Carbon dioxide electroreduction provides a useful source of carbon monoxide, but comparatively few catalysts could be sustained at current densities of industry level. Herein, we construct a high-yield, flexible and self-supported single-atom nickel-decorated porous carbon membrane catalyst. This membrane possesses interconnected nanofibers and hierarchical pores, affording abundant effective nickel single atoms that participate in carbon dioxide reduction. Moreover, the excellent mechanical strength and well-distributed nickel atoms of this membrane combines gas-diffusion and catalyst layers into one architecture. This integrated membrane could be directly used as a gas diffusion electrode to establish an extremely stable three-phase interface for high-performance carbon dioxide electroreduction, producing carbon monoxide with a 308.4 mA cm$^{-2}$ partial current density and 88% Faradaic efficiency for up to 120 h. We hope this work will provide guidance for the design and application of carbon dioxide electro-catalysts at the potential industrial scale.
The excessive emission of greenhouse gas CO₂ results in a chain of serious climate problems. As a consequence, it is intensely urgent to convert CO₂ into beneficial fuels or chemicals, and simultaneously alleviate atmospheric CO₂ level. Especially, electrochemical reduction of CO₂ has tremendous potential to contribute value-added compounds directly from atmospheric CO₂, coupling with renewable-energy electricity. This research area has evolved remarkably in the past few decades, making significant progress towards industrial application. CO₂ is highly stable molecule, which needs proper catalysts to activate. Therefore, catalyst development has played a crucial part with a steady stream of novel electro-catalysts, i.e., metals, metal oxides, metal complexes, transition-metal phosphides or carbon-based materials. Some catalysts could even exhibit nearly 100% selectivity for specific products (i.e., single-atom Co sites). On the other hand, CO₂ electoreduction takes place around the three-phase interface, hence the diffusion and adsorption of CO₂ molecules contribute to the reaction rate to a large extent. A majority of catalytic materials for CO₂ electoreduction are still developed, characterized and evaluated in typical H-type electrochemical cells. In these H-type cells, the catalyst layer is fully immersed and CO₂ gas is generally bubbled into the conductive electrolyte. Accordingly, the current densities of CO₂ conversion would be largely limited by the low solubility and slow diffusion of CO₂ in aqueous systems, and hardly satisfy the industrial-relevant current densities (>100 mA cm⁻²) in CO₂ reduction. To solve these problems, electrochemical reactors like flow cell) using gas-diffusion electrode have been seriously investigated in recent years. Instead of aqueous electrolyte, CO₂ is supplied from the gaseous phase through a gas-diffusion layer to construct a gas–liquid interface adjoined to the solid catalyst layer. This unique architecture has the potential to deliver enough CO₂ directly onto the catalyst surface and reach very high current densities. To produce a usable gas-diffusion electrode, catalysts for CO₂ electoreduction were usually adhered or deposited on a specific substrate. Unfortunately, because of the weak contacts between substrate and electro-catalyst, the electroactive species become detached and easily fall off, thus reducing the long-term performance of CO₂ electoreduction.

Herein, we report an effective method for the preparation of a high-yield and self-supported Ni single-atom/porous carbon fiber membrane catalyst (NiSA/PCFM) via electrospinning method for high-efficiency CO₂ electoreduction. The extensive single-atom Ni sites distributing throughout carbon nanofiber could play a decisive role in activation of CO₂. The porous and interconnected carbon nanofibers of NiSA/PCFM provide substantial channels for CO₂ diffusion and electron transport. In addition, NiSA/PCFM has favorable mechanical strength and flexibility, which can be easily tailored to specific shapes or thickness. Therefore, NiSA/PCFM could combine gas-diffusion and catalyst layers into a single architecture. This integrated NiSA/PCFM membrane could be immediately utilized as GDE compartment for high-performance CO₂ reduction, avoiding the weak connection between the substrate and electro-catalyst.

**Results**

**Synthesis and characterization of NiSA/PCFM catalysts.** The fabrication method of NiSA/PCFM membrane is described in Fig. 1a, which involves electrospinning process and heated treatment. The primary fibers were firstly electrospun from the mixture solution of ZIF-8 nanoparticles (Supplementary Fig. 1), Ni(NO₃)₂·6H₂O and polyacrylonitrile (PAN). Notably, ZIF-8 nanoparticles did not contain Ni ions and mainly acted as pore former in our experiments. Ni ions were introduced by Ni(NO₃)₂·6H₂O and would be reduced by carbonized organic polymers of PAN in the following heated treatment, leaving atomically dispersed Ni atoms anchored on nitrogen-doped porous carbon. It also deserves to be mentioned that the NiSA/PCFM membranes yielded in our laboratory apparatus are at least ~280 cm⁻² in size (Fig. 1b). The manufacturing method is uncomplicated and easily adapted to larger size (or thickness) membranes. The NiSA/PCFM membrane is robust and flexible shown in Fig. 1b, which could retain the initial structure during bending stress (Supplementary Fig. 2). As disclosed by the stress–strain curve in Fig. 1c, NiSA/PCFM membrane also can endure ~1.3 MPa tensile strength. Besides, the water contact angle (CA) of NiSA/PCFM catalyst was very close 0°, indicating the hydrophilic property (Supplementary Fig. 3). If a little amount of NaF mixture was spread on the surface of NiSA/PCFM, water CA was measured to be 142.7° ± 3.0° (Fig. 2a), implying the excellent hydrophobic property. This hydrophobic property could guarantee that NiSA/PCFM is not easily flooded in aqueous solution, which is crucial for GDE devices.

NiSA/PCFM was also characterized with morphological and gas adsorption experiments. The field-emission scanning electron microscopy (FE-SEM, Fig. 2b, c, Supplementary Fig. 4a, b) and high-resolution transmission electron microscopy (HRTEM, Fig. 2d) showed that the nanofibers of NiSA/PCFM created a three-dimensionally interconnected architecture with visible and well-distributed macropores (ca. 100 nm) throughout these nanofibers. N₂ sorption isotherms (Fig. 3a) furtherly verifies that NiSA/PCFM has a 714 m² g⁻¹ specific surface area and hierarchical structure. The pore size distribution peaks at ca. 100 nm and ca. 1 nm (~3.6 times of the dynamic diameter of CO₂ molecules) reveal both macropores and micropores. The macropore channels could enhance the gas transportation to micropores where CO₂ molecules are mainly captured, causing the remarkable CO₂ adsorption capacity of NiSA/PCFM under atmospheric pressure (Fig. 3b). This exceptional structure could accelerate the diffusion and adsorption of CO₂, resulting in CO₂ enrichment around the active sites. If no ZIF-8 nanoparticles but only Ni(NO₃)₂ were added into the precursor solution for electrospinning, as-synthesized NiSA/CFM presents typical microporous structure and no macropores were observed on the nanofibers (Supplementary Fig. 5, 6a–b), which sharply reduced the CO₂ adsorption capacity (Fig. 3b) even with a higher specific surface area (789 m² g⁻¹, Supplementary Table 1) than that of NiSA/PCFM. PCFM has the similar hierarchical structure (Supplementary Figs. 6c, d, 7) and CO₂ adsorption capacity as NiSA/PCFM.

In addition, based on the orientation of the distorted graphene layers in HR-TEM images (Supplementary Fig. 4d), a longer range orientation of the graphene unit is seen in NiSA/PCFM. This might affect the conductivity and redox reactions, which involve electron transfer through the electrode surface. Moreover, the Ni, N, and C elements were homogenously distributed on the NiSA/PCFM, water contact angle (CA) of NiSA/PCFM catalyst was very close 0°, indicating the hydrophilic property (Supplementary Fig. 3). If a little amount of NaF mixture was spread on the surface of NiSA/PCFM, water CA was measured to be 142.7° ± 3.0° (Fig. 2a), implying the excellent hydrophobic property. This hydrophobic property could guarantee that NiSA/PCFM is not easily flooded in aqueous solution, which is crucial for GDE devices.

As presented in Fig. 1f and Supplementary Fig. 8, the white spots distributed on the black background of NiSA/PCFM which can be indexed to the (002) and (100) planes of carbon. The diffraction peaks belonging to metallic Ni or Ni oxides are absent in all three...
patterns (Supplementary Fig. 9). The relatively sharp and intensive peaks at 26.2° and 44.0° for NiSA/PCFM, compared with that of NiSA/CFM, indicate that the additional ZIF-8 nanoparticles result in a higher degree of crystallization and graphitization. Furthermore, X-ray photoelectron spectroscopy (XPS, Supplementary Table 2) N 1s spectrum of NiSA/PCFM (Fig. 3c) could deconvolute into pyridine N (398.2 eV), Ni–N (399.4 eV), pyrrole (400.5 eV), graphitic (401.3 eV) and oxidized (403.0 eV) N species. The binding energies of Ni 2p3/2 in NiSA/PCFM (Supplementary Fig. 10) were reflected by the peak at 854.8 eV, locating between metallic Ni0 (853.5 eV) and Ni2+ (855.8 eV), suggesting that the Ni atoms in NiSA/PCFM is likely to be in a low-valent state21. The coordination environment of Ni species would be intensively investigated via X-ray absorption fine spectroscopy (XAFS). Figure 3d exhibits the X-ray absorption near-edge structure (XANES) spectrum in the Ni K-edge of NiSA/PCFM, using Ni foil and NiO as the references. The near-edge spectra of NiSA/PCFM locates between Ni foil and NiO, implying that the valence state of those isolated Ni atoms are between metallic (Ni0) and oxidized (Ni2+) status22,23. The Fourier transform (FT) k3-weighted χ(k) function of the extended X-ray absorption fine structure (EXAFS) spectra of NiSA/PCFM exhibited a dominant peak at 1.42 Å for Ni–N coordination (Fig. 3e). As contrast, Ni foil shows a typical Ni-Ni pair at 2.20 Å, and the Ni–O interaction of NiO sits around 1.62 Å. According to the fitting results, the proposed local structure of NiSA/PCFM involves coordination by four N atoms (Ni–N4, Fig. 3f, Supplementary Table 3)24–27. NiSA/CFM also possess Ni–N4 structure, indicated by XPS (Supplementary Fig. 11) as well as XAFS spectra (Supplementary Fig. 12). No Ni element was observed in XPS patterns of PCFM (Supplementary Fig. 13). The Ni amount of NiSA/PCFM and NiSA/CFM were 1.3 wt% and 1.2 wt%, revealed by inductively coupled plasma optical emission spectroscopy.

CO2 electro-reduction with H-type cell. On the basis of above characterizations, as-prepared NiSA/PCFM composites possess excellent physical properties and massive single-atom doping, which may facilitate enhanced activity for CO2 electroreduction. The CO2 electro-reduction activity, NiSA/PCFM, NiSA/CFM, and PCFM powders were firstly drop-cast onto a carbon paper using typical powdering and binding method for LSV tests and electrolysis in H-type cell. These powdered samples were named as P-NiSA/PCFM, P-NiSA/CFM and P-PCFM, easy to distinguish. The anode and cathode sections were separated by a
Nafion-117 proton exchange membrane. The cathodic compartment was continuously bubbled with CO2 and injected directly into gas chromatography (GC) for quantitative detection of gaseous products. Liquid-phase products were determined by 1H NMR spectroscopy. The primary products catalyzed by three catalysts are only CO and H2, and no trace of any liquid products were observed in 1H NMR spectroscopy (Supplementary Fig. 16). Clearly, the P-NiSA/PCFM and P-NiSA/CFM maintain excellent faradic efficiency (FE) for CO generation at working potentials from −0.5 to −1.0 V RHE, and P-NiSA/PCFM could achieve a maximum value of 95% at −0.7 V RHE (Fig. 4a, Supplementary Fig. 15a, b). In contrast, P-PCFM catalyst without Ni composition showed very limited FEs for CO production throughout the potential range (Supplementary Fig. 15c), indicating that the Ni–N4 structure played a dominating role in CO2 electroreduction to CO. As mentioned above, NiSA/PCFM membrane has excellent mechanical strength and could directly act as working electrode in H-type cell system. As displayed in Fig. 4c, the best CO Faradaic efficiency obtained by NiSA/PCFM is 96% at −0.7 V RHE, similar to that of P-NiSA/PCFM (95%). However, NiSA/PCFM brought forth a 56.1 mA cm−2 CO partial current density at −1.0 V RHE (Fig. 4d), significantly higher than those of P-NiSA/PCFM, P-NiSA/CFM and P-PCFM (Fig. 4b).

To understand the observed activity difference of three catalysts, density functional theory (DFT) calculations were performed on the well-defined N–C and Ni–N4−C structures (Supplementary Fig. 16, Table 4). The corresponding free energy profiles (ΔG) are summarized in Fig. 5a. According to the simulations, the conversion of CO2 to adsorbed COOH∗ intermediate on N–C and Ni–N4−C structures are uphill in the free energy profiles, demonstrating that the first electronic step is the rate-determining step for both systems28. Especially, the Ni–N4−C has more favorable ΔG (0.7 eV) than N–C (1.3 eV) for this step. The lower ΔG requires less energy to form *COOH intermediate, consistent with the much smaller Tafel slope of NiSA/PCFM (117 mV/dec) and NiSA/CFM (156 mV/dec), than that of PCFM (204 mV/dec, Fig. 5b)29,30. Notably, the partial CO current density catalyzed by P-NiSA/PCFM and NiSA/PCFM are higher than those of PCFM and NiSA/CFM (Fig. 4b). NiSA/PCFM also presented a much larger overall current density in the linear sweep voltammetry (LSV) tests (Supplementary Fig. 17). Since NiSA/PCFM and NiSA/CFM possess very close amount of Ni single atoms, the differences in CO2 electroreduction activity may also be ascribed to the hierarchically porous structures10,31,32. Different from the typical microporous structure of NiSA/CFM, NiSA/PCFM shows hierarchically porous structures, which can accelerate the diffusion and adsorption of reactants, lead to remarkable CO2 adsorption capacity (Fig. 3b) and a higher electrochemical active surface areas (ECSA, Supplementary Fig. 18). In a word, the free-standing and
CO₂ electro-reduction with GDE cell. All the above electrolysis were conducted in a conventional H-type cell, whereby the catalyst layer is completely soaked in a conductive CO₂-saturated electrolyte. The current densities were severely restricted by mass transportation as a result of the limited solubility of CO₂ in aqueous solution. Therefore, we deployed a GDE device (see Supplementary Fig. 20 and Fig. 5c) and tested the NiSA/PCFM catalyst to advance toward industrial current densities (>100 mA cm⁻²). Firstly, NiSA/PCFM, NiSA/CFM, and PCFM were loaded onto a gas diffusion layer (SIGRACET) via typical methods as the cathode compartment (denoted as P-NiSA/PCFM) in flow cell for LSV tests. As presented in Supplementary Fig. 17d, e, the current densities of three samples observed in flow cell are all far greater than those in H-type cell, indicating the superiority flow cell device.

Figure 4c, d displays the CO FE and partial current densities for three catalysts at various cathode potentials in H-type cell. e CO faradaic efficiencies, d partial current densities of NiSA/PCFM at various cathode potentials in different cells. e Long-term stability tests in GDE cell and f H-type cell at -1.0 V_RHE, respectively.

CO₂ electro-reduction with GDE cell. All the above electrolysis were conducted in a conventional H-type cell, thereby the catalyst layer is completely soaked in a conductive CO₂-saturated electrolyte. The current densities were severely restricted by mass transportation as a result of the limited solubility of CO₂ in aqueous solution. Therefore, we deployed a flow cell equipped with GDE device (see Supplementary Fig. 20 and Fig. 5c) and tested the NiSA/PCFM catalyst to advance toward industrial current densities (>100 mA cm⁻²). Firstly, NiSA/PCFM, NiSA/CFM, and PCFM were loaded onto a gas diffusion layer (SIGRACET) via typical methods as the cathode compartment (denoted as P-NiSA/PCFM) in flow cell for LSV tests. As presented in Supplementary Fig. 17d, e, the current densities of three samples observed in flow cell are all far greater than those in H-type cell, indicating the superiority flow cell device.

Figure 4c, d displays the CO FE and partial current densities plotted against the RHE-corrected cathode potential. H₂ and CO accounts for up to 99% of the transferred charge on NiSA/PCFM membrane using GDE cell, while no liquid product could be detected (Supplementary Fig. 15d), similar to those in H-type cell. In GDE device, It is also apparent that the NiSA/PCFM can be performed exceeding its optimized range of cathode potentials in the above H-type cell measurements. The CO FE of NiSA/PCFM begins to reduce obviously at more negative potentials than −0.7 V in H-type cell, but more than 80% FEs could still be obtained even at −1.2 V_RHE in GDE cell (Fig. 4c). Besides, partial CO current densities with GDE device were significantly higher. For example, with the same 88% CO FEs, NiSA/PCFM exhibits 308.4 mA cm⁻² CO partial current density in GDE cell at −1.0 V_RHE, which is more than 5 times larger than that of H-type cell (56.1 mA cm⁻²). In addition, the partial CO current density obtained in GDE cell could reach an even higher value of 336.5 mA cm⁻² with 83% CO FE at −1.2 V_RHE. To the best of our knowledge, accounting for the electrochemical conversion of CO₂ to CO, the catalytic activity of NiSA/PCFM is comparable to these state-of-the-art electro-catalysts in previously reports (Supplementary Table 5), especially at industrial-relevant current densities.

To explore the stability of the NiSA/PCFM membrane, long-term tests were conducted using NiSA/PCFM catalysts at −1.0 V_RHE electrode potential where the best CO FE was acquired. CO FEs and partial current densities against time are plotted in Fig. 4e. Both CO FEs and partial current densities of the NiSA/PCFM membrane showed only a negligible drop during 120 h tests, retaining more than 95% of initial value. More than
100 h stability of NiSA/PCFM membrane could also be achieved in two-electrode system (Supplementary Fig. 21)\textsuperscript{34,35}. Moreover, the CO FE\textsubscript{s} obtained with P-NiSA/PCFM was very close to those of NiSA/PCFM using cathode potentials from $-0.3$ V\textsubscript{RHE} to $-1.2$ V\textsubscript{RHE}, attaining an optimal value of 87\% at $-1.0$ V\textsubscript{RHE} (Supplementary Fig. 22a). Furthermore, P-NiSA/PCFM could preserve a lower but steady current density of 70.5 mA cm\textsuperscript{-2} at $-0.6$ V\textsubscript{RHE} cathode potential for 2 h. However, current densities dropped significantly at more negative $-0.8$ or $-1.0$ V\textsubscript{RHE} cathode potential within the short time of 1 h (Supplementary Fig. 22b, d), which is noticeably worse than the long-term stability of NiSA/PCFM. In a typical GDE device, the catalysts (e.g., P-NiSA/PCFM) were usually spray-coated (or deposited) onto a gas diffusion layer using polymer binders (GDL, Fig. 5c). During electrolysis procedure, massive gas products would pour out from the active sites along with the large current densities at relatively negative cathode potentials. Consequently, the electro-active species would easily fall off and reduce the long-term performance of CO\textsubscript{2} reduction, due to the weak contacts between catalyst and substrate\textsuperscript{16}. If P-NiSA/PCFM was tested in H-type cell, the current-time curve also shows a distinct drop after 10 h electrolysis (Fig. 4f). As mention early, NiSA/PCFM has abundant Ni single atoms and cross-linked carbon nanofibers, which combines both gas-diffusion and catalyst layers. Hence, NiSA/PCFM membrane was readily used as cathode compartment (Fig. 5d, e), instead of spray-coating the catalyst ink onto a specific gas-diffusion layer. The integrated structure and fine mechanical strength of NiSA/PCFM membrane could guarantee the long-time stability under high current densities.

Besides, the hydrophobicity of gas-diffusion and catalyst layer also have vital impact on the long-term stability\textsuperscript{18}. The thickness of catalyst layer is always in nanoscale and this thin catalyst layer could easily be flooded by electrolyte and destroy the three-phase interface around the active sites\textsuperscript{4}. In our system, the thickness of NiSA/PCFM membrane is in the range of hundreds of micrometers (Fig. 2b), which is not easily to be totally soaked by electrolyte and establish an extremely stable three-phase interface for high-performance CO\textsubscript{2} electroreduction. Furthermore, the architecture of NiSA/PCFM membrane still remained unchanged after electrolysis (Supplementary Fig. 23, 24) and could be readily utilized for next run with simple oven dry.

**Discussion**

In conclusion, we prepared an effective approach for the relatively large-scale production of single-atom Ni doped porous carbon...
membrane. We started from the screening tests in typical H-type cell, which could produce CO with maximum 96% faradaic efficiency at \(-0.7\) V_{RHE}. DFT calculations based on *COOH intermediate provide a molecular understanding of the observed high efficiency on single-atom Ni sites. In addition, the outstanding structure of NiSA/PCFM could generate substantial effective Ni single atoms actually participating in CO2 reduction. Moving to GDE device, this NiSA/PCFM membrane can combine gas-diffusion as well as catalyst layer to create a reactive and provide stable three-phase interface for high-performance CO2 electroreduction. Remarkably, NiSA/PCFM membrane could manifest CO product with a commercially relevant partial current density of 308.4 mA cm\(^{-2}\) as well as a high Faradaic efficiency of 88\% at \(-1.0\) V_{RHE} for at least 120 h, which is even comparable to those state-of-the-art electro-catalysts for CO2 reduction to CO. This work may also give some reference and guidance for the design and preparation of integrated electro-catalysts utilized in gas-diffusion device.

Methods

Chemicals and Characterizations. All reagents were used as received. LSV\(^{7}\) tests were performed using an electrochemical workstation (Princeton Applied Research 263 A) with a three-electrode system in typical divided electrochemical cell separated by Nafion\(^{\circledast}117\) (Dupont) membrane between cathode and anode. Gas phase products were analyzed by Gas Chromatography (SRI 8610 C). Microstructure images were taken using field emission-scanning electron microscope (FEI JEO-L78000F). High-resolution TEM images as well as element mapping were obtained using transmission electron microscope (JEM-2100F). Aberration-corrected HAADF-STEM characterization was carried on a JEM JEM-AR2000F TEM/STEM. X-ray diffraction spectra were taken by a Rigaku MiniFlex 600 powder diffractometer. \(N_{2}\) adsorption-desorption was performed using a Micromeritics ASAP 2460 instrument. CO2 adsorption was carried out using Quantachrome Autosorb-IQ2-MP. X-ray photoelectron spectra were taken with ThermoVG Scientific ESCALAB 250 X-ray photoelectron spectrometer. The X-ray absorption fine structure spectra, including XANES and EXAFS at Ni K-edge, were collected at 1W1B station in Beijing Synchrotron Radiation Facility.

Synthesis of ZIF-8. ZIF-8 crystals were prepared via rapidly pouring an aqueous solution (16 mL deionized water) of Zn(NO\(_3\))\(_2\)-6H\(_2\)O (7.9 mmol, 2.34 g) into an aqueous solution (160 mL deionized water) of 2-methylimidazole (553 mmol, 45.4 g). Then, the mixed solution magnetically stirred for 5 min at room temperature. After stirring, the mixture was filtered by filter paper. The liquid product was collected by centrifuge (7000 revolutions per minute, 30 min) and thoroughly cleaned by deionized water for at least five times. The product was dried at 60 °C for 12 h in a vacuum drying oven.

Synthesis of NiSA/PCFM, NiSA/CFM, and PCFM. NiSA/PCFM were synthesized using an electropinning process. First, 1.5 g of polyacrylonitrile (PAN, Mw = 130,000), 1.5 g of ZIF-8 and 30 mg of Ni(NO\(_3\))\(_2\)-6H\(_2\)O powder were dissolved in 15 mL of N,N-dimethylformamide (DMF) with ultrasonic stirring to get a homogenous mixture. The transparent mixture was diverted into a plastic syringe with a stainless needle at the tip. After electropinning process, the as-spun raw fibers were firstly preoxidized in air at 250 °C for 1 h and immediately carbonized under argon gas at an optimum temperature of 900 °C for 2 h. The heating rate was 5 °C min\(^{-1}\). Then, the catalyst membrane was cooled to room temperature under argon gas flow. Then, the resultant materials were immersed into H\(_2\)SO\(_4\) solution (3.0 M) for 10 h to remove the remaining Zn species, Ni or NiO nanoparticles. NiSA/PCFM membrane was thus obtained.

The transparent solution containing 30 mg Ni(NO\(_3\))\(_2\)-6H\(_2\)O and no ZIF-8 was also electropun and carbonized under 900 °C to act as control for the comparison. The resulting product was referred to as NiSA/CFM.

CO2 reduction procedure. H-type cell. Electrolysis and LSV\(^{7}\) tests were performed in a in conventional H-type electrochemical cell separated by Nafion\(^{\circledast}117\) membrane between cathode and anode, consisted of Pt foil as the counter electrode and an Ag/AgCl as reference electrode in CO2 or N\(_2\) saturated 0.5 M KHCO\(_3\) solution as electrolyte, which were separated by a piece of Nafion\(^{\circledast}117\) membrane. Electrolytes were cycled at 50 mL min\(^{-1}\). The CO2 gas was supplied at rate of 20 mL min\(^{-1}\) to the cathode and was flown through the NiSA/PCFM membrane.

Data availability

The data that support the findings of this study are available within the paper and its Supplementary Information file or are available from the corresponding authors upon request.

Received: 27 August 2019; Accepted: 17 December 2019; Published online: 30 January 2020

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Acknowledgements
We sincerely appreciate the financial support from Shenzhen Government’s Plan of Science and Technology (JCYJ20170817095641221 and JCYJ20170818091657056), the Natural Science Foundation of Guangdong Province (2017A040405066), National Natural Science Foundation of China (51902209, 21975162). We also acknowledge the help of Electron Microscopy Centre of Shenzhen University for testing the aberration-corrected HADDF-STEM.

Author contributions
C.H. conceived the project and idea. H.Y. designed the experiments and wrote the manuscript. Q.L., C.Z., X.Y., Z.C., G.L. and Q.H. carried out the synthesis, material characterizations and electrochemical measurements. X.R., Q.Z. and J.L. analysed the data from these experiments and commented on the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41467-020-14402-0.

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Peer review information Nature Communications thanks the anonymous reviewers for their contribution to the peer review of this work. Peer reviewer reports are available.

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