

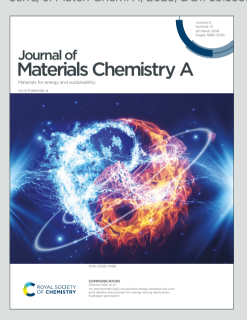
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Maximizing Oxygen Permeation via Catalytic Functionalization under Oxyfuel Conditions

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Abstract

Oxygen transport membranes (OTMs) offer a promising route for high-efficiency, cost-effective oxygen supply in energy and chemical industries, with the potential to significantly reduce CO₂ and NO_x emissions when integrated into oxy-combustion processes. However, conventional OTMs suffer from poor chemical stability in CO₂-rich environments, prompting the development of dual-phase membranes that, while more stable, typically exhibit lower oxygen permeation rates. In this study, we address this limitation by enhancing the surface exchange kinetics of Fe₂NiO₄-Ce_{0.8}Tb_{0.2}O_{2-δ} (NFO-CTO) membranes by surface modification with various oxygen oxidation-reduction reaction (OORR) catalysts, including Ce, Pr, Sm, Tb, Co, Nb, Zr, and Al oxides, and Pr-based binary oxides. Comprehensive characterization using electrochemical impedance spectroscopy, oxygen isotopic exchange, and gas permeation measurements revealed a substantial improvement in surface reaction kinetics. Catalyst activation led to a six-fold increase in oxygen flux under standard conditions and up to a 2.5-fold enhancement under harsh environments containing CO₂ and SO₂ at 850 °C, mimicking oxyfuel combustion conditions. This work demonstrates that rational catalyst selection and integration can overcome fundamental surface limitations in dual-phase membranes, offering a viable strategy to advance oxygen separation technologies for sustainable energy applications.

1. Introduction

Power generation and industry sectors account for more than 42% of global CO₂ emissions. In 2023, global CO₂ emissions from these sectors reached around 34.5 Gt (GtCO₂), with a year-on-year increase of about 0.9%. Although this is a slight improvement from previous years, the upward trend continues due to persistent demand for energy, particularly in regions like Asia.[1–3] Amongst the existing technologies, oxyfuel combustion (fuel combustion with pure O₂) is a leading option for implementing carbon capture and storage (CCS) strategies. This process enhances CO₂ capture efficiency by producing a flue gas that consists mostly of CO₂ and water, making CO₂ separation easier.[4,5] With regard to the O₂ supply, the use of Oxygen Transport Membrane (OTM) modules offers an alternative to traditional cryogenic O₂ production,[6] permitting its integration into most existing medium-to-small scale installations as a promising and cost-effective solution.[7–11]

In recent years, significant advances have been made in producing CO₂ and SO₂-stable mixed ionic-electronic conducting (MIEC) materials for oxyfuel applications.[12–16] One of the most promising approaches is the use of dual-phase composites that integrate an electronic conductor

 $Ce_{0.8}Gd_{0.2}O_{2-\delta}$ – $FeCo_2O$ iew Article Online with conductor, such as $Fe_2NiO_4-Ce_{0.8}Tb_{0.2}O_{2-\delta}$ $Ce_{0.8}Gd_{0.15}Cu_{0.05}O_{2-\delta}-SrFeO_{3-\delta}, \quad Ce_{0.85}Gd_{0.1}Cu_{0.05}O_{2-\delta}-La_{0.6}Ca_{0.4}FeO_{3-\delta}. \quad Particularly, \quad the \quad 60\%$ Fe₂NiO₄ – 40% Ce_{0.8}Tb_{0.2}O_{2.δ} (NFO-CTO) system[17–20] presented outstanding stability and high performance. However, the obtained oxygen permeability should be further improved to meet the techno-economic target of 10 mL·min⁻¹·cm⁻².[18] To approach this goal, membrane thickness can be reduced; for instance, high O₂ permeation was achieved under oxyfuel conditions using a 10 µm-thick NFO-CTO membrane.[17] Since reducing thickness increases the influence of surface exchange reactions on oxygen permeability, additional surface modifications can be employed to further enhance oxygen flux, particularly when surface exchange becomes the ratelimiting step.[21,22] This limitation is particularly significant at lower temperatures or when the membrane thickness is reduced.[23] One common approach to improving surface exchange reactions and increasing the oxygen flux is the addition of porous layers to the membrane surface, consisting of the same dual-phase material with the same or different ratios, or a new material.[24-26] These modifications primarily boost oxygen permeation by increasing the surface-specific area, providing more active sites, and extending the triple phase boundary (TPB) where the oxygen oxidation/reduction reaction (OORR) takes place. However, the performance of these systems remains limited by the reaction kinetics involved in the adsorption, dissociation, and recombination of oxygen. Adding active elements with redox properties and strong adsorption capabilities can enhance these surface exchange reactions, improving O₂ permeation. Several lanthanide and metal oxides are candidates that could meet these criteria.[27-29] In this work, a screening study was carried out to evaluate the catalytic activation of porous NFO-CTO layers with different single-element oxides and Pr-based binary formulations. These elements were selected based on their reported ability to enhance surface oxygen exchange kinetics (e.g., Ce, Pr, Sm, Tb via high oxygen mobility and redox activity), provide electronic conductivity and catalytic sites (e.g., Co), or improve structural and chemical stability under reactive environments (e.g., Nb, Zr, Al). The catalytic performance was assessed by electrochemical impedance spectroscopy and isotopic exchange to quantify improvements in surface exchange kinetics and oxygen ion transport. These kinetic parameters were then correlated with oxygen permeation measurements to establish structure-property relationships and identify the most effective catalytic promoters.

2. Material and methods

2.1. Materials and membranes fabrication

NFO-CTO (60% Fe₂NiO₄ - 40% Ce_{0.8}Tb_{0.2}O₂₋₈) composite powders were synthesized by a one-pot Pechini method To this end, stoichiometric amounts of metal nitrates (Ce(NO₃)₃·6H₂O, and Fe(NO₃)₃·9H₂O provided by Sigma Aldrich, Tb(NO₃)₃·6H₂O, and Ni(NO₃)₃·6H₂O by ABCR GmbH) were dissolved in distilled water. Subsequently, citric acid (Sigma Aldrich) and ethylene glycol were added as chelating and gelating agents in a molar ratio of metal:citric acid:ethylene glycol 1:2:4. After drying up to 270 °C, the obtained gel was fired at 600 °C to obtain finely dispersed powders with the correct crystalline phases. Dense NFO-CTO ceramics were obtained by uniaxial pressing, followed by sintering of the obtained disks in air at 1400 °C for 10 h (2 °C min⁻¹ heating rate, 5 °C min⁻¹ cooling rate). For oxygen permeation measurements, membranes with a diameter of 15 mm and a thickness of 0.6 mm were obtained from the sintered NFO-CTO disks after grinding and polishing with sandpaper.

Ce_{0.8}Gd_{0.2}O_{1.9} (CGO) dense electrolytes were selected for electrochemical impedance Article Online spectroscopy (EIS) studies. The CGO powders were synthesized using the co-precipitation method and co-doped with cobalt to improve sinterability. [30,31] This was done by impregnating a 2% molar of Co(NO₃)₂·6H₂O dissolved in deionized water. The obtained powders were calcined in air at 800 °C for 5 h. Dense disks (CGO-Co) were obtained by uniaxial pressing, followed by sintering in air at 1480 °C for 4 h (2 °C min⁻¹ heating rate, 5 °C min⁻¹ cooling rate). The densities of all samples, checked by Archimedes' method and SEM, exceeded 98%.

For EIS and oxygen permeation measurements, 30 µm-thick porous layers of NFO-CTO were screen-printed on both sides of CGO electrolyte disks and NFO-CTO membranes, respectively. Inks for screen-printing were made by mixing NFO-CTO and terpineol in a 1:2 weight ratio in a three-roll mill. After deposition, the obtained samples were sintered in air at 1100 °C for 2 h. Catalytic activation of the porous NFO-CTO scaffold layers was accomplished by infiltration of well-dispersed catalyst nanoparticles. To this end, 2 M solutions of the element precursors Ce, Pr, Sm, Tb, Co, Nb, Zr, and Al oxides and Pr-based binary formulations (typically nitrates) were prepared in a 1:1 ethanol-water volume ratio and dripped onto the porous composite scaffold layers. [32] After infiltration by capillary force, the samples were dried at 80 °C for 1 h and calcined in air at 850 °C for 2 h.

NFO-CTO composite powders for Pulse Isotopic Exchange (PIE) measurements were obtained by crushing dense sintered samples, and sieving the obtained powder through a 100 µm mesh. These composite powders were activated with catalyst nanoparticles by incipient wetness impregnation using the abovementioned procedure. Ce, Pr, Al, Ce-Pr, and Pr-Al cations were selected as representative catalysts from the screening results from the EIS study.

2.2. Membrane characterization

The crystalline phases of the prepared samples were identified by using a PANalytical Cubix fast diffractometer with $CuK\alpha1$ radiation ($\lambda=1.5406$ Å) and an X'Celerator detector, operating in Bragg-Brentano geometry. X-ray diffraction (XRD) patterns recorded in the 2θ range from 25° to 90° were analyzed using X'Pert Highscore Plus software. Cross-sectional analysis of the sintered samples, both before and after the tests, was conducted by performing scanning electron microscopy (SEM) and energy dispersive X-ray (EDX) characterization using a ZEISS Ultra55 field emission scanning electron microscope. Additionally, backscattered detector (BSD) imaging was used to provide compositional contrast for distinguishing grains and elemental distributions.

PIE measurements were conducted using a homemade setup described elsewhere.[33] The powder samples were loaded in the centre of a quartz tubular micro-reactor with an inner diameter of 2 mm. Quartz wool plugs were used to secure the packed powder bed. The length and mass of the packed bed were typically 10 mm and 0.08 g, respectively. $^{16}O_2$ mixed with Ar was used as carrier gas and fed through the reactor with a total flow rate of 30 ml min⁻¹ (NTP). Gases were dried using Agilent gas clean moisture filters before entering the reactor. Oxygen isotope gas was purchased from Cambridge Isotope Laboratories, Inc. (> 97 atom% $^{18}O_2$). A six-port valve with a 500 μ l sample loop was used for injection of the $^{18}O_2/N_2$ pulse into the $^{16}O_2/Ar$ carrier gas, with the pulse having the same pO_2 as the carrier gas. The diluent N_2 in the pulse was used for internal calibration of the mass spectrometer (Omni Star TM GSD 301 Pfeiffer-Vacuum). The mean residence time of the reactor varied between 10 and 30 ms, depending on temperature. Prior to measurements, the samples were pre-treated at 850 °C for 2 h to remove possibly adsorbed water and CO_2 , then cooled to 50 °C at a rate of 5 °C min⁻¹. Measurements were performed at a pO_2 of

0.21 atm. Averaged values of the $^{18}O_2$ and $^{16}O^{18}O$ effluent fractions in three pulse experiments. An Article Online a given temperature, were used to calculate the exchange rate. The reactor was equilibrated prior to each measurement before data acquisition. The overall surface exchange rate, \mathfrak{R}_0 [mol (O) m⁻² s⁻¹], was calculated from:

$$\Re_{0} = \frac{2F_{\rm m}}{S} \ln \left(\frac{f_{\rm i}^{18}}{f_{\rm e}^{18}} \right) \tag{1}$$

Where f_i^{18} and f_e^{18} are the 18 O isotope fractions in the pulse at the reactor's inlet and outlet, respectively, F_m is the molar flow rate of oxygen through the packed bed, and S is the total surface area of the sample. The fraction f^{18} was calculated from:

$$f^{18} = 0.5f^{34} + f^{36} \tag{2}$$

Symmetrical NFO-CTO/CGO/NFO-CTO cells for impedance measurements were prepared by coating 30 µm-thick porous NFO-CTO layers, with and without catalyst-impregnation, onto both sides of 0.8 mm-thick CGO-Co electrolyte disks. Measurements were performed with an excitation voltage of 20 mV in the frequency range $0.01 - 3 \cdot 10^5$ Hz, using a Solartron 1470E multichannel potentiostat and a 1455A frequency response analyzer. Measurements were performed at 850 °C, under different atmospheres (air, 5% O_2 in N_2 , 5% O_2 in CO_2 , and 250 ppm in 5% O_2 in CO_2). In all cases, the total flow remained constant (100 ml min⁻¹). The impedance spectra were analysed using $ZView2^{\circ}$ software. The impedance spectra were fitted using the equivalent circuit LR_e (R_1Q_1)(R_2Q_2)(R_3Q_3), where L is an inductance, R_e the apparent ohmic resistance, R_i (i = 1,2,3) the different contributions of the total polarization resistance (R_P), and Q_i (i = 1,2,3) the corresponding constant phase element. Table 1 presents the various test environments and durations analyzed using electrochemical impedance spectroscopy (EIS).

Table 1: EIS test conditions studied for the different symmetrical NFO-CTO/CGO/NFO-CTO cells.

	Gas composition	Duration
Feed side	$21\% O_2$ in N_2	90 min
Sweep side (clean conditions)	$5\% O_2$ in N_2	90 min
Sweep side (oxyfuel conditions)	$5\% O_2$ in CO_2	90 min
Sweep side (oxyfuel conditions)	250 ppm SO ₂ , 5% O ₂ in CO ₂	250 min
Sweep side (recovery, clean conditions)	$5\% O_2$ in N_2	10 h

Permeation tests were performed on a gastight NFO-CTO composite disk sintered at 1400 °C for 10 h, resulting in a final membrane thickness of 0.6 mm. NFO-CTO membranes were screen-printed on both sides with a 30 μm-thick porous NFO-CTO layer. After deposition and drying, the layers were calcined at 1100 °C in air for 2 h. The permeation tests were conducted in a lab-scale reactor consisting of a cylindrical two chambers quartz reactor, in which synthetic air (21%, vol. O₂) was fed into the oxygen-rich chamber, while Ar, 30% vol. CO₂ in Ar and 250 ppm SO₂ in 30% vol. CO₂ in Ar were used as sweep gases on the permeate side chamber, in a 4-end mode configuration. Both streams were fed at atmospheric pressure. Inlet gases were preheated in order to ensure proper contact temperature with the membrane surface. This is particularly important when high gas flow rates are employed. All streams were individually controlled by mass flow controllers. The temperature was measured by a thermocouple attached to the membrane.

Membrane gas leak-free conditions were achieved using rings of tailored alloys. The permeatew Article Online was analyzed at steady state by online gas chromatography using a micro-GC Varian CP-4900 equipped with Molsieve5A, Pora-Plot-Q glass capillary, and CP-Sil modules. Leak-tight sealing and membrane integrity were ensured by continuously monitoring the N₂ concentration in the product gas stream. Sealing was considered acceptable when the ratio between the oxygen flow leak and the oxygen flux remained lower than 3%. The reported data were collected at steady state after at least 1 h in the reaction stream. Each test was repeated three times, yielding an analytical error below 0.5%.

3. Results and discussion

3.1. Structural characterization

XRD patterns of the impregnated NFO-CTO catalytic powders are presented in the Supporting Information (SI). These powders were subjected to the same thermal cycling and catalyst loading as the infiltrated porous NFO-CTO backbones used in electrochemical impedance spectroscopy, oxygen isotopic exchange, and gas permeation measurements.

Peaks associated with the NFO-CTO phase, along with peaks associated with the different oxide catalysts, are visible in the XRD patterns. Fig. S1 shows characteristic peaks for ZrO₂, Co₃O₄, Tb₇O₁₂, Sm₂O₃, and Nb₂O₃, while Fig. S2 shows peaks for CeO₂, Pr₆O₁₁, Al₂O₃, Pr-Al, Ce-Pr. In the particular cases of Al, Ce, and Ce-Pr catalysts, it was not possible to detect the corresponding oxide catalyst phase(s). On the one hand, the resulting catalyst oxide particles could be CeO₂ and CeO₂-PrO₂, masked by the CTO fluorite phase from the backbone, or the concentration of catalyst nanoparticles being below the detection limit of XRD.

Scanning Electron Microscopy (SEM) cross-sectional images of the NFO-CTO backbone and all the infiltrated NFO-CTO backbones (Co, Sm, Al, Nb, Tb, Zr, Ce, Pr, Ce-Pr, and Pr-Al) after calcination in air at 850 °C are shown in Fig. 1. A good dispersion of the composite particles in the backbone is easily identifiable in the backscattered detector (BSD) image, with NFO appearing as dark grey and CTO as light grey, indicating a well-distributed composition and rounded particle morphology. Furthermore, nano-sized catalyst particles showed different particle morphologies with particle sizes below 100 nm, depending on the specific infiltrated metal oxide. Most catalyst particles show spherical morphology, whereas Pr₆O₁₁ and Al₂O₃ show needle- and sheet-like morphologies, respectively.

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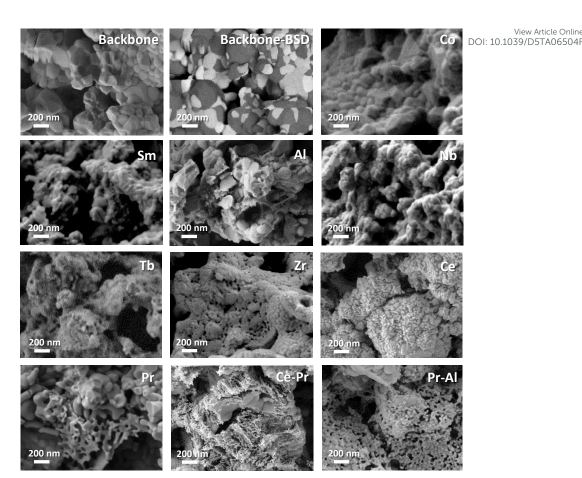


Fig. 1. SEM and BSD cross-sectional images of a pure NFO-CTO backbone and infiltrated (Co, Sm, Al, Nb, Tb, Zr, Ce, Pr, Ce-Pr, and Pr-Al) NFO-CTO backbones after sintering at 850 °C.

3.2. Electrochemical characterization

The electrochemical performance of the different backbones with and without catalyst activation on CGO-Co electrolytes was studied at 850 °C in streams of 21% O₂ in N₂ and 5% O₂ in N₂, mimicking feed and clean sweep side environments, respectively. Fig. 2 shows the total polarization resistance (R_p) values obtained for each catalytically-activated membrane after 90 min on-stream under both conditions (21% O₂ and 5% O₂ in N₂) at 850 °C. As can be observed, the performance of the infiltrated backbones is enhanced relative to the non-activated pure NFO-CTO, except for the backbones infiltrated with Al and Nb catalysts. Notably, the Pr-activated electrode shows a significant enhancement, achieving a 10-fold reduction in R_p from 1.85 to 0.17 Ω·cm² in 21% O₂. Previous studies have demonstrated strong and stable promotion of the oxygen oxidation/reduction reaction (OORR) kinetics by Pr activation, particularly using Pr₆O₁₁ nanoparticles infiltrated into the porous backbones. [27,28] Due to this notable effect, Pr-based binary formulations were explored, specifically (i) Ce-Pr in a 1:1 molar ratio and (ii) Pr-Al in a 4:1 molar ratio. The Ce-Pr oxide binary formulation was selected due to previously reported outstanding catalytic properties.[28,34], while the Pr-Al-based catalyst was selected to combine the beneficial properties of Pr with the acidic character of Al, potentially improving the OORR kinetics under oxyfuel environments due to lower CO2 and SO2 adsorption on the active sites. The distinct stoichiometric ratios of Pr-Al and Ce-Pr catalysts were tailored to balance catalytic activity and surface properties, aiming to achieve the optimal OORR performance by compensating for the activity of Al₂O₃ and maximizing the Ce-Pr synergistic effect. As shown in

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Fig 2a, both binary catalysts outperform most studied catalysts, although the Pr catalyst still yields Article Online the lowest R_p values. The Nyquist and Bode plots depicted in Fig. 2a and b show that the observed R_p reduction in the activated samples is due to a significant lowering of the imaginary component of the impedance at medium and low frequencies (10^{-1} - 10^3 Hz), corresponding to enhanced OORR kinetics at the catalyst surface. Nb and Al catalysts exhibit higher impedance values than pure NFO-CTO in this region, likely due to the lack of redox properties, which hinders oxygen surface exchange. The catalysts producing the most substantial reduction in the R_p values are Pr, Pr-Al, and Ce-Pr. Conversely, pure NFO-CTO and Zr-, Ce-, and Sm-based catalysts exhibit increased impedance contributions at higher frequencies (10^4 Hz), which may be attributed to partial blocking of pathways for electronic conduction.

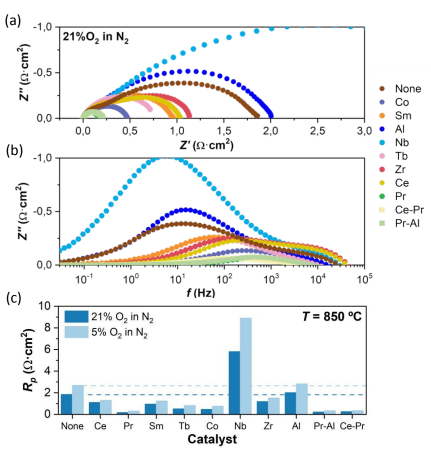


Fig. 2. (a) Nyquist, (b) Bode plots in 21% O₂ in N₂ and (c) Polarization resistance of NFO-CTO backbones, both pure and infiltrated with different catalysts (None, Co, Sm, Al, Nb, Tb, Zr, Ce, Pr, Ce-Pr, and Pr-Al) in 21% O₂ in N₂ and 5% O₂ in N₂, mimicking clean feed and sweep conditions at 850 °C.

Further tests were performed under oxyfuel-like conditions, where the presence of CO_2 and SO_2 is a critical factor, to evaluate performance in such challenging environments. Data of EIS measurements in Fig. 3 show the electrochemical behavior of the different activated backbones under 5% O_2 in CO_2 and 250 ppm SO_2 , 5% O_2 in CO_2 . Switching the gas atmosphere from 5% O_2 in O_2 in

(e.g., via in situ XPS or DRIFTS) was not performed in this study, the combination of reversible Article Online R_P changes, the affected frequency region, and supporting literature provides strong evidence that SO₂ affects the oxygen surface exchange reaction. Despite this detrimental effect, the backbones infiltrated with Zr, Sm, Ce, Pr, Pr-Al, and Ce-Pr catalysts still outperform the non-activated NFO-CTO backbone, with Ce-Pr showing the most pronounced improvement. Specifically, Ce-Pr infiltration reduces the polarization resistance under SO₂ exposure from 7.1 Ω ·cm² (for pure NFO-CTO) to 2.5 Ω ·cm². After SO₂ exposure, the R_p recovery was monitored for 10 h under 5% O₂ in N₂ atmosphere. Table 2 presents the R_p values in 5% O₂ in N₂ before and after SO₂ exposure. The non-activated backbone shows the slightest change in R_p , while the backbones infiltrated with Pr, Tb, Co, and Pr-Al exhibit the largest increases. With the exception of Nb and Al, all infiltrated samples maintain lower R_p values than the pure backbone after the recovery. Notably, Ce-Pr and Ce-Al activated backbones display particularly low R_p values of 0.40 and 0.78 Ω ·cm², respectively, even after SO₂ exposure.

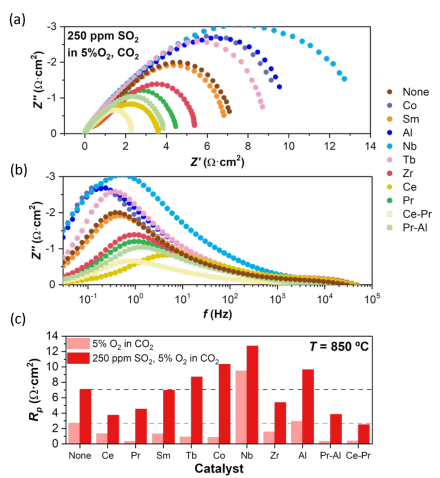


Fig. 3. (a) Nyquist, (b) Bode plots in 250 ppm SO₂, 5% O₂ in CO₂ and (c) Polarization resistance of NFO-CTO pure backbones and infiltrated with different catalysts (None, Co, Sm, Al, Nb, Tb, Zr, Ce, Pr, Ce-Pr, and Pr-Al) in 5% O₂ in CO₂ and 250 ppm SO₂, 5% O₂ in CO₂, mimicking oxyfuel conditions at 850 °C.

Table 2. R_p values obtained from EIS measurements for the different activated backbones under the View Article Online different conditions tested at 850 °C.

	$R_p\left(\Omega\cdot\mathrm{cm}^2 ight)$					
Catalyst	21% O ₂ in N ₂	5% O ₂ in N ₂	5% O ₂ in CO ₂	250 ppm SO ₂	5% O2 in N2*	5% O ₂ in N ₂
None	1.85	2.69	2.71	7.09	2.62	-3%
Ce	1.10	1.31	1.33	3.73	1.53	17%
Pr	0.17	0.32	0.35	4.53	1.06	236%
Sm	0.96	1.25	1.31	6.96	2.10	68%
Tb	0.52	0.82	0.91	8.70	1.97	141%
Co	0.47	0.76	0.84	10.34	1.56	103%
Nb	5.81	8.91	9.47	12.72	10.33	16%
Zr	1.19	1.51	1.56	5.38	1.86	23%
Al	2.01	2.82	2.92	9.63	3.35	19%
Pr-Al (4:1)	0.22	0.34	0.35	3.83	0.78	132%
Ce-Pr (1:1)	0.25	0.35	0.38	2.49	0.40	29%

^{*:} After being exposed to SO₂.

The polarization resistances of the Pr, Ce, Al, Pr-Ce, and Pr-Al activated backbones, as well as the bare NFO-CTO electrode, were obtained by fitting the impedance spectra to an equivalent circuit consisting of three R||CPE elements.[16,24] The resistances at high (R_{HF}) , medium (R_{MF}) , and low frequencies (R_{LF}) are plotted for the four different studied atmospheres. As shown in Fig. S4, the resistance at HF (>1 kHz) remains almost unaltered across all tested conditions. However, when comparing the measured values of R_{HF} for all catalysts, including the bare material, the highest value of R_{HF} is observed for Ce, Ce-Pr, and the non-infiltrated backbone. This indicates that infiltration of Ce does not enhance the high-frequency performance, which is attributed to the lower conductivity of CeO₂ [35–37]. Incorporation of Al₂O₃ increases the resistance at medium frequencies (10-10³ Hz). Considering that the value of R_{MF} is usually linked with the TPB length, the higher R_{MF} of the Al-infiltrated electrode layer (compared to the bare electrode) can be attributed to the low catalytic activity of Al and, therefore, to the reduction of the TPB length. The limited influence of CO₂ in the inlet stream may indicate negligible competitive adsorption between O₂ and CO₂ at the active sites. Finally, exposure of the activated electrode layers to 250 ppm SO₂ has a pronounced detrimental impact on the polarization resistance at both medium and low frequencies. These frequency regions (<10 Hz) are typically associated with surface and gas-phase diffusion processes, [24,27]. The larger size and higher polarizability of SO_2 -relative to O_2 - facilitate its strong adsorption on active sites, and its dissociation can lead to the formation of sulfate species. These species not only block active sites for oxygen adsorption but also obstruct gas pathways within the porous structure, thereby simultaneously hindering surface exchange reactions and gas diffusion, resulting in a marked increase in polarization resistance.

Overall, the R_P values measured in this work fall within the range reported for state-of-the-art air/O₂ electrode materials, confirming their relevance for application. The invariance of R_P upon switching from O₂/N₂ to O₂/CO₂ indicates superior CO₂ stability compared to BSCF and LSCF, which typically degrade under such conditions. [38,39] Additionally, the materials demonstrated tolerance to SO₂ exposure, indicating robust performance in oxyfuel-relevant environments.

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3.3. Pulse isotopic exchange studies

Pulse Isotopic Exchange (PIE) measurements were performed to further investigate the surface electrochemistry of the bare and catalyst-activated NFO-CTO backbones with Pr. Ce, Al, Ce-Pr, and Pr-Al. Fig. 4 shows that the non-activated reference material exhibits a relatively high activation energy of 136 ± 8 kJ/mol. Infiltration with Pr, a rare earth metal known for its catalytic activity in the OORR, improves the surface exchange rate by more than one order of magnitude. Furthermore, the low activation energy found for the Pr-infiltrated material renders it more suitable for use at intermediate temperatures. In the case of Ce, there is a slight enhancement relative to bare material, which is attributed to the redox properties of CeO₂ nanoparticles.[40,41] Impregnation of the NFO-CTO composite with Al leads to a two-fold Arrhenius behavior of the surface exchange rate, with a higher activation energy at elevated temperatures. The poor performance observed for the composite after infiltration with Al is expected since alumina (Al₂O₃) exhibits poor catalytic activity for the OORR. The presence of these oxide particles on the composite surface reduces the effective active surface area, thereby reducing the surface exchange rate. Conversely, samples impregnated with binary catalysts, Ce-Pr or Pr-Al, show a substantial increase in the exchange rates, consistent with the trends observed in the EIS measurements. For comparison, PIE data of reference materials including BSFC, STF35, GDC, LSCF, and LSC20 are also presented in Fig. S5, allowing a direct assessment of the composites studied to commonly investigated OTM and solid-oxide electrochemical cells (SOC) materials. The results indicate that the investigated composites exhibit surface exchange kinetics and activation energies within the range reported for leading OTM/SOC candidates, underscoring their potential applicability in oxygen transport and electrochemical devices.[42–44]

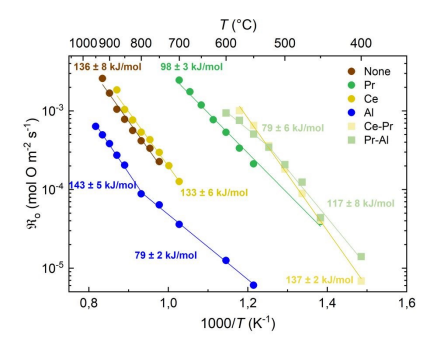


Fig. 4. Surface exchange rate of non-activated and activated NFO-CTO composite powders as a function of inverse temperature.

3.4. Oxygen permeation studies

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Oxygen permeation tests were conducted on 0.6 mm-thick NFO-CTO membranes (bare), coated with a 30 µm-thick porous NFO-CTO backbone (none), and infiltrated with Pr, Ce, Al, Ce-Pr, and Pr-Al catalysts. The O₂ permeation was studied under an Air/Ar gradient in the temperature range from 850 to 700 °C. The evolution of the oxygen flux, $J(O_2)$, with temperature for the different activated cases is depicted in Fig. 5a. The addition of a porous NFO-CTO layer enhances $J(O_2)$ by a factor of 2.5 at 850 °C, which is even more significant as the temperature is lowered. Several publications have reported an increase in the permeation by surface modification [45,46]. This enhancement can be ascribed to the increase in the membrane surface specific area and, thus, to the formation of additional active sites for oxygen exchange. The infiltration of Ce, Pr, and the binary combinations Ce-Pr and Pr-Al into the porous NFO-CTO backbones increases $J(O_2)$. Conversely, $J(O_2)$ decreases after activation with Al, consistent with its poor catalytic activity for the oxygen surface exchange reaction The results in Fig. 5a show that the most significant improvement in flux is obtained after activation with Ce-Pr, resulting in a 6-fold increase of $J(O_2)$, at 850 °C, relative to the non-infiltrated NFO-CTO-coated membrane. These results are in excellent agreement with data from EIS and PIE, confirming the role of the catalyst nanoparticle in activation of the OORR and the importance of the surface exchange reactions in governing O₂ permeation at temperatures below 850 °C.

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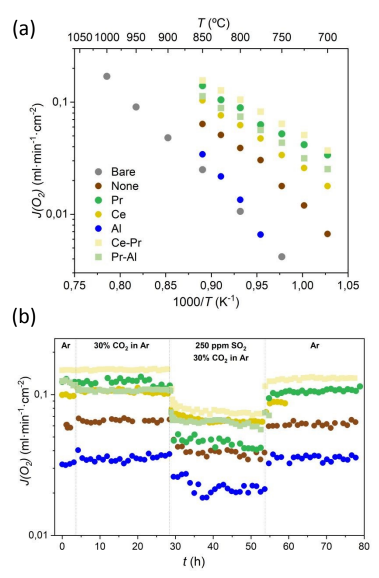


Fig. 5. (a) Oxygen permeation of NFO-CTO membranes (bare), coated with an NFO-CTO backbone (none), and infiltrated with Pr, Ce, Al, Ce-Pr, and Pr-Al catalysts in dependence on temperature ($Q_{\text{feed}} = 100 \text{ ml} \cdot \text{min}^{-1}$, $Q_{\text{sweep}} = 150 \text{ ml} \cdot \text{min}^{-1}$). (b) Oxygen permeation evolution in dependence on time of different activated membranes under different sweep environments involving CO₂ and SO₂ ($Q_{\text{feed}} = 100 \text{ ml} \cdot \text{min}^{-1}$, $Q_{\text{sweep}} = 150 \text{ ml} \cdot \text{min}^{-1}$) at 850 °C.

In view of the above results, Ce, Pr, Pr-Al, and Ce-Pr catalysts are identified as most promising for catalytic activation of membranes subjected to oxyfuel environments. Short-term stability tests of 24 h were performed on NFO-CTO activated membranes by monitoring the evolution of $J(O_2)$ under different environments at the sweep side (Argon, 30% CO₂ in Argon, 250 ppm SO₂ in 30% CO₂ in Argon) while feeding with synthetic air at 850 °C. The results in Fig. 5b demonstrate stable behavior for the coated NFO-CTO membrane under all tested conditions. A slight drop in $J(O_2)$ is observed under SO₂-containing environment, but the flux recovers slowly to its initial value when switching back to clean conditions. This recovery demonstrates the stability of NFO-CTO in oxyfuel atmospheres and indicates that the performance loss is due to SO₂ adsorption, which hinders the oxygen surface exchange reaction without causing permanent structural degradation.

The activation of the NFO-CTO-coated membrane with Al yields lower $J(O_2)$ across all conditions used in this study, confirming its poor activity in the OORR. In contrast, all other

catalysts enhance performance under the given conditions, as expected. Among them, the Ce-Prw Article Online catalyst yields the highest oxygen flux, reaching 0.15 and 0.07 ml·min⁻¹·cm⁻² under CO_2 and SO_2 conditions, respectively. This represents a 2.5-fold improvement with respect to the non-infiltrated NFO-CTO membrane under the same harsh conditions. The Ce and Pr-Al catalysts perform exceptionally well when exposed to SO_2 -containing atmospheres.

Finally, a long-term stability test was conducted using a Ce-Pr-activated NFO-CTO membrane for 100 h at 850 °C. Fig. 6 depicts the oxygen flux when exposed to 250 ppm SO₂, 30% CO₂ in Argon. As can be seen when switching from clean conditions to a sweep gas containing SO₂, $J(O_2)$ drops to 0.07 ml·min⁻¹·cm⁻². Next, $J(O_2)$ declines at a constant rate of 0.00013 ml·min⁻¹·cm⁻² (0.14%) per hour, reaching a value of 0.057 ml·min⁻¹·cm⁻² after 100 h. After the stability test, the feed was switched again to clean Ar sweeping conditions, reaching a value of $J(O_2)$ of 0.11 ml·min⁻¹·cm⁻² after 5 h with a slow flux recovery, which is lower than the initial value of 0.15 ml·min⁻¹·cm⁻² before SO₂ exposure. This type of degradation and recovery determined by SO₂ desorption kinetics has also been observed for other spinel/fluorite membranes.[16,47]

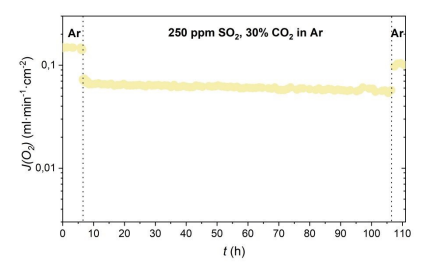


Fig. 6. Oxygen permeation evolution over time for the Ce-Pr activated membrane under different sweep environments involving CO_2 and SO_2 ($Q_{feed} = 100 \text{ ml} \cdot \text{min}^{-1}$, $Q_{sweep} = 150 \text{ ml} \cdot \text{min}^{-1}$) at 850 °C.

3.5. Post-mortem characterization

XRD analysis of the symmetrical cells was conducted after EIS measurements were performed in O₂, CO₂, and SO₂ environments on both the non-activated and activated porous NFO-CTO backbones. The diffraction patterns correspond to the peaks associated with the NFO-CTO phase (porous layer) and the different oxide catalysts (none, Tb₇O₁₂, Nb₂O₃, Co₃O₄, Pr₆O₁₁, Sm₂O₃, ZrO₂, CeO₂, Al₂O₃). The Au signal is due to the gold contacts used as current collectors. No sulfates were detected after SO₂ exposure (Fig. 7), suggesting that the irreversible performance loss observed in the membranes may be attributed to the formation of minor phases or to catalyst nanoparticles coarsening during time on stream.

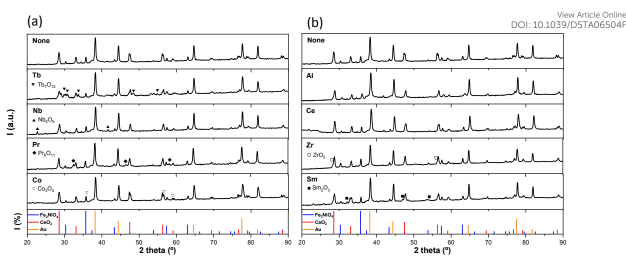


Fig. 7. XRD patterns of the NFO-CTO backbones, non-activated (none) and activated (a) Tb, Nb, Pr, Co, and (b) Al, Ce, Zr, and Sm layers after the EIS tests performed in O₂, CO₂, and SO₂ environments.

A SEM image of a fresh NFO-CTO membrane coated with a porous NFO-CTO backbone infiltrated with Ce-Pr (Fig. 8) is compared with the same cell after the 100 h oxygen permeation stability test in SO_2 - and CO_2 -containing atmospheres. The post-test images reveal a loss of catalyst dispersion and significant particle growth, which likely contribute to the performance degradation over time, particularly the reduced $J(O_2)$ observed upon returning to clean conditions. EDX post-mortem analysis on the tested sample (Fig. S6) reveals no formation of sulfur species, either in the catalytic layer or on the membrane surface exposed to SO_2 , thus confirming the chemical stability of NFO-CTO and CeO_x -PrO_x after long-term CO_2 and SO_2 exposure.

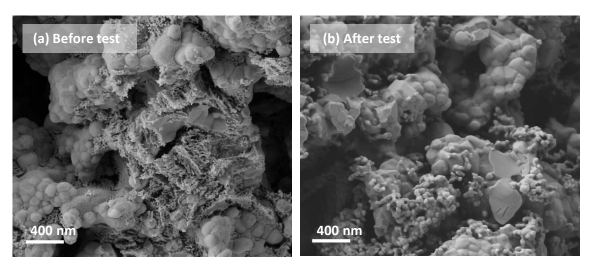


Fig. 8. SEM images of the NFO-CTO membrane coated with a porous NFO-CTO backbone infiltrated with Ce-Pr (a) before and (b) after the 100 h oxygen permeation stability test with SO₂ and CO₂ atmospheres.

4. Conclusions

Targeted catalyst activation strategies enabled a substantial enhancement in the performance of Fe₂NiO₄-Ce_{0.8}Tb_{0.2}O_{2-δ} (NFO-CTO) membranes for high-temperature oxygen separation

applications. Various oxide catalysts, comprising individual lanthanides (Ce, Pr, Tb, Sm) w Article Online transition metals (Co, Nb, Al, Zr), and binary combinations (Ce-Pr and Pr-Al), were systematically screened with the aim of enhancing the oxygen surface exchange kinetics. Through comprehensive electrochemical characterization using electrochemical impedance spectroscopy (EIS) and pulsed isotopic exchange (PIE) techniques, Pr, Ce, Pr–Al, and Ce–Pr were consistently identified as the most effective catalyst systems. These activations significantly enhanced the oxygen surface exchange reaction, correlating with improvements in the overall membrane performance. Activation of the porous NFO-CTO membrane overlayers with Pr, in particular, yielded a 10-fold reduction in the polarization resistance, decreasing from 1.85 Ω·cm² to $0.17 \Omega \cdot \text{cm}^2$ in air. Under simulated combustion environments containing 250 ppm SO₂ and CO₂, Ce–Pr activation demonstrated the highest activity and stability, achieving an R_p of 2.49 $\Omega \cdot \text{cm}^2$. Oxygen permeation measurements further supported the electrochemical findings, with Pr-, Ce-, Pr-Al-, and Ce-Pr-activated membranes consistently exhibiting the highest oxygen fluxes. Under clean conditions at 850 °C, a six-fold increase in oxygen production was achieved with Ce-Pr activation, while a 2.5-fold improvement was observed under harsh SO₂/CO₂ conditions. Longterm stability tests conducted over 100 h indicated low performance degradation, with a sustained oxygen flux decline of only 0.14% per hour. Microstructural analysis suggested that the degradation mechanisms were primarily associated with nanoparticle coarsening and partial loss of catalyst dispersion.

These results highlight the critical role of tailored catalyst activation in simultaneously improving the activity and durability of mixed ionic–electronic conducting membranes for high-temperature oxygen separation technologies. Under oxyfuel conditions, achieving high oxygen permeation requires not only the enhancement of surface exchange kinetics but also robust resistance against performance degradation caused by gases present in oxyfuel conditions, such as CO₂ and SO₂. In this study, the NFO-CTO membrane was used as a model material, and the insights gained were applied to the design and selection of catalyst activation strategies. The resulting membranes exhibit high catalytic activity combined with excellent stability under harsh CO₂/SO₂-containing environments. These findings are expected to be transferable to other dual-phase composite systems.

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Conflicts of interest

There are no conflicts to declare.

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- ‡ Footnotes relating to the main text should appear here. These might include comments relevant to but not central to the discussion, limited experimental and spectral data, and crystallographic data.
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