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Transition metal-nitrogen-carbon catalysts for oxygen reduction reaction in neutral electrolyte



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ABSTRACT

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Keywords: ORR PGM-free Current production Electron transfer Neutral pH Platinum group metal-free (PGM-free) catalysts based on M-N-C types of materials with M as Mn, Fe, Co and Ni and aminoantipyrine (AAPyr) as N-C precursors were synthesized using sacrificial support method. Catalysts kinetics of oxygen reduction reaction (ORR) was studied using rotating ring disk electrode (RRDE) in neutral pH. Results showed that performances were distributed among the catalysts as: Fe-AAPyr > Co-AAPyr > Mn-AAPyr > Ni-AAPyr. Fe-AAPyr had the highest onset potential and half-wave potential. All the materials showed similar limiting current. Fe-AAPyr had an electron transfer involving $4e^-$ with peroxide formed lower than 5%. Considering H_2O_2 produced, it seems that Co-AAPyr, Mn-AAPyr and Ni-AAPyr follow a 2 × 2 e^- mechanism with peroxide formed during the intermediate step. Durability test was done on Fe-AAPyr for 10,000 cycles. Decrease of activity was observed only after 10,000 cycles.

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1. Introduction

Oxygen reduction reaction (ORR) at the cathode is often the limiting steps in the reduction reactions happening generally into FCs dealing with fuels like hydrogen [1–2]. Those limitations have been studied deeply in acidic and alkaline environments but not in neutral media [2–5]. ORR follows different patterns in function of the pH environment in which the reaction occurs. H^+ and OH^- are required to complete the reaction in acidic and alkaline media respectively [6–7]. Neutral media (pH = 7) has a low concentration (10⁻⁷ M) of H⁺ and OH⁻ that affects negatively the ORR kinetics. High overpotentials are shown in the existing literature [2–5].

Microbial fuel cells (MFCs) operate at neutral conditions and room temperature in order to allow bacteria activity and survivor and this lowers the ORR performances. Catalysts are then used to enhance the reaction rate. Three different typologies of catalysts are investigated in MFC [3–5].

The first one is based on platinum group metals as inheritance of the most studied acidic or alkaline fuel cells. This choice cannot be justified due to the high catalyst cost compared to the low power output produced. Moreover, MFCs operates in very polluted environment and interaction of anions with Pt is known to deactivate the catalytic activity in short time [8–10].

The second choice has been introduced in the past years and it concerns the utilization of carbonaceous-based materials but unfortunately overpotentials are very high and kinetics remains weak [11–12].

The third choice is the utilization of M-N-C types of catalysts in which M can be platinum group metal (PGM) [13–16] or completely PGM-free with M being an earth abundant transition metal like Mn, Fe, Co and Ni. M-N-C catalysts were deeply studied in acidic and alkaline media [17–20] and recently the most pursued in neutral media [3–5,21–25]. PGM-free catalysts showed high performances and durability in long terms operations compared to Pt [9–10,26].

Only few mechanistic studies showing catalysts kinetics in neutral media are presented in literature focusing on Fe-based [24,27–29] or carbonaceous catalysts [30–31]. None of the studies faced the mechanistic activity and the kinetics of M-N-C with M as Co, Ni and Mn that are the other earth abundant metals together with Fe mostly used to substitute the more expensive Pt. Kinetics behavior of those catalysts is quite elusive and not well understood. Moreover, H_2O_2 yield production, electron transfer mechanisms and loading effect on kinetic current density have not been shown for any catalysts working in neutral media.

In this work, we studied the kinetics of four PGM-free catalysts based on the same synthetic process and the same precursors (aminoantipyrine) using rotating ring disk electrode (RRDE) method. Disk current, Tafel plots, H_2O_2 production, electron transfer and loading effect on kinetic current density are discussed. Durability test (10,000 cycles) on Fe-AAPyr is presented and a comparison with Pt is shown. This work enhances the understanding of PGM-free catalysts working in neutral conditions.

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2. Materials and method

2.1. Catalysts preparation

Sacrificial support method (SSM) was used to synthesis the catalysts investigated. Aminoantipyrine (AAPyr) was used as organic precursor for the synthesis. (Fe(NO₃)₃ × 9H₂O, Co(NO₃)₂ × 6H₂O, Mn(NO₃)₂ × 4H₂O, Ni(NO₃)₂ × 6H₂O) were the metallic salts mixed with AAPyr and impregnated with fumed silica (~250 m² g⁻¹). Ball milling was used to ground the mixed materials. Heat treatment was then applied in a constant flow (100 mL min⁻¹) of UHP nitrogen. Temperature was raised till 950 °C (25 °C min⁻¹) and stabilized for 30 min. After pyrolysis, the mixture was cool down at room temperature and silica was etched using 20 wt% HF for 12 h. The catalyst was washed and then dried overnight. Obtained catalysts were named in function of the metal used that was Mn, Fe, Co, Ni.

2.2. Electrochemical measurements and analysis

Rotating ring disk electrode (RRDE) technique on Mn-AAPyr, Fe-AAPyr, Co-AAPyr, Ni-AAPyr inks was used to evaluate the catalysts kinetics. The inks were prepared uniformly suspending 5 mg of each catalyst into 0.075% of 1100 EW Nafion solution (FuelCellStore, USA) and then sonicating for three times (30 s) for a correct dispersion. Different loadings (50, 100, 200, 400 and $600 \,\mu \text{g cm}^{-2}$) were tested. Experiments were done in neutral conditions with electrolyte solution composed by potassium phosphate buffer (0.1 M) and KCl (0.1 M). The solution was purged with pure oxygen for over 30 min. RRDE setup allows to measure the disk current produced by the catalysts and also the peroxide produced as intermediate of the 4e⁻ transfer through the ring current. Linear sweep voltammetry (LSV) was run from 1.08 V to 0.18 V (vs RHE) at scan rate of 5 mV s⁻¹ in separate triplicates. Disk (j_D) and ring (j_R) currents densities are correlated by Eq. (1) in which the electrons transferred are estimated.

$$n = \left| \frac{4 \times j_D}{j_D - j_R} \right| \tag{1}$$

The estimated electrons transferred are also used for identifying the H_2O_2 produced (%) during the ORR process as showed by Eq. (2):

$$x = \frac{4-n}{2} \times 100 \tag{2}$$

Durability tests were done to Fe-AAPyr (loading 600 μ g cm⁻²) cycling 10,000 times from 1.08 V to 0.18 V (vs RHE) at scan rate of 50 mV s⁻¹. The acquisition of the reported LSV (10, 100, 1000, 3000 and 10,000 cycles) was done at scan rate of 5 mV s⁻¹.

Koutecky-Levich equation Eq. (3) was used and $|i_d|^{-1}$ was plot against ω^{-1} . i_k and *n* can be extrapolated from the system.

$$\frac{1}{j_d} = \frac{1}{j_k} + \frac{1}{0.62nFC_{02}D_{02}^{2/3}\nu^{-1/6}\omega^{1/2}A}$$
(3)

 i_k is the electrode potential dependent kinetic current density of the ORR, n is the average number of electrons transferred per catalytic event (4 is the maximum), F is the Faraday's constant (96,487 C mol⁻¹), C_{02} is the concentration of O_2 in the electrolyte (1.117e⁻⁶ mol mL⁻¹), D_{02} is the O_2 diffusion coefficient in aqueous media (1.9e⁻⁵ cm² s⁻¹), and v is the kinetic viscosity of the electrolyte (0.01073 cm² s⁻¹), ω is the angular momentum in rad·s⁻¹, and A is the electrode sectional area.

3. Results and discussion

3.1. Disk current, e^- transfer mechanism and H_2O_2 production

Disk currents (Fig. 1a), electrons transferred (Fig. 1b) and peroxide yields (Fig. 1c) are here presented for every catalyst. Different trends can be noticed into the disk current (Fig. 1a). Higher onset potential was measured by Fe-AAPyr (0.98 V (vs RHE)) followed by Co-AAPyr (0.892 V (vs RHE)), Mn-AAPyr (0.864 V (vs RHE)) and Ni-AAPyr (0.802 V (vs RHE)). The onset potential trends followed the current densities produced from the kinetic limited region until reaching the transport limited plateau. Fe-AAPyr had the highest disk current followed by Co-AAPyr, Mn-AAPyr and Ni-AAPyr with the latter as the lowest value measured (Fig. 1a). Fe-AAPyr, Co-AAPyr and Mn-AAPyr reached a similar diffusion limited current at 0.185 V (vs RHE) quantified as \approx 4.5 mA cm⁻². Ni-AAPyr had current of \approx 4 mA cm⁻² at that same potential. As for the half-wave potential, the trend is as follows: 0.807 V for Fe, 0.782 V for Co, 0.716 V Mn and 0.571 V for Ni. The electrons transferred at 0.185 V (vs RHE) are estimated to be between 3.5 and 4 with exact number of 3.96 for Fe-AAPyr, 3.84 for Co-AAPyr, 3.67 for Mn-AAPyr and 3.64 for Ni-AAPyr (Fig. 1b). Interestingly, at higher potentials Co-AAPyr and Ni-AAPyr had lower electron transferred of 3.00 and 2.87 respectively (Fig. 1b). Also Mn-AAPyr had slightly lower number of electron transferred (3.50) at higher potential (Fig. 1b). At the contrary, Fe-AAPyr was the only catalysts showing the same electrons transferred (3.96) along the range investigated (Fig. 1b). Peak peroxide yield of 50%, 24% and 56% was formed by Co-AAPyr, Mn-AAPyr and Ni-AAPyr respectively at lower overpotentials investigated (Fig. 1c). Fe-AAPyr had low production of peroxide quantified in 2-3%. Peroxide data are necessary for further explaining the electron transfer mechanisms involving the catalysts investigated. In fact, Fe-AAPyr seems to have an apparent direct 4e⁻ transfer mechanism, evidenced by low peroxide production. This mechanism is preferred since the maximum number of electrons is electrochemically transferred during the reaction. In contrast, the high peroxide produced at low overpotentials for Co-AAPyr, Mn-AAPyr and Ni-AAPyr indicates that the peroxide intermediate is produced and then electrochemically oxidized to water at higher overpotentials. This allows to speculate that probably those catalysts have $2 \times 2e^{-}$ transfer mechanisms during ORR. Peroxide production in MFCs is an undesired process since peroxide can negatively affect the electroactive bacteria on the anode electrode. Those results indicated a superior electrocatalytic activity of Fe-AAPyr compared to the other PGM-free catalysts investigated. The origin of such selectivity towards a 4e⁻ mechanism for the Fe based catalyst versus the $2 \times 2e^-$ transfer mechanism for the other metals has been modeled by density functional theory calculations [32-34].

3.2. Effect of loading on ORR kinetics

The effect of the catalyst loading on the ORR kinetics was studied for Fe-AAPyr, as it was the best performing catalysts investigated (Fig. 2). Limiting current densities enhanced substantially increasing the loading from 50 to 200 μ g cm⁻² (Fig. 2a). This indicates that the reaction mechanism of Fe-AApyr follows a 2 × 2e⁻ transfer process, as an increase in loading increases the limiting current densities and decreases the per-oxide yield (Fig. 2c). This is due to a fast H₂O₂ reduction within the catalytic layer, conclusion that is supported by the 4e⁻ transfer mechanism seen at all potentials and loadings (Fig. 2b). H₂O₂ peroxide production was always lower than 10% for all the loading investigated. Peroxide decrease with increase of loading is due to H₂O₂ entrapment inside a thicker catalyst layer, being further reduced before reaching the platinum ring (Fig. 2c).

Fig. 2d displays the kinetic current densities calculated by the Koutecky-Levich analysis for each loading. The increase in kinetic current density between the loadings of 50 and 200 μ g cm⁻² demonstrated that higher availability of active sites within the catalytic layer turn into faster oxygen reduction. This supports the idea of a two-steps electron

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