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Tuning electrochemical performance and interfacial compatibility of oxygen electrodes in proton-conducting solid oxide electrolysis cells†

Saroj Karki, Shuanglin Zheng, Idris Temitope Bello, Allison Le and Hanping Ding 🕩



The growing demand for clean energy and the urgent need to reduce carbon emissions have accelerated the development of alternative energy solutions, with solid oxide electrochemical cells standing out owing to their efficiency in energy conversion between renewable energies and hydrogen. However, the slow reaction kinetics of its oxygen electrode, particularly at intermediate temperatures, impose a significant obstacle in optimizing their performance, reversibility, and durability. To address these challenges, this study introduces a new A-site deficient perovskite oxide as a potential electrode material for reversible protonic ceramic electrochemical cells. The cation deficiency effectively triggered the formation of oxygen vacancies and proton defects after hydration to facilitate multiple charge carrier conduction for enhanced electrode activity. After investigating the effects of cationic deficiency on structure and electrode polarization in a symmetric cell configuration of praseodymium nickel cobaltite perovskite $(Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta})$, the optimal composition was confirmed and used for integration into full cells. The electrochemical performances in fuel cell and electrolysis modes were studied, and reversible operation and short-term stability test were conducted to understand the improved behaviors, providing a pathway to facilitate excessive proton conductivity for enhanced reaction activity in oxygen electrodes.

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Introduction

The depletion of fossil fuels and global efforts to reduce carbon emissions have made it essential to seek alternative clean energy technologies to meet future energy needs. Due to the intermittent nature of solar and wind energy, hydrogen is widely considered a key energy carrier with the potential to power the world by generating electricity for driving vehicles, fueling industries, and powering homes, all with net-zero carbon emissions. In recent years, reversible protonic ceramic electrochemical cells (R-PCECs) have attracted considerable interest from academia and industry because of their transformative potential.2 R-PCECs are unique in their ability to effortlessly alternate between two operational modes: functioning as a protonic ceramic fuel cell (PCFC) for electricity generation and as a protonic ceramic electrolysis cell (PCEC) for hydrogen production.3 R-PCECs offer impressive advantages over traditional oxide ion-conducting ceramic cells, making them a compelling choice for sustainable energy systems. R-PCECs only require significantly lower activation energy for proton conduction, enabling them to operate more efficiently and produce ultra-pure hydrogen in electrolysis mode, which is

School of Aerospace and Mechanical Engineering, University of Oklahoma, Norman, OK, USA. E-mail: hding@ou.edu

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crucial for cleaner energy solutions. In fuel cell mode, they promote optimal fuel utilization, thereby maximizing system output.4-6 Moreover, to improve the energy efficiency of R-PCECs, excellent materials systems are required for achieving durable high-performance operation. Oxygen electrodes, where complicated elemental reactions occur, are considered critical components that enable faster water-splitting reactions. However, the improper use of oxygen electrodes could result in slow reaction kinetics and degradation under realistic reversible conditions.7,8

An effective oxygen electrode must exhibit high catalytic activity for the oxygen reduction reaction (ORR) in fuel cell mode and for the water oxidation reaction (WOR) in electrolysis mode. It should also have low polarization resistance (R_p) and be highly compatible with the electrolyte, ensuring lower interfacial resistance.9-11 Mixed ion-electron conducting (MIEC) cathode materials have been extensively studied because they can incorporate oxygen ions into the cathode bulk and diffuse them across the entire cathode area. 12,13 However, these electrodes are unable to conduct protons, restricting the water formation reaction to the triple-phase boundary (TPB)—the junction where gas, ions, and electrons converge (Fig. 1).14,15 Under polarization, significant accumulation of water at the TPB slows down the rate of subsequent reactions. 16 Interestingly, this limitation can be overcome by triggering proton conductivity in oxides to form a triple-conducting oxide (TCO). Theoretically, the use of a TCO electrode would significantly

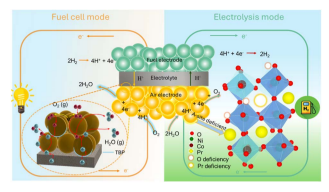


Fig. 1 Schematic of an R-PCEC with a triple-conducting oxygen electrode designed to enhance electrochemical reaction kinetics and optimize crystal structure through A-site modifications in perovskite oxides, improving charge carrier conduction properties. (Left) TCO electrode enables efficient transport of protons, oxygen ions, and electrons across the electrode layers, thereby increasing active sites for oxygen reduction and water oxidation reactions. (Right) Creation of oxygen vacancies through A-site deficiency in the crystal structure of the $Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta}$ electrode.

increase the reaction sites because the protons can move freely via surface or bulk.17

In our previous studies, we developed a series of TCO oxygen electrodes based on PrCoO3 for PCECs. Among these, the PrNi_{0.7}Co_{0.3}O_{3-δ} composition emerged as a promising candidate after structural optimization, exhibiting excellent electrochemical performance. Comprehensive characterizations unveiled the underlying mechanisms, revealing that the induced formation of oxygen vacancies facilitated effective hydration, resulting in a reasonable concentration of proton defects. Remarkably, this electrode significantly enhanced activity under dynamic operating conditions. Furthermore, a series of thermal cycling and accelerated stress tests validated its exceptional mechanical stability at the electrode/electrolyte interface, establishing PrNi_{0.7}Co_{0.3}O_{3-δ} as a benchmark oxygen electrode for PCECs.

However, further optimization of this composition was necessary to improve its performance at reduced temperatures (400-500 °C) and enhance system reliability. Lately, it was found that A-site deficiency on the lattice structure of double perovskite $(PrBa_{0.8}Ca_{0.2})_{0.95}Co_2O_{6-\delta}$ can trigger more formation of oxygen vacancies and deform the local lattice for better proton diffusion. 18 The non-stoichiometry of A-site cations can increase oxygen vacancy concentration due to charge neutrality in the lattice (Fig. 1) and it can transform the chemical and physical properties such as thermal expansion coefficient, electrical conductivity, and polarization resistance. 19-23 Contemporary density functional theory studies proved that A-site deficiency reduces vacancy formation energy and promotes charge compensation through the formation of oxygen vacancies. This structural reform also induces a downward shift in the Fermi level, resulting in ligand hole formation that boosts p-type electronic conduction.24 Investigations into the A-site-deficient $BaCo_{0.4}Fe_{0.4}Zr_{0.1}Y_{0.1}O_{3-\delta}$ cathode demonstrate that increased A-site deficiency reduces electrical conductivity due to hindered

electron and hole hopping caused by elevated oxygen vacancy concentrations. These vacancies play a pivotal role in enhancing oxygen mobility, surface exchange, and bulk diffusion. Remarkably, the area-specific resistance and activation energy progressively decrease with optimized deficiency levels, as evidenced by the superior performance of Ba_{0.90}Co_{0.4}Fe_{0.4}Zr_{0.1}- $Y_{0.1}O_{3-\delta}$ compared to stoichiometric BaCo_{0.4}Fe_{0.4}Zr_{0.1}Y_{0.1}O_{3-δ}.²⁵ These findings underscore the potential of fine-tuning A-site deficiency to significantly improve cathode catalytic activity and ORR kinetics, offering a promising strategy for advancing electrocatalyst design.

Here, we implemented state-of-the-art A-site deficiency engineering to tailor the defect chemistry and phase stability of $Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta}$ (x = 0, 0.1, 0.2, and 0.3) perovskites in PCECs. Our rigorous examination of the structural and electrochemical properties of the material series revealed that $Pr_{0.8}Ni_{0.7}Co_{0.3}O_{3-\delta}$ (PNC20) is the optimal composition. PNC20 exhibits a synergistic balance of proton transport, catalytic activity, and long-term durability. It delivers a remarkable peak power density of 0.826 W cm⁻² in fuel cell mode and an exceptional current density of 1.07 A cm⁻² at 1.3 V in electrolysis mode at 600 °C, significantly outperforming the myriad of conventional MIEC and TCO electrodes. Moreover, transient and long-term durability experiments show that PNC20 is resilient under both fuel cell (FC) and electrochemical cell (EC) modes, with steady performance over 20 cycles of voltage cycling between FC and EC modes and no detectable degradation after 100 h of continuous electrolysis operation at 1.4 V. These findings establish A-site engineering as a powerful strategy for designing next-generation R-PCEC electrodes, positioning PNC20 as a benchmark material for highly efficient and durable protonic ceramic energy conversion technologies.

Experimental

Cathode powder for symmetric and fuel cells

The $Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta}$ series electrode powders were created using a wet-chemistry method. Thermo Fisher-sourced $Pr(NO_3)_3 \cdot 6H_2O$ (99.9% purity), $Ni(NO_3)_2 \cdot 6H_2O$ (99% purity), and Co(NO₃)₂·6H₂O (99% purity) were dissolved in deionized water. Ethylenediaminetetraacetic acid (EDTA) and citric acid (CA) were utilized as complexing agents and fuels to improve combustion. The molar ratios of EDTA, citric acid, and cations were maintained at 1:1.5:1. The resultant precursor solution was repeatedly stirred with a magnetic stirrer while heated on a hot plate. The solution became gel-like as the water completely evaporated. After about 3 h of heating at 300 °C, the gel auto-ignited, releasing a considerable volume of powder ash. This ash was then combusted in a muffle furnace at 1000 °C for 5 h, yielding a crystalline perovskite.

Electrolyte powder for fuel cell

 $BaCe_{0.7}Zr_{0.1}Y_{0.1}Yb_{0.1}O_{3-\delta}$ (BCZYYb7111) electrolyte was synthesized using a sol-gel method with Ba(NO₃)₂, Ce(NO₃)₃·6H₂O, ZrO(NO₃)₂·xH₂O, Y₂O₃, and Yb₂O₃ as precursors, while CA and EDTA were employed as complexing and chelating agents,

respectively. First, Y_2O_3 and Yb_2O_3 were dissolved in a 20 wt% nitric acid solution at 120 °C, and $ZrO(NO_3)_2 \cdot xH_2O$ was dissolved in deionized water to form clear solutions. Concurrently, stoichiometric amounts of $Ba(NO_3)_2$, $Ce(NO_3)_3 \cdot 6H_2O$, and EDTA were dissolved in deionized water, and the pH was adjusted to 7 using ammonium hydroxide (NH₄OH). After achieving a pH of 7, $ZrO(NO_3)_2 \cdot xH_2O$, Y_2O_3 , and Yb_2O_3 solutions were mixed with the $Ba(NO_3)_2$ – $Ce(NO_3)_3$ -EDTA solution and CA, with the pH further adjusted to \sim 7 by adding NH₄OH. The mixture was stirred for 30 min at room temperature to promote complexation, followed by heating at 250 °C to form a gel that auto-ignited. The resulting powder was calcined at 1000 °C for 5 h in the air to produce the BCZYYb7111 electrolyte.

Anode powder for fuel cell

The anode powder consisted of sol-gel-derived BCZYYb7111, NiO, and corn starch, mixed in a weight ratio of 2:3:1.

Anode functional layer power for the fuel cell

The anode functional layer powder was prepared using a sol-gel method similar to that employed for the synthesis of the BCZYYb7111 electrolyte, with the addition of NiO. The BCZYYb7111 and NiO were mixed in a weight ratio of 2:3 to form the final composition.

Electrolyte powder for symmetric cells

The BaCe $_{0.7}$ Zr $_{0.1}$ Y $_{0.1}$ Yb $_{0.1}$ O $_{3-\delta}$ electrolyte powder for symmetric cells was produced using solid-state techniques. The precursors BaCO $_3$ (99.8% purity), ZrO $_2$ (99.7% purity), CeO $_2$ (99.5% purity), Y $_2$ O $_3$ (99.9% purity), and Yb $_2$ O $_3$ (99.9% purity), all obtained from Thermo Fisher Scientific, were thoroughly mixed by ball milling in ethanol at 300 rpm for 24 h. After drying, the mixture was calcined at 1100 °C for 10 h in an air atmosphere. The calcined powder was then ball-milled at 300 rpm in ethanol for another 12 h before being dried and calcined at 1100 °C for 10 h to yield a pure-phase electrolyte powder.

Preparation of symmetric cells

To fabricate symmetrical cells, BCZYYb7111 electrolyte pellets were formed by uniaxially pressing the powders (BCZYYb7111 + 2 wt% nickel oxide (NiO)) to obtain green pellets. The pellets were subsequently sintered at 1450 °C for 10 h to densify the product. For PNC electrode slurries, the electrode powders were ball milled with a binder solution (2% ethyl cellulose in terpineol) and ethanol for 10 min. The slurry was mixed and defoamed with a planetary centrifugal mixer (Thinky mixer) until it was thick and sticky. To produce symmetrical cells, an electrode slurry was brush-painted on both sides of the electrolyte pellet and calcined at 920 °C for 5 h to evaporate the organic solvents and binders, yielding full symmetric cells with an active area of 0.178 cm².

Preparation of half-cells

Half-cell pellets were prepared by sequential hydraulic pressing of the NiO-BCZYYb7111-starch anode, functional layer, and

electrolyte powder prepared via sol-gel method with a 12 mm die. Half-cell pellets were then sintered at 1425 °C for 10 h to obtain the dense structure. The PNC20s electrode slurry was brush-painted over the electrolyte surface with an active area of 0.178 cm² and then calcined at 920 °C for 5 h to remove organic solvents and binders from the cathode slurry and generate complete cells.

Electrochemical performance measurement

The ASR measurements of $Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta}$ series electrodes were conducted using open circuit voltage (OCV) conditions under the supply of wet oxygen ($\sim 3\%~H_2O$). The desired amount of humidity was maintained by supplying the gas through the lab-made humidifier. For the PNC20 ASR tests with varying partial pressures of water, the humidifier was maintained at different temperatures to adjust the partial pressure of water. Additionally, ASR measurements were performed under varying oxygen partial pressure by controlling the oxygen and argon (Ar) gas flow rates. The electrochemical stability of the cell was assessed at 600 °C under 3% H_2O and a current density of $-100~mA~cm^{-2}$.

During full-cell testing, the PNC20 oxygen electrode was exposed to oxygen. The mixed hydrogen (H_2 , 7 mL min⁻¹) and argon (13 mL min⁻¹) gas was fed to the testing fixture for reducing NiO to metallic nickel. The increase in the OCV was used to track the reduction process. This gas combination was gradually adjusted by increasing the hydrogen flow rate while decreasing the argon flow until the ultimate circumstances were H_2 at 20 mL min⁻¹ and no argon flow. When a steady OCV above 1.0 V was achieved, fuel cell mode testing was conducted, and current–voltage (I–V) curves were obtained between 450 °C to 600 °C. Electrochemical impedance spectroscopy (EIS) was measured at OCV, with a frequency range of 10^6 to 10^{-1} Hz and an AC signal amplitude of 20 mV.

In electrolysis testing mode, pure $\rm H_2$ was fed to the fuel electrode and humidified oxygen (20% water) was supplied to the air electrode side. The EIS experiments in electrolysis mode were performed at an applied voltage of 1.3 V with a frequency range from 10^6 to 10^{-1} Hz and an AC amplitude of 20 mV using VersaStudio software. For a comprehensive analysis of the EIS spectra, the distribution of relaxation times (DRT) was conducted with Gamry Echem analysis software.

Characterizations

The crystalline phase structure of $\Pr_{1-x}\operatorname{Ni}_{0.7}\operatorname{Co}_{0.3}\operatorname{O}_{3-\delta}$ series powders were analyzed using X-ray diffraction (XRD, Rigaku SmartLab) with a Cu rotating anode source. Characterization of the fuel cell was performed using a scanning electron microscope (TFS Quattro S) to analyze the cross-section in secondary electron and backscattered electron modes. Thermogravimetric analysis (TGA, STA 509 Jupiter® Supreme instrument) was conducted to understand the oxygen non-stoichiometry through the study of weight loss by heating the samples in the air from room temperature to 650 °C with a ramping rate of 10 °C min $^{-1}$. X-ray photoelectron spectroscopy (XPS) was

conducted using a PHI Quantera system to analyze the O 1s spectra of electrodes.

Results and discussion

Phase structure

The wet-combustion method was used to synthesize the Pr_{1-x} $Ni_{0.7}Co_{0.3}O_{3-\delta}$ series (x = 0, 0.1, 0.2, and 0.3, noted as PNC, PNC10, PNC20, and PNC30, respectively). After calcination at 1000 °C, XRD was conducted in these samples at room temperature (Fig. 2a). All the samples exhibit a single pure perovskite orthorhombic structure which is indexed to be a Phnm space group.26 No discernible impurities were observed, indicating the formation of pure crystal structures. Furthermore, the absence of impurity peaks in the XRD of a homogeneously mixed PNC series oxides and electrolyte powder (Fig. S1†) confirms their excellent chemical compatibility, ensuring a stable electrode-electrolyte interface without the formation of secondary phases or undesired reactions. Additionally, XRD analysis of the post-test PNC20 cathode confirms the stability of the material under reversible operation, with no detectable impurity phase indicating the material's structural integrity.

When these electrodes are integrated into the symmetric cells, the electrode polarization resistance can be measured based on the difference between the high and low-frequency intercepts on the real axis to study the catalytic activity towards WOR and ORR.27 Fig. 2b depicts the area-specific resistance (ASR) of each composition measured at the OCV condition. As the A-site deficiency increases, ASR continuously drops until x = 0.2 across all intermediate temperature measurements. Beyond this point, an additional deficit causes an increase in ASR. At 600 °C, PNC20 has a R_p of 2.53 Ω cm², which is 24%, 13.4%, and 9% lower than PNC, PNC10, and PNC30, respectively. Also, at 400 °C, PNC20 stays to outperform the other compositions, with R_p values 16.8%, 6.65%, and 2.7% lower, respectively. This result indicates that introducing Pr deficiency within a definite range (here, x = 0.2) improves catalytic activity by uplifting the formation of oxygen vacancies,

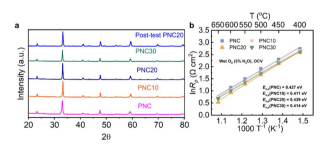


Fig. 2 Structural analysis of $Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta}$ and its impact on electrode polarization resistance. (a) XRD patterns $Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-\delta}$ (x = 0.1, 0.2, and 0.3), along with post-test PNC20 samples, showing phase purity and structural variations as a function of Pr-site deficiency. (b) Arrhenius plot of electrode polarization resistance from symmetric cells under open-circuit conditions in wet oxygen (\sim 3% H₂O), demonstrating temperature-dependent changes in polarization resistance.

improving oxygen dissociation and adsorption on the cathode surface, and facilitating ORR. However, further Pr deficiency can adversely reduce electrode activity. Thus, PNC20 demonstrates the lowest resistance across the entire intermediate temperature range, underscoring its superior electrochemical performance. Furthermore, the activation energies of all the cathode compositions are calculated by using the following equation:26

$$R_{\rm p} = R_{\rm p}^{\rm o} \times \left(\frac{E_{\rm a}}{K_{\rm B}T}\right) \tag{1}$$

where $K_{\rm B}$ is the Boltzmann constant 1.38 \times 10⁻²³ J K⁻¹; $E_{\rm a}$ is the activation energy; T is the temperature.

The calculated activation energy is 0.427 eV for PNC, 0.411 eV for PNC10, 0.439 eV for PNC20, and 0.414 eV for PNC30. The alterations in activation energies of all the compositions are insignificant with variation of ± 0.013 . The lower R_p values of PNC20 across the temperature spectrum accentuate its efficient catalytic activity of charge transfer kinetics and reduced operational losses.

Electrochemical performance of the symmetric cell

The electrochemical performance of the PNC20 electrode was evaluated utilizing EIS on symmetric cells. Fig. 3a shows the EIS measurements carried out at different applied voltages, demonstrating a distinct reduction in ASR with increasing voltage. The ASR value was measured to be 0.118 Ω cm² at an applied voltage of 0.7 V, while it sharply declined to 0.01 Ω cm² at 1.2 V. This substantial reduction in ASR values shows that the ORR becomes increasingly kinetically favorable at higher potentials, thereby reducing the polarization resistance. Moreover, EIS measurements were also performed under varying water partial pressures (Fig. 3b). At 3%, 10%, 20%, and 30% H_2O , the ASR values were 0.47, 0.36, 0.17, and 0.12 Ω cm², respectively. The dramatic reduction in ASR with increasing

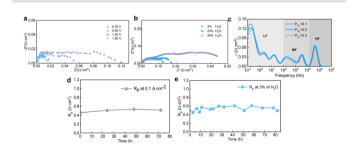


Fig. 3 Electrochemical performance of symmetric cells at 600 °C under varying operational conditions. (a) EIS measurements under different applied voltages, highlighting voltage-dependent impedance responses. (b) EIS measurements at varying partial pressures of water, showing the effect of water content on cell performance. (c) Distribution of relaxation time analysis at different oxygen partial pressures, revealing oxygen-dependent reaction mechanisms. (d) Polarization resistance stability over time at 3% water content, demonstrating longterm cell durability. (e) Stability of polarization resistance as a function of applied current density (0.1 A cm⁻²) at 3% water content, showcasing the cell's performance under operational loads.

water partial pressure indicates that hydration at the cathode in a humidified atmosphere facilitates the formation of protonic defects through the hydration of oxygen vacancies described by the Stotz–Wagner mechanism (eqn (2)), thereby enhancing the proton conduction, improving charge transfer and overall electrochemical performance.²⁸

$$H_2O + V_O^* + O_O^* \rightleftharpoons 2OH_O^*$$
 (2)

The increase in oxygen vacancy concentration in A-site deficient PNC20 was further confirmed by XPS analysis. As illustrated in Fig. S2,† the O 1s XPS spectra of the samples were deconvoluted into four distinct sub-peaks located at approximately 528, 529, 531, and 533 eV. These peaks correspond to lattice oxygen (O²⁻), highly oxidative oxygen (O₂²⁻/O⁻), adsorbed oxygen (OH-/O2), and adsorbed H2O, respectively. The higher percentage of ${\rm O_2}^{2-}/{\rm O}^-$ species in PNC20 compared to PNC implies higher oxygen vacancy concentration which results in high proton uptake during hydration reaction when exposed to a wet atmosphere.²⁹⁻³³ The oxygen non-stoichiometry between PNC and PNC20 was also revealed via a thermogravimetric analysis test (TGA). As depicted in Fig. S3,† PNC20 experiences a more significant weight loss than PNC. This pronounced weight reduction directly correlates with the increased concentration of oxygen vacancies in the PNC20 electrode. 18,34 This finding corroborates our hypothesis that the introduction of A-site deficiency in PrNi_{0.7}Co_{0.3}O_{3-δ} promotes the formation of oxygen vacancies, which can subsequently facilitate proton conduction and enhance the electrochemical performance of the electrode.

Additionally, EIS measurements were conducted by changing oxygen partial pressures to better distinguish between ORR and the WOR occurring at the PNC20 electrode. DRT analysis was then used to extract particular electrochemical subprocesses. To identify these specific processes, the distinctive frequency range of the EIS spectra $(10^6-10^{-1} \text{ Hz})$ was used, as illustrated in Fig. 3c. The low-frequency (LF, 10^{-1} – 10^2 Hz) peaks are generally related to gas diffusion or surface adsorption/desorption phenomena; the mid-frequency (MF, 10²–10⁴ Hz) peaks are associated with bulk diffusion processes and surface oxygen exchange; and the high-frequency (HF, 10⁴– 10⁶ Hz) peaks are associated with charge transfer reactions.^{34–38} The area under the peaks in each frequency range correlates with the polarization resistance of the respective electrochemical step. It is evident from Fig. 3c that the MF and LF regions contribute the most to the total polarization resistance under different oxygen partial pressures, suggesting that surface exchange, gas diffusion, and bulk diffusion processes dominate the electrode's polarization resistance.³⁹ The MF and LF resistances decrease upon introducing oxygen, indicating improved surface mass transfer and bulk ion transport contributing to the acceleration of the ORR kinetics. 40,41

The long-term electrochemical stability of the PNC20 electrode was evaluated by monitoring the R_p over time at a current density of 0.1 A cm⁻² and 3% H_2O (Fig. 3d and e). Remarkably, no significant decrease was observed in ASR after 82 h of operation in 3% H_2O . Furthermore, even after 72 h at

a sustained current density of 100 mA cm⁻², the ASR exhibited only a modest increase of 12.45%, underscoring the electrode's robust performance stability.

Electrochemical performance of the reversible protonic ceramic cell

The enhanced electrochemical characteristics of PNC20 as a cathode material for reversible protonic ceramic cells were further validated by single-cell performance testing. In fuel cell mode, a cell with a PNC20 cathode and an electrolyte of BCZYYb7111 was assessed at temperatures between 450 °C and 600 °C. Fig. 4a demonstrates the temperature-dependent I-V-P characteristics of the cell in FC mode. The cell achieved a peak power density of 826 mW cm⁻² at 600 °C, 522 mW cm⁻² at 550 ° C, 402 mW cm⁻² at 500 °C, and 260 mW cm⁻² at 450 °C, surpassing the performance of the same single cell configuration with PNC as air electrode (Fig. S4†) which underscores PNC20's exceptional performance across a range of operating conditions. This further establishes PNC20's enhanced electrocatalytic ORR, and positions it amongst the highestperforming electrodes reported in the literature. 42-50 EIS results further support the improved ORR activity of the PNC20 electrode (Fig. 4b). At 600 °C, 550 °C, and 500 °C, the electrolyte resistance (R_0) values were 0.237, 0.313, and 0.4 Ω cm², respectively, whereas the R_p was 0.144, 0.228, and 0.4 Ω cm², respectively. It is conspicuous that electrolyte resistance largely controls the total cell resistance across a considerable portion of the intermediate temperature range, which is principally triggered by the electrolyte's conductivity and thickness.

To gain a more comprehensive understanding of the polarization mechanisms, DRT analysis of EIS was conducted to assess the individual electrochemical subprocesses. As illustrated in Fig. 4c, four distinct peaks, denoted as P_1 , P_2 , P_2 , and P_4 , emerge when moving from high frequency to low frequency. The LF ($<10^2$ Hz) peaks predominantly reflect gas diffusion limitations and surface adsorption/desorption dynamics. MF

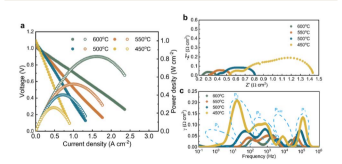


Fig. 4 Electrochemical electrolysis performance evaluations under different temperatures. (a) I-V-P characteristics in FC mode at 450 °C, 500 °C, 550 °C, and 600 °C, demonstrating the temperature-dependency of power density and current density. (b) Electrochemical impedance spectra measured at different temperatures to study respective contribution from each cell component. (c) Distribution of relaxation time analysis based on ElS data, providing insight into the frequency-dependent polarization resistances and reaction kinetics under FC mode operation.

(10²-10⁴ Hz) features are indicative of bulk ion transport and surface oxygen exchange kinetics. Meanwhile, HF (10⁴–10⁶ Hz) peaks capture the rapid charge transfer events at the electrodeelectrolyte interface.51 Conspicuously, the peak intensity exhibits a pronounced decline with increasing temperature. Given that the area under these peaks in DRT analysis directly correlates with the polarization resistance of individual electrochemical processes, this trend unequivocally indicates a lower polarization resistance at 600 °C compared to 450 °C, where the peaks are more intense. The high-frequency peak (P_1) shifts towards lower frequencies, while the peaks in the MF and LF ranges (P2, P3, and P4) shift towards higher frequencies as the temperature increases. Below 550 °C, the appearance of an additional peak (Padd) in the mid-frequency region, associated with bulk ion transport and surface oxygen exchange, signifies the transition of a previously non-rate-limiting step into a ratelimiting one, introducing an additional energy barrier that must be overcome at this temperature.⁵² The figure further reveals that the LF peak (P_3) and MF peak (P_2) dominate as temperature decreases, with P3 being the most pronounced. This suggests that the oxygen adsorption and dissociation processes represent key rate-limiting steps.³⁹ Hence, further optimization of the cathode, potentially through surface modification, could substantially enhance performance and lead to higher power outputs and performances.

Complementing the above discussions, transient voltage experiments on the PNC20 fuel cell were conducted to evaluate its resilience under operating stress. As seen in Fig. 5a, no discernible deterioration of the cell current density was observed over 100 cycles when the applied voltage was swept between 0.5-0.8 V. However, the cell showed a modest decrease in current density under lower voltages, which was probably caused by temporary oxygen starvation.53 Nevertheless, subsequent EIS data (Fig. 5b) demonstrated a reduction in polarization resistance following the transient voltage test, indicating

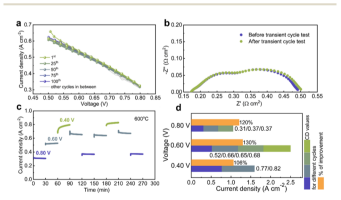


Fig. 5 Comprehensive analysis of cell performance under transient voltammetry cycling. (a) Transient voltammetry analysis conducted over 100 cycles, examining the stability and degradation of the cell's electrochemical performance. (b) EIS spectra recorded before and after the transient voltammetry test, highlighting changes in impedance and reaction kinetics. (c) Current density response under stepwise voltage increase and decrease in the number of voltage cycles, assessing the dynamic response of the cell. (d) Bar graph depicting the trend of current density after each cycle, exhibiting the cumulative effect of cycling on electrochemical performance.

that oxygen starvation did not cause irreversible damage or compromise the long-term stability or functionality of the cell.

Dynamic gains in current density were found by conducting a stepwise voltage test (Fig. 5c), which involved cycling the applied potential from 0.8 V to 0.6 V and then to 0.4 V at 30minute intervals over two cycles at a fixed temperature of 600 ° C. For example, as depicted in Fig. 5d, the current density increased by 20% from 0.31 A cm⁻² in the first step to 0.372 A cm⁻² in the third step cycle at 0.8 V. Comparable patterns were noted at 0.6 V and 0.4 V, where the current density increased by 6% and 30%, respectively, following two and four voltage steps.

The performance of the R-PCEC was further assessed in the electrolysis mode. The cell's current-voltage (I-V) characteristics, recorded at temperatures between 450 °C and 600 °C, are shown in Fig. 6a where the fuel electrode was exposed to dry hydrogen and the air electrode was supplied with humidified air (20% H₂O). At the applied cell voltage of 1.3 V, current densities of 1.07, 0.656, 0.376, and 0.184 A cm⁻² were achieved at 600, 550, 500, and 450 °C, respectively. These outcomes pinpoint the substantial activity of the PNC20 cathode toward the oxygen evolution reaction and the WOR in the intermediate temperature range.

Impedance spectroscopy (Fig. 6b) further revealed that the total cell resistance is predominantly governed by the electrolyte resistance, which is influenced by the electrolyte's thickness and conductivity. The R_0 values were determined to be 0.216, 0.297, 0.400, and 0.550 Ω cm² at 600 °C, 550 °C, 500 °C, and 450 °C, respectively, exhibiting an increasing trend of R_0 with decreasing temperature. Alternatively, the R_p , largely contributed by the electrode at a bias of 1.3 V, was determined to be 0.048, 0.1, 0.239, and 0.548 Ω cm² across the same temperature spectrum. These results prove that R_p is less dominant in the overall resistance, suggesting that reducing electrolyte thickness could significantly lower total resistance, thereby enhancing performance under electrolysis conditions.

The reversible operation of the R-PCEC between electricity generation and hydrogen production was validated through transient testing conducted over a 20-cycle period, with a rapid voltage transition between the electrolysis (1.4 V) and fuel cell (0.7 V) modes. In Fig. 7a, no discernible degradation was seen in the current density during these cycles in both modes, demonstrating the cell's exceptional dynamic response to

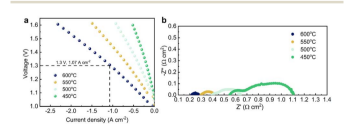


Fig. 6 Electrochemical electrolysis performance evaluations under different temperatures. (a) I-V characteristics in EC mode at 450 °C, 500 °C, 550 °C, and 600 °C, demonstrating the temperature-dependency of current density. (b) Electrochemical impedance spectra measured at different temperatures to study the respective contribution from each cell component.

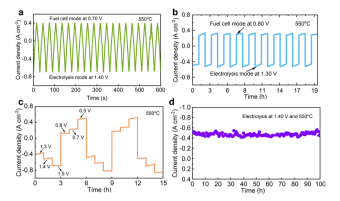


Fig. 7 Reversible operation and stability assessment in reversible operation modes for the as-prepared electrochemical cells. (a-c) Current density responses during reversible operation, illustrating the transient behavior as the working mode alternates between EC and FC modes. (d) Short-term stability evaluation in electrolysis mode upon constant applied voltage of 1.40 V.

varying operating conditions. Additionally, the cycling experiment was conducted at 550 °C by alternating between fuel cell mode (0.8 V) and electrolysis mode (1.3 V), with each mode lasting for one hour over a total period of 20 hours (Fig. 7b). The cell exhibited stable performance throughout the test, confirming its high reversibility.

Moreover, the cell was tested under cycles of different electrolysis voltages (1.3 V, 1.4 V, and 1.5 V) and fuel cell voltages (0.8 V, 0.7 V, and 0.5 V) for 15 hours as depicted in Fig. 7c. Stable current density was witnessed in fuel cell mode, while an increasing trend in electrolysis mode was perceived. Further, The R-PCEC was measured through 100 h of continuous operation at 1.4 V in electrolysis mode at 550 °C to assess its longterm durability Fig. 7d. It showed sustained performance with only 0.017% degradation per hour, highlighting its robustness and potential for use in reversible systems.

After the electrochemical testing, scanning electron microscopy (SEM) analysis was performed to investigate the interface between the PNC20 electrode and the electrolyte membrane (Fig. 8a). The PNC20 electrode exhibited strong adhesion to the electrolyte and maintained a porous structure, which facilitates efficient gas diffusion across the entire electrode surface. The

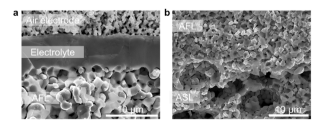


Fig. 8 Microstructural analysis of the post-test cell after electrochemical measurements using SEM. (a and b) Scanning electron microscopy (SEM) images revealing the microstructural features of the cell after testing, focusing on changes in morphology, surface integrity, and possible degradation of materials resulting from operational stress

absence of visible cracks, delamination, or other structural defects highlights the excellent thermal compatibility between PNC20 and the electrolyte, enhancing the mechanical stability and durability of the system.

Furthermore, the 8 µm-thick electrolyte membrane exhibits a dense and defect-free morphology, devoid of open pores or pinholes, effectively preventing fuel crossover and ensuring optimal electrochemical performance. As depicted in Fig. 8b, the post-test SEM analysis highlights the microstructural integrity of both the anode functional layer (AFL) and the anode support layer (ASL). Notably, the AFL, measuring approximately 10 μm, undergoes complete reduction to metallic nickel along with ASL, significantly enhances proton transport. The presence of this AFL intermediate layer plays a crucial role in minimizing resistance associated with electrochemical proton incorporation at the anode-electrolyte-gas triple-phase boundaries, thereby facilitating more efficient charge transfer and boosting overall cell performance.54,55 Moreover, the fully reduced nickel phase within the anode is complemented by a well-maintained porosity, ensuring efficient hydrogen diffusion pathways essential for sustained and high-efficiency operation.

Conclusions

The introduction of A-site Pr deficiency in Pr_{1-x}Ni_{0.7}Co_{0.3}O_{3-δ} as a TCO electrode offers remarkable improvements for R-PCECs. The PNC20 electrode showed exceptional electrochemical performance in both FC and EC modes, making it an excellent candidate for operation at reduced temperatures. The efficiency of catalyzing ORR and WOR contributes to reduced polarization resistance, leading to higher power densities and long-term durability.

The improved electrochemical performances of the PNC20 electrode in R-PCEC are attributed to several key findings. In fuel cell mode, PNC20 achieves a peak power density of 826 mW cm⁻² at 600 °C, surpassing many previously reported values in the literature. Similarly, the current density reaches a high current density of 1.07 A cm⁻² at 1.3 V in the electrolysis mode. This is because the formation of additional oxygen vacancies induced by Pr deficiency can enhance both surface oxygen exchange and bulk diffusion processes. These oxygen vacancies enable faster ORR kinetics and improve overall charge transfer, making PNC20 the optimal candidate among the tested compositions. DRT analysis of EIS spectra in the intermediate temperature range identifies oxygen adsorption and dissociation as critical rate-limiting processes. This underscores the potential of advanced surface modifications to significantly boost electrochemical performance and achieve greater power output, presenting a promising pathway for further optimization. The long-term durability of PNC20 is validated through transient voltage cycling and reversible operations, where no significant degradation in performance is observed. The electrode's stability across various humidified and oxygen-rich environments emphasizes its robustness for R-PCEC. Additionally, stepwise voltage tests indicate dynamic gains in current density, demonstrating its resilience under fluctuating operational conditions.

In summary, the incorporation of A-site deficiency in Pr_{1-x} Ni_{0.7}Co_{0.3}O_{3- δ} (x = 0.2) significantly improves the performance, making PNC20 a promising electrode for the next generation R-PCECs. The material's high activity and excellent stability highlight its potential for advancing this technology.

Data availability

All the relevant data are included in the manuscript.

Conflicts of interest

There are no conflicts to declare.

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