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Pencil Graphite Rods Decorated with Nickel and Nickel-Iron as Low-Cost Oxygen Evolution Reaction Electrodes

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ABSTRACT

Society is demanding clean energy to substitute the greatly pollutant carbon-based fuels. As an alternative, the green hydrogen produced by electrocatalysis constitutes a nice strategy as its products and reactants are not toxic to the environment. However, the use of scarce materials and the high overpotentials to accomplish the Oxygen Evolution Reaction (OER) make electrocatalysis an uncompetitive process. To solve these challenges, a low-cost procedure for the preparation of earth-abundant Ni, Fe and NiFe decorated electrodes has been developed. For this purpose, Pencil Graphite Rods have been selected as highly porous substrates. A reasonable performance is achieved when they are employed for OER. Furthermore, for the first time, a detailed analysis of Impedance Spectroscopy allows the association of the Ni redox transitions Ni\textsuperscript{2+}/Ni\textsuperscript{3+} and Ni\textsuperscript{3+}/Ni\textsuperscript{4+} (including the identification of the hydrated \(\alpha-\gamma\) and the non-hydrated \(\beta\) phases) with an electrochemical redox capacitance response. Additionally, the Ni\textsuperscript{3+}/Ni\textsuperscript{4+} redox peak capacitance together with a quick decrease in the charge transfer resistance remarks the implication of Ni\textsuperscript{4+} in the OER. These results show the utility of Impedance Spectroscopy as a non-destructive and non-invasive technique to study in detail of these electrochemical systems in operating conditions.
KEYWORDS

Water splitting, NiFe, Ni, OER, Impedance Spectroscopy

1. INTRODUCTION

Limitations in the use of fossil fuels like oil, carbon or natural gas are one of the main challenges that humanity must face in the next few decades. Dependence of the economy upon cheap energy urges environmentally friendly energy sources to satisfy the continuous growth in energy demand around the world. Renewable energies are increasing their importance in the global share of energy production very quickly, but problems regarding the continuity in their availability still limit their spread to attain a carbon-free society, with an affordable and scalable way of storing the energy needed. One of the lines under development, to make this scenario possible, is focused on the use of molecular hydrogen as an energy storage system. $\text{H}_2$ is not only the most efficient energy source (due to its large energetic power per unit mass) but also one of the greenest technologies as its sub-products, essentially water, have no impact on human health and the environment.

Electrocatalysis is regarded as an appropriate technique to produce pure hydrogen by water splitting. To do this, external electric power is employed which, ideally, may be obtained from natural sources, e.g. solar or wind power. To have an efficient energy conversion, the overpotentials needed to produce the oxygen evolution reaction (OER) and the hydrogen evolution reaction (HER) must be minimized. From a thermodynamic and kinetic point of view, OER is the limiting reaction as it involves four electrons and a bigger overpotential. Traditionally, noble metals oxides such as RuO$_x$, IrO$_x$ have been employed as anodes due to their high performance for the OER. Recently, RuO$_x$ has been deposited on Pencil Graphite Rods (PGR) to obtain high-performance porous electrodes. However, the cost of these noble metals is high as they are not earth-abundant, which makes water splitting an uncompetitive process if using them. In addition, RuO$_x$ catalyst shows low stability in alkaline solutions. In this study, high OER activity was achieved using only inexpensive metal-based materials without using expensive precious metal-based materials.

Earth-abundant Ni-based materials have risen as an attractive alternative to noble metal for water oxidation. Among the different Ni electrocatalysts, nickel oxide NiO$_x$ and Ni oxide hydroxide (NiOOH) have attracted great attention mainly due to their efficiency.
and robustness. Since Corrigan discovered the higher activity and the decrease in the overpotential of OER when Fe was incorporated onto a NiOOH layer, a noteworthy effort has been made to understand and design new electrodes with better OER performances. Thus, becoming Ni$_{1-x}$Fe$_x$OOH layered double hydroxides (LDH) the best catalyst based on earth-abundant materials for water oxidation in alkaline media since today. However, the performance of this material is strongly dependent on different issues, such as the chemical and electronic structure of the electrode, the electrochemical environment, the electrode preparation method, etc. It has recently been reported a graphene-nanoplatelets-supported (Ni,Fe) metal–organic framework (MOF) with outstanding performance for water splitting in alkaline media and high stability. It is well-known that NiFe catalysts are more stable in alkaline aqueous solutions than other noble metal catalysts such as RuO$_x$. It has been shown that the combination of Ru and Ni, in a compressed metallic Ru-core and oxidized Ru-shell with Ni single atoms (SAs), led to low overpotentials and high current densities in strong acidic media for water oxidation.

In this paper, we study the electrochemical response of Pencil Graphite Rods decorated with nickel and nickel-iron alloys to perform the oxygen evolution reaction. Techniques such as flame annealing (FA) or electrodeposition were used to decorate the PGRs obtaining reasonable results for OER. Structural analysis was used to identify oxidation states and contents of nickel and iron in the samples while electrochemical measurements allowed us to associate the differences in electrical response of the Ni$^{2+}$/Ni$^{3+}$ transition to the presence of hydrated and non-hydrated phases and to show-up the activation of OER after Ni$^{4+}$ formation.

2. EXPERIMENTAL SECTION

PGR pre-deposition treatment. Pencil Graphite Rods (PGRs, 4B hardness, 2.0 mm diameter and 13.0 mm length, Mitsubishi Pencil Co., Ltd., Japan) were used as supporting electrode. To control the area of the PGRs, tips were polished with a polishing machine (LaboPol Struers) until the surface was completely flat and homogeneous. PGRs have a coating polymer on the surface which makes it stronger. In order to remove this coating, the whole rod was burned for 1 min until it became red because of the heat. This process was repeated 10 times and led to a porous structure of carbon with traces (0.2-0.4 %) of iron as checked by SEM. Using this technique, we prepared the so-called FA@PGRs.
**Catalyst (co)-deposition on PGR by flame annealing.** Flame annealing (FA) deposition was based on the thermal procedure proposed by Tsuji et al.\(^9\) To prepare Ni and Fe decorated PGRs, a 10 mg/mL precursor solution (NiCl\(_2\)x6H\(_2\)O for Ni-based electrocatalyst and FeCl\(_3\)x6H\(_2\)O for Fe-based electrocatalysts) in ethanol was prepared. The thermal treatment process consists of three steps which are 10 times repeated: (1) First, the PGR was flame heated until it became red. (2) Immediately, PGR was retired from the flame and two seconds after it was dipped for five seconds into a vial which contained 8 mL of the precursor solution. (3) Finally, flame annealing was applied to the electrode for 20 s. To anneal, the PGR is slid in the direction of the rod axis, then turned 180° to the left and right by hand so that the catalyst can be annealed evenly. Using this technique, we prepared FA@Ni/PGR and FA@NiFe/PGR electrodes.

**Ni electrodeposition on PG.** The electrodeposition (ED) of nickel on PGR electrodes was performed by cyclic voltammetry. A mixture of 0.1 M Na\(_2\)SO\(_4\) and 0.02 M NiCl\(_2\)x6H\(_2\)O in water was used as the electrolyte solution and Ag/AgCl and Pt were used as reference and counter electrode, respectively. After PGR pre-deposition treatment, cyclic voltammaries, consisted of 25 cycles from -1.2 to 0.2 V vs V\(_{Ag/AgCl}\), with a scan rate of 50 mV·s\(^{-1}\), were performed. Using this technique, we prepared ED@Ni/PGR electrodes.

**Scanning electron microscopy (SEM).** The morphological characterization of the samples was performed in a Field Emission Scanning Electron Microscopy with a JSM-7000F JEOL FEG-SEM system (Tokyo, Japan), equipped with an INCA 400 Oxford EDS analyser (Oxford, U.K.) and operating at 15 kV.

**X-ray photoemission spectroscopy (XPS).** A photoelectron spectroscopy apparatus installed at a beamline 07B end station of New SUBARU synchrotron radiation facility performed the chemical states and surface characterization by XPS technique. The XPS peaks were fitting by attributing the oxidation state found in the bibliography to the corresponding binding energy. Ni (2p) was adjusted by 6 peaks at 855.5 eV, 856.9 eV, 873.5 eV, and 875 eV, 852.7 eV and 871.4 eV, which were attributed to Ni\(^{2+}\) (2p\(^3 / 2\)), Ni\(^{3+}\) (2p\(^3 / 2\)), Ni\(^{2+}\) (2p\(^1 / 2\)), Ni\(^{2+}\) (2p\(^1 / 2\)), Ni\(^0\) (2p3/2) and Ni\(^0\) (2p1/2), respectively. Fe (2p) peak was adjusted by 4 peaks at 725.4 eV, 723.5 eV, 714 eV and 711 eV, which were attributed to Fe\(^{3+}\) (2p\(^3 / 2\)), Fe\(^{2+}\) (2p\(^3 / 2\)), Fe\(^{3+}\) (2p\(^1 / 2\)), Fe\(^{2+}\) (2p\(^1 / 2\)). O (1s) was adjusted by 5 peaks at 529.4 eV, 530.7 eV, 530 eV, 531.5 eV and 533.3, which were attributed to nickel oxide, nickel hydroxide, iron oxide, C = O and C-O, respectively.
**Electrochemical measurements.** All electrochemical measurements were performed on a PGSTAT302N potentiostat (Methrom-Autolab), equipped with a FRA32 Module. The electrochemical performance and electrochemical characterization were carried out by using a homemade polypropylene one-compartment three electrode electrochemical cell (EC). No corrosion effects were observed on the material after use. In the EC, an aqueous Ag/AgCl electrode was used as a reference electrode and Pt mesh was employed as counter electrode. To ensure high basicity during the experiments 8 M KOH in water was used as electrolyte. Cyclic Voltammetries (CV) were taken at a scan rate of 25 mV·s⁻¹. Impedance Spectroscopy (IS) measurements were carried out at selected bias with a perturbation of 20 mV and a frequency range from 1 MHz to 10 mHz. CV and IS measurements are represented as a function of the overpotential: \( \eta = V_{\text{app}} \text{ vs RHE} – 1.23 \text{ V} \), and the interfacial overpotential (i.e. the applied potential corrected by the voltage drop at the external series resistances \( \eta – \text{IR}_{\text{s}} \)).

The CV and IS measurements of the ED@Ni/PGR sample was performed under Fe free KOH electrolyte. The KOH employed in this experiment was treated by following the procedure reported by Boettcher et. al.²⁴ Ni(NO₃)₂ x 6H₂O salt was dissolved in high purity KOH 1M solution in order to precipitate Ni(OH)₂. Once the salt was precipitated, the solution was centrifugated and the supernatant was decanted. This procedure was repeated three times until Ni(OH)₂ was completely pure. Then, the pure Ni(OH)₂ was redissolved in high purity 8 M KOH and stirred for 1h. Finally, the purified electrolyte was centrifuged and decanted into a polypropylene bottle for its electrochemical use.

**Electrochemically Active Surface Area (ECSA).** ECSA of PGR was calculated by Impedance Spectroscopy following equation 1.

\[
\text{ECSA} = \frac{C_{\text{dl}}}{C_s} \quad (1)
\]

where \( C_s \) is the capacitance of a planar glassy carbon microelectrode that was used as reference and \( C_{\text{dl}} \) stands for the double-layer capacitance of the PGR electrodes (\( C_{\text{PGR}} \)) obtained at an overpotential of -0.20V vs \( V_{\text{RHE}} \), where neither redox process nor Faraday current were observed. Both capacitances were normalized to the geometric area of the electrode.

3. **RESULTS AND DISCUSSION**
Pencil Graphite Rods (PGR) are made up of a graphite structure mixed by a polymer assembly which makes them harder (Fig. 1a). After a flame annealing (FA) treatment, part of the polymer is removed and the FA@PGR is transformed in a network of branches of crystalline graphite (Fig. 1b). The space between graphite branches provides the high porosity of the rods. Previous data by SEM and BET showed that flame annealing treatment improves the porosity of PGRs.

![Figure 1. Schematic structure of a) PGR, b) graphite branches, c) detail of graphite branches decorated with catalyst for OER](image)

We tested FA@PGRs for OER, however, they were not good catalysts for this reaction. For this reason, in order to improve the performance of PGRs, we developed new electrodes by the deposition of Ni and NiFe on the PGR (Fig. 1c). Ni was deposited following two different methodologies, namely flame annealing (FA) and electrodeposition (ED), obtaining the corresponding electrodes FA@Ni/PGR ED@Ni/PGR (see the experimental section for experimental details). To prepare the PGR decorated with NiFe we followed only the FA procedure (FA@NiFe/PGR).
The morphology and the composition of the decorated PGRs were studied by SEM and EDX techniques (see Fig. S1, S2 and S3 in supporting information [SI]). The SEM image of the ED@Ni/PGR sample showed an ordered and homogeneous surface composed of Ni particles of variable size, around 200 nm on top of the PGR. Compared to the ED sample, FA samples present smaller particles on the top of a disordered and non-homogeneous PGR surface. From the microanalysis made at SEM, we could conclude that the amount of Ni deposited by ED is higher than by FA (see Fig. S1 and S2 in SI). Moreover, we could also determine the Ni/Fe ratio of each sample. For the FA@NiFe/PGR, the Ni:Fe ratio was ~3:2. In the case of FA@Ni/PGR and ED@Ni/PGR iron was also detected (Ni:Fe ratio 9:1 and 24:1, respectively). The presence of iron in these two samples was due to contamination of iron of the bare PGR (~0.2%)\(^9\). The presence of iron, even in small amounts, has implications for the performance of the electrodes as will be explained.

XPS measurements were performed to investigate the chemical environment in the electrodes at the surface level (see Fig. S4 in SI). The analysis of the high resolution XPS spectra and the data fitting of Ni (2p) peak in each sample confirms the non-homogeneity environment of these samples. The Ni (2p) peak in FA@Ni/PGR and FA@NiFe/PGR electrodes show the presence of Ni in two different oxidation states, Ni\(^{2+}\) and Ni\(^{3+}\).\(^{26,27}\) The proportion of each oxidation state was quantified as a function of the peak area (Table S2, SI) and not differences were observed between this two samples. By contrast, Ni (2p) peak in the ED@Ni/PGR sample points to the presence of Ni in the electrode as a mixed phase between metallic and oxidized Ni. The quantification of each state indicates the same concentration of Ni\(^{2+}\) species, pointing that ED led to Ni\(^0\) species instead of Ni\(^{3+}\).

The presence of Fe could also be observed by XPS as a Fe (2p) peak. The presence of Fe in the samples was detected as oxidized state species Fe\(^{2+}\) and Fe\(^{3+}\). This peak was already found for the bare PGR treated by FA (FA@PGR), confirming that impurities of Fe came from the PGR. Additionally, Fig. S4c shows the high resolution XPS spectra of O (1s) for the electrodes. The presence of C-O interactions either as a double or singlet bond dominates in the sample response. The differences in the O (1s) spectra in the samples have been attributed to Ni-O and Fe-O interactions.\(^{28,29}\)
Electrochemical oxygen evolution reaction on Ni and NiFe PGR.

We performed studies of the performance in the OER of FA@PGR electrodes and the corresponding electrodes decorated with Ni and NiFe (ED@Ni/PGR, FA@Ni/PGR and FA@NiFe/PGR, respectively (Fig. 1c). These tests were performed with 8M KOH solution. The reason why the KOH concentration is 8M is that the activity of water electrolysis is higher in a higher concentration of KOH aqueous solution. We sought to obtain higher catalytic activity by testing at a concentration of 8M KOH higher than 6M. Fig. S5 in supporting information shows the cyclic voltammetry for the oxidation of water using FA@NiFe/PGR in 1M and 8M. It can be observed how the activity and onset for this reaction is improved by increasing the pH of the electrolyte. Fig. 2 shows the cyclic voltammograms and the Tafel plots obtained. For a better analysis of the electrode behaviour, the plots in Fig. 2 have been corrected from potential drop at series resistance.

![Cyclic voltammograms and Tafel plots](image)

**Figure 2.** a) Cyclic voltammograms of FA@PGR, ED@Ni/PGR, FA@Ni/PGR and FA@NiFe/PGR and (b) Tafel Plots of FA@PGR, ED@Ni/PGR, FA@Ni/PGR and FA@NiFe/PGR electrodes measured in KOH 8M. Scan rate: 25 mV·s⁻¹. Tafel plots were obtained from J-V curves in stationary conditions.

The analysis of FA@PGR sample in Fig. 2a shows a hysteretic behaviour in the flat area of the CV. This capacitive behaviour could be associated with the microporous structure of the rods and the ability of carbon to absorb small cations. In the case of the catalyst decorated samples, the hysteresis cannot be clearly observed mainly due to the presence of the Ni(OH)₂/NiOOH redox peak, which appears preceding to the region corresponding to OER.
The potential at which the Ni²⁺/Ni³⁺ redox peak appears, depends on both the Fe content in the catalyst and the Ni phase segregation. It has been reported that the presence of more than 10 % of Fe produces that Ni redox peak appear close to OER. By contrast, lower amount of Fe in the Ni electrode separates both phenomena. Thereby, the Ni redox peak is displaced to lower overpotentials while, at the same time, the OER overpotential is higher. In our case, as can be seen in Fig. 2a, we observe the same trend. Thus, for ED@Ni/PGR with a Ni:Fe ratio of 24:1 (estimated by EDX) the wide redox peak for Ni²⁺/Ni³⁺ appears at an overpotential of 0.12 V vs RHE in the forward direction (0.08V vs RHE in the reverse direction) with the onset overpotential for the OER occurring at 0.26 V. However, for FA samples, FA@Ni/PGR and FA@NiFe/PGR with Ni:Fe ratios 9:1 and 3:2, respectively (see Fig. S2 and S3 in SI), both the redox peak and the OER onset are so close that the two phenomena approach each other and overlap in the CV. From the steady-state J-V curves of the electrodes in Fig. 5d, (and Fig. S6d in SI), we could determine with better accuracy both the onset of OER and, after subtracting the Faradaic contribution, the peaks of the overpotentials of the redox states. Thus, for the two FA samples that have Ni:Fe ratios above 9:1, the same values are obtained for the Ni²⁺/Ni³⁺ redox peak, ~0.20V vs RHE in forward (and 0.09V in the reverse direction) and for the onset overpotential for OER, 0.22 V vs RHE.

Table 1. Overpotential at the current density of 10 mA·cm⁻² was found in this paper and compared with previous studies. In brackets the value of η @10 mA·cm⁻² after correcting the potential drop at the Rs.

| Catalyst | Electrode Support | Electrolyte | η @10 mA·cm⁻² (mV) | References |
|----------|-------------------|-------------|--------------------|------------|
| NiFe     | PGR               | 8 M KOH     | 252 (240)          | this work  |
| Ni       | PGR               | 8 M KOH     | 290 (270)          | this work  |
| RuOₓ     | PGR               | 1 M KOH     | 312                | (9)        |
| NiFeₓFe₀.ₓ- AHNA | NiFe NW | 1 M KOH     | 180                | (14)       |
| NiFe     | EG                | 1 M KOH     | 214                | (31)       |
| NiFe     | DG                | 1 M KOH     | 310                | (32)       |
| NiFe     | Ni Foam           | 1 M KOH     | 240                | (27)       |
| NiFe     | Pt                | 1 M NaOH    | 340                | (33)       |
Focusing now on the performance of the electrodes, the interfacial overpotential needed to deliver 10 mA·cm\(^{-2}\) (of the geometrical area) is ~270 mV for ED@Ni/PGR and ~240 mV for FA@NiFe/PG, see Table 1 and Table S1. These values are in line with the ones reported for NiFe deposited onto graphene and exfoliated graphite as supporting electrode, and improve some of the published values shown in Table 1.\(^{31,32}\) Therefore, these materials provide top performance electrodes for OER while keeping an easy and low-cost processing method. We believe that the performance of our electrodes is due to their high surface area and the interaction between Ni and Fe. As can be seen from the data in Table 1, there is still room to improve the performance of the electrodes, as for example minimizing series resistance by including carbon doping to improve its conductivity.

Tafel plots in Fig. 2b also showed the increase in the OER catalytic activities of PGR by the addition of the catalysts. In addition to the decrease of the onset potential observed for the decorated samples vs the bare one, the Tafel slope is reduced ~60 mV·dec\(^{-1}\) when the catalyst is deposited on top of the PGR, which has a slope 120 mV·dec\(^{-1}\) when uncoated. Tafel slopes of 60 mV·dec\(^{-1}\) have been associated with proton-coupled electron transfer mechanism for OER.\(^{34}\)

**Impedance Spectroscopy on Ni and NiFe PGRs.**

We completed the study of FA@PGR electrodes and the corresponding electrodes decorated with Ni and NiFe (ED@Ni/PGR, FA@Ni/PGR and FA@NiFe/PGR, respectively) by Impedance Spectroscopy (IS) measurements. In order to determine the electrocatalytic active surface area (ECSA) of PGR during the FA, through IS, we compared the capacitance of glassy carbon and PGR at low voltages (~0.2V), where neither redox nor faradaic currents were performed. The resulting ECSA obtained shows a huge increase in the electrocatalytic active area of around 700 and 1000 times in ED and FA samples, respectively (Table S1), being these ECSA values responsible for the improvement in the OER performance achieved by PGR after FA treatment.
For a better understanding of the performance of Ni and NiFe on PGR electrodes, IS analysis in the area of Ni$^{2+}$/Ni$^{3+}$ redox peak and the OER onset was performed. Fig. 3 shows the Nyquist and Bode plot data of FA@PGR and FA@NiFe/PGR obtained at overpotentials before and after OER onset (-0.05 and 0.35V vs RHE). The plots obtained for ED@Ni/PGR are omitted here as they have minimum differences with the FA@NiFe/PGR sample. IS shapes are characterized by two features. At high frequencies (Fig. 3b), a 45° line appears which is followed by an arc at medium-low frequencies. This behaviour is described with a transmission line in two different situations: (i) when transport resistance ($R_t$) of electrons in the porous material is smaller than charge transfer resistance ($R_{ct}$) towards the solution, the transmission line presents the shape obtained at low potentials (Fig. 3, line black and blue); (ii) in the opposite case, when $R_t > R_{ct}$, the transmission line has the shape shown at high voltages (Fig. 3, line gray and cyan) which is given by a Gerischer element.\textsuperscript{35–38}

The first situation is observed at low potentials when there is no current flow to the electrolyte. In this case, $R_{ct}$ is very large (\textgtrsim R_t) and the two samples show very similar impedance. When the current flow is activated at higher overpotentials (0.35V vs RHE), $R_{ct}$ becomes smaller than $R_t$ and the arc width ($=[R_t\cdot R_{ct}]^{1/2}$) becomes smaller. The incorporation of the catalysts on the PGRs favours the OER performance, being this effect also observed on the IS measurements by the smaller arc in the NiFe case, indicating a better charge transfer (smaller $R_{ct}$) for the decorated PGRs.
Figure 3. Experimental (a), (b) and (c) NyQuist and (d) Ohmic Drop corrected Bode plot obtained by impedance spectroscopy of OER in KOH 8M using FA@NiFe/PGR and FA@PGR as working electrodes. The dark lines show IS at low potential (-0.05 V vs RHE) and the light lines at high potential (0.35 V vs RHE).

Detailed analysis of IS measurements at overpotentials between -0.2 and 0.4 V vs RHE was performed by fitting the experimental data with the equivalent circuits (EC) proposed in Fig. 4. The EC suggested for these porous electrodes consists of a transmission line, where electrons are regularly distributed around the PGR (Fig. 4a). However, when the metal catalyst is incorporated into the PGR, the electron conduction occurs along the PGR catalyst line, i.e. where the catalyst is in contact with the PGR. In this new EC, (Fig. 4b) a new capacitance is incorporated to describe the OER in the not catalyst covered area. The elements employed in these two EC are described as:

- $R_s$ is the series resistance, which includes the resistance of the carbon rod out of the electrolyte together with the resistances of the bulk of the electrolyte and at the contacts.
- $r_{tr}$ is the electron transport resistance per unit length of the PGR electrode immersed in the electrolyte, which yields a total transport resistance $R_{tr} = r_{tr} \cdot L$. 


• $r_{ct,G}$ is the charge transfer resistance at the graphite/electrolyte interface. In the case of the ED@Ni/PGR and FA@NiFe/PGR samples, it accounts for the charge transfer at the surface of the PGR which is not coated by the catalyst. At the macroscopic level, the total charge transfer at this interface is given by $R_{ct,G} = r_{ct,G}/L$.

• $c_G$ is the capacitance given by the graphite/electrolyte interface. In the case of the decorated samples, it accounts for the uncoated PGR/electrolyte surface. The total capacitance of the graphite is given by $C_G = c_G \cdot L$.

• $r_{ct,cat}$ is the charge transfer resistance associated to Ni or NiFe catalysts. The total contribution to the total charge transfer resistance of the catalyst is given by $R_{ct,cat} = r_{ct,cat}/L$.

• $c_{cat}$ is the capacitance associated to the catalysts (Ni and NiFe) deposited on the PGRs and includes the contribution of their redox states. The total contribution of the catalyst capacitance is given by $C_{cat} = c_{cat} \cdot L$.

• $Z_D$ is the impedance diffusion associated with the reactive species at the diffusion layer in the solution.

With these definitions, the charge transfer resistance and the capacitance of the PGR are given by the parallel combination of the graphite and the catalyst contributions, $R_{ct}^{-1} = R_{ct,cat}^{-1} + R_{ct,G}^{-1}$ and $C_{PGR} = C_{cat} + C_G$, which are the effective values we measure.
Figure 4. Transmission line based equivalent circuits selected for IS analysis. (a) FA@PGR, (b) ED@Ni/PGR, and FA@NiFe/PGR electrodes. The meaning of the different elements is described in the main text.

The parameters obtained from the IS fitting are represented as a function of the overpotential (Fig. S6) and the interfacial overpotential (Fig. 5). Focusing on the capacitance data of the FA@PGR sample (Fig. 5a, black line), a nearly constant capacitance of \(~180\) mF·cm\(^{-2}\) corresponding to the PGR is observed, with a small peak at an \(\eta\) of 0.35 V, the overvoltage at which charge transfer activates in this sample. We associate this peak with the iron contents of FA@PGR, as a low amount of Fe was detected in this electrode by SEM microanalysis. When purposely we deposited Fe on PGR, the same peak with higher intensity was observed (Fig. S7). The baseline capacitance observed for ED@Ni/PGR is \(~150\) mF·cm\(^{-2}\), and for FA@NiFe/PGR is \(~180\) mF·cm\(^{-2}\), being both attributed to the PGR. The higher Ni coverage of PGR in the ED sample causes a decrease in the PGR capacitance, as indicated in ECSA measurement.
When the Ni and NiFe are deposited on the PGR, three capacitance peaks are observed, what suggest a more complex explanation for the peaks observed in the cyclic voltammetry of Fig. 2a. In fact, and according to the literature, the two peaks observed at low overpotentials are attributed to two different Ni$^{2+}$/Ni$^{3+}$ redox transitions. As commented before, the energy of these transitions depends on the Fe content but also to the phase segregation in the sample, which is a function of the deposition environment and the flame annealing treatment. There are two possible phases for Ni(OH)$_2$ molecules present at Ni surface: the $\beta$–Ni(OH)$_2$, which is the normal and stable phase and the $\alpha$–phase, a hydrated form of the nickel hydroxide, 3Ni(OH)$_2$·2H$_2$O. The formation of the $\alpha$–phase is related to the ability of incorporating water between the layers during the nickel deposition on the PGR. We expect that the thermal annealing conduct to a larger proportion of de-hydrated phase while electrodeposition, which is not thermally treated, becomes more hydrated. In any case, $\alpha$ to $\beta$ conversion phase could happen as a consequence of aging and temperature.

The oxidation of Ni$^{2+}$ to Ni$^{3+}$ by the application of a potential lead to the formation of $\gamma$– and $\beta$–NiOOH (hydrated and de-hydrated), respectively. According to literature, $\alpha$–Ni(OH)$_2$ oxidizes to $\gamma$–NiOOH ($\alpha/\gamma$) at $\eta = 0.12$ V ($E^0 = 1.35$ V vs RHE) and $\beta$–Ni(OH)$_2$ oxidizes to $\beta$–NiOOH ($\beta/\beta$) at $\eta = 0.20$ V ($E^0 = 1.43$ V vs RHE), which match very well with the peaks found in the capacitance of the ED@Ni/PGR electrode in Fig. 5a. This result confirms that the small amount of Fe still present in the ED sample, has a minimal effect in the Ni oxidation states, as discussed above. The larger height on the first peak indicates that the $\alpha/\gamma$ transition is the most important one in this sample and dominates the peak in the CV. The $\beta/\beta$ transition also occurs, causing the wide peak measured in the CV curve.

The third peak, that appears in Fig. 5a at the overpotential of $\sim 0.26$ V ($E^0 = 1.49$ vs RHE), corresponds to the redox transformation between Ni$^{3+}$/Ni$^{4+}$, which some studies have associated with the formation of NiOO after deprotonation of NiOOH. This peak is strongly affected by the iron concentration of the electrode and marks the starting of the quick decrease of $R_\alpha$ observed in Fig. 5c. $R_\alpha$ is related to the activation of the charge transfer mechanism that yields to the onset of the OER in the CV and the $J$-$V$ curve in Figs. 2a and 5d, respectively.
Figure 5. Results from the IS measurements data for FA@PGR (Black), ED@Ni/PGR (Green) and FA@NiFe/PGR (Blue) as a function of the interfacial overpotential (a) Electrode capacitance, (b) transport resistance, (c) charge transfer resistance and (d) J-V curve.

For the FA@NiFe/PGR, a clear peak at an overpotential of 0.19 V vs RHE is observed. We associated this peak to the β/β transitions of Ni$^{2+}$/Ni$^{3+}$ which is much larger than in the case of the ED and slightly displaced to more negative potentials. In this case, the data suggest that the α/γ transition contribution to the capacitance is much smaller than in the ED@Ni/PGR case, and only produces a distortion of the Ni$^{2+}$/Ni$^{3+}$ peak. Consequently, the peak observed in the CV of Fig. 2 is displaced towards positive values. This result matches well with results obtained in many previous papers. Therefore, these data suggest that the effect of Fe rather than displacing the Ni$^{3+}$/Ni$^{2+}$ peaks is favouring the presence of the β phases of Nickel rather than the α/γ.
For the third peak of the capacitance associated with Ni$^{3+}$/Ni$^{4+}$, now we see a reduction in the overpotential needed to make the redox transition to the Ni$^{3+}$/Ni$^{4+}$ which now occurs at 0.23 V, see Fig. 5d. Consequently, OER activates at an overpotential 30 mV smaller, in good agreement with literature data. Associated with this effect, we can see that the abrupt drop of $R_{ct}$ matches very well with the OER activation in the CV (Fig 5d). The analysis of $R_{ct}$ shows even more clearly how the onset decrease in $R_{ct}$ is displaced towards smaller overpotentials as we move from the FA@PGR sample to the ED@Ni/PGR and then to the FA@NiFe/PGR, in perfect agreement with the J-V response observed in Fig. 5d.

Non-electrochemical techniques have been used to correlate the detection of the different Ni phase transitions for Ni(OH)$_2$/NiOOH ($\alpha$/\$\gamma$ and $\beta$/\$\beta$) and the Ni$^{3+}$/Ni$^{4+}$ transition with electrochemical techniques.$^{41,45}$ To the best of our knowledge, the IS analysis made here have successfully demonstrated for the first time the suitability of this technique for that purpose, showing that IS is a low-cost, easy and useful tool to characterize these kind of samples.

Finally, the transport resistances of FA@NiFe/PGR and FA@PGR present similar values at low voltages, while the ED@Ni/PGR sample presents slightly larger values, see Fig. 5b. This result suggests that larger Fe concentrations produce larger conductivity of the Ni/PGR. Deeper analysis is needed to fully understand this behaviour, which is outside the scope of this paper. At the potentials of the Ni$^{2+}$/Ni$^{3+}$ redox transition, $R_{tr}$ diminishes until a valley is formed, which we associate to the contribution of the redox species to the overall conductivity of the film.

4. CONCLUSIONS

We present herein a low-cost procedure to obtain porous Ni-decorated electrodes based on graphite rods, that provided a reasonable performance. For the study of their electrochemical response, we proposed an impedance model based on a transmission line and made a detailed analysis of the capacitance. This allowed to deconvolute the contributions of $\alpha$-$\gamma$ and $\beta$-$\beta$ phases in the Ni$^{2+}$/Ni$^{3+}$ redox transitions. Through this analysis, we could identify the most prominent phases present in each of the electrodes measured here. Thus, we found that in the electrodeposited sample the predominant phases are $\alpha$-Ni(OH)$_2$ and $\gamma$-NiOOH associated with hydrated Ni, while in the case of the sample with flame annealing treatment, the dominant are the non-hydrated $\beta$ phases.
Rather than the Fe content, the dominant Ni-phase in the electrode is the origin of the position of the Ni$^{2+}$/Ni$^{3+}$ redox peaks found. Finally, we show the direct relationship between the capacitance peak associated to Ni$^{3+}$/Ni$^{4+}$ redox transition and the activation of charge transfer resistance and, consequently the oxygen evolution reaction.

- ASSOCIATED CONTENT

Supporting Information:

ECSA data, SEM, XPS and complementary impedance spectroscopy plots are included in supporting information

- AUTHOR CONTRIBUTIONS

YK and RA fabricated the samples and made the measurements, YK and YH made XPS analysis. RA contributed with cyclic voltammetry and impedance spectroscopy analysis. RT, HM and found the improvement attained by NiFe for flame-annealed catalysts. SI provided the initial idea of the paper, completed later by FFS and EMM. FFS made interpretation of impedance spectroscopy data. EMM and FFS directed the experiments done and selected the contents for the paper and RA wrote first version of the paper. All authors revised and improved the paper with their contributions.

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