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Recent advances in the understanding of Ni-based catalysts for the oxidation of hydrogen-containing fuels in alkaline media

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ABSTRACT.

Nickel is a very abundant transition metal in the Earth crust, and finds numerous applications in electrochemical processes where metallic Ni or its oxides are thermodynamically stable, particularly in alkaline environments. This contribution addresses electrocatalytic properties of Ni-based catalysts in reactions of fuel oxidation in alkaline media. It firstly details the electrochemical behavior of Ni in alkaline media and approaches to determine the active surface area of Ni electrodes. Secondly, the electrocatalytic activities of Ni-based electrocatalysts for the alkaline hydrogen oxidation reaction are described (an endeavor for the development of anion exchange membrane fuel cells), along with a detailed analysis of the strategies put forward to improve them. It is notably shown that the state of Ni surface (oxidized or reduced) largely determines its electrocatalytic activity. This state of the surface also conveys a pivotal importance regarding the activity of Ni for the oxidation of complex fuels (borohydride, boranes and hydrazine). Finally, emphasis is made on the durability of Ni-based catalysts in alkaline environments. It is shown that in such media, the materials durability of Ni-based electrodes can be high, but this does not necessarily warrant stable electrocatalytic activity, owing to possible deactivation following surface oxide or bulk hydride formation in operation.

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1. Introduction

Recent years have witnessed tremendous increase in the interest towards electrocatalysis in alkaline medium. This is largely due to the success in the development of more potent anion exchange membranes, but also to the desire to replace scarce and expensive noble metals by more abundant platinum group metal (PGM)-free materials in fuel and electrolysis cells. ^{1–4} Ni is a ubiquitous element widely applied in both heterogeneous catalysis and electrocatalysis due to its low cost, high abundance and high corrosion stability in alkaline media.⁵ In heterogeneous catalysis Ni is mostly used for hydrogenation and reforming of hydrocarbons, largely owing to its ability to dissociatively adsorb (similar to Pd and Pt) hydrogen providing M-H_{ad} active species, while being a much cheaper alternative.⁶ Ni is indispensable in energy storage/conversion systems, such as batteries and electrochemical capacitors, ^{7–9} Li-ion batteries, ^{10,11} solar cells ¹² where it is used in the form of (oxy)hydroxides, simple and complex oxides. In electrocatalysis, Ni-based materials are used for catalyzing oxygen ^{13–16} and hydrogen evolution ^{17–19} reactions. Ni is also widely encountered as a current collector/substrate for catalyst' deposition (see *e.g.* Refs.^{20–22}).

This review considers electrocatalytic properties of Ni as a promising anode material for fuel cells fed either with hydrogen or complex fuels (borohydride, boranes and hydrazine) in alkaline medium. We start by discussing electrochemical properties of Ni in alkaline medium. Under ambient conditions, Ni is prone to oxidation resulting in the formation of ca. 1-2 nm thick surface oxide layer composed of NiO and Ni(OH)₂.^{23–25} The presence of oxides on the Ni surface can greatly affect its catalytic properties, especially in low-temperature electrocatalysis, where, contrary to high temperature heterogeneous catalysis, their reduction may not be facile. Section 2.1. is devoted to the electrochemical oxidation of Ni in different potential intervals as well as hydrogen adsorption/absorption. Section 2.2. discusses the determination of the surface area of Ni electrocatalysts. This issue is of paramount importance for heterogeneous catalysis and electrocatalysis, since it forms the basis for benchmarking Ni-based electrocatalysts. Section 3 of the

review summarizes recent advances in the understanding of electrocatalysis of the hydrogen oxidation reaction (HOR) on Ni, with a specific emphasis on how it is affected by the surface state of Ni. It considers the development of Ni and Ni-based electrocatalysts for the HOR and strategies used to prepare highly active materials for the anodes of alkaline fuel cells (AFC) and anion exchange membrane fuel cells (AEMFCs). Meanwhile, the hydrogen evolution reaction (HER) on Ni-based electrodes, which has been widely discussed and reviewed in the past ^{17,18,26–28}, is only briefly discussed, primarily in the potential region in the vicinity of the equilibrium potential of the hydrogen electrode, when it enters in competition with the oxidation of complex fuels. Section 4 is devoted to the oxidation of complex fuels, such as borohydride, boranes and hydrazine, which can be used in direct liquid fuel cells (DLFC). Finally, the stability issues of Ni-based electrocatalysts are briefly presented in Section 5.

2. Electrochemical properties of Ni electrocatalysts

2.1. Electrochemical behavior of Ni in alkaline media

Electrochemical properties of Ni electrodes have been widely studied since the beginning of the XXth century, starting from the works of Emelianova,²⁹ Frumkin et al.,^{30,31} Conway et al.,^{32–39} Breiter et al.,^{40,41} Bockris and Potter ⁴² as well as some others. The behavior of Ni in alkaline media has been recently summarized by the groups of MacDougall ^{43,44} and Jerkiewicz,^{45–47} who discussed in detail various potential-dependent processes taking place on the surface of Ni electrodes. In addition, Oshchepkov et al. have recently pointed out to the influence of the surface state of Ni electrodes on their HOR/HER activity in the potential interval -0.2 < E < 0.5 V vs RHE.^{48–52} Below the most relevant information regarding electrochemistry of Ni electrodes in alkaline media is briefly summarized.

In order to facilitate the discussion about characteristic features of cyclic voltammograms (CVs) of a Ni electrode (Figure 1a) as well as its electrocatalytic properties in a number of oxidation reactions, it is

convenient to mark out two potential intervals, depending on the reversibility of the Ni oxidation. The latter include the "*low-potential region*" at $-0.2 \le E \le 0.5$ V vs RHE, where the surface of a Ni electrode can be reversibly oxidized and reduced, and the "*high-potential region*" at $0.5 \le E \le 1.6$ V vs RHE, in which the formation of irreversible, passivating, Ni oxides occurs, the degree of irreversibility depending on the anodic potential limit. The *high-potential region* can be additionally split into two sub-regions, namely $0.5 \le E \le 1.2$ V vs RHE and $1.2 \le E \le 1.6$ V vs RHE depending on the degree of Ni oxidation, as discussed further in the text. Below E = 0 V vs RHE and above ca. E = 1.5 V vs RHE the evolution of hydrogen and oxygen take place, respectively. In what follows, the processes taking place on Ni electrodes in the *low-* and the *high-potential regions* will be briefly discussed, mainly focusing on the former one, due to controversies existing in the literature.



Figure 1. (a) CVs of a polycrystalline Ni electrode in N₂-saturated 0.1 mol L⁻¹ NaOH at a sweep rate $v = 50 \text{ mV s}^{-1}$ and $T = 25 \,^{\circ}$ C. The first CV in panel a (dash-dotted blue curve) covers the potential range from E = -0.2 to 0.5 V vs RHE, while two other CVs (first scan – solid black curve and second scan – dashed red curve) cover the potential range from E = -0.3 to 1.6 V vs RHE. The figure is inspired by some previous works 47,53-56 (b) CVs obtained for metallic and partially oxidized polycrystalline Ni electrodes in N₂-saturated 0.1 mol L⁻¹ NaOH at a sweep rate $v = 20 \text{ mV s}^{-1}$ and $T = 25 \,^{\circ}$ C. Adapted from Ref. 48 with permission from Springer.

<u>*a-Ni(OH)*</u>² formation

It is generally accepted that the main process occurring in *the low-potential region* is the reversible oxidation of Ni with the formation of surface α -Ni(OH)₂ during the anodic sweep, followed by its reduction in the cathodic sweep.^{43,45,53,55,57} The common use of the term ' α -Ni(OH)₂' (suggesting formation of a 3D structure with H₂O molecules incorporated between Ni(OH)₂ layers ⁴³) is explained by the fact that it likely grows on the Ni surface through formation of 3D islands of α -Ni(OH)₂. This has been evidenced by the use of *in situ* scanning tunneling microscopy ^{58,59} as well as by the analysis of CVs at low (< 263 K) temperatures. ⁵⁵ According to the place-exchange model proposed by Conway et al. ⁶⁰ and later adapted by Alsabet et al., ⁴⁵ the formation of α -Ni(OH)₂ proceeds through three consecutive steps: firstly, the discharge of OH⁻ anions at the Ni surface and formation of adsorbed OH species (Ni-OH_{ad}), secondly, the incorporation of OH into subsurface between the first and the second layer of Ni atoms ((OH-Ni)_{quasi-3D lattice}) and thirdly, chemisorption of the second monolayer of OH groups on the (OH-Ni)_{quasi-3D lattice}:

$$Ni + OH^{-} \rightleftharpoons Ni - OH_{ad} + e^{-}$$

$$(2.1, fast)$$

$$(2.2 - 4 - 1.4 - 1$$

$$Ni-OH_{ad} \rightleftharpoons (OH-Ni)_{quasi-3D \ lattice}$$
 (2.2, rate determining step)

$$(OH-Ni)_{quasi-3D \ lattice} + OH \rightleftharpoons Ni(OH)_{2,ad} + e^{-1}$$
 (2.3, fast)

The existence of intermediate Ni-OH_{ad} species was suggested on the basis of electrochemical (mainly cyclic voltammetry) ^{41,61} and *in situ* scanning tunneling microscopy ⁵⁹ measurements.

The influence of crystallographic orientation of the Ni surface on its electrochemical behavior has been studied in several works, both in acid and in alkaline media $^{40,41,54,59,62-65}$. While early studies 40,54,64 did not find noticeable differences in the α -Ni(OH)₂ formation for low-index Ni planes (111), (110), (100), recent work of Esau et al. clearly demonstrates the differences in the CV profile depending on the crystallographic orientation of the Ni surface.⁶⁵ In particular, the anodic peak potential of α -Ni(OH)₂ formation shifts positive following the order (100) < (110) < (111), and an additional (pre-)peak at lower potential values shows up in the case of Ni (100) and Ni (110) planes (with a less intense pre-peak in the latter case). This observation agrees with DFT calculations,^{66,67} which point to structure-sensitive adsorption of OH and other species on the Ni surface (*vide infra*). Meanwhile, the potential of the cathodic peak is very similar for the three Ni single crystal planes.

In addition to α -Ni(OH)₂ discussed above, some works mention possible formation of NiO species sandwiched between the metal and the outer layer of α -Ni(OH)₂. ^{23,68,69} The amount of NiO formed during potential cycling in the *low-potential region* can be considered insignificant, as evidenced by *in situ* Raman ⁵⁷ and IR spectroscopy⁷⁰ as well as *ex situ* X-ray photoelectron spectroscopy (XPS).⁷¹ Note however, that NiO might be also formed on the surface of Ni electrode upon contact with oxygen species, followed by partial transformation of this NiO layer into α -Ni(OH)₂ after either contact with humid air or immersion in an aqueous alkaline electrolyte, thus resulting in a three-layer structure Ni | NiO | α -Ni(OH)₂.²³ Finally, it is worth mentioning that, contrary to acidic medium, anodic dissolution and cathodic redeposition of Ni, which might be expected from Pourbaix diagram,⁷² have not been observed in alkaline electrolytes in the low-potential interval.^{55,69} It is remarkable that for a Ni electrode, whose surface is partially covered by stable Ni oxides (*e.g.* NiO or β -Ni(OH)₂), the latter formed either through oxidation of Ni in air (*i.e.* chemical oxidation) or via excursions to high anodic potentials (*i.e.* electrochemical oxidation), the shape of CV changes substantially, an additional anodic peak showing up for 0.05 < E < 0.15 V vs RHE (Figures 1a and 1b). Such changes are reproducible and have been reported in several publications.^{41,64,68,73–79} The peak potential and the corresponding current density significantly depend on the surface state of the electrode, as well as on the cathodic potential limit of the CV. ^{50,76,79} The origin of that second anodic peak is still debated, some authors attributing it to the formation of Ni–OH_{ad},^{48,64} the other – to the oxidation of adsorbed/absorbed hydrogen,^{41,73,74,80} as discussed below.

Hydrogen adsorption

Hydrogen adsorption is of paramount importance for the HOR/HER on metal electrodes (*vide infra*), and thus has widely been discussed in the literature (see Refs.^{81–83} and references therein). On Pt and other PGM electrodes, there has been convincing evidence for the existence of two types of phenomena, usually addressed as "under-potential" deposition of hydrogen (H_{upd}) at potentials above E = 0 V vs RHE and "over-potential" deposition of hydrogen (H_{opd}) at potentials below E = 0 V vs RHE.^{81,82} While the presence of H_{opd} on Ni is commonly accepted, there are significant controversies in the literature regarding the existence of H_{upd} on Ni electrodes. Some researchers insist that, above E = 0 V vs RHE, a Ni electrode is covered by adsorbed oxygenated species (OH_{ad} / OH⁻_{ad} / H₂O_{ad}) which prevent adsorption of H_{ad}.^{45,69,81,84,85} Other publications present evidence for the existence of H_{upd} on the surface of Ni electrodes (even if it is still being debated). More specifically, Hu and Wen⁷³ investigated electrocatalytic hydrogenation of p-nitroaniline in alkaline medium, and found out that the reaction occurs on Ni and on Pt electrodes roughly in the same potential interval, that is below ca. E = 0.2 V vs RHE. Considering formation of H_{upd} on Pt in this potential interval, they inferred the existence of adsorbed hydrogen on the

surface of Ni under these conditions as well. Note that the authors of Ref.⁷³ demonstrated that prior oxidation of the Ni electrode surface at T = 400 °C leads to complete suppression of p-nitroaniline hydrogenation, and attributed this to the blocking of hydrogen adsorption by the formed passive layer of NiO, which cannot be electrochemically reduced due to likely insufficient electronic conductivity. In addition, several authors documented intersection (named "isopotential") points in the forward and backward scans of CVs recorded until different positive potential limits [for example, Hu and Wen,⁷³ Katic et al.⁸⁶], which they attributed to α -Ni(OH)₂ formation and hydrogen adsorption/desorption occurring in the same potential interval (above E = 0 V *vs* RHE) and competing for the Ni surface sites. This is in agreement with the adsorption isotherms calculated by Pshenichnikov for the hydrogen and oxygen adsorption from the gas phase.⁷⁴ Finally, some of us ^{48–51} studied the HOR on Ni by combining electrochemical measurements with microkinetic modeling, and concluded that assuming the existence of H_{upd} on Ni allows the most consistent explanation of a large amount of experimental data for the HOR on Ni electrodes subjected to various surface treatments (reductive *vs* oxidative), as discussed in more detail in section 3 of this review.

Along with experimental works, there has been significant number of computational studies devoted to the adsorption of H, OH, O and other species on Ni. The binding energy values computed using density functional theory (DFT) and reported by different research groups, vary depending on the density functional used for the calculations. However, some general trends can be identified. Firstly, it is widely accepted that oxygen atoms (O_{ads}) adsorb much stronger that the hydrogen atoms.^{66,87,88} In the meantime, comparison of H and OH binding energies is less straightforward, some studies reporting stronger adsorption of OH compared to H, while the other presenting similar binding energy values. For example, Bai et al. ⁸⁸ computed the following binding energy values (in eV) for H and OH on different sites on Ni(111) using self-consistent GGAPW91 functional (at a ¹/₄ monolayer coverage): H (-2.89 (fcc); -2.88

(hcp)); OH (-2.47 (top); -2.92 (bridge); -3.01 (fcc); -2.91 (hcp)). Thus, one may see that the binding energy strongly depends on the type of surface site, and that the H and OH adsorption energy is comparable. Taylor at al.⁸⁹ performed *ab initio* calculations in order to probe influence of water and surface charge on the hydrogen adsorption on Ni(111). Based on calculated Gibbs energies of adsorption, the authors constructed a hydrogen over Ni(111) potential-pH diagram, which corroborates hydrogen adsorption on Ni above E = 0 V vs RHE (hence, H_{upd}), and suggests that equilibrium potential of the H_{ad}/H^+ couple shifts positive with pH. Besides, it is worth noting that according to theoretical calculations, adsorption energies for H_{ad} and OH_{ad} demonstrate different coverage dependence. Indeed, according to Greeley and Mavrikakis, the strength of H_{ad} to Ni(111) only slightly decreases from 0 to 1 monolayer coverage.90 Meanwhile, Juares et al. 87 found that because of their high negative charge, OH_{ad} on Ni strongly repel each other, leading to a significant decrease of the adsorption strength with the coverage. As a result, under certain conditions, adsorption of hydrogen may become more favorable compared to that of OH_{ad} species. Note also that, according to the DFT calculations,^{91,92} the state of the Ni surface (namely the presence of adsorbed oxygen in the vicinity of Ni⁰ sites) largely affects the adsorption energy of the H_{ad}. This conclusion is also supported by microkinetic modeling^{48–50} of the HOR on Ni electrodes as discussed in the following sections.

In the end, if one acknowledges the existence of H_{upd} on Ni in the potential interval from E = 0 to 0.2 V vs RHE, the anodic CV peak at ca. E = 0.3 V vs RHE in Figure 1a should not be interpreted as α -Ni(OH)₂ formation on free Ni surface sites but rather as H_{upd} replacement by OH_{ad} followed by the conversion of the latter into a surface α -Ni(OH)₂ layer (Eqs. 2.2 and 2.3). In this context, it is instructive to recall hydrogen UPD on Pt, which until recently had been considered as a well-understood phenomenon. Meanwhile, new evidence provided by electrochemical experiments, DFT calculations and surface science studies (see Janik et al.⁹³ and references therein) confirms that certain peaks observed in the

potential interval of hydrogen UPD in fact do not correspond to H_{upd} , but rather to the replacement of adsorbed H atoms by OH on low-co-ordinated Pt sites.

Hydrogen absorption

There are numerous evidence of hydrogen **ab**sorption in the Ni lattice, which occurs at potentials significantly below E = 0 V vs RHE, especially upon prolonged water electrolysis conditions.^{43,69,75,77,78,94–96} Since hydrogen diffusion in the Ni lattice under ambient conditions is slow, in a transient (*e.g.* fast CV) experiment the release of these hydrogen species from the Ni lattice may be detected as an anodic current at potentials above E = 0 V vs RHE, ^{43,95,97} that is in the potential interval of α -Ni(OH)₂ formation, further complicating interpretation of the CVs. Such a release of **ab**sorbed hydrogen at positive potentials has been evidenced by electrochemical quartz crystal microbalance measurements⁶⁹ and *in situ* measurement of surface stress of a thin Ni film.⁹⁷ It has also been noticed that oxidation of the Ni surface facilitates hydrogen **ab**sorption and related formation of Ni hydrides ^{41,64,75,77,78} (in essence Ni-oxides are more disordered and thus less dense than Ni-metal, and therefore facilitate hydrogen insertion). However, in most cases, the contribution of **ab**sorbed hydrogen to the anodic currents at potential above E = 0 V vs RHE is negligible, especially when the cathodic limit of CVs lies above E = -0.2 V vs RHE.

Hydrogen absorption in Ni lattice has also been studied in surface science and heterogeneous catalysis. For example, surface science studies of Comsa et al. provided a convincing evidence for the absorption and bulk diffusion of hydrogen (deuterium) into the Ni lattice. ⁹⁸ More recently, Ceyer's group performed high precision surface science experiments in order to imbed hydrogen atoms in the Ni lattice, detect their formation, and study their reactivity.⁹⁹ An excellent overview of the literature, published before 2001 and related to the hydrogen absorption in metals was performed by Ceyer.¹⁰⁰ In addition, Greeley and Mavrikakis performed first-principles study and calculated the phase diagram of the hydrogen/Ni system, which suggest the formation of one monolayer of subsurface hydrogen at high H₂ gas pressure.⁹⁰ Considering this phase diagram and by using the Nernst equation, it is possible to estimate an equilibrium potential for the conversion of hydrogen from the adsorbed into a sub-surface state. At 300 K this potential lies around E = -0.15 V vs RHE. This estimation is in agreement with the electrochemical studies evidencing hydrogen absorption in the Ni lattice at high cathodic polarization.

α -/ β -Ni(OH)₂ and Ni(OH)₂/NiOOH transition

In *the high-potential region*, increase of the potential from E = 0.5 to 1.2 V vs RHE leads to dehydration of α-Ni(OH)₂ and its irreversible transformation into β-Ni(OH)₂, along with the formation and/or thickening of the NiO.^{23,43,44,46,47,57,59,101} Formation of a stable β-Ni(OH)₂ layer leads to an attenuation of the cathodic peak in *the low-potential region* (Figure 1a), firstly followed by its shift towards more negative values, ^{45,61,102} and finally, once β-Ni(OH)₂ fully covers the electrode surface (red curve on Figure 1a), complete disappearance of the peak in *the low-potential region*.^{43,77,79} Above E = 1.2 V vs RHE, α/β-Ni(OH)₂ undergoes further transformation into γ/β-NiOOH, resulting in an emergence of the anodic peak around E = 1.4 V vs RHE (Figure 1a). The corresponding cathodic peak appears around E = 1.3 V vs RHE and at low sweep rates is often split into two components supposedly due to the presence of two distinct (β and γ) phases of the surface NiOOH oxyhydroxide ¹⁰³⁻¹⁰⁵. At potentials above E = 1.5 V vs RHE, the surface of the Ni electrode is covered by NiOOH and NiO as evidenced by *in situ* Raman spectroscopy ^{101,106-109} and XPS measurements (both *ex situ* and *in situ*) ^{71,110,111}. The crystal plane-orientation dependence of Ni oxidation in *the high-potential region* has not been observed, likely because of the fast structure disordering at high potentials.⁶⁵

2.2. Determination of the active surface area of Ni

Correct determination of the electrochemical surface area (ECSA) of an electrode is of paramount importance for evaluation of its intrinsic electrocatalytic activity and comparison between various materials. However, facile oxidation of Ni and possible presence of various oxides (reversible or

irreversible, vide supra) on its surface complicate its ECSA determination. Thus, there is still no generally accepted way to determine the ECSA, the latter depending on several factors discussed below. For electrocatalytic reactions occurring in *the low-potential region* (for example, the HOR), the adsorption of reagents typically takes place at metallic Ni sites, whose number is thus essential for determination of the specific electrocatalytic activity. As a consequence, the surface fraction covered by stable oxide species (such as β -Ni(OH)₂ or NiO) may be discarded for the determination of the ECSA value. In contrast, for reactions occurring in *the high-potential region* (for example oxidation of various organic molecules as well as the oxygen evolution reaction), the active sites generally comprise oxidized Ni. In this case, utilization of the Ni/α-Ni(OH)₂ potential region for the ECSA determination (as it is frequently done in the literature) may result in its significant underestimation, if the electrode surface has been (at least partially) covered by stable Ni oxides. This is evident from the comparison of CVs of a Ni disk (Figure. 1a): after partial oxidation of the Ni electrode, the charge under the anodic peak of α -Ni(OH)₂ formation as well as the double layer capacitance between E = 0.5 and 1.2 V vs RHE significantly decrease, while the charge under the peaks corresponding to the Ni(OH)₂/NiOOH transition changes only slightly, due to a small modification of the Ni surface roughness caused by cycling in a wide potential range ^{43,112}.

At present, one of the most suitable approaches developed to estimate the ECSA of metallic Ni is based on the use of the full anodic charge under the peak of α -Ni(OH)₂ formation^{45,75} and the corresponding specific charge of 514 µC cm⁻² experimentally determined with the aid of low index single crystal Ni(hkl) electrodes⁵⁴. This value assumes transfer of 2 e⁻ per Ni surface site and hence corresponds to the formation of a monolayer of α -Ni(OH)₂. The cathodic charge of α -Ni(OH)₂ reduction may also be considered to determine the ECSA^{113–115}, but this requires correct subtraction of currents corresponding to the HER (which is not an easy task). In order to minimize the influence of side processes on the estimation of ECSA, it is recommended to record CVs with a cathodic potential limit between *E* = -0.06 and -0.12 V vs RHE (to minimize the formation of Ni hydrides) and anodic potential limit between E = 0.40 and 0.50 V vs RHE (to avoid formation of irreducible β -Ni(OH)₂), at sweep rates between v = 10 and 50 mV s⁻¹. Note however that the described procedure has some inherent uncertainties. Firstly, the α -Ni(OH)₂ layer may grow beyond a monolayer thickness, and its growth depends on the experimental conditions (sweep rate, upper potential limit, temperature, etc.), which affects the accuracy of the surface area determination ⁴⁵. Furthermore, assuming that formation of surface α -Ni(OH)₂ occurs via replacement of H_{upd} (see Section **2.1**), the charge under the anodic peak in the *low-potential region* must also involve hydrogen desorption, further questioning the accuracy of the described procedure of the Ni surface area determination.

Another way to estimate the ECSA of metallic Ni is based on the measurement of the electrochemical double layer capacitance (C_{DL}), either by electrochemical impedance spectroscopy (EIS) or by CV in a narrow potential interval (< 100 mV) slightly positive of E = 0 V vs RHE. Specific capacitance values of metallic Ni electrodes range from 20 to 25 µF cm⁻².^{41,80,114,116–118} Once the surface of a Ni electrode is (partially) oxidized, the C_{DL} value changes. Thus, Grdeń et al. estimated the C_{DL} value of a fully-oxidized Ni surface to be around 8 µF cm⁻².¹¹⁴ in the potential interval between E = -0.2 and -0.25 V vs RHE. However much higher values of 40 to 60 µF cm⁻² have been reported at positive potentials (0.8 < E < 1.2 V vs RHE).^{15,119–121} Such significant uncertainties in the specific capacitance values raise a question regarding the applicability of the estimated C_{DL} values to measure the ECSA.^{15,122}

In order to determine the total surface area of a Ni electrode (*i.e.* including contribution both from metallic and from oxidized Ni sites), Grdeń et al. have suggested to complement CV (or CV plus EIS) measurements with XPS experiments.¹¹⁴ In their approach the XPS data are used to determine the ratio between the surface areas of oxidized and metallic Ni ($r = A_{ox}/A_m$), which is then applied to calculate the entire surface area of Ni. Note however that *ex situ* XPS data might not necessarily reflect the surface state of a Ni electrode established *in situ* after conditioning in a specific potential region. Besides, to the

best of the authors' knowledge, XPS does not allow to differentiate between α -Ni(OH)₂ and β -Ni(OH)₂, the former species being reducible in the *low-potential region*, while the latter – not.

When Ni electrocatalysts are used for reactions occurring in the high-potential region (organics oxidation or oxygen evolution reaction), the active sites consist of Ni(II)/Ni(III) species. In this case, the most straightforward way to estimate the ECSA of a Ni electrode could be the determination of the charge associated with the Ni(OH)₂/NiOOH transition. The difficulty, however, lies in the NiOOH layer growth beyond a monolayer, its thickness depending on the growth conditions.⁴⁷ To overcome this complication, Hall et al.,¹²³ suggested to use oxalate salt (acting as a stabilizing agent due to the formation of an oxalate ad-layer) and sufficiently high sweep rate $v \ge 150$ mV s⁻¹ to limit the thickness of surface Ni(OH)₂ and NiOOH to a single layer. This approach was later adopted to measure the ECSA of a Ni foam.¹¹² In this case sharp and narrow peaks for the Ni(OH)2/NiOOH oxidation-reduction processes were observed in the CVs, resulting in higher precision of the determined ECSA value of Ni assuming a specific charge of 195 μ C cm⁻². However, the latter method can only be applied either after measuring the catalytic activity of an electrode, or in a separate experiment, since the presence of the oxalate salt in the electrolyte will most likely affect the electrocatalytic properties of Ni. Alternatively, Ho and Piron⁹⁵ and later Lyons and Brandon⁹⁴ suggested to galvanostatically charge a Ni electrode in the potential interval of the oxygen evolution reaction (occurring on the NiOOH-covered surface) and then calculate charge passing during a cathodic current-time transient. This method assumes a specific charge of 420 µC cm⁻², although, as mentioned by the authors, the exact charge passing during the desorption of a full monolayer of the oxygen evolution reaction intermediates is uncertain, and may vary, in particular, depending on the type of the NiOOH surface phase (β or γ) at stake, which present somewhat different unit cell parameters and symmetries ^{44,124} (not to mention that the nature of active oxygen evolution intermediates is still debated). Recently, Watzele et al. ¹²⁵ proposed a yet another strategy to determine the ECSA of metal oxide catalysts

(including Ni) by measuring the so-called adsorption pseudocapacitance at the onset of the oxygen evolution reaction.

Finally, chemical methods to measure the surface area, while not giving the exact information about the ECSA, are still useful as they allow to calculate the full surface area of Ni. Among various techniques described by Trasatti and Petrii,¹²⁶ CO and H₂ chemisorption measurements for metallic Ni, or N₂ physisorption measurements for Ni oxides can be mentioned.^{122,127–129}

3. Electrocatalytic properties of Ni-based electrocatalysts in the HOR in alkaline media

3.1. Introduction: electrocatalysis of hydrogen electrode reactions on Ni

The HOR/HER are among the most widely studied electrode processes. The mechanism of the hydrogen electrode reactions on metal electrodes ("M" in the Eqs. below stands for a metal surface site) includes Volmer, Heyrovsky and Tafel steps, which in alkaline media can be written as follows ^{26,130}:

$$H_2 + 2 M \rightleftharpoons 2 M - H_{ad}$$
(3.1, Tafel step)

$$H_2 + OH^2 + M \rightleftharpoons M - H_{ad} + H_2O + e^2$$
 (3.2, Heyrovsky step)

$$M-H_{ad} + OH^{-} \rightleftharpoons H_{2}O + e^{-} + M$$
 (3.3, Volmer step)

Since atomically adsorbed hydrogen (H_{ad}) acts as a reaction intermediate, the electrocatalytic activity in hydrogen electrode reactions is closely related to its strength of adsorption. ^{131–134} There has been convincing evidence that on Pt and other PGM electrodes, H_{upd} , which is adsorbed in multi-fold hollow sites, cannot act as the HOR/HER intermediate due to its strong binding to the surface, and that it is rather weakly adsorbed hydrogen (H_{opd}), likely adsorbed on-top, which participates in hydrogen electrocatalysis.^{81–83} For the most catalytically-active metals like Pt and Pd, the Gibbs energy of adsorption of this weakly-adsorbed hydrogen is close to zero.^{131,135,136} On Ni, hydrogen in multi-fold hollow sites is adsorbed stronger than either on Pt or Pd. ^{131,135,136} Santos et al., using DFT, compared strongly- and weakly-adsorbed hydrogen on Ni, Pt and some other surfaces.¹³³ According to these calculations, and in agreement with other studies, strongly-adsorbed hydrogen binds to Ni(111) stronger than to Pt(111). In the meantime, for weakly-adsorbed hydrogen on Ni(111), adsorbed on top of a monolayer of the strongly-adsorbed hydrogen, $\Delta G_{\rm H} \approx 0.9$ eV, which makes its participation in the HOR rather unlikely. In what follows, when discussing the HOR on Ni, it will be assumed that it is strongly-adsorbed hydrogen, which acts as a reaction intermediate on Ni, and it will simply be denoted as H_{ad}. While this conjecture requires additional confirmation, it agrees with an increase of the HOR on Ni-based electrodes with the decrease of the strength of hydrogen adsorption, the fact widely acknowledged in the literature.^{48,50,137–142}

Besides, experimental data (collected for PGM-based electrodes) evidence much slower hydrogen electrode kinetics in alkaline media than in acid.^{143–145} To explain this experimental observation, the so called 'bifunctional' mechanism has been proposed,^{18,146} which postulates that in order to be active in the HOR/HER in alkaline media, an electrode must provide two types of sites: one for H_{ad} adsorption and another one for water dissociative adsorption to form H_{ad} and OH_{ad}. We will come back to the feasibility of this 'bifunctional' mechanism on Ni.

Owing to the use of alkaline water electrolysis at the industrial level, studies of Ni-based electrocatalysts have been primarily carried out in the potential interval of the HER, while, until recently, the interest in the HOR on Ni has been rather limited. HER starts with the Volmer step, which provides hydroxyl ion and an adsorbed H_{ad} . Then, hydrogen molecule is produced via either recombination of two H_{ad} atoms (Tafel step) or interaction of H_{ad} and a water molecule (Heyrovsky step). The contribution of individual steps (3.1 – 3.3) to the overall reaction mechanism depends on many factors such as the Ni surface state, the presence of a second element (metal or nonmetal) as well as the applied overpotential.

In particular, the HER on polycrystalline Ni follows Volmer-Heyrovsky mechanism if the electrode surface is dominated by metallic Ni sites 30,56,75,118 , while the Volmer-Tafel mechanism was proposed in some publications where Ni electrodes were used likely in a partially-oxidized state of the surface. 85,147,148 The latest progress in the field of the HER on Ni and Ni-based electrocatalysts, including the reaction mechanism, is summarized in several reviews 17,18,26,27 and is beyond the scope of this paper. In this review we are primarily interested in the HOR and will mention the HER only when relevant (*e.g.* when it enters in competition with reactions of interest for this work). The HOR comprises the same elementary steps as the HER but in the reverse direction, the H₂ molecule first dissociating with the formation of surface H_{ad} (through the Heyrovsky and/or Tafel step), the latter undergoing electrochemical oxidation with the formation of H₂O (Volmer step). It should be noted however that the nature of the rate-determining step depends on the electrode potential, thus resulting in differences in the detailed reaction mechanisms for the HOR and the HER if considered at high anodic or cathodic potentials, correspondingly.

3.2. HOR on monometallic Ni electrodes

Literature data regarding the kinetics of the HOR on Ni (and HER at low cathodic overpotentials) are very contradictory,^{41,42,64,85,118,147,149–151} which may be attributed to the strong influence of the electrode pre-treatment and thus the state of the surface on the reaction kinetics.^{48–50,146,152} In the vicinity of the equilibrium potential, a purposely-reduced metallic Ni electrode (see black solid trace in Figure 2b) shows a very small HOR current, which then increases peaking at ca. E = 0.25 V vs RHE (*i.e.* close to the anodic peak position observed under N₂ atmosphere), and then decays again. Oshchepkov et al. attributed this low activity of metallic Ni to the slow kinetics of the Volmer step likely due to the strong adsorption of H_{ad} species already mentioned above.^{48,49,131,132} Indeed, according to the results of kinetic modeling, surface coverage of metallic Ni with H_{ad} (θ_H) is high under N₂ atmosphere, and further increases in the presence of H₂, adsorbed hydrogen atoms dominating on the surface until ca. E = 0.2 V vs RHE (Figures 2a and 2b, for details a reader is referred to Ref. ⁴⁹). Note that the obtained data are in a good agreement with hydrogen adsorption isotherm (from the gas phase) reported by Pshenichnikov.⁷⁴ Besides, according to Taylor et al. at pH = 13 the equilibrium potential for H_{ad}/H^+ is similar to 0.2 V, which is in agreement with the data of Figure 2a.⁸⁹ At higher potential values, formation of α -Ni(OH)₂, which proceeds through the OH adsorption (see θ_{OH} and $\theta_{(OH)2}$ traces in Figures 2), results in blocking of the Ni active sites.^{45,48,66,92} Thus, considering the kinetics of the HOR on Ni electrocatalysts also requires accounting for the processes of the Ni surface hydroxide formation and reduction, since they occur in the same potential range as the HOR. Besides, the coexistence of both Ni-H_{ad} and Ni-OH_{ad} species on the electrode surface might result in their recombination with the formation of water through the following reaction ('bifunctional' mechanism)¹⁴⁶:

$$Ni-H_{ad} + Ni-OH_{ad} \rightleftharpoons H_2O + 2 Ni$$
 (3.4)

However, contribution of the latter step to the mechanism of the HOR has been shown to be unlikely on a pure metallic Ni surface because of the very strong adsorption of H_{ad} on Ni and thus the high activation barrier for the recombination reaction.⁵¹ Nevertheless, the recombination step might contribute to the HOR on bimetallic NiM electrocatalysts and/or partially oxidized Ni electrodes, when the adsorption energies of H_{ad} and OH_{ad} are weakened.^{50,51}



Figure 2. Experimental anodic scans of CVs (black solid curves) for (a, b) metallic and (c, d) partially oxidized polycrystalline Ni rod in (a, c) N₂- and (b, d) H₂-saturated 0.1 mol L⁻¹ NaOH at $v = 5 \text{ mV s}^{-1}$ and T = 25 °C. Dashed curves represent the simulated data of either CVs or coverage of the electrode surface by various adsorbates. Note that only Ni metal sites free from stable oxide species (NiO_x surface coverage is estimated to be ca. 14 %) were considered for calculations. Data are replotted from Ref. ⁴⁹ with permission from Elsevier.

The catalytic activity of metallic ('reduced') Ni electrodes being rather low, several approaches are currently proposed to improve the performance of Ni and Ni-based electrocatalysts in the HOR. They target either (i) decreasing the Ni-H_{ad} binding energy and hence increasing the Volmer step kinetics, or

(ii) increasing the kinetics of the recombination step, or (iii) making Ni less sensitive to the oxide blocking thus extending the operating potential window of an AEMFC with a Ni anode. In section 3.3 of the review, we summarize recent advances in the development of nanoparticulated Ni and Ni-based electrocatalysts for the HOR and outline three main strategies, which have been put forward to achieve high catalytic activity of Ni.

3.3. Strategies for improving the activity of Ni electrocatalysts in the HOR *Effect of the surface state of Ni and heteroatom doping*

As highlighted in section 2 of this review, Ni is prone to oxidation already at ambient conditions resulting in the formation of surface Ni oxides, some of them (addressed as 'stable' surface oxides) surviving in the potential intervals of the HOR/HER and coexisting with metallic Ni sites at the electrode surface. It has been shown that the presence of surface NiO_x species (presumably serving as water adsorption sites) along with metallic Ni sites (serving as H_{ad} adsorption sites) results in a synergistic effect leading to significant enhancement of the HOR currents on partially-oxidized Ni electrodes.^{48–50} The HOR activity depends on the NiO_x surface coverage, with an optimum observed at ca. 30 to 50 % of NiO_x depending on the type of Ni electrodes (*e.g.* polycrystalline Ni ⁵⁰ or Ni/C nanoparticles ¹⁵³). On the basis of DFT calculations as well as microkinetic modeling, it has been concluded that NiO_x species weaken the strength of adsorption of H_{ad} at nearby-located Ni sites and likely increase the strength of H₂O adsorption at the surface, both leading to an enhancement of the kinetics of the Volmer step (Figure 2).^{50,91} As a result, the HOR starts at lower overpotential, as evidenced by an emergence of the second anodic peak (labeled as a_2 in Figure 3a) in CVs in H₂ atmosphere. This peak shifts towards E = 0 V *vs* RHE with the NiO_x coverage and an ensuing decrease of Ni-H_{ad} binding energy (Figure 3). It is instructive to

compare $\theta_{\rm H}$ for a metallic and partially oxidized Ni surface (Figure 2). Indeed, $\theta_{\rm H}$ significantly decreases in the latter case (NiO_x surface coverage is estimated to be ca. 14 % based on the data presented in Ref.⁴⁹), which goes along with the increase of the HOR current. While further (preferably *in situ* spectroscopic) studies are required to unveil the surface state of Ni electrodes during the HOR and thus unambiguously confirm its influence on the reaction kinetics, it should be emphasized that the model proposed by Oshchepkov et al. currently presents the most consistent explanation of the available literature data.^{48–52}

Armed with this knowledge, Oshchepkov et al. have recently reported a partially oxidized Ni-NiO_x/XC-72 electrocatalyst prepared by electrodeposition technique, which possesses HOR specific activity at the level of a commercial Pd/C electrocatalyst (Table 1).¹³⁷ It has been demonstrated that the existence of surface NiO_x species (their estimated coverage varies between 48 and 58%) enhances the surface- and mass-normalized HOR exchange current densities from 6.2 to 56 μ A cm_{Ni}⁻² and from 5.7 to 22.4 A g_{Ni}⁻¹, respectively, compared to a reduced Ni/C electrocatalyst. The HOR kinetic current density of Ni-NiO_x/XC-72 normalized by mass of Ni was estimated as 32.1 A g_{Ni}^{-1} at E = 50 mV vs RHE, which makes it the current state-of-the-art in the field of PGM-free electrocatalysts for the HOR. An improved HOR specific activity was also reported for Ni/NiO/C composites derived from the Ni-based metalorganic frameworks and ascribed by the authors to the optimal binding energies of both H_{ad} and OH_{ad} at the Ni/NiO interface as well as to the high level of carbon graphitization, the latter resulting in high conductivity.¹⁴¹ It has also been demonstrated, that the Ni/NiO/C composite possesses much better CO tolerance than a 20 wt.% Pt/C catalyst, the HOR activity of the latter in the presence of CO decaying below that of Ni/NiO/C within a couple of hours. Since most of hydrogen is currently produced through natural gas reforming, the presence of CO in the fuel stream cannot be fully avoided, not to mention that the carbon support itself can lead to CO-like species adsorption at the metal nanoparticles (see Section 5), hence to their "poisoning" at low potentials. Therefore, synthesis of CO-tolerant electrocatalysts is an important task in the development of AEMFCs.



Figure 3. Comparison of the anodic scans of (a) experimental and (b) simulated CVs for Ni/NiO_x electrodes with different degree of oxidation of their surfaces (characterized by the corresponding NiO_x coverage) in H₂-saturated 0.10 mol L⁻¹ NaOH at $v = 5 \text{ mV s}^{-1}$. (c) Schematic illustration of the Ni active sites with various adsorption energies depending on the number of closely located oxide species. Influence of the NiO_x coverage on (d) the percentage of Ni active sites affected by oxygen species (peak a₂), (e) the Gibbs energy of the adsorbed H_{ad} intermediate, (f) the Volmer rate constant k_V^0 , and (g) the potential of Ni-OH_{ad} formation. Reproduced from Ref. ⁵⁰ with permission from Springer.

Besides modification of the Ni surface state, several authors reported successful examples of introducing heteroatoms in the structure of Ni nanoparticles. It has been demonstrated that formation of a Ni₃N phase provides significant enhancement of the activity in the HOR with respect to a pure Ni

electrocatalyst (Table 1).^{139,140,142} DFT calculations suggest that in the presence of an interstitial N dopant, the absolute value of $\Delta G_{\rm H}$ decreases, hence approaching zero (*i.e.* an optimal value for the HOR).^{139,142} In addition, it was proposed that the activation energy for both water formation and water dissociation decrease at the interface between Ni₃N and Ni ^{139,142}. As a result, the reported carbon-supported Ni₃N electrocatalysts provide very high mass-normalized HOR kinetic current densities at E = 50 mV vs RHE (24.38 and 29.75 A g_{Ni} ⁻¹, respectively from the Refs. ¹⁴⁰ and ¹³⁹), thus being close to the state-of-the-art Ni-NiO_x/XC-72 electrocatalyst (32.1 A g_{Ni}⁻¹).¹³⁷ Notably, the Ni₃N nanoparticles supported on either carbon or on Ni foam demonstrated very high resistance towards CO poisoning (Figure 4) ^{139,142}. Moreover, the Ni₃N/C shows a limiting HOR current densities up to E = 0.26 V vs RHE, which is ca. 0.1 V higher the values typically observed with Ni-based electrocatalysts. ^{137–142,154} On the basis of ultraviolet photoemission spectroscopic measurements, this was attributed to a weaker oxygen adsorption on Ni₃N as compared to pure Ni.¹⁴⁰ A decrease of the propensity of Ni towards oxidation was also reported by Gao et al. for a Ni core-hexagonal boron nitride (h-BN) shell (Ni/h-BN/KB electrocatalyst).¹²⁹ This was achieved via formation of a few layers of a defective hexagonal BN shell, which protected the Ni core from an excessive oxidation in air, yet allowing H, OH, and H₂O to interact with the Ni surface. In addition, it was suggested that the h-BN shell weakened H_{ad} adsorption on the Ni surface, resulting in a six times higher HOR exchange current density measured for Ni/h-BN/KB as compared to pure Ni/KB sample (Table 1).



Figure 4. Electrocatalytic performance of Ni₃N/Ni/NF, Ni/NF, and Pt/NF (NF is a Ni foam) catalysts in the HOR. (a) Steady-state polarization curves of Ni₃N/Ni/NF, Ni/NF, and Pt/NF in H₂-saturated 0.1 mol L⁻¹ KOH. (b) Chronoamperometry curves of Ni₃N/Ni/NF in H₂- or Ar-saturated 0.1 mol L⁻¹ KOH measured at E = 0.09 V vs. RHE. (c) Comparison of the HOR polarization curves of Ni₃N/Ni/NF and Pt/NF in 0.1 mol L⁻¹ KOH saturated with H₂ or a mixture of H₂ and 2% CO (v/v). (d) Chronoamperometry curves of Ni₃N/Ni/NF and Pt/NF in 0.1 mol L⁻¹ KOH saturated with a mixture of H₂ and 2% CO (v/v) measured at E = 0.09 V vs. RHE. All results were collected without iR compensation. The current densities were calculated on the basis of the geometric electrode areas. Reproduced from Ref. ¹⁴² with permission from Springer Nature

Bimetallic NiM electrocatalysts

Another (and the most popular) approach to modify the catalytic properties of materials and, in particular, Ni nanoparticles is based on alloying them with other metals. In order to take full advantage of the AEMFC concept, noble metal additives to Ni electrocatalysts should be avoided. Besides, alloying Ni with metals like Pt, Pd and Ir does not allow to decipher the influence of the noble metal (being more active than Ni) on its activity in the HOR. Instead, Ni serves as an additive helping to further improve the activity of noble metals in the HOR, presumably by providing surface sites for adsorbing hydroxyl species ^{155–157} or by tuning the hydrogen binding energy.¹³⁴ Considering the above-mentioned reasons, in what follows, we will only discuss NiM systems with a non-precious second component (Table 1).

Oshchepkov et al. have studied NiCu/XC-72 electrocatalysts with a Cu fraction from 0.05 to 0.50 prepared by the incipient wetness impregnation. In this series, samples containing 5 wt.% of Cu showed the highest specific and mass activities in the HOR, which was at least 3 times higher compared to a pure Ni/C electrocatalyst prepared by the same method.¹²⁷ By using freeze-drying, the same group prepared more homogeneous alloyed NiCu/XC-72 electrocatalysts with relatively narrow particle size distribution, in which the highest activity was again observed for the Ni_{0.95}Cu_{0.05}/XC-72 sample.¹⁵⁸ The specific activity of Ni_{0.95}Cu_{0.05}/XC-72 was found to be the same in both series (ca. 14 μ A cm_{Ni}⁻²), while the mass activity of the sample prepared using freeze-drying increased by a factor of 1.5 (up to 2.5 A gm⁻¹) due to a smaller average particle size. Similar electrocatalyst composition was also explored by Roy et al. ¹⁵⁹, who reported two times higher exchange current density (25 μ A cm_{Ni}⁻²) for their Ni₉₅Cu₅/Ketjenblack electrocatalyst at however 2.5 times lower mass-normalized activity (Table 1). The latter is likely arising from the low ECSA of NiCu nanoparticles in Ref. ¹⁵⁹ The authors also evaluated the performance of an AEMFC

consisting of a Ni₉₅Cu₅/Ketjenblack anode and a Pd/XC-72R cathode separated by a Tokuyama A201 membrane, possessing the power density of ca. 160 and 350 mW cm⁻² at 70 and 80°C, respectively, and 100 % relative humidity.¹⁵⁹ In addition, Wang et al. have studied Ni-Cu alloy films with different composition prepared using a combinatorial magnetron co-sputtering method.¹⁶⁰ The authors have found that the highest (4 times higher compared to a pure Ni sample) exchange current density in the HOR (34 μ A cm_{Ni}⁻²) is reached for NiCu alloys with ca. 40 % Cu content.¹⁶⁰ The large difference in the optimum composition of NiCu nanoparticles and NiCu alloy films towards the HOR may be related to the particle-size dependent segregation of Cu atoms to the surface.^{127,161} Indeed, Monte Carlo simulations revealed massive segregation of Cu to the surface of NiCu nanoparticles.⁵¹ For example, for ca. 18 nm diameter NiCu nanoparticles with the 0.04 Cu fraction in the bulk, Cu surface fraction was as high as 0.5.

Another additive used to tailor Ni activity in the HOR is molybdenum. Roy et al. reported a 50% Ni₉Mo₁/Ketjenblack electrocatalyst, which demonstrated high activity in the HOR in terms of both surface- (27 μ A cm_{Ni}⁻²) and mass-normalized (4.5 A g_M⁻¹) exchange current densities.¹⁶² The authors assigned the high activity to the decrease of the binding energy of the hydrogen intermediate to the NiMo surface as compared to that of metallic Ni. In addition, they reported power density of 120 mW cm⁻² for an AEMFC with the Ni₉Mo₁/Ketjenblack anode and 30 wt.% Pd/XC-72R cathode separated by Tokuyama A201 membrane at 70 °C and 70 % relative humidity. Note however that *post-mortem* XPS analysis revealed almost total disappearance of Mo (initially dominating the electrocatalyst surface) from the surface of a membrane electrode assembly after the AEMFC test. The fact that Mo leaching did not result in a noticeable decrease in the fuel cell performance raises doubts regarding the role of Mo in the electrocatalyst and the actual nature of active sites. Leaching Mo from NiMo samples has been also observed by Schalenbach et al. ¹⁶³, who studied NiMo alloys with different composition prepared by arc melting (more emphasis on the durability of Ni-based catalysts will be given in Section 5). Sheng et al.

reported a ternary $Co_{0.17}Ni_{4.49}Mo_1$ alloy prepared by electrodeposition with the HOR exchange current density of 15 μ A cm_{Ni}⁻².¹⁶⁴ The much higher activity of the latter compared to pure Ni electrocatalyst has been attributed to a favorable Ni-H_{ad} binding energy, while the effect of OH_{ad} species on the HOR was considered as insignificant. It has also been proposed that MoO₂ species (if surviving at the surface of a Ni/MoO₂ electrocatalyst) should accelerate the Volmer step of the HOR by promoting the OH_{ad} formation, while the nearby Ni sites serve for the hydrogen adsorption.¹⁶⁵

Tang et al. reported Ni-Ag alloys with various compositions prepared by electron-beam vapor codeposition.¹⁶⁶ DFT calculations predict a decrease of the H_{ad} binding energy on Ni₃Ag hollow sites as compared to pure Ni, which should result in an enhanced kinetics of the hydrogen electrode reactions. The latter was indeed observed by the authors for the Ni_{0.75}Ag_{0.25} sample in the HER, while the HOR current densities of NiAg alloys were found to be either similar (for Ni-rich samples) or inferior (for Agrich samples) to the pure Ni sample. Finally, by using CeO₂(r)-Ni/XC-72 electrocatalyst prepared through decoration of Ni particles with partially reduced CeO₂ during hydrothermal synthesis, Yang et al. investigated the influence of the oxygen vacancies in CeO₂ on the kinetics of the HOR on Ni.¹¹⁵ The CeO₂(r)-Ni/XC-72 electrocatalyst demonstrated an improved activity in the HOR with the values of exchange current density (38 μ A cm_{Ni}⁻²) and mass-normalized kinetic HOR current at E = 50 mV vs RHE (12.8 A g_{Ni}^{-1}) being at least 2 times higher than the ones obtained for the reference Ni/C sample (Table 1). On the basis of DFT calculations the authors attributed the enhanced HOR performance of the $CeO_2(r)$ -Ni/XC-72 electrocatalyst to a bifunctional mechanism at the interface between Ni and CeO₂ (Figure 5). It was suggested that CeO₂ moieties on the surface serve as active sites for OH_{ad} adsorption (due to the presence of oxygen vacancies), while the nearby Ni sites are characterized by a weaker adsorption of H_{ad} as compared to pure Ni. Both of these factors should facilitate the HOR kinetics.¹¹⁵



Figure 5. Schematic representation of the $CeO_2(r)$ -Ni hydrogen oxidation catalyst (the symbol 'r' points on a partial reduction of CeO_2 during the catalyst preparation). (a) Representation of electron transfer between $CeO_2(r)$ and Ni in the layered $CeO_2(r)$ -Ni hybrid. The calculated (b) ΔG_H and (c) ΔG_{OH} adsorption energies on Ni and $CeO_2(r)$ -Ni hybrid. (d) Schematic illustration of role of the oxygen-vacancy-containing CeO_2 during the HOR. Reproduced from Ref. ¹¹⁵ with permission from Willey.

Effect of support

Modification of support rather than Ni nanoparticles opens up a new route for preparing Ni-based electrocatalysts with an improved activity in the HOR. Zhuang et al. have shown that Ni nanoparticles supported on N-doped carbon nanotubes (N-CNT) possess high surface- (28 μ A cm_{Ni}⁻²) and mass-normalized (9 A g_{Ni}⁻¹) exchange currents, which are at least 20 and 3 times higher than the values obtained for unsupported and CNT-supported Ni nanoparticles, respectively.¹¹³ Using DFT calculations, the authors suggested that the N-doped carbon support stabilizes Ni nanoparticles against reconstruction and

affects d-orbitals of Ni leading to weakening of the Ni-H_{ad} binding energy. Similar enhancement of the HOR activity of Ni through metal-support interaction was observed by Yang et al., who systematically evaluated performance of Ni nanoparticles supported on N-, B- and S-doped XC-72 carbon.¹⁶⁷ The activity trend was found to follow the sequence of Ni/SC > Ni/NC > Ni/BC > Ni/C (Table 1) and was ascribed by the authors to two main effects caused by heteroatom-doping: (i) formation of Ni nanoparticles with a smaller and more uniform particle size; (ii) metal-support interaction which weakens Ni-H_{ad} bonding resulting in an enhanced specific activity in the HOR.

Table 1. Comparison of the HOR activity of PGM-free Ni-based electrocatalysts in alkaline media reported between 2014 and 2019. Notations: d_{NiM} (TEM) is the mean size of Ni(M) nanoparticles determined by TEM, T is the temperature (RT means room temperature), *ECSA* is the electrochemically active surface area normalized by the mass of Ni, j_0 and i_0 are the surface- and mass-normalized exchange currents, $j_k^{50\text{mV}}$ and $i_k^{50\text{mV}}$ are the surface- and mass-normalized HOR kinetic currents at E = 50 mV vs RHE

	Electro- catalyst	Loading, mg _{NiM} cm _{geom} ⁻²	d _{NiM} (TEM), nm	Electrolyte	<i>Т</i> , °С	$\begin{array}{c} \text{ECSA,} \\ \text{m}^2 \ \text{g}_{\text{Ni}}^{-1} \end{array}$	<i>j</i> ₀ , μA cm _{Ni} -2	$\dot{i}_0,$ A \mathbf{g}_{M} -1	$i_k^{50mV}, \mathbf{A} \cdot \mathbf{g}_{Ni}^{-1}$	j_k^{50mV} , $\mu \mathbf{A} \cdot \mathbf{cm}_{Ni}^{-2}$	Ref.
and	42% Ni/XC-72	0.07 ± 0.01	10	0.1 mol L ⁻¹ NaOH	25	92.4 ± 6.3	6.2 ± 0.6	5.7 ± 0.2	-	-	137
e state toms	42% Ni- NiO _x /XC-72	0.07 ± 0.01	10	0.1 mol L ⁻¹ NaOH	25	46.7 ± 8.6	56 ± 10	22.4 ± 4.3	32.1 ± 4.8	83 ± 12	137
surface neteroa	63.2% Ni/NiO/C	0.32	0-60	0.1 mol L ⁻¹ KOH	-	-	26	-	5	-	141
of the g by h	Ni ₃ N	0.28	-	0.1 mol L ⁻¹ KOH	-	-	1.73	-	-	-	140
ience (dopir	51.4% Ni ₃ N/XC-72R	0.16	4.6 ± 1	0.1 mol L ⁻¹ KOH	-	-	14	12	24.38	-	
Influ	71.7% np- Ni ₃ N/C	0.16	5-20	0.1 mol L ⁻¹ KOH	RT	-	-	10.3	29.75	-	139
of a t	Ni NPs	0.25	-	0.1 mol L ⁻¹ KOH	RT	-	1.3	0.15	0.28	2.4	
uence uppor	70% Ni/CNT	0.25	-	0.1 mol L ⁻¹ KOH	RT	10.4	9	1	1.9	18	113
Infl	70% Ni/N-CNT	0.25	20	0.1 mol L ⁻ ¹ KOH	RT	12.4	28	3.5	9.3	75	

	27% Ni/KB	-	10-15	0.1 mol L ⁻¹ NaOH	25	15.34 (H ₂ chemisorp tion)	4	0.61	-	-	129
	32.5% Ni/h- BN/KB	0.25	10-15	0.1 mol L ⁻¹ NaOH	25	15.20 (H ₂ chemisorp tion)	23	3.5	-	-	
	45% Ni/C	0.17	33.1	0.1 mol L ⁻¹ KOH	30	7.1±0.8	13.8 ± 5.0	0.97 ± 0.25	-	-	167
	54% Ni/BC	0.17	17.7	0.1 mol L ⁻¹ KOH	30	15.4±0.5	13.0 ± 4.7	1.99 ± 0.73	-	-	
	55% Ni/NC	0.17	16.5	0.1 mol L ⁻¹ KOH	30	16.1±0.3	29.9 ± 5.1	4.82 ± 0.83	-	-	
	55% Ni/SC	0.14	14.2	0.1 mol L ⁻¹ KOH	30	16.9±0.9	40.2 ± 2.8	7.44 ± 0.88	11	-	
	60% Ni/KB	0.06	12	0.1 mol L ⁻¹ NaOH	25	24.5 ± 0.6	28.6 ± 0.3	7.02 ± 0.1			128
	52.3% Ni/XC- 72	0.16	15.1	0.1 mol L ⁻¹ KOH	30	-	16 ± 2	-	5.89±0.71	-	115
	46.1% CeO ₂ (r)- Ni/XC-72	0.14	11.9	0.1 mol L ⁻¹ KOH	30	-	38 ± 2.2	-	12.8±0.27	-	
	50% Ni ₉ Mo ₁ /KB	0.26	5-50	0.1 mol L ⁻¹ NaOH	25	16.9 ± 1.1	27 ± 2	4.5 ± 0.2	-	-	162
	Co _{0.17} Ni _{4.49} Mo ₁	-	-	0.1 mol L ⁻¹ KOH	20	-	15	-	-	44 ± 5	164
				0.1 mol I -			10.0.01				
tal	25% Ni/XC-72	0.025	36.5	NaOH	25	5.1 ± 0.6	10.8 ± 0.1	0.55 ± 0.06	-	-	
cond metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72	0.025 0.025	36.5 29.8	0.1 mol L NaOH 0.1 mol L ⁻¹ NaOH	25 25	5.1 ± 0.6 20.6	10.8 ± 0.1 14 ± 1	0.55 ± 0.06 2.5 ± 0.2	-	-	158
of a second metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72 50% Ni _{0.95} Cu _{0.05} /KB	0.025 0.025 0.51	36.5 29.8 22±9	0.1 mol L ⁻¹ NaOH 0.1 mol L ⁻¹ NaOH 0.1 mol L ⁻¹ KOH	25 25 RT	5.1 ± 0.6 20.6 3.7 ± 0.2	10.8 ± 0.1 14 ± 1 25 ± 1	0.55 ± 0.06 2.5 ± 0.2 0.9 ± 0.1	-	-	158
ence of a second metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72 50% Ni _{0.95} Cu _{0.05} /KB Ni	0.025 0.025 0.51 -	36.5 29.8 22±9	0.1 mol L NaOH 0.1 mol L ⁻¹ NaOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹	25 25 RT 25	5.1 ± 0.6 20.6 3.7 ± 0.2 1.81 cm ²	10.8 ± 0.1 14 ± 1 25 ± 1 16	0.55 ± 0.06 2.5 ± 0.2 0.9 ± 0.1	-	-	158
Influence of a second metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72 50% Ni _{0.95} Cu _{0.05} /KB Ni Ni Ni ₆ Cu ₄	0.025 0.025 0.51	36.5 29.8 22±9 - -	0.1 mol L NaOH 0.1 mol L ⁻¹ NaOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹ KOH	25 25 RT 25 25	5.1 ± 0.6 20.6 3.7 ± 0.2 1.81 cm^2 3.3 cm^2	10.8 ± 0.1 14 ± 1 25 ± 1 16 34	0.55 ± 0.06 2.5 ± 0.2 0.9 ± 0.1 -	-	-	158 159 160
Influence of a second metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72 50% Ni _{0.95} Cu _{0.05} /KB Ni Ni ₆ Cu ₄ 43% Ni/VXCMAX2 2	0.025 0.025 0.51 - - 0.09	36.5 29.8 22±9 - -	0.1 mol L ⁻¹ NaOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹ KOH	25 25 RT 25 25 25	5.1 ± 0.6 20.6 3.7 \pm 0.2 1.81 cm ² 3.3 cm ² 1.3 \pm 0.2	10.8 ± 0.1 14 ± 1 25 ± 1 16 34 39	0.55 ± 0.06 2.5 ± 0.2 0.9 ± 0.1 - - 0.83	-	- - - - -	158 159 160
Influence of a second metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72 50% Ni _{0.95} Cu _{0.05} /KB Ni Ni ₆ Cu ₄ 43% Ni/VXCMAX2 2 37% Ni ₃ Fe ₁ / VXCMAX22	0.025 0.025 0.51 - - 0.09 0.07	36.5 29.8 22±9 - - - ca. 10	0.1 mol L NaOH 0.1 mol L ⁻¹ NaOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹ KOH 1 mol L ⁻¹ KOH 1 mol L ⁻¹ KOH	25 25 RT 25 25 25 25 25	5.1 ± 0.6 20.6 3.7 ± 0.2 1.81 cm^2 3.3 cm^2 1.3 ± 0.2 1.2 ± 0.6	$ \begin{array}{c} 10.8 \pm 0.1 \\ 14 \pm 1 \\ 25 \pm 1 \\ 16 \\ 34 \\ 39 \\ 60 \\ \end{array} $	0.55 ± 0.06 2.5 ± 0.2 0.9 ± 0.1 - 0.83 0.96			158 159 160 154
Influence of a second metal	25% Ni/XC-72 25% Ni _{0.95} Cu _{0.05} /XC- 72 50% Ni _{0.95} Cu _{0.05} /KB Ni Ni ₆ Cu ₄ 43% Ni/VXCMAX2 2 37% Ni ₃ Fe ₁ / VXCMAX22 45% Ni ₃ Cu ₁ / VXCMAX22	0.025 0.025 0.51 - - 0.09 0.07 0.09	36.5 29.8 22±9 - - - ca. 10 -	0.1 mol L NaOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹ KOH 0.1 mol L ⁻¹ KOH 1 mol L ⁻¹ KOH 1 mol L ⁻¹ KOH 1 mol L ⁻¹ KOH	25 25 RT 25 25 25 25 25 25 25	5.1 ± 0.6 20.6 3.7 ± 0.2 1.81 cm^2 3.3 cm^2 1.3 ± 0.2 1.2 ± 0.6 1.2 ± 0.2	10.8 ± 0.1 14 ± 1 25 ± 1 16 34 39 60 51	0.55 ± 0.06 2.5 ± 0.2 0.9 ± 0.1 - 0.83 0.96 0.98		- - - - - - - -	158 159 160 154

3.4.Perspectives

Data presented in section 3.3 revealed significant progress in the development of PGM-free Ni-based electrocatalysts for the HOR in alkaline media within the last 5 years. The activities of the best materials approach those reported for much more expensive Pd/C electrocatalysts, while being only 10 times lower compared to those of Pt/C electrocatalysts. The most popular strategies used to enhance the performance of Ni in the HOR are based on its alloying with a second metal or incorporation of heteroatoms in either the Ni structure or in the carbon support. In both strategies the electronic effect of the additives on Ni is

proposed as the key factor, which leads to the enhanced HOR activity due to weakening of the Ni-H_{ad} bond. At the same time, the values of the surface- and especially mass-normalized HOR activities of these alloyed and/or heteroatom-doped materials are inferior to those of the Ni-NiO_x/XC-72 electrocatalyst with an optimized surface state of the Ni electrode.¹³⁷ This suggests that addition of a second metal and/or dopant, along with a possible electronic effect, might indirectly affect the energy of hydrogen and oxygen bonding to the surface through stabilization of a particular Ni-NiO_x surface state, which possesses high activity in the HOR. In order to better distinguish between these two possibilities, *in situ* studies for monitoring the Ni surface state along with the measurement of the HOR activity are highly desirable.

4. Oxidation of complex fuels on Ni-based electrocatalysts

Section 3 of this review made it clear that Ni (and its alloys/composites) is a very popular electrocatalyst in hydrogen oxidation and evolution reactions, and that its activity greatly depends on the state of the Ni surface. Considering this, one may expect similar behavior of Ni in other electrocatalytic reactions occurring on the surface of metallic Ni sites and/or involving adsorbed H_{ad} species in the reaction mechanism. In the current section we discuss more "exotic" oxidation reactions in basic electrolytes, such as the oxidation of complex fuels: borohydride, boranes and hydrazine. All these molecules contain hydrogen in their structure and have been studied as promising fuels for DLFCs.

4.1. Borohydride

The borohydride (BH₄⁻) oxidation reaction (BOR, Equation 4.1) is thermodynamically possible 0.41 V negative of the equilibrium potential of the hydrogen electrode (Equation 4.2). This conveys large theoretical energy density and specific energy to direct borohydride fuel cells (DBFC), but requires

complete valorization of the BH_4^- fuel (recovering 8 electrons per BH_4^- species) at potentials close to its reversible potential. In practice, this requires electrocatalysts which are very active for the BOR (Equation 4.1), but at the same time are poorly active for the heterogeneous hydrolysis of BH_4^- (Equation 4.3 and 4.4) and for the HER (Equation 4.2).

$$BH_4^- + 8 OH^- \rightleftharpoons BO_2^- + 6 H_2O + 8 e^ E^0 = -0.41 V vs RHE$$
 (4.1)

$$H_2 + 2 \text{ OH}^- \rightleftharpoons 2 H_2 0 + 2 e^ E^0 = 0 \text{ V } vs \text{ RHE}$$
 (4.2)

$$BH_4^- + H_2O \rightleftharpoons BH_3OH^- + H_2 \tag{4.3}$$

$$BH_3OH^- + H_2O \rightleftharpoons BO_2^- + 3 H_2 \tag{4.4}$$

The first study dealing with the borohydride oxidation reaction at Ni electrodes was published by Elder in 1962.¹⁶⁸ As for the two other complex fuels mentioned above (hydrazine and boranes), the practical interest of such studies was linked to electroplating reactions.¹⁶⁹ Two decades later, the group of Suda published the first papers dealing with the BOR at Ni electrodes (based on commercial µm-size Ni powders mixed with a PTFE binder pasted on a Ni foam) in the context of the DBFC development.^{170,171} They immediately spotted the main advantages and drawbacks of the BOR at Ni: "*The open-circuit potential was found to be about 0.15-0.2 V more negative than the hydrogen potential, depending on the concentration of borohydride. The results of polarization measurements indicated that a high power density can be achieved for the borohydride/Ni system. However, the coulombic efficiency was found to be 50% or less due to hydrogen evolution (…) hydrogen gas was not only generated from the hydrolysis reaction, but also from the electrochemical reaction. The actual anodic reaction of borohydride on the Ni electrode was proved to be a four-electron process rather than an eight-electron one." ¹⁷⁰ Wang et al.*

later on confirmed these findings ¹⁷²: Ni fails to completely valorize the BH₄⁻ fuel and leads to non-negligible *hydrogen escape* ¹⁷³ in the course of the BOR.

Based on this statement, there have been multiple studies exploring Ni-based (sometimes nanostructured) catalysts for the BOR. Many focused on alloying Ni with other metals, like lanthanum in hydrogen storage alloys,^{174,175} boron,¹⁷⁶ cobalt,^{177,178} coper,¹⁷⁹ cerium ¹⁸⁰ or other rare Earths,¹⁸¹ ruthenium,^{182,183} palladium, either in composites ^{184,185} or alloys,¹⁸⁶ gold, either in composites,^{21,187} coreshell ¹⁸⁸ or alloyed ¹⁸⁹ nanoparticles, and platinum in composite ^{185,190} or alloyed ^{191–196} nanoparticles. Most of these strategies have been reviewed in Refs. ^{197,198}, but none of them enabled to reach significant BOR currents at potentials below that of the reversible hydrogen electrode, and in many cases (except for NiCo alloys), the co-metal was governing the overall catalyst's behavior. Besides, the issue of Ni passivation upon incursion to positive (E > 0.4 V vs RHE) potentials or contact with oxygen (either during the electrode storage/preparation or while in an electrolyte) was often ignored, even for pure Ni catalysts,¹⁹⁹ which renders comparison between different materials inconclusive. In essence, the existing literature prior to 2019 did not really enable to state whether nanostructured Ni could be a viable catalyst for DBFC anode.

The picture recently changed, when Oshchepkov *et al.* demonstrated that the state of surface of electrodeposited nanostructured Ni electrodes (Ni_{ED}/C) had a tremendous influence on their BOR activity.²⁰⁰ The authors took benefit of the slow HER at a reduced Ni surface (see Section 3), to exploit electrocatalytic activity of carbon-supported electrodeposited Ni_{ED}/C towards the BOR: in other words, the net BOR oxidation current at potentials below E = 0 V vs RHE is larger when the electrode is less HER active (Figure 6). In result, significant BOR currents are monitored below E = -0.2 V vs RHE on a reduced metallic Ni_{ED}/C (curve 3 in Figure 6b and 6a), the limiting current being already reached at E = 0 V vs RHE (a potential where most of the catalysts listed above as well as their PGM-based counterparts
barely initiate the BOR). On the contrary, a partially-oxidized Ni_{ED}/C demonstrates significantly lower BOR activity, which goes along with its higher HER activity (curve 2 in Figure 6b and 6a). Finally, a strongly-oxidized Ni_{ED}/C is neither BOR nor HER active. This observation essentially explains the very contradictory results reported for Ni-based electrodes in the literature: in essence, most of these catalysts had an uncontrolled (likely strongly oxidized) state of the surface and could not lead to consequent BOR currents below E = 0 V vs RHE.

To better understand the origin of high BOR activity of metallic Ni, DFT calculations were used to determine binding of key BOR intermediates on Ni(111) and compare them with those of Pt(111), Au(111) and Pd(111).²⁰⁰ It was found that the binding of adsorbed BH₄ with the metallic Ni surface is intermediate to that of Pt and Au. It is strong enough to allow for the B-H bond breaking at low overpotential but not too strong to result in surface poisoning by BH_{ad} species as on the Pt surface.¹⁷³

These promises of reduced Ni_{ED}/C catalysts were then confirmed in a DBFC configuration (Figure 7); Ni nanoparticles could be electrodeposited directly on a carbon-based commercial gas diffusion electrode (GDE), mounted as the anode in a membrane electrode assembly with a Nafion cation-exchange membrane and a commercial Pt-based cathode, and tested in a DBFC unit cell.²⁰⁰ In these tests, the Ni_{ED}/GDE outperformed a Pt_{black} anode in identical operating conditions: the Ni-based DBFC presented a higher open circuit voltage (OCV = 1.2 V against 1.0 V for the Pt-based DBFC), hereby confirming the possibility to catalyze BOR at potential below E = 0 V vs RHE, but also led to larger current densities at a given cell voltage. Overall, the power density of the Ni-based DBFC was ca. 2.5 times superior to that of the Pt-based DBFC, but larger hydrogen escape was monitored, owing to the inability of Ni to completely valorize produced H₂. More recently Braesch et al. have further improved the DBFC performance by preparing Ni-based anodes with an open structure to facilitate mass-transfer of either borohydride or evolved hydrogen.²⁰¹ For this purpose, the authors used electrochemically-assisted etching

in order to depassivate the surface of an initially-oxidized (because of storage under air) Ni felt (NFT) support. The latter was further modified by electrodeposition of Ni nanoparticles providing high active surface area and ensuring its metallic state (Figure 8). The maximum power density of the DBFC achieved using Ni_{ED}/eNFT (Ni nanoparticles electrodeposited on an etched NFT) anode was as high as 470 mW cm⁻² at *ca*. 0.58 A cm⁻², while the current density at the cell voltage of 1.0 V was more than two times higher compared to Ni_{ED}/GDE anode (23 mA cm⁻² *vs* 11 mA cm⁻² after 12 hours, respectively).

These results clearly demonstrate that Ni-based catalysts, provided that they are duly-prepared and employed, can outperform PGM-based ones for the oxidation of complex fuels in alkaline conditions, which is promising for the development of PGM-free DLFCs.



Figure 6. (a, b) CVs and (c) chronoamperommetry curves obtained for various Ni_{ED}/C catalysts with a similar loading on a rotating disc electrode but different surface state (as indicated in panel a) under N_2 atmosphere at the following conditions: (a) 1.0 mol L⁻¹ NaOH, v = 20 mV s⁻¹, $\omega = 0$ rpm, (b)

1.0 mol L^{-1} NaOH + 5 mmol L^{-1} NaBH₄, v = 20 mV s⁻¹, $\omega = 1600$ rpm, (c) 1.0 mol L^{-1} NaOH + 5 mmol L^{-1} NaBH₄, $\omega = 1600$ rpm, E = 0.1 V vs RHE. Reproduced from ²⁰⁰ with permission from the American Chemical Society.



Figure 7. Characterization of Ni_{ED}/GDE samples. SEM images of pure carbon cloth (SGL carbon, SIGRATEX) (a) and after electrodeposition of Ni nanoparticles at low (Ni_{ED}/GDE-L, c and e) and high (Ni_{ED}/GDE-H, d and f) metal loading; (b) XRD patterns obtained for the corresponding samples; (g) Ohmic-drop corrected I-U and I-P curves registered in fuel cell configuration using either Ni_{ED}/GDE or Pt-black/GDE anodes and Pt-black/GDE cathode; (h) Durability study of Ni_{ED}/GDE-H anode at the

potentiostatic hold at U = 1.0 V. Reproduced from ²⁰⁰ with permission from the American Chemical Society.



Figure 8. (a) CVs obtained for various Ni-based DBFC anodes at $v = 20 \text{ mV s}^{-1}$ in 1 mol L⁻¹ NaOH, $T = 25^{\circ}$ C for (geometric surface area 8 cm²; ECSA values indicated in the plot): pristine Ni felt (NFT), etched Ni felt (eNFT), Ni nanoparticles electrodeposited on the etched Ni felt (Ni_{ED}/eNFT). (b) DBFC performance of the electrodes of panel (a) compared to Ni_{ED}/GDL and Pt black/GDL from Ref. ²⁰⁰ and Pt/GDL, $T = 60^{\circ}$ C, flow rate 50 mL min⁻¹. (c) Stability measurements at 1 V for the four different electrodes. (d) Hydrogen escape (%) during DBFC test measured using a H₂ pump connected to the outlet of the cell. Reproduced from ²⁰¹ with permission from Willey.

4.2. Boranes

Studies of borane electrooxidation on Ni electrodes were firstly motivated by the need to understand electroplating reactions.²⁰² These works were pioneered by Burke et al., who studied the behavior of dimethylamine-borane (DMAB) at mechanically-polished Ni.²⁰³ Ni was found to be very active for the reaction in its reduced state until the onset of β -Ni(OH)₂ formation (-0.3 < *E* < 0.4 V *vs* RHE), above which it gets passivated and hence inactive for the DMAB oxidation reaction (DMABOR). Such low (*E* = -0.3 V *vs* RHE) potential for the reaction onset suggests that BH₃OH⁻ is the reactive species (eq. 4.5), as had been demonstrated for gold surfaces two decades earlier.²⁰⁴

$$BH_{3}OH^{-} + 6 OH^{-} \rightleftharpoons BO_{2}^{-} + 5 H_{2}O + 6 e^{-}$$
(4.5)

Since then, to the authors' knowledge, only two papers have dealt with borane oxidation at Ni-based catalysts. Zadick et al. studied a series of Ni₃M/Ketjenblack catalysts (M = Pd, Ag and Co) for the ammonia borane (AB) oxidation reaction (ABOR).²⁰⁵ All the catalysts were active for either the direct ABOR or for the decomposition of AB to produce hydrogen, the latter being then oxidized in the HOR. The noblest catalyst (Ni₃Pd) exhibited the most positive ABOR onset potential (a clear drawback), because it essentially promoted fast hydrogen generation (by decomposition of AB or by evolution from water through the HER) and oxidation reactions and not the ABOR. The lowest ABOR onset potential was reached for the non-noble material (Ni₃Co), the least active for AB decomposition and H₂ generation (an advantage). Unfortunately, this material was also the less easy to depassivate (Ni/Vulcan XC72 being completely inactive in similar conditions, owing to its strongly oxidized surface state upon the synthesis method chosen then), but in the end, enabled the best compromise in terms of cost *vs* activity/durability (this material leads to a low onset potential and reasonable ABOR activity at fuel cell anode-relevant

potentials, Figure 9). Later on, Wang et al. ²⁰⁶ studied Ni_{1-x}M_xSe₂ (M = Fe, Co, Cu) nanowires grown on carbon fibers (denoted as CFP) for the ABOR and observed near-similar catalytic activities for the ABOR to those of Zadick et al. ²⁰⁵: the most active of their catalysts, Ni_{1-x}Cu_xSe₂/CFP, exhibited a slightly lower ABOR onset potential (*ca.* E = -0.15 V *vs* RHE versus E = -0.1 V *vs* RHE for the Ni₃Co/KetjenBlack of Zadick et al.) and equivalent kinetic current density to that of Zadick et al. An asset of these two catalysts (Ni_{1-x}Cu_xSe₂/CFP and Ni₃Co/Ketjenblack) is that they are poor HOR/HER catalysts, which enables low-potential oxidation of the AB fuel. In both cases, the activity in ABOR operation is rather stable, showing insignificant signs of poisoning of the catalyst surfaces by reaction intermediate species/products, which leaves hope to develop practical Ni-based catalysts for the ABOR in a direct borane fuel cell.



Figure 9. Comparison of the ABOR activity measured in 5 mmol $L^{-1} AB + 0.1 \text{ mol } L^{-1} \text{ NaOH at } E = 100 \text{ mV vs.}$ RHE, T = 25 °C and $\omega = 1600 \text{ rpm}$ for the Ni₃Ag/C, Ni₃Pd/C and Ni₃Co/C catalysts (C = Ketjenblack). Reproduced from ²⁰⁵ with permission from Elsevier.

4.3. Hydrazine

The use of Ni-based electrocatalysts (namely Ni salts and Ni boride "impregnated on carbon") for hydrazine oxidation dates back to 1970.²⁰⁷ Note that as the oxidation of hydrazine for fuel cell applications

is studied in aqueous electrolytes, this reactant is likely oxidized as hydrazine hydrate (hence the notation hydrazine hydrate oxidation reaction, HHOR, eq. 4.6).

$$N_2H_4 + 4 \text{ OH}^- \rightleftharpoons N_2 + 4 H_2O + 4 e^ E^0 = -0.33 \text{ V} \text{ vs RHE}$$
 (4.6)

In his seminal paper, Wiesener noted that the HHOR performance surpassed that of PGM catalysts, provided the Ni surface was "appropriately" activated (treatment with gaseous NH₃ was the best solution according to the author). Many studies followed this initial report, aiming at optimizing the HHOR performance. Meibuhr et al. evaluated NiB/Ni electrodes for the HHOR ²⁰⁸ and found that surfaces reduced for 1 hour in a mixture of 6% H₂ in Ar at mild temperature (93°C) were the most durable in operation, which was explained by their optimal retention of boron in such a reduced NiB layer.²⁰⁹ Considering the above-cited study of the BOR (see section 4.1.), strong dependence of the HHOR activity on the extent of oxidation of the Ni surface might represent an alternative explanation. Ni (studied as a mechanically-polished Ni rod) is itself active for the HHOR, and shows two domains of activity: the first at low potentials around the Ni/ α -Ni(OH)₂ transition (0 < *E* < 0.4 V *vs* RHE), and the second one at higher potentials (above *E* = 1 V *vs* RHE) ²⁰³. This suggests that reversible redox transitions of Ni/Ni(II) and Ni(II)/Ni(III) are involved in the HHOR at low and high potentials, respectively (the second domain being useless for electricity generation in a direct hydrazine fuel cell).

Because practical fuel cell electrodes are necessarily porous, it soon became obvious that nanostructured Ni-based electrodes were mandatory. So, several studies evaluated carbon-supported Ni-based electrodes for the HHOR in either neutral ²¹⁰ or alkaline ²¹¹ conditions, and it became clear that the irreversible oxidation of Ni was a severe impediment for nanostructured materials. As such, many strategies have been employed to make Ni-based HHOR catalysts more robust. Amongst these strategies, forming composites (*e.g.* carbon-supported Ni + Pd ²¹²) or alloys (*e.g.* unsupported NiZn ²¹³, NiCo ²¹⁴, NiLa ²¹⁵ and NiMo ²¹⁶ alloys) could be promising solutions. Their success was explained by a retardation

of the Ni passivation ²¹²; for example, in the case of the NiZn alloy, Ni alloying with Zn creates a Lewis base catalyst in which Ni acts as an electron-pair donor inducing the adsorption and partial dehydrogenation of hydrazine, which was proposed to prevent (or at least retard) its passivation ²¹³ (it could also be suggested that Zn provides anodic protection of Ni). For the NiCo alloy, the authors noted that Ni was more reduced at a given potential than in a monometallic Ni electrode, here again pointing to the importance of the state of the Ni surface for electrocatalysis of oxidation reactions. The importance of initially metallic Ni for high-activity HHOR was confirmed by Jeon et al. ²¹⁷ and Finkelstein et al. ²¹⁸. In the case of NiLa "alloys", a positive influence of some La(OH)₃ phase (the material is therefore a composite, at least in part) enhances the reaction kinetics by providing OH-species at lower potential values. Finally, for the NiMo catalysts, the non-negligible (< 15 at.%) molybdenum content was proposed "*to stabilize the hydrazine N-N bond, preventing the chemical decomposition of hydrazine into ammonia, and improving the catalyst selectivity towards the complete (and desired) hydrazine oxidation into N₂ gas", ²¹⁶*

This selected literature review demonstrates that there is some merit to combine Ni with other elements, notably to improve its capability to remain in "reduced" state of surface. The strategy was further followed by preparing binary or ternary Ni-based catalysts in a combinatorial chemistry approach,²¹⁹ or oxide-based ^{220,221} catalysts, but the latter materials showed more positive reaction onset potentials and did therefore not enable to surpass catalysts where Ni was in its reduced state.

Then, many other materials were developed for the HHOR, including strategies to enhance the bubble release (N₂ is the final product of the HHOR), via shape optimization (super-hydrophobicity was targeted) of the electrode material,^{222,223} or combination of Ni with Se,^{224,225} noble metal,^{226,227} phosphorus,^{228,229} or sulfur.^{224,230} As a matter of fact, none of these strategies resulted in catalysts which really outperform those prepared by the more conventional routes presented above. As such, real direct hydrazine fuel cell

tests were performed in automotive conditions with the unsupported Ni-Zn catalysts, moreover, a demonstration vehicle was launched with this catalyst ²³¹, clearly highlighting that Ni-based catalysts are practical and good candidates to replace PGM ones for automotive applications.

5. Durability of Ni-based catalysts in alkaline environments

Studies dealing with the degradation of Ni-based catalysts in alkaline environments are scarce. In the 1980s, Ross et al. performed systematic studies of corrosion of carbon black substrates to be used in oxygen evolution anodes in concentrated potassium hydroxide at a moderate temperature ($T = 50^{\circ}$ C). They noticed that while the presence of Co oxides (which were used as oxygen evolution catalysts) accelerated irreversible oxidation of an acetylene black support into CO and/or soluble organic compounds,²³² the presence of Ni oxide nanoparticles led to little or no effect on the corrosion rate of a graphitized carbon black.²³³

More recently, Davydova et al. investigated chemical and electrochemical stability of tailored Vulcan (VXCMAX22) carbon-supported monometallic Ni and bimetallic Ni₃M (M = Co, Fe, Cu, Mo) nanoparticles in supporting alkaline electrolytes, using a combination of *in situ* on-line ICP-MS (inductively coupled plasma mass spectrometry) with a flow cell and *ex situ* TEM (transmission electron microscopy), XPS (X-ray photoelectron spectroscopy) and XRD (X-ray diffraction).²³⁴ The authors noticed different behavior of the studied materials (Figure 10a). On the one hand, two catalysts have shown non-negligible instability in potentiodynamic conditions: Mo intensely dissolves from Ni₃Mo/C, while Cu in Ni₃Cu/C is stable below E = 0.4 V *vs* RHE but undergoes noticeable electrochemical transient dissolution above this potential. On the other hand, pure Ni/C, Ni₃Co/C and Ni₃Fe/C showed negligible dissolution up to E = 0.7 V *vs* RHE. This absence of dissolution does however not grant stability of their HOR performance, since all catalysts lose their HOR activity upon incursions to the potential interval

above E = 0.5 V vs RHE (Figure 10b, for Ni₃Fe/C), owing to the passivation of metal nanoparticles. The authors concluded that fuel starvation events in AEMFCs, which would result in high anodic potentials, require adapted cell activation and operation strategies to prevent any catalyst passivation and an ensuing depreciation of their HOR performance.



Figure 10. (1) Characterization of the dissolution of Ni/C and of several Ni₃M/C catalysts upon potentiodynamic cycling in the range -0.05 < E < 0.7 V vs RHE in 0.05 mol L⁻¹ KOH supporting electrolyte for M = Fe (green), M = Co (light blue), Ni (dark blue), M = Cu (orange), and M = Mo (red). The light gray traces represent the original dissolution data, and the colored components were smoothed for clarity. (11) HOR polarization curves (right panel) for Ni₃Fe/C before (blue) and after (red) various degradation protocols shown in the left panel: 1000 cycles in the potential window of (a) 0 < E < 0.3 V vs RHE and (b) 0 < E < 0.7 V vs RHE, as well as (c) before and after 30 days at open-circuit potential.

0.05 mol L⁻¹ KOH, T = 25 °C, $v = 1 \text{ mV s}^{-1}$, $\omega = 1600 \text{ rpm}$. Reproduced from ²³⁴ with permission from the American Chemical Society.

These results confirmed those of Zadick et al., obtained on a similar class of materials (Ni₃M/ Ketjenblack 600J nanoparticles, with M = Pd, Co and Ag).²⁰⁵ The Ni-based nanoparticles showed neither sign of consequent metal dissolution nor of severe modification of their shape or of detachment from the carbon support, as revealed by identical-location transmission electron microscopy after potentiostatic (at E = 100 mV vs RHE in a 5 mmol L⁻¹ AB +0.1 mol L⁻¹ NaOH electrolyte) or potentiodynamic (between 0.1 < E < 1.23 V vs RHE in 0.1 mol L⁻¹ NaOH) polarization at $T = 25^{\circ}$ C (Figure 11), for Ni₃Co/ Ketjenblack 600J nanoparticles). While the borane oxidation activity did decrease after incursion of the electrodes to high potential values, it could be restored provided a proper re-activation procedure (incursion to low potential values in presence of the strong borane reducer) was adopted. This shows that combining a suitable reducing-potential and the presence of strong reducer in the electrolyte can depassivate Ni₃Co nanoparticles (and even more easily noble-based Ni₃Ag and Ni₃Pd catalysts); the comparison with the results of Davydova et al. ²³⁴, shows that molecular hydrogen is not a sufficiently strong reducer to enable depassivation of this class of non-noble carbon-supported Ni-based catalysts (nanostructured Ni₃M, M = Co, Fc, Cu, etc.).

In any case, carbon-supported Ni-based nanoparticles do not seem to suffer consequent nanoparticles detachment from their carbon support, owing to the rupture of their binding to the carbon substrate by nanoparticles-assisted local corrosion of the carbon into carbonate species. While this mechanism of degradation seriously hampers the long-term stability of PGM-based nanoparticles in alkaline electrolytes,^{235–238} it seems rather less operant in interface with an anion-exchange polymer electrolyte,²³⁹ but no such data exists for Ni-based catalysts, at least to the authors' knowledge.



Figure 11: Representative ILTEM micrographs of a Ni₃Co/Ketjenblack 600J catalyst obtained before (top three micrographs) and after several electrochemical treatments, (i) activation + 6-h-long steadypolarization at E = 100 mV vs. RHE in a 5 mM AB +0.1 M NaOH electrolyte at T = 25 °C (middle three micrographs); (ii) 1000 CV cycles of AST (bottom three micrographs); the AST consisted of repeated cycling between 0.1 < E < 1.23 V vs. RHE in 0.1 M NaOH at T = 25 °C. Reproduced from ²⁰⁵ with permission from Elsevier.

Extended and unsupported Ni surfaces, also show some degree of instability upon electrochemical operation in alkaline electrolytes. As detailed in Section 2, Ni surfaces are subjected to complementary adverse effects, which can affect their electrocatalytic properties ⁴³: below E = 0 V vs RHE, they form Ni hydrides (α -NiH_x and β -NiH_x depending on the applied conditions) ^{96,240}; above E = 0 V vs RHE (and in particular at open-circuit conditions), Ni (hydr)oxides are formed (*e.g.* reversible α -Ni(OH)₂, which mostly reduces back to Ni on subsequent cathodic polarization, but also β -Ni(OH)₂, which cannot be electrochemically reduced back into metallic Ni, and at higher potentials undergoes reversible oxidation into β/γ -NiOOH ^{45–47,114}), which may deactivate the electrode if uncontrolled. Sections 3 and 4 presented the means to optimize the electrode activity for the HOR ^{48,137} and the BOR ²⁰⁰ by controlling the extent of Ni surface oxidation. Appropriately cycling a Ni electrode at high/low potential values could enable to prevent detrimental hydride or oxide formation,²⁴¹ but repeated oxidation and reduction periods (*e.g.* experienced in transient or intermittent operation) may lead to strain in the Ni catalyst, which can induce mechanical failure.⁴³

Such mechanical failure was observed by Schulze and Gülzow ²⁴² when using Raney Ni electrodes for the HOR in an alkaline fuel cell (pasted Raney Ni + PTFE binder electrodes were studied). On the one hand, hydrogen embrittlement caused disintegration of the crystallites/grains of the Ni electrodes, hence their destructuration (also owing to PTFE gradual decomposition). On the other hand, irreversible Ni oxide formation led to their progressive deactivation, overall gradually depreciating the HOR performance in operation. Raney Ni is particularly used for the HER in alkaline environments.^{243,244} This catalyst, obtained from dealloying NiAl or NiZn alloys, is known to deactivate owing to the formation of hydrides when maintained in prolonged hydrogen evolution regime ^{245,246}; the leaching of their residual impurities (*e.g.* Al and Zn) when the electrode is maintained at open-circuit (*i.e.* in discontinuous HER operation) or positive (*e.g.* HOR) polarization also accounts for their gradual loss of activity over time.²⁴⁷ To counter these adverse effects, several doped Raney Ni alloys ^{243,244,248,249} and composites between Raney Ni and metal hydride alloys ^{250,251} have been tested, as well as addition of metal ions/complexes in the electrolyte (*e.g.* iron cations from the electrolyte deposit at the Ni surface and inhibit Ni hydrides formation,^{246,252} while vanadium salts in the electrolyte enhance Ni hydride decomposition,²⁵³ a positive effect in both cases). To date, these strategies were only partly successful, and there remains room for improving the durability and long-term activity of Ni-based electrodes in operation.

Finally, although Ni electrodes show some material's stability in operation in alkaline environments, their progressive deactivation (e.g. by surface oxide or hydride formation) may pose a problem. Future studies should therefore aim at stabilizing such Ni surfaces for real applications, *i.e.* in conditions where the materials could be subjected to oxygen contact, alternation between low and high potential values, open-circuit holds, etc.

6. Conclusions

In this paper, the electrochemical properties of Ni electrocatalysts have been reviewed. Since Ni and its (oxy)(hydr)oxides are stable at high pH (while being unstable at low pH), the electrochemical behavior of Ni in alkaline media was particularly emphasized. As one moves from low to high potentials, the Ni surface is (i) covered by adsorbed hydrogen (H_{ad}) in the hydrogen evolution and likely also in the hydrogen oxidation potential region, (ii) covered by surface α -Ni(OH)₂ and then (iii) β -Ni (OH)₂, the latter being (iv) further oxidized into γ/β -NiOOH in the oxygen evolution region. The occurrence of Ni/ α -Ni(OH)₂ and α/β -Ni(OH)₂/ γ/β -NiOOH transitions in the "low" and "high" potential regions provide practical tools to determine the active surface area of Ni. Besides, these different potential-dependent states of the surface

largely determine the activity of Ni electrodes towards fuel oxidation reactions. This is the case for the HOR in alkaline media, which is strongly accelerated on partially-oxidized Ni surfaces (mixed Ni/NiO_x surface). Alloying/combining Ni with other elements or supporting Ni on well-chosen substrates, can also lead to a better HOR performance, and both strategies have been practically demonstrated in AEMFCs. Oxidation of complex fuels (borohydride, boranes, hydrazine) is also possible on Ni-based electrocatalysts, paving the way to PGMs-free direct liquid alkaline fuel cells. In these cases, Ni is interesting because it is possible to tune its surface state so that it catalyzes the desired reaction (*e.g.* the borohydride oxidation reaction) without promoting possible competing reactions (*e.g.* the hydrogen evolution reaction or the hydrolysis of the fuel species). It should be noted however that much work still needs to be done by applying *in situ* spectroscopic tools (*e.g.* FTIR), *ab initio* calculations, and kinetic modeling in order to better understand the nature of adsorbed species on Ni surface depending on the applied potential and mechanisms of oxidation of these fuel molecules on Ni electrodes, and propose viable solutions for the development of more active catalysts.

Lastly, this selected review also addresses the durability of Ni-based catalysts in alkaline environments. It is notably shown that, unlike carbon-supported PGM-based electrocatalysts, Ni/C and NiM/C materials are rather robust against the detachment of the metal nanoparticles from the carbon support, and, depending on their composition, can be resistant to metal dissolution. Stability of Ni-based electrodes is however strongly dependent on the explored potential domain, and more work is necessary to clarify whether they are sufficiently stable under the AEMFCs and DLFCs operation conditions. In addition, it is clear that materials stability of Ni-based catalysts must not be confused with the performance stability of the corresponding electrodes. Indeed, passivation by Ni oxides or hydride formation is possible if the Ni-based electrode is brought to "inadequate" potential intervals during the operation. This means that,

although Ni catalysts do really present strong assets for use in fuel oxidation reactions, their practical deployment in long-lasting real devices still requires research and development.

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Author Contributions

All authors contributed to the analysis of the referenced papers, writing and reviewing of the manuscript.

Notes

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