMFC is the lowest. At 0.5–1 mol L⁻¹ methanol solution,

Table 2 Maximum power density for different works						
Work	Concentration (mol L^{-1})	Temperature (°C)	Maximum power density (mW cm ⁻²)			
Xue <i>et al.</i> ³²	3.00	25.00	23.80			
Chao et al. ³³	3.00	22.00-24.00	50.00			
Aricò et al.34	5.00	21.00	20.00			
Sudaroli et al.1	1.00	60.00	32.00			
This work	1.00	70.00	49.53			

the power density enhancement of the hole-type flow field μ DMFC is significantly higher than that of the grid-type flow field μ DMFC. As the concentration of methanol solution increases, the power density of μ DMFC starts to decrease. The main reason is that as the concentration of methanol solution increases, the electrochemical reaction rate and the methanol permeation increase, ³⁵⁻³⁷ resulting in the power density of the μ DMFC decreasing. The decrease of the power density of the μ DMFC is more visible at 1–4 mol L⁻¹ methanol solution. And the μ DMFC is more alleviated at 4–5 mol L⁻¹ methanol solution. High methanol solution concentration destroys less for the power density of the μ DMFC. This indicates that the excess methanol permeation severely affects the reduction reaction on the cathode side of μ DMFC.

3.2 Analysis of electrochemical impedance spectroscopy

To further investigate different flow field structures and materials, the EIS measurements of the μ DMFC at 70 °C in a methanol solution concentration of 1 mol L⁻¹ were performed.^{38,39} Fig. 9 gives the EIS at a discharge current density of 80 mA cm⁻2. In the EIS, the size of the semicircular in the highfrequency region indicates the magnitude of the Charge Transfer Resistance (CTR) of the μ DMFC. The larger this semicircular is, the greater the CTR.¹⁴ The size of the semicircular in the low-frequency region indicates the magnitude of the mass transfer impedance of the μ DMFC. The larger the semicircular is, the higher the mass transfer impedance. In other words, the CTR and mass transfer impedance is small, the corresponding semicircular almost disappears.⁴⁰

In the Nyquist plot, the total impedance of the HFSS- μ DMFC is the smallest, and the GFT- μ DMFC is the largest. The CTR of

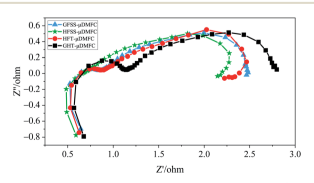


Fig. 9 Discharge of EIS with a current density of 80 mA cm^{-2} .

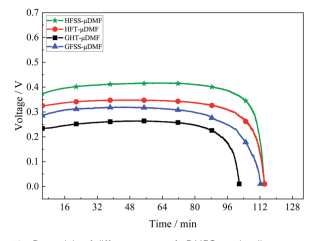


Fig. 10 Potentials of different types of µDMFC at a loading current of 80 mA cm⁻²

HFSS-µDMFC is the smallest, and the GFT-µDMFC is the largest. The total impedance of the GFSS-µDMFC is higher than that of the HFT-µDMFC, but the CTR is the same. The transfer impedance of the HFSS-µDMFC is lower than that of the HFTµDMFC. The transfer impedance of the GFSS-µDMFC is lower than the GFT-µDMFC.

Under the same flow field structure, the total impedance of the FSS µDMFC is smaller than that of the FT µDMFC. Among them, the difference in the CTR between the FSS cathode collector µDMFC and the FT cathode collector µDMFC is higher. But the difference in transfer impedance is lower. It is mainly because the charge transfer conductivity of the FSS is higher than that of the FT. The value of the interfacial capacitance due to the charge accumulation effect at the interface between the electrode substrate and electrolyte affects the CTR. The larger the interfacial capacitance, the greater the CTR. The larger the CTR value, the more severe the polarization during the electrode reaction,35 and the polarization loss of the FT is greater than that of the FSS. The selected current collectors all have a three-dimensional porous structure with the same effective opening ratio, but the contact angle of FT is lower than that of FSS. At a discharge process, the water generated by the cathode is easily infiltrated into the microporous structure of the FT, which reduces the oxygen channel of the cathode and increases the impedance in the oxygen mass transfer process. It implies that the oxygen transfer rate of µDMFC of the FSS cathode current collector is better than that of the FT cathode current collector. The FSS has better hydrophobicity, the H₂O produced by the cathode being less to clog the micropores on the current collector. But the experiment is conducted at 70 °C, the water is easy to volatilize, resulting in the phenomenon of clogging the microporous is alleviated. Then, the difference is reduced between the mass transfer impedance of the FT cathode collector µDMFC and the FSS cathode collector µDMFC. The HFSS-µDMFC has the highest velocity constant and the charge conduction reaction rate. The high-frequency semicircular arc of the HFSS-µDMFC disappears. It implies that the charge reaction rate is faster than the mass transfer rate. At this

moment, the porous dynamics process is dominated by diffusion processes.

Under the same material, the total impedance of the holetype flow field µDMFC is smaller than that of the grid-type flow field µDMFC. This is because the oxygen transfer rate of the hole-type flow field µDMFC is better than that of the gridtype flow field µDMFC. The hole-type flow field has more even oxygen distribution.41-43 The hole-type flow field is more suitable than the cathode flow field structure. Among them, the CTR and mass transfer impedance of the hole-type flow field µDMFC are smaller than those of the grid-type flow field µDMFC. The flow field of the hole-type is distributed more uniformly.22 It causes the distribution of the electrochemical reaction area and the charge conduction reaction to be greater. Compared with the grid-type flow field, the cathode oxygen distribution in the holetype flow field is more uniform, resulting in the oxygen transfer process having less resistance.

3.3 Constant current discharge

A constant current discharge test was conducted at a current density of 80 mA cm $^{-2}$, and the test results are as in Fig. 10. The experimental temperature was 70 °C, the volume of methanol solution was 2 ml, and the concentration was 1 mol L^{-1} .

At the discharge process, the losses at the same current density are different due to the different total impedance magnitudes of the µDMFC, resulting in different durations.

The HFSS-µDMFC has the highest discharge voltage and the longest discharge duration, and the GFT-µDMFC has the lowest discharge voltage and the lowest discharge duration. The total impedance of the GFT-µDMFC during the discharge process is highest while the duration and voltage are lowest. It is mainly because the GFT-µDMFC has a higher loss due to impedance. The total impedance of HFSS-µDMFC is the least, the loss due to impedance is lower, the longest discharge time and the highest voltage during the discharge process. The voltage and duration of the HFT-µDMFC during the discharge process are better than those of the GFSS-µDMFC. The total impedance and CTR of the HFT-µDMFC and the GFSS-µDMFC are the same, and the difference in transfer impedance is higher slightly. During the discharge process, the transfer loss of the GFSS-µDMFC is higher due to the influence of the cathode on the oxygen mass transfer efficiency of the current collector. Thus, the voltage and duration of the HFT-µDMFC during the discharge process are better than those of the GFSS-µDMFC.

Conclusions 4

In this work, the effects of different types of cathode current collectors on the performance of the µDMFC were investigated. The impact of various materials and flow field structures on the gas-liquid two-phase flow of the µDMFC cathodes were analyzed by performing polarization characteristics, EIS, and constant current discharge test on the µDMFC.

For both materials, the FSS has a contact angle of 96.525°, which is hydrophobic, and the FT has a contact angle of 70.751°, which is hydrophilic. The maximum power density of the HFSS- μ DMFC is 49.53 mW cm⁻², which is 70.15% higher than that of the HFT- μ DMFC. The maximum power density of the GFSS- μ DMFC is 22.60 mW cm⁻², which is 11.99% higher than that of the GFT- μ DMFC. The hydrophobic FSS prevents water generated by the cathode from adhering to the foam metal. This reduces the concentration polarization and mass transfer impedance and enhances the oxygen transfer rate.

For both flow fields, the maximum power density of the HFSS- μ DMFC is 119.16% higher than that of the GFSS- μ DMFC. The maximum power density of the HFT- μ DMFC is 44.25% higher than that of the GFT- μ DMFC. The hole-type flow field has a more uniform structural distribution and oxygen distribution. This improves the effective current collection efficiency of the cathode collector. During the discharge process, the HFSS- μ DMFC has the highest discharge voltage and the longest discharge duration. By reducing the total impedance of the μ DMFC, energy losses can be reduced, thus improving the μ DMFC performance.

Author contributions

Conceptualization; Zhengang Zhao and Fan Zhang; methodology, Zhengang Zhao and Fan Zhang; software, Fan Zhang and Yanhui Zhang; validation, Fan Zhang and Yanhui Zhang; formal analysis, Dacheng Zhang; investigation, Fan Zhang and Yanhui Zhang; resources, Zhengang Zhao and Dacheng Zhang; data curation, Fan Zhang; writing – original draft preparation, Fan Zhang and Yanhui Zhang; writing – review and editing, Zhengang Zhao and Dacheng Zhang; visualization, Chuan Luo; supervision, Zhengang Zhao and Dacheng Zhang; project administration, Zhengang Zhao and Dacheng Zhang; funding acquisition, Zhengang Zhao and Dacheng Zhang.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work is partly supported by National Natural Science Foundation of China (Grant No. 62162035, 62103174) and Applied Basic Foundation of Yunnan Province (Grant No. 202001AU070046, 202101AT070131).

Notes and references

- 1 B. M. Sudaroli and A. K. Kolar, Energy, 2016, 98, 204-214.
- 2 G. Ramasamy and S. Somasundaram, *Processes*, 2020, **8**, 796. 3 T. Noor, S. Pervaiz, N. Iqbal, H. Nasir and E. Pervaiz,
- Nanomaterials, 2020, **10**, 1601.
- 4 C. E. Damian-Ascencio, A. Saldaa-Robles, A. Hernandez-Guerrero and S. Cano-Andrade, *Energy*, 2017, **133**, 306–316.
- 5 L. Rostami, P. Nejad and A. Vatani, *Energy*, 2016, 97, 400-410.
- 6 M. Herranen, U. Wiklund, J. O. Carlsson and S. Hogmark, *Surf. Coat. Technol.*, 1998, **99**, 191–196.
- 7 R. C. Makkus, A. Janssen, F. Bruijn and R. Mallant, *J. Power Sources*, 2000, **86**, 274–282.

- 8 D. Zhang, L. Duan, G. Lu and W. H. Tuan, *Int. J. Hydrogen* Energy, 2010, 35, 3721–3726.
- 9 L. Yang, X. Zhang, W. Yuan, Y. Zhang and X. Liu, *Energy*, 2018, 157, 599–607.
- 10 H. Choi, O. H. Kim, M. Kim, H. Choe, Y. H. Cho and Y. E. Sung, ACS Appl. Mater. Interfaces, 2014, 6, 7665–7671.
- 11 A. Kumar and R. G. Reddy, J. Power Sources, 2003, 114, 54-62.
- 12 J. E. Park, J. Lim, S. Kim, I. Choi, C. Y. Ahn, W. Hwang, M. S. Lim, Y. H. Cho and Y. E. Sung, *Electrochim. Acta*, 2018, S001346861830238X.
- 13 K. S. Dong, H. Y. Jin, G. K. Dong and S. K. Min, *Renewable Energy*, 2017, **115**, 663–675.
- 14 X. Sun, X. Xie, S. Wu, Z. Liu and K. Jiao, *Int. J. Green Energy*, 2021, **18**, 1–12.
- 15 D. Spernjak, A. K. Prasad and S. G. Advani, *J. Power Sources*, 2010, **195**, 3553–3568.
- 16 B. R. Friess and M. Hoorfar, *Int. J. Hydrogen Energy*, 2012, **37**, 7719–7729.
- 17 N. Guo, C. L. Ming and U. O. Koylu, *Int. J. Hydrogen Energy*, 2013, **38**, 6750–6761.
- 18 T. Wilberforce, Z. E. Hassan, E. Ogungbemi, O. Ijaodola, F. N. Khatib, A. Durrant, J. Thompson, A. Baroutaji and A. G. Olabi, *Renewable Sustainable Energy Rev.*, 2019, **111**, 236–260.
- 19 X. Li and I. Sabir, Int. J. Hydrogen Energy, 2005, 30, 359-371.
- 20 S. A. Atyabi, E. Afshari, S. Wongwises, W. M. Yan and M. S. Shadloo, *Energy*, 2019, **179**, 490–501.
- 21 Z. Yuan, W. Fu, Y. Zhao, Z. Li, Y. Zhang and X. Liu, *Energy*, 2013, **55**, 1152–1158.
- 22 Z. Zhao, F. Zhang, Y. Zhang and D. Zhang, *Energies*, 2021, 14, 6608.
- 23 A. M. Oladoye, J. G. Carton and A. G. Olabi, *J. Mater.*, 2014, 2014, 1–10.
- 24 S. S. Hsieh, C. C. Ho and L. C. Hung, *Renewable Energy*, 2016, **90**, 28–37.
- 25 I. Sharma and M. Ghangrekar, *Water Sci. Technol.*, 2018, 77, 999–1006.
- 26 L. Li, W. Fan, J. Xuan, M. K. H. Leung, K. Zheng and Y. She, *Energy Procedia*, 2017, **105**, 1557–1563.
- 27 N. V. Demeneva, D. V. Matveev, V. V. Kharton and S. I. Bredikhin, *Russ. J. Electrochem.*, 2016, **52**, 678–684.
- 28 Y. Wei, T. Yong, Q. Wang and Z. Wan, *Appl. Energy*, 2011, **88**, 1671–1680.
- 29 T. Yong, Y. Wei, M. Pan, B. Tang, Z. Li and Z. Wan, *J. Power Sources*, 2010, **195**, 5628–5636.
- 30 Y. Wei, T. Yong, Z. Wan and M. Pan, *Int. J. Hydrogen Energy*, 2011, **36**, 2237–2249.
- 31 C. C. Roberts, R. R. Rao, M. Loewenberg, C. F. Brooks, P. Galambos, A. M. Grillet and M. B. Nemer, *Lab Chip*, 2012, 12, 1540–1547.
- 32 R. Xue, Y. Zhang and X. Liu, Energy, 2017, 139, 535-541.
- 33 X. Chao, A. Faghri, X. Li and T. Ward, *Int. J. Hydrogen Energy*, 2010, **35**, 1769–1777.
- 34 A. Aricò, V. Baglio, A. Stassi and V. Antonucci, *Adv. Crop Sci. Technol.*, 2010, **72**, 271–276.
- 35 Y. Zhang, X. Rui, X. Zhang, J. Song and X. Liu, *Energy*, 2015, **91**, 1081–1086.

- 36 T. S. Zhao, C. Xu, R. Chen and W. W. Yang, *Prog. Energy Combust.*, 2009, 35, 275–292.
- 37 Z. Yuan and J. Yang, J. Power Sources, 2015, 285, 318-324.
- 38 A. Frignani, C. Monticelli, M. Tassinari and G. Trabanelli, AEIT Conference-from Research to Industry: the Need for A More Effective Technology Transfer, 2002.
- 39 D. Chun, D. Kim, Z. R. Williamson, T. Lee and C. W. Squibb, *Appl. Therm. Eng.*, 2013, **50**, 293–301.
- 40 T. Chasen, Y. Q. Liang, X. Xie, L. I. Lincai, Z. Liu, D. U. Qing and K. Jiao, *Sci. China: Technol. Sci.*, 2021, **64**, 13.
- 41 I. Alaefour, S. Shahgaldi, A. Ozden, X. Li and F. Hamdullahpur, *Fuel*, 2018, **230**, 98–103.
- 42 M. Ashrafi, H. Kanani and M. Shams, *Energy*, 2018, **147**, 317–328.
- 43 B. H. Lim, E. H. Majlan, W. Daud, M. I. Rosli and T. Husaini, *Int. J. Hydrogen Energy*, 2016, **42**, 9210–9218.