Ni Promotion by Fe: What Benefits for Catalytic Hydrogenation?

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Abstract: Metallic nickel is known to efficiently catalyze hydrogenation reactions, but one of its major drawbacks lies in its lack of selectivity, linked to side-reactions of hydrogenolysis and over-hydrogenation. More selective hydrogenations can be obtained upon the introduction of a second metal in combination with Ni. Fe is an interesting choice, as it is a cheap and abundant metal. This review aims at discussing the advantages and constraints brought by the preparation procedures of bimetallic supported Ni–Fe nanoparticles, and at analyzing the benefits one can draw by substituting Ni–Fe supported catalysts for Ni monometallic systems for the catalytic hydrogenation of organic molecules. Specific formulations, such as Ni₇₅Fe₂₅, will be singled out for their high activity or selectivity, and the various hypotheses behind the roles played by Fe will be summarized.

Keywords: Nickel; iron; bimetallic catalysts; hydrogenation; hydrogenolysis

1. Introduction

Compared to noble metals, non-noble metals such as Fe, Co, Ni and Cu are attractive alternatives for catalytic hydrogenations because of their low cost. However, Fe is often the least active metal of the series and deactivates easily, Co is the most expensive one, and Cu usually exhibits a low activity due to the difficulty of having it highly dispersed on a support. Metallic nickel stands out as the best compromise. Raney nickel has been used for almost one century as a catalyst in organic chemistry, in the laboratory and industry [1]. However, one of its most important drawbacks is its pyrophoricity, because of which it must be handled under an inert atmosphere. Another disadvantage of nickel is its fast deactivation due to the deposition of carbon and formation of carbide. Finally, a major problem with Ni-based catalysts is their possible lack of selectivity in the hydrogenation of oxygen- or nitrogen-containing functions, as they are highly active both in the hydrogenation of carbon–carbon bonds and of aromatic rings, and in hydrogenolysis side-reactions.

The addition of a second metal as a promoter is known to effectively modify the properties of Ni catalysts [2]. Bimetallic Ni-based catalysts exhibit specific properties, via a synergistic or bifunctional effect, which differ from those of the parent metals and offer the opportunity to design catalysts with an activity and selectivity that exceed those achievable with monometallic nickel. Because the size and composition of bimetallic particles affect their catalytic properties, these parameters should be precisely characterized and optimized in order to correlate the catalyst formulation and its performances in reaction.

Iron has been cited as a promising promoter of nickel in bimetallic Ni–Fe combinations because of its low cost, of its abundance and of its effective enhancement of catalytic performances (promoter is understood here in a general sense of boosting catalytic performances, which does not necessarily imply that Fe is added in minor proportions). Since the 1980s, supported Ni–Fe catalysts have been
described as promising systems in three major domains: The catalytic reforming of methane or of organic molecules, for which Ni–Fe systems present a gain of stability compared with monometallic Ni, linked to a slower rate of carbidization [3–8]; the hydrogenation of CO and CO₂, to methane or C₂ hydrocarbons, for which an increase of activity and selectivity to CH₄ or short olefins, depending on the reaction conditions, has been noted [9–16]; finally, the selective hydroconversion of organic molecules. Since the first results published in 1992 by Kumbhar et al. on the selective hydrogenation of ketones and nitriles [17], about 20 papers have appeared so far on the hydroconversion of organic molecules on Ni–Fe supported catalysts, 18 since 2013 and 10 since 2017, which emphasizes the potential of these systems for this family of applications. However, a survey of the literature also shows that standard preparation methods for Ni–Fe bimetallic catalysts, such as co-impregnation, may yield metal particles of different size or composition [5,10,17,18]. As a consequence, owing to a lack of control of the nature of the individual nanoparticles (alloyed vs. segregated metals, disparities in metal content), the existence of a general correlation between the nanoparticles composition and the best catalytic results in the hydrogenation of organic molecules, as for now, remains an open question. The present review aims at discussing the advantages and constraints brought by different preparation procedures of bimetallic supported Ni–Fe nanoparticles, and at analyzing the benefits the literature has reported upon substituting Ni–Fe supported catalysts for Ni systems for the selective hydrogenation of organic molecules. We will summarize the various hypotheses explaining the action of Fe with respect to these catalytic reactions, identify the most promising formulations, and suggest which lines of research appear as key issues for the future.

In order to make comparisons easier between the different articles, the formulation of the catalysts described in the literature will be written in the following way: (Total metal content, wt%) NiₓFe(100-x)/support, where x and (100-x) represent the respective molar proportions of the two metals in the catalyst formulation.

2. Structure, Size, Composition and Formation of Ni–Fe Supported Nanoparticles

The nominal proportions of Ni and Fe, the preparation method of Ni–Fe catalysts, the temperature of reduction, the nature of the support, all have an influence on the structure of supported Ni–Fe nanoparticles, on their size and on their actual metals content, which are likely to affect the performances of the catalyst in the hydrogenation of organic molecules. This section provides elements for a better understanding of the relationships existing between these experimental parameters and the structural characteristics of Ni–Fe nanoparticles.

2.1. Influence of the Metals Content on the Structure of Ni–Fe Nanoparticles

According to the simplified phase diagram published in 1937 by Bradley et al. (Figure 1a) [19], the Ni–Fe bulk alloy displays two different crystal structures, depending on the Fe and Ni contents, and on the temperature. The α phase, containing less than 8 at% of Ni, exhibits a body-centered cubic structure (bcc), like metallic Fe, while the γ phase, containing more than 32 at% of Ni, exhibits a face-centered cubic structure (fcc), like monometallic Ni. Solubility between Fe and Ni does not extend to the whole composition range, and compositions between 8 and 32 at% lead to a mixture of bcc and fcc phases after cooling. Except for the lowest Ni contents, for which the alloy retains the bcc structure, the fcc phase dominates above 400 °C. More complex phase diagrams based on the study of meteorites include the existence of intermediate phases in the γ domain, such as awaruite, of average composition FeNi₅, as shown in Figure 1b [10].

In the literature dedicated to Ni–Fe catalysts, experimental results generally follow the trend given by the simpler diagram. Matsuyama et al. reported that Fe-rich alloys (<5 at% Ni) exhibit the bcc structure, while Ni-rich alloys (>40 at% Ni) exhibit the fcc structure, the intermediate range corresponding to a mixture between particles of the two varieties [20]. This co-existence on supported catalysts has indeed been noted for nominal formulations NiₓFeₘ and NiₓFeₙ [14,17,21–24]. De-mixing has sometimes been mentioned to occur for preparations based on nominal formulations expected to fall in the γ domain, such as NiₓFeₘ or NiₓFeₙ [21,24,25]. The identification of the ordered intermediate NiₓFeₘ phase has been mentioned in a few instances [26,27].
Fe is a slightly larger atom than Ni (empirical atomic radius: 0.126 vs. 0.124 nm). As a consequence, the lattice parameter of the fcc cell in the $\gamma$ domain decreases when the Ni content increases. This trend globally follows a linear Vegard’s law, as has been verified by several teams [4,8,28]. The evidence for alloying, and the composition of Ni–Fe particles, can thus be derived from X-ray diffraction (XRD) measurements.

2.2. Synthesis Routes for Supported Ni–Fe Nanoparticles: Influence on the Particles Size and Composition

Three easy-to-scale methods have been principally used for the preparation of supported Ni–Fe bimetallic catalysts: Co-impregnation, and, to a lesser extent, sol–gel routes and precipitation, with different outcomes in terms of particle size and composition. We will list below the characteristics of these three methods, their advantages and their limits.

2.2.1. Incipient Wetness Impregnation

Incipient wetness impregnation (IWI) is a classical way to prepare Ni–Fe supported catalysts. The porous support is wetted by an aqueous solution containing the Fe and Ni precursors (usually Fe(III) and Ni(II) nitrates) in the proper Fe/Ni ratio. The volume of the solution is equal to the pore volume of the support. Drying, calcination in oxidative atmosphere and reduction steps are subsequently performed to obtain reduced Ni–Fe particles.

The phases detected by XRD and $^{57}$Fe Mössbauer spectroscopy after calcination are usually the two monometallic metal oxides, FeO$_x$ and NiO. Ni–Fe mixed oxides have also been identified. Despite the detection of $\alpha$-FeO$_x$ by XRD in Ni–Fe/SiO$_2$ and Fe/SiO$_2$ catalysts, a difference in the half linewidths of the Mössbauer peaks, significantly broadened for Ni–Fe/SiO$_2$ compared with Fe/SiO$_2$, suggested that iron ions had nickel second-shell neighbors. The oxide was thus designated as $\alpha$-Fe$_{\gamma}$.Ni$_{\gamma}$O$_{\gamma}$, for Ni-doped FeO$_x$. A low intensity of NiO XRD reflections also suggested that most of Ni was in fact present in substitution in iron oxides [23]. In the second case, the NiFeO$_x$ phase was identified by XRD [29]. IWI thus does not ensure a complete association of the two metals during the first stages of the catalyst preparation. However, doping of iron oxides by Ni$^{2+}$ ions (or, conversely, of cubic NiO by iron ions) probably occurs, but can be hardly evidenced by routine characterization techniques such as XRD. Because they do now allow an early association between the two metals, successive impregnations may be less successful than co-impregnation [24,30,31].
The reduction temperature of the oxidic phases usually ranges between 300 and 500 °C. A usual drawback from IWI, as has been recently reviewed by Tomishige et al. [5], is that the particle size distribution of reduced Ni–Fe nanoparticles may be broad or bimodal, with a co-existence between particles smaller and larger than 10 nm (Figure 2A) [18,25,33]. Mild drying under vacuum prior to reduction has been reported to limit the growth of the particles [25], possibly because anhydrous nitrates do not melt and diffuse upon heating over the support surface.

An alternative way to form Ni–Fe particles on a carbonaceous support, without reduction in H₂, was proposed by Chieffi et al. [34]. Carbon-supported Ni–Fe nanoparticles were prepared by carbothermal reduction of a cellulose filter paper impregnated with Fe(III) and Ni(II) nitrates, at 800 °C for 2 h under nitrogen. The shape and size of the cellulose fibers was retained after heat-treatment. The crystallite size of Ni–Fe nanoparticles was comprised of between 20 and 30 nm.

Another problem with IWI is the possible disparities in the composition of Ni–Fe particles on a given catalyst, which can supposedly result from the non-uniform composition of the oxidic phases formed upon calcination. It is difficult to identify the precise origin of the problem, which can derive from the characteristics of support and/or of the conditions of drying. For example, Wang et al. prepared a series of Ni–Fe/α-Al₂O₃ catalysts with different Fe/Ni ratios [18]. For a Ni₆₇Fe₃₃ formulation (Fe/Ni ratio = 0.5), the actual Fe/Ni ratio measured by energy-dispersive X-ray spectroscopy (EDX) widely varied in the range 0.06–2.6 (Figure 2a). Kumbhar et al. also found that 20 wt% Ni₆₇Fe₃₃/SiO₂ and 20 wt% Ni₁₀₀Fe₁₀₀/SiO₂ catalysts both presented a mixture of fcc and bcc phases, implying the presence of nanoparticles with very different compositions, in particular Fe-rich nanoparticles [17]. These disparities may make the analysis and interpretation of catalytic data tricky, as each class of nanoparticles may display its own catalytic performances. Catalysts are often characterized by XRD, which provides at best an average view over the composition of the nanoparticles, as the XRD peaks position changes within a very small range of a few tenths of degree upon modifying the metals content across the whole γ domain. The composition of individual nanoparticles is not always studied nor reported. Especially for IWI preparations, scanning transmission electron microscopy (STEM)–EDX mappings [34] are necessary to confirm the alloying between the two metals within bimetallic nanoparticles (Figure 2B), and the absence of heterogeneities of composition.

### 2.2.2. Sol–Gel Routes

In sol–gel routes, a sol is formed by adding an acid to a solution containing an alkoxide precursor of the support (for example, aluminium isopropoxide when alumina is selected as the future support). After addition of Fe and Ni precursors (such as nitrates or acetates), the system gradually evolves towards the formation of a gel containing the three metallic elements [12,13,35,36]. Systems with an ordered porosity can be prepared by addition of a porogen and slow evaporation of the solvent (evaporation-induced self-assembly, EISA) [36]. Calcination and reduction lead to Ni–Fe alloyed nanoparticles dispersed onto the support.
Figure 3. TEM images and size histograms of Ni(–Fe)/ordered mesoporous alumina (OMA) catalysts prepared by the evaporation-induced self-assembly (EISA) method and reduced at 700 °C: (a) 8 wt% Ni/OMA; (b) 9 wt% Ni9Fe9/OMA; (c) 9 wt% Ni8Fe16/OMA. Reprinted with permission from Reference [36]. Copyright 2017 American Chemical Society.

Because of their early insertion in what will become the catalyst support, the resulting solid can be considered as more homogeneous than in the case of IWI. But phases obtained upon calcination are a mixture of several metal oxides, some of them being formed by insertion of the Fe and Ni ions into the support: NiAl2O4, FeAl2O4, AlFeO3 [12,13,35]. Metal ions present in spinel and perovskite phases are difficult to extract and reduce, thus requiring a high reduction temperature; for example, a temperature of 700 °C has been selected by Hwang et al. [12,13], and Liu et al. [36]. However, sintering is limited upon reduction. The average particle size is reported to be smaller than 8 nm with narrow size distributions (Figure 3) [12,13,36], and in this respect sol–gel routes appear to provide a better control than IWI. The drawback of the sol–gel route is the cost of the alkoxide precursor, and its sensitivity to humidity.

2.2.3. Co-Precipitation and Deposition–Precipitation

A procedure that associates a simple implementation like IWI, and the early insertion of the two metals in the solid precursor as is achieved with the sol–gel route, is co-precipitation. Layered double hydroxides of the hydrotalcite-type can be conveniently obtained by basification, with ammonia or carbonate ions, of an aqueous solution containing Fe, Ni, Al and Mg precursors [8,37–39]. This method has become favored for the preparation of Al2O3-supported catalysts for catalytic reforming or CO/CO2 methanation [3–5,8,16]. Co-precipitation has also been reported in association with an organic substrate like melamine, which after carbonization allows protection of the metal nanoparticles [40]. Ni–Fe particles of homogeneous size (3–7 nm) and composition are formed upon reduction (Figure 4), but, as in the case of the sol–gel route, a high temperature, up to 800 °C, is sometimes necessary for complete reduction [8].

Deposition–precipitation (DP) is a related preparation method that leads to the deposition of a Fe- and Ni-containing compound either onto a pre-existing support, or as the result of dissolution–reprecipitation reactions involving the support. Nickel and iron salts are introduced into an aqueous solution with the support, such as SiO2 or Al2O3, and the pH of the solution is progressively increased by addition of a base, till precipitation takes place. A homogeneous increase of pH can be obtained by decomposing urea above 80 °C. Ni–Fe nanoparticles are formed upon reduction in H2.
Figure 4. TEM characterization of a 10 wt% Ni₈₀Fe₂₀/magnesium aluminate catalyst prepared by co-precipitation and reduced at 800 °C: (a) Bright-field TEM image; (b) STEM–EDX analysis, including the Ni/Fe atomic ratio for individual nanoparticles (nominal ratio: 4); (c) linear STEM–EDX scanning across one Ni–Fe nanoparticle. Reprinted with permission from Reference [8]. Copyright 2017 American Chemical Society.

Mutz et al. have used Ni(II) and Fe(III) nitrates as precursors for a DP process with urea on alumina [41]. A narrow distribution around a small average size (3.9 nm) and a high homogeneity of chemical composition (Ni₇₅Fe₂₅) were achieved after reduction at 500 °C. Because of the different valences of the ions, iron (III) hydroxide should deposit at a lower pH than nickel (II) and iron (II) hydroxides (pH 2–3 vs. 5–6 for dilute solutions). The homogeneity of the solid obtained by Mutz et al. can be attributed to the use of alumina as support, and to the likely precipitation not of a hydroxide, but of an Al-based hydrotalcite phase apt to accommodate Fe³⁺ ions.

Using a different support may require a change of strategy. In order to avoid the early, separate precipitation of Fe(III)-containing compounds, precursors to Fe-rich particles, Shi et al. rather selected Ni(II) and Fe(II) sulfates for the preparation of Ni–Fe/SiO₂ catalysts, and eliminated all sources of oxidation from the preparation protocol (use of sulfuric acid instead of nitric acid, degassing of the solution by Ar) [28]. After 22 h of DP with urea under inert atmosphere, the resulting solid was a 1:1 ill-organized (Ni(II), Fe(II)) phyllosilicate containing a slight excess of Ni compared with the initial composition of the solution. Iron ions would later oxidize in air to Fe(III), though without any consequence on the subsequent formation of the Ni–Fe particles.

Like on co-precipitated catalysts, reduced Ni–Fe nanoparticles on catalysts prepared by DP present a size often smaller than on catalysts prepared by IWI (3–7 nm) [28,41,42]. When the reduction temperature is low, the metals may not be fully reduced, and a high reduction temperature may be necessary because of the presence of phases formed between metal and support (silicates). Though an increase of the particle size has been reported (up to 15.2 nm [7]), nanoparticles prepared by DP can sustain a high reduction temperature without undergoing severe sintering. Shi et al. have shown that on silica, the increase of the average Ni–Fe particle size is moderate between 500 and 700 °C (5.1 vs. 5.4 nm), without major change in the size histogram. The composition of the particles was reported to be homogeneous: The standard deviation calculated from EDX measurements carried out on a population of individual particles was 8 at%Fe for Ni₆₉Fe₃₁ nanoparticles, and no monometallic or Fe-rich particles were detected [28].

Alternatively, van de Loosdrecht et al. prepared Ni–Fe catalysts by implementing the deposition–precipitation of iron–nickel cyanide complexes onto a TiO₂ support [43,44]. A potassium iron cyanide solution was injected into a suspension of TiO₂ in a solution of nickel nitrate kept at pH 5. The resulting solid was calcined in air at 300 °C to remove the cyanide ligands. The total reduction of the metals was reached only at 900 °C according to temperature-programmed reduction (TPR) profiles, because of the formation of nickel or iron titanate. The size of the Ni–Fe particles after reduction of 900 °C, calculated from XRD data, turned out to be quite large, 15–20 nm. Co-precipitation of oxalates over a melamine substrate has been recently described, leading to 38 nm bimetallic nanoparticles after carbothermal reduction [27].

In conclusion, the phases obtained after incipient wetness impregnation and calcination are based on the two monometallic metal oxides, which shows that the two metals are not associated at
this stage of the preparation, but doping of the monometallic oxides by the other metal is likely to occur. Phases detected after calcination when sol–gel or co-precipitation methods are implemented are a mixture of monometallic and mixed oxides, some of them involving the support (NiAl₂O₄, FeAl₂O₄), which may require a high temperature to achieve complete reduction. Ni–Fe nanoparticles prepared by procedures based on co-precipitation exhibit a high level of homogeneity in terms of size and composition. The particle size obtained by sol–gel route, co-precipitation or deposition–precipitation is usually smaller, and the size distribution narrower, than those obtained by incipient wetness impregnation, which for the sake of convenience still remains a standard procedure that can be applied to any porous support.

2.3. Reducibility of Ni–Fe Supported Oxidic Phases and Formation of Ni–Fe Nanoparticles

The reduction stage is crucial to form Ni–Fe nanoparticles since metal oxides do not bear the active sites for hydrogenation catalysis. Globally speaking, Fe₂O₃ and NiO reduce in the lower temperature range (below 500 °C) if the interaction between the oxidic phases and the support surface is weak (as is the case for IWI when oxidic particles are large), and higher temperatures are required for reduction completion (up to 900 °C) when metal ions form oxidic phases with the support (IWI, sol–gel route, DP). In one of the earliest studies on the reducibility of the Ni–Fe system, Unmuth et al. have investigated by TPR the reducibility of Fe₂O₃ and NiO on SiO₂ independently, with different temperature rates [23]. Two stages of reduction were evidenced for iron oxide: An “α peak” was assigned to the reduction of Fe₂O₃ to Fe₃O₄ at about 320 °C, while a “β peak” was assigned to the reduction of Fe₃O₄ to Fe at about 450 °C. The reduction of bulk NiO to Ni was found to take place at about 300 °C. A shoulder on the peak was thought to be characteristic of very small particles of oxide strongly interacting with silica. Therefore, Fe was found to be the more difficult metal to reduce of the two, notwithstanding particle size and support effects that may change the game, and make a phenomenological interpretation only based on TPR profiles difficult.

The presence of Ni on bimetallic Ni–Fe catalysts is widely considered to favor the reduction of Fe, even at low Ni contents. As early as 1982, Brown et al. published a series of TPR profiles evidencing the gradual shifts to lower temperatures of reduction peaks when increasing the Ni content of co-precipitated Ni–Fe catalysts, with a significant effect noticed for Ni₄₀Fe₆₀ (lower limit of the fcc domain) and Ni-richer formulations (Figure 5A) [45]. In the Fe-rich domain, Li et al. found that while the reduction temperature for monometallic Fe was 570 °C, a small shift to 550 °C appeared for the Ni₁₃Fe₈₇ formulation, and the reduction temperature for the Ni₃Fe₇₉ formulation was lowered to 510 °C [38].

We will detail here only a few examples from the literature, selected to show that this trend is verified for a variety of solids. A dramatic decrease of reduction temperature for Ni–Fe/SiO₂ catalysts compared with Fe/SiO₂ was found by Yu et al. [46]. A first reduction peak at 260–510 °C for Fe/SiO₂ was attributed to the reduction of Fe₂O₃ to Fe₃O₄ and a second peak above 930 °C was ascribed to the reduction of Fe₃O₄ to Fe, but should most likely be attributed to the reduction of iron silicates (Figure 5B). The main reduction peak at about 390 °C for Ni/SiO₂ was assigned to the reduction of NiO to Ni. In contrast with Fe/SiO₂, Ni–Fe/SiO₂ calcined solids containing the same amount of Ni (8 wt%) and increasing amounts of Fe did not display any reduction peak above 530 °C, implying the simultaneous reduction of iron oxide and nickel oxide even when Ni was the minor metal.

In the case of systems in which the two metals are initially inserted in an oxidic matrix, Oemar et al. have studied the reducibility of perovskites containing Fe and Ni. No peak was observed on the TPR profile of LaFeO₃ up to 900 °C (Figure 5C) [47]. However, a reduction peak appeared at 850 °C after adding a small amount of Ni (LaFe₃₈Niₓ₆₂O₉), which indicates that some chemical interplay existed between Ni and Fe ions inside the structure. The peak shifted to lower temperatures, down to 540 °C, when further increasing the Ni content up to LaNiO₃.
The reducibility of the 17 wt% Ni<sub>75</sub>Fe<sub>25</sub>/Al<sub>2</sub>O<sub>3</sub> catalyst prepared by DP by Mutz et al. was investigated by TPR and compared to the profile obtained for a monometallic Ni catalyst [41]. The H<sub>2</sub> consumption profile of the Ni catalyst revealed one main peak at 573 °C, referring to the reduction of NiO to metallic Ni. The shoulder around 750 °C was ascribed to smaller NiO particles exhibiting stronger interactions with the alumina support. The main reduction peak shifted to lower temperatures for the Ni<sub>75</sub>Fe<sub>25</sub>/Al<sub>2</sub>O<sub>3</sub> catalyst (from 573 to 555 °C). The two reduction peaks of Ni<sub>75</sub>Fe<sub>25</sub>/Al<sub>2</sub>O<sub>3</sub> were assigned to the reduction of Fe<sub>2</sub>O<sub>3</sub> to Fe<sub>3</sub>O<sub>4</sub> in the ternary Al–Ni–Fe oxide (350 °C), and to the reduction of Fe<sub>3</sub>O<sub>4</sub> to Fe and NiO to Ni (555 °C).

More recently, Shi et al. followed by in situ X-ray absorption and <sup>57</sup>Fe Mössbauer spectroscopies the kinetics of reduction of a 1:1 (Fe(III, Ni(II)) phyllosilicate formed by DP with urea [28]. They could evidence a succession of stages that confirms the classical interpretations of TPR thermograms for Ni–Fe systems: Reduction of Fe(III) to Fe(II) between 275 and 360 °C; between 360 and 500 °C, simultaneous reduction of Ni(II) to Ni(0) and Fe(II) to Fe(0), which confirms the decisive role of Ni in the reduction of Fe when the two metals are closely associated in the solid; completion of the reduction of Fe(II) to Fe(0) between 500 and 700 °C, and migration of metallic Fe into the core of the just-formed Ni–Fe fcc nanoparticles.

It derives from these results that the higher the reduction temperature, the more Ni–Fe alloyed nanoparticles are formed and the higher the extent of Fe reduction, with a lower risk of particle sintering when metal particles are stabilized by interactions with the support, such as for sol–gel or
precipitations routes. Globally speaking, a SiO₂ or a TiO₂ support provides a higher reducibility compared with an Al₂O₃ support, because of the easy formation of spinels on Al₂O₃, in which case some Ni- or Fe-containing phases have been reported to remain unreduced even after reduction at 900 °C [45]. Evidence for this support effect was clearly shown by Jiang et al. using four catalysts prepared by co-impregnation in similar conditions, two on TiO₂ and two on Al₂O₃ [48]. The two Ni–Fe/TiO₂ catalysts appeared as almost fully reduced at 440 °C, as their ⁵⁷Fe Mössbauer spectrum presented the singlet peak characteristic of Ni–Fe alloys, while their Ni–Fe/Al₂O₃ counterparts, though prepared in the same conditions, still contained a significant amount of unreduced Fe²⁺ ions, as shown by the signal at 2 mm/s, that constitutes one part of the quadrupole split component associated to these ions (Figure 5D). The late reduction of Fe and the trapping of unreduced Fe ions inside oxidic phases may be the source of further variance of metal proportions within the Ni–Fe particles, compared with the nominal formulation of the catalyst.

3. Ni–Fe Catalysts for the Hydrogenation of Organic Molecules

Since the 1980s, several authors have noted that Ni–Fe bimetallic catalysts are interesting candidates for the selective hydrogenation of organic molecules, with gains in terms of activity and selectivity compared to Ni catalysts. In this section, we will aim at identifying the main characteristics of Ni–Fe catalysts in terms of catalytic performances, and if specific formulations have consistently provided promising results.

3.1. Hydrogenation of Double and Triple Carbon–Carbon Bonds

One of the first papers in which Ni–Fe catalysts have been tested for the hydrogenation of organic molecules concerns the hydrogenation of ethylene to ethane. Matsuyama et al. compared three families of unsupported Ni-based alloys, Ni–Cu, Ni–Fe and Ni–Co [20]. At 0 °C, Ni–Fe alloys from the fcc domain, reduced at 385 °C, provided the highest normalized rate of reaction: 1 min⁻¹m⁻² for the Ni₅₀Fe₅₀ and Ni₆₀Fe₄₀ bulk formulations, three times more than Fe-rich bcc phases and monometallic Ni (Figure 6A). We will come back to this seminal paper and to Figure 6A in Section 4.4, as they give an important key to understand the reactivity of the Ni–Fe system.

The hydrogenation of propyne was carried out by Kojima et al. on unsupported Ni–Fe alloys prepared by the reduction of a mixed hydroxide obtained by co-precipitation from an ethylene glycol solution [49]. Ni–Fe catalysts were activated under H₂ at 400 °C before reaction. The catalytic performances were measured between room temperature and 160 °C. Ni₅₀Fe₅₀ alloys appeared as less active than monometallic Ni (the reaction rate normalized per m² decreased by a factor 10), but a dramatic decrease of selectivity to propane, and correlated increase of selectivity to propene, were noted, even at full conversion (Figure 6B). Decreasing the Ni content led to a slight decrease of the normalized reaction rate, but the selectivity to the alkene, propene, was maintained above 60% over the whole temperature range. Reaction orders did not change compared with Ni, but the activation energy was seen to decrease from 49 kJ mol⁻¹ for Ni, to 36 kJ mol⁻¹ for Ni₅₀Fe₅₀. The increase to 42 kJ mol⁻¹ noted for Ni₃₃Fe₆₇ can be attributed to the presence of bcc Fe-rich and fcc Ni-rich particles for this formulation, the only one that falls in the de-mixing domain (Section 2.1).

A quantitative hydrogenation of the alkene or alkyne function in styrene and phenylacetylene to ethylbenzene has been obtained by Chieffi et al. at 50 °C in ethanol, using a 56 wt% Ni₅₀Fe₅₀/carbon catalyst prepared by impregnation of cellulose and carbothermal reduction (alloy particle size: 20–30 nm, XRD, TEM) [34]. No deactivation was observed over a period of 80 h. This was attributed to the protection of the metal particles by carbon layers.
Figure 6. (A) Normalized rate of hydrogenation of ethylene as a function of the bulk composition (circles, dotted line) and of the surface composition (circles, full line) of Ni–Fe alloys; solubility of hydrogen in the alloys (triangles). Reprinted from Reference [20], copyright 1986, with permission from Elsevier; (B) conversion and selectivity to propane of (a) Ni and (b) a Ni60Fe40 alloy, in the hydrogenation of propyne. Reprinted from Reference [49].

Finally, using 56 wt% Ni80Fe20/carbon and 25 wt% Ni80Fe20/CeO2 catalysts, respectively, a high activity was achieved for the selective hydrogenation of C=C bonds from two more complex molecules, α-terpinene [34] and 2-cyclohexen-1-one [26] at a respective reaction temperature of 125 °C (conversion of α-terpinene: 99%, selectivity to 1-isopropyl-4-methylcyclohexene: 56%) and 130 °C (conversion of 2-cyclohexen-1-one: 100%, selectivity to cyclohexanone: 98%).

3.2. Selective Hydrogenation of Oxygenates

The largest range of organic molecules tested for selective hydrogenation on Ni–Fe catalysts is the family of oxygenates: Ketones, aldehydes, acids, esters, monosaccharides. Given their specificities, furanic compounds will be presented separately in Section 3.6.

Table 1 summarizes the results obtained by Chieffi et al. [34], and Putro et al. [26], on a variety of oxygenates, including the reaction conditions, conversions and selectivities. The characteristics of the Ni–Fe catalyst used by Chieffi et al. have been presented above (Section 3.1). Putro et al. prepared both unsupported and supported Ni–Fe catalysts by co-precipitation followed by a hydrothermal treatment and reduction between 300 and 500 °C. XRD results were interpreted on the basis of the formation of both ordered Ni3Fe and Ni nanoparticles (average crystallite size for unsupported Ni67Fe33: 16 nm, XRD; particle size on TiO2-supported systems: 18−22 nm, TEM). Support effects will be detailed below. Only one Ni–Fe formulation was tested in each work.

Table 1 shows that while the reduction of aromatic aldehydes to aromatic alcohols is quantitative, the results in the hydrogenation of unsaturated carboxyls are contrasted and seem to depend on the relative position of the C=C and C=O bonds. Conversion and selectivity to the unsaturated alcohols are often high for molecules in which the C=C and C=O bonds are not close to each other, and they are significantly lower for α,β-unsaturated aldehydes or ketones for which the hydrogenation of the C=C bond is sometimes favored.

Chieffi et al. have also explored the hydrogenation of levulinic acid to γ-valerolactone, and of monosaccharides (glucose, xylose) to the corresponding polyols (sorbitol, xylitol), with in this case a quantitative conversion in water, for glucose concentrations comprised of between 0.1 and 0.5 mol L\(^{-1}\). This result confirms the early findings by Gallezot et al. on Fe-promoted Raney Ni (six-fold increase of initial reaction rate when comparing Ni80Fe20 to Ni catalysts), which were troubled though by the leaching of Fe into the 2.3 mol L\(^{-1}\) aqueous solution of glucose upon successive runs [50]. Increasing the concentration to 1 mol L\(^{-1}\) was indicated by Chieffi et al. to reduce the sorbitol yield to 40%.
Table 1. Hydrogenation of oxygenates using a 56 wt% Ni₈Fe₄/carbon catalyst prepared by carbothermal reduction [34] or a 25 wt% Ni₈Fe₃/TiO₂ catalyst reduced at 300 °C [26]: Experimental conditions, conversion (Conv.) and selectivity (S) in the mentioned product.

| Reference | Substrate | Product | P(H₂) (bar) | T (°C) | Solvent | Conv. (%) | S (%) |
|-----------|-----------|---------|-------------|--------|---------|-----------|-------|
| [34]      |            |         | 10          | 150    | EtOH    | >99       | 84    |
| [26]      |            |         | 10          | 150    | iPrOH   | 100       | 99    |
| [26]      |            |         | 10          | 150    | iPrOH   | 100       | 87    |
| [26]      |            |         | 10          | 150    | iPrOH   | 98        | 98    |
| [26]      |            |         | 10          | 150    | iPrOH   | 92        | 95    |
| [26]      |            |         | 10          | 150    | iPrOH   | 99        | 95    |
| [26]      |            |         | 10          | 150    | iPrOH   | 95        | 92    |
| [26]      |            |         | 10          | 150    | iPrOH   | 56        | 87    |
| [26]      |            |         | 10          | 150    | iPrOH   | 50        | 95    |
| [34]      |            |         | 50          | 150    | EtOH    | 84        | 99    |
| [34]      |            |         | 10          | 100    | EtOH    | >99       | >99   |
| [26]      |            |         | 10          | 150    | iPrOH   | 4         | 44    |
| [26]      |            |         | 10          | 150    | iPrOH   | 4         | 41    |
| [34]      |            |         | 50          | 150    | EtOH    | 99        | 60    |
| [34]      |            |         | 50          | 150    | H₂O     | >99       | >99   |
| [34]      |            |         | 50          | 150    | H₂O     | >99       | >99   |
The role of the oxidic support (Al₂O₃, TiO₂, CeO₂) on the hydrogenation of 2-cyclohexen-1-one as investigated by Putro et al. cannot be analyzed simply, as selectivities have been compared at very different conversions [26]. The least active catalysts, supported on Al₂O₃ and TiO₂, would favor the hydrogenation to 2-cyclohexen-1-ol at low conversion (hydrogenation of the ketone function), while the most active catalysts, unsupported or supported on CeO₂, would favor the formation of cyclohexanone at high conversion (hydrogenation of the C=C bond).

The influence of the Ni–Fe proportions on catalytic performances were reported in the first paper published on the application of Ni–Fe catalysts to the hydrogenation of oxygenated functions. Kumbhar et al. found that the highest activity for the hydrogenation of acetophenone to α-phenylethanol in methanol at 135 °C was obtained with a 20 wt% Ni₇₅Fe₂₅/SiO₂ catalyst, prepared by co-impregnation and reduced at 450 °C (particle size determined by the Scherrer equation: 13 nm) (Figure 7a) [17]. However, the comparison with Ni/SiO₂ is ambiguous, as the Ni catalyst presents larger Ni nanoparticles, and the ratio between the reaction rates is similar to the ratio between the two metal dispersions. The lack of activity of the other Ni–Fe catalysts is also difficult to comment, because unlike Ni₇₅Fe₂₅/SiO₂, the catalysts bear a mixture of bcc and fcc particles. Hydrogenolysis to ethylbenzene is mentioned to occur above 70% of conversion whatever the catalyst formulation.

The hydrogenation of acids and esters has been described in a few papers. Stearic acid was selectively hydrogenated into stearic alcohol in dioxane at 250 °C on 67–88 wt% Ni–Fe/carbon catalysts prepared by co-precipitation of metal oxalates on melamine followed by carbothermal reduction at 800 °C under N₂ (particle size: 38 nm, TEM) [27]. The nickel content was kept constant while the Fe content was varied. The ordered Ni₇₅Fe₂₅ phase, possibly protected by carbon layers, was identified by XRD for the Ni₇₅Fe₂₅/carbon nominal formulation, and by STEM–EDX, though with some variance on chemical composition, for two Fe-richer catalysts, and was associated to a full conversion. Monometallic Ni rather favored several secondary reactions of coupling: Esterification between stearic acid and stearic alcohol, formation of stearone by ketonization and decarboxylation of two molecules of stearic acid. Similarly, a 16 wt% Ni₆₇Fe₃₃/SiO₂ catalyst prepared by co-impregnation and reduced at 450 °C (average alloy particle size: 5 nm, TEM) provided a 50% selectivity to dodecanol in the hydroconversion of methyl laurate at 300 °C in the gas phase, while in the same conditions, an 8 wt% Ni/SiO₂ catalyst mostly yielded hydrogenolysis products [46].
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Figure 8. Influence of reaction time on the conversion of ethyl levulinate (EL) to ethyl 4-hydroxy pentanoate (4-HPE) and γ-valerolactone (GVL), using a 10 wt% Ni$_{67}$Fe$_{33}$/carbon catalyst reduced at 400 °C (T = 100 °C, P(H$_2$) = 40 bar, solvent: Water). Reprinted from Reference [42], copyright 2017, with permission from Elsevier.

The optimized hydrogenation of ethyl levulinate (conversion: 100%, selectivity to γ-valerolactone (GVL): 99%) was achieved at 100 °C in water on a 10 wt% Ni$_{67}$Fe$_{33}$/carbon catalyst prepared by co-precipitation on activated carbon and reduced at 400 °C (alloy particle size: 6–10 nm, average size 7.1 nm, XRD, TEM) [42]. The hydrogenation of the ketone function to a secondary alcohol function was followed by an intramolecular transesterification reaction, with release of an ethanol molecule (Figure 8). With the increase of Fe content at constant Ni content, the conversion and yield increased and reached a maximum for Ni$_{67}$Fe$_{33}$. A further increase led to a decrease of conversion and yield. A 7 wt% Ni/carbon catalyst provided a 51.7% yield of γ-valerolactone at a conversion of 53.5%; the nature of the side-products was not given. A stability test was carried out, aided by the easy magnetic separation of the Ni$_{67}$Fe$_{33}$/carbon catalyst. Conversion and selectivity to GVL decreased along the four successive runs. A limited recovery was obtained by regeneration by calcination, and a loss of 3.46% of Ni and 6.78% of Fe was detected. The lack of stability of the catalyst was thus attributed both to the deposition of carbonaceous species and to the leaching of metals in water.

3.3. Selective Hydrogenation of Nitriles

The hydrogenation of the C≡N bond in nitriles, such as benzonitrile, butyronitrile, cinnamionitrile and crotononitrile, was performed at 150 °C in methanol [17,51]. The 20 wt% Ni$_{56}$Fe$_{42}$/SiO$_2$ catalyst presented in Section 3.2 for the hydrogenation of acetophenone was found to represent the optimum formulation (Figure 7b). The articles mention that primary amines were mainly formed along with some secondary amines, and that only saturated nitriles and amines were obtained, but the products were not named. In contrast, it is the secondary amine, N,N-dibenzylamine, that Chieffi et al. reported to be the main product of hydrogenation of benzonitrile at 150 °C in ethanol, using the 56 wt% Ni$_{56}$Fe$_{42}$/carbon catalyst formerly seen in the sections above (conversion: >99%, selectivity: 40%) [34].
3.4. Selective Hydrogenation of Nitroaromatics

The selective hydrogenation of nitroaromatics to the corresponding aromatic amines has been reported in three occurrences. Chieffi et al. have carried out the quantitative hydrogenation of nitrobenzene to aniline at 100 °C in ethanol with 56 wt% Ni$_{58}$Fe$_{42}$/carbon (see Section 3.1) [34]. Ni–Fe/N-containing carbon catalysts (38–60 wt%) (constant Fe content, Ni content varying between Ni/Fe ratios = 2 to 4) were prepared by co-precipitation, contact with melamine and carbothermal reduction under N$_2$ at 600 °C [40]. The particle size determined by XRD and TEM was in the 18–26 nm range, and the homogeneity of the particle composition was verified by STEM–EDX. A quantitative reduction of ortho-chloronitrobenzene to ortho-chloroaniline was achieved at 80 °C in ethanol for a Ni$_{75}$Fe$_{25}$/carbon catalyst. It must be noted though that the catalyst mass was kept constant for the kinetic measurements involving the different catalysts, and that the Ni$_{75}$Fe$_{25}$/N-containing carbon catalyst was the one possessing the highest loading of metals. The other formulations were less active, but almost as selective as Ni$_{75}$Fe$_{25}$, including the analogue monometallic Ni catalyst. Reported conversion and selectivities for other nitroaromatics were similarly elevated.

Ni–Fe nanoalloys prepared by carbothermal reduction in N$_2$ at 800 °C on coal have finally been tested in the hydrogenation of para-nitrophenol by NaBH$_4$ in water [52]. A 6.5 wt% Ni$_{45}$Fe$_{55}$/carbon catalyst appeared as 5.4 times more active than the monometallic Ni analogue system. Interpretation is however difficult as both Ni and Fe nanoparticles were detected by XRD on the bimetallic catalyst.

3.5. Hydroconversion at Higher Temperatures: The Case of Aromatic Rings

At high temperatures (300–400 °C) and in the gas phase, hydrogenolysis reactions start to predominate. Unlike Ni catalysts, Ni–Fe systems have been shown to prevent the concomitant hydrogenation of aromatic rings. A first hint was given by Guerrero-Ruiz et al. as early as 1992. They showed that the activation energy of a 10 wt% Ni$_{39}$Fe$_{61}$/carbon catalyst was 69 kJ mol$^{-1}$ for benzene hydrogenation, compared with 55 kJ mol$^{-1}$ for 5 wt% Ni/carbon [30,31]. The Ni–Fe catalysts were prepared by successive impregnations, which after reduction at 400 °C led to the co-existence of Fe and alloyed particles (2–6 nm) as evidenced by $^{57}$Fe Mössbauer spectroscopy and TEM. As a Fe/carbon catalyst proved totally inactive in benzene hydrogenation, the increase of activation energy was assigned to the bimetallic nanoparticles.

Leng et al. showed that 20 wt% Ni$_{75}$Fe$_{25}$/Al$_2$O$_3$ catalysts prepared by co-impregnation and reduced at 400 °C (average particle size: 4.8 nm, TEM) provided an almost quantitative hydrodeoxygenation (HDO) of furfuryl alcohol, benzyl alcohol and ethyl octanoate to, respectively, 2-methylfuran, toluene and heptane, at atmospheric pressure and at T = 400 °C [53]. Fang et al. studied the HDO of guaiacol at 300 °C over Ni–Fe catalysts supported on carbon nanotubes (CNT), prepared by co-impregnation and reduced at 400 °C (particle size: 5–8 nm, XRD, TEM, alloying evidenced by XRD and STEM–EDX) [54]. Monometallic 7 wt% Ni/CNT exhibited a 79% conversion and 60% selectivity to cyclohexane, the product of full HDO and hydrogenation (Figure 9). An 83.4% selectivity to cyclohexane at 96.8% of conversion was obtained with 7 wt% Ni$_{83}$Fe$_{17}$/CNT. But when increasing the Fe content, the hydrogenation of the aromatic ring was toned down, as conversion decreased and selectivity was shifted to the production of phenol, such as with 7 wt% Ni$_{7}$Fe$_{93}$/CNT (conversion of guaiacol: 50%, selectivity to phenol: 77%). The Ni–Fe nanoparticle size and the reduction temperature were also investigated. Increasing the reduction temperature from 400 to 600 °C increased the size of Ni$_{7}$Fe$_{93}$/CNT nanoparticles from 7.7 to 11.2 nm, leading to a decrease of the selectivity to cyclohexane from 85.4% to 55.8%, and an increase of the selectivity to cyclohexanol from 12.7% to 31.9% at full conversion. No phenol was produced. This indicates that the preservation of the aromatic ring is heavily linked to the Fe content of the nanoparticles.
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Figure 9. Catalytic performances of Ni–Fe/carbon nanotube catalysts in the hydrodeoxygenation of guaiacol in the gas phase, as a function of the Ni and Fe molar proportions (T = 300 °C, P(H2) = 30 bar). Conversion: Triangles, selectivities: Bars. Adapted from Reference [32], copyright 2017, with permission from Elsevier.

The effect of Fe has also been surveyed by Nie et al. for the HDO of meta-cresol at 300 °C, and similar tendencies were found [22]. The predominant product for a 15 wt% Ni/SiO2 catalyst was the product of hydrogenation and tautomerization, 3-methylcyclohexanone (conversion: 16.2%, selectivity: 33.3%). Instead, for a 15 wt% Ni50Fe50/SiO2 catalyst prepared by co-impregnation and reduced at 450 °C (average particle size: 10 nm, XRD), the predominant product at a conversion of 13.7% was toluene (selectivity: 52.6%). Toluene was the only product detected with a Ni50Fe50/SiO2 catalyst that contained both alloyed and Fe-rich nanoparticles (XRD). Density functional theory (DFT) studies and microkinetic modelling suggest that the aromatic ring is preserved from hydrogenation on Ni–Fe catalysts, which ensure the direct hydrogenolysis of the C–O bond, whereas Ni would rather transform the unsaturated ketone tautomer of meta-cresol [54]. A more complex route to toluene, involving the dehydration of the unsaturated alcohol resulting from the selective hydrogenation of the ketone tautomer, had been formerly postulated to explain the selectivity of the Ni–Fe catalysts [22].

3.6. Hydrogenation of Furanic Compounds

Given their specific reactivity and the multiplicity of transformations to which this family of molecules can be subjected, results obtained in the hydroconversion of furfural and 5-hydroxymethyl furfural will be presented separately in this section.

Furfural holds an aromatic furan ring, which can undergo hydrogenation and ring-opening reactions, and a side-group containing an aldehyde function –CH=O, which can be hydrogenated, hydrogenolyzed, or removed by decarbonylation (Figure 10) [55]. The selective hydrogenation of the aldehyde group of furfural leads to furfuryl alcohol (noted by FOL below), an intermediate in the production of fine chemicals and synthetic fibers. Tetrahydrofurfuryl alcohol (THFOL), a key intermediate for the synthesis of vitamin C, plasticizers, lubricants and lysine, is produced by hydrogenation of the furan ring of FOL. The hydrogenolysis of the C–O bond of the alcohols yields fuel additives, 2-methyl-furan (MF), from FOL, or 2-methyltetrahydrofuran (MTHF), from THFOL (the hydrogenation of the ring of MF being another possible route to MTHF). In a competing route, decarbonylation of furfural leads to furan, which can be hydrogenated into tetrahydrofuran (THF).
Ni–Fe bimetallic catalysts prepared by co-impregnation and reduced at 450 °C (average particle size: 10 nm, XRD; 16–17 nm, TEM) have been explored by Sitthisa et al. for the hydroconversion of furfural in the gas phase between 210 and 250 °C (Figure 11) [21]. Furan, the decarbonylation product, was the main product of hydroconversion over a 5 wt% Ni/SiO₂ catalyst at 250 °C (conversion of furfural: 98%, yield of furan: 50%). On Ni–Fe bimetallic catalysts, a high conversion was also obtained (96.3% for 7 wt% Ni₇/Fe₂₉/SiO₂), but the yield of 2-methylfuran greatly increased (from 5% at 250 °C for 7 wt% Ni/SiO₂ to 39.1% for 7 wt% Ni₇/Fe₂₉/SiO₂) while the yields of furan decreased (from 50% for 7 wt% Ni/SiO₂ to 12.1% for 7 wt% Ni₇/Fe₂₉/SiO₂). The yield of FOL was comprised between 22% and 32% at 210 °C both for the Ni and Ni–Fe catalysts, but became low at 250 °C. At that temperature, 7 wt% Ni/SiO₂ did not yield any FOL, while a yield of 10% was obtained for 7 wt% Ni₇/Fe₂₉/SiO₂. It decreased when further increasing the Fe content (conversion of furfural for 7 wt% Ni₇/Fe₅₀/SiO₂ <30%, yield of FOL: 6%). C₄ and C₅ ring-opening products were detected in large amounts for the Ni and the Ni–Fe catalysts in these experimental conditions. In summary, the addition of Fe suppressed the decarbonylation properties of Ni, while promoting both C=O hydrogenation and C–O hydrogenolysis reactions. Complementary experiments carried out with FOL and benzyl alcohol.
confirmed that Ni–Fe catalysts are more selective catalysts than Ni for the hydrogenolysis of the C–O bond of alcohols. These trends have been recently confirmed by Wang et al. (10 wt% Ni50Fe50/carbon catalysts prepared by co-impregnation and reduced at 500 °C, large distribution of particle sizes, average value: 12 nm, TEM) [56].

In the liquid phase (2-propanol) at 150 °C, Putro et al. obtained a 95% furfural conversion with 97% selectivity to FOL using a 20 wt% Ni50Fe50/Al2O3 catalyst (preparation and characteristics described in Section 3.2) [26]. The higher dispersion of Ni50Fe50 nanoparticles on CeO2, Al2O3 and TiO2 supports enhanced the catalyst intrinsic activity (conversion: 99%, 95% and 100%, respectively) and selectivity (93%, 97% and 91%, respectively), compared to unsupported Ni50Fe50 and Ni (respective conversion: 90% and 100%, and selectivity: 92% and 7%). The main product formed with Ni was THFOL, the product of hydrogenation of the furan ring (selectivity: 61%). The values were also much higher than those measured on SiO2, taeniolite or hydrotalcite supports (conversion: 21%, 22% and 42%; selectivities: 95%, 77% and 90%) which all increased the selectivity to THFOL (5%, 23% and 10%).

Chieffi et al. also found that in ethanol at 150 °C, Ni58Fe42 particles supported on carbon (described in Section 3.1) exhibited a high conversion of furfural (99%) and selectivity to FOL (90%) [34]. The same tendency was evidenced by Shi et al. [28]. A 52 wt% Ni9 Fe59/SiO2 catalyst prepared by DP with urea and reduced at 700 °C (particle size: 4–7 nm, TEM) appeared to be more selective to FOL in 2-propanol at 150 °C (yield >80% at full conversion) than the homologue Ni/SiO2 catalyst that favored the hydrogenation of the furan ring to THFOL at lower furfural concentrations, and the hydrogenolysis to MF, and hydrogenolysis/hydrogenation of the furan ring to MTHF at higher concentrations. Side-reactions of etherification between the furanic alcohols and the solvent were also evidenced. These reactions appeared as prominent with a Ni50Fe50/SiO2 catalyst that was incompletely reduced.

Using heptane as a solvent, Halilu et al. showed that at 250 °C, when increasing the Ni content from 0.05 wt% to 0.5 wt% in a magnetic Ni–Fe2O3–SiO2 nanocatalyst (Ni2+ ions loaded on the surface of a Fe2O3–SiO2 core–shell structure, reduced at 500 °C), the conversion of furfural increased from 55% to 94% while the selectivity to FOL was kept at ~100% in the whole series [57]. With 0.5 wt% Ni/SiO2, the conversion of furfural was only 50%, the selectivity to FOL 25%, and a wide range of products was obtained due to decarbonylation and ring opening reactions, such as furan (selectivity: 48%), butanal (selectivity: 13%), butanol (selectivity: 3%) and butane (selectivity: 11%).

The selective hydrogenation and hydrogenolysis of a molecule close to furfural, 5-hydroxymethylfurfural, were investigated by Yu et al. using carbon nanotube-supported Ni–Fe catalysts (Ni–Fe/CNT) prepared by co-impregnation and reduced at 400 °C (average alloy particle size: 6.5 nm, TEM, STEM–EDX) [58]. Two side-groups are held by 5-hydroxymethylfurfural: An aldehyde function like furfural, and a –CH2OH function (Figure 12a). Monometallic 10 wt% Ni/CNT catalyst displayed a high conversion at 110 °C in butanol (100%), but a selectivity to the diol, 2,5-furandimethanol, of only 76.4%. The byproducts of the reaction included decarbonylation (5-methylfurfural), ring opening (1,2-hexanediol), etherification products (grouped as “others” on Figure 12b), and hydrogenolysis and ring hydrogenation products (5-methyl-2-furanmethanol, 2,5-dimethylfuran, 2,5-dimethyltetrahydrofuran). Even a small addition of Fe to Ni increased the selectivity to 2,5-furandimethanol (87.7% for 10 wt% Ni5Fe5/CNT) while keeping the conversion at 100%. A selectivity of 97.5% to 2,5-furandimethanol at 100% of conversion could be achieved with 10 wt% Ni5Fe5/CNT. Chieffi et al. also found that in ethanol at 150 °C, Ni50Fe50/carbon supplied a high conversion of 5-hydroxymethylfurfural (99%) and a high selectivity to 2,5-furandimethanol (88%) [34].
Figure 12. (a) Hydroconversion of 5-hydroxymethylfurfural: Reaction pathways favored by a Ni/carbon nanotubes catalyst and by a Ni<sub>67</sub>Fe<sub>33</sub>/carbon nanotube catalyst (FDM: 2,5-furandimethanol, MF: 5-methyl furfural, MFM: 5-methyl 2-furanmethanol, DMF: 2,5-dimethylfuran, DMTHF: 2,5-dimethyl tetrahydrofuran, HD: 1,2-hexanediol). Reprinted from Reference [58], copyright 2015, with permission from Wiley; (b) selectivities obtained as a function of the Ni molar proportion (T = 110 °C, P(H<sub>2</sub>) = 30 bar, solvent: n-butanol; others: Ethers, decarbonylation product, ring-opening products, humin). Based on data from Reference [58].

The products distribution was completely changed to hydrogenolysis products when the reaction temperature was increased to 200 °C. The main product was then 2,5-dimethylfuran (selectivity: 46.3%) for monometallic Ni/CNT catalyst at 100% conversion (selectivity to 2,5-furandimethanol: 13.4%). After increasing the Fe content up to Ni<sub>67</sub>Fe<sub>33</sub>/CNT, the selectivity to 2,5-dimethylfuran was 91.3% at 100% conversion and no 2,5-furandimethanol was formed. Fe thus appeared to increase the catalyst selectivity associated to the aldehyde and alcohol side-groups, both toward selective hydrogenation products at lower temperature, and to hydrogenolysis products at higher temperature, while preventing reactions involving the furan ring. A stability test over seven runs showed that the full conversion was retained while the selectivity to 2,5-dimethylfuran (DMF) decreased from 90% to 75%. Deactivation was ascribed to the deposition of carbonaceous molecules, rather than to metal leaching or sintering.

This survey from the literature clearly shows that whatever the support, fcc Ni–Fe phases in the range of formulation Ni<sub>x</sub>Fe<sub>y</sub>–Ni<sub>50</sub>Fe<sub>50</sub>–Ni<sub>80</sub>Fe<sub>20</sub> often provide the best results in terms of activity and selectivity. The Ni<sub>x</sub>Fe<sub>y</sub> formulation is also the one that has become favored for CO and CO<sub>2</sub> hydrogenation [9,10,14,15,41,59,60], while the Ni<sub>x</sub>Fe<sub>y</sub> formulation was found to be the most active for dry and steam reforming [3,8]. It must be noted though that comparison between different catalysts, or between different works, should be done with caution for several reasons:

- The nature of the Ni-Fe nanoparticles on the catalyst is not always homogeneous (mixture of bcc and fcc phases, of particles with different compositions).
- The temperature of reduction is usually lower than 500 °C: There is a, sometimes demonstrated, risk that Fe, and even Ni, reduction is not complete after activation in H<sub>2</sub> (X-ray photoelectron spectroscopy (XPS) measurements [32,42]; FTIR measurements [22]).
- SiO<sub>2</sub>, and various types of carbon, have been used as supports in most of the articles, but specific effects involving supports such as Al<sub>2</sub>O<sub>3</sub> and TiO<sub>2</sub> have been reported.
- Even for a given support and for a similar procedure of preparation (co-impregnation or precipitation), discrepancies are large between the different articles in terms of particle size. Some size-dependent effects on catalytic performances have however been reported [26,32].
- Selectivities and yields may be given only at high or full conversion, which does not allow a clear comparison between different systems, and makes discussion difficult. A kinetic study carried out as a function of reaction time, or at increasing conversion, is often lacking.
• A limited number of Ni–Fe formulations is usually tested.
• In some cases, the metals loading varies along with the nanoparticle formulation (when the content in one of the metals is kept constant while the other one increases), while the mass of catalyst used for catalytic test remains the same, possibly skewing comparisons of activities.

Nevertheless, several tendencies emerge when Ni–Fe catalysts are compared with Ni catalysts:
• The hydrogenation of C=C, C=O and C≡N bonds has been reported to be faster for the bimetallic formulations.
• Ni–Fe catalysts are more selective than Ni for the hydrogenation of C=C and C=O bonds, to alkenes vs. alkanes, and to alcohols vs. secondary products, respectively. In particular, the hydrogenation of aromatic rings, and the opening of furanic rings, are often toned down on Ni–Fe systems. At higher temperatures, the hydrogenolysis of C–O bonds is avoided on optimal formulations.
• The result of the hydrogenation of a molecule that contains both a C=C and a C=O bond depends on the relative localization of the two bonds. A distance of several bonds between the two functions seems to result in the preferential hydrogenation of the C=O bond, while the outcome is less clear-cut for α,β-unsaturated carbonyls.

4. The Role of the Two Metals in the Reactivity of Ni–Fe Nanoparticles: Hypotheses and Open Questions

To account for the specificities of Ni–Fe catalysts, four categories of explanations have been proposed so far. They are respectively based on the structure of Ni–Fe particles; on their global electronic properties; on the interaction between adsorbates and the surface of the Ni–Fe nanoparticles; finally, on the existence of an excess of Fe atoms at the surface of the nanoparticles.

4.1. Structural and Geometric Effects

Matsuyama et al. have first proposed that fcc Ni–Fe phases are more active than bcc phases in the hydrogenation of ethylene because of a higher solubility of hydrogen in their core [20]. However, crystal structure and chemical composition of the alloys are correlated parameters, and it is not possible to consider them separately.

Kojima et al. have interpreted the reactivity of propyne on Ni–Fe surfaces as the result of a simple balance between Ni domains and Ni–Fe ensembles, directly derived from the molar composition of the bulk alloy [49]. Fewer Ni domains would lead to less available hydrogen and to a catalyst that limits the hydrogenation of the alkyne to the alkene. Kumbahr et al. and Yadav et al. also recognized the higher activity of fcc phases in the hydrogenation of acetophenone and benzonitrile. They would present the right balance between Ni domains active in the chemisorption of H₂, and Ni–Fe ensembles presumably involved in the activation of the organic substrate [17,51].

4.2. Electronic Effects

Several teams have gone beyond mere geometric effects, and have attributed the performances of Ni–Fe systems to global electronic properties. Ray et al. have associated the activity of Ni₇₅Fe₂₅ surfaces in CO₂ methanation to a minimal number of Ni d density of states at the Fermi level [61,62]. In contrast, Yu and Chen have demonstrated that covering Ni with a monolayer of Fe atoms would lead to an increased density of states at the Fermi level, that would explain the stronger binding energy of adsorbates on Fe/Ni(111) compared with Ni(111) [63].
Figure 13. Right: (A) Correlation between (a) the activation energy in CO hydrogenation ($E_a$), (b) the measured activity in CO hydrogenation, and the dissociation energy of CO evaluated by DFT ($E_{\text{diss}}$), evidencing the close properties of the Ni$_{50}$Fe$_{50}$ and Ni$_{75}$Fe$_{25}$ surfaces with respect to Co and Ru. Reprinted from Reference [9], copyright 2006, with permission from Elsevier; Left: Measured reaction rates in the hydrogenation of CO at 225 °C for (B) Ni–Fe/MgAl$_2$O$_4$ and (C) Ni–Fe/Al$_2$O$_3$ catalysts. Reprinted from Reference [10], copyright 2007, with permission from Elsevier.

Actually, the adsorption of NO on Ni–Fe nanoparticles followed by infrared spectroscopy revealed a high proportion of bent-type “NO” species ($1820$–$1827$ cm$^{-1}$), with respect to linear-type “NO” species ($1874$–$1878$ cm$^{-1}$), characteristic of an increase of electron donation toward the anti-bonding orbitals of the adsorbates, with a maximum found for the Ni$_{50}$Fe$_{50}$ composition [64]. It should be noted though that the linear type, and the bent type, could not be observed on monometallic catalysts containing Fe, and Ni, respectively, and that this interpretation does not necessarily reveal synergistic effects between the two metals.

The existence of a stronger back-donation toward the anti-bonding orbitals of the adsorbate is in line with the findings from Andersson et al., who correlated the superior activity of Ni$_{50}$Fe$_{50}$ and Ni$_{75}$Fe$_{25}$ alloy surfaces in the hydrogenation of CO to the dissociation energy of CO ($E_{\text{diss}}$) used as a relevant descriptor (Figure 13A) [9]. Compared with the optimal value of $E_{\text{diss\;optimal}} = 0.06$ eV determined by DFT for Co and Ru, the calculated gap $\Delta E_{\text{diss}} = E_{\text{diss}} - E_{\text{diss\;optimal}}$ for Ni$_{50}$Fe$_{50}$ and Ni$_{75}$Fe$_{25}$ was calculated to be lower than 0.25 eV, while it was significantly higher for Fe (0.9 eV) and Ni (0.4 eV). The relevance of this descriptor for CO hydrogenation was experimentally confirmed by Hwang et al. using temperature-programmed surface reactions [12,13]. Experimental measurements in the hydrogenation of CO confirm the superior performances of the Ni-rich alloyed phases for different supports and metal loadings (Figure 13B and C) [10]. We should note though that these correlations remain global, and are not based on the formation and reactivity of intermediates, or on kinetic parameters.

4.3. Specificity of the Interactions between Ni–Fe Surfaces and Adsorbates

The explanations presented above omit the description of surface active sites, and do not provide a precise definition of the role of the two metals in the adsorption of the substrate to hydrogenate. In
this respect, several interpretations concur in showing that Ni and Fe actually play different and complementary roles on the alloyed surfaces.

Fe is a more electropositive metal compared to Ni. It was supposed by Yadav and Kharkara that a charge transfer from Fe to Ni took place on Ni$_{75}$Fe$_{25}$ particles, with an increase of electron density on Ni that should lead to strengthened metal–carbon bonding, and weakening of adsorbed polar functions (ketone, nitrile) [51].

Fe also displays a stronger affinity to oxygen than Ni. The oxophilicity of Fe with respect to Ni has in fact been listed as a key factor for the performances of Ni–Fe catalysts in the reforming of organic molecules and CO$_2$ hydrogenation. Li et al. have stated that in reforming reactions, Ni atoms act as adsorption sites for the organic molecules, while Fe activates water [4]. Kim et al. have shown by operando characterization that under dry reforming conditions, small de-mixed domains of FeO reversibly form at the surface of the nanoparticles, and enable carbon oxidation and removal [8]. Pandey et al. have considered that oxidized Fe may be beneficial to the hydrogenation of CO$_2$, as it facilitates its adsorption [14,15].

These two properties of Fe can be brought together to explain the specific adsorption mode of polar functions on Ni–Fe surfaces. In the hydrogenation of furfural in the gas phase, the effect of Fe was attributed by Sitthisa et al. to the increase of the stability of $\eta^2$(C, O) surface intermediates, in which the O atom of the aldehyde function interacts with oxophilic Fe atoms [21]. Compared with Ni, a longer C–O bond was found on a Ni$_{50}$Fe$_{50}$ surface, indicating a favored electron back-donation from the metals to the anti-bonding $\pi^*$ orbital of the carbonyl function. In contrast, monometallic Ni interacts primarily with the carbon atom from the aldehyde function, preparing the cleavage of the C–C bond between the furan ring and the side group, and the decarbonylation of furfural to furan.

These observations were corroborated by Yu et al., who used DFT calculations and surface science studies on Fe/Ni(111), alloyed Ni–Fe and Ni(111) surfaces to determine the most favorable reaction pathways involving furfural [65]. Furfural was found to bond onto Fe-rich surfaces more strongly through the carbonyl group than onto Ni surfaces, with a longer C–O bond in the adsorbed molecule and a shorter O–M bond. The furan ring was tilted away from the Ni–Fe surface, preventing its hydrogenation. The same repulsive effect may not be effective for $\alpha,\beta$-unsaturated carbonyls, as the proximity between the C=C and the C=O bonds tends to favor the hydrogenation of the two bonds, or of the C=C bond [26].

A dual role assigned on the one hand to Fe for the optimal adsorption of carboxylic groups and ester functions, and on the other hand to Ni for the adsorption of H$_2$, was postulated by Kong et al. to account for the performances of the Ni$_{75}$Fe$_{25}$ phase in the hydrogenation of stearic acid [27], and by Yu et al. for the hydroconversion of methyl laurate [46]. Using propanoic acid as a model reactant (Figure 14), Kong et al. have demonstrated that the kinetically-favored pathway is the adsorption and activation of the acid via the carboxylic group on a Ni–Fe pair, followed by its deoxygenation to a CH$_3$–CH$_2$–CO$^*$ intermediate which finally undergoes hydrogenation to propanol.

Besides, the reaction order with respect to H$_2$ in the hydrogenation of stearic acid was measured to increase from 0.6 on a Ni catalyst to 1.6 on the bimetallic phase, indicating that Ni ensembles on Ni–Fe nanoparticles present specificities with respect to H$_2$ activation. Fang et al. have supposed that the adsorption of guaiacol takes place preferentially at the surface of oxophilic Fe domains, and that Ni would be required for the dissociation of H$_2$; the production of phenol on Fe-rich surfaces was linked to an insufficient amount of available activated hydrogen atoms to allow the hydrogenation of the aromatic ring [32].

While confirming the role of Fe in the adsorption of oxygenates on Ni–Fe catalyst, Li et al. have rather emphasized the action of oxidized Fe species, through their Lewis acidity, instead of metallic Fe atoms [42]. This point should not be neglected, as Fe may appear as ionic species if its reduction is not complete.
4.4. A Surface Excess of Fe on Ni–Fe Nanoparticles?

Various results from the literature tend to show that the surface of Ni–Fe is poorly ordered, and that Fe atoms may be present in a significant excess in the first atomic layers. The DFT models proposed by Sitthisa et al. for a Ni₅₀Fe₅₀ (111) surface [21] were based on alternating rows of Ni and Fe atoms. The later model developed by the same team included the possibility that Fe atoms could be grouped (Figure 15) [22]. Comparison with Ni, Fe or ordered Ni–Fe surfaces showed that it is in fact the presence of patches of Fe atoms that would cause a maximum repulsion between an aromatic ring and the metal, explaining the poor activity of Ni–Fe system in its hydrogenation.

This hypothesis is in line with several characterization results. According to surface science studies carried out by Vasiliev and Gorodetsky [66], the surface of Ni₅₀Fe₅₀ single crystals becomes preferentially terminated by Fe atoms upon heating above 350 °C. Several teams have reported that the extended X-ray absorption fine structure (EXAFS) coordination number of Fe in reduced fcc Ni–Fe nanoparticles prepared by co-impregnation is smaller than that of Ni [17,18,24,65,67]. Moreover, when exposed to oxidizing conditions, Fe was shown be more oxidized than Ni, that retained its metallic state and coordination number as if it were preferentially located in the core of the nanoparticles [18,67]. Mössbauer data supplied by Shi et al. also pointed to the presence of non-magnetic Fe at the surface of nanoparticles prepared by deposition–precipitation [28]. Ishihara et al. observed that the desorption profile of H₂ from a Ni₅₀Fe₅₀/SiO₂ catalyst was close to that of monometallic Fe, and very different from monometallic Ni [33]. On some of their catalysts, Yu et al. observed by STEM–EDX measurements an enrichment in Fe at the periphery of Ni–Fe nanoparticles [46]. Fang et al. have presented a low-energy ion scattering (LEIS) spectrum confirming a surface enrichment in Fe [32].

A direct link to catalysis was indeed given by Matsuyama et al. in 1986 [20]. By quantifying β-rays emitted by ⁶³Ni, they demonstrated that the surface concentration of Ni was much lower than what the bulk composition of unsupported alloys would indicate. The maximum activity in the hydrogenation of ethylene was reached for a bulk composition of 50% Ni, but this corresponded to a surface composition of only 20% Ni vs. 80% Fe (Figure 6A). Also in connection with catalysis, Shen and Dumesic have demonstrated that the deposition of Fe atoms onto the surface of Ni nanoparticles by decomposition of Fe(CO)₅ progressively lowers the activation energy of the catalyst in the methanation of CO, and increases the reaction order with respect to H₂ [68], two characteristics that we have seen before.
5. Conclusions and Perspectives

Compared to monometallic Ni catalysts, catalysts based on Ni–Fe formulations present four characteristics for hydrogenation reactions, both in the gas and in the liquid phase: Higher reaction rates for the hydrogenation of C=C, C=O and C≡N bonds; higher selectivity for the hydrogenation of C≡C bonds or polar functions, with low levels of full hydrogenation, ring-opening or hydrogelenolysis side-reactions; in particular, a demonstrated selectivity for the hydrogenation of polar functions, such as that of oxygenates (aldehydes, ketones, furanic compounds, acids, esters, monosaccharides) to alcohols; inhibited hydrogenation of aromatic rings.

Whatever the support, fcc Ni–Fe alloyed phases in the range of formulation Ni₅₀Fe₅₀–Ni₇₅Fe₂₅ tend to provide the best results in terms of activity and selectivity. Formulations close to Ni₅₀Fe₅₀ have often been singled out for their activity and selectivity, similarly to what has been found for CO/CO₂ hydrogenation or reforming reactions. However, it is currently difficult to firmly conclude that their
specificity simply derives from the existence of ordered phases exhibiting this composition, present in the Ni–Fe phase diagram, because several questions remain open.

Conclusions purely based on the alloy structure and overall electronic properties are insufficient, as surface effects seems to dominate. Geometric effects may explain, at a simple level, the lower availability of H₂ compared to monometallic Ni, and the properties of Ni–Fe in selective hydrogenation. A consensus has been currently reached on the dual role of surface Fe and Ni atoms, based in their different electronegativity and oxophilicity: Stabilization of specific intermediates onto Ni–Fe pairs for polar molecules, strong back-donation into the antibonding orbitals of the adsorbates resulting in a facilitated hydrogenation. Nevertheless, the investigation of transition states, in order to explain why activation energies are lowered even for simple, non-polar molecules like ethylene or propylene, is still necessary. How Ni–Fe catalysts modify the reactivity of hydrogen is also unclear. Microkinetic modelling will certainly be a useful tool, even if modelling is more difficult for liquid-phase reactions, as solvation effects may be important. Given the evidence for an enrichment of Fe atoms at the surface of Ni–Fe systems, models should preferably include patches of Fe atoms, that may be involved in the repulsion of aromatic rings. Globally speaking, more quantitative measurements are needed to precisely characterize surface compositions.

Before comparing catalytic results, it should also be remembered that on Ni–Fe catalysts, the nanoparticles size is not always homogeneous, and sometimes their composition is not homogeneous either. As a synthesis procedure, incipient wetness impregnation is easy to implement, but with some risks in terms of heterogeneities between Ni–Fe nanoparticles. Co-precipitation is currently favored for the preparation of supported Ni–Fe catalysts displaying homogeneous nanoparticles, but high reduction temperatures may be required to complete the reduction of the two metals. The heterogeneity depends on the preparation method, but also on the support used: Alumina might trap metal ions (mostly Fe), and change the proportion of metals in the nanoparticles compared with the expected formulation. Support roles have been demonstrated for some selective hydrogenations, but they are difficult to rationalize as they can be accompanied by changes in the particle size or composition. The role of unreduced Fe ions in the activation of polar functions is also unclear. The thermal history of the catalyst is probably of importance, as it may play both on the reduction degree of the two metals, and on the Fe surface content. Further progress in the understanding of Ni–Fe catalysts in selective hydrogenation will require the maximum details to be taken into account (homogeneity of particles in size and composition, reduction temperature, degree of reduction of the metals) in order to compare catalytic results obtained at similar levels of conversion or in similar experimental conditions, to identify the right balance in terms of Ni vs. Fe surface composition, and to build realistic DFT and kinetic models.

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