From Iron to Copper: The Effect of Transition Metal Catalysts on the Hydrogen Storage Properties of Nanoconfined LiBH$_4$ in a Graphene-Rich N-Doped Matrix

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Abstract: Incipient wetness impregnation was employed to decorate two N-doped graphene-rich matrixes with iron, nickel, cobalt, and copper nanoparticles. The N-doped matrix was wetted with methanol solutions of the corresponding nitrates. After agitation and solvent evaporation, reduction at 800 $^\circ$C over the carbon matrix promoted the formation of nanoparticles. The mass of the metal fraction was limited to 5 wt. % to determine if limited quantities of metallic nanoparticles catalyze the hydrogen capture/release of nanoconfined LiBH$_4$. Isotherms of nitrogen adsorption afforded the textural characterization of the matrixes. Electronic microscopy displayed particles of definite size, evenly distributed on the matrixes, as confirmed by X-ray diffraction. The same techniques assessed the impact of LiBH$_4$ 50 vol. % impregnation on nanoparticle distribution and size. The hydrogen storage properties of these materials were evaluated by differential scanning calorimetry and two cycles of volumetric studies. X-ray diffraction allowed us to follow the evolution of the material after two cycles of hydrogen capture-release. We discuss if limited quantities of coordination metals can improve the hydrogen storage properties of nanoconfined LiBH$_4$, and which critical parameters might restrain the synergies between nanoconfinement and the presence of metal catalysts.

Keywords: nanoconfined LiBH$_4$; mesoporous carbon matrix; coordination metal catalyst

1. Introduction

The ever-growing anthropogenic demand for energy is still primarily quenched with fossil fuels. Inevitably, this appetite will unleash two banes: their finitude and the greenhouse effect. Sustainable energies should supersede this dependence, but their alternative nature complicates their implementation into the existing energy matrix. Hydrogen is considered an ideal energy vector, except for its low density [1]. Solid-state metal storage systems based on metal hydrides benefit from their high gravimetric potential and improved security but may suffer from their slow hydrogen release/capture kinetics [2]. The rise of nanomaterials proposed 1D, 2D, and 3D materials to bypass these limitations [3–5].

Lithium borohydride (LiBH$_4$) is a prime example of the potential and limitations of metal hydrides [6]. LiBH$_4$ presents high theoretical gravimetric (18 wt. %) and volumetric (121 kg m$^{-3}$) hydrogen density [7,8], but its application suffers from the high temperatures (500 $^\circ$C) of hydrogen release and high pressure (100 bars) of uptake [6,9,10]. Nanocoexpression is a promising route to improve the kinetic and thermodynamic properties of LiBH$_4$ [11,12]. Reducing the particle size of this hydride to the nanoscale improves mass transport rates, and the contact/tension of the matrix also alters phase transformations.
of LiBH$_4$ [13–18] and reaction pathways [19,20]. The nanoconfinement of LiBH$_4$ lowers the orthorhombic to hexagonal phase transition temperature and greatly enhances species mobility, the hydride behaving as an ionic liquid from transition temperature (circa 120 °C) [21–23]. Yet, nanoconfinement alone did not solve all the issues of LiBH$_4$ hydrogen storage application: it is still difficult to release all available hydrogen below 300 °C, and reversibility hardly exceeds 60% [6,24,25]. Indeed, the mechanism of pore wetting by LiBH$_4$ during impregnation and the expulsion of LiH during hydrogen release are pivotal to the material’s life cycle [26,27]. The own mass of the matrix also quickly limits LiBH$_4$ specific interest by lowering the material’s hydrogen mass capacity; for example, a matrix presenting 1.0 cm$^3$/g pore-volume filled at 50% suffers a 14 to 3.5 wt. % mass capacity loss. Reducing pore size and pore filling generally lowers the onset of hydrogen release, but unfortunately, it also reduces the available pore volume, thus the mass of LiBH$_4$ (i.e., of H$_2$) per mass of material. Even worse, the reversibility of hydrogen uptake/release may suffer from this strategy [6,28–30]. In a recent study, we demonstrated that the gradual release of hydrogen from nanoconfined LiBH$_4$ can be related to the gradual filling of the pores, the smaller being filled first and releasing hydrogen at a lower temperature [31]. The fast exchange rate of the mobile nanoconfined LiBH$_4$ is probably responsible for the gradual hydrogen release in a matrix constituted of both micro- and mesopores [21,31].

Thus, even if nanoconfined LiBH$_4$ presents much better performance than its bulk counterpart, it might not be enough to attain technological applications. Considering this, the strategies used to improve bulk LiBH$_4$ could help to enhance further the properties of nanoconfined LiBH$_4$ [6,24,25]. Many illustrated how combining carbon nanomaterials and metallic nanoparticles improved hydrogen release/capture. For example, cobalt-decorated carbon nanotubes presented a synergistic effect over MgH$_2$ sorption properties [32], while palladium displayed an improved hydrogen uptake capacity when decorating N-doped graphene [33]. Regarding specifically the nanoconfinement of LiBH$_4$ in a carbon matrix, Ngene et al. decorated a porous matrix with nickel nanoparticles [34], Xian et al. doped a carbon network with carbon nanotubes and TiO$_2$ nanoparticles [35], and Wang et al. presented N-doped carbon nanospheres [36]. Exceptionally, Chen et al. replaced the carbon matrix by a reactive Ni-based porous materials [37]. However, carbon derivatives remain a major focus, and graphene in particular is a widely studied carbon nanomaterial with outstanding properties and wide potential applications that have attracted great interest [38]. For instance, graphene wrapping promoted a synergistical activation of LiBH$_4$ in the presence of catalysts [39,40]. Accordingly, our group proposed to take advantage of this nanomaterial by its sequential inclusion within a carbon matrix [28], its doping with heteroatoms [29], and decoration by coordination metals [30]. The latter advanced materials presented nickel, cobalt, and their mixture, improving hydrogen release and capture, but it was unclear if this was a catalytic or a mass effect. Indeed, we chose to decorate a matrix of relatively wide pores (9 nm) with coordination metals at 5 vol. % of the mesopore volume, 0.75 cm$^3$/g, representing more than 100% of the actual mass of LiBH$_4$ in the 70 vol. % LiBH$_4$ filled material (almost three times the mass of LiBH$_4$ in the 30 vol. % filled material). The disproportion between LiBH$_4$ (0.67 g/cm$^3$) and coordination metal (around 9 g/cm$^3$) densities strongly limits both the technical application of the material (worsening the hydrogen’s mass capacity) and the physicochemical interaction of the species (the metal volume being much lower than that of its hydride counterpart). The ability to disperse smaller metallic particles all over the matrix would improve their interaction with the hydride without compromising the mass capacity of the system. It is crucial to homogeneously decorate the inner parts of the pores (where the activated LiBH$_4$ is), not only the surface of the matrix. Recent theoretical studies highlight the interest of nickel, cobalt, copper, and iron in catalyzing the dehydrogenation of LiBH$_4$ [41], but it is still difficult to simulate the behavior of hundreds to thousands LiBH$_4$ units in a 2–5 nm cavity [42]. Hence, it is unclear if the kinetic effect of a metal dopant over H-desorption can translate from the bulk to the nanoconfined space within a matrix activated by a Lewis base [43].
Here, we present the effects of four coordination metals (Fe, Ni, Co, Cu) decorating at 5 wt. % two N-doped matrixes of small pore diameters (4 and 6 nm) filled at 50 vol. % with LiBH$_4$. We aimed to determine if a catalytic effect of a coordination metal can enhance the reversible hydrogen release properties of nanoconfined LiBH$_4$ in an N-doped matrix. Regarding our previous work, the method proposed here emphasizes reducing the size of the metallic nanoparticles decorating the matrix and improving their homogeneity. We also employed matrixes of smaller pores with different nitrogen doping to determine if synergies enhance both effects. The textural parameters of those materials were determined by isotherms of nitrogen adsorption. The coverage of the matrixes by metallic nanoparticles was estimated by SEM (scanning electron microscopy) with elemental mapping. To evaluate the impact of the material’s life on the crystallographic nature of the nanoparticles, PXRD (powder X-Ray diffraction) was performed before impregnating the matrixes by LiBH$_4$, just after impregnation, and after two cycles of hydrogen release/uptake. The influence of the nanoparticles over the release/uptake of hydrogen from nanoconfined LiBH$_4$ was assessed with DSC (differential scanning calorimetry) and volumetric experiments.

2. Results
2.1. Chemical and Morphological Characterizations of the Hydride-Free Matrixes
2.1.1. Textural Characterization of the Matrixes
- Non-decorated N-doped matrixes

The hydrogen release/uptake properties of nanoconfined LiBH$_4$ are profoundly dependent on the matrix pore-size and its filling value: the smaller the pores and the lower their filling, the lower the temperature needed to release hydrogen [31]. Here, we focused on matrixes of relatively small mesopores (from 4 to 6 nm wide), with a non-negligible proportion of micropores, enriched with graphene and doped with nitrogen [29]. X-ray photoelectron spectroscopy (XPS) of these materials can be found in our previous work. Figure 1 presents the isotherms of nitrogen adsorption of the non-decorated, non-impregnated matrixes. Both matrixes were obtained with distinct concentrations of N-dopant, but employing the same concentrations of resin precursors, affording sharp pore-size distribution according to BJH (inset). Two distinct peaks were obtained depending on the proportion of ethylene–diamine crosslinker in the graphene hydrogel. As previously observed, the material doped with more nitrogen (G2N) presented pores of smaller size (3.8 nm) than the material doped with less nitrogen (6.1 nm, GN), probably because nitrogen acts as a base-catalyst for the reticulation of the precursors [29]. Despite the peaks’ sharpness, a non-negligible proportion of the total pore volume was constituted by micropores (0.16 cm$^3$ in both cases, representing 23% of GN total pore volume, up to 43% of G2N pore volume, see Table 1). As discussed before, the micropores are likely to be filled faster than the mesopores at a given filling value, enhancing the nanoconfined behavior of the impregnated LiBH$_4$. We increased the proportions of graphene (+50 wt. %) and N-dopant (+25 mol. %) relative to our previous work, but it did not markedly affect the definition of the peaks [29].

Table 1. Textural parameters of matrixes GN and G2N with and without decoration by metallic nanoparticles before their impregnation by LiBH$_4$.

| Matrix Type | Metallic NP | $S_{BET}$ | $S_{ext}$ | $S_{micro}$ | $V_{tot}$ | $V_{meso}$ | $V_{micro}$ | $D_{max}$ |
|-------------|-------------|------------|-----------|-------------|-----------|-----------|-----------|-----------|
| GN          | None        | 720        | 330       | 390         | 0.68      | 0.48      | 0.16      | 6.1       |
|             | Fe          | 670        | 300       | 360         | 0.63      | 0.46      | 0.15      | 6.0       |
|             | Co          | 610        | 280       | 330         | 0.60      | 0.45      | 0.14      | 6.1       |
|             | Ni          | 660        | 300       | 370         | 0.63      | 0.45      | 0.15      | 6.1       |
|             | Cu          | 690        | 300       | 390         | 0.65      | 0.42      | 0.16      | 6.1       |
|             | NP *        | 660 ± 30   | 290 ± 10  | 360 ± 20    | 0.63 ± 0.02 | 0.45 ± 0.02 | 0.15 ± 0.01 | 6.1      |
would improve the hydrogen capacity of the material, limit particle size, and determine if a catalytic activity is present. Reducing the relative mass of the metallic element would improve the hydrogen capacity of the material, limit particle size, and determine if a catalytic activity is present.

Table 1. Cont.

| Matrix Type | Metallic NP | \( S_{\text{BET}} \) \(^a\) (m\(^2\)/g) | \( S_{\text{exp}} \) \(^b\) (m\(^2\)/g) | \( S_{\text{micro}} \) \(^c\) (m\(^2\)/g) | \( V_{\text{tot}} \) \(^d\) (cm\(^3\)/g) | \( V_{\text{meso}} \) \(^e\) (cm\(^3\)/g) | \( V_{\text{micro}} \) \(^c\) (cm\(^3\)/g) | \( D_{\text{max}} \) \(^e\) (nm) |
|-------------|-------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|
| G2N         | None        | 580 ± 190               | 390 ± 150               | 0.35 ± 0.01              | 0.17 ± 0.03              | 0.16 ± 0.04              | 3.8                      |
|             | Fe          | 360 ± 210               | 150 ± 30                | 0.31 ± 0.02              | 0.23 ± 0.02              | 0.06 ± 0.01              | 3.8                      |
|             | Co          | 360 ± 180               | 180 ± 90               | 0.31 ± 0.02              | 0.22 ± 0.02              | 0.08 ± 0.01              | 3.8                      |
|             | Ni          | 520 ± 180               | 340 ± 20               | 0.32 ± 0.02              | 0.17 ± 0.02              | 0.14 ± 0.02              | 3.8                      |
|             | Cu          | 500 ± 170               | 330 ± 90               | 0.31 ± 0.01              | 0.16 ± 0.01              | 0.13 ± 0.01              | 3.8                      |
|             | NP \(^*\)  | 430 ± 90               | 180 ± 20               | 250 ± 90                | 0.31 ± 0.01              | 0.20 ± 0.03              | 0.10 ± 0.04              | 3.8                      |

Values determined by \(^a\) BET, \(^b\) BET-t-plot, \(^c\) t-plot, \(^d\) Gurvich, \(^e\) BJH. \(^*\) Average and standard deviation of the parameter obtained from all four metal-decorated matrices.

Figure 1. Nitrogen isotherms of GN (A) and G2N (B), with corresponding pore-size distribution obtained by BJH (inset).

- N-doped matrixes decorated with nanoparticles

The evolution of the textural parameters when decorating the matrixes is presented in Table 1 and in the Supplementary Information (Figure SI 1). Regarding the material with less nitrogen (GN), the limited decrease of the mass-related parameters is attributable to the mass increase due to the addition of 5 wt. % of metal to the matrix (Table 1). Yet, in the case of the material doped with more nitrogen (G2N) and decorated with Fe or Co, an increase of the mesopore volume (+35%) was observed at the expense of the micropore volume (−60%); while pore-size and overall pore volume were mostly unaffected, the specific area of the material (highly dependent on the micropores) declined notably (−40%). Remarkably, the concerned peak at 3.8 nm was always present in our material but usually in limited proportions. Further characterization techniques will step up the specificity of these matrixes (G2N Fe and Co). It is noteworthy that reducing the micropore volume should raise the onset temperature of the material [31].

2.1.2. Characterization of Nanoparticle’s Size and Distribution

In previous studies, we highlighted that the presence of nanoparticles affects the performance of the hydride [30]. Reducing the relative mass of the metallic element would improve the hydrogen capacity of the material, limit particle size, and determine if a catalytic quantity of metal can positively affect the hydride’s degradation temperature. Previously, we observed Ni, Co, and NiCo particles of limited size (22 ± 6, 27 ± 9, 22 ± 6 nm, respectively) but the presence of large metallic domains translated into a larger average particle size according to the Scherrer’s equation (58, 60, 45 nm, respectively). We hoped to reduce the extent of metallic domains by employing incipient wetness impregna-
tion of the saline solution instead of manual milling of the salt. We chose methanol as it is an excellent solvent of nitrates, its low surface tension is more likely to wet the inner pores of the matrix, and its weaker hydrogen bonding makes it easier to evaporate than water or ethanol.

- SEM observations of N-doped matrixes decorated with nanoparticles

Figure 2 presents the typical distribution of Fe, Co, Ni, and Cu nanoparticles over GN and G2N matrixes. Elemental mapping confirmed the attribution of the particles to their corresponding element (see the “SI M Map” files, where M = Fe, Co, Ni, Cu). Cu-decorated materials (D, H) present much bigger particles (>100 nm) than other metals (~20 nm); the larger Cu crystals (with well-defined facets) develop upward longitudinally past 100 nm. The manual measure of at least 300 nanoparticles from five or more representative pictures afforded their statistical size, presented in Table 2 and distribution histograms (Figure SI 2). These data and the careful observation of higher-magnification images (Figure SI 3) indicate that the materials with more nitrogen (G2N) present bigger particles with broader distribution and a less defined shape. By comparing Figure 2D,H, the latter presented particles of less defined outer shape, suggesting that the nitrogen-metal interaction is higher and restrains the formation of Cu crystals (see also Figures SI 3D,H and SI 4). The same phenomenon seemed to occur for smaller particles of Fe, Co, and Ni, even if not as obvious. Contrarily to our previous observations [30], we did not observe areas covered with extensive metallic structures, indicating that reducing the [metal mass]/[matrix surface] ratio and/or the incipient wetness impregnation favors a more homogeneous repartition of the metal over the matrix. Yet, if the presence of wide crystals was prevented here, the average observed particle size remained similar. Elemental analysis by EDS (energy dispersive microscopy) of the surface of our materials at distinct energies (3 KeV, 15 KeV) indicated that Fe, Co, or Ni metals were present at values very close to the expected 5 wt. % (see the corresponding SI M Map files for the EDS). Given the penetration of the beam is highly dependent on its energy (50 nm at 3 keV, 1 um at 15 keV), and as similar results were obtained at distinct energies, the repartition of the metal is likely homogeneous through the thickness of the matrix. It suggests that incipient wetness impregnation efficiently distributes those metals within the whole sample, at least up to the penetrating value of the beam of higher energy. On the other hand, for Cu-decorated matrixes, EDS at 3 keV revealed a mass concentration closer to 20 wt. %, while at 15 keV this value dropped to 5 wt. % As Cu nanoparticles tend to agglomerate to form much larger structures, they are likely less able to remain within a constricted pore.

Table 2. Crystallographic parameters of the NPs obtained by XRPD of the free matrixes.

| Matrix Type | Metallic NP | Average Size a (nm) | De b (nm) | Peak Position (°) c | Attributed Species |
|-------------|-------------|---------------------|----------|---------------------|--------------------|
| GN Fe      | 17 ± 9      | 19                  | 35.6; 62.6 | Fe₂O₃/Fe₃O₄        |
| Co          | 21 ± 9      | 13                  | 44.2      | CoO                 |
| Ni          | 20 ± 7      | 11                  | 36.5; 42.5 | NiO                |
| Cu          | 160 ± 60    | 103                 | 43.3; 50.4; 74.1 | CuO |
| G2N Fe     | 20 ± 10     | 23                  | 35.4; 62.5 | Fe₂O₃/Fe₃O₄        |
| Co          | 30 ± 20     | 11                  | 44.2      | CoO                 |
| Ni          | 21 ± 7      | 37                  | 44.5; 51.9 | NiO                |
| Cu          | 170 ± 50    | 105                 | 43.3; 50.4; 74.1 | Cu          |

a Average particle size determined by manual counting on SEM images, b average crystallite size determined with the peak of highest intensity according to Scherrer’s formula, c peak of highest intensity of a given species is indicated in bold.
Figure 2. SEM observations of the non-impregnated GN (A–D) and G2N (E–H) matrixes decorated with Fe (A,E), Co (B,F), Ni (C,G), Cu (D,H) nanoparticles.

- Structural characterization of N-doped matrixes with and without nanoparticles

The nature and average size of the nanoparticles were determined by powder X-ray diffraction measurements, presented in Figure 3. Except for the matrixes decorated with Cu, broad peaks were observed, a good indication that large crystal domains were absent. The average crystallite obtained by Scherrer's equation (Table 2) appeared very close to the one obtained from manual counting. It indicates large crystals are absent, in opposition to our previous work [30], depicting the interest of the current method. Only in the case of metallic
nickel, the average particle size is a bit higher according to the equation than from counting, but it is still much closer in values than previously. While the diffractograms of Cu-, Co-, and Ni-decorated matrixes displayed the expected metallic peaks, almost every PXRD pattern (except for G2N Cu) also presented a non-negligible fraction of their corresponding oxide. In the case of GN Fe and G2N Fe, only Fe$_3$O$_4$ (or possibly Fe$_2$O$_3$) could be observed, despite a shoulder being present close to 42.8°, accounting for possible narrow metallic particles. It was not the case in our previous work for Ni and Co nanoparticles, albeit then the diffractograms were obtained in the air over several hours, revealing the formation of an oxide layer did not occur rapidly at room temperature. In the current work, an air-tight dome was employed for each sample (accounting for the broad peak between 15–30°), and the samples were exposed to the air for only a short time (less than 5 min) between pyrolysis and vacuum activation. Two factors might have promoted the formation of these oxides: the preponderance of smaller, more reactive particles (namely, because less metal was employed) or the use of methanol. To discriminate between both factors, the same amount of nickel nitrate was manually ground with the matrix, submitted to pyrolysis, and vacuum activated. This material was submitted to PXRD, and no trace of oxide was observed, implying methanol was responsible for the presence of oxides (Figure SI 5A). While the methanol was carefully removed by roto-evaporation and the samples were dried overnight under a flux of dry nitrogen, some solvent likely coordinated with the salt or wet the matrix, affecting the reduction of the metal during the pyrolysis. We were surprised to observe such differences with the ground material; indeed, in any case, we employed nitrate hydrates, the matrix quickly moistened due to air humidity, and the oxides typically formed during pyrolysis were supposed to be fully reduced by the carbon matrix under high temperature in an inert flow. Still, to prevent the formation of oxides, the powders could be placed under a high vacuum at 150 °C (below nitrate melting temperature) for several hours to remove any trace of solvent. In any case, two points must be remembered here: (i) the impregnation of the material is realized under high pressures of H$_2$, and we ensured the oxides were not present after this step (see Figure SI 6); (ii) oxides effectively catalyze the liberation of H$_2$ from LiBH$_4$, so their presence in the decorated material might not necessarily be prejudicial to the final hydrogen storage material [35,39,44–46].

![Figure 3. PXRD patterns of non-impregnated GN (A) and G2N (B) matrixes (black) decorated with Fe (orange), Co (red), Ni (green) and Cu (blue) nanoparticles. Crystalline structures are marked for Fe (▲), Co (▼), Ni (●), Cu (▲). Respective oxides are indicated with empty symbols (◇◇◇). A possible graphite species is marked for G2N Fe and G2N Co (●).](image-url)
Another intriguing point raised by PXRD was the appearance in specific samples of an unexpected peak at 26.0°. Whilst we could not definitively attribute this peak, its position is similar to the peak of graphite [47–49]. It appeared only for G2N matrixes and was more intense for Fe and Co (it was absent from Cu and a shoulder was barely observable with Ni). Interestingly, the same materials presented distinct nitrogen adsorption isotherms, indicating that both the structure and texture of these matrixes were affected during the pyrolysis.

2.2. Chemical and Morphological Characterizations of the LiBH$_4$-Impregnated Matrixes

- SEM observations of just-impregnated N-doped matrixes decorated with nanoparticles

Particles on matrixes just impregnated with LiBH$_4$ appeared slightly larger (Figure 4), with a broader distribution (Figure SI 7), denoting an interaction between the metal and the hydride. This interaction was particularly impressive in the case of G2N Cu (Figure 4H), with splatter-like nanoparticles (see elemental attribution in Figure SI Cu 3). Higher magnification (Figure SI 8) indicated that this process could occur with other metals, as the shape of their respective nanoparticles was less defined. Unfortunately, elemental mapping does not allow us to efficiently differentiate boron from carbon, especially at this magnification. Still, single point EDS indicated that boron was present on metallic nanoparticles. The increase in particle size might account for an external layer of boron derivative coating the metal. Similar to the non-impregnated samples, bigger and less-defined nanoparticles were observed for higher proportions of N-dopant.

In some cases, a binder appeared between the nanoparticles (Figure 4D and Figures SI 8H and SI 9B,D) that can be attributed to the nanoparticles melting together during pyrolysis or LiBH$_4$ melting and gluing these during its impregnation. It should be stated here that the presence of LiBH$_4$ between the nanoparticles might account for an affinity between the metal and the hydride. This being so, this affinity might compete with the insertion of the hydride within the matrix. Molten structures were also observed (Figure SI 10B,C) and in a few cases, we were able to observe extensive spilling at the surface of some chunks of matrix (Figures SI 9C and SI 10E). Once, flat structures were observed outside of the matrix of G2N50 Fe (Figure SI 12I). EDS suggested that higher proportions of boron were present when these structures were observed (Figures SI Co 6, SI Cu 5 and 6, SI Fe 5 and 7). It should be noted that our samples were exposed to air for a very short time (<2 min) when inserting the sample in the SEM vacuum chamber. Elemental analysis indicates hydrolysis of LiBH$_4$ occurred, as higher proportions of oxygen were measured with boron. Given LiBH$_4$ degradation products occupy a larger volume than the fresh hydride, it might explain how boron derivatives were observed outside of the matrixes even if filled at 50 vol. %. The observation of cracks in the matrix when spilled boron was observed also suggested the same expulsion mechanism, so the just-impregnated sample might not necessarily present notable amounts of LiBH$_4$ outside of the matrix, as we will see by DSC.

Some intriguing behavior observed on boron-rich particles, that are far beyond the scope of this article, are presented in the Supplementary Information for the reader’s curiosity (Figures SI 11 and SI 12).
Figure 4. SEM observations of the just-impregnated GN50 (A–D) and G2N50 (E–H) matrixes decorated with Fe (A,E), Co (B,F), Ni (C,G), Cu (D,H) nanoparticles.

• Structural characterization of N-doped matrixes with and without nanoparticles

To gain further insight into the impact of LiBH$_4$ impregnation on decorated matrixes, we present the PXRD patterns in Figure 5, and Table 3 presents the characteristic peaks of our materials. While LiBH$_4$ impregnated 50 vol. % of each matrix, none of its typical crystallographic peaks ($2\theta = 17.7^\circ; 23.7^\circ; 24.7^\circ; 25.6^\circ$) was present in any of our samples. It is specific to LiBH$_4$ impregnated in small pores to lose its crystalline long-range order. Yet, when the nanoparticles were absent, a peak was observed at 12.6$, typical of our N-doped
matrixes once impregnated [29]. We already proposed that this peak might appear by removing elements of symmetry of the orthorhombic cell of LiBH₄. We are not sure if this could be related to the flat structures observed for G2N50 Fe (Figure SI 12I), as this peak was missing when metallic nanoparticles were present, as already observed for Ni and Co [30].

![Figure 5. PXRD traces of just-impregnated GN50 (A) and G2N50 (B) matrixes (black) decorated with Fe (orange), Co (red), Ni (green) and Cu (blue) nanoparticles. Crystalline structures are marked for Co (♣), Ni (♦), Cu (♥). Impregnated LiBH₄ is marked in GN50 and G2N50, assuming crystal order.](image)

**Figure 5.** PXRD traces of just-impregnated GN50 (A) and G2N50 (B) matrixes (black) decorated with Fe (orange), Co (red), Ni (green) and Cu (blue) nanoparticles. Crystalline structures are marked for Co (♣), Ni (♦), Cu (♥). Impregnated LiBH₄ is marked in GN50 and G2N50, assuming crystal order. Impregnated LiBH₄ is marked in GN50 and G2N50, assuming crystal order.

| Matrix Type | Metallic NP | Average Size a (nm) | Dc b (nm) | Peak Position (°) c | Attributed Species |
|-------------|-------------|---------------------|-----------|---------------------|-------------------|
| GN50        | Fe          | 18 ± 6              | -         | -                   | -                 |
|             | Co          | 17 ± 7              | -         | -                   | -                 |
|             | Ni          | 24 ± 9              | -         | -                   | -                 |
|             | Cu          | 160 ± 90            | 98        | 43.3; 50.4          | Cu                |
| G2N50       | Fe          | 20 ± 10             | 10        | 42.5; 45.0          | Fe₂B              |
|             | Co          | 30 ± 10             | 33        | 44.4                | Co                |
|             | Ni          | 22 ± 6              | 37        | 44.5                | Ni                |
|             | Cu          | 170 ± 80            | 89        | 43.3; 50.4; 74.1    | Cu                |

a Average particle size determined by manual counting on SEM images, b average crystallite size determined with the peak of highest intensity according to Scherrer’s formula, c peak of highest intensity of a given species is indicated in bold.

As previously [30], the peak associated with transition elements suffered a loss in intensity once LiBH₄ impregnated the matrix, confirming an interaction between the hydride and the nanoparticles. In the case of GN, only Cu was observed, while in G2N Co and Ni were still observed. It must be highlighted that (i) oxides were not observed, (ii) Cu particles were less affected by LiBH₄ impregnation as they were larger, (iii) G2N50 Fe presented two peaks that could be related to Fe₂B species, (iv) the peak at 26.0° remained mostly unchanged. While metal particles of comparable size (but more dispersed) were observed by SEM after the impregnation with LiBH₄, the PXRD patterns of these samples indicated the loss of their crystalline structure.
2.3. Evaluation of Hydrogen Release and Characterization of Material’s Evolution

2.3.1. Differential Scanning Calorimetry

DSC is a convenient technique for the rapid evaluation of nanoconfinement over hydrogen release. Figure 6 summarizes the impact of N-doping and nanoparticle decoration over the distinctive peaks of nanoconfined LiBH$_4$: phase transition, melting, and hydrogen release. Usually, the loss of long-range order causes these peaks to be flattened and shifted toward lower temperatures. For GN50 (Figure 6A) those three peaks were still relatively well defined, as expected for 50 vol. % filled matrixes. Those peaks were almost unaffected by the addition of metallic nanoparticles, the main differences being: (i) the fusion peak was a bit less intense, (ii) the apparition of a diminutive transition peak around 117 °C, which might account for “core” LiBH$_4$° was a bit less intense, (ii) the apparition of a diminutive transition peak around 117 °C, which might account for “core” LiBH$_4$, filling mesopores, but not in close contact with the matrix [29,30]. This could be in line with our SEM observations that the metal might restrict LiBH$_4$ from going deep within the matrix where there are smaller pores; still, it should not be related to “bulk” LiBH$_4$ plainly outside of the matrix, as this species should present a sharp fusion peak at 275 °C [30]. For G2N50, the situation is a bit more appealing, as the peaks behaved distinctly. There the fusion peak was almost undistinguishable, and the peak of decomposition was lowered and flattened in the presence of Ni and Cu (while they were sharper and shifted toward higher temperatures for Co and Fe). Furthermore, the trace obtained with Ni did not display a peak of transition, which is a good indication of strong nanoconfinement effects. G2N50 Ni exhibited our lowest peak of decomposition (309 °C) at this filling value (Table 4).

![Figure 6](image_url)

**Figure 6.** Differential scanning calorimetry from GN (A) and G2N (B) filled with LiBH$_4$ at 50% (vol) without (black) and with Fe (orange), Co (red), Ni (green), and Cu (blue) nanoparticles.

| Matrix Type | Metallic NP | $T_1^a$ | $T_1^b$ | $T_d^c$ |
|-------------|-------------|---------|---------|---------|
| GN50        | None        | 104     | 239     | 335     |
|             | Fe          | 103 (116)| 238     | 334     |
|             | Co          | 104 (116)| 242     | 335     |
|             | Ni          | 103 (116)| 240     | 332     |
|             | Cu          | 103 (117)| 240     | 335     |
| G2N50       | None        | 98 (115)| 219     | 333     |
|             | Fe          | 103     | -       | 335     |
|             | Co          | 105     | 252     | 339     |
|             | Ni          | 99      | -       | 309     |
|             | Cu          | 99      | -       | 320     |

$^a$ Temperature of orthorhombic to hexagonal transition (°C). $^b$ Melting temperature (°C). $^c$ Temperature of decomposition of the hydride (°C).
2.3.2. Volumetric Study

- First hydrogen release

The volumetric studies are presented in Figure 7 to evaluate the functional behavior and reversibility of our materials. GN50 and G2N50 behaved particularly well, considering they are 50 vol. % filled materials, behaving even better than our previous N-matrixes filled at 30 vol. % (we recall that we augmented the proportion of graphene and N-dopant) [29]. Indeed, G2N50 liberated 1 wt. % \( \text{H}_2 \) (versus m LiBH\(_4\)) at 229 °C (15 ° lower than our previous best), and at 325 °C, it liberated 9.3 wt. % \( \text{H}_2 \) (a 0.7 wt. % increase). Unfortunately, in most cases, the presence of metallic nanoparticles did not improve the functional properties of our hydrogen storage material (Table 5). At best, in the case of GN50 Fe, the presence of metallic nanoparticles did not affect the first and second cycle of hydrogen release, while for the same matrix in the presence of Co, Ni, and Cu nanoparticles to attain 1 wt. % of \( \text{H}_2 \) released, the temperature had to be increased by 16 °C; for those same metals at 325 °C, the proportion of released \( \text{H}_2 \) was reduced by 1 wt. %. In the case of G2N, the impact of metallic nanoparticles was surprisingly negative, reducing the onset temperature by 30 to 40 °C and lowering the \( \text{H}_2 \) release at 325 °C by more than 2 wt. % in average. In any case, the hydrogen released per mass of LiBH\(_4\) was very close to its theoretical value (13.6 wt. %), which is in line with the oxides being reduced during the impregnation process as presented in Figure SI 6. If this was not the case, the available oxides were present in limited quantities (supposing all 5 wt. % Fe was oxidized to Fe\(_2\)O\(_3\), in G2N at maximum 1 mmol of atomic O might be present per 5 mmol of LiBH\(_4\), reducing by 2.5 wt. % the liberated \( \text{H}_2 \); this value decreases to 0.8 wt. % for GN Ni with 50% of oxidized Ni). Still, it cannot be ignored that a very slight decrease in hydrogen capacity was observed when the matrixes were covered by nanoparticles.

![Figure 7](image)

**Figure 7.** Volumetric studies of GN50 (A,C) G2N50 (B,D) without (black) and with Fe (orange), Co (red), Ni (green), and Cu (blue) nanoparticles during the first (A,C) and second (B,D) dehydrogenation cycles.
Table 5. Hydrogen release capacity of GN50 and G2N50 with and without decoration.

| Matrix type | Metallic NP | T1% a | Wt325 % b | Wtrev % c |
|-------------|-------------|-------|-----------|-----------|
| GN50        | None        | 255   | 8.0       | 5.6       |
|             | Fe          | 258   | 8.3       | 5.3       |
|             | Co          | 274   | 6.8       | 5.5       |
|             | Ni          | 271   | 6.9       | 5.8       |
|             | Cu          | 268   | 7.3       | 5.1       |
|             | None        | 229   | 9.3       | 5.4       |
|             | Fe          | 272   | 6.8       | 4.6       |
|             | Co          | 268   | 7.0       | 4.6       |
|             | Ni          | 263   | 7.6       | 5.5       |
|             | Cu          | 260   | 7.4       | 4.4       |

a Temperature (°C) to attain 1 wt. % of liberated H2. b Amount of liberated H2 liberated at 325 °C with respect to infiltrated LiBH4. c Maximum value of H2 released at 400 °C during second dehydrogenation.

- Second hydrogen release

Considering the second cycle, the impact of nanoparticles over hydrogen release was limited for GN, and a bit negative for G2N with a decrease of 0.6 wt. %. As in the previous study, matrices decorated with Ni nanoparticles behaved better than other metals and inclusively better than the metal-free matrices.

2.3.3. Structural Impact of Material Cycling

The impact of two hydrogen release cycles over the material was evaluated by PXRD (Figure 8) and summarized in Table 6. No noticeable peaks were observed from cycled GN50 and G2N50, which is usual for LiBH4 degradation products in matrixes of reduced pore-size [20]. Noteworthily, the peak at 26° observed for G2N Fe and G2N Co was still present but far less intense. Wang et al. reported a decrease in the intensity of this peak under specific conditions [50]. Cu derivatives present very little change regarding their non-impregnated and just-impregnated relatives. In the case of Fe, Co, and Ni derivatives, cycling promoted the appearance of poorly resolved peaks attributed to the formation of Fe2B, CoB, and Ni2B species, as observed in our previous work [30]. According to Scherrer’s equation, those peaks correspond to crystals of limited size (<20 nm), except for the case of Ni2B (46 nm) and metallic Cu (100 nm). In the case of cGN50 Ni, a very weak peak was observed at 43.5°, attributed to a limited fraction of NiO.

Figure 8. PXRD traces of cycled GN50 (A) and G2N50 (B) matrixes (black) decorated with Fe (orange), Co (red), Ni (green) and Cu (blue) nanoparticles. Crystalline structures are marked for Cu (♦). Nickel oxides are indicated (◦). Metal borides are noted for Fe2B (◆), CoB (♣) and Ni2B (✈). A possible graphite species is marked for G2N Fe and G2N Co (♦).
Table 6. Crystallographic parameters of the NPs obtained by PXRD of the cycled matrixes.

| Matrix Type | Metallic NP | Dc $^a$ (nm) | Peak Position ($^\circ$) $^b$ | Attributed Species |
|-------------|-------------|--------------|-----------------------------|-------------------|
| cGN50       | Fe          | 4            | 37.3; 45.6                  | CoB               |
|             | Co          | 46           | 36.1; 42.7; 46.0            | Ni$_2$B           |
|             | Ni          | 21           | 43.5                        | NiO               |
|             | Cu          | 112          | 43.3; 50.4; 74.1            | Cu                |
| cG2N50      | Fe          | 14           | 42.5; 45.0                  | Fe$_2$B           |
|             | Co          | 21           | 37.3; 45.6; 41.1; 49.2      | CoB               |
|             | Ni          | 16           | 46.0                        | Ni$_2$B           |
|             | Cu          | 98           | 43.3; 50.4; 74.1; 89.9       | Cu                |

$^a$ average crystallite size determined with the peak of highest intensity according to Scherrer’s formula, $^b$ peak of highest intensity of a given species is indicated in bold.

3. Discussion

The decoration of GN matrixes by incipient wetness impregnation afforded Fe, Co, and Ni nanoparticles of 17 to 21 nm (20 to 30 nm on G2N matrixes) evenly distributed on the surface and throughout the thickness of the matrix, with no evidence of large metallic domains (unlike with manual grinding of the salt). Nevertheless, the same method afforded 150 nm Cu nanoparticles concentrated on the surface of the matrix. It illustrates that if a metal displays interesting theoretical features (as with Cu), its application in the real material might present difficulties not considered (such as aggregation) by the thermodynamics of reaction. Whereas incipient wetness impregnation displayed some advantages over manual grinding of the salt, it also promoted the formation of oxides in the cases of Fe, Co, and Ni. By impregnating the matrix with LiBH$_4$ the particles appeared slightly larger and SEM observations indicated this could be due to the formation of a very thin LiBH$_4$ film covering these. If it is the case, one should consider if the competition between matrix or nanoparticle wetting by LiBH$_4$ might limit the performance of the system. Higher proportions of LiBH$_4$ might also promote the clustering of nanoparticles.

The interaction between the nanoparticles and LiBH$_4$ was confirmed by PXRD, with the formation of Fe$_2$B, CoB and Ni$_2$B. Interestingly, no metal borides were observed with Cu nanoparticles. Phase separation is a major limitation for the reversible degradation of LiBH$_4$, as LiH and B segregate, particularly in carbon matrixes. It should be determined how metal borides and lithium hydride are locally distributed along several hydrogen release/capture cycles.

N-doped matrixes displayed interesting behavior, with notably low temperatures of hydrogen release. Raising the proportion of nitrogen and graphene dopants improved the material’s performance, suggesting that there is some room to improve our matrixes with nitrogen derivatives. On the other hand, while the matrix’s decoration with metallic nanoparticles was performed with good control over particle size and distribution, the impact on performance was negligible at best, antagonistic in some instances. It refutes our previous observations, where Ni, Co, and NiCo nanoparticles reproducibly improved N-doped matrixes [30]. Several factors should be highlighted to discuss these contradictions:

1. Here, the proportion of metal was limited to 5 wt. %, while in our previous work, the proportion of nanoparticles was set to 5% of the mesopore volume, reaching a 27 wt. %
2. The matrixes’ pore size was lowered, and the proportion of N-dopant was increased, which are conditions more favourable to hydrogen release;
3. Oxides were observed in the present work.

Factor 1 is crucial as, if a metal promotes hydrogen liberation, higher proportions of this metal are likely to display more impact, up to coalescence of the particles. Thus, metal catalysts usually present an optimum of activity at an intermediate weight percentage, where the particles are small enough to present a high surface/volume ratio, with enough mass to display noticeable activity. Nevertheless, higher proportions of dense metal
will drastically reduce the hydrogen capacity of our materials. Considering the overall performance, it was crucial to determine if the activity of the added metallic dopant was catalytic or massive. The results showed that lowering a metal’s mass at the surface of the matrixes lessened activity, suggesting that catalytic quantities of metal are not enough to improve the material’s performance. The discrepancy between this work and the previous one [30] can highlight that a substantial amount of metal is required to observe an enhancement, thus posing the question of the method’s viability as mass capacity is the central parameter of this topic.

Regarding factor 2, in our previous work [30] the N-doped matrixes presented larger pores of broader distribution (9 nm), and its decoration with metallic nanoparticles lowered the onset of hydrogen liberation. Here we employed N-doped matrixes with well-defined pores of 4 to 6 nm, but their decoration with distinct nanoparticles was not beneficial for the performance of the material. This suggests that the effect of metallic nanoparticles does not synergize well with reducing the size of the pores, one effect taking dominance over the other. Moreover, if we compare this work to our previous N-doped matrixes, it appears that the results are quite comparable, even better in the present publication, probably because a higher proportion of nitrogen was doping the matrixes [29]. It is likely that hydrogen dissociation is a limiting factor for LiBH₄ decomposition that can be tackled by nanoconfinement, nitrogen doping and/or metal catalyst. Nevertheless, if in matrixes of reduced pore size the hydrogen dissociation is not a limiting factor anymore, then the metal catalyst would lose its specific interest.

The relevance of factor 3 is more debatable as the oxides mostly disappeared after the impregnation process. Secondly, the presence of oxides, while detrimental to the reversibility of the hydrides, is not necessarily limiting their kinetics, as there is proof of faster hydrogen release in the presence of oxides [35,39,44–46]. To settle the origin of those oxides, we decided to decorate our matrix by manual grinding instead of incipient wetness impregnation. It appeared that by reproducing the manual grinding of the matrix with nickel nitrates we observed only the peaks of metallic nickel, as previously (Figure SI 5). This evidenced that the oxides originate from dissolving the nitrate in methanol. We are not sure how dissolving an aqueous nitrate in methanol can result in such a drastic change in our material after drying and pyrolysis, but we must assume that some coordination complexes were formed that hindered the complete reduction during the pyrolysis. Further calorimetric and volumetric characterizations of our solid-grinded material were performed, and very few differences were observed compared to the wet-impregnation protocol (Figure SI 13), discarding de facto the role of oxides over the poor performances of the nanoparticle’s decoration strategy.

Yet, if the lack of positive results observed for GN matrixes can be attributed to the insufficient quantities of metal and reveals the absence of synergies with the porous structure, we were surprised that, in the case of G2N, the metal-decorated materials performed even worse than their non-decorated counterparts. SEM observations and PXRD patterns illustrated the existence of an interaction between the metallic nanoparticles and just-impregnated LiBH₄. In our ultimate work, we demonstrated that LiBH₄ fills micropores first and these had much better performances than mesopores during the first cycle [31]. G2N matrixes present a huge proportion of micropores (43%) and our best performance so far. We were curious to see if the decline of the material’s performance could be related to LiBH₄ wetting the matrix. In Figure SI 14 we compare the wetting behavior of G2N with G2N Ni after impregnating it with LiBH₄ at 50 vol. %. While non-impregnated G2N and G2N Ni display very similar textural characteristics, just-impregnated G2N50 displays many fewer empty micropores than its G2N50 Ni counterpart. It convincingly attests that the presence of nickel nanoparticles might hinder the complete wetting of the smaller pores of the matrix, as the metal is competing with the matrix for LiBH₄’s contact. Interestingly, Gross et al. improved the wetting of their matrix by decoration with nickel nanoparticles, but what is noteworthy, they confined metallic magnesium instead of lithium borohydride, so the affinity of the impregnated material for the decorating particles differed.
widely [51]. It is also worth pointing out that, if decorated by Co or Fe, G2N displayed a decrease of the micropore volume (related to the appearance of graphitic domains at 2θ = 26°) that could be related to their poor overall performance.

4. Materials and Methods

4.1. Synthesis of Materials

4.1.1. Chemicals

All chemicals were employed as received, without purification. Graphite flakes were supplied by Aldrich (St. Louis, MO, USA, product number 33246-1) and hydrogen peroxide (30%), formaldehyde (40%), and ethylenediamine (98.0%) by Biopack (Buenos Aires, Argentina). Ascorbic acid (99.0%), potassium permanganate (99.0%), sulfuric acid (98%), hydrochloric acid (35%), diethyl ether (98.0%), ethanol (99.5%), and resorcinol (98.5%) were purchased from Cicarelli (San Lorenzo, Argentina) and phosphoric acid (85%) from Merck (Kenilworth, NJ, USA). Sodium carbonate decahydrate (99.99%) was provided by Timper. Co(NO$_3$)$_2$·6H$_2$O was provided by Biopack (98.0%), while Ni(NO$_3$)$_2$·6H$_2$O, Fe(NO$_3$)$_3$·9H$_2$O, and Cu(NO$_3$)$_2$·XH$_2$O were supplied by Sigma-Aldrich (97.0%). LiBH$_4$ (Sigma-Aldrich, 90%) was milled prior to use. Every hydride material was handled within a glovebox to avoid air contact (content of oxygen and water < 5 ppm).

4.1.2. Wet Synthesis

In brief, graphene oxide was prepared from graphite flakes according to the Tour method, employing potassium permanganate as oxidant in a mixture of sulfuric and phosphoric acid [52]. Later, the reaction mixture was poured at room temperature (RT) over ice with hydrogen peroxide to solubilize any manganese derivative. Then, it was washed, filtered, and centrifuged several times with Milli-Q water, diluted hydrochloric acid, ethanol, and finally coagulated with diethyl ether. After drying, a 6 mg/mL graphene oxide suspension in Milli-Q water was dispersed in an ultrasonic bath for 1 h and stored at 4 °C.

Each gel and following resin was prepared in a polypropylene bottle, screw-tapped before heating, to avoid any mechanical perturbation. N-doped hydrogels were formed by adding 10 mL of a recently sonicated (30 min) graphene oxide suspension (6 mg/mL) to a 1/5 diluted aqueous ethylenediamine solution (0.75 and 1.5 mmol for GN and G2N resp.) under magnetic stirring [53]. After a quick homogenization (1 min) the mixture was placed in an oven at 85 °C for 5 h. This temperature promoted gel shrinkage by 10%. The gel was washed several times with water at RT, then wet every 4 h with 3 g of the resorcinol—formaldehyde sol over 24 h, and the supernatant was discarded prior to the addition of the following aliquot. The sols were prepared with a mixture of water (5.43 g), resorcinol (13.05 g), formaldehyde (19.52 g), and sodium carbonate (71.9 mg). After the last washing, the gel was placed in an oven at 50 °C for 24 h and then at 90 °C for 72 h. Once cooled, the resins were washed with water and acetone and allowed to dry.

4.1.3. Solid-State Synthesis

The resins were broken to smaller pieces (approx. 1 cm$^3$ each, any red part without graphene was visually discarded) and pyrolyzed at 800 °C (3 °C/min from RT to 600 °C, 60 min dwelling at this temperature then heated to 800 °C at 3 °C/min and staying at this temperature for 6 h, then allowed to cool to RT at 3 °C/min over 3 h) to afford black chunks. Before impregnation, the resins and the LiBH$_4$ were separately milled by employing a P6 Pulverisette planetary device, with an 80 cm$^3$ milling chamber and five stainless balls, under an argon atmosphere. LiBH$_4$ (1.5 g) were ball-milled for 300 min at 400 rpm, with a sequence of 10 min milling and 10 min pause. To reduce morphological impact and Fe contamination, the resins were milled for a shorter time: five reverse repetitions of 2 min milling at 200 rpm with a 1 min pause. The powders obtained from the resins were activated under reduced pressure (5 °C/min ramp then 3 h at 400 °C) prior to their introduction in the glove box.
To decorate the carbon material with metallic NPs, 10 mL of nitrate solutions (9.0 mM for Fe, 8.5 mM for Co, 8.5 mM for Ni, and 7.9 mM for Cu) were poured over 0.095 g of resin powder and agitated at RT (in the air) for 2 h. The amount of salt was calculated so that the mass of reduced metal would equal 5 wt. % of the matrix. The methanol was removed in a rotavapor at 65 °C until the powder looked dry on the glass. Then the powder was placed in a quartz tube that was degassed with N2 at 50 mL/min overnight. The mixture was heated at 800 °C (4 °C/min from RT to 200 °C, 2 h dwelling at this temperature then heated to 800 °C at 4 °C/min and staying at this temperature for 5 h, then allowed to cool to RT at 3 °C/min over 3 h) to promote the melting of the nitrate and its following reduction by the carbon surrounding material [54,55] These powders were activated at 400 °C under reduced pressure (5 °C/min ramp then 3 h at 400 °C) prior to their introduction in the glove box. To limit any oxidation of the metal nanoparticles, the resins were transferred quickly (<1 min) from the quartz tube of the furnace to the vacuum tube, which was opened after activation in the glove box.

The powders of resins (either decorated or not) and LiBH4 were manually mixed within the glovebox for 30 min using a mortar and pestle at 50 vol. %. The amount of LiBH4 employed to fill the resins at any given volume percentage was determined according to the volume of micropores + mesopores obtained from the nitrogen desorption isotherms.

4.1.4. Melt Impregnation

The mixed powders of resins and LiBH4 were placed in a reactor within an autoclave and heated to 300 °C under 60 bar H2 for 30 min. At this temperature, the solid LiBH4 melts and the liquid LiBH4 wets the resin and fills its pores, while the high hydrogen pressure ensures that no hydrogen is liberated by the hydride. Once the system cooled back to RT, two options were taken: (i) the sample’s hydrogen capacity was directly evaluated using a modified Sieverts-type device and (ii) the impregnated sample was stored in the glove box to be submitted to other techniques.

4.2. Material Characterization

Textural parameters of the samples were studied using a Micromeritics ASAP 2020 analyzer. After surface cleaning in vacuum overnight, N2 adsorption/desorption isotherms were collected at −196 °C on 0.1 g of sample. For resins without LiBH4, this was done at a temperature of 300 °C, while if LiBH4 was present, the temperature was limited to 150 °C. The surface area and pore size distribution were obtained by the application of the Brunauer−Emmett−Teller (BET) and the Barrett−Joyner−Halenda (BJH) models, respectively. BET was determined for 0.03 < P/P0 < 0.12 with positive values of C. The mesopore volume was calculated according to BJH, and the total pore volume was determined by the Gurvich method at P/P0 = 0.96. The micropore volume and the external surface were estimated with the t-plot method. In this case, we used the standard reference t-curve for carbonaceous materials (carbon black) proposed by Magee [56].

Morphological and agglomerate size distribution analyses of the samples were performed by scanning electron microscopy (SEM, SEM-FIB, Zeiss, Crossbeam 340), the powders were dispersed over a carbon tape. Elemental analyses of the materials were also performed by energy-dispersive X-ray spectroscopy (EDXS) on SEM.

X-ray powder diffraction (PXRD) was realized with an air-tight chamber filled in a glove box, and the trace was recorded on a Bruker D8 ADVANCE apparatus using Cu Kα radiation.

The thermal desorption behavior of the hydride phases was studied by differential scanning calorimetry (DSC, TA q2000 calorimeter), using a heating rate of 5 °C/min and an argon flow rate of 122 mL/min. The samples were placed in a closed aluminum holder within the glovebox to minimize air contact. For each plot, the heat flow was normalized with respect to the mass of LiBH4.

Hydrogen sorption kinetic measurements were obtained using modified Sieverts-type equipment coupled with a mass flow controller. The sample was placed in a stainless reactor,
within an autoclave that was connected to the Sieverts device. Dehydrogenation curves were obtained by heating up to 400 °C with a hydrogen back pressure of 0.5 ± 0.1 bar. The amount of absorbed/desorbed hydrogen is expressed as the H₂/LiBH₄ mass ratio and is determined with a relative error of ± 5%. Rehydrogenation was performed at 400 °C with a sudden increase of the pressure to 60 bar and keeping the sample overnight and measuring the evolution of pressure (typically higher than 50 bar). After cooling, a second dehydrogenation was performed and the sample was collected for further characterization (noted “c”).

In the ensuing discussion, the samples are presented accordingly to the following codes: non-decorated (GN and G2N, black), decorated with Fe (orange), Co (red), Ni (green), Cu (blue). The samples just impregnated with LiBH₄ are noted by a number indicating the volumetric percentage of LiBH₄ filling the matrix (for example GN50). The cycled samples (submitted to dehydrogenation/rehydrogenation/dehydrogenation) are highlighted by a “c” prefix (for example cGN50).

5. Conclusions

Two N-doped matrixes presenting micropores and small mesopores (3.8 and 6.1 nm) were decorated at 5 wt. % by incipient wetness impregnation using Fe, Co, Ni, and Cu nitrates in methanol. Fe-, Co-, and Ni-decorated matrixes presented particles of limited size (20 nm) homogeneously distributed over and within the matrix, while Cu nanoparticles were larger (160 nm) and concentrated on the surface. Only metallic Cu was observed by PXRD, while the matrixes decorated with Fe, Co, and Ni also presented oxides, attributed to the incipient wetness impregnation. The oxides disappeared after LiBH₄ was impregnated at 50 vol. % and metal borides were observed after two hydrogen/release cycles. According to DSC studies, the impact of metal nanoparticles on the calorimetric behavior of the material was limited. Volumetric experiments revealed that the presence of metallic nanoparticles did not improve the hydrogen release properties of GN matrixes and induced even worse behavior for G2N. We proposed that the low proportion of metal and their lack of synergies with pores of reduced size might make it harder to improve the already good behavior of N-doped matrixes. In the case of micropore-rich G2N, the competition between LiBH₄ pore-wetting and LiBH₄ metal-wetting could be responsible for the observed decrease in the material’s performance.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/molecules27092921/s1, Figure SI 3; Figure SI Co 1–6; Figure SI Cu 1–7; Figure SI Fe 1–7; Figure SI Ni 1–4. Figure SI 1. Nitrogen isotherms (A,B) and corresponding pore size distribution obtained by BJH (C,D) of resins GN (A,C, black) and G2N (B,D, black) decorated with Fe (orange), Co (red), Ni (green) and Cu (blue); Figure SI 2. Size distribution histograms of metallic nanoparticles observed by SEM of non-impregnated matrixes GN (light) and G2N (dark) decorated with Fe (orange, brown), Co (red, wine), Ni (green, dark green), Cu (blue, deep blue); Figure SI 3. High magnification (50k x) SEM observations of GN (A–D) and G2N (E–H) free matrixes decorated with Fe (A,E), Co (B,F), Ni (C,G), and Cu (D,H); Figure SI 4. Very high magnification (90 k x) SEM observations of Cu nanoparticles decorating GN (A) and G2N (B) matrixes; Figure SI 5. PXRD traces of free (A), just-impregnated (B) and cycled (C) GN matrixes decorated with Ni nanoparticles prepared by incipient wetness impregnation in methanol (green) versus manual grinding (teal) of nitrates. The symbols are attributed to the cubic metal (♦), its oxide (♣), and the metal boride (♠); Figure SI 6. PXRD traces of GN Ni just prepared by incipient wetness impregnation in methanol (green) and after 30 min at 300 °C, 60 atm H₂ (dark green); Figure SI 7. Size distribution histograms of metallic nanoparticles observed by SEM of just-impregnated matrixes (cyan) GN50 and G2N50 with respect to their non-impregnated counterparts decorated with Fe (orange, brown), Co (red, wine), Ni (green, dark green), Cu (blue, deep blue); Figure SI 8. High-magnification (50k x) SEM observations of GN50 (A–D) and G2N50 (E–H) just-impregnated matrixes decorated with Fe (A,E), Co (B,F), Ni (C,G), and Cu (D,H); Figure SI 9. Cu decorated GN50 (A–C) and G2N50 (D–F) just-impregnated matrixes; Figure SI 10. GN50 (A–C) and G2N50 (D–F) just-impregnated matrixes decorated with Fe (A,D), Co (B,E) and Ni (C,F) displaying excessive LiBH₄ derivative with nanoparticles visible.
by transparency; Figure SI 11. Just-impregnated decorated matrix with excessive LiBH₄ derivative observed by secondary electrons (A,C) and in lens (B,D) modes; Figure SI 12. Evolution of excess LiBH₄ degradation products in just-impregnated matrices (A,C,E,G,I), and after several minutes of observation (B,D,F,H,J); Figure SI 13. Calorimetric (A) and volumetric (B: first cycle, C: second cycle) studies of GN50 without (black) and with Ni nanoparticles formed by incipient wetness impregnation (green) and manual grinding (teal); Figure SI 14. (A) Isotherms of G2N (black) and G2N Ni (green) before (up, thin) and after (down, bold) impregnation at 50 vol. % with LiBH₄, (B) pore-size distribution obtained by BJH of G2N (black) and G2N Ni (green) before (up, thin) and after (down, bold) impregnation at 50 vol. % with LiBH₄; Table SI 1. Textural parameters of G2N before and after impregnation of LiBH₄ at 50 vol. % with and without Ni nanoparticles. Elemental mapping of the decorated matrices before and after their impregnation with LiBH₄ are presented to sustain the appropriate metallic nature of the particles. They are entitled SI Map X, where X is the metal decorating the matrix (X = Fe, Co, Ni, and Cu).

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