

Cite this: *Energy Adv.*, 2024,  
3, 495

# Co-Prussian blue analogue supported on graphene-based materials as an electrocatalyst for OER at neutral pH†

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The development of efficient, robust, and low-cost electrocatalysts for the oxygen evolution reaction (OER) under mild conditions is a sustainable pathway for energy conversion and storage technologies. Cobalt Prussian blue analogue (Co-PBA) has been emerging as a potential candidate to be engineered and match such requirements. In this study, the catalytic activity performance of Co-PBA was improved by depositing over different types of graphene-based materials that act as supports, resulting in hybrid composites. The Co-PBA + r-GO N<sub>2</sub>, which is Co-PBA deposited over a reduced graphene oxide doped with N atoms (urea as N source), showed the best OER catalytic improvements in a neutral medium (0.5 mol L<sup>-1</sup> KNO<sub>3</sub>), reducing the overpotential in 54 mV and 137 mV to reach current densities of 10 (*J*<sub>10</sub>) and 50 (*J*<sub>50</sub>) mA cm<sup>-2</sup>, respectively, in comparison to the sole electrocatalyst Co-PBA. These results indicate that the interaction between the Co-PBA and graphene-based supports plays a pivotal role in effectively enhancing the OER activity of the electrocatalyst.

Received 29th September 2023,  
Accepted 25th December 2023

DOI: 10.1039/d3ya00483j

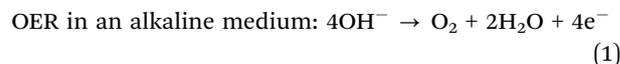
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## Introduction

The generation of renewable electricity from natural sources (such as solar, wind, and ocean) can potentially attend to the global energy demand.<sup>1</sup> One of the main issues associated with promoting energy transition from fossil fuel to intermittent renewable energy sources is mainly related to energy storage.<sup>2–4</sup> Green hydrogen emerges as a critical enabler in this energy transition scenario, where renewable electricity can be stored in chemical bonds through the electrochemically driven water-splitting reaction.<sup>2–4</sup>

The cathodic hydrogen evolution reaction (HER) and anodic oxygen evolution reaction (OER) are the two half-reactions that comprise electrochemical water splitting.<sup>5,6</sup> Despite the product of interest being generated at the cathode, the intrinsic sluggish kinetics of the OER, which is a complex multistep process that involves four proton-coupled electron transfers and O<sub>2</sub> bond formation, hamper accomplishing the overall process.<sup>6,7</sup> In addition to its application in the water splitting field, the OER is a strategic reaction in energy conversions and storage technologies, such as solar energy,<sup>8,9</sup> rechargeable metal-air batteries,<sup>10–12</sup> and regenerative fuel cells.<sup>13</sup> To minimize the

higher overpotential and kinetic barrier associated with OER, various electrocatalysts have been designed to operate under extremely alkaline pH conditions. The OER is a pH-dependent reaction that profits from the abundant OH<sup>-</sup> active species according to eqn (1).<sup>1,14–16</sup>



Nevertheless, the energy transition should be preferentially aligned with green chemistry to avoid harsh conditions that are not environmentally friendly. Moreover, performing water splitting under mild conditions may enable the use of water from natural sources and significantly reduce the corrosion of the electrolysis system.<sup>17,18</sup> Therefore, it is extremely important to develop cost-effective electrocatalysts with high activity for the OER in near-neutral media.

Cobalt-based catalysts for the OER in a neutral medium have attracted considerable appreciation for their activity, stability, and simple process of preparation from low-cost materials.<sup>5,12</sup> Cobalt-Prussian blue analogue (Co-PBA) is one example that has been displaying great performance towards the OER under mild conditions.<sup>5,6,19</sup> Despite its advantages, Co-PBA exhibits a major drawback of a low concentration of exposed electroactive cobalt sites.<sup>20</sup> A strategy to circumvent this shortcoming is the use of a Co-PBA/graphene-based composite where the graphene-based materials act as a support providing a boost to the electroactive sites.<sup>21</sup> Additionally, nitrogen doping on

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3ya00483j>



graphene-based materials can effectively impact their properties such as conductivity and charge transfer. Accommodating electrocatalysts over nitrogen-doped graphene-based supports has been reported as an effective way to improve the distribution of metal nanoparticles, enhancing mass transfer and boosting the electrocatalytic performance toward water splitting.<sup>22,23</sup>

Herein, we report a simple and facile strategy to improve the performance of the Co-PBA electrocatalyst toward the OER at neutral pH by depositing over different types of graphene-based materials that act as a support, resulting in hybrid composites. Furthermore, we aim to understand the role of the graphene-based supports in the electrocatalyst activity and how we can modulate this interaction to achieve outstanding performances in the resulting hybrid composite.

## Results and discussion

### Graphene-based supports

**Elemental CHN analysis.** In a brief description of the synthesis strategy applied in this work, graphene oxide (GO) was first synthesized as a precursor for all graphene-based samples. The other 4 derived samples were obtained by subjecting GO to hydrothermal treatment in an autoclave. This treatment simultaneously provides a partial reduction process for GO and a method for doping with N atoms. Therefore, from GO, we synthesized r\_GO, which is GO partially reduced, and r\_GO N1, r\_GO N2, and r\_GO N3, which are GO partially reduced and doped with nitrogen from different sources.

All the graphene-based samples were subjected to CHN elemental analysis to determine their composition and further confirm the presence of N atoms in their structures. Table 1 lists the elemental compositions of all graphene-based samples. As expected, GO has the highest oxygen content (57.96 wt%) and lowest carbon content (38.53 wt%). Upon hydrothermal reduction, the carbon content of the derived samples tended to increase while their oxygen content decreased, as evidenced by their C/O ratios (Table 1). These results confirm that the hydrothermal treatment partially removed some of the oxygen functionalities. In addition, the analysis confirms the successful doping of N atoms on r\_GO N1, r\_GO N2, and r\_GO N3, which were used stoichiometrically with 3 different sources of nitrogen (ammonium hydroxide, urea, and hydrazine, respectively). As can be seen in Table 1, all 3 N-doped samples have relatively similar nitrogen content.

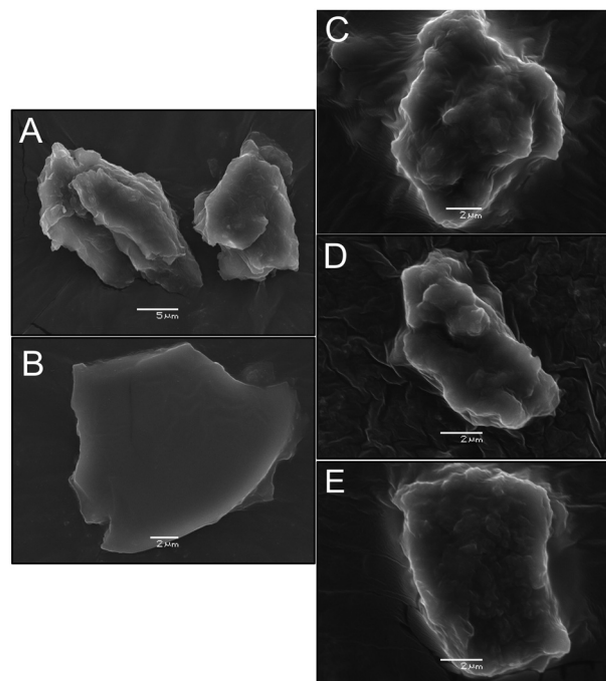
**Table 1** Elemental quantification of graphene-based materials from CHN elemental analysis

Sample	C (wt%)	O <sup>a</sup> (wt%)	N (wt%)	H (wt%)	C/O	C/N	Nitrogen source
GO	38.53	57.96	—	3.51	0.66	—	—
r_GO	67.73	30.73	—	1.54	2.20	—	—
r_GO N1	63.11	27.21	8.31	1.37	2.33	7.59	NH <sub>4</sub> OH
r_GO N2	63.64	26.89	7.66	1.81	2.35	8.31	(NH <sub>2</sub> ) <sub>2</sub> CO (urea)
r_GO N3	69.27	22.3	7.42	1.01	3.10	9.33	N <sub>2</sub> H <sub>2</sub>

<sup>a</sup> By difference.

Although r\_GO N1 has the highest nitrogen content, its elemental composition is similar to that of the r\_GO N2. A trend observed in the N-doped materials is that a higher nitrogen content results in a lower C content, and consequently a lower C/N ratio. These results also indicate that starting from the same precursor (GO) and varying the type of nitrogen source leads to the formation of different materials. r\_GO N3 is noteworthy due to its highest carbon content and lowest oxygen and nitrogen contents. We believe that this is related to the source of nitrogen used in the r\_GO N3 synthesis, hydrazine, which is a strong reducing agent. As in the synthesis of the N-doped graphene-based materials, the precursor, GO, was previously stirred for 2 hours with the respective nitrogen source before the hydrothermal treatment; thus, r\_GO N3 was first subjected to a chemical environment that lowered the oxygen content prior to the hydrothermal treatment. This prior removal of oxygen functionalities over the precursor material may also hinder the insertion of N atoms since it is expected that the N atoms are incorporated by the carbon lattice when they react with oxygenated functionalities during the hydrothermal treatment.<sup>24</sup> Thus, in this approach, r\_GO N3 already underwent the hydrothermal treatment with a relatively small number of oxygenated groups, leading to the material with the highest degree of reduction and lowest N content.

**Scanning electron microscopy (SEM).** The morphologies of the graphene-based materials were obtained by SEM and are presented in Fig. 1. GO (Fig. 1A) showed characteristic exfoliated structures in concordance with previous studies.<sup>25</sup> It is also possible to evidence that GO exhibits overlapped structures



**Fig. 1** SEM images of the graphene-based samples. The scale bars are shown in each figure. Images are (A) GO; (B) r\_GO; (C) r\_GO N1; (D) r\_GO N2; (E) r\_GO N3.



and wrinkled sheet morphology. r\_GO (Fig. 1B) showed fewer stacked layers with a considerably smoother aspect than all other samples. All samples of the N-doped materials (Fig. 1C–E) showed wrinkled areas, with r\_GO N1 and r\_GO N2 (Fig. 1C and D) exhibiting more kinked areas.

**Spectroscopic analysis.** FTIR analysis was performed to investigate the presence of oxygenated groups and the extent of the reduction and insertion of nitrogen atoms in the graphene-based materials. Their FTIR spectra are presented in Fig. 2. Several characteristic bands of oxygenated groups at  $3400\text{ cm}^{-1}$  (O–H stretching),  $1735\text{ cm}^{-1}$  (C=O stretching),  $1402\text{ cm}^{-1}$  (OH deformation), and  $1224\text{ cm}^{-1}$  (epoxy C–O stretching) can be observed mainly in GO.<sup>26,27</sup> The samples that were subjected to hydrothermal treatment exhibited smaller intensities of the bands associated with oxygenated groups, mostly  $3400\text{ cm}^{-1}$ , which agrees with the reduction process. The bands at  $2930$  and  $2853\text{ cm}^{-1}$  correspond to the  $\text{CH}_2$  symmetric and anti-symmetric stretching vibration respectively,<sup>26</sup> while the bands at  $1630\text{ cm}^{-1}$  can be correlated to the skeletal vibration of graphitic domains.<sup>28</sup> For the samples that were N-doped, the bands located at  $1570\text{ cm}^{-1}$  correspond to the carbon–nitrogen bond.<sup>26</sup>

Raman spectroscopy analyses were done to investigate the structural features of graphene-based materials. Their Raman spectra (Fig. 3) exhibit two distinct bands, D ( $1346\text{--}1349\text{ cm}^{-1}$ ) and G ( $1592\text{--}1600\text{ cm}^{-1}$ ). The D band arises due to disorder and structural defects caused by the  $\text{sp}^3$  hybridized carbon atoms, while the G band is commonly observed in all graphitic structures and occurs due to the doubly degenerate phonon mode of the  $\text{sp}^2$  carbon network.<sup>29</sup>

The intensity ratio of the D and G bands ( $I_d/I_g$ ) is associated with the amount of structural defects, disorder, and edge plane exposure of the graphene sheets.<sup>30</sup> The increase in the  $I_d/I_g$  values shown in Fig. 3 indicates that the hydrothermal reduction and nitrogen doping possibly resulted in materials with a higher  $\text{sp}^3/\text{sp}^2$  carbon ratio, more defects, disorder, and edge sites in comparison to the precursor material.<sup>30,31</sup> Also, the Raman spectra evaluated the positions of the D and G bands exhibited in Table S1 (ESI†). The downshift observed in

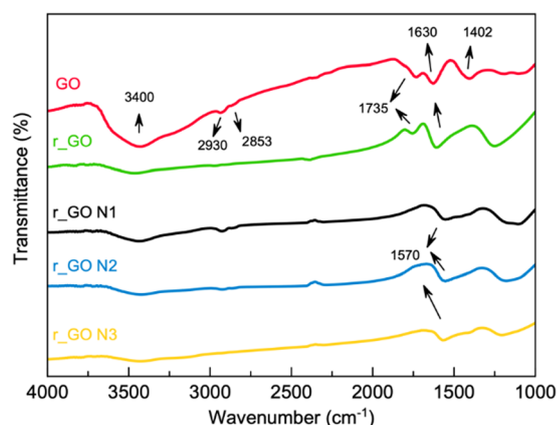


Fig. 2 IR spectra of the graphene-based samples.

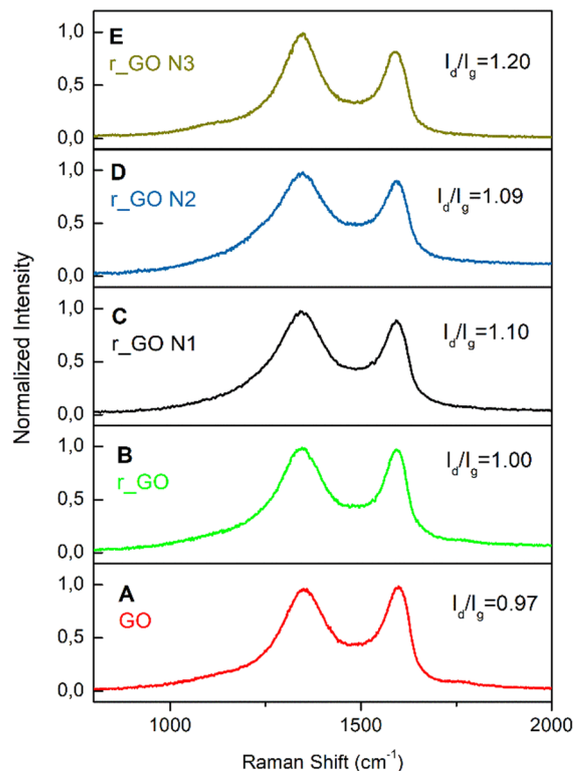


Fig. 3 Raman spectra of the graphene-based samples with their respective  $I_d/I_g$  ratios. These spectra correspond to (A) GO; (B) r\_GO; (C) r\_GO N1; (D) r\_GO N2; (E) r\_GO N3.

the G band from the precursor to the derived samples agrees with previous studies and is indicative of reduction and/or doping.<sup>32</sup> This downshift may be explained by the lower content of oxygen bonded to the carbon skeleton, providing the carbon atoms a bond less rigid in its stretch mode and consequently lower energy.

### Co-PBA electrocatalyst

**Scanning electron microscopy (SEM).** The morphology of the as-prepared Co-PBA material was investigated by SEM. The SEM images (Fig. 4) reveal the occurrence of the Co-PBA particles with sizes ranging from  $1\text{--}2\text{ }\mu\text{m}$ , a cubic-shaped structure, and some of them eventually exhibit a disordered morphology.

**Catalyst structure.** The nanostructure of the cobalt Prussian blue analogue (Co-PBA) was analyzed by X-ray powder diffraction (XRD). Fig. 5 shows the diffraction pattern of the Co-PBA synthesized and we can observe characteristic diffraction peaks at  $17.1$ ,  $24.3$ ,  $34.9$ ,  $39.2$ ,  $43.2$ ,  $50.3$ ,  $53.6$ , and  $56.8^\circ$  corresponding to the (200), (220), (400), (420), (422), (440), (422) and (620) planes of the face-centered cubic (fcc) structure of Prussian blue (PDF number 72-1431), respectively. Furthermore, the pattern analysis of the Co-PBA, using Bragg's law, allows the determination of the interlayer distance to be about  $0.518\text{ nm}$  for the (200) crystallographic plane. In addition, the crystalline dimension ( $L_{hkl}$ ) was calculated using the Scherrer equation,<sup>33</sup> and the thickness was found to be about  $14.4\text{ nm}$



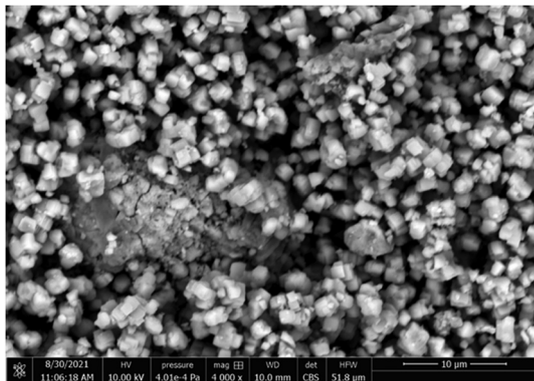


Fig. 4 SEM image of Co-PBA.

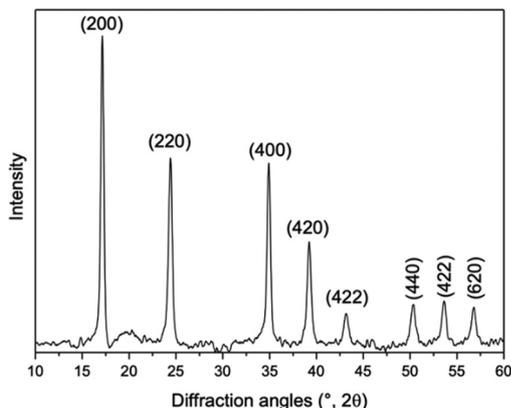


Fig. 5 X-ray powder diffraction (XRD) pattern of the cobalt Prussian blue analogue (Co-PBA) with the crystallographic planes (PDF number 72-1431).

for the most intense plane. Fig. S1 (ESI<sup>†</sup>) shows a schematic of the structure of Co-PBA.

**Spectroscopic analysis.** Raman and infrared spectroscopies were also used to characterize the Co-PBA material (Fig. 6). The interpretation of the stretching  $\nu(\text{CN})$  ligand plays a major role in identifying Prussian blue analogues since these ligands act as linkers between 2 metallic centers. This characteristic band commonly occurs in the interval range between 2100–2200  $\text{cm}^{-1}$  in both Raman and infrared spectra.<sup>34</sup> By analyzing both spectra in Fig. 6, we confirm the characteristic band of Prussian blue analogues. In detail, the Raman spectra of the bands at 2201  $\text{cm}^{-1}$  and 2183  $\text{cm}^{-1}$  are assigned to the symmetric stretching mode ( $A_{1g}$ ) and  $E_g$  symmetry, respectively, of  $\text{Co}^{\text{III}}\text{-CN-Co}^{\text{II}}$ .<sup>6</sup> In the low-frequency region, the band at 491  $\text{cm}^{-1}$  refers to the deformation  $T_{2g}$  modes of the  $\text{Co}^{\text{III}}\text{-CN-Co}^{\text{II}}$  system.<sup>6</sup> In the infrared spectra, the band around 2167  $\text{cm}^{-1}$  is assigned to the stretching mode ( $T_{1u}$ ). The band at 1418  $\text{cm}^{-1}$  is correlated to the C–O vibrations of carboxyl groups, suggesting the existence of coordinated citrate in the vacant sites that are usually occupied by the water molecules,<sup>35</sup> while the band around 447  $\text{cm}^{-1}$  is correlated to the stretching mode M–C.<sup>6,34</sup>

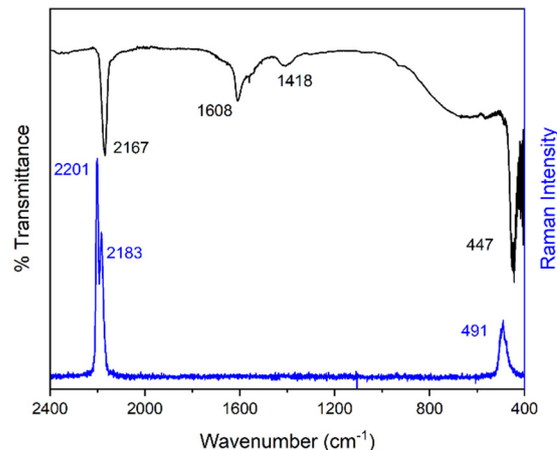


Fig. 6 Raman and infrared spectra of Co-PBA.

### Electrochemical parameters

To evaluate the electrochemical parameters of the materials, cyclic voltammograms (CV) of the modified GCE with the described materials were obtained using a solution containing 5  $\text{mmol L}^{-1}$   $[\text{Ru}(\text{NH}_3)_6]^{3+}$  and 0.1  $\text{mmol L}^{-1}$   $\text{KNO}_3$  as the redox probe and support electrolyte respectively. The voltammograms and parameters obtained from the experiments are presented in Fig. S2 (ESI<sup>†</sup>) and Table 2, respectively.

Among all the materials, Co-PBA exhibited the highest  $k_{\text{obs}}^\circ$ , which was expected due to the presence of the metallic centers. Regarding the comparison of the graphene-based materials, GO and r\_GO N3 had higher values for  $k_{\text{obs}}^\circ$ , as more defined peaks can be seen in their voltammograms (Fig. S2(A) and (F), ESI<sup>†</sup>). We suppose that this is correlated to the higher amount of oxygen functionalities present on GO, as O atoms have relatively high electronegativity and easily get a partial negative charge, thus increasing the interaction and approximation with the redox probe,  $[\text{Ru}(\text{NH}_3)_6]^{3+}$ , which is a positive species. However, the electrochemical behavior of r\_GO N3 could be caused by the highest intensity ratio of the D and G bands ( $I_d/I_g$ ), which indicates that the material may have more edge sites, where electronic transfer is faster.<sup>36,37</sup> For the other materials, r\_GO, r\_GO N1 and r\_GO N2, their voltammograms (Fig. S2(C)–(E), ESI<sup>†</sup>) reveal capacitive features.

Linear sweep voltammetry (LSV) in a 0.1  $\text{mol L}^{-1}$   $\text{KNO}_3$  solution at pH 7 was performed to analyze the oxygen evolution reaction (OER) activity of the materials (Fig. 7A). The electrocatalytic information obtained from the LSV curves is presented in Table 2. As expected, Co-PBA has the lowest overpotentials of 590 mV and 980 mV to reach current densities of 1  $\text{mA cm}^{-2}$  ( $J_1$ ) and 10  $\text{mA cm}^{-2}$  ( $J_{10}$ ), respectively. In the graphene-based group, GO and r\_GO N3 exhibited lower overpotentials to reach  $J_{10}$ , mostly according to the same trend observed in their  $k_{\text{obs}}^\circ$  values.

Tafel plots were used to assess the kinetic behavior of the materials toward OER. This allowed us to obtain information regarding how the current density increased with the applied overpotential.<sup>7</sup> The Tafel plots and slopes are presented in



Table 2 Electrochemical parameters for Co-PBA and graphene-based materials

Sample	Electrochemical properties		OER activity		
	ECSA $10^{-2}$ (cm <sup>2</sup> )	$K_{\text{obs}}$ $10^{-3}$ (cm s <sup>-1</sup> )	Tafel (mV dec <sup>-1</sup> )	$\eta$ (mV) $j = 1$ mA cm <sup>-2</sup>	$\eta$ (mV) $j = 10$ mA cm <sup>-2</sup>
Co-PBA	4.94	3.01	188	590	980
GO	5.26	1.70	174	670	1040
r_GO	3.99	0.62	196	660	1060
r_GO N1	3.90	0.77	210	700	1070
r_GO N2	3.81	0.91	217	770	1090
r_GO N3	3.89	2.01	208	660	1050

Fig. 7B and Table 2, respectively. The kinetic behavior of the materials was evaluated in the interval range between  $J_1$  and  $J_{10}$ , where the analyzed variables behaved linearly. Examining the Tafel slope values in Table 2, GO displays the lowest value followed by Co-PBA, indicating that both materials have faster kinetics and a relatively similar mechanism. Also, the Tafel slope values reveal that subjecting GO to hydrothermal treatment and doping with N atoms results in materials with a slower mechanistic pathway. As can be verified in Table 2, higher values of the Tafel slope were obtained for all materials obtained from GO.

### OER activity of the hybrid composites

In an attempt to improve the performance of the Co-PBA electrocatalyst, we first deposited the graphene-based materials

onto the surface of the GCE to act as a support, followed by the deposition of Co-PBA over the graphene-based depositions. Thereby, the Co-PBA remains on the interface with the electrolytic solution to act as the electrocatalyst. For further comparison with the previous electrochemical experiments, the total amount of material deposited on the GCE surface was maintained. The results and electrochemical parameters obtained for the hybrid composites are presented in Fig. 8A–C and Table 3.

The LSV curves (Fig. 8A) reveal that all the hybrid composites could reach  $J_{10}$  and  $J_{50}$  at lower potentials than the sole Co-PBA. The ECSA of Co-PBA was used to normalize the current density of all hybrid composites due to the presence of Co-PBA in all hybrid composites and in direct contact with the electrode/solution interface. Table 3 summarizes the variation between the hybrid composites and sole Co-PBA to reach  $J_{10}$  and  $J_{50}$ . We represent the term  $\Delta J_{10}$  as the difference between the composite and sole Co-PBA to reach  $J_{10}$  ( $\Delta J_{10} = \text{voltage at } J_{10} \text{ of the composite} - \text{voltage at } J_{10} \text{ of the sole Co-PBA}$ ). Almost all hybrid composites could reach  $J_{10}$ , needing less than 50 mV in comparison to sole Co-PBA, except Co-PBA + r\_GO N1, which required less than 39 mV. Moreover, we could achieve significant progress at a larger current density of 50 mA cm<sup>-2</sup> ( $J_{50}$ ), as listed in Table 3. Except for Co-PBA + r\_GO N1, all the hybrid composites could be reduced by more than 100 mV to reach  $J_{50}$ . Notably, Co-PBA + r\_GO N2 and Co-PBA + r\_GO N3 exhibited the best performances, reducing 137 and 136 mV, respectively.

These results clearly indicate that the graphene-based supports play a pivotal role in the electrocatalytic performance of Co-PBA. The lower overpotentials observed for the hybrid composites to reach  $J_{10}$  and  $J_{50}$  may indicate a lower energy barrier. In electrocatalytic processes that use cobalt-based catalysts, it is assumed that Co<sup>3+</sup> is the main active site involved in the catalytic steps such as adsorption, activation, and desorption.<sup>38</sup> However, the Co<sup>3+</sup> active site on Co-PBA is obtained from the oxidation of Co<sup>2+</sup> present in its sphere of external coordination, Co<sub>3</sub>[Co(CN)<sub>6</sub>]<sub>2</sub>, where is also located some vacancies on the structure mainly occupied by water molecules, a feature of Prussian blue analogues.<sup>39</sup> The Co<sup>3+</sup> in the sphere of internal coordination is more related to providing stabilization to the structure. Therefore, when the GCE was modified with the hybrid composite, the presence of the graphene-based support assisted in obtaining the Co<sup>3+</sup> active sites on Co-PBA at lower potentials in comparison to the sole Co-PBA, as can be seen in Fig. 8B. Also, it is noteworthy

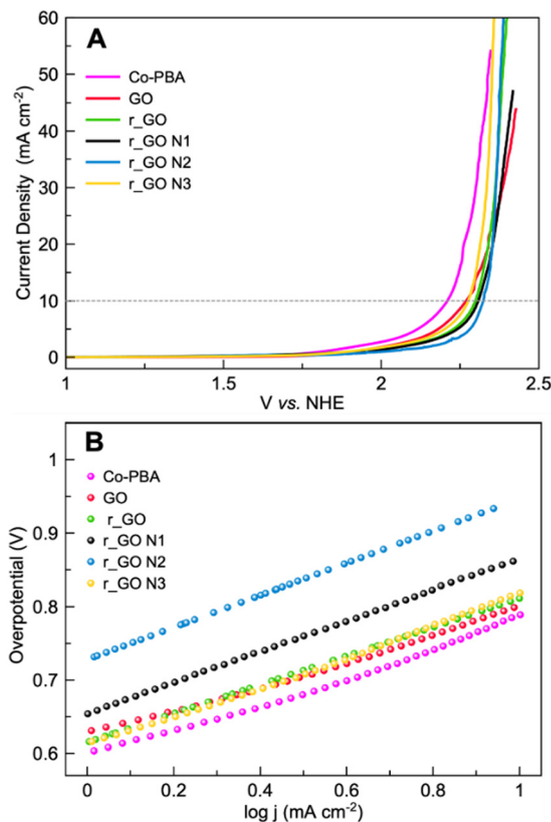


Fig. 7 Electrochemical experiments of Co-PBA and graphene-based materials performed in 0.5 mol L<sup>-1</sup> KNO<sub>3</sub> at a scan rate of 5 mV s<sup>-1</sup>. (A) Linear sweep voltammetry (LSV) and (B) Tafel plot.



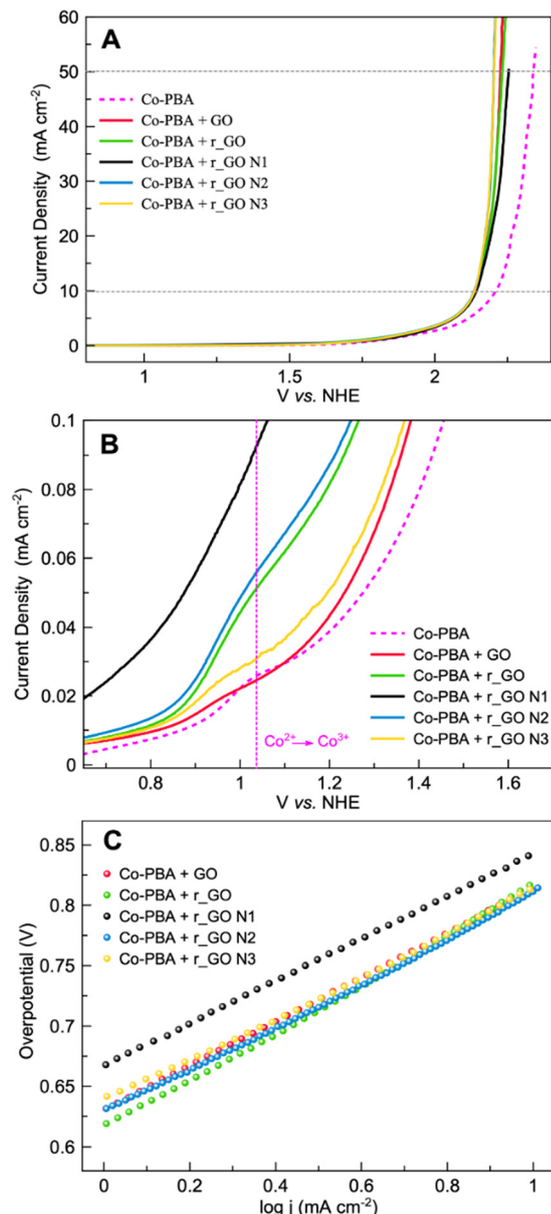


Fig. 8 Electrochemical experiments of the hybrid composites performed in  $0.5 \text{ mol L}^{-1} \text{ KNO}_3$  at a scan rate of  $5 \text{ mV s}^{-1}$ . (A) Linear sweep voltammetry (LSV), (B) the redox process  $\text{Co}^{2+}/\text{Co}^{3+}$  and (C) the Tafel plot.

Table 3 Electrochemical parameters of the hybrid composites

	Support	$\Delta J_{10}$ (mV)	$\Delta J_{50}$ (mV)	Tafel ( $\text{mV dec}^{-1}$ )
Co-PBA	GO	-52	-115	183
	r_GO	-51	-107	200
	r_GO N1	-39	-85	175
	r_GO N2	-54	-137	180
	r_GO N3	-51	-136	175

$\Delta J_{10}$  = (voltage of the composite – voltage of Co-PBA) at  $J_{10}$ .  $\Delta J_{50}$  = (voltage of the composite – voltage of Co-PBA) at  $J_{50}$ .

that the hybrid composites start polarization earlier than the sole Co-PBA, especially the supports derived from GO, which is

related to the capacitive properties of these graphene-based materials. As the potential is applied, these supports accumulate charge behaving similarly to a capacitor. Thus, the charged supports interact with the electrocatalyst resulting in a boost in the active sites and lowering the potential of the  $\text{Co}^{2+}/\text{Co}^{3+}$  redox process, which may help the water molecules to undergo the nucleophilic attack to proceed with the OER.

According to the Tafel slopes obtained for the hybrid composites (Fig. 8C and Table 3), the presence of graphene-based supports can also improve the kinetics of the reaction. As can be seen, most hybrid composites had lower Tafel slope values than the sole Co-PBA (except Co-PBA + r\_GO). In addition to these Tafel slopes being lower, they are relatively close to the Tafel slope of sole Co-PBA, which may indicate that even in the presence of the support, the kinetics of the reaction could continue through a similar mechanism performed by the sole Co-PBA. To compare the performance of the hybrid composites used in this work, Table S2 (ESI<sup>†</sup>) exhibits the electrochemical parameters of some electrocatalysts reported in the literature under mild conditions.

## Conclusions

In summary, an effective way to improve the electrocatalytic performance of Co-PBA toward the OER in a neutral medium is presented. Hydrothermal treatment could partially reduce graphene oxide and dope with nitrogen stoichiometrically considering different sources of nitrogen (ammonium hydroxide, urea, and hydrazine), resulting in different materials. These graphene-based materials were designed to act as supports to boost the electroactive sites of the electrocatalyst. Overall, all hybrid composites could reduce the potential to reach  $J_{10}$  and  $J_{50}$  compared to sole Co-PBA. Moreover, by analyzing the kinetic parameters, all hybrid composites with only one exception, Co-PBA + r\_GO, could still improve the kinetics of the reaction. This improvement, especially at  $J_{50}$ , may be seen as progress in the field of the OER at neutral pH (mild conditions) since the materials used in the hybrid composites are relatively cheap and easy to synthesize/work. In addition, it is worth mentioning that by using hybrid composites, we could reduce the amount of the electrocatalyst by 50% and still obtain better results. Further research is still necessary to better understand the interaction between the support and electrocatalyst; however, this approach has proven to be an efficient way to help the electrocatalyst achieve outstanding OER performance in a neutral medium.

## Experimental materials

Graphite powder (100%) was obtained from Fischer Chemicals (Fair Lawn, NJ, USA). Sulfuric acid,  $\text{H}_2\text{SO}_4$  (95–98.0%), potassium permanganate,  $\text{KMnO}_4$  (99.0%), phosphoric acid,  $\text{H}_3\text{PO}_4$  (85.0%), hydrogen peroxide aqueous solution,  $\text{H}_2\text{O}_2$  (29.0%), hydrochloric acid,  $\text{HCl}$  (37–38.0%), sodium hydroxide,  $\text{NaOH}$  (97.0%), urea,  $\text{CH}_4\text{N}_2\text{O}$  (99.5%), ammonium hydroxide,  $\text{NH}_4\text{OH}$



(28.0%) and tri-sodium citrate dihydrate,  $C_6H_5Na_3O_7 \cdot 2H_2O$  (99.0%) were purchased from Synth (Diadema, SP, Brazil). Potassium nitrate,  $KNO_3$  ( $\geq 99.0\%$ ), potassium hexacyanocobaltate(III),  $K_3[Co(CN)_6]$  ( $> 97.0\%$ ), cobalt acetate tetrahydrate,  $Co(CH_3COO)_2 \cdot 4H_2O$  ( $\geq 99.0\%$ ), Nafion perfluorinated (5 wt% in lower aliphatic alcohols and water), hexaammineruthenium(III) chloride,  $[Ru(NH_3)_6]Cl_3$  (98.0%), hydrazine monohydrate,  $N_2H_4$  64–65.0%,  $N_2H_4 \cdot H_2O$  (98.0%) and cellulose membrane (avg. flat width: 76 mm, MW 12.121) were provided by Sigma-Aldrich (USA). Ethanol P.A. was obtained from LS Chemicals. All aqueous solutions were prepared using ultrapure water (18.2 M $\Omega$  cm), obtained from a Milli-Q Plus system (Merck Millipore, USA). All the above chemicals were of analytical grade and used without any further purification.

### Synthetic procedures

**Graphene oxide (GO).** Graphene oxide (GO) solution was synthesized from graphite powder by the improved Hummers' method<sup>40</sup> and stored in a diluted solution ( $\approx 2$  mg mL<sup>-1</sup>). To obtain GO powder, the collected volume of the diluted GO solution was left to dry at 40 °C for 24 h.

**Reduced graphene oxide (r\_GO) and N-doped reduced graphene oxides (r\_GO N1, r\_GO N2 and r\_GO N3).** GO was the precursor in all the syntheses of graphene-based materials. To prepare N-doped reduced graphene oxide samples (r\_GO N1, r\_GO N2 and r\_GO N3), 3 different sources of nitrogen ( $NH_4OH$ ,  $CH_4N_2O$ , and  $N_2H_4 \cdot H_2O$  respectively) were used stoichiometrically. First, 200 mg of GO was dispersed in 19 mL of water and sonicated for 1 hour. Then, 4 mL of ammonium hydroxide solution was added and the mixture was stirred for 1 hour. The reaction mixture was sonicated again for 10 minutes and transferred to a 25 mL autoclave for hydrothermal reaction at 150 °C for 24 h. The resulting solid was collected by centrifugation and washed once with 5% HCl (v/v). Then, it was washed several times with DI water until the pH reached 7.0, and the mixture was left to dry at 65 °C for 12 h. To prepare r\_GO N2 and r\_GO N3, we used the same procedure mentioned above, with urea (6.20 g) and hydrazine monohydrate (3.85 mL) as nitrogen sources, respectively. The pH was adjusted to 10 with the addition of drops of a 0.5 mol L<sup>-1</sup> NaOH solution. To prepare r\_GO N1, it was unnecessary to adjust the pH. To prepare r\_GO, we used the same procedure described above but with no further addition of any nitrogen source.

**$Co_3[Co(CN)_6]$  cobalt Prussian blue analogue (Co-PBA).** Co-PBA was prepared following a typical synthesis used in the literature.<sup>35</sup> First, 2.5 mmol of  $Co(CH_3COO)_2 \cdot 4H_2O$  was dissolved in 25 mL of DI water in the presence of 5 mmol of sodium citrate ( $C_6H_5Na_3O_7 \cdot 2H_2O$ ), forming solution A. In this approach, the molar ratio of citrate/ $Co^{2+}$  was 2. Meanwhile, 1.67 mmol of  $K_3[Co(CN)_6]$  was dissolved in 25 mL of DI to prepare Solution B. Solution A was then slowly added to Solution B with magnetic stirring for 3 h. The precipitate was centrifuged, washed (3 times) with DI water and absolute ethanol, and dried at 65 °C for 12 h.

### Characterization methods

The morphologies of the graphene-based materials and Co-PBA were characterized using a Jeol JSM-6360 LV scanning electron

microscope and a Quanta 250 field emission scanning electron microscope, respectively. X-ray diffractograms (XRD) were obtained using a Shimadzu 7000 XRD diffractometer with  $CuK\alpha_1$  radiation ( $\lambda = 1.5418$  Å) in the  $2\theta$  range of 10–60° at a scan rate of 2° min<sup>-1</sup>. Raman spectra were obtained using a Confocal T64000 spectrometer (Jobin Yvon, USA) with a solid-state sapphire laser of 532 nm. Elemental analyses (CHN) were carried out on a PerkinElmer 2400 series ii instrument based on the Pregl–Dumas method. The oxygen content was estimated from the difference between 100% and the sum of the C + H + N%, assuming that no ash was present. ATR-FTIR spectra were recorded on an Agilent Cary 630 in a range of 400 to 4000 cm<sup>-1</sup>, and the FTIR spectra of the graphene-based samples were obtained using standard KBr pellets (0.5 mg sample + 100 mg KBr).

### Working electrode modification

The glass carbon electrode (GCE) was polished with 1.0, 0.5 and 0.3 μm alumina slurry on a polishing cloth, rinsed with DI water, and then immersed in ethanol for 10 minutes. The suspensions were prepared with 4.5 mg of the samples dispersed in 1.1 mL of ethanol, 360 μL of DMF, and 30 μL of Nafion, resulting in a concentration of  $\approx 3$  mg mL<sup>-1</sup>. The prepared sample dispersions were sonicated for 2 hours and subsequently, 4 depositions of 2 μL of suspension were dropped onto the surface area of GCE (area = 0.071 cm<sup>2</sup>, resulting in a catalyst load of  $\approx 0.34$  mg cm<sup>-2</sup>). The interval between each deposition was 30 minutes and then the samples were allowed to dry at room temperature for over 12 h. To obtain the hybrid composite, we performed the same procedure as described above with the first 2 depositions of a graphene-based material to act as a substrate, followed by 2 depositions of the Co-PBA catalyst.

### Electrochemical measurements

All electrochemical experiments were performed at room temperature (25 °C  $\pm$  2 °C) using a Metrohm Autolab potentiostat PGSTAT10302N, connected to a PC. A standard three-electrode electrochemical cell was used with modified GCE as the working electrode, saturated calomel electrode (SCE) as the reference electrode, and platinum wire as the counter electrode. Prior to the experiments, the electrolyte solutions were purged with  $N_2$  for 15 minutes.

### Calculation methods

The calculations used to determine the electrochemical properties and electrocatalytic parameter activities are described next.

The ECSA of the samples were obtained according to the Randles–Ševčík equation,<sup>41–43</sup> where  $I_p$  is the peak current,  $n$  is the number of electrons involved in the electrochemical process,  $A_{ECSA}$  is the electrochemical surface area,  $D$  is the diffusion coefficient ( $9.10 \times 10^{-6}$  cm<sup>2</sup> s<sup>-1</sup> for  $[Ru(NH_3)_6]^{3+}$ ),  $C$  is the concentration of the redox probe and  $\nu$  is the scan rate:

$$I_p = 2.69 \times 10^5 n^{\frac{3}{2}} A_{ECSA} D^{\frac{1}{2}} C \nu^{\frac{1}{2}}$$



$K_{\text{obs}}$  was obtained by the Nicholson method<sup>41–43</sup> using the following equation:

$$K_{\text{obs}} = \left[ 2.18 \left( \frac{D\alpha n F \nu}{RT} \right)^{\frac{1}{2}} \right] \exp \left[ -\frac{\alpha^2 n F}{RT} \right] \Delta E_p$$

where  $D$  is the diffusion coefficient,  $\alpha$  is the charge transfer coefficient (assumed to correspond to 0.5),  $n$  is the number of electrons involved in the electrochemical process,  $F$  is the Faraday constant,  $\nu$  is the scan rate,  $R$  is the gas constant,  $T$  is the temperature and  $\Delta E_p$  is the peak-to-peak separation:

The  $iR$  drop correction of 90% was carried out using the positive feedback compensation method, provided by the electrochemical workstation in the LSV experiments used to evaluate the OER activity. The Tafel plots were extracted from the LSV curves with 100%  $iR$  correction. The reference electrode (SCE) potentials were converted into normal hydrogen electrode (NHE) potentials according to the following equation:

$$E \text{ vs. NHE} = E \text{ vs. SCE} + 0.224$$

## Author contributions

Rafael G. Yoshimura – investigation, formal analysis, data curation, writing – original draft, review & editing; Thiago V. de B. Ferraz – formal analysis, writing – review & editing; Priscilla J. Zambiasi – formal analysis, writing – review & editing; Juliano A. Bonacin – supervision, resources, visualization, writing – review & editing.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

All authors are grateful for financial support of Brazilian Funding Agencies. This study was financed in part by the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior – Brasil (CAPES) – Finance Code 001 and Process 88887.339135/2019-00, Conselho Nacional de Desenvolvimento Científico e Tecnológico CNPq (308203/2021-6) and Fundação de Amparo à Pesquisa do Estado de São Paulo, FAPESP (grant# 2017/11986-5, grant# 2021/05976-2).

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