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Efficient electrosynthesis of hydrogen peroxide in neutral media using boron and nitrogen doped carbon catalysts†

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The two-electron electrocatalytic oxygen reduction reaction ($2e^-$ ORR) is a promising, sustainable, and efficient route for on-site and small-scale production of hydrogen peroxide (H_2O_2) compared to the conventional anthraquinone oxidation process. Although heteroatom doped carbon materials are widely studied for the $2e^-$ ORR, few studies have been conducted on the synergistic relationship between multiple doping elements on H_2O_2 performance. Moreover, there is a lack of efficient and stable catalysts for H_2O_2 electrosynthesis in neutral media. In this work, a series of boron and nitrogen co-doped carbon (BNC) catalysts are synthesized and assessed for the $2e^-$ ORR in neutral pH electrolyte. Through X-ray photoelectron and absorption spectroscopy analyses, we characterize B–N–C moieties and correlate them with H_2O_2 production performance. In rotating ring disk electrode measurements, BNC catalysts pyrolyzed at 900 °C exhibit H_2O_2 Faradaic efficiency of ~70% and demonstrate excellent long-term stability over 10 hours of continuous operation. In a three-electrode flow cell, this catalyst achieves ~78% H_2O_2 Faradaic efficiency and $0.9\text{ mmol cm}^{-2}\text{ h}^{-1}$ productivity at 0.20 V vs. RHE, surpassing most previously reported carbon-based catalysts in neutral media. This work contributes fundamental insights into the role of boron and nitrogen heteroatom dopants toward $2e^-$ ORR and highlights the practical viability of these catalysts for real world applications.

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1. Introduction

Hydrogen peroxide (H_2O_2) serves as an eco-friendly oxidant employed across diverse industries, including chemical synthesis, paper and pulp, semi-conductor, environmental remediation, and others.^{1–3} The global demand for H_2O_2 is rapidly growing, especially after the recent COVID-19 pandemic where it was used as a disinfectant in the healthcare industries.^{4,5} Current industrial production of H_2O_2 is achieved via the multi-step anthraquinone oxidation process, which is not only energy-intensive, but also environmentally hazardous, producing significant amounts of organic by-product waste.⁶ In addition, this process relies on a centralized infrastructure and requires expensive and hazardous transport and storage of concentrated H_2O_2 solutions.⁷ Thus, the development of a low-cost and decentralized H_2O_2 production method is highly desirable to reduce the costs and safety concerns associated with H_2O_2 synthesis, storage, and transportation.⁸

At the forefront of a potential sustainable and efficient route to the on-site and small-scale production of H_2O_2 is the two-

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electron electrocatalytic oxygen reduction reaction ($2e^-$ ORR).^{9,10} At present, state-of-the-art $2e^-$ ORR catalysts are based on noble metals and their alloys (e.g., Pd, Au, and Pt–Hg).^{8,11–14} However, such catalysts have considerable drawbacks, namely, scarcity and high cost, which impede their implementation on a large-scale. Recently, carbon-based materials emerged as promising alternatives for the $2e^-$ ORR due to their relative abundance, facile preparation, low cost, and excellent stability.^{15–18} More importantly, their tunable surface and structural properties make it possible to modify the electrochemical performance of these carbon-based electrocatalysts.¹⁹ Among various modification methods, nitrogen doping has been reported to generate efficient active sites with favorable electrochemical properties.^{20,21} However, many of these studies with high selectivity toward $2e^-$ ORR are performed in alkaline conditions, while much higher ORR overpotentials are reported in acidic or neutral pH conditions.^{20,22} Production of H_2O_2 in proton exchange membrane electrolyzers is preferable over hydroxide conducting membranes, primarily due to their superior energy efficiency, improved stability against H_2O_2 , and greater scalability.^{23–25} Thus, it is desirable to develop carbon-based catalysts with improved H_2O_2 selectivity and activity in neutral or weakly acidic pH conditions.^{12,26,27}

The introduction of boron dopants into nitrogen doped carbon frameworks is a broadly reported strategy to enhance the ORR activity vs. pure nitrogen doped carbons.^{28–36} Boron dopants have been hypothesized to serve as active sites or to impact the electronic properties of surrounding carbon atom active sites upon incorporation into the carbon matrix.²⁹ Several studies have investigated boron and nitrogen co-doped carbon catalysts (BNC), employing a variety of synthetic approaches, including chemical vapor deposition and thermal annealing, in attempts to induce the most active and selective defect-site structures.^{37,38} However, despite significant progress in the development of BNC catalysts, many of these studies focused on the $4e^-$ ORR pathway to H_2O in alkaline media.^{36,39–41} There are very few reports on high-performance toward the $2e^-$ ORR pathway to H_2O_2 , especially in neutral pH media.⁴² Moreover, detailed information of the nature of the active B–N–C moieties and understanding their role in the origin of ORR activity and selectivity is lacking.

In the present work, we synthesize a series of BNC catalysts and study their activity and selectivity toward $2e^-$ ORR to H_2O_2 in neutral pH solutions. We report a facile preparation method for the BNC catalysts and tune the content and nature of B and N dopants *via* varied pyrolysis temperatures. Spectroscopic characterizations confirm the presence of diverse B–N, B–N–C, and N–C dopant structures in the carbon matrix. The catalyst performance toward H_2O_2 activity and selectivity are investigated in three different reactor geometries, using a rotating ring disk electrode (RRDE), a gas diffusion electrode (GDE) flow cell, and a dual membrane electrode assembly (MEA) solid electrolyte (SE) flow cell. Benefiting from the high efficiency of active sites and abundant interfacial B–N–C moieties, BNC pyrolyzed at 900 °C (900-BNC) demonstrates the best performance in each of these reactor geometries. The 900-BNC catalyst reaches H_2O_2 selectivity and Faradaic efficiency up to 80% and 70%,

respectively, in an RRDE geometry, and achieves 78% H_2O_2 Faradaic efficiency at a production rate of $0.9 \text{ mmol cm}^{-2} \text{ h}^{-1}$ in a GDE flow cell. While similar production rates have been achieved in strongly alkaline conditions, this production rate is ~ 1.75 times greater compared to previously reported non-metal carbon-based electrocatalysts in similar reactors in neutral media.⁴³ This work provides collective insights into the development of heteroatom-doped carbon catalyst materials for the electrosynthesis of H_2O_2 and enables the design of improved materials toward other applications, such as electrochemical capacitors⁴⁴ and alkali metal batteries.^{45,46}

2. Experimental section

2.1. Materials and chemicals

Zinc nitrate hexahydrate, 2-methylimidazole, potassium bicarbonate, phenylboronic acid, potassium sulfate, hydrogen peroxide (30 wt%), hydrochloric acid, isopropyl alcohol, and cobalt sulfate heptahydrate were purchased from Sigma Aldrich. All electrolytes and deionized (DI) water were prepared from Millipore-quality water ($18.2 \text{ M}\Omega \text{ cm}$ at 25 °C). All gases used (O_2 and N_2) in this work had a purity of $>99.995\%$ (Airgas).

2.2. Synthesis of catalysts

The zeolitic imidazolate framework 8 (ZIF-8) precursor was prepared according to previous literature.^{47,48} Briefly, 1.47 g of $Zn(NO_3)_2 \cdot 6H_2O$ was dissolved in 100 mL methanol; a second solution consisting of 3.24 g of 2-methylimidazole in 100 mL methanol was prepared in parallel. The Zn^{2+} solution was rapidly poured into the 2-methylimidazole solution, and after 3 hours of stirring at room temperature, ZIF-8 nanocrystals were recovered from the mother liquor by centrifuging at 8000 rpm for 15 minutes and washing three times with fresh methanol. The final product was then dried in an oven overnight at 80 °C in air.

To prepare the BNC catalyst materials, 50 mg of the prepared ZIF-8 precursor was suspended in 2 mL of ethanol, followed by the addition of 5.4 mg phenylboronic acid. The resultant mixture was sonicated for 4 hours and subsequently dried in an oven overnight at 80 °C in air. These BNC pre-catalysts were then pyrolyzed at different temperatures ranging from 700 °C to 1100 °C for 3 hours under 100 s.c.c.m N_2 with a heating rate of $10 \text{ }^\circ\text{C min}^{-1}$. Once cooled to room temperature, the resultant powders were acid treated in 5 mL 1 M HCl for 1 hour with sonication, centrifuged, washed several times with water, and dried overnight at 80 °C. The boron and nitrogen co-doped carbon catalysts are denoted as Y-BNC, where Y represents the pyrolysis temperature. For comparison, nitrogen doped carbon pyrolyzed at 900 °C (900-NC) was synthesized in a similar manner without addition of the boron precursor.

2.3. Material characterization

Scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDS) elemental mapping were performed on a Hitachi SU8030 with an Oxford Aztec X-max 80 SDD EDS detector at 10 and 20 keV, respectively. X-ray photoelectron



spectroscopy (XPS) was conducted with a Thermo Scientific ESCALAB 250Xi with a monochromatic Al K α source and 500 μ m spot size. All XPS spectra were fitted using CasaXPS software and electron binding energy spectra were calibrated to adventitious carbon C 1s peak set to 284.8 eV. Raman spectra were collected on a laser confocal microscopy Raman with laser excitation at 532 nm.

Powder X-ray diffraction (PXRD) was conducted using a Rigaku Ultima IV with Cu K α radiation source ($\lambda = 0.1540$ nm). A scan rate of 5° min^{-1} was used to record the PXRD patterns for a 2θ range $10\text{--}90^\circ$. Inductively coupled plasma-mass spectrometry (ICP-MS) was conducted on a Thermo iCapQ ICP-MS (Thermo Fisher Scientific, Waltham, MA, USA) operating in KED mode and equipped with an ESI SC-2DX PrepFAST auto-sampler (Omaha, NE, USA). N_2 adsorption-desorption curves were performed at liquid nitrogen temperature (77 K) on a Micromeritics 3Flex instrument. Brunauer-Emmett-Teller (BET) surface area was calculated based on the adsorption data ($p/p^0 = 0.05\text{--}0.25$). Pore size distributions were calculated using non-local density functional theory assuming slit carbon pores. Electrode hydrophobicity was determined by the measurement of water contact angle which was performed using a contact angle goniometer (Ossila). *Ex situ* soft X-ray absorption spectroscopy (XAS) spectra at the K-edges of carbon, oxygen, nitrogen, and boron were collected at beamline 8-2 at Stanford Synchrotron Radiation Lightsource (SSRL). Total electron yield (TEY) data were acquired for samples in a single load at room temperature and under ultrahigh vacuum (10^{-9} torr).

2.4. Electrochemical RRDE measurements

The RRDE experiments were performed in a four-electrode configuration using a bipotentiostat (Biologic VSP) in a heart-shaped electrochemical cell equipped with a gas dispersion tube, a silver/silver chloride (Ag/AgCl sat'd KCl, Fisher) reference electrode, a Pt wire counter electrode, and two parallel working electrodes (a glassy carbon (GC) disk with deposited catalyst and a Pt ring, Pine Research Instrumentation, AFE7R9GCPT). The RRDE collection efficiency was determined experimentally to be $\sim 37\%$ using a ferrocyanide/ferricyanide half reaction system.⁴⁹ Catalyst ink for the GC disk working electrode was prepared by mixing 3.3 mg of catalyst, 1.00 mL of isopropyl alcohol (ThermoFisher Scientific) and 10 μ L of cation-exchanged Nafion solution (5%, Sigma-Aldrich). Then, the resultant ink was sonicated for three hours and 7.6 μ L of the catalyst ink was drop-casted onto a clean and polished GC disk substrate (0.246 cm^2) to obtain a catalyst loading of 0.1 mg cm^{-2} .

Prior to the ORR measurements, the Pt ring electrode was cleaned by successive cyclic voltammetry (CV) in N_2 -saturated $0.5 \text{ M K}_2\text{SO}_4$ electrolyte within the potential range 0.05 to 1.2 V vs. RHE at 20 mV s^{-1} until steady CVs were obtained. Baseline electrochemical assessment of the GC disk/catalyst included measuring background (capacitive) current between 0.8 V to 1.0 V vs. RHE , followed by electrochemical impedance spectroscopy (EIS) at open circuit voltage to determine the series resistance (R_s) of the liquid electrolyte for the subsequent iR compensation. All potentials were controlled with respect to a Ag/AgCl reference

electrode in saturated KCl and converted to an RHE scale using the expected redox potential and pH shift *via* $E(\text{V vs. RHE}) = E(\text{V vs. Ag/AgCl}) + 0.197 (\text{V}) + 0.059 (\text{V dec}^{-1}) \times \text{pH}$.

To assess catalytic activity, electrolyte was purged with O_2 for 30 minutes and then linear sweep voltammetry (LSV) measurements were carried out between 1.1 to 0.05 V vs. RHE at 5 mV s^{-1} with an RRDE rotation speed of 1600 rpm . To obtain Faradaic currents on the disk electrode, the capacitive current was subtracted from the total measured current. The amount of H_2O_2 produced at the disk was quantified by holding the Pt ring at 1.2 V vs. RHE to oxidize the H_2O_2 species transported from the disk electrode *via* rotation. The molar selectivity of H_2O_2 , which represents the ratio between O_2 consumption rate toward H_2O_2 and total O_2 consumption rate at the disk electrode, and the Faradaic efficiency of H_2O_2 , which represents the electron efficiency towards driving $2e^-$ vs. $4e^-$ ORR, were calculated according to the following eqn (1) and (2):⁴⁹

$$\text{Molar selectivity of } \text{H}_2\text{O}_2(\%) = 200 \times \frac{\frac{I_R}{N}}{|I_D| + \frac{I_R}{N}} \quad (1)$$

$$\text{Faradaic efficiency of } \text{H}_2\text{O}_2(\%) = 100 \times \frac{\frac{I_R}{N}}{|I_D|} \quad (2)$$

where I_R is the ring current, I_D is the disk Faradaic current, and N is the collection efficiency (0.37 , measured with a ferrocyanide/ferricyanide half reaction, determined as the ratio of anodic ring limiting current to the cathodic disk limiting current).

Tafel slopes ($\frac{\alpha F}{2.3RT} \equiv b$) can be calculated using the Tafel equation (eqn (3)) obtained from the linear relationship between the overpotential (η) vs. log kinetic disk current density ($j_{\text{disk,kin}}$).

$$\eta = \frac{2.3RT}{\alpha F} \times \log(j_0) + b \times \log(j_{\text{disk,kin}}) \quad (3)$$

Here, α is the transfer coefficient, F is the Faraday's constant (96485 C mol^{-1}), R is the ideal gas constant (8.314 J mol^{-1}), T is temperature (298 K), and j_0 is the exchange current densities. The kinetic disk current densities ($j_{\text{disk,kin}}$) can be determined according to the Koutecky-Levich equation (eqn (4)):⁵⁰

$$\frac{1}{j_{\text{disk}}} = \frac{1}{j_{\text{disk,kin}}} + \frac{1}{j_{\text{disk,lim}}} \quad (4)$$

where j_{disk} and $j_{\text{disk,lim}}$ are the measured current density and the diffusion-limited current density, respectively.⁵¹ In addition, $j_{\text{disk,lim}}$ can be calculated using the Levich equation (eqn (5)):

$$j_{\text{disk,lim}} = 0.62 \times n \times F \times D^{\frac{2}{3}} \times \nu^{-\frac{1}{6}} \times \omega^{\frac{1}{2}} \times C \quad (5)$$

where n is the number of electrons transferred, F is the Faraday's constant (96485 C mol^{-1}), C is the bulk concentration of oxygen ($1.2 \times 10^{-6} \text{ mol cm}^{-3}$), D is the diffusion coefficient of oxygen ($1.9 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$), ν is the kinetic viscosity ($0.01 \text{ cm}^2 \text{ s}^{-1}$), and ω is the angular velocity of the disk ($\omega = 2\pi N$, N is the linear rotation speed).^{21,52}



2.5. Electrochemical GDE flow cell measurements

As shown in Fig. 4A and S1,[†] a custom-made three-electrode GDE flow cell was constructed to enhance reactant mass transport and mimic geometry of an applied reactor design that could be used for larger-scale electrochemical generation of H₂O₂. The catalyst ink for the working electrode was prepared by mixing 4.5 mg of catalyst with 900 μ L isopropanol and 45 μ L Nafion binder. The resulting catalyst ink was ultrasonicated for 1 hour to obtain a homogeneous dispersion of catalyst powder. Afterward, the catalyst ink was spray coated onto a 3 $\times$ 3 cm² hydrophobic carbon cloth (Fuel Cell Store, W1S1011) resulting in a mass loading of $\sim$ 0.5 mg cm⁻². After drying overnight at room temperature, the spray coated carbon cloth was cut into four 1.5 $\times$ 1.5 cm² electrodes. The anode catalyst for the oxygen evolution reaction was an Ir-based electrocatalyst on Ti felt (platinized, fuel cell store) prepared by a dip coating and decomposition method.⁵³ Ti felt was first etched in 0.5 M oxalic acid (anhydrous 98%, Fisher) at 80 $^{\circ}$ C for 1 hour. Next, the electrode was dipped into a solution containing 30 mg IrCl₃·xH₂O (TCI), 1 mL concentrated HCl, and 9 mL isopropyl alcohol. The as-prepared Ti felt was then dried at 100 $^{\circ}$ C for 5 minutes followed by calcination at 500 $^{\circ}$ C for 15 minutes in air. This process was repeated 5 times. A fresh piece of IrO_x/Ti anode was used for each electrochemical experiment. The GDE flow cell was assembled with catalyst layers facing the flowing liquid electrolyte; the exposed geometric surface area of each catalyst was 1 cm². The cathode and anode compartments were separated with a Nafion-115 proton exchange membrane (fuel cell store) and accommodated 1 mL electrolyte each. Prior to operation, the Nafion-115 membrane was pre-treated with 5% (v/v) H₂O₂ for 1 hour at 80 $^{\circ}$ C, followed by 10% (v/v) H₂SO₄ for 1 hour at 80 $^{\circ}$ C, then DI water for 1 hour at 80 $^{\circ}$ C. Membranes were stored in DI water. Both catholyte and anolyte liquid flow rates were set to 3 mL min⁻¹ using a peristaltic pump (BT100L with dual YZ15 pump heads, Golander) from respective 10 mL reservoirs containing 0.5 M K₂SO₄. 20 s.c.c.m. of O₂ was supplied to the cathode side *via* a mass flow controller (MKS Instruments). ORR was conducted using a potentiostat (Biologic VMP3) with a Ag/AgCl reference electrode. All potentials were corrected to the RHE potential scale and manually compensated for iR to 100%. CVs were performed at a scan rate of 20 mV s⁻¹ with a cell voltage range from 1.1 to 0.05 V *vs.* RHE. Experiments were carried out using chronoamperometry (CA) at 0.20 V *vs.* RHE to evaluate the H₂O₂ production rates by measuring H₂O₂ concentration after set reaction times (See Section 2.7).

2.6. Electrochemical MEA flow cell measurements

H₂O₂ electrosynthesis was further evaluated using a two-electrode setup in a dual MEA SE flow cell with Ti flow fields (5 cm² formic acid electrolyzer, Dioxide Materials), comprising a porous SE compartment sandwiched by a cation and an anion exchange membrane (CEM = Nafion-115 and AEM = Sustainion X37-50 (Grade RT), respectively) on the anode and cathode side, respectively. Outside of these membranes, the anode/cathode catalyst layers and flow plates are compressed (with the anode adjacent to the CEM and the cathode adjacent to the AEM).

Polytetrafluoroethylene (PTFE) gaskets were used to seal the electrolyzer. The cell configuration is illustrated in Fig. S2.[†] The middle chamber SE material was chosen as a styrene-divinylbenzene sulfonated copolymer Amberchrom 50WX8 hydrogen form cation conductor (50–100 mesh, Sigma-Aldrich). Prior to use, the SE was washed in a beaker with 0.5 M K₂SO₄ several times until the pH reached approximately 5. In the SE chamber, Milli-Q water was flowed in a single pass at 0.5 mL min⁻¹ to wash out H₂O₂ produced, while the anode side was circulated with 1 M H₂SO₄ at 3 mL min⁻¹. The O₂ flow rate to the cathode flow plate was supplied with 20 s.c.c.m. of humidified O₂ controlled by a mass flow controller and water/electrolyte flow rates were controlled by a peristaltic pump. A Biologic VMP3 potentiostat with 10 A booster was employed to record the electrochemical response. Two pieces of copper tape (3 M) were used to connect the electrolyzer to potentiostat leads. The electrolyzer was first cycled between open circuit potential and 2.5 V, then held at 2.5 V for 10 minutes for activation. Then, 40 minutes chronopotentiometry (CP) holds in increasing current densities from 10 to 200 mA cm⁻² were applied to the electrolyzer. The product streams of the first 10 minutes of each CP hold were discarded, and product streams from the next 30 minutes were collected to examine the H₂O₂ concentration produced.

2.7. H₂O₂ product quantification

The produced H₂O₂ was quantified using a cobalt-carbonate (Co/CO₃) assay coupled with UV-vis spectrophotometry (Agilent Cary 5000).⁵⁴ The Co/CO₃ assay contains a mixture of KHCO₃ and cobalt sulfate heptahydrate (CoSO₄·7H₂O) both of which dissociate in water and exhibit a light pink color solution. H₂O₂ oxidizes Co²⁺ in the presence of bicarbonate likely forming a 'carbonatocobaltate' complex (Co(CO₃)₃³⁻) which strongly absorbs at 290 nm and weakly in the visible region (440 and 635 nm), exhibiting a dark green color. The measured absorbance of the Co(CO₃)₃³⁻ complex is linearly proportional to the concentration of H₂O₂ present; the 290 nm feature was used for quantification unless otherwise specified. The Co/CO₃ assay was prepared by first mixing 1.35 mL of 2 M KHCO₃ and 35.8 μ L of 70 mM CoSO₄·7H₂O in a cuvette. Next, 35.8 μ L of our H₂O₂ sample (known or unknown concentration) was added to the cuvette and mixed. Immediately afterward, 1.425 mL of 2 M KHCO₃ was added to the cuvette and mixed. The absorbance of these samples was measured in the wavelength range 200–800 nm with 1 nm resolution and scanning rate of 2 nm s⁻¹. The UV-vis spectra were background-subtracted using a baseline with 35.8 μ L of H₂O added instead of H₂O₂-containing samples. A typical calibration curve was plotted by linear fitting the absorbance values at wavelength of 290 nm for various known concentrations of H₂O₂ from 20 ppm to 750 ppm (Fig. S3[†]). The Faradaic efficiency for H₂O₂ production is calculated using the following equation:

H₂O₂ Faradaic efficiency (%) =

$$100 \times \frac{C_{\text{H}_2\text{O}_2}(\text{mol L}^{-1}) \times 2 (\text{mol e}^-) \times V(\text{L}) \times 96485 (\text{C mol}^{-1})}{I(\text{A}) \times \text{time}(\text{s})} \quad (6)$$



where C is concentration of H_2O_2 , V is the volume of electrolyte, I is the current applied, and time is the total reaction time.

3. Results and discussion

3.1. Structural characterization

ZIF-8 was chosen as the precursor for the catalyst preparation due to their superior thermal and chemical stability compared to other metal-organic-frameworks (MOFs). During the carbonization process, ZIF-8 decomposes to form microporous carbon frameworks, which provide abundant active sites and high surface areas for electrocatalytic reactions. Furthermore, the high nitrogen content in the ZIF-8 precursor results in incorporation of nitrogen into the carbon framework during the carbonization process, making ZIF-8 attractive as both the carbon support and nitrogen precursor.⁵⁵ As illustrated in Fig. 1A, a two-step method to prepare the BNC catalysts was adopted. Briefly, phenylboronic acid, the chosen boron source, was mixed with ZIF-8 in ethanol overnight, followed by pyrolysis under N_2 for 3 hours, then acid washed to remove any residual Zn from the ZIF-8 support. Remaining Zn in the structure forms $\text{Zn}(\text{CN})_2$ during calcination, which further decomposes and vaporizes at temperatures above

800 °C.⁵⁶ Further acid treatment would deactivate Zn towards ORR, as Zn has not been shown to be an effective ORR catalyst. Due to the different formation energies of the B–N, B–C and N–C species, the B–N–C configurations can be tuned by adjusting the pyrolysis temperature. To systematically study the effects of pyrolysis temperature and identify the optimal conditions to support H_2O_2 production, catalyst materials were carbonized at temperatures ranging from 700 °C to 1100 °C. For comparison, nitrogen doped carbon pyrolyzed at 900 °C (900-NC) was prepared in a similar manner without the addition of the boron precursor.

To characterize the morphology of the BNC catalyst materials, SEM was employed, as shown in Fig. 1B for 900-BNC and in Fig. S4† for all other catalysts. SEM images for all the catalysts show similar nanoparticulate morphologies and highly porous structures which are favorable for facilitating mass transport of reactants and products. The EDS elemental mapping of the as-prepared catalyst materials (Fig. 1C and S5†) exhibits uniform distribution of C, N, B, and O elements, validating the successful doping of B and N atoms into the carbon framework.

Raman spectra were also acquired to examine the degree of graphitization, the defect content, and extent of disordered

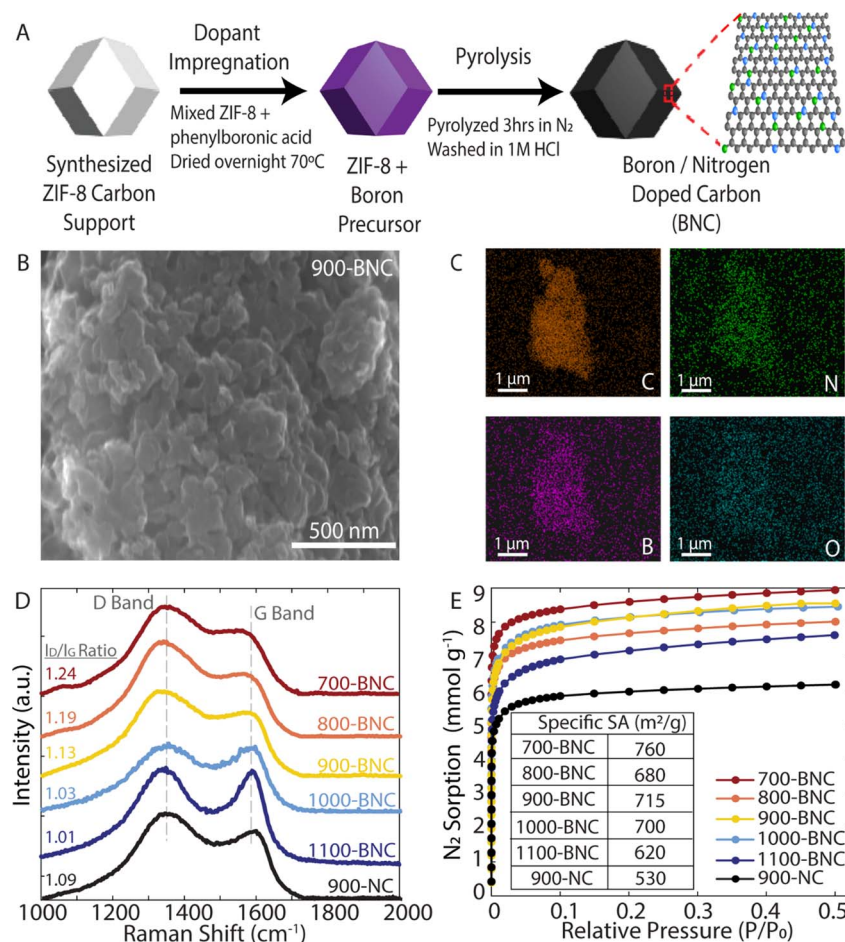


Fig. 1 Synthesis and characterization of the catalysts. (A) Schematic illustration of the BNC catalyst synthesis. (B) SEM image of the 900-BNC catalyst and (C) corresponding EDS mapping images. (D) Raman spectra. (E) N_2 adsorption/desorption isotherms including the corresponding calculated specific surface areas.



structures. As shown in Fig. 1D, all materials exhibit typical D band (1350 cm^{-1}) and G band (1580 cm^{-1}) peaks; the D band is related to disordered graphitic structures and the G band reflects the degree of graphitization and presence of sp^2 -hybridized carbon.⁵⁷ Therefore, the extent of defect formation in the BNC catalysts can be quantified by using the intensity ratio of the D and G bands (I_D/I_G). The I_D/I_G ratio (Table S1†) decreases with increasing pyrolysis temperature, indicating an improved degree of graphitization. Comparing 900-NC and 900-BNC catalysts, the intensity ratio of D band increases with the addition of boron, which we attribute to the creation of additional defects in the carbon matrix. The structural properties of the samples were further characterized by PXRD. As displayed in Fig. S6,† all samples show two broad diffraction peaks at 25° and 44° , corresponding to the (002) and (100) lattice planes, respectively.^{58,59} This indicates that the structures are mainly amorphous, which is typical for nanoporous carbon materials, also supported by the absence of peaks from crystalline hexagonal boron nitride (h-BN) in the as-prepared samples.⁶⁰ We further observe the (002) and (100) peaks becoming more prominent and less broad with increasing pyrolysis temperatures, indicating a higher degree of graphitization. Prior studies have reported that the graphitic carbon moieties formed during pyrolysis increase the conductivity of carbon materials (*i.e.*, charge transport),⁶¹ which may lead to a boost in ORR performance.

Nitrogen adsorption-desorption isotherms were measured to determine specific surface area and pore size characteristics. As displayed in Fig. 1E, all BNC catalysts display higher BET surface areas than 900-NC, indicating that the introduction of boron is beneficial to the enhancement of surface area. However, we find that BET surface area and total pore volume generally decrease with increasing pyrolysis temperature from 700-BNC ($760\text{ m}^2\text{ g}^{-1}$; $0.31\text{ cm}^3\text{ g}^{-1}$) to 1100-BNC ($620\text{ m}^2\text{ g}^{-1}$; $0.26\text{ cm}^3\text{ g}^{-1}$) consistent with the carbon structures becoming more graphitic, and in agreement with our Raman and XRD results (Fig. 1D and Table S2†). All catalysts showed typical type I isotherms indicative of a characteristic micropore morphology, similar to other previously reported ZIF-8 derived carbon materials.⁶² These results match the patterns of pore size distributions in Fig. S7† where the dominant pore size is between *ca.* 1–1.5 nm.^{63,64} Combining this characterization and analysis, it can be concluded that the incorporation of boron is favorable for enhancing the surface area and inducing formation of a highly defective structure, while tuning pyrolysis temperature allows for more fine tuning of the graphitic carbon structure, thereby providing advantages for further catalytic applications.

X-ray photoelectron spectroscopy (XPS) was conducted to examine the electronic structure and chemical composition of the BNC catalysts and 900-NC. As shown in Table 1 and Fig. S8,† all BNC materials primarily consist of C, N, O, and B elements at their surface, with residual amounts of Zn remaining from the ZIF-8 precursor. Although pyrolysis was performed under a N_2 gas environment, oxygen content is observed in the BNC materials, likely originating from either oxidation of NC support after being in contact with air or residual phenylboronic acid

Table 1 Elemental analysis of the catalyst materials obtained by different temperature treatments derived from XPS measurements

Sample	C (at%)	O (at%)	Zn (at%)	N (at%)	B (at%)
700-BNC	64.6	6.9	2.6	25.2	0.71
800-BNC	68.6	5.4	3.2	21.7	0.98
900-BNC	75.4	7.5	1.6	13.8	1.78
1000-BNC	85.0	7.3	0.4	6.3	1.01
1100-BNC	85.6	7.9	0.2	4.7	1.63
900-NC	80.9	5.6	1.8	11.7	—

form the B precursor.⁶⁵ Continuous loss of N content was observed with increasing pyrolysis temperature from 25.2% for 700-BNC followed by 21.7%, 13.8%, 6.3%, and 4.7% for 800-BNC, 900-BNC, 1000-BNC, and 1100-BNC, respectively. This is due to the nitrogen in the ZIF-8 being removed *via* the N_2 gas flow when the framework is destroyed at elevated temperatures. We note that the N contents for these materials are higher than that of typically reported N doped carbon materials under the same temperature conditions.⁶⁶ Based on previous reports, this likely arises due to the synergistic effect of co-doping with boron and nitrogen, where boron dopants alter the electronic properties of the carbon structure, thereby promoting additional nitrogen incorporation.³⁹ On the other hand, C concentrations trend closely with increasing pyrolysis temperature from 64.6% to 85.6% for 700-BNC and 1100-BNC, respectively. For boron composition, the total B content generally increases from low to high pyrolysis temperature, although 900-BNC uniquely displays the highest atomic percent (1.78%). We postulate the decrease in B content above 900 °C may be due to structural changes (*e.g.*, fewer defects) or surface chemical changes (*e.g.*, lower N content) at elevated temperatures.

Different bonding configurations can be formed in the carbon framework by tuning pyrolysis temperature and incorporating boron atoms. As shown in Fig. 2A, the high-resolution XPS B 1s spectra can be deconvoluted into three peaks at 190 eV, 191 eV and 192 eV assigned to B–N–C (*e.g.*, BN_2C , BNC_2), B–N, and B–O bond structures, respectively.^{67–70} We note that the FWHM peak widths for the B 1s spectra range between 1.9 and 2.5 eV, which are significantly greater than h-BN FWHM value of 1.5 eV. This broadening may be due to the superposition of multiple peaks in the spectra, in agreement with previous reports.⁷¹ No significant features are observed between 187–188.7 eV; previous work has assigned peaks in this region to B–C bonding similar to boron carbide (B_4C).⁷² At lower pyrolysis temperatures, the boron atoms present are predominately bonded with oxygen (B–O), consistent with the O 1s XPS spectra (Fig. S11B†). It is noteworthy that the proportion of B–N–C and B–N configurations, previously correlated with improved 2e^- ORR performance,^{32,42} gradually increases with increasing pyrolysis temperature, while B–O slightly decreases (Table S4†). More specifically, for the 900-BNC catalyst, the B–N, B–N–C, and B–O configurations account for approximately 44%, 29%, and 27%, respectively. Interestingly, the sum of the B–N and B–N–C in 900-BNC accounts for $\sim 73\%$ of the total boron configurations and is the highest among the catalysts studied.



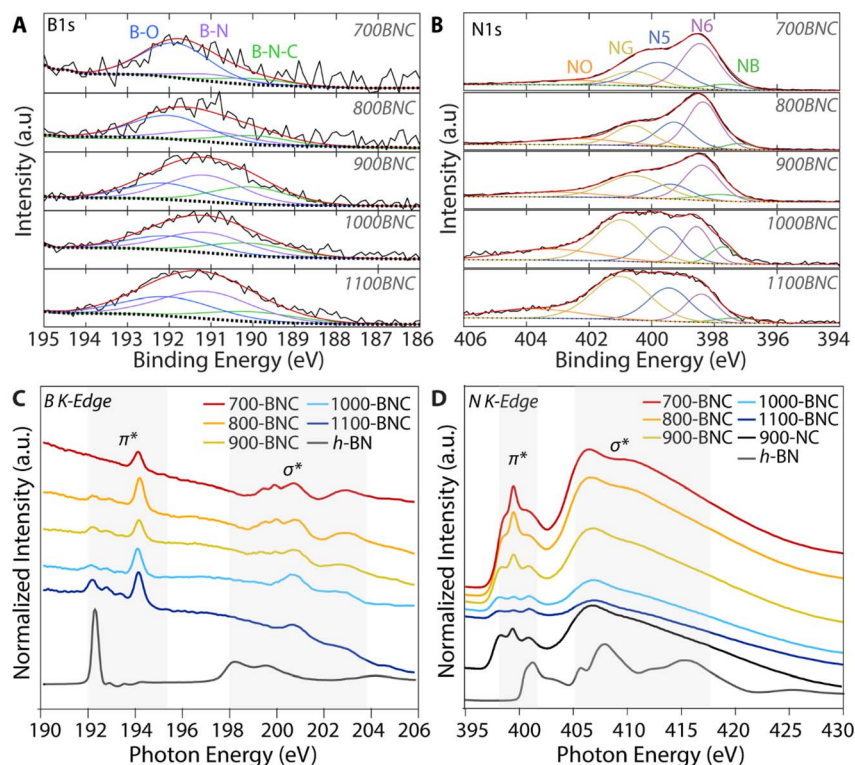


Fig. 2 Composition investigation of the catalysts. (A) High-resolution B 1s and (B) N 1s XPS spectra. (C) Boron K-edge and (D) nitrogen K-edge X-ray absorption spectroscopy (XAS).

Similarly, the N 1s spectra were deconvoluted and peaks were identified at 397.7 eV, 398.5 eV, 399.5 eV, 400.7 eV, and 403 eV, which are assigned to N-B, pyridinic-N (N6), pyrrolic-N (N5), graphitic-N (NQ), and oxide-N (NO), respectively (Fig. 2B).^{32,34,73,74} We acknowledge the exact chemical state of the N-B species cannot be clearly defined due to difficulty in deconvoluting the spectra. Based on previous reports, the N-B species likely comprise a B-N-C type structure or a B-N-B moiety similar to h-BN (Fig. S9†).⁶⁸ While the surface boron atomic percent and the ratio of total B to total N are both small (such that the absolute content of N-B moieties is small), 900-BNC exhibits the highest relative N-B content (7.5%), followed by 1000-BNC (6.1%), 1100-BNC (5.6%), 800-BNC (4.8%), and 700-BNC (4.0%). In addition, we observe in Fig. S10 and Table S5† that, as total nitrogen content decreases with increasing pyrolysis temperature, the relative amount of pyridinic-N decreases whereas the graphitic-N increases, since the stability of graphitic-N is superior to that of pyridinic-N at higher carbonization temperatures.⁷⁵ Prior studies have shown that a higher ratio of graphitic-N and N-B species is correlated with improved H₂O₂ selectivity, suggesting these species may act as active sites for the ORR catalysis.^{21,39}

We also deconvolute the high-resolution XPS C 1s spectra, but it is difficult to discern between different types of C in these materials because C species such as C=C, C-C, C-OH, C=O, and O=C-OH exhibit similar chemical shifts.^{16,43} For the purpose of simplifying the analysis, the C 1s spectra are divided into four different peaks centered at 283.9 eV, 284.5 eV, 286 eV,

and 288 eV, corresponding to C-B, sp² carbon (C=C), C-N, and C=O, respectively (Fig. S11A and Table S6†).^{43,74} Based on this fitting, we observe that the percentage of C=C bonds increases from ~38% to ~49% with the rise in pyrolysis temperature, which is consistent with the increase in sp² hybridized carbon observed from Raman. Additionally, the contribution of C-B increases gradually with increasing pyrolysis temperature, while the C-N appears to decrease. On the other hand, C=O does not appear to trend, consistent with the O 1s spectra in Fig. S11B.† These XPS results demonstrate that by tuning the pyrolysis temperature, one can control the formation of different N-B, B-N-C, and N-C bonds, and therefore, the nature and content of the B-N-C moieties.⁷⁶

To provide a more in-depth understanding of the local geometric structure of the B-N-C moieties in the as-prepared carbon catalysts, soft X-ray absorption spectroscopy (XAS) measurements were performed at the respective elemental K edges. For comparison, h-BN was included as a reference material to provide insight on the types of dopant bond motifs. As shown in Fig. 2C, the B K-edge spectra of h-BN exhibits the typical intense peak at 192.3 eV, with two smaller peaks at 193.0 eV and 193.6 eV. The peak at 192.3 eV is a characteristic B 1s → π* transition in h-BN, reflecting sp² hybridization of boron trigonally coordinated with nitrogen (BN₃). The BNC materials pyrolyzed above 700 °C exhibit three peaks at these same energy positions as h-BN; however, the relative intensities are much more even, and the total intensity of these features increases with pyrolysis temperature, coinciding with



a decrease in total N content (Table 1). As discussed previously, a more complete carbonization process at higher temperatures generates additional sp^2 hybridized carbon and the boron incorporates into the carbon matrix forming a trigonally coordinated boron. Similarly to h-BN, these features in the BNC catalyst spectra are attributed to the π^* transitions of B–N bonds as in h-BN (e.g., BN_3), but also include signal from trigonal BN_2C and BNC_2 moieties.^{69,77,78} At slightly higher energy, the BNC materials display a sharp peak at 194.4 eV that is absent in h-BN, corresponding to B–O bonding consistent with reports of bulk B_2O_3 . However, the absence of the σ^* feature at ~ 202 eV observed in B_2O materials suggests the formation of a mixture of BO_3 species that are distinct from bulk B_2O_3 on the surface of the BNC catalysts.⁷⁹ At even higher energies, we observe several peaks in the σ^* region between 199–202 eV for BNC materials pyrolyzed at lower temperatures (700–900 °C) which are ascribed to the central boron atom binding to species in the carbon matrix forming sp^3 hybridization with a tetrahedral structure. Moreover, the broad peak around 203 eV represents contributions from both trigonal and tetrahedral boron, in agreement with previous reports.⁸⁰ These features suggest that, at lower carbonization temperatures, the BO_3 species and its derivatives are highly embedded in the carbon matrix and likely bind with additional species (e.g., N, C) forming a tetrahedral coordination. Furthermore, the peaks located at 199–202 eV and 203 eV in the σ^* region decrease in intensity indicating less boron sp^3 hybridization with increasing pyrolysis temperature. Consistent with the B 1s XPS results, no features assigned to boron carbide are observed; when present in other reported materials, these boron carbide species are characterized by a $1s \rightarrow \pi^*$ transition at lower photon energy (~ 190.9 eV).⁶⁷ The lack of a feature in this energy region indicates a low concentration or simply undetectable distinct B–C bonds in the BNC catalyst materials.

Further insights were obtained from the N K-edge XAS spectra of the BNC catalysts. As shown in Fig. 2D, h-BN displays a strong π^* resonance at 400.3 eV and two features at 407 eV and 414.5 eV in the σ^* region, respectively, validating typical h-BN domains with sp^2 hybridized N–B bonding.³² As noted previously from the XPS results, the B : N ratio is quite small in the BNC materials, such that B–N moieties contribute a significant signal to B K XAS spectra, but do not necessarily contribute a significant signal to N K XAS spectra, relative to intensity of other N-containing species (especially N–C). For 900-NC and all the BNC catalyst materials, we observe three distinct peaks in the π^* region between 397–402.5 eV. Compared to the h-BN feature at 400.3 eV, these π^* transition features are shifted to lower photon energy indicating formation of an N bonding structure that is weaker than N–B in a h-BN moiety. The two lower energy peaks at 398.4 eV and 399.5 eV are in close agreement with the π^* transitions from pyridinic-N and pyrrolic-N, respectively.^{81,82} The feature at ~ 400 eV is present in both the 900-NC catalyst and h-BN; while this feature likely represents another N–C species for the 900-NC sample (such as graphitic-N), this may represent B–N–C type bonding (BN_2C or BNC_2) for the BNC materials.^{32,42,83,84} Previous computational simulations suggest that a decreased BN domain size in carbon

could also lead π^* peaks to shift to a lower energy, which is attributed to the increased ratio of interfacial N–B–C species compared to that of N–B in h-BN.⁸⁴ Focusing on the higher energy features, we observe shifts in energy position and broadening of the σ^* resonances (which becomes even more pronounced at higher pyrolysis temperatures) for the BNC materials vs. h-BN; these differences are representative and have been identified as a “fingerprint” of N–B–C bonding.^{77,85} Therefore, based on these XAS results, it is evident that the N–B–C coordination structure (e.g. N_2BC , NBC_2) becomes more dominant with increasing pyrolysis temperature resulting in weaker N–C structure.

In the C K-edge XAS spectra, we observe a characteristic π^* resonance from sp^2 -like carbon ($C=C$) at 285.4 eV and σ^* resonance centered around 293 eV (Fig. S12†).^{69,86,87} The intensity of these features increase with carbonization temperature demonstrating an enhanced graphitic network at elevated temperatures, in agreement with Raman and XPS results. In addition, the features at 287.8 and 288.8 eV are consistent with C atoms that are attached to O containing groups, such as C–O bonds or carbonyls ($C=O$).⁸⁸ Interestingly, the lack of the C–B feature in the lower photon energy region, as observed in the XPS results, suggests that B is most likely to be in the form of B–N or B–N–C moieties. The feature at 287 eV is consistent with C–N bonds,^{89,90} which appears to decrease for catalysts pyrolyzed at higher temperatures, as the C is more likely to graphitize and lose N content.

3.2. Electrocatalytic ORR performance

The ORR electrocatalytic activity and selectivity of the BNC catalysts were first evaluated using a three-electrode RRDE in O_2 -saturated 0.5 M K_2SO_4 (pH = 7). Fig. 3A shows linear sweep voltammogram (LSV) curves for the disk with the associated ring currents, corresponding to the total ORR current (i.e., both H_2O_2 and H_2O production) and H_2O_2 production current, respectively. The disk onset potential (defined in this study as the potential when disk current density reaches -0.20 mA cm^{-2}) shifts to less cathodic potentials (lower overpotentials) with increasing pyrolysis temperature such that 1100-BNC demonstrates the least cathodic onset potential (0.42 V vs. RHE) and 700-BNC demonstrates the most cathodic onset potential (0.17 V vs. RHE). Additionally, 1100-BNC shows the highest total disk current density (-1.55 mA cm^{-2}) at 0.1 V vs. RHE, followed by 1000-BNC (-1.47 mA cm^{-2}), 900-BNC (-1.41 mA cm^{-2}), 800-BNC (-0.96 mA cm^{-2}), and 700-BNC (-0.36 mA cm^{-2}), indicating overall ORR activity improves with increasing pyrolysis temperature. We postulate that this improvement in overall electrocatalytic activity is due to the higher degree of disordered graphitic structure (Fig. S19A†) and an increase in the B : N ratio (Fig. S19B†), which allows for more facile electron transfer and enhances the accessibility of active sites.⁶¹

With respect to the selective $2e^-$ ORR activity, or the onset of the ring current density, a different activity trend is observed with 900-BNC demonstrating the least cathodic onset potential, followed by 1000-BNC, 1100-BNC, 900-NC, 800-BNC, and 700-BNC which all demonstrated higher overpotentials for onset. To



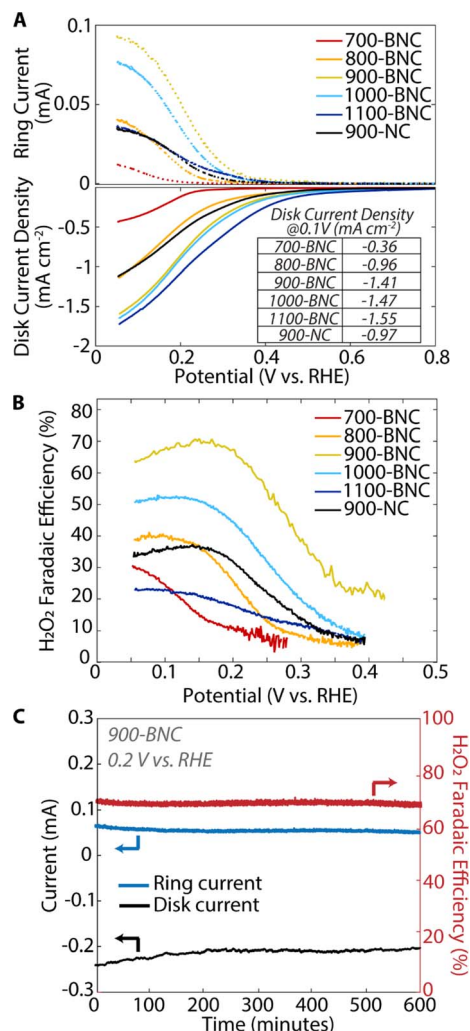


Fig. 3 ORR performance using a RRDE of the catalysts in neutral pH solutions. (A) Linear sweep voltammetry (LSV) curves at 1600 rpm for the disk electrode (solid lines) and simultaneous H₂O₂ detection currents at the ring electrode (dashed lines) in 0.5 M K₂SO₄. (B) Calculated H₂O₂ Faradaic efficiency (%) over the applied potential range derived from RRDE data. (C) Chronoamperometric responses of 900-BNC at 0.20 V vs. RHE for 10 hours.

understand these trends, we first examined the accessibility of active sites for these BNC materials. We note that although BET surface area generally decreased at elevated temperatures, these surface areas may not accurately reflect the accessible active sites in an electrocatalytic reaction. Moreover, the pore size and volumes of microporous and mesoporous structures show no correlation with H₂O₂ electrosynthesis performance (Fig. S20†). Thus, we also estimated the electrochemically active surface area (ECSA) by measuring the electrochemical double-layer capacitance of all catalysts in N₂-saturated electrolyte (Fig. S13–18†). As displayed in Fig. S19C,† the estimated ECSA of BNC catalysts follows the order 900-BNC > 1100-BNC > 1000-BNC > 900-NC > 800-BNC > 700-BNC; this surface area trend is similar to the order of selective 2e⁻ ORR activity, suggesting that a high ECSA plays an important role in improving H₂O₂ performance. Interestingly, 700-BNC shows a similar BET surface area as the 900-BNC but displays a much lower ECSA.

Based on prior studies, we found this poor charge transfer for 700-BNC is related to the high N content and lower degree of graphitization.⁹¹ Additionally, despite a higher ECSA value, 1100-BNC shows inferior activity compared with 1000-BNC, suggesting that ECSA is not the only decisive factor accounting for 2e⁻ ORR activity.

The calculated H₂O₂ Faradaic efficiencies (Fig. 3B) and molar selectivities (Fig. S22†) reveal a slightly different behavior for 2e⁻ ORR performance. Notably, the 900-BNC catalyst delivers the highest H₂O₂ Faradaic efficiency and molar selectivity of these catalysts, at 70% and 80%, respectively, demonstrated at 0.20 V vs. RHE. The other BNC catalysts demonstrate H₂O₂ Faradaic efficiency between 20–50% at the same potential with 700-BNC and 1100-BNC exhibiting the overall lowest 2e⁻ ORR performance. Moreover, 900-BNC also demonstrated the highest partial kinetic current density at 0.20 V vs. RHE of 0.82 mA cm⁻² (Table S7†). In addition to activity and selectivity, the electrochemical stability of the 900-BNC catalyst was evaluated using chronoamperometry (Fig. 3C). After 10 hours of continuous electrocatalytic H₂O₂ production (constant voltage of 0.20 V vs. RHE), 900-BNC retained ~90% of its initial activity demonstrating excellent overall stability under these reaction conditions. In a solid membrane electrode assembly electrolyzer, 900-BNC catalyst maintained its activity over 50 h of operation, producing a pure H₂O₂ stream with average H₂O₂ concentration above 300 mM (11 000 ppm) (Fig. S28†). Overall, the 900-BNC catalyst demonstrates a superior 2e⁻ ORR performance in neutral pH compared to other reported carbon-based catalysts (Table S9†), exhibiting a relatively high H₂O₂ activity, high selectivity, and excellent stability.

We postulate that the differences in ORR activity and selectivity observed for the BNC catalyst materials may be attributed to the synergistic effects of the chemical bonding between B and N dopants in the carbon matrix. From combined XAS and XPS analyses, we found that by changing the pyrolysis temperature, the relative formation of B–N, N–C and B–N–C dopant structures can be tuned. With increasing pyrolysis temperature from 700–900 °C, there is an energetic driving force for the formation of local BNC dopant structures, consisting of sp³ hybridized boron and nitrogen atoms in the carbon matrix (e.g., BN₂C or BNC₂) with tetrahedral coordination. Previous work investigating the ORR energetics using density functional theory for boron-carbon nitride materials demonstrated that the B–N–C dopant structures, namely the BNC₂ configuration, exhibits an optimal charge state of B sites and OOH* adsorption energy, thus favoring H₂O₂ production.⁹² In contrast, samples obtained at higher pyrolysis temperatures above 1000 °C, form distinct B–N and graphitic-N bonds, similar to h-BN (e.g., BN₃) and N-doped carbon, which comprise the dominant dopant bond motifs. The formation of these isolated B and N graphitic dopants likely leads to a reduction in H₂O₂ performance, thus shifting to the 4e⁻ pathway to H₂O. Therefore, our results indicate that the higher content of local B–N–C dopant configurations in the 900-BNC catalyst likely contributes to its enhanced electrocatalytic performance (Fig. S19D†).

Moreover, based on the previous N content analysis, we postulate that the moderate N doping of the 900-BNC catalyst is



more favorable to the $2e^-$ ORR (Fig. S19E†). Conversely, excessive N doping and lower degree of carbon graphitization, particularly in the 700-BNC and 800-BNC catalysts, may lead to lower ORR rates due to poor charge transfer and selectivity shift towards the $4e^-$ pathway, as reported for other N-doped carbon materials.⁹¹ On the other hand, the very low N content of the 1100-BNC catalyst suggests a deficiency in catalytic ORR active sites, which is unfavorable for achieving a high catalytic current. Moreover, the specific N configurations may contribute to the improved H_2O_2 performance. Pyrrolic and pyridinic moieties were found to impede H_2O_2 formation, similar to previous reports (Fig. S19F and G†) while graphitic moieties improve H_2O_2 production (Fig. S19H†). Graphitic-N has been reported to be favorable for the $2e^-$ pathway due to the lower energy barrier for O_2 adsorption compared to pyridinic-N.²⁰ Notably, pyrrolic-N is widely regarded to be less active for the $4e^-$ ORR, and more recently, has been correlated with high activity for the $2e^-$ ORR in alkaline conditions.⁸⁸ In this work, the 900-BNC catalyst possesses primarily graphitic-N as well as the highest N-B content, further confirming the important role of B-N-C moieties in catalyzing the $2e^-$ ORR to H_2O_2 .

3.3. H_2O_2 electrosynthesis in flow cell

To investigate the as-prepared BNC catalysts for practical applications, we employed a three-electrode flow cell equipped with a gas diffusion electrode (GDE) in neutral pH electrolyte (Fig. 4A). A GDE cell design alleviates the intrinsic mass transport limitations by utilizing a hydrophobic gas diffusion layer (GDL, carbon cloth) to support formation of a solid-liquid-gas triple phase boundary with the catalyst, electrolyte, and gaseous O_2 reactant, enabling improved mass transport.^{93–95} The bare GDE support (carbon cloth) exhibits a water contact angle of $135 \pm 2^\circ$ which is critical to ensure efficient gas transport and prevent electrode flooding under reaction conditions.^{93,96,97} After BNC catalysts are deposited on the carbon paper, the electrodes exhibit water contact angles of 110 ± 2 to $125 \pm 2^\circ$ (Fig. S23†); the 900-BNC electrode presents moderately higher hydrophobicity than the other electrodes, which may be beneficial in enhancing the accessibility of O_2 to the catalyst surface.^{98,99}

To evaluate H_2O_2 electrosynthesis performance, we conducted chronoamperometry at 0.20 V vs. RHE for two hours in 0.5 M K_2SO_4 and measured the corresponding current response (Fig. S24†). In this set-up, the electrolyte and H_2O_2 product are continuously recycled such that the product concentration builds up over time (Fig. 4B). For all BNC catalyst materials, H_2O_2 concentration increases linearly with time. Remarkably, 900-BNC delivers superior performance reaching ~ 180 mM H_2O_2 concentration, 78% H_2O_2 Faradaic efficiency, and $0.9 \text{ mmol cm}^{-2} \text{ h}^{-1}$ H_2O_2 production rate, outperforming most of the previously reported non-metal carbon-based catalysts in neutral pH (Table S10†). Following electrochemical testing, the catholyte composition was sampled *via* ICP-MS to examine the catalyst stability. As displayed in Table S8,† very little B content dissolved in the catholyte after the two-hour measurement, demonstrating the stability of BNC materials under these ORR conditions. The H_2O_2 production performance of 900-BNC was

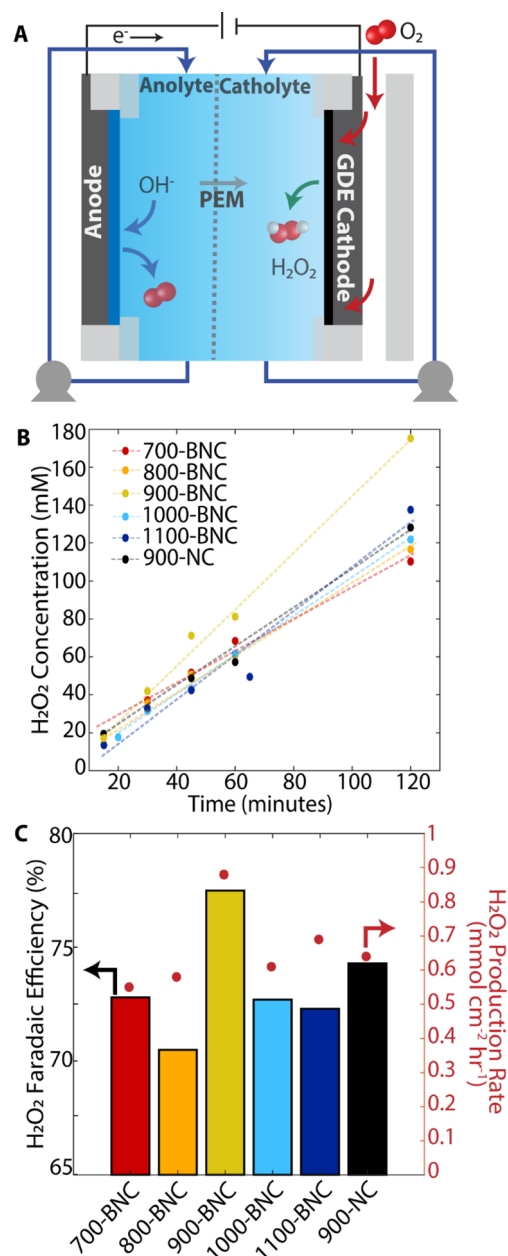


Fig. 4 Practical application of H_2O_2 production in flow cell device. (A) Schematic of electrochemical flow cell set-up. (B) H_2O_2 concentration measured over time. (C) H_2O_2 Faradaic efficiencies (right axis) and H_2O_2 production rate (left axis) for different catalysts. All measurements performed at 0.20 V vs. RHE for 2 hours in 0.5 M K_2SO_4 .

further evaluated *via* chronopotentiometric measurements, by applying constant current densities ranging from -10 to -100 mA cm^{-2} for 20 minutes (Fig. S25A†). H_2O_2 production rates (Fig. S25B†) and H_2O_2 concentrations (Fig. S25C†) increased with increasing applied current densities reaching $40 \text{ mM } H_2O_2$ and $1.6 \text{ mmol cm}^{-2} \text{ h}^{-1}$ production rate at -100 mA cm^{-2} . Notably, this high current ORR process exhibits a H_2O_2 Faradaic efficiency of 85% (Fig. S23B†), further verifying the excellent performance for practical H_2O_2 electrosynthesis.

The stability of 900-BNC was further examined under a constant current density of 100 mA cm^{-2} for 10 hours



(Fig. S26†). The measured H_2O_2 concentrations over this time suggest a slight decrease in H_2O_2 Faradaic efficiency (Fig. S26A†) and decline in H_2O_2 production rate (Fig. S26B†) from ~ 1.6 to $\sim 1.3 \text{ mmol cm}^{-2} \text{ h}^{-1}$. However, this could result from a reduction in catalytic performance or from self-decomposition of H_2O_2 into H_2O and O_2 , due to the high H_2O_2 concentrations. Interestingly, we also find that the final H_2O_2 solution pH is alkaline ($\sim \text{pH } 10$), which may exacerbate the self-decomposition of H_2O_2 leading to a reduction in measured H_2O_2 concentrations.¹⁰⁰ Despite the possible self-decomposition, the H_2O_2 produced at this current density reaches a concentration of 1290 mM (4.39 wt%), which is more than sufficient to be used for disinfection or wastewater treatment ($<1 \text{ wt\%}$ typically used for these applications).¹⁰¹

One potential drawback of this three-electrode GDE flow cell is that the H_2O_2 generated is mixed with an electrolyte, which would require additional separation steps to enable further use. To avoid the extra cost and energy of such separation steps, H_2O_2 electrosynthesis can be performed in a dual membrane electrode assembly SE electrolyzer where pure H_2O_2 solutions can be obtained as illustrated in Fig. S3.† CP measurements were performed from 10 mA to 200 mA cm^2 for 30 minutes each using the 900-BNC catalyst, as shown in Fig. S27.† At 200 mA cm^2 , H_2O_2 concentration reaches 600 mM ($\sim 2 \text{ wt\%}$), with a corresponding 72% H_2O_2 Faradaic efficiency and 2.7 mmol $\text{cm}^{-2} \text{ h}^{-1}$ production rate, further demonstrating the practical applicability of this H_2O_2 electrosynthesis technology.

4. Conclusions

In summary, we successfully synthesized a boron and nitrogen co-doped carbon catalyst that achieves high activity and selectivity for H_2O_2 production under neutral pH conditions. By tuning the pyrolysis temperature, we controlled the structural and electronic properties, chemical species, and bonding motifs of the BNC materials. Among the catalysts, 900-BNC demonstrated the highest H_2O_2 performance with 78% Faradaic efficiency and 0.9 mmol $\text{cm}^{-2} \text{ h}^{-1}$ production rate in a three electrode GDE flow cell as well as 72% H_2O_2 Faradaic efficiency and 2.7 mmol $\text{cm}^{-2} \text{ h}^{-1}$ production rate in a dual MEA SE flow cell. We attribute the superior performance of 900-BNC in part to its large electrochemically active surface area, higher degree of graphitization, and enhanced hydrophobicity. Moreover, combined XAS and XPS reveals a key factor to the enhanced H_2O_2 performance for 900-BNC may be the formation of a relatively high content of BNC moieties in the carbon matrix in the form of interfacial B–N–C bonds. An improved understanding of these catalyst material properties in neutral pH electrolyte can provide insights into the design and optimization of electrochemical systems for various applications, including energy conversion, water treatment, and chemical synthesis.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

BR contributed to the conceptualization, data curation, formal analysis, investigation, methodology, and writing of the original draft of this work. XL contributed to the collection of XRD experiment, design of electrochemical flow cell experiments, analysis, and review and editing of this work. BL contributed to materials synthesis and review and editing of this work. AD contributed to the design of electrochemical flow cell experiments and review and editing of this work. DN supported soft XAS experiments and contributed to data analysis. JMN contributed valuable insights to project conception and experimental planning. LCS contributed to the conceptualization, methodology, resources, supervision, and review and editing of this work. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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