Facilitating the acidic oxygen reduction of Fe–N–C catalysts by fluorine-doping†‡

Xiafang Tao, §ab Ruihu Lu, §b Lingmei Ni, ‡d Vladislav Gridin, ‡d Samir H. Al-Hilfi, b Zijie Qiu, b Yan Zhao, c Ulrike I. Kramm, ‡db Yazhou Zhou ab and Klaus Müllen ab

As the alternatives to expensive Pt-based materials for the oxygen reduction reaction (ORR), iron/nitrogen co-doped carbon catalysts (FeNC) with dense FeN₄ active sites are promising candidates to promote the commercialization of proton exchange membrane fuel cells. Herein, we report a synthetic approach using perfluorotetradecanoic acid (PFTA)-modified metal–organic frameworks as precursors for the synthesis of fluorine-doped FeNC (F-FeNC) with improved ORR performance. The utilization of PFTA surfactants causes profound changes of the catalyst structure including desorption of intermediates at FeN₄ active sites and increases the BET surface area (up to 1085 m² g⁻¹), as well as dense FeN₄ sites. The F-FeNC catalyst exhibits an improved ORR activity with a high E₁/₂ of 0.83 V (vs. RHE) compared to the pristine FeNC material (E₁/₂ = 0.80 V). A fast decay occurs in the first 10 000 potential cycles for the F-FeNC catalyst, but high durability is still maintained up to another 50 000 cycles. Density functional theory calculations reveal that the strongly withdrawing fluorine atoms doped on the graphitic carbon can optimize the electronic structure of the FeN₄ active center and decrease the adsorption energy of ORR intermediates.

New concepts

Iron and nitrogen co-doped carbon catalysts (FeNC) play an important role as potential alternatives to expensive Pt-based catalysts in proton exchange membrane fuel cells. The relatively strong adsorption of oxygen reduction reaction (ORR)-related species on FeN₄ sites largely limits the ORR kinetics. The electron-withdrawing heteroatoms in FeNC catalysts is regarded as a promising strategy to improve the kinetic activity. This effect should be even more pronounced in the case of fluorine (F) as F possesses the highest electronegativity. To date, studies about F-modified FeNC catalysts (F-FeNC) of high ORR activity have been little reported. Here, we developed a new synthetic approach by pyrolysis of perfluorotetradecanoic acid (PFTA)-modified metal–organic frameworks for improving the ORR performance of F-FeNC in acidic solution. The utilization of PFTA was proven to be beneficial to achieve the increased content of micropores, high Brunauer–Emmett–Teller surface area (1083 m² g⁻¹), dense FeN₄ active sites, as well as fluorine-doping (F-doping). Density functional theory calculations revealed that the strongly electron-withdrawing F atoms doped into carbon can optimize the electronic structure of the FeN₄ active center and decrease the adsorption energy of ORR intermediates.

Introduction

Hydrogen-based sustainable energy technologies, particularly proton exchange membrane fuel cells (PEMFCs), hold promise for achieving carbon neutrality and reducing the threats of climate change.¹⁻⁴ While the development of PEMFCs has been significantly advanced over the past decades, their commercialization is still hindered by the need for platinum-based catalysts, which are expensive and scarce.⁵⁻⁷ Therefore, searching for non-noble metals to catalyze the oxygen reduction reaction (ORR) in PEMFCs is highly rewarding.⁸⁻¹⁰

Transition metal and nitrogen co-doped carbon materials (M–N–C, M = Fe, Co, Mn, etc.) have been developed as catalysts for ORR, in which atomically dispersed MN₄ moieties on carbon supports are generally considered as the main active sites to directly absorb O₂ and catalyze the subsequent ORR kinetics.¹¹⁻¹⁷ In particular, the Fe–N–C catalysts demonstrate an encouraging ORR performance in acidic solution and are promising alternatives to Pt-based catalysts in PEMFCs.¹⁸⁻²¹

Previous studies have revealed that heteroatom doping (e.g. N, O, S, and P) into Fe–N–C catalysts or binding with Fe atoms can determine the ORR performance by balancing the adsorption and desorption of intermediates at FeN₄ sites.²⁰,²²⁻²⁵ Recently, in
Mun’s work, the ORR activity of the FeN₄ catalytic center could be tailored by introducing S-functionalities into carbon black.²⁶ Li et al. demonstrated a higher activity of the N-FeN₄ than for the FeN₄ moiety.²⁷ Han et al. reported that due to its electronegativity, chlorine atoms vertically coordinated with the FeN₄ center that can facilitate electron transfer from Fe atom to oxygen intermediates during the ORR.²⁴ This effect should be even more pronounced in the case of fluorine (F) as F possesses the highest electronegativity.²⁸,²⁹ However, F-modified Fe–N–C catalysts are rarely studied in the literature.³⁰ Surface modification of carbon black supports by grafting with trifluoromethylphenyl groups is reported to improve the stability, however, lower the activity of Fe–N–C catalysts.³¹ This is caused by the increased hydrophobicity of carbon black, which keeps the oxidants from reaching the triple-phase boundary.³¹ The fluorination of Fe–N–C samples with fluorine gas has led to the poisoning of the Fe active sites and dramatic ORR degradation.³² Attempts at F-doping into carbon supports without poisoning active sites can therefore affect ORR performance.

In this work, we explore an approach to disperse the FeNₓ active sites on N, F co-doped graphitic carbon materials (F-FeNC) for improved ORR performance. This F-FeNC catalyst was prepared by pyrolysis of perfluorotetradecanoic acid (PFTA)-modified Fe-doped zeolitic imidazolate frameworks (PFTA/Fe-ZIFs), wherein PFTA served as fluorine source. The PFTA is proven to be beneficial to maintain the microporous structure of ZIFs, achieving the high Brunauer–Emmett–Teller (BET) surface area (1085 m² g⁻¹). The surface and bulk characterizations reveal that the F-FeNC catalyst possesses densely embedded single-atom Fe sites and fluorine-doping into graphitic carbon. As a result, the prepared catalysts deliver a superior ORR activity with a half-wave potential of 0.83 V versus the reversible hydrogen electrode (vs. RHE) in acidic solution as compared to the pristine FeNC material.

Results and discussion

Synthesis and characterization of F-FeNC catalysts

As illustrated in Fig. 1a, the F-FeNC catalysts were prepared by high-temperature pyrolysis of PFTA/Fe-ZIFs. Details of the method are described in the (ESI†). In brief, a methanol solution containing zinc nitrate hexahydrate, ferric nitrate nonahydrate, 2-dimethylimidazole, and PFTA surfactant in a molar ratio of 75 : 1 : 190 : 2 was hydrothermally treated at 60 °C overnight to synthesize the PFTA/Fe-ZIFs. The PFTA surfactant was in situ introduced into the frameworks of PFTA/Fe-ZIFs through the coordination between carboxyl groups of PFTA and the metal atom. The presence of carboxyl groups and C–F bonds in PFTA/Fe-ZIFs precursors is proven by X-ray photoelectron spectroscopy (XPS) (Fig. S1, ESI†). Notably, Fe-ZIFs and PFTA/Fe-ZIFs showed similar diffraction patterns (Fig. S2a, ESI†), uniform morphological features (Fig. S3, ESI†), and similar pore size distributions (Fig. S4, ESI†), suggesting that
the introduction of a limited amount of PFTA does not change the crystalline structure of ZIFs. The PFTA/Fe-ZIFs precursors were then thermally annealed in an argon atmosphere at 1000 °C to obtain a black powder. During high-temperature pyrolysis, the hydrocarbons of PFTA/Fe-ZIFs were carbonized into partially graphitic carbon, as evidenced by powder X-ray diffraction (PXRD) patterns (Fig. S2b, ESI‡) and Raman spectra (Fig. 2a). The inclusion of nitrogen dopants into the graphitic carbon provided a large number of anchoring sites for bonding Fe atoms. Samples prepared by adding different amounts of PFTA surfactants were denoted as F-FeNC-1, F-FeNC-2, and F-FeNC-3 based on the molar ratio of PFTA to 2-dimethylimidazole (169 : 1, 85 : 1, and 68 : 1, respectively) (see Method, ESI‡). For comparison, the F–NC sample without Fe and the FeNC sample without fluorine were synthesized in an analogous fashion.

The morphologies of the new catalysts were characterized by transmission electron microscopy (TEM) and high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) revealing carbon nanoparticles with 100 nm size in F-FeNC-2 (Fig. 1b and c). No obvious Fe-based particles with crystalline phases were observed in the high-resolution STEM (HR-STEM) image, which was also supported by selected area electron diffraction (SAED, Fig. S5f, ESI‡) analysis. Therefore, Fe species were highly dispersed as single atoms. Distinguishable signals for N, F, and Fe were detected in corresponding elemental mapping images and electron energy loss spectroscopy (EELS) across the carbon nanoparticles, indicating uniform distributions of these elements in the F-FeNC-2 sample (Fig. 1d–g). The bright spots in the aberration-corrected HAADF-STEM image corresponded to single heavy atoms. This gave evidence that the Fe atoms seem mainly atomically dispersed within the graphitic carbons (Fig. 1h). Similar morphologies were observed for the reference materials (Fig. S5 and S6, ESI‡).

The effect of PFTA surfactants on the characteristics of graphitic carbon in the catalysts was analyzed using Raman spectroscopy. Two prominent peaks located at around 1345 and 1585 cm⁻¹ were attributed to the D band and the G band, respectively (Fig. 2a). The ratio of the intensity of D and G bands (I_D/I_G) and the crystallite size of the carbon plane (L_a) were correlated with the amount of PFTA surfactant in FeNC, F-FeNC-1, F-FeNC-2, and F-FeNC-3 (Fig. 2b and Table S4, ESI‡). The I_D/I_G values, which are commonly utilized to determine the density of defects increased significantly with higher amounts of PFTA surfactants, indicating that more defects were created in the F-FeNC catalysts by F-doping. In the meanwhile, the increased density of defects induced the formation of smaller carbon planes, as evidenced by the decreased L_a value.

The porosity of the catalysts was assessed by nitrogen-sorption analysis and a non-local density functional theory (NLDFT) model was used to determine the pore size distributions. Type IV isotherms with a steep increase in V_ads at relatively low and high N₂ pressures indicated the preferred existence of micropores and large outer surfaces in all the samples (Fig. 2c). The BET surface area of the F-FeNC-2 catalysts was 1085 m² g⁻¹, higher than that of the FeNC catalyst (809 m² g⁻¹) (Table S5, ESI‡). The larger surface area originated from the increased micropore surface area (S_{micropore}) of 665 m² g⁻¹ compared with that of FeNC (362 m² g⁻¹), which was further confirmed by the pore size distributions in which ultramicropore and micropore were centered at 0.6 and 1.5 nm, respectively (Fig. 2d). When using higher contents of PFTA,
the $S_{\text{Micropore}}$ of F-FeNC-3 decreased (561 m$^2$ g$^{-1}$), which was attributed to the carbonization of excessive PFTA which may block the microporosity. Notably, all the F-FeNC samples showed the high $S_{\text{Micropore}}$, suggesting that the addition of PFTA is beneficial for the formation of micropores.

The surface-near elemental composition of FeNC and F-FeNC were obtained from XPS. The high-resolution C 1s XPS spectra of all samples displayed similar C 1s peaks (Fig. S7a, ESI†). The weak C–F peaks located at 685.5–688.0 eV in the high-resolution F 1s XPS spectra (Fig. S7b, ESI†) suggested that fluorine atoms were doped into graphitic carbon.35,36 The percentage of Fe$^{3+}$ species was increased from 27% for the FeNC sample to 48% for F-FeNC-3 based on fitting results of Fe 2p$^{3/2}$ spectra (Table S1, ESI†). The higher oxidation state of the Fe site was ascribed to the surrounding fluorine heteroatoms with high electronegativity. The high-resolution N 1s spectra in Fig. S8 (ESI†) were deconvoluted into five types of N species with binding energies of 402.1, 401.2, 399.4, and 398.4 eV which could be assigned to oxidized-, graphitic-, pyrrolic-N, FeN$_x$, and pyridinic-N, respectively.34,39 All the F-FeNC catalysts displayed a higher percentage of FeN$_x$ (11–16%) than that in the FeNC (~9.2%). This finding suggested the formation of more FeN$_x$ active sites by PFTA surfactant addition (Table S2, ESI†). The fluorine contents measured by XPS were 0.1, 0.1, and 0.2 at% for F-FeNC-1, F-FeNC-2, and F-FeNC-3, respectively (Table S3, ESI†). The Fe content was determined to be 1.68 wt% for F-FeNC-2 by inductively coupled plasma atomic emission spectroscopy (ICP-AES), higher than that of the FeNC reference material (1.49 wt%) and other F-FeNC catalysts.

To gain deeper insight into the nature of Fe-related species, $^{57}$Fe Mössbauer spectroscopy was employed for both F-FeNC-2 and FeNC catalysts. The Mössbauer spectrum of FeNC was fitted with two major doublets of D1 and D2 that correspond to FeN$_4$ species (Fig. S9a and Table S6, ESI†), whereas three doublets of D1, D2, and D3 were observed for the F-FeNC-2 sample (3a). Based on the fact that similar D3 sites were already found in other fluorine-free FeNC catalysts14,15, it may conclude that a fluorine coordination to iron can be excluded. This is in agreement with the previous conclusion that F atoms in F-FeNC were present as F-doping into the graphitic carbon. In addition to the doublets, the FeNC sample exhibited a sextet and singlet, whereas no sextet was found in the F-FeNC-2 sample (3b), suggesting a possible formation of more Fe active sites in F-FeNC.

Further information on the binding situation of Fe atoms was obtained from X-ray absorption spectroscopy (XAS). The Fe $K$-edge X-ray absorption near-edge structure (XANES) displayed a shift towards higher energy from FeNC to F-FeNC-2, confirming the increased oxidation state of Fe due to the F-doping in the surrounding carbon (Fig. 3c),41,42 which is consistent with the XPS analysis. More detailed information on the mode of coordination of the Fe site was obtained from the Fe $K$-edge extended X-ray absorption fine structure (EXAFS) analysis. A prominent peak was observed at ~1.48 Å in the radial distribution function obtained for the Fourier transformed (FT) EXAFS for the F-FeNC (Fig. 3d), which can be assigned to the Fe–N (O) coordination. The signal of Fe–Fe bonds with typical bond distances of around 2 Å were very weak in both
FeNC and F-FeNC catalysts pointing toward more abundant single-atom Fe sites. Two backscattering paths of Fe–N and Fe–O were performed for fitting EXAFS spectra (Fig. 3e and f). For the Fe–N path, the coordination number of Fe atom was calculated to be 5.4 ± 0.4 with a Fe–N distance of 2.00 ± 0.02 Å (Table S7, ESI†). The EXAFS fitting for the F-FeNC could also be obtained by a mixture of Fe–N and Fe–O paths, with coordination numbers of 3.9 ± 0.9 and 1.1 ± 0.7, respectively. Based on this fitting, we propose two structures for the Fe center, i.e. an in-plane FeN₄ with either an axial nitrogen ligand (FeN₄–N, Fig. 3e inset) or an axial oxygen-containing ligand (e.g. as FeN₄–O), respectively. In contrast, the FeNC samples possess mainly FeN₄ sites (Table S7, ESI†).

ORR performance

The electrocatalytic ORR performance of F-FeNC and control FeNC samples was, first, investigated in an O₂-saturated 0.5 M H₂SO₄ electrolyte using a rotating disk electrode (RDE) technique. A commercial Pt/C catalyst (20 wt% Pt, Fuelcell store) as a reference was evaluated in 0.1 M HClO₄ solution. All potentials were calibrated against a hydrogen electrode (RHE). ORR polarization curves clearly revealed the effect of fluorine incorporation (Fig. 4a). The FeNC sample demonstrated an ORR onset potential (Eₒ) of 0.93 V (Eₒ: the potential at current density of 0.1 mA cm⁻²) and a half-wave potential (E½) of 0.80 V (vs. RHE). By using PFTA, the ORR activity of F-FeNC-1 was enhanced. When increasing the use of PFTA, the F-FeNC-2 achieved the best ORR activity with a high E½ of 0.83 V which was only 20 mV lower than that of the Pt/C reference catalyst (E½, 0.85 V). At 0.85 V, the F-FeNC-2 material gave a three times higher kinetic current density (Jk) than that of the bare FeNC sample (1.63 vs. 0.55 mA cm⁻², Fig. 4b). Moreover, the Tafel slopes of all samples fell into the range from 66 to 70 mV dec⁻¹ (Fig. S10 and Table S8, ESI†). Further increasing the PFTA caused a decline in the activity of F-FeNC-3. Due to a high affinity of SCN⁻ anion to Fe ion, the SCN⁻ can poison FeN₄ active sites and lead to a significant suppression of the ORR activity.²⁶,⁴³ The poisoning test using an electrolyte containing SCN⁻ ions and the comparison with the ORR polarization curve of the Fe-free F–NC revealed that the ORR activity was mainly stemming from the Fe₇ sites (Fig. 4f).²⁶,⁴³ Compared to reference samples, the F-FeNC-2 exhibited a higher density of FeN₄ active sites because of the larger surface area of micro-pores and the higher contents of FeN₄, achieving the enhanced ORR activity.⁴²

To examine the ORR mechanism of the catalysts, the rotating ring-disk electrode (RRDE) technique was applied for monitoring the formation of H₂O₂ during the tests. As shown in Fig. 4c, the H₂O₂ yield for F-FeNC-2 was below 1.7% in the potential range from 0.2 to 0.7 V, which was lower than that of the FeNC sample. The low H₂O₂ yield and the calculated electron-transfer number of 3.98 were firm evidence of a four-electron pathway for oxygen reduction.³³ The stabilities of F-FeNC and FeNC catalysts were evaluated using accelerated stress tests (ASTs) by cycling the potential from 0.6 to 1.0 V (vs. RHE) in an O₂-saturated 0.5 M H₂SO₄ electrolyte (Fig. 4d). The changes of E½ and H₂O₂ yield

Fig. 4  (a) Steady-state ORR polarization plots recorded with a rotation speed of 900 r.p.m. and a scan rate of 5 mV s⁻¹. (b) Jk at 0.85 V and E½ and (c) calculated H₂O₂ yield for FeNC, F-FeNC-1, F-FeNC-2, F-FeNC-3, and commercial Pt/C catalysts, (d) ORR polarization plots of the F-FeNC-2 and (e) the changes of E½ and H₂O₂ yield for FeNC and F-FeNC catalysts during the durability tests. (f) ORR polarization curves of F–NC and F-FeNC and results of poisoning test with SCN⁻ of F-FeNC.
were calculated after every 10,000 cycles (Fig. 4e). After the first 10,000 cycles, both catalysts showed a steep activity degradation with $E_{1/2}$ losses for FeNC and F-FeNC of 35 and 27 mV due to the removal of unstable sites in acid solution. After continuing the potential cycling for another 50,000 cycles, a significant activity decline with another 73 mV loss in $E_{1/2}$ and increased H$_2$O$_2$ product were observed for FeNC (Fig. 4e and Fig. S11, ESI) due to the carbon corrosion and demetalation of FeN$_x$ via Fenton [like] reactions between the Fe centers and the intermediate product of H$_2$O$_2$. In contrast, the F-FeNC-2 presented improved durability with a small loss of only 13 mV in $E_{1/2}$. The involved F-doing on graphitic carbon can enhance the durability of Fe–N–C catalyst by the mitigation of the oxidative corrosion of the carbon support, which is one of the major decay mechanisms of FeNC catalysts. The F-FeNC-2 catalyst was further studied as the cathode in a PEMFC in an H$_2$-air system (1 bar back pressure for H$_2$ and air), and compared with commercial Pt/C catalyst. The F-FeNC-2 catalyst demonstrated a peak power density of 122 mW cm$^{-2}$ (Fig. S12a, ESI). A stability test at a constant voltage of 0.5 V with F-FeNC-2 catalyst as cathodic test is shown in Fig. S12b (ESI), exhibiting much poorer stability performance than in the RRDE test. The reasons for this difference are still a matter of debate but observed for several Fe–N–C catalysts.

**Theoretical study of F-FeNC on ORR**

To understand the effect of F-doping, we investigated the adsorption behavior and corresponding free-energy evolution on the Fe active sites by first-principles calculations (see Computational method, ESI). The appropriate structure, denoted as FeN$_4$-aX-F$_x$, was constructed based on EXAFS fitting and XPS analyses and consisted of a porphyrin-like FeN$_4$ with an axial ligand (X represents C-, N-, and O-containing axial ligands), and fluorine atoms doped in graphene layers ($x = 2, 4$, and 8) (Table S9, ESI). The control structures of FeN$_4$, FeN$_4$-F, and FeN$_4$-aX without F-doping, were considered for comparison. The optimized ORR intermediates ($^*$OH, $^*$OOH, and $^*$O) adsorbed Fe sites were also included. The Gibbs free energy profiles for the ORR process on F-FeNC catalysts were investigated under zero potential ($U = 0$ V), highest potential ($U =$ biggest output potential), and equilibrium potential ($U = 1.23$ V) (Fig. S13–S16, ESI). The calculated free energy changes suggest that the reduction of $^*$OH is the rate-determining step (RDS) for the ORR process on most of FeN$_4$-aX-F$_x$ catalysts, whereas the RDS of FeN$_4$-aO-F$_x$ and FeN$_4$-aO-F$_8$ catalysts is the formation of $^*$OOH. The adsorption energies of $^*$OH, $^*$OOH, and $^*$O display linear correlations (Fig. 5a). This scaling relations lead to a volcano relationship between the ORR catalytic activity and the adsorption energy of $^*$OH ($\Delta G_{^*OH}$) (Fig. 5b). The principle for enhancement of intrinsic ORR activity is to balance the $^*$OH adsorption on the Fe active site that is neither too strong nor too weak. The FeN$_4$ exhibits a high $\Delta G_{^*OH}$ of 0.50 eV and an overpotential of 1.73 V. The activity of FeN$_4$ site is mostly limited by the $^*$OH reduction due to a strong $^*$OH adsorption. The FeN$_4$-F$_x$ and FeN$_4$-aX-F$_x$ structures located at the right side of FeN$_4$ point toward a reduced $\Delta G_{^*OH}$ on the FeN$_4$, certifying that electron-withdrawing F-doping weakens the intermediates adsorption on active sites. The FeN$_4$-aO-F$_2$ with a low content of F-doping reaches the top of the volcano plot and is thus identified as the most active ORR catalyst.

Due to the low contents of fluorine in the catalyst, it can be questioned if the enhanced performance of Fe-FeNC catalysts is only related to the fluorine doping. Indeed, it might not be only governed by the fluorine content but maybe cooperatively by one or all of the following three aspects. Firstly, PFTA plays an important role in maintaining the microporous structure of ZIFs, achieving a higher surface area of micropores and BET surface area. More micropores improve the accessibility of FeN sites for ORR. Secondly, the F-FeNC-2 sample possesses a high density of active sites due to the high loading of single Fe atoms. Thirdly, the fluorine atoms doped into graphitic carbon can balance the adsorption of oxygen intermediate, facilitating the intrinsic activity on FeN sites. Overall, different from the fluorinated FeNC catalysts in literature, our experimental and theoretical results indicate that the use of PFTA-modified ZIFs as precursors can influence both the Fe active sites and carbon structure of the F-FeNC catalyst, thus leading to very good ORR performance in acidic media.

**Conclusions**

In summary, a new synthetic approach using PFTA-modified ZIFs has been developed to prepare microporous Fe–N–C catalysts. Compared with the FeNC without fluorine incorporation, the prepared F-FeNC catalyst exhibited strongly enhanced ORR performance in terms of the onset and have-wave potentials, as well as durability. This enhancement is associated with the larger surface area of micropores (665 vs. 362 m$^2$ g$^{-1}$) and BET surface area (1085 vs. 809 m$^2$ g$^{-1}$), abundant single Fe atoms, and strong electron-withdrawing F-doping compared to F-free FeNC. We believe that the findings demonstrated here could be extended to fabricate other F-modified M–N–C catalysts with high performance for the hydrogen/oxygen evolution reaction, electrochemical carbon dioxide reduction reaction, nitrogen fixation reaction, and other (electro-) catalytic applications.
Author contributions

Y. Z and K. M conceived and designed the experiments. X. T and Y. Z conducted the synthesis, characterization and RDE measurements of catalysts. U. I. K and L. N recorded $^{57}$Fe Mössbauer transmission spectra and did the data analysis. U. I. K and V. G performed the PEM fuel cell investigations. R. L and Y. Z carried out computational studies. X. T, Z. Q, U. I. K, Y. Z and K. M co-wrote the paper. The research was supervised by U. I. K, Y. Z and K. M.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was financially supported by the Max Planck Society and the Natural Science Foundation of China (51702129 and 51972150). Y. Z acknowledges the support from the China Postdoctoral Science Foundation (2018M630527) and the China Scholarship Council for financial support (201708320150). X. T thanks the scholarship from Jiangsu University. U. I. K, L. N and V. G thank the German federal ministry of education and research (BMBF) for financial support via the NanoMatFutur young research group FeNC–StRedO (03XP0092). We thank Umicore for providing the P/C catalyst for FC testing. Open Access funding provided by the Max Planck Society.

References

1 I. Staffell, D. Scammann, A. Velazquez Abad, P. Balcombe, P. E. Dodds, P. Ekins, N. Shah and K. R. Ward, Energy Environ. Sci., 2019, 12, 463–491.
2 M. K. Debe, Nature, 2012, 486, 43–51.
3 X. X. Wang, M. T. Swihart and G. Wu, Nat. Catal., 2019, 2, 578–589.
4 G. Karanfil, Int. J. Energy Res., 2020, 44, 4–25.
5 L. L. Chen, X. L. Xu, W. X. Yang and J. B. Jia, Chin. Chem. Lett., 2020, 31, 626–634.
6 H. A. Gasteiger, S. S. Kocha, B. Sompalli and F. T. Wagner, Appl. Catal., B, 2005, 56, 9–35.
7 Y. Z. Zhou, C. H. Yen, Y. H. Hu, C. M. Wang, X. N. Cheng, C. M. Wai, J. Yang and Y. H. Lin, J. Mater. Chem. A, 2016, 4, 18628–18638.
8 A. A. Gewirth, J. A. Varnell and A. M. DiAscro, Chem. Rev., 2018, 118, 2313–2339.
9 B. P. Setzler, Z. B. Zhuang, J. A. Wittkopf and Y. S. Yan, Nat. Nanotechnol., 2016, 11, 1020–1025.
10 X. Wan, X. F. Liu, Y. C. Li, R. H. Yu, L. R. Zheng, W. S. Yan, H. Wang, M. Xu and J. L. Shui, Nat. Catal., 2019, 2, 259–268.
11 M. Chen, Y. He, J. S. Spendelow and G. Wu, ACS Energy Lett., 2019, 4, 1619–1633.
12 H. T. Chung, D. A. Cullen, D. Higgins, B. T. Sneed, E. F. Holby, K. L. More and P. Zelenay, Science, 2017, 357, 479–483.
13 A. Zito, V. Goellner, V. Armel, M. T. Sougrati, T. Mineva, L. Stieveno, E. Fonda and F. Jauuen, Nat. Mater., 2015, 14, 937.
14 U. I. Kramm, M. Lefevre, P. Bogdanoff, D. Schmeisser and J. P. Dodelet, J. Phys. Chem. Lett., 2014, 5, 3750–3756.
15 U. I. Kramm, M. Lefevre, N. Larouche, D. Schmeisser and J. P. Dodelet, J. Am. Chem. Soc., 2014, 136, 978–985.
16 U. I. Kramm, L. M. Ni and S. Wagner, Adv. Mater., 2019, 31, 1805623.
17 L. Ni, C. Gallenkamp, S. Paul, M. Kübler, P. Theis, S. Chabbra, K. Hofmann, E. Bill, A. Schnegg, B. Albert, V. Krewald and U. I. Kramm, Adv. Energy Sustainab. Res., 2021, 2, 2000064.
18 S. T. Thompson and D. Papageorgopoulos, Nat. Catal., 2019, 2, 558–561.
19 H. Zhang, H. T. Chung, D. A. Cullen, S. Wagner, U. I. Kramm, K. L. More, P. Zelenay and G. Wu, Energy Environ. Sci., 2019, 12, 2548–2558.
20 Y. Zhou, X. Tao, G. Chen, R. Lu, D. Wang, M. X. Chen, E. Jin, J. Yang, H. W. Liang, Y. Zhao, X. Feng, A. Narita and K. Mullen, Nat. Commun., 2020, 11, 5892.
21 J. Li, M. T. Sougrati, A. Zito, J. M. Ablett, I. C. Oğuz, T. Mineva, I. Matanovic, P. Atanassov, Y. Huang, I. Zenyuk, A. Di Cicco, K. Kumar, L. Dubau, F. Maillard, G. Dražič and F. Jauuen, Nat. Catal., 2020, 4, 10–19.
22 X. H. Yan, X. L. Li, C. H. Fu, C. Lin, H. M. Hu, S. Y. Shen, G. H. Wei and J. L. Zhang, Prog. Nat. Sci.: Mater. Int., 2020, 30, 896–904.
23 K. Hu, L. Tao, D. D. Liu, J. Huo and S. Y. Wang, ACS Appl. Mater. Interfaces, 2016, 8, 19379–19385.
24 Y. H. Han, Y. G. Wang, R. R. Xu, W. X. Chen, L. R. Zheng, A. J. Han, Y. Q. Zhu, J. Zhang, H. B. Zhang, J. Luo, C. Chen, Q. Peng, D. S. Wang and Y. D. Li, Energy Environ. Sci., 2018, 11, 2348–2352.
25 K. J. Chen, K. Liu, P. D. An, H. J. W. Li, Y. Y. Lin, J. H. Hu, C. K. Jia, J. W. Fu, H. M. Li, H. Liu, Z. Lin, W. Z. Li, J. H. Li, Y. R. Lu, T. S. Chan, N. Zhang and M. Liu, Nat. Commun., 2020, 11, 4173.
26 Y. Mun, S. Lee, K. Kim, S. Kim, S. Lee, J. W. Han and J. Lee, J. Am. Chem. Soc., 2019, 141, 6254–6262.
27 X. N. Li, C. S. Cao, S. F. Hung, Y. R. Lu, W. Z. Cai, A. I. Rykov, S. Miao, S. B. Xi, H. B. Yang, Z. H. Hu, J. H. Wang, J. Y. Zhao, E. E. Alp, W. Xu, T. S. Chan, H. M. Chen, Q. H. Xiong, H. Xiao, Y. Q. Huang, J. Li, T. Zhang and B. Liu, Chem, 2020, 6, 3440–3454.
28 X. J. Sun, Y. W. Zhang, P. Song, J. Pan, L. Zhuang, W. L. Xu and W. Xing, ACS Catal., 2013, 3, 1726–1729.
29 S. G. Peera, A. Arunchander and A. K. Sahu, Nanoscale, 2016, 8, 14650–14664.
30 Y. Chang, J. X. Chen, J. C. Jia, X. Hu, H. J. Yang, M. L. Jia and Z. H. Wen, Appl. Catal., B, 2021, 284.
31 Y. C. Wang, P. F. Zhu, H. Yang, L. Huang, Q. H. Wu, M. Rauf, J. Y. Zhang, J. Dong, K. Wang, Z. Y. Zhou and S. G. Sun, ChemElectroChem, 2018, 5, 1914–1921.
32 G. X. Zhang, X. H. Yang, M. Dubois, M. Herrera, R. Chermitz, M. Lefevre, M. Cherif, F. Vidal, V. P. Gilbin, S. H. Sun and J. P. Dodelet, Energy Environ. Sci., 2019, 12, 3015–3037.
33 J. Z. Li, M. J. Chen, D. A. Cullen, S. Hwang, M. Y. Wang, B. Y. Li, K. X. Liu, S. Karakalos, M. Lucero, H. G. Zhang, C. Lei, H. Xu, G. E. Sterbinsky, Z. X. Feng, D. Su, K. L. More, G. F. Wang, Z. B. Wang and G. Wu, *Nat. Catal.*, 2018, 1, 935–945.

34 H. G. Zhang, S. Hwang, M. Y. Wang, Z. X. Feng, S. Karakalos, L. L. Luo, Z. Qiao, X. H. Xie, C. M. Wang, D. Su, Y. Y. Shao and G. Wu, *J. Am. Chem. Soc.*, 2017, 139, 14143–14149.

35 Y. Z. Zhou, C. H. Yen, S. F. Fu, G. H. Yang, C. Z. Zhu, D. Du, P. C. Wo, X. N. Cheng, J. Yang, C. M. Wai and Y. H. Lin, *Green Chem.*, 2015, 17, 3552–3560.

36 Y. Z. Zhou, J. Yang, C. Z. Zhu, D. Du, X. N. Cheng, C. H. Yen, C. M. Wai and Y. H. Lin, *ACS Appl. Mater. Interfaces*, 2016, 8, 25863–25874.

37 P. Z. Chen, T. P. Zhou, S. B. Wang, N. Zhang, Y. Tong, H. X. Ju, W. S. Chu, C. Z. Wu and Y. Xie, *Angew. Chem., Int. Ed.*, 2018, 57, 15471–15475.

38 G. Panomsuwan, N. Saito and T. Ishizaki, *J. Mater. Chem. A*, 2015, 3, 9972–9981.

39 Y. Li, Y. Z. Zhou, H. J. Wen, J. Yang, C. Maouche, Q. Q. Liu, Y. Y. Wu, C. Cheng, J. Zhu and X. N. Cheng, *Dalton Trans.*, 2018, 47, 14992–15001.

40 S. Wagner, H. Auerbach, C. E. Tait, I. Martinaiou, S. C. N. Kumar, C. Kubel, I. Sergeev, H. C. Wille, J. Behrends, J. A. Wolny, V. Schunemann and U. I. Kramm, *Adv. Mater.*, 2020, 32, 1906764.

41 Q. H. Li, W. X. Chen, H. Xiao, Y. Gong, Z. Li, L. R. Zheng, X. S. Zheng, W. S. Yan, W. C. Cheong, R. A. Shen, N. H. Fu, L. Gu, Z. B. Zhuang, C. Chen, D. S. Wang, Q. Peng, J. Li and Y. D. Li, *Adv. Mater.*, 2018, 30, 1800588.

42 Q. Wang, Y. Q. Yang, F. F. Sun, G. B. Chen, J. Wang, L. S. Peng, W. T. Chen, L. Shang, J. Q. Zhao, D. Sun-Waterhouse, T. R. Zhang and G. I. N. Waterhouse, *Adv. Energy Mater.*, 2021, 11.

43 Q. Wang, Z. Y. Zhou, Y. J. Lai, Y. You, J. G. Liu, X. L. Wu, E. Terefe, C. Chen, L. Song, M. Rauf, N. Tian and S. G. Sun, *J. Am. Chem. Soc.*, 2014, 136, 10882–10885.

44 G. Bae, M. W. Chung, S. G. Ji, F. Jaouen and C. H. Choi, *ACS Catal.*, 2020, 10, 8485–8495.

45 C. H. Choi, H. K. Lim, M. W. Chung, G. Chon, N. R. Sahraie, A. Altin, M. T. Sougrati, L. Stievano, H. S. Oh, E. S. Park, F. Luo, P. Strasser, G. Drazic, K. J. J. Mayrhofer, H. Kim and F. Jaouen, *Energy Environ. Sci.*, 2018, 11, 3176–3182.

46 D. Banham, S. Ye, K. Pei, J. Ozaki, T. Kishimoto and Y. Imashiro, *J. Power Sources*, 2015, 285, 334–348.

47 J. K. Norskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. R. Kitchin, T. Bligaard and H. Jonsson, *J. Chem. B*, 2004, 108, 17886–17892.