Nanoimpact Electrochemistry to Quantify the Transformation and Electrocatalytic Activity of Ni(OH)₂ Nanoparticles: Toward Sizeactivity Relationship at High Throughput

Mathias Miranda Vieira, Jean-François Lemineur, Jérôme Médard, Catherine Combellas, Frédéric Kanoufi, Jean-Marc Noël*

Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France.

AUTHOR INFORMATION

Corresponding Author

* jean-marc.noel@u-paris.fr

ABSTRACT

The fast establishment of structure-reactivity relationships is crucial to identify the most appropriate nanoparticles (NPs) for a given application. This requires the development of methodologies allowing, simultaneously, to unravel the NPs geometry and screen their reactivity. Herein nanoimpact electrochemistry (NIE) allows quantifying the transformation and measuring the electrocatalytic activity for the oxygen evolution reaction (OER) of >100 Ni(OH)₂ NPs of a wide range of size (NP radii from 25 to 100nm). This is achieved by scanning electrochemical microscopy in a generation/collection like mode, one electrode being used to electro-generate by local precipitation colloidal Ni(OH)₂ NPs and the second one to collect them by NIE. It allows i)

quantifying the reductive and oxidative conversion of the $Ni(OH)_2$ NPs and, ii) separating the electrochemical conversion and the OER electrocatalysis, leading to the evaluation of a structure-activity relationship.

TOC GRAPHICS



KEYWORDS Scanning electrochemical microscopy, single entity electrochemistry, electrosynthesis, electrocatalysis, insertion/disinsertion.

Transition metal oxides and hydroxides are promising abundant materials for energy conversion and storage.^{1–3} In the context of electrocatalytic reactions including the oxygen evolution reaction (OER),^{4–6} and ion insertion^{7–10}, nickel hydroxides present interesting physicochemical properties that can be optimized when the material is nanostructured.¹¹ Since those properties are closely linked to the composition, size and morphology of the nanoscale objects, methodologies allowing fast establishment of structure-activity relationships are essential to identify the most appropriate nanomaterial for a given application.

Single entity nanoimpact electrochemistry (NIE) has emerged as an easily implementable analytical technique to study at high throughput the reactivity of single nano-objects in solution, such as nanoparticles (NPs).^{12–17} The analysis of the current spikes (shape, and amplitude) arising from the collisions of individual NPs with a biased ultramicroelectrode (UME) allows extracting information about the colloidal system under investigation including nickel based NPs.¹⁸⁻²⁰ Generally, NIE experiments are carried out at an UME plunging in a large volume of diluted colloidal NPs. This allowed discriminating different NPs populations,²¹ deciphering mechanisms of NPs transformation,^{13,22} quantifying electrocatalytic reaction rates^{19,23-26} or physicochemical processes such as NPssubstrate^{21,27} or NPs-electrolyte²⁸ interactions. However, NIE remains rarely employed to establish structure-activity relationships (except *i.e.* anodic-cathodic nano-collisions at fast pulse potentials²⁹ and correlative methods combined with optical microscopies³⁰⁻³³). Herein, we propose a new NIE approach allowing to i) quantify at high throughput the conversion of electrosynthesized Ni(OH)₂ NPs and, ii) discriminate the electrochemical conversion of the NPs and their reactivity against OER, leading to a fast establishment of sizeactivity relationships. This is performed in an electrochemical generation/collection mode of scanning electrochemical microscopy (SECM).³⁴ This configuration, already employed to investigate NPs transformation³⁵ or dissolution,³⁶ uses one electrode to electrogenerate colloidal Ni(OH)₂ NPs from solution of Ni salts whereas a second one collects and probes their activity by NIE during their formation (Figures 1a and SI1). This offers several advantages: (i) the generating electrode allows on demand production of NPs with tuned concentration and size;³⁷ particularly a wide distribution of size is possible enabling to explore size-activity relationships and, (ii) the collecting microelectrode allows a non-destructive probing of the NPs at high

throughput since, thanks to their transport, only a limited but statistically relevant population of the NPs is titrated by NIE.³⁵



Figure 1. (a) NIE-SECM configuration to investigate the transformation and reactivity of $Ni(OH)_2$ NPs during their electrosynthesis. (b) CVs at a C-UME held at 25µm from an Au-ME in aqueous solution containing 0.1M KCl, 5mM NiCl₂ and 10mM citrate, before (black) and after (blue) polarizing the Au-ME at E_{Au} =-1.3V vs Ag-QRE. v=50mV.s⁻¹. (c) and (d): Zoom in the anodic and cathodic parts of the CVs, respectively

Microscopic evidence of Ni(OH)₂ **formation.** Details can be found in Sections 2 and 3 in SI. The NIE-SECM configuration uses a 500µm diameter gold microelectrode (Au-ME) generator to increase locally the pH of a solution containing 0.1M KCl, 5mM NiCl₂ and 10mM citrate by applying a potential to reduce H_2O (Equation (1)) and produce colloidal Ni(OH)₂ NPs in the diffusion layer of the Au-ME (Equation (2)).

$$4H_20 + 4e^- \to 4H0^- + 2H_2 \tag{1}$$

$$2HO^{-} + Ni^{2+} \rightarrow Ni(OH)_2 \tag{2}$$

The collector is a 7μ m carbon fiber ultramicroelectrode (C-UME) facing the Au-ME; it probes in its reaction layer the transformation of the NPs.

The formation of Ni(OH)₂ was first demonstrated by cyclic voltammetry (CV) at the C-UME held at 25µm from the Au-ME in the Ni²⁺ solution (Figure 1b). While the CV displays no Faradic current before applying any potential to the Au-ME, (Figure 1b, black), right after application of a constant potential, E_{Au} =-1.3V vs Ag-QRE for 100s (Figure 1b and c, blue), the CV shows a quasi reversible redox process centred at $E^{0} \approx 0.5$ V vs Ag-QRE. The corresponding oxidation and reduction peaks are characteristic of the oxidation of Ni(OH)₂ to Ni(OOH) and of the reverse reduction, respectively (Equation (3)).³⁸⁻⁴¹

$$Ni(OH)_2 \rightleftharpoons Ni(OOH) + H^+ + e^-$$
(3)

This oxidation is followed by another one at more anodic potential that is attributed to OER (Equation (4)) known to be catalysed by Ni(OOH) in alkaline media.^{38–41} A rather weak and not well-defined reduction peak is present between -0.3 and -0.9V that can be associated to the reduction of Ni(OH)₂ to metallic Ni (Figure 1d, Equation (5)).⁴² The

poor detection of this peak may be related to the partial reduction of $Ni(OH)_2$ thin films when probed by CV at scan rates > $0.2mV.s^{-1}.^{42}$

$$2H_20 \to 0_2 + 4H^+ + 4e^-$$
 (4)

$$Ni(OH)_2 + 2e^- \rightarrow Ni + 2HO^-$$
(5)

This experiment attests the presence of $Ni(OH)_2$ on the C-UME resulting from the formation of $Ni(OH)_2$ within the diffusion layer of the Au-ME. Moreover, "noisy" stochastic current spikes on the CV suggest the presence of free NPs colliding stochastically the C-UME.⁴³

Ni(OH)₂ NPs characterization by reductive nanoimpacts. The formation of Ni(OH)₂ NPs was characterized *in situ* by reductive NIE. Chronoamperograms were recorded at a C-UME polarized at E_{C} =-0.6V vs Ag-QRE for 120s, while the Au-ME was polarized at E_{Au} =-1.3V vs Ag-QRE (Figure SI2a). The current at the C-UME (Figures 2a and SI2a) reveals reduction current spikes immediately after ca. 0.8s (Figure SI2a). These spikes showing a sharp rise of absolute current followed by a slower return to baseline (inset of Figure 2a) are characteristic of NPs transformation,⁴⁴ here attributed to the reduction of individual Ni(OH)₂ NPs to Ni NPs (Equation (5), sketched in Figure 2b). The reduction spikes last for few tens of milliseconds (see nanoimpacts inset of Figure 2a and histogram of Figure SI2b). This rapid reduction at the scale of individual NPs is in contrast to the slow reduction during CV for adsorbed Ni(OH)₂.⁴² The enhanced activity of colloids during electrochemical collisions compared to adsorbed ones has been already

highlighted.^{25,33} The charge, Q_{red} , associated to each current spike (inset of Figure 2c), allowed estimating an apparent size of the electrochemically reduced Ni(OH)₂ NPs, defined as an equivalent NP radius, $r_{EC,red}$ (Equation (6)):

$$r_{EC,red} = \sqrt[3]{\frac{3Q_{red}M}{4\pi z F \rho}}$$
(6)

with M and ρ the molar mass and density of Ni(OH)₂, F the Faraday constant and z the number of electron exchanged.



Figure 2. Ni(OH)₂ NPs characterization by reductive NIE. (a) Chronoamperogram at the C-UME biased at E_{C} =-0.6 V vs Ag-QRE, held at 25µm from the Au-ME polarized at E_{Au} =-1.2V vs Ag-QRE. Inset: zoom of one spike. (b) Mechanism of the reduction of Ni(OH)₂ NPs to Ni. (c) Size distribution obtained from the nanoimpacts (green) and from the SEM images (black). Inset: example of the charge considered to evaluate the size of one NP.

The radius distribution is centred at $r_{EC,red}$ =37±7nm (Figure 2c, green), in good agreement with the one obtained by SEM (black histogram in Figure 2c) suggesting a complete electrochemical reduction of the Ni(OH)₂ NPs. SEM reveals spherical-like NPs attributed to Ni(OH)₂ NPs (Figure SI3a). The good agreements between both distributions coupled to the long time period observed between each collision event (>100ms for >90% of NPs) suggests that one collision corresponds to one NP rather than multiple collisions of one single NPs.^{33,45–48}

Ni(OH)₂ NPs characterization by oxidative nanoimpacts. In another set of experiments, the C-UME, placed at 50 μ m from the Au ME, was biased at E_C=+1V vs Ag-QRE. The current transient at the C-UME (Figure 3a) reveals oxidation spikes. Their shape is significantly different from that of the reductive ones since the peak-shaped transient is immediately followed by a current step before return to baseline (inset of Figure 3a). A fast EC response followed by slow activity was interpreted as a consequence of NPelectrode interaction for Ag NPs oxidative dissolution.²¹ However, here, the EC charge of the current transient leads to a large electrochemical overestimation of the NP size $(r_{EC,ox}=105\pm34$ nm, see Figure 3b) compared to the geometrical size distribution established from SEM images (r_{SEM,ox}=71±20nm, see Figure SI4) considering a oneelectron oxidative full conversion of Ni(OH)₂ into Ni(OOH).³⁸⁻⁴¹ Noteworthy, the highest size distribution recorded under this experimental condition is expected as the NPs size should increase with the distance between both electrodes.³⁵ It demonstrates the potentiality of such configuration to probe the reactivity of a wide distribution of NPs sizes.

The discrepancy between the distribution observed by NIE and SEM reveals that the electrochemical transient contains not only the contribution of the EC conversion of

Ni(OH)₂ into NiOOH, but also another contribution. The current step following the current spike is typical of electrocatalytic NIE,^{24,49,50} and is attributed to the newly formed NiOOH NPs that drive the electrocatalysis of water oxidation (Figure 3c). Indeed at the 1V vs Ag-QRE electrode polarization, both Ni(OH)₂ oxidation and electrocatalysis of OER by the resulting NiOOH NPs can occur (Figure 1b). Additionally, looking precisely to the current spikes for some of them (~10%), multi steps within ~50ms can be visualized for which the current spike always appears at the first step and is followed by discontinued current steps (Figure SI5). This allows distinguishing multiple collision events of individual Ni(OH)₂ NPs for which the first step corresponds to their oxidation followed by their catalytic activity.



Figure 3. Ni(OH)₂ NPs characterization by oxidative nanoimpacts. (a) Chronoamperogram at the C-UME biased at $E_C=1V$ vs Ag-QRE, held at 50µm from the Au-ME polarized at $E_{Au}=-1.2V$ vs Ag-QRE. Inset: zoom of one spike. (b) Histogram showing the size distribution obtained from nanoimpacts (red) and from SEM images (black). Inset: example of the charge considered to evaluate the size of one NP. (c) Mechanism proposed for Ni(OH)₂ NPs oxidation followed by OER.

Even if both contributions are somehow interrelated, a first simplification was made by assuming that the fast current spike corresponds to Ni(OH)₂ conversion into NiOOH while the longer current step corresponds to electrocatalysis. The NPs sizes was estimated by considering only the charge Q_{ox} under the current spike (red part in inset of Figure 3b), which gives an electrochemical NP radius distribution, $r_{EC,ox}=74\pm18$ nm in excellent agreement with that obtained by SEM (Figure 3b).

Therefore, the current steps were interpreted as the electrocatalysis of OER by NiOOH NPs (Figures 4a and SI6a, purple area). Such electrocatalysis was analyzed for different NP sizes ($r_{EC,ox}$ extracted from Q_{ox}). For N=140 EC spikes, the electrocatalytic activity lasts for 80±5ms and is independent of the NPs size (Figure SI6c). It ends by an abrupt current drop, in line with the reported OER activity on CoFe₂O₄ NPs,²⁴ likely due to a rapid electrical disconnection of the NP rather than its progressive deactivation due to O₂ gas accumulation around the NP. Indeed, similarly to what has been observed,²⁴ there is no correlation between the amount of O₂ produced (charge of the current step, Q_{cat}) and the size of the NPs (Figure SI6c).

Structure-activity relationship of Ni(OH)₂ conversion and electrocatalysis. Firstly, the dynamics of the NP conversion was analysed for the oxidative and reductive processes. In both cases the amplitude and duration of the nanoimpact current spikes increase with the NPs radius, r_{EC} (Figure SI7a,b and SI7c,d, respectively). Noteworthy, the reduction of Ni(OH)₂ at more reductive potential (-0.8V) reveals chronoamperogram (Figure SI8a) and distribution of nanoimpact charge, current amplitude and duration similar to those at -0.6V (Figure SI8b). It indicates that the reduction process is limited by mass transfer rather than by electrochemical kinetics. Owing to the relatively large NP size (>50nm), a limitation by the solid-state diffusion of HO⁻ is expected within the NP during its transformation (inset of Figure 4b, green). Assuming that the size variation of the NP is not significantly altered during the conversion (<15% NP size variation expected from Ni(OH)₂ to Ni(OOH) ⁵¹), the peak duration, Δt , should be related to the initial NP radius, r_0 , (Equation (8)):⁵²

$$\Delta t = \frac{r_0^2}{3V_M^{Ni(OH)_2} D_{HO} - C_{max}}$$
(8)

with D the diffusion coefficient of HO⁻ within the Ni(OH)₂ NP and C_{max} its maximum concentration (mol.cm⁻³) $\approx 2/V_M^{Ni(OH)_2}$ in the Ni(OH)₂ NP. This yields $r_0^2 \approx 6D_{HO} \Delta t$.



Figure 4: (a) Analysis of an individual current signal for oxidative and reductive nanoimpacts. (b) NP radius vs the square root of peak duration for $E_{C UME}$ =-0.6 V (green) and +1V (red). Dotted lines: linear fits to determine the diffusion coefficients of HO⁻ (green) and H⁺ (red) within the NPs using Equation (8). (c) Turnover frequency (TOF) of individual NPs as a function of their electrochemical radius $r_{EC,ox}$.

 r_{EC} , evaluated at -0,6V evolves linearly with the square root of the peak duration (Figure 4b, green), confirming a diffusional limitation. The same relationship is found at -0.8V (Figure SI8e). By fitting the scatterplots by Equation (8), the diffusion coefficient of HO⁻ ions in Ni(OH)₂ NPs during their reduction is $D_{HO} \approx 6 \times 10^{-11} \text{ cm}^2 \text{ s}^{-1}$, a value within the range of solid phase ion diffusion.

The same analysis for the oxidation of Ni(OH)₂ NPs (Figure 4b, red) evidences similar linear relationship with a higher slope. The limiting process is then the disinsertion of H⁺ within the NPs (inset of Figure 4b). By fitting the data with Equation (8) and considering the maximum H⁺ concentration that can be removed from the structure, C_{max,H^+} (mol.cm⁻ $^{3})\approx 1/V_{M}^{Ni(OH)_{2}}$, the diffusion coefficient of H⁺ is $D_{H^+}\approx 5 \times 10^{-10}$ cm².s⁻¹, a value within the range of those already reported.^{41,53} D_{H^+} is one order of magnitude faster than D_{HO}extracted from the analysis of the reduction spike, which is attributed to the lower sterically hindrance of H⁺ compared to HO⁻. This involves a faster disinsertion process for oxidation of Ni(OH)₂ than for reduction.

Secondly, the catalytic charge, Q_{cat} , exchanged during the current step was measured for each nanoimpact. From Q_{cat} , a turnover frequency, *TOF*, was evaluated providing the catalytic activity for OER at the single NP level using equation (9), (Section 3 in SI):

$$TOF = \frac{Q_{cat}}{4Q_{ox}\Delta t_{cat}} \qquad (9)$$

with Q_{ox} the charge for the oxidative spike for each nanoimpact, (see Figure 4a) and Δt_{cat} the duration of the current step attributed to the catalytic process. This assumes that the entire volume of the NP is involved in the OER electrocatalysis, allowing a comparison with the previous work performed on individual Ni(OH)₂ NPs immobilized on a nanoelectrode.⁴¹ The *TOF* values decrease with the NP radii (Figure 4c) and are of the same order of magnitude as those previously reported for single NPs electrodeposited on a nanoelectrode.⁴¹

To summarize SECM was used in a generation/collection configuration to simultaneously electrosynthesize Ni(OH)₂ NPs (NPs radii ranging from 25 to 100nm) and investigate their electrochemical conversion and reactivity during their formation. Such SECM-NIE strategy allows analyzing the reductive transformation of Ni(OH)₂ NPs into metallic Ni and their oxidative conversion into NiOOH NPs. Both conversions are controlled by HO⁻ or H⁺ ions transport within the NPs. The faster NPs oxidative conversion compared to the reductive one suggests a ~10 times faster diffusion coefficient of H^+ with respect to $HO^$ in the Ni-based solid oxide. The oxidative nanoimpact transients carry more electrochemical information. Indeed, a ca. 100 ms longer current tail provides valuable information on the electrocatalysis of OER by NiOOH NPs, as confirmed by the TOF values that decrease when increasing the NP size in agreement with previous study for Ni(OH)₂ NPs immobilized on nanoelectrodes.⁴¹ The separation of electrochemical conversion and electrocatalysis allowed, for the first time, determining size-reactivity approach relationships. Both the simple electrosynthetic and the complete characterization of the NPs in the NIE-SECM configuration could be extended to other

types of oxides NPs, implementable in microfluidic platforms. Besides, the double process characterization, here exemplified for OER, opens new ways toward the determination of structure-activity relationships at high throughput for other NPs involved in anodic or cathodic electrocatalysis.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at XXX.

Material and methods (electrosynthetic procedure, SEM, electrochemical experiments, SEM), dark field optical microscopy experiments, characterization of the NPs by nanoimpacts and determination of the turn over frequency.

AUTHOR INFORMATION

Corresponding author

Jean-Marc Noel - Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France.

orcid.org/0000-0001-8278-7748

Email: jean-marc.noel@cnrs.fr

Authors

Mathias Miranda Vieira - Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France.

Jean-François Lemineur - Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France.

Jérôme Médard - Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France.

Catherine Combellas - Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France.

Frédéric Kanoufi - Université Paris Cité, ITODYS, CNRS, F-75013 Paris, France. 0000-0002-9784-2380

Notes

The authors declare no competing financial interests.

ACKNOWLEDGMENT

We thank CNRS, the ANR JCJC program (PIRaNa project, ANR-20-CE42-0001) and the Emergence call from the Université de Paris within the Investissement d'Avenir program under reference ANR-18-IDEX-0001 for financial support.

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