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Multi-Walled Carbon Nanotubes Supported Pd(II) Complexes: A Supramolecular Approach towards Single-Ion Oxygen Reduction Reaction Catalysts

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Abstract: Lowering the platinum group metal content of oxygen reduction reaction catalysts is among the most prevalent research focuses in the field. This target is herein approached through supported Pd(II) complexes. Starting from a commercial macrocycle, a new ligand is synthesized, its solution behavior and binding properties briefly explored (potentiometry, UV-Vis) and then used to prepare a new catalyst. A supramolecular approach is used in order to obtain homogeneous decoration of carbon nanotubes surfaces, fostering novel possibilities to access single-ion active sites. The novel catalyst is characterized through X-ray photoelectron spectroscopy and scanning transmission electron microscopy and its promising oxygen reduction reaction performance is evaluated via rotating ring-disk electrode and rotating disk electrode in half-cell studies.

Keywords: heterogeneous catalysis; surface adsorption; palladium catalyst; supramolecular interactions; ORR; RRDE; CNTs

1. Introduction

Climate change evidence [1], pollution [2], and recent data concerning the improvement of air quality connected with the decrease of human industrial activities during COVID-19 lockdowns [3] are raising the awareness of public opinion on the importance of transitioning to cleaner energy. At the same time, as Mendeleev suggested a long time ago, there are much smarter possible usages for non-renewable resources, such as hydrocarbons, than burning them for energy production.

Fuel cells and hydrogen technology appear as a possible solution: provided that H₂ is produced from clean sources, its use as energy vector would be ideal, as its combustion is highly exothermic and produces no waste residuals beyond water [4,5]. Currently, beyond superable pragmatic issues (the conversion of fuel supply chain, safe high-pressure portable tanks for vehicles, etc.) [4,6], technological obstacles are still limiting large scale H₂ usage: these mainly include effective electrocatalysts for fuel cell and water hydrolysis which are able to maintain high performances over time at affordable costs [7–9]. Here, we remark on the crucial dependence of most up-to-date solutions on so-called PGM (platinum group metals), which are not only expensive but also, again, finite resources (i.e., not fundamentally better than hydrocarbons to critically rely upon).

How to design a better catalyst?
This is the question that led chemists and material scientists to explore a plurality of tentative answers. Metal nanoparticles [10–15], non-noble metals [10,11,16], and even metal-free catalysts [16–19] have been proposed, with each of these solutions coming with its own merits and shortcomings.

Our contribution to the field goes in yet another direction. Regardless of which metal we will use as catalyst, if any, it is important to make a strategical use of available resources: reuse and recycle are good things [20], using less is even better. In terms of catalysts, this implies, at its theoretical limit, using a single ion or metal atom as a catalytic active site. The general concept of single-atom catalysts (SACs) has been introduced and discussed elsewhere [21–23]. In the case of Pd, which is the element that we currently focus on (face-centred cubic, lattice constant 3.859 Å), the difference between supported small nanoparticles (a perfect cubic small NP with a 3.859 nm side consists of 4000 atoms) and a SAC catalyst, could mean lowering the required amount of Pd of about three orders of magnitude. At market prices above 60 $/g Pd, this can, in principle, have a huge impact on the economic feasibility of any catalysts: a crucial aspect for oxygen reduction reaction (ORR).

We make use of supramolecular chemistry and supported metal complexes to achieve this type of catalyst, rather than doping strategies [24–26]. A smart modular design of organic ligands possessing a polyaminic macrocyclic unit for Pd(II) binding and a grafting unit for non-covalent spontaneous functionalization of graphitic surfaces (activated carbon, multi-walled carbon nanotubes (MWCNT), graphene), bridged by a spacer, allows for relatively easy preparation of heterogenous catalysts possessing a polyaminic macrocyclic unit for Pd(II) binding and a grafting unit for non-covalent functionalization. Linear commercial ligands, despite interesting, tend to form metal complexes of various character, could, in principle, also be replaced, we tried to improve the stability of our catalysts by adding a second identical pyrimidine moiety for each metal complex in order to achieve a firmer chemisorption. Linear commercial ligands, despite interesting, tend to form metal complexes of various stoichiometry, which complicate the picture and conflict with the idea of single identical catalytic active sites [36]. A more stable catalyst adsorbed via two pyrimidine moieties per metal center has been recently reported for the Sonogashira cross-coupling: no desorption issues were observed; yet, as the complex nature of the non-commercial (and thus sub-optimal, price-wise) polyaminic portion of the ligand offered further binding sites stabilizing Pd(0) NPs, deactivation still happened over time due to Pd(II) reduction [28].

Proofs of concept are important, but a rational catalysts design, we believe, should factor in all relevant costs, including those of starting materials. Herein we propose a novel catalyst based on the novel ligand H₃L (Figure 1), built around the commercial macrocyclic [9] aneN₃ (1,4,7-triazacyclononane) core, featuring three pyrimidine grafting moieties that are bridged by
simple ethylamine spacers (again from commercial reagents). This prosecutes the course of our investigation while shifting it further towards affordable costs and fostering stronger chemisorption (three grafting moieties).

![Figure 1. The H₃L ligand.](image)

### 2. Materials and Methods

#### 2.1. Synthesis of H₃L

2,2',2''-(((1,4,7-triazonane-1,4,7-triyl)tris(ethane-2,1-diyl))tris(azanediyl))tris(6-amino-3-methyl-5-nitrosopyrimidin-4(3H)-one (H₃L) has been synthesized according to Scheme 1.

N,O-bis(p-toluenesulfonyl)ethanolamine (1) had been previously prepared from commercial ethanolamine and p-toluenesulfonylchloride. Its cyclization afforded N-(p-toluenesulfonyl)aziridine (2) in 85% yield according to literature procedures [37]. The reaction of [9]aneN₃ (3) with three equivalents of 2 in CH₃CN under N₂ atmosphere afforded the tosylated polyamine 4 (N,N',N''-((1,4,7-triazonane-1,4,7-triyl)tris(ethane-2,1-diyl))tris(4-methylbenzenesulfonamide) in high yield (86%). Detosylation with concentrated sulfuric acid afforded the hydrogen sulfate salt of the polyamine 5 (2,2',2''-((1,4,7-triazonane-1,4,7-triyl)tris(ethane-2,1-diyl))tris(4-methylbenzenesulfonamide), which was converted to its free amine form with a Dowex 1 x 8 ionic exchange column that was activated with NaOH (global yield of 5: 79%) [38]. 5 is then reacted with the pyrimidine precursor 6. This molecule is accessible from commercial products on a multigram scale [39]. 0.5 g of 5 (1.94 mmol) was reacted with 1.15 g of 6 (6.19 mmol, i.e., 3.2 equivalents with respect to 5) in 500 cm³ of MeOH. After adding solid 6 in small portions within 3 h, the solution is heated to reflux for 1 h. Excess pyrimidine is eliminated by adding five drops of NH₃aq (33%), which converts it to 2,4-diamino-1-methyl-5-nitroso-6-oxopyrimidine, insoluble in MeOH. After storing overnight in a fridge, the resulting suspension has been filtered, and the MeOH solution evaporated to dryness under vacuum at r.t. to obtain H₃L as a deep purple solid compound. Yield 87%.

Elemental analysis calculated for C₂₇H₄₂N₁₈O₆: C, 45.37; H, 5.92; N, 35.27; O, 13.43. Found: C, 45.1; H, 5.6; N 34.9. ¹H NMR (D₂O, pH 1, 25 °C, 400 MHz) δ 3.95 (m, 6H), 3.45 (s, 9H), 3.41 (m, 12 H), 3.36 (m, 6H).

All of the solvents and reagents were purchased from Sigma–Aldrich (Milan, Italy).
2.2. Potentiometric Measurements

Ligand protonation constants were determined through potentiometric (pH-metric) titrations. The experiments were performed in 0.1 M NaCl at 298.1 ± 0.1 K using previously described procedures and instrumentation [40,41]. Emf data acquisition was performed with the software PASAT [42,43]. The Metrohm 6.0262.100 combined electrode was calibrated as a H⁺ concentration probe by titration of previously standardized amounts of HCl with CO₂-free NaOH solutions. The determination of the equivalent point was performed by Gran’s method [44], thus obtaining standard potential, E°, and ionic product of water (pKₐ = 13.78(1) in 0.1 M NaCl at 298.1 K). The software HYPERQUAD [45] was employed in order to calculate the stability constants from emf data. Ligand concentration was about 1 × 10⁻³ M in all experiments. The studied pH range was 2.0–11. At least three measurements were performed for H₃L protonation constants determination.
2.3. Spectroscopic Measurements

The $^1$H NMR spectra were recorded in D$_2$O at 25 °C on a Bruker AV300 spectrometer (Bruker Italia, Milano, Italy). Absorption spectra were recorded at 298 K on a Jasco V-670 spectrophotometer (Jasco Europe, Cremella, LC, Italy). Ligand concentration for UV–Vis spectra was 1.2 × 10$^{-5}$ M. In the case of Pd(II) complexes, batchwise UV titration is conducted, as reaching equilibrium can require up to several (3–4) days. X-ray photoelectron spectroscopy (XPS) spectra were recorded while using Al Kα radiation and a pass energy of 44 eV using previously reported instrumentation and methods [46–48], namely using a model TA10 X-ray source and a model HA100 hemispherical analyzer (VSW Scientific Instrument Limited, Manchester, UK) with a 12 channels detector. The C1s transition at 284.8 eV was used as a reference for obtaining the heteroatoms binding energies. It was not possible to carry out catalyst dismounting from the electrochemical cell and introduction in the XPS chamber maintaining the sample excluded from air.

2.4. Preparation of MWCNTs/H$_3$L and MWCNTs/H$_3$L-Pd(II) Catalysts

The preparation of said materials has been performed according to reported procedures [24–30,34]. In brief, MWCNTs (graphitized, 99.9% C, outer diameter 8–15 nm, inner diameter 3–5 nm, specific surface area 117 m$^2$/g, NanoAmor, TX, USA) were put in contact with a 1.0 mM aqueous solution of the ligand maintaining a 1 mg MWCNT/1 mL of solution ratio and gently shaken during three days. This is merely a precautionary timeframe, as the adsorption of polyamines functionalized with the same pyrimidine on MWCNTs happens via a Langmuir-type adsorption in a few (2–3) h [24,27,28,31,34]. The pH of the solution was conveniently adjusted to 6 (in the 5–8 pH range a single species exists, H$_4$L$^+$, cf. Figure S1). After the said time, solid was recovered and amount of adsorbed ligand quantified by measuring residual ligand in solution (exploiting the 300 nm isosbestic point characteristic of this pyrimidine chromophore). MWCNTs/H$_3$L was recovered as a solid product by filtration, briefly washed with milliQ water, and then dried in vacuum at r.t. to constant weight. Normally, imperfect adsorption (e.g., on edges or defect sites rather than on MWCNT surface) of similar ligands implies that a certain % of the adsorbed ligands (5–8%, generally) can be displaced; therefore, desorption of the obtained MWCNT/H$_3$L material was performed. MWCNT/H$_3$L material was placed in contact again with milli-Q water at pH 6 with a 1 mg/1 mL solid to solution ratio. After three days at r.t. under gentle shaking, the amount of desorbed ligand was quantified again, the solid recovered by filtration and dried. Desorption was repeated further on a portion of the material showing no further ligand desorption. The final amount of ligand adsorbed was 0.360 mmol H$_3$L/g MWCNT.

The metal decoration of MWCNTs/H$_3$L can also be easily achieved from aqueous solution. Accordingly, a weighted amount of MWCNTs/H$_3$L was suspended in a solution containing one equivalent of K$_2$PdCl$_4$ at pH 5 for six days. Given the high thermodynamic stability of Pd(II) complexes (simple ethylenediamine already affords complexation constants > 10$^{20}$, i.e., free energy change of the complexation reaction exceeds −115 kJ/mol [49]) all Pd(II) was sequestered. Despite this expected behavior, Vis spectra were still checked for PdCl$_4^{2−}$ bands, which were not present anymore after metal coordination. The resulting material contains 0.272 mmol of Pd/g, i.e., it has a Pd content of 2.90% by weight.

2.5. Preparation of the Modified Electrodes

Immobilization of the catalysts onto a rotating glassy carbon (GC) working electrode was achieved by preparing an ink consisting of MWCNT/H$_3$L-Pd(II) dispersed in a polymeric membrane (Nafion), which was then cast on the GC disk. The samples were prepared according to reported procedures [24]. In brief, composition of wet ink is as follows: catalyst 2%, water 52%, EtOH 26%, Nafion® ion exchange resin 20%.
2.6. Electrochemical Characterization

The rotating ring disk electrode (RRDE) experiments were performed using a Pyrex™ (Princeton Applied Research, Oak Ridge, TN, USA) glass cell that was filled with 0.1 M KOH solution. The reference electrode was a commercial (Princeton Applied Research) Ag/AgCl/sat. KCl with a potential of +197 mV with respect to the normal hydrogen electrode (NHE). A platinum gauze enclosed in a glass tube with porous bottom constitutes the counter electrode. A model 636A rotating ring-disk electrode and a PGSTAT 100N bi-potentiostat were employed. A Pine™ (Durham, NC, USA) ring-disk electrode (E6 Series), with platinum ring electrode and GC disk electrode covered with the catalyst layer (see above), was the working electrode. The size of electrodes: GC disk insert, \( \varnothing \text{5 mm} \) (\( \text{A} = 0.196 \text{ cm}^2 \)); Pt ring (\( \text{A} = 0.11 \text{ cm}^2 \)). Ring and disk currents were recorded simultaneously. \( \text{K}_3\text{Fe(CN)}_6 \) calibration in \( \text{N}_2 \)-saturated KOH 0.1 M was employed in order to determine the collection \( \text{HO}_2^- \) efficiency (\( N = I_{\text{ring}}/I_{\text{disk}} \)) of the ring-disk electrode. The fractional yields of \( \text{HO}_2^- \) in the ORR were calculated from the RRDE currents as \( X(\text{HO}_2^-) = (2I_{\text{ring}}/N)/(I_{\text{disk}} + I_{\text{ring}}/N) \). RRDE measurements were carried out at a scan rate of 5 mVs\(^{-1}\) with a 1600 rpm rotation rate, while the Pt ring electrode was held at +0.50 V versus Ag/AgCl/sat. KCl. \( \text{N}_2 \), or \( \text{O}_2 \) were used to purge the solution to achieve oxygen-free or oxygen-saturated electrolyte solution. Still, CVs in \( \text{N}_2 \) and \( \text{O}_2 \) are reported in the Supplementary Material (Figure S2).

For the optimal catalyst loading, sample 3, the number of exchanged electrons was also evaluated by working in RDE (rotating disk electrode) mode at a different scan speed and using the Koutechy–Levich (KL) equation. The general procedure has already been described elsewhere [18]. All of the onset potentials (\( E_{\text{on}} \)) reported are quantified with a general methodology that we previously reported [24]; one example is given in the Supplementary Material (Figure S3).

2.7. Scanning Transmission Electron Microscopy (STEM) Characterization

Electron microscopy characterization was done using a state of the art 200kV STEM microscope (JEOL, Tokyo, Japan, Model 2200FS) that was equipped with a probe and image aberration correctors (CEOS, Heidelberg, Germany). HAADF (high angle annular dark field) STEM images were acquired while using an 80 mrad inner collection semi-angle and an outer collection semi-angle of 180mrad. The samples for STEM analysis were prepared by finely crushing grains of MWCNT powder between glass slides and tapping holey carbon films supported by standard Cu TEM grids directly on to the thinner areas of the crushed powders. TEM grids were thermally pre-treated following the procedure that is described in ref [50]. Excess powder was then removed by flicking the grids onto the edges of the glass slides under a fume hood before transfer to single tilt TEM sample holders. The sample was kept in the TEM holder in a vacuum rig overnight before measurements to reduce moisture at \( 5 \times 10^{-7} \text{ mbar} \) vacuum pressure.

3. Results and Discussion

3.1. Ligand Protonation Equilibria

Table 1 lists equilibrium constants that were determined for ligand protonation by means of potentiometric (pH-metric) titrations (aqueous solution; 0.1 M NaCl; 298.1 K; pH range 2.0–11).

| Equilibrium | \( \log K \) |
|-------------|-------------|
| \( \text{H}_3\text{L}^2^- + 2\text{H}^+ = \text{H}_3\text{L} \) | 21.65(1) |
| \( \text{H}_3\text{L} + \text{H}^+ = \text{H}_4\text{L}^+ \) | 9.84(1) |
| \( \text{H}_4\text{L}^+ + \text{H}^+ = \text{H}_5\text{L}^{2+} \) | 3.42(3) |
| \( \text{H}_5\text{L}^{2+} + \text{H}^+ = \text{H}_6\text{L}^{3+} \) | 2.45(3) |
| \( \text{H}_6\text{L}^{3+} + 2\text{H}^+ = \text{H}_6\text{L}^{5+} \) | 4.54 (4) |
The species distribution diagram as a function of pH is reported as Supplementary Materials (Figure S1). In the studied pH range, \( \text{H}_3\text{L} \) behaves as a heptaprotic molecule. In agreement with the behavior of analogous polyamines functionalized with the same pyrimidine derivative \([27–36]\), the secondary amines directly connected to the pyrimidine residues can be deprotonated at \( \text{pH} \approx 11 \), while the pyrimidine nitroso groups are protonated below pH 2.5. These protonation equilibria are generally easily followed through UV-Vis measurements, as they significantly alter \( \pi-\pi^* \) transitions of the chromophore. The characteristic absorption wavelengths have been commented elsewhere \([27,28,31,34,36,51,52]\), and they are only affected slightly by the nature of appended polyamine. UV-Vis spectra of the ligand at different pHs, as reported in Figure 2, clearly show three absorption bands at 328, 271, and 230 nm. Combining potentiometric and spectroscopic measurements by superimposing species distribution diagram (derived from potentiometric data) and UV bands trends with pH (Figure 3), it is possible to rationalize the ligand protonation. This allows for understanding which of the observed protonation processes affect the chromophores, altering UV bands, and which the spectrophotometrically silent macrocycle. The most deprotonated species in solution is \( \text{HL}^{2-} \). The complete deprotonation to \( \text{L}^{3-} \) is not achieved in the 2.0–11.0 pH range of potentiometric measurements (extending pH range would offer no benefit due to large alkaline errors in a \( \text{Na}^+ \) medium). However, UV spectra confirm that, between pH 11 and 12, spectral changes are still observed, suggesting a \( \log K > 11.5 \) for \( \text{L}^{3-} \) protonation. The first observed two proton equilibrium (\( \log K = 21.65 \)) mainly involve the remaining equivalent anionic N-pyr sites, as demonstrated by important changes in the UV spectra. The entrance of the fourth proton has no further effect on UV-Vis bands, as it involves the macrocycle (\( \log K = 9.84 \), i.e., still a base of remarkable strength). \( \text{H}_4\text{L}^+ \) is the most persistent form of the ligand in solution, dominating the distribution diagram for at least four pH units: this is convenient for achieving MWCNT functionalization with a single species/protonation state. Formation of \( \text{H}_5\text{L}^{2+} \) (\( \log K = 3.42 \)) does not alter spectroscopic properties either, as the fifth proton is still localized onto the macrocycle. Starting from \( \text{H}_6\text{L}^{3+} \) (\( \log K = 2.45 \)) absorption spectra are again affected, signaling that protonation happens on pyr-NO groups. The further addition of two protons to give \( \text{H}_8\text{L}^{5+} \) (\( \log K = 4.54 \) for \( 2\text{H}^+ \)) has a profound effect on absorption spectra, as it happens on equivalent pyr-NO groups. The eventual last protonation of the macrocycle is not observed, as it most likely only happens in extremely acidic pH ranges. In the end, of the nine expected acid-base equilibria, seven can be fully characterized and 2 lay beyond the extremes of explored pH range, although all of them are fully rationalized, owing to associated spectral variations. The characteristic isosbestic point at around 300 nm is also particularly convenient for ligand quantification in solution.

![Figure 2. Acidic (left) (pink pH 0.83 to blue pH 4.2) and alkaline (right) (pink pH 4.2 to violet pH 12.3, first differing spectrum pH 9.1) branches of \( \text{H}_3\text{L} \) UV-Vis absorption spectra ([\( \text{H}_3\text{L} \]) = 1.2 \times 10^{-5} \text{ M}].
3.2. Pd(II) Complexes Formation

As mentioned in the experimental section, Pd(II) complexes of simple polyamines generally possess high thermodynamic stability [49]. The firm complexation of the metal cation is self-evident, as chelating and macrocyclic ligands afford further stabilization of such complexes. This being said, as determination of very large stability constants present inherent difficulties, which are further aggravated by Pd(II) kinetic inertness, the identification of all complex species formed by this metal ion under different pH conditions and the determination of the corresponding stability constants are generally impossible.

Nevertheless, batch spectrophotometric pH titration of a 1:1 Pd(II):H₃L 1.2 × 10⁻⁵ solution and comparison of molar attenuation coefficient with respect to H₃L alone (Figure 4) reveals complex formation. It appears that the Pd(II) complex is already formed in very acidic medium, in accordance with the high stability of the complex and, hence, of the ineffective proton competition, even at low pH, and it remains stable all the way to alkaline conditions, again in accordance with expected stability. As the ligand never regains its free spectrum, the complex appears to be resistant towards demetallation over the explored pH range. Full spectra are available in supporting material, as in Figure S4. This information, as for other related ligands [24], is useful as preliminary assessment to check that the complex can sustain (as found, cf. Sections 3.3 and 3.4) pH conditions that are relevant to ORR experiments.
3.3. Electrochemical Evaluation of Catalytic Performance

The results of the electrochemical studies are summarized in Table 2 and Figure 5.

Table 2. Catalytic performances of our catalyst.

| Sample | Ink Loading (mg) | \( n_{\text{theo}} \) (−0.5 V) \(^a\) | \( n_{\text{exp}} \) (−0.5 V) \(^b\) | \( N_{\text{exp}} \) \(^c\) | \( E_{\text{on}} \) (V) \(^d\) | \( E_{1/2} \) (V) \(^e\) |
|--------|-----------------|-----------------------------------|-----------------------------------|----------------|----------------|----------------|
| 1      | 2.4             | 4.00                              | 4.00                              | 0.229          | −0.0670        | −0.19          |
| 2      | 3.3             | 3.55                              | 3.93                              | 0.200          | −0.0288        | −0.11          |
| 3      | 8.7             | 3.68                              | 3.96                              | 0.233          | −0.0066        | −0.11          |
| 4      | 12.5            | 3.69                              | 3.88                              | 0.197          | −0.0605        | −0.15          |

\(^a\) number of exchanged electrons computed with theoretical N value at −0.5 V vs. reference electrode; \(^b\) number of exchanged electrons computed with experimentally determined N values at −0.5 V vs. reference electrode; \(^c\) experimentally determined collection efficiency number; \(^d\) Onset potential; \(^e\) Half-wave potential.

Increasing quantities of ink were initially deposited on the GC electrode and tested in order to screen for optimal electrode preparation in terms of catalyst loading. Although the deposition of small ink amount (2.4 mg, 1) was already sufficient for appreciating the catalytic effect, it was found that the deposition of an increased amount of catalyst would translate into superior catalytic performances, with optimum response at about four times the initial amount (8.7 mg, 3). The deposition of larger ink quantities resulted instead in being detrimental. This could be due to a non-significant increase of catalytic sites, which might remain non-accessible to gases, accompanied by a change of the texture of the electrode, potentially causing diffusion issues.

In the abovementioned optimal conditions, sample 3, the system is observed to display promising catalytic activity in the half cell studies, featuring an \( E_{\text{on}} \) of only −0.0067 V with respect to reference electrode and a number of exchanged electrons (\( n \)) close to four (3.96), i.e., promoting almost selectively direct water formation.

Calibration with \( K_3[Fe(CN)]_6 \) has been used to determine the experimental collection efficiency number in each case, so that accurate values for number of exchanged electrons could be derived.
We deemed it important to further verify the number of exchanged electrons, especially for sample 3, with an independent method, i.e., RDE experiments and Koutechy–Levich (KL) equation. Such a treatment allows for an independent indirect determination of $n$. To do so, LSVs at different spin rates have been performed (Figure 6), and meaningful information extracted from the double reciprocal $J_{\text{disk}}^{-1}$ vs. $\omega^{1/2}$ plot (with $\omega$ spin rate expressed in rad s$^{-1}$) by linear regression (Figure S5) (cf. Materials and Methods section). This is important in a preliminary study, such as ours, since the generation and accumulation of peroxide, via a concurrent $2e^-$ process, can be detrimental to the long-term viability of the catalyst. It was found that KL treatment affords, in this case, a very similar result ($n = 3.92, R^2$ for KL fitting 0.999) with respect to RRDE value (3.96).

In order to complete the half-cell characterization of the new catalyst, time viability has been probed. For this task the catalyst was left working for 1.5 h at $-0.4$ V vs. Ag/AgCl/sat. KCl (galvanostatic mode), with currents (and hence RRDE number of exchanged electrons) continuously monitored. This test showed no sign of drift of monitored parameters (Figure S6).
Figure 6. LSVs (linear sweep voltammograms) recorded at different spin rates (400; 800; 1200; 1600; 2000 rpm) at 20 mV/s scan rate of sample 3.

3.4. Before and After-Use Analysis

After the galvanostatic test, the catalyst was collected. XPS and STEM investigation were performed on the as-prepared and after-use catalysts.

XPS analysis mainly focused on the Pd region (Figure 7). This is the most diagnostic region for our system, potentially allowing to detect irreversible alterations due to the electrochemical treatment, the most common of which involve Pd oxidation state. In our conditions (alkaline media, reducing potential), both the replacement of N donor groups with O donors (i.e., transition from Pd(II) complexes to PdO) as well as reduction to Pd(0) are possible outcomes. XPS spectra reveals that the main oxidation state of the metal is $^{+2}$ and that it is mostly maintained even after use. The as prepared sample contains a small amount ($\approx 12\%$, as evaluated from XPS spectra) of more reduced Pd, giving rise to the minor shoulder at 333.4 eV (Figure S7). Similar behavior was showed by a graphene-supported Pd(II) complex prepared by the same route, and it is perhaps connected to preparation method and/or reducing character of leftover C impurities or Pd interactions with surface defects [28]. The main modifications that are introduced by the electrochemical treatment of the sample are the appearance of 348 eV peak and the broadening of Pd main peak at 337.5 eV. The analysis of Pd peak reveals a growth of Pd reduced component (now about 40%, Figure S7) with respect to Pd(II) main component and its shift towards typical Pd(0) values (observed 334.7 eV). The 348 eV is most likely spurious, and it has been attributed to a Ca impurity (typical peak at 351.1 eV is also observed). Ca is easily detected by XPS and little impurities might come from water samples and/or glassware (in consideration of the 0.1 M KOH environment). Both relative robustness of similar Pd(II) complexes towards redox processes [24] and cases featuring Pd(0) NPs growth [28] had previously been reported, depending on the nature of the ligand used and its ability to protect the metal’s oxidation state.
Figure 7. Before and after use (1.5 h −0.4 V vs. Ag/AgCl/sat. KCl) Pd XPS spectra of the catalyst.

As-prepared MWCNT/H₃L-Pd(II) can be seen in the STEM images shown in Figure 8. The MWCNTs are bundled together, forming grains of several micron size (Figure 8a). Images reveal the structure of the nanotubes and the distribution of the Pd. After deposition, the nanotubes maintain their crystallinity, as shown in the HR-bright field STEM images in Figure 8b–d. The typical contrast of graphitic planes of the nanotubes multi-walls is visible edge-on as well as the in plane in Figure 8d. HAADF STEM images, where the intensity is roughly proportional to the square of the atomic number of the elements in the field of view (I ∝ Z²), are able to highlight the presence of heavy elements over the carbon material. Pd is well distributed on to the nanotubes surface. Some larger clusters, with particle size of 2–3 nanometers Figure 8c–e), are distributed on the sample. In addition, in-between them, a finer dispersion of smaller Pd-based clusters of sub-nanometer size can be appreciated, as indicated by the white arrows in Figure 8e).
After use, the sample has a more compact structure, with MWCNTs embedded in amorphous material (See Figure 9a). Although the resolution is reduced by the presence of the amorphous material and its overall thickness, graphitic planes of the nanotubes walls are still clearly visible on the edges of the grains (Figure 9b). The brighter contrast in the HAADF STEM image in Figure 9c shows a Pd rearrangement on the surface, forming a network-like structure rather than single clusters. It could be concluded, given the limited impact that this rearrangement has on electrochemical performance, that the obtained Pd nanostructured network, as stabilized by the ligand, still allows a superior accessibility of the active sites.
Figure 9. STEM images of sample after use. (a) Overview BF image showing a more compact structure of MWCNT; (b) HR-BF STEM image showing MWCNTs imbedded into amorphous material. (c) HAADF STEM image of the same area, as in (a), Pd redistributed in a network-like structure.

4. Conclusions

Our approach to graphitic surface decoration afforded supported Pd(II) complexes with overall low (2.90% w/w) Pd content. The obtained catalyst displays considerable catalytic activity ($E_{on}$ up to $-0.0066$ V vs. Ag/AgCl/sat. KCl) and an almost exclusive ($\approx98\%$) $4e^-$ process. These results, especially $E_{on}$ values, surpass those that we already reported for MWCNT/HL2-Pd(II) [24] (Sample 3 $E_{on} = -0.0066$ V vs. MWCNT/HL2-Pd(II) $E_{on} = -0.014$ V [24]), both vs. Ag/AgCl/sat. KCl), which, in turn, exceeded the performance of a solid commercial Pt polycrystalline electrode. STEM investigation shows textural modification after use, still the comparison of before and after use catalyst (STEM, XPS, electrochemical) demonstrates the substantial preservation of catalytic performance. This can be already regarded as a success for an emerging strategy for catalyst preparation. In this sense, our green approach (water, r.t.) has been further improved by replacing custom polyazamacrocycles with commercial alternatives, cutting the catalyst cost. In terms of performance, the positive results of this preliminary half-cell study could support the catalyst as a good candidate for further investigation addressing its behaviour in a complete cell. Accessing true and stable SACs through a supramolecular route instead, while it is true that most Pd remained in its +2 state and its partial reduction to Pd(0) did not affect the catalytic study, still requires efforts in the ligand design stage in order to better prevent undesired reductive processes.

Supplementary Materials: The following are available online at http://www.mdpi.com/1996-1073/13/21/5539/s1, Figure S1. Species distribution diagram for the H3L ligand; Figure S2: sample still CVs in N2/O2; Figure S3: graphical definition of $E_{on}$ as employed in this study; Figure S4: H3L:Pd(II) 1:1 solution UV spectra.; Figure S5: KL regression plot; Figure S6 Time viability of the catalyst; Figure S7: XPS analysis showing different Pd peaks analysis.

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Abbreviations: List of Used Abbreviations (Alphabetical Order)

CV  Cyclic Voltammogram
E_{1/2}  Half-Wave Potential
E_{on}  Onset Potential
GC  Glassy Carbon
HAADF  High Angle Annular Dark Field
KL  Koutechy-Levich
LSV  Linear Sweep Voltammogram
MWCNTs  Multi-Walled Carbon Nanotubes
N  Collection Efficiency Number
n  Number of Exchanged Electrons
ORR  Oxygen Reduction Reaction
PGM  Platinum Group Metals
RDE  Rotating Disk Electrode
RRDE  Rotating Ring Disk Electrode
SAC  Single Atom Catalyst
STEM  Scanning Transmission Electron Microscopy
XPS  X-ray Photoelectron Spectroscopy

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