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Enhanced electrocatalytic performance triggered by atomically bridged boron nitride between palladium nanoparticles and carbon fibers in gas-diffusion electrodes

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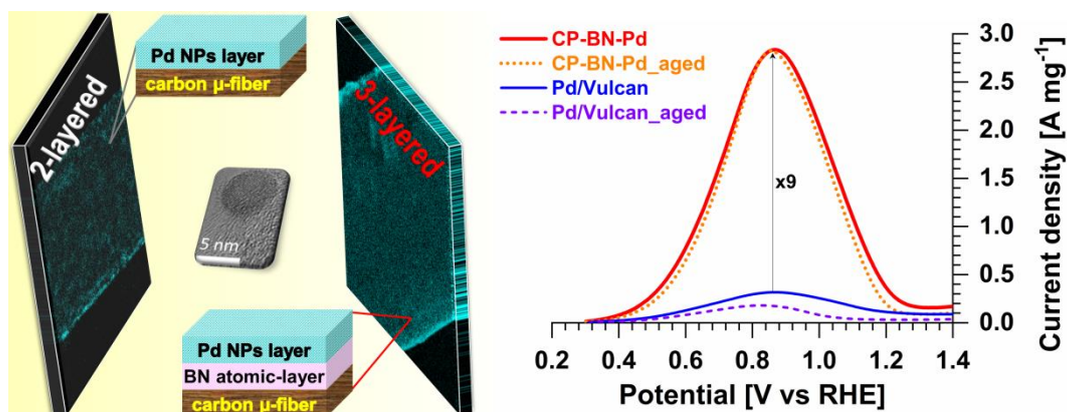
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Graphical Abstract



Highlights:

- Decrease catalysts loading below ten micrograms per square centimeter of electrode
- Bridging boron nitride between palladium and carbon foster kinetics and stability
- Multifunctional catalyst for both oxygen reduction and C2 alcohol electrooxidation
- Atomic layer deposition enables to engineer highly active palladium nanocomposites

Abstract:

Lowering significantly the amount of precious metals in catalysts while keeping the same or even increasing performances is a major contemporary challenge in the community of heterogeneous electrocatalysis. Herein, we report advanced carbon paper-boron nitride-palladium (CP-BN-Pd) electrocatalytic electrodes fabricated using the atomic layer deposition (ALD) route, that exhibit a drastic enhancement of kinetics and stability. The nanocatalysts are based on highly dispersed Pd nanoparticles (NPs) of 3 to 7 nm supported on a tunable 1-4 nm BN film covering each microfibr of a carbon paper (CP) as gas-diffusion layer electrodes (GDL). The strategy enables the fabrication of freestanding electrocatalysts easy-to-handle

and ready-to-use without any additional steps of catalytic ink preparation involving the addition of binders. The presence of the BN nanolayer leads to electrodes of $5.3 \mu\text{g}_{\text{Pd}}/\text{cm}^2$ presenting significantly enhanced performance towards the reactions of oxygen reduction (ORR) and C2 alcohols oxidation (ethanol, ethylene glycol). The results indicate that CP-BN-Pd outperforms the reported data for those alcohols in alkaline media, reaching a peak current of 17 amps per milligrams of Pd in 1 M NaOH + 1 M ethanol. Importantly, the accelerated ageing reveals that the electrochemically active surface area (ECSA) of CP-BN-Pd varies from 54 (1st cycle) to 46 (1000th cycle) m^2/g . Specifically, in the presence of BN, ethanol oxidation activity shows a negligible decay of 1% while the sample without BN has a significant decay of 11% and commercial Pd/C a prominent activity decay of 44%. The enhanced performance is due to the presence of the BN nanolayer sandwiched between active Pd NPs and the excellent electrical conducting carbon microfibers of GDL. This layer effectively creates a heterogeneous active interface as well as excellent active site-supporting material interaction and reduces aggregation/detachment of palladium NPs. Those results open up prospects in heterogeneous catalysis for the development of low-metal content nanocatalysts.

Keywords: atomic layer deposition, boron nitride, palladium nanoparticles, oxygen reduction reaction, ethanol electrooxidation reaction

1. Introduction

The electrochemical energy conversion and storage devices of fuel cells (FCs) and electrolysis cells (ECs) are rapidly growing as part of X-to-Power and Power-to-X energy transition scenarios [1-3]. The systems utilizing anion exchange membrane (AEM) have emerged in recent years as possible low-cost energy devices resulting from the fact that alkaline media overcomes the existing cost barriers of proton exchange membrane (PEM) [4-8]. Oxygen reduction reaction (ORR) is the key cathode reaction for both FCs and metal-air batteries. The anode oxidation of biomass-based compounds such as ethanol and glycerol (less toxic, cheaper, easier to store and handle) emerges as a sustainable and attractive alternative in FCs to produce electricity and ECs for low-energy consumption H₂ production [9-12]. The sluggishness of the electrochemical kinetics of both cathode and anode reactions is so important that platinum group metals (PGM) are considered to be the essential cornerstone catalytic species. The diminution of the loadings of these precious metals (below 0.1 mg cm⁻²) while keeping or even increasing performance is a key challenge [6,13]. To date, almost all the synthesis routes focus on the decomposition of precursors in the presence of surfactants or stabilizers to engineer catalytic materials with tunable surface structure and composition, ranging from nanocages, nanowires, nanoframes, nanosponges, mesoporous materials, nanoalloys to highly-faceted materials [2,3,9,14]. Hence, the nanoparticles (NPs) do not take the advantage of the metal-support interaction [15,16] and have to be stabilized in a liquid or dispersed in a powder before using a binder such as Nafion for immobilization on a substrate (mostly glassy carbon) during various post-synthesis steps. While the use of such ionomer contributes to the proton transport, the process adds a layer of complexity to the fabrication process, the isolation of some NPs is common and the degradation during the long-term catalysis results in the undesired loss of active sites and subsequent decrease in catalytic performance [17]. In addition, those methods result in high loading of precious

metals. To avoid the need to collect and immobilize NPs after their synthesis, the ideal solution is to directly prepare them on their own support of FCs and ECs that is the three-dimensional (3D) gas-diffusion layer electrodes (GDL) [18]. Among the solutions for the advantageous direct growth of NPs on GDL electrodes, atomic layer deposition (ALD) appears as a promising route, as materials can be prepared on a layer-by-layer basis at the atomic-level thickness control as well as an excellent uniformity and conformity over the substrate surface, which makes this technique applied in a myriad of applications [18-27].

Using ALD to elaborate electrocatalysts enables to maximize the metal-support interaction and impact the electronic properties of active sites, thus changing the adsorption strength of reactants and intermediates, resulting in improved reactivity and stability [15,16]. Assaud *et al.*[21,22] have shown that, in alkaline media, the electrochemically active surface area (ECSA) and ethanol's electrocatalysis of ALD-synthesized Pd NPs is much improved. Recently, our group has shown that the increase of the glycerol activity by an order of magnitude in comparison to the Pd/C reference is reachable by decorating microfibers of a carbon felt electrode with Pd NPs of 4-6 nm at the reduced loading of $96 \mu\text{g}_{\text{Pd}} \text{cm}^{-2}$ [23]. It has been reported that boron nitride (BN) film enables improving the efficiency of number of heterogeneous catalysis reactions [28-32] substantiated by theoretical studies [28,33,34]. However, to the best of our knowledge, there is no report in the literature on the performance of ALD grown Pd NPs on BN surfaces, despite the great potential of this combination. It is also missing the electrochemical properties of BN-Pd coated large surfaces of GDL such as those used in energy technologies. Conclusively, a fundamental knowledge is missing about the electrocatalytic properties of Pd NPs supported on GDL electrodes through a nano-bridge of BN.

To address all those challenges and establish a performance trend, we report herein the design, synthesis and assessment of the electrocatalytic properties of those new classes of

ALD-grown GDL-BN-Pd freestanding electrodes with tunable BN's thickness towards four model-reactions of $\text{Fe}(\text{CN})_6^{3-/4-}$ redox probe, ORR, ethanol and ethylene glycol oxidation. When BN bridges Pd NPs and carbon microfibers, the resulting electrode of $5 \mu\text{g}_{\text{Pd}} \text{cm}^{-2}$ showed surprisingly high electrochemical kinetics for all tested reactions. The catalysts also exhibited excellent durability results. It is expected that the developed strategy can be readily extended to other type of low metal content catalysts thus re-enforcing the possible widespread deployment of ALD-based materials in heterogeneous catalysis.

2. Experimental

2.1. Reagents and materials

GDL-based carbon paper (CP, AvCarb MGL190, $190 \mu\text{m}$ thickness, Fuel Cell Earth LLC), ethanol (EtOH, VWR, 100%), sodium hydroxide (NaOH, Prolabo, 99%) and potassium hydroxide (KOH, Prolabo, 85%) were used as received. Isopropanol (ACS reagent, $\geq 99.8\%$), palladium hexafluoroacetylacetonate ($\text{Pd}(\text{hfac})_2$, 95%), potassium nitrate (KNO_3 , 99%), Nafion® suspension (5 wt%), ethylene glycol (EG, $\geq 99\%$), boron tribromide (BBr_3 , 99.9%), formalin (37 wt.% in H_2O , contains 10-15% methanol as stabilizer) and potassium hexacyanoferrate ($\text{K}_3[\text{Fe}(\text{CN})_6]$, $\geq 99.0\%$) were purchased from Sigma-Aldrich and used as received. Gases (O_2 , N_2 , Ar, NH_3) and water (Milli-Q Millipore, $18.2 \text{ M}\Omega \text{ cm}$ at $20 \text{ }^\circ\text{C}$) were ultrapure. Commercial catalyst is Pd/C (20 wt%, 3-4 nm, Premetek Co., USA).

2.2. Nanocatalysts synthesis by ALD

The GDL substrates were 5 cm high, 1 cm width, and 0.19 mm thickness. All depositions have been carried out in a low pressure hot-wall ALD reactor [35]. The BN process was based on BBr_3 precursor and NH_3 as co-reactant. Both BBr_3 and NH_3 lines were directly connected to the reactor through gate valves and heated at 100°C to avoid condensation. The deposition chamber was set at $750 \text{ }^\circ\text{C}$. The typical ALD cycle consists of 0.1 s pulse of BBr_3 , 5 s

exposure, and 15 s purge, followed by a 3 s pulse of NH_3 , 5 s exposure and 15 s purge with Ar. The as-fabricated samples with different thicknesses of BN are referred to as CP-BN $_x$ ($x = 10, 20, 40 \text{ \AA}$). The Pd process was based on $(\text{Pd}(\text{hfac})_2)$ and formalin as co-reactant. The formalin was kept at room temperature and the $\text{Pd}(\text{hfac})_2$ was heated at $70 \text{ }^\circ\text{C}$ to have sufficient vapor pressure. The lines were heated at $100 \text{ }^\circ\text{C}$ to avoid any condensation and the deposition chamber was set at $220 \text{ }^\circ\text{C}$. The typical ALD cycle consists of 5 s pulse of $\text{Pd}(\text{hfac})_2$, 15 s exposure, and 20 s purge, followed by a 1 s pulse of formalin, 15 s exposure and 60 s purge with Ar. The synthesized electrodes are labeled to as CP-Pd or CP-BN-Pd.

2.3. Physicochemical characterization

Transmission electron microscopy (TEM) was performed using JEOL 2200FS (200 kV) and JEOL ARM-200F (200kV). Inductively coupled plasma optical emission spectrometry (ICP-OES) analysis was performed on a spectrometer Optima 2000 DV (PerkinElmer). X-ray photoelectron spectroscopy (XPS) was fulfilled on an ESCALAB 250 spectrometer (Thermo Electron, monochromatic radiation source Al $K\alpha = 1486.6 \text{ eV}$). Survey spectra were recorded at a step of 1 eV (transition energy: 150 eV) and high-resolution spectra were recorded at a step of 0.1 eV (transition energy: 20 eV). The measurement of binding energy (BE) was corrected on the basis of the energy of C1s at 284.4 eV, and the quantification was carried out from the corresponding XPS peak area after correction with a suitable sensitivity factor.

2.4. Electrochemical and catalytic measurements

The half-cell electrochemistry was studied in a conventional three-electrode cell. Ag/AgCl/KCl (3 M) was used as reference electrode and was separated from the solution by a Haber-Luggin capillary tip. However, control experiments in alkaline solution were performed using Hg/HgO/Na(K)OH. Most of potentials were scaled versus the reversible hydrogen electrode (RHE) according to the calibration relationship $E(\text{V vs RHE}) = E(\text{V vs Ag/AgCl}) + \Delta E$, with $\Delta E = 0.959 \text{ V}$ (0.1 M NaOH) and 1.010 V (1 M NaOH). The calibrating

with respect to RHE was performed in the high-purity H₂-saturated electrolyte (Fig. S1). A slab of glassy carbon (12.4 cm²) was used as counter electrode. The working electrode consists of a CP-based electrode cut into L-shape of 1 cm high and 0.5 cm width, leading to an estimated area of 1 cm² and enough space on the top for electrical wiring with gold. The catalytic ink preparation for the commercial material follows standard procedures. Typically, 180 μL of isopropanol and 20 μL of Nafion® suspension were ultrasonically mixed. Then, 2.4 mg of Pd/C powder was added. Finally, 7 μL of the homogeneous ink was drop-casted onto each face of a bare L-shape CP electrode of 1 cm high and 0.5 cm width and dried at room temperature under N₂ flow. The Pd value assessed by ICP being 23.5 wt.%, the loading is 39 μg_{Pd} cm⁻². Electrochemical characterization by the redox probe Fe(CN)₆³⁻/Fe(CN)₆⁴⁻ was performed in 1 M KNO₃. Given the Pd NPs ability to absorb hydrogen into their crystal lattice, we used the PdO reduction peak (Fig. S2) by considering a monolayer charge density of 424 μC cm⁻² [36,37]. EtOH and EG oxidation reactions were investigated by the methods of cyclic voltammetry (CV), chronoamperometry (CA) and potentiostatic electrochemical impedance spectroscopy (EIS). ORR polarization curves were obtained with O₂-saturated electrolyte. Since the as-synthesized materials are freestanding (in the form of a sheet), the classical rotating (ring)-disk electrode R(R)DE setup could not be used to routinely overcome the diffusion process setup and run ORR under different rotation speeds. The polarization was thus collected by gently stirring the solution. Preliminary EIS was performed at OrigaStat OGS100 potentiostat (Orignalys) scanning frequencies from 1 kHz to 25 mHz (10 mV amplitude). Then, SP150 potentiostat (Biologic Science Instruments) was used for frequencies of 200 kHz to 25 mHz (10 mV amplitude). All the voltammograms are iR-free, *i.e.*, corrected by the “potential drop” between the working and reference electrodes according to the relationship $E_{\text{real}} = E_{\text{applied}} - R_{\Omega} \times I$. R_{Ω} is obtained by EIS at the intersection between the

Nyquist curve and x -axis at high frequencies, herein $R_{\Omega} \sim 20 \Omega$ in 0.1 M NaOH and $\sim 3 \Omega$ in 1 M NaOH.

3. Results and discussion

3.1. Electrochemical performance of the developed BN nanolayers at the surface of a gas-diffusion electrode

In the following sections, the overall process of ALD assisted fabrication of the electrode materials and electrochemical tests have been validated more than three times the studies and the conclusion is that the process is reproducible. To determine if the surface of CP would be favorable to the deposition of a layer of BN and Pd NPs or not, we first carried out a systematic physicochemical investigation. We especially aimed to study the hydrophobic/hydrophilic character, as that is the presumed starting point to have a good wettability, a prerequisite for use in electrocatalysis in aqueous media. Indeed, in some cases, it is necessary to perform an activation step before NPs deposition or obtaining significant electrocatalysis [23]. As-shown in Fig. 1a, the pristine CP electrode (190 μm thickness) is composed of $\sim 7 \mu\text{m}$ diameter microfibers, which means than the entire electrode could be described as a stack of $190/7 = 27$ layers of fibers. XPS survey (Fig. 1b) shows O signals and the high resolution spectrum of the C1s core level (Fig. 1c) demonstrates the presence of oxygenated species on the surface, thus making the surface hydrophilic. Performed water-contact angle-measurements have upheld the wettability of the material. Those results confirm that CP can be used in aqueous media without any further pretreatment steps.

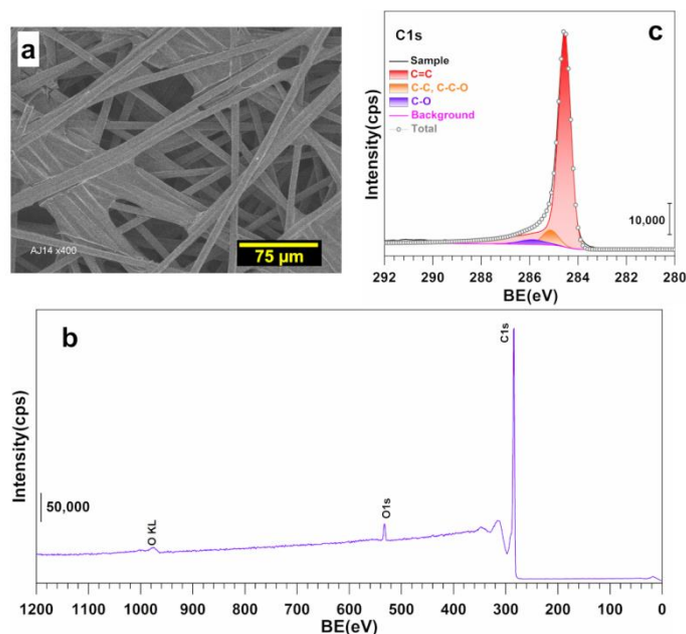


Fig. 1. (a) SEM image of the bare CP electrode. (b) Survey XPS spectra of the CP electrode. (c) High-resolution XPS spectrum of the C1s core level for the bare CP electrode.

To gain a deep understanding on the effects of B, N co-modification on the performance, we first prepared a series of CP-based materials by varying the BN atomic layer thickness. The optimized ALD process allows for the saturated growth of conformal BN films with a steady-state growth of $\sim 0.8 \text{ \AA/cycle}$ with no visible nucleation delay [35,38]. Several analytical methods have been used to characterize the films, and the growth-per-cycle, C and O contents, roughness and mass density values are given in the Table S1. In the present study, we applied different cycles in order to coat the pristine CP substrate with an ultra-thin film and referred to as CP-BN x ($x = \text{thickness in \AA}$). Then, the samples were electrochemically probed in the presence of $\text{Fe}(\text{CN})_6^{3-}/\text{Fe}(\text{CN})_6^{4-}$ as the quasi-reversible redox probe, herein used to evaluate electrochemical kinetics. In order to provide a better qualitative and quantitative analysis of the electrochemical parameters, we used two different methods [39-42], namely the peak-to-peak separation (ΔE_p) between the anodic and cathodic peaks on a CV and the EIS. The control experiments using a Pt RDE and the collected results are

reported in Fig. S3-S6. Fig. 2a shows the steady-state profiles of CVs in the presence of the substrate while those recorded in the electrolyte are displayed within Fig. S7. The blank voltammograms indicate an increase of the capacitive current when BN thickness augments, which is in agreement with the presence of BN that is a lesser electron conductor than CP. However, we unexpectedly found from Fig. 2a that the introduction of BN atomic layer increases significantly, from +20 to 70%, the peak current density associated to the oxidation and reduction waves assigned to $\text{Fe}(\text{CN})_6^{4-}$ and $\text{Fe}(\text{CN})_6^{3-}$, respectively. Please note that the background current is highly negligible (Fig. 2a versus Fig. S7). From the ΔE_p modeling curve of Fig. S6, we specifically determine the standard rate constant (k°), charge transfer resistance (R_{ct}) and exchange current density. $k^\circ(10^{-3} \text{ cm s}^{-1}) = 1.7, 2.0, 6.2, \text{ and } 10.5$ for CP, CP-BN10, CP-BN20, and CP-BN40, respectively (Table S2). Those data correspond to a significant enhancement of the electrochemical kinetics by factors of 1.2-6.3 upon the deposition of BN. It features the ability to regulate the kinetics by the means of BN's ALD. Specifically, the trend can be translated to as a decrease of R_{ct} . Fig. 2b shows EIS data at open circuit potential (OCP) and the corresponding R_{ct} and k° determined by using $R_{\Omega+Q_{CPE}}/(R_{ct}+W)$ as a representative equivalent electrical circuit (EEC) wherein W is the Warburg impedance (to model the mass transfer, *i.e.*, diffusion) and Q_{CPE} is the constant phase element (to model "imperfect" capacitors) [43,44]. This second method confirms the observed trend for the metrics of R_{ct} , k° , and j_0 . R_{ct} (inversely proportional to k° , thus j_0), is the ability of electron-transfer and the difficulty of an electrode reaction for driving larger current density with a less driving force [45]. For a given electrode, the bigger R_{ct} is, the lower is the kinetic rate and lesser is the driving force. It can thus be concluded that the presence of BN improves the transfer of one electron from the catalytic surface to the reactant, while the reverse is true. With nearly one order of magnitude increase of k° for BN-modified CP, one

can come up with a beneficial effect of the atomically bridged heteroatoms on the electrochemical kinetics of one electron-transfer process of a quasi-reversible [46,47].

The above outcomes demonstrate only our ability to regulate the kinetics of one electron-transfer process that is $\text{Fe}(\text{CN})_6^{3-} + \text{e}^- = \text{Fe}(\text{CN})_6^{4-}$. Suitability of modified electrodes for applications in heterogeneous electrocatalysis may require larger number of electrons. So, we next theorized a same trend for much higher number of electrons. To validate such hypothesis, we investigated ORR in alkaline media which is of paramount importance in electrochemical energy conversion and storage technologies. The performance in 0.1 M NaOH is displayed in Fig. 2c-2d for LSVs and EIS at $E_{\text{appl}} = 0.81$ V vs RHE. The inset shows the fitted EIS data by an EEC of $R_{\Omega} + Q_{\text{CPE}} // R_{\text{ct}}$ (extended data in Table S3). It should be noted that we firstly determined the depth of ORR by using our routine hydrogen peroxide method that combines chronoamperometry and UV-Vis [48]. After chronoamperometry, samples are then acidified toward pH 3.5 ($\text{p}K_{\text{a}}(\text{H}_2\text{O}_2 / \text{HO}_2^-) = 11.75$, any produced H_2O_2 from the incomplete ORR is a form of HO_2^-), and the resulting H_2O_2 was quantified by spectrophotometry, using potassium titanium (IV) oxalate as a colored indicator via a calibration curve obtained from standard solutions. Then, the absorbance of the yellow pertitanic acid complex between H_2O_2 and potassium titanium oxalate in acidic solution is measured at $\lambda = 400$ nm. We did not observe any significant amount of intermediate species for $E \geq 0.8$ V vs RHE. Thus, it can be argued that ORR at the entire electrodes is a 4-electron process ($\text{O}_2 + 2\text{H}_2\text{O} + 4\text{e}^- \rightarrow 4\text{OH}^-$) so that kinetics parameters can be accurately well compared. Chen *et al.*[28] have shown that ORR starts at 0.8 V vs RHE for a catalytic ink of BN materials drop-casted onto RDE (by mixing catalytic powder, solvent and Nafion ionomer) in a 2-electron major process. The high selectivity observed on the present materials can be explained by synthesis procedure leading to surfaces that foster the perpendicular binding of O_2 and results in its cleavage. The polarization curves in Fig. 2c highlight a

prominence shift of the potential towards higher potential, *i.e.*, lower overpotential. Similar improvement of the electrochemical performance of carbon substrates modified by B and N elements has been reported for other reactions and assigned to the heteroatom-carbon interface that acts as reactive sites [28,30-32,34,49-51]. From EIS data, the high activity originates from the decrease of R_{ct} , about 60-70% upon the deposition of BN layer, which accelerate the rate of the catalytic reaction. For a potential deployment as an ORR's cathode material for fuel cells or metal-air batteries purposes, the electrode should show large current density for potential higher than 0.8 V vs RHE. As the efficiency is below the threshold, the only possibility is its use as scaffolds for the deposition of ultra-low amount of active Pd metal, ideally lower than $50 \mu\text{g cm}^{-2}$ [4,6,13].

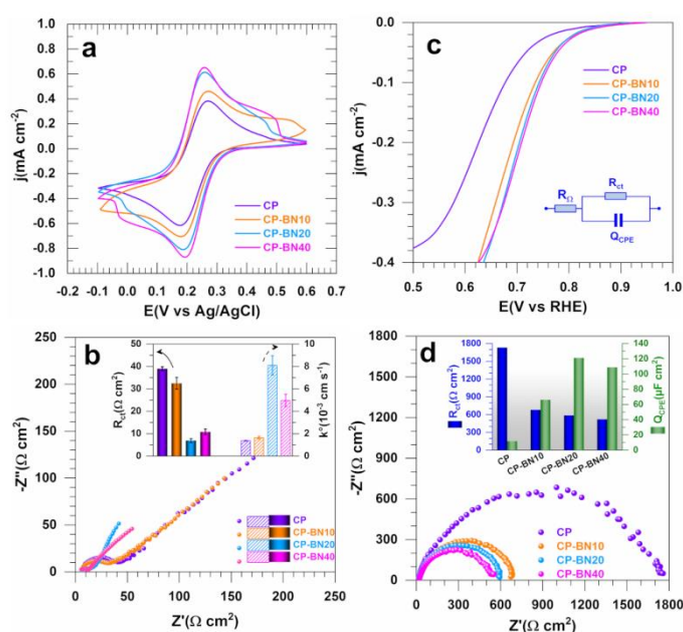


Fig. 2. (a) iR -free steady-state CVs recorded at 10 mV s^{-1} for pristine CP and as-fabricated electrodes in $10 \text{ mM K}_3[\text{Fe}(\text{CN})_6] + 1 \text{ M KNO}_3$. (b) Complex-plane Nyquist impedance plots at $E_{\text{appl}} (=OCP) = 0.3 \text{ V vs Ag/AgCl}$: the inset shows the fitted R_{ct} . (c) iR -free ORR LSVs recorded on pristine CP and as-fabricated CP-BN in O_2 -saturated 0.1 M NaOH at 5 mV s^{-1} : the inset shows the used EEC for fitting. (d) Complex-plane Nyquist impedance plots at $E_{\text{appl}} = 0.81 \text{ V vs RHE}$ in O_2 -saturated 0.1 M NaOH : the inset shows the fitted R_{ct} (left Y-axis) and

Q_{CPE} (right Y -axis). The temperature is 25 °C. Error bars represent one standard deviation ($n = 3$).

3.2. Electrochemical probing and physicochemical characterization of Pd modified CP-based electrodes.

Having demonstrated the ability to use ALD to grow BN nano-layers directly onto a GDL support, we sought to study the modification by electrocatalytic active Pd NPs more carefully. Indeed, the suitability of electrode materials for their application as cathodes may not require the presence of Pd, but for anode catalyst for fuel oxidation, the presence of those elements termed as PGMs is mandatory to achieve high current densities at lower overpotentials. Based on the previous results, we selected CP-BN20 for comparative studies with CP to fabricate by 100 cycles of Pd ALD, CP-Pd and CP-BN-Pd electrodes. Then, their electrochemical behaviors were confronted to the commercial Pd/C (20 wt.%, “C” denotes carbon Vulcan XC-72) presenting similar Pd NPs size (3-4 nm). Fig. 3a shows the profiles of the steady-state CVs in the presence of the redox probe, $\text{Fe}(\text{CN})_6^{3-}/\text{Fe}(\text{CN})_6^{4-}$. The extracted kinetics data of k° , R_{ct} and j_0 from the peak-to-peak separation method are gathered in Table S2 wherein $j^\circ(\text{mA cm}^{-2})$ is 0.3, 0.9 and 1.0 for CP-Pd, CP-BN-Pd and Pd/C, respectively. The values of R_{ct} and k° extracted from the second method of EIS performed at $E_{\text{appl}}(=\text{OCP}) = 0.3 \text{ V vs Ag/AgCl}$ are shown in Fig. 3b and confirm the accelerated electrochemical kinetics when BN is intercalated between carbon surface and Pd NPs. The elementary analysis by ICP-OES shows that the Pd content in CP-Pd and CP-BN-Pd is about 30 and 5 $\mu\text{g cm}^{-2}$, respectively. Furthermore, it can be deduced from TEM images (for this purpose, single fiber was prepared) of Fig. 3c (CP-Pd) and 3d (CP-BN-Pd) that the Pd NPs are crystalline with a size distribution of $5 \pm 2 \text{ nm}$. A deeper analysis indicates that the NPs of quasi-spherical shape are well dispersed all over the surface of the BN coated fibers with a longer distance between

individual NPs. This enables to diminish the total amount of the precious metal. As the BN layer is turbostratic, the strength of interaction between the BN layer and the Pd clusters is difficult to determine. Many competing surface reactions occur during ALD, and these reactions are both very dynamic and complex. The BN surface is expected to be terminated by hydrogen atoms. The initial growth of ALD of palladium at the BN surface is likely to start with the chemisorption reaction between the Pd(hfac)₂ precursor and the BH or NH groups present at the surface, resulting in the deposited Pd atoms and volatile H(hfac) species. The strength of the chemical bond between the Pd atoms and the BN surface can be determined by theoretical calculations. From theoretical studies carried out in the past, it has been found that in all cases, the B atom is closer than the N atom to the palladium surface, and the h-BN-palladium bonding energy is around 0.4 eV. Taken together, those set of results demonstrate that the presence of a ~2 nm BN layer enables lowering significantly the loading of Pd while getting high electrochemical kinetics. The BN surface presents a certain number of atoms in direct contact with the palladium NPs, thus forming an atomically bridged region between the carbon and the noble metal material. The electrons are able to readily cross this thin BN nanolayer bridge to reach the carbon fiber, and this bridged region is likely to explain the superior electrochemical performance measured. In fact, due to special structure of BN which is similar to graphene with small voids, it can act as a bridge for small species when deposited as an ultrathin film.

Given the high amount of Vulcan (carbon black of nanoscale domains [52]) in the Pd/C material, its contribution to the kinetics of the studied one electron-transfer process ($\text{Fe}(\text{CN})_6^{3-} + e^- = \text{Fe}(\text{CN})_6^{4-}$) cannot be completely ruled out. So, we have decided to next interrogate the electrocatalytic abilities towards reactions in potential windows where this contribution is much negligible.

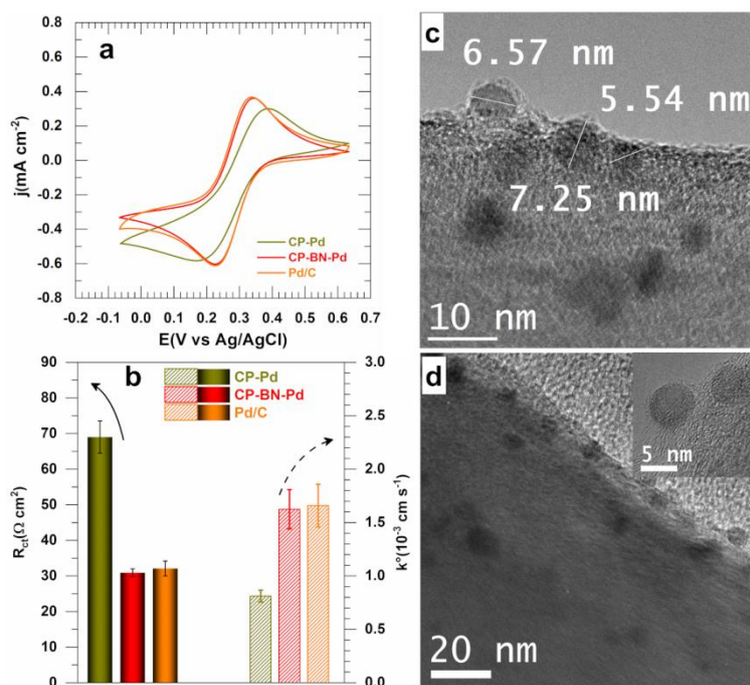


Fig. 3. (a) iR -free steady-state CVs recorded at 10 mV s⁻¹ in N₂-saturated 10 mM K₃[Fe(CN)₆] + 1 M KNO₃: The temperature is 25 °C. (b) Extracted kinetics data of R_{ct} (left Y -axis) and k° (right Y -axis) from EIS performed at E_{appl} (\approx OCP) = 0.3 V vs Ag/AgCl. (c) TEM of CP-Pd. (d) TEM of CP-BN-Pd. Error bars represent one standard deviation ($n = 3$)

3.3. Electrocatalytic performance towards ORR.

In order to clarify the actual electrocatalytic trend observed experimentally by the quasi-reversible redox probe molecule, we further considered ORR in 0.1 M NaOH (Fig. 4a) and 0.1 M KOH (Fig. 4b). The voltammograms clearly show that CP-BN-Pd has superior electrochemical performance than other tested catalysts of CP-Pd and Pd/C. Specifically; the results reveal that, on all electrodes, ORR starts at ca. 1 V vs RHE, ultimately approaching the theoretical equilibrium potential of 1.18 V vs RHE. The observed positive shift of 60 mV (0.1 M NaOH) and 25 mV (0.1 M KOH) at CP-BN-Pd electrocatalyst in comparison to CP-Pd at $j = 3$ mA cm⁻² can be translated to as an optimal balance between the available active sites for hydroxyl species (the main ORR performance descriptors in alkaline electrolytes [2,3,53]) and

O₂ species. Given the configuration of the present materials (freestanding), it is not possible to use an (R)RDE and extract the “pure kinetic current”. So we use the Tafel plot with the “mass activity” (A per mg of Pd) as reported in Fig. 4c. At 0.9 V vs RHE, the CP-BN-Pd shows a drastic activity increase of one and two order of magnitude compared to the material without BN and the commercial, respectively. Overall, the outcomes highlights that CP-BN-Pd aspires to be a potential active cathodic electrocatalyst for ORR in either FCs or metal-air batteries. We further used the technique of EIS to interrogate ORR at the ALD-synthesized samples, firstly at OCP, and secondly at 0.87 V vs RHE in both alkaline electrolytes. As shown in Fig. S8, the complex-plane Nyquist impedance plots recorded in O₂-saturated 0.1 M KOH at OCP can be fitted by an EEC of $R_{\Omega}+Q_{CPE}/R_{ct}$ to extract the data. From the collected data in 0.1 M KOH (Table S4), $R_{ct} = 2761$ and $33 \Omega \text{ cm}^2$ for CP-Pd at OCP and 0.87 V vs RHE, respectively. For CP-BN-Pd, the values are 1416 and $28 \Omega \text{ cm}^2$. The same trend was observed in NaOH electrolyte. These results demonstrate that at OCP where the reaction needs much driving force, the presence of BN nanolayer permits to lower by a factor of nearly two, the energy input to activate O₂ molecules at the catalytic surface. Since R_{ct} is correlated to the number of electrons that are transferred from the catalytic surface to the reactant(s) as well as intermediate(s) formation inside the double layer, it can be argued that the presence of BN creates an interface that modifies the electronic properties of Pd and/or C to weaken the substrate binding and regulate the accumulation of intermediates.

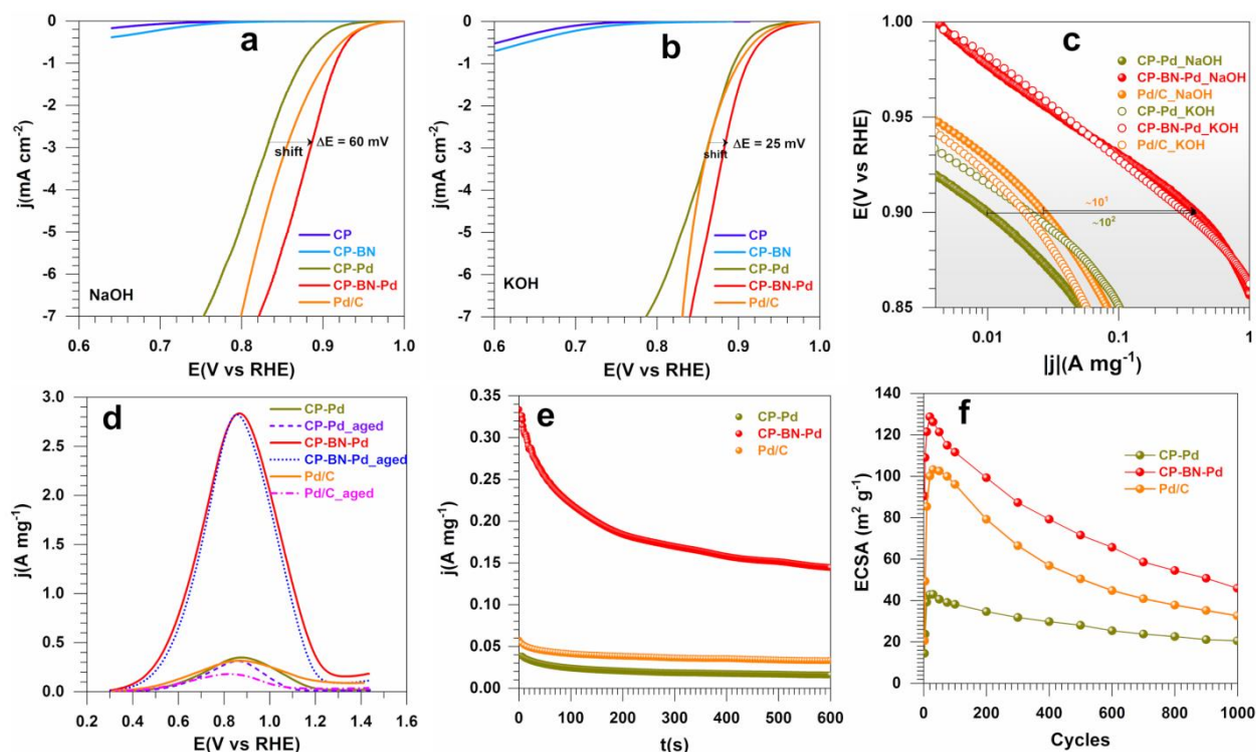


Fig. 4. (a) iR-free ORR LSVs recorded in O₂-saturated 0.1 M NaOH at 5 mV s⁻¹. (b) iR-free ORR LSV recorded in O₂-saturated 0.1 M KOH at 5 mV s⁻¹. (c) Tafel plots from panels (a) and (b). (d) iR-free steady-state forward CVs recorded at 50 mV s⁻¹ in the presence of 0.1 M EtOH before (dotted and dashed lines) and after (solid lines) the accelerated ageing. (e) Chronoamperometry in the presence of 0.1 M EtOH at $E_{\text{appl}} = 0.6$ V vs RHE. (f) ECSA of Pd behavior during the accelerated ageing. The temperature is 25 °C.

3.4. Electrocatalytic performance toward C2 alcohols electrooxidation reaction.

The previous outcomes demonstrate that we were able to manipulate the ALD synthesis towards electrodes with different electrocatalytic profiles. Except the use of Pd/C commercial material considered to as the benchmark catalyst in alkaline electrolytes [1,4,54], no rational comparison with the literature is feasible for ORR given the configuration of our electrodes. So, we next aimed to study and compared with the literature the newly described materials as electrode materials. For that, we investigated the electrocatalytic properties by considering two model anode reactions in alkaline media that are ethanol and ethylene glycol oxidation.

Since stability is an important parameter to be considered when designing catalysts, we also set an electrochemical accelerated ageing program for a dual examination of the ability of the catalysts in maintaining significant ECSA and activity under practical operations. The program consists of cycling the electrode potential from 0.3 to 1.45 V vs RHE (forwards and backward) at 50 mV s^{-1} for a total of 1000 cycles in 0.1 M NaOH (roughly 13 h of continuous operation). The lower potential limit has been fixed to 0.30 V vs RHE to suppress the irreversible insertion of hydrogen in the Pd crystal lattice while the higher potential limit has been set to 1.45 V vs RHE to avoid the formation of superior oxide of PdO₂ thus keeping the monolayer charge at $424 \mu\text{C cm}^{-2}$ [36,55-57]. The electrochemical performance is shown in Fig. 4d-4f in terms of CVs, CA and ECSA. Typical full CVs in the absence or presence of EtOH as well as the recorded voltammograms without current normalization are reported in Fig. S9 while extended CA, CVs at different concentrations of EtOH or scan rate, and ageing CVs are shown within Fig. S10-S14. During the positive-direction scan in the blank electrolyte, the palladium surface oxidation starts at about 0.5 V vs RHE followed by the reduction of the formed PdO during the backward scan at 0.6 V vs RHE on the as-synthesized CP-Pd and CP-BN-Pd catalysts and 0.6-0.5 V vs RHE at Pd/C. From Fig. 4, Table 1 for current density and Fig. S9b for current, one can see that CP-BN-Pd is undoubtedly the best performing catalyst with excellent stability. Specifically, in the presence of BN, the activity at the peak current displays a negligible decay of 1% while the sample without BN has a significant decay of 11% and commercial Pd/C a prominent activity decay of 44%. Looking deeply at the available ECSA, the values in the beginning of the ageing are 14, 90 and $21 \text{ m}^2 \text{ g}^{-1}$ for CP-Pd, CP-BN-Pd and Pd/C, respectively. The measured activity towards EtOH referred to as “before ageing” was recorded after 20 cycles of surface pre-treatment in electrolyte, corresponding to ECSA of 43, 129 and $100 \text{ m}^2 \text{ g}^{-1}$ for the same trend. Those values become 21, 46 and $33 \text{ m}^2 \text{ g}^{-1}$ at 1000th cycle. The increase of ECSA during the first

30th cycles is rationally assigned to the traditional surface rearrangement phenomenon upon the application of an electrical field between the working and the counter electrodes through the electrode potential.

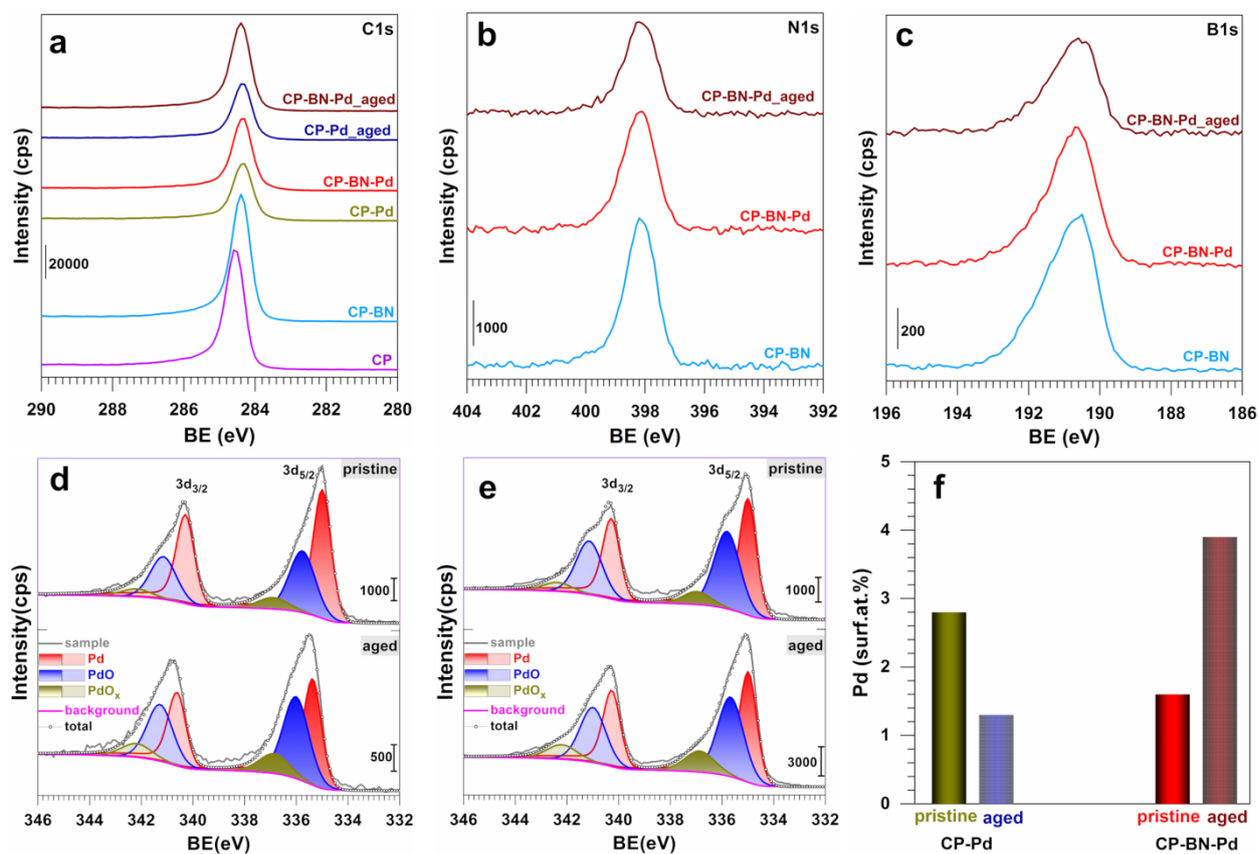


Fig. 5. (a-c) High-resolution XPS spectra of the C1s (a), N1s (b), and B1s (c) core levels for the different materials. (d-e) High-resolution XPS spectra of Pd 3d core level before (top) and after (bottom) the accelerated ageing for the as-fabricated CP-Pd and CP-BN-Pd electrode materials, respectively. (f) Quantitative data from the high-resolution XPS measurements of Pd 3d core level of the freshly synthesized (pristine) and after the accelerated ageing program (aged).

The achieved ECSA of 90 m² g⁻¹ at the 3th cycle or 129 m² g⁻¹ at 20th cycle clearly outperforms the previous reports of ALD-synthesized Pd onto carbon felt electrode [23] and more importantly, those of (electro)chemically-synthesized NPs or self-supported Pd-based

electrocatalysts wherein ECSA is about 10-70 m² g⁻¹, approaching 100 m² g⁻¹ in bimetallic structures [1,21,22,36,54,58-71]. The high values measured for ECSA in the presence of BN can be assigned to the high dispersion of Pd NPs on the electrode surface (previously shown by TEM) and to the absence of any organic surfactants on the surface of the nanocatalysts as it happens in conventional synthesis by colloidal procedures. The recorded peak current density (j_p) of 17 A mg⁻¹ (~84 mA cm⁻²) in 1 M NaOH + 1 M EtOH is about 2-10 times higher than relevant reports [10,21,22,64,66-68,70-74] of ethanol electrocatalysis at noble metals mono and bimetallic based on Pd, Au and Pt which are known to provide best up to date values in both alkaline/acidic media (Table S5). We further evaluated the electrocatalytic performance towards ethylene glycol oxidation, another potential candidate for organic molecule-fuelled devices to produce electricity or H₂. The peak current densities are gathered in Table 1 while CVs and CA curves are reported in Fig. S15. The value of j_p is 0.38 and 3.39 A mg⁻¹ in 0.1 M EG for CP-Pd and CP-BN-Pd, *i.e.*, about 9-times increase of performance; thus underpinning the higher electrochemical activity induced by BN intercalation between carbon surface and Pd NPs. It is anticipated that at higher EG concentration, CP-BN-Pd should largely outperform world-class nanoparticles electrocatalysts, j_p of 3-6 A mg⁻¹ in 0.5 M KOH + 0.5 M EG, [61,62] and 0.1-2 A mg⁻¹ under other conditions [75-80].

The origin of the present exceptional activity results from several parameters of structural, geometric and electronic order that are well-established in heterogeneous electrocatalysis [15]. Specifically, our catalysts were obtained in the configuration of simple deposition of Pd NPs on carbon-based substrate, while in most previous studies, NPs are dispersed inside a porous matrix of Vulcan carbon powder to maximize the surface area of the NP catalysts and improve the electron-transfer process. This is followed by further steps of mixing with organic solvents and Nafion suspension as a binder to insure a strong attachment of catalytic species to the surface. So, we believe that the simplicity of the developed approach herein

leads to bare uncontaminated surface of nanomaterials in contrast to conventionally-synthesized ones where the surface is blocked by organic byproducts coming from the precursor's decomposition to form NPs. The second reason that might explain the improvement is related to the strong interaction between **Pd NPs** and the carbon support whose electronic properties have been modified upon the deposition of BN atomic layers.

Table 1. Performance of EtOH and EG alcohols electrooxidation from CVs at 50 mV s⁻¹.

Current density j_p (A per mg of Pd)		CP-Pd	CP-BN-Pd	Pd/C
0.1 M NaOH + 0.1 M EtOH	before ageing	0.35	2.84	0.32
	after ageing	0.31	2.82	0.18
1 M NaOH + 1 M EtOH		-	16.73	-
0.1 M NaOH + 0.1 M EG		0.38	3.39	-

3.5. Compositional and surface characterizations of the as-grown Pd NPs onto CP and CP-BN electrodes.

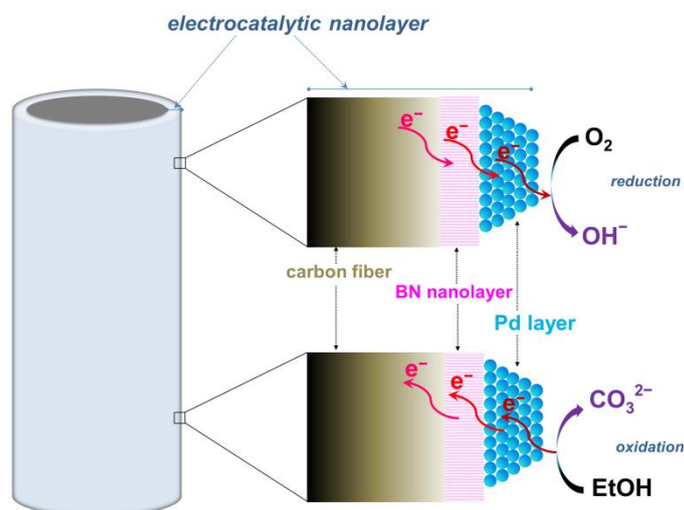
Having demonstrated the ability to use jointly ALD of BN and Pd to fabricate electrodes with high electrocatalytic properties, we sought to study the surface composition and morphology more carefully. To this end, we employed the surface technique of XPS. Fig. 5 shows the set of results while extended data from XPS spectra fitting are reported in Tables S6-S7. As shown in Fig. 5a and Fig. S16-S17 (survey spectra), the signal of C1s decreases upon the ALD processes indicating that the surface of each individual fiber is being covered by BN film (2 nm) and **Pd clusters**. Fig. 5b-5c highlight the presence of B1s (BE = 190.7 eV) and N1s (BE = 398.1 eV) whose surface compositions are about 6 and 5 at.% before ageing and 4 and 3 at.% after ageing test. The B/N ratio is quasi-constant meaning that the ageing might not affect the structure of BN nanolayer. For the high-resolution XPS spectra of Pd3d core level of the ALD-synthesized Pd NPs of 5 ± 2 nm (Fig. 5d-5e), the comparison with the chemically synthesized Pd/C of similar particles size [52] reveals that **Pd NPs** at the surface

are much less oxidized. The identified species are pure Pd metal, PdO, and PdO_x ($x > 1$) [52,81-83]. It is worth of mentioning that, when a metal is exposed to ambient air, it intuitively undergoes a chemical oxidation process because of the oxygen in air. This leads to the formation of a very thin layer, so-called passivation layer, which protects the bulk metal from deep oxidation process. The observed doublets are related to the spin-orbit splitting ($j = l \pm \frac{1}{2}$, quantum number $l = 2$ for d band). The metallic state of Pd 3d region is characterized by the two spin-orbital components 3d_{5/2} (335.2 eV) and 3d_{3/2} (340.5 eV). The table S7 given in SI shows the extended data from XPS spectra fitting. The measured Pd 3d_{5/2} binding energy peak (BE=335.2 eV) level for Pd is the same before and after ageing and corresponds to the position of metallic palladium, which indicates that the Pd NPs are in the Pd(0) state and are stable in the media. From Fig. 5f, one can see that the surface atomic amount of Pd decreases in CP-Pd while that of CP-BN-Pd increases. Since both B and N amounts have decreased in CP-BN-Pd, it is normal to expect an augmentation of total Pd. Looking closely at the SEM image of CP in Fig. 1a, and based of the fact that XPS probes micro-scale domains, it is highly possible that the analyzed area after ageing domains is composed of the interconnected regions wherein Pd NPs are much important. This can thus rationally explain the increase of Pd3d signal. The augmentation of Pd at the surface of BN coated fibers after potential cycling can lastly be explained by the presence of trace by-products of ALD process of hexafluoroacetylacetonate and formalin that were still remaining on the surface of Pd NPs so that the first voltammetry cycles act as “cleaning” and further re-organization of Pd NPs into a morphology that exposed much atoms at the surface. This is a common observation in electrochemical science and is in agreement with the profiles of ECSA and ethanol oxidation activity. Furthermore, the slight increase of oxidized Pd species in the CP-BN-Pd material in comparison to the unmodified CP-Pd might be explained by the fact that boron nitride attracts electron from Pd similar to previous report [52]. We finally argue that the presence of BN

reinforces the tethering of Pd NPs to the surface which can also explain the loose of activity of Pd NPs or their agglomeration into bigger one, i.e., less surface atoms in the CP-Pd material. Conclusively, those outcomes corroborate the findings from electrochemical characterization and catalytic measurements and consolidate our starting hypothesis that atomic layer of boron nitride brings positive effects on the catalytic activity of Pd NPs. In Table S5, the reported performance of nanocatalysts towards the ethanol (EtOH) electrooxidation in alkaline media is given, in order to have a good comparison with relevant data from literature. The results presented showed a 16-times increase for CP-BN-Pd compared to the commercial material increase of performance, and an overall better performance of the system presented when compared to relevant data from literature.

3.6. Mechanistic scheme of electrocatalytic reduction and oxidation processes at CP-BN-Pd nano-interface.

To resume these forays toward the deployment of electrocatalytic electrodes derived from the BN and Pd ALD, we finally combined the outcomes to sketch the process in Scheme 1. Since with a nano-bridge of 20-40 Å, electrons coming from the electrode reach O₂ substrate (reduction) and those released by the ethanol befall the electrode (oxidation), it is stipulated that the scaffolding created by B and N heteroatoms at the interstitial region are acting as electron relays. Hence, the BN nanolayer plays the role of “bifunctional interface” that regulates the accumulation of intermediates and exchange of electrons.



Scheme 1. Proposed reaction sketch of the electrocatalysis mechanisms taking place in CP-BN-Pd electrode (left side) during electro-reduction reaction, O_2 , (right side, top) and electro-oxidation reaction, EtOH, (right side, bottom).

4. Conclusion

In conclusion, we have presented a versatile ALD approach to grow nanostructured palladium (Pd) directly on carbon fibers of a gas-diffusion electrode (GDL). We have relied heavily on electrochemical techniques of cyclic voltammetry (CV), chronoamperometry (CA), electrochemical impedance spectroscopy (EIS) as well as transmission electron microscopy (TEM) and X-ray photoelectron spectroscopy (XPS) to provide viable quantitative parameters. By using boron nitride (BN) nano-bridge, we were able to tightly regulate the electrochemical kinetics of the electrodes by the means of the thickness (x). Specifically, the rate constant of the redox probe molecule $Fe(CN)_6^{3-}/Fe(CN)_6^{4-}$ has been found to be $k^o(10^{-3} \text{ cm s}^{-1}) = 1.7, 2.0, 6.2$ and 10 for $x = 0, 10, 20,$ and 40 \AA , respectively. This enhancement of the kinetics was confirmed by oxygen reduction reaction (ORR) with a charge transfer resistance (R_{ct}) of $1733, 680, 587$ and 520 \Omega cm^2 in 0.1 M NaOH at 0.81 V vs RHE for the same trend of x .

It was also demonstrated that the presence of 2 nm BN film enables not only lowering significantly the loading of Pd NPs on carbon paper (CP) electrodes of tunable dimensions but also controlling the dispersion to reach exceptional activity and durability for ORR, ethanol oxidation (EtOH), and ethylene oxidation (EG) reactions. In the start-up stage, electrochemically active surface area (ECSA) of 14, 90 and 21 m² g⁻¹ for CP-Pd, CP-BN-Pd and Pd/C, respectively, represents 4-times increase compared to the benchmark catalyst. After extended electrochemical ageing tests, it was found that the sample without BN losses 54 at.% of Pd and significant activity decay towards ethanol oxidation which was not the case in the presence of BN. A peak current density of $j_p = 0.31, 2.82, 0.18$ A mg⁻¹ for CP-Pd, CP-BN-Pd and Pd/C indicates a 16-times increase for CP-BN-Pd compared to the commercial material. Importantly, the catalytic efficiencies of this advanced CP-BN-Pd electrode towards EtOH and EG electrocatalysis outperform the produced nanocatalysts in the literature. These outstanding outcomes encapsulate definitely the potential interest of boron nitride in electrocatalysis and open prospects for the use of ALD to elaborate bimetallic NPs with even superior catalytic performance.

Supplementary material. Tables S1-S5 and Fig. S1-S17.

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