# REVERSIBLE HYDROGEN SORPTION IN THE COMPOSITE MADE OF MAGNESIUM BOROHYDRIDE AND SILICA AEROGEL

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# Abstract

Magnesium borohydride  $Mg(BH_4)_2$  is a promising hydrogen storage material as it releases high hydrogen storage capacity at mild desorption temperatures, but it is still limited by slow hydrogen release kinetics and by the harsh conditions required to re-hydrogenate this compound. In this work, composites made of commercial Mg(BH<sub>4</sub>)<sub>2</sub> and synthesized silica aerogel microparticles were prepared by thermal treatment in hydrogen under 120 bar and 200°C. As a result, the sorption properties of the hydride are improved: calorimetric measurements show that decomposition temperature is reduced by 60°C, and the typical 3-step decomposition mechanism of  $Mg(BH_4)_2$  changes to a single-step mechanism in range of 220-400°C. The kinetics of the first dehydrogenation at 300°C was two times faster in Mg(BH<sub>4</sub>)<sub>2</sub>- $SiO_2$  composites than in the case of bulk  $\gamma$ -Mg(BH<sub>4</sub>)<sub>2</sub>. Additionally, the re-hydrogenation of this material at comparatively moderate conditions of 390°C and 110 bar is presented for the first time, achieving cyclability with a reversible release of hydrogen up to 6wt%. Different amounts of hydrogen were exchanged depending on the temperature of desorption (300°C or 400°C) and the presence or absence of silica aerogel. This result indicates that silica aerogel chemically interacts with Mg(BH<sub>4</sub>)<sub>2</sub>, acting as an additive, which can result in different hydrogenationdehydrogenation routes in which different amounts and types of intermediates are formed, influencing the kinetics and the cyclability.

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## 1. Introduction

Hydrogen society, which is related to the use of hydrogen as an energy vector, could be a solution to current environmental and economic problems associated to the use of fossil fuels [1][2]. Due to the low density of hydrogen gas, an important challenge for onboard applications is the search of an effective technology to reversibly store hydrogen in a compact system [3].

Among currently studied solid state hydrogen storage materials, complex metal hydrides, such as LiBH<sub>4</sub> [4] or NaAlH<sub>4</sub> [5], are very attractive due to their high hydrogen storage capacity, but they are limited by their kinetics and thermodynamics. In this work, magnesium borohydride Mg(BH<sub>4</sub>)<sub>2</sub> is also considered as effective storage material due to its high gravimetric (14.8% wt H<sub>2</sub>) and volumetric capacity (0.112 kg/L) [6], which exceeds the ultimate targets from Department of Energy (DoE) for 2020. Because of these outstanding properties, it could be also of interest in batteries applications, since Mg metal holds better volumetric capacity and is more abundant than Li. Battery-related applications of this complex hydride has been recently reviewed in [7].

Moreover,  $Mg(BH_4)_2$  starts the decomposition at 270°C, and although this decomposition temperature still is too high, it is lower than the decomposition temperature of similar alternative compounds. However, due to the formation of stable intermediates [6], rehydrogenation still remains as the main challenge for use of  $Mg(BH_4)_2$  or their composites.

High conditions of pressure and temperature are necessary in order to obtain a high conversion grade for Mg(BH<sub>4</sub>)<sub>2</sub> from decomposed products. Reversibility of 11wt% of hydrogen [9] was experimentally shown starting from MgB<sub>2</sub> to Mg(BH<sub>4</sub>)<sub>2</sub> at 950 bar H<sub>2</sub> and 400°C during 108 hours. In another work of Newhouse et al. [10], MgB<sub>2</sub> obtained after thermal decomposition at 600°C was partially rehydrogenated, getting 9.7 wt% of hydrogen by treatment at similar conditions of pressure and temperature (900 bar H<sub>2</sub> and 390°C) for 3 days. Li et al. [11][12]reported 6.1wt % hydrogen absorbed after rehydrogenation at 400 bar H<sub>2</sub> and 270°C for 48 h through the formation of the stable MgB<sub>12</sub>H<sub>12</sub> intermediate. However, when mild pressure and temperature conditions (134 bar H<sub>2</sub> and ~300°C) were used during rehydrogenation, only 2.7-3.1 wt% [6] or 3.5wt% H<sub>2</sub> [13] was achieved which most probably corresponded to the formation of MgH<sub>2</sub>. Also in [14] a triborane Mg(B<sub>3</sub>H<sub>8</sub>)<sub>2</sub> in addition to MgH<sub>2</sub> was reabsorbed at 250°C and 120 bar for 48 hours. In a recent work [15], the rehydrogenation was performed at similar conditions of 280 °C and 120 bar hydrogen for 14hours, reabsorbing only 2.5 wt% suggesting the formation of two Mg-B-H phases during

desorption, only one of which being reversible and could explain the limited absorption. The other intermediate compound could be too stable and prevent the complete rehydrogenation.

Different strategies have been tested in order to improve not only the reversibility but also the kinetics and the temperature of the desorption reactions [16] and recently reviewed in [8]. The addition of different additives or catalysts such as NbF<sub>5</sub>, TiO<sub>2</sub>, CoCl<sub>2</sub>, CoF<sub>2</sub> or TiCl<sub>3</sub> among others [14-15], the confinement in porous materials (mainly in carbon materials) [19–21], their combinations [23] or the formation of composites [24] are some of the proposals that have been studied.

Fitchner et el. [20], successfully confined around 44wt% of the hydride in activated carbon (BET area of 860 m<sup>2</sup>/g and pore volume of 0.61 g/cm<sup>3</sup>) using wet impregnation as method and diethyl ether as solvent. As a result, a composite with lower decomposition temperature and lower activation energy was obtained. Wahab et al. [23] infiltrated 45 wt% Mg(BH<sub>4</sub>)<sub>2</sub> in ordered mesoporous carbon CMK-3 scaffold (BET area of 1499m<sup>2</sup>/g and a volume of pores of 1.63cm<sup>3</sup>/g) with addition of 5 wt% of Ni. In this case, a lowering on 200°C in the decomposition temperature was obtained and kinetics of hydrogen release were 10 times faster compared to the bulk Mg(BH<sub>4</sub>)<sub>2</sub> due to the synergic effect of nanoconfinement and of the addition of the Ni catalyst.

Since decomposition tends to occur before melting point, melt infiltration is not a suitable approach for nanoconfinement in this case [19][25]. The only possibility is to confine  $MgH_2$  in a support via melt infiltration and then treat the composite with  $B_2H_6$  in order to obtain  $Mg(BH_4)_2$  infiltrated in the host [21].

In the case that composites are formed [24], 3.6wt% of hydrogen was reversible in 1h at 90bar hydrogen, after 12h of decomposition process at 180 °C for 20 cycles due to the formation of polyborane intermediates which could transform to [BH<sub>4</sub>], and make the whole system reversible under mild conditions.

In this work, we report the experimental trial to confine  $Mg(BH_4)_2$  in microparticles of silica aerogel by temperature treatment in hydrogen atmosphere. Microparticles of silica aerogel, which were used for the first time with this complex hydride, have been produced by supercritical CO<sub>2</sub> drying, a technique that enables to produce a material with outstanding surface properties. The drying method employed is a key aspect that determines the textural properties of this porous host. If the solvent is removed by evaporation or lyophilization, the capillary stresses associated to the formation of vapor-liquid interfaces inside the pores of the support are responsible of partial collapses of the pore structure of the material. In case of SiO<sub>2</sub> matrixes, the materials obtained by these drying methods usually show pore volumes below 0.5 - 1.0

 $cm^{3}/g$ , such as SBA-15 or MCM-41 mesoporous silica. In contrast, if pressurized or supercritical carbon dioxide is used to extract the solvent, the collapse of the pore structure is avoided or minimized, because under these conditions carbon dioxide is completely miscible with the organic solvent, and therefore the extraction proceeds without formation of gas-liquid interfaces[26] and without capillary stresses, thus resulting in pore volumes up to 4  $cm^{3}/g$  [27].

The prepared Mg(BH<sub>4</sub>)<sub>2</sub>–SiO<sub>2</sub> composites were characterized by SEM, XRD and coupled calorimetric-manometric technics. Their kinetic curves were obtained during several hydrogen desorption-absorption cycles and compared to those of the bulk  $\gamma$ -Mg(BH<sub>4</sub>)<sub>2</sub>.

#### 2. Experimental methods

# 2.1 Reactants

The commercial  $\Upsilon$ - Mg(BH<sub>4</sub>)<sub>2</sub> powder (Sigma Aldrich, 95%), thereafter referred to as MBH, was constituted by prismatic particles of around 100 µm (see SEM images reported in Section 3).

Tetramethylorthosilicate (TMOS, 98.0% purity) and ammonium hydroxide (NH<sub>4</sub>OH, 28.0-30.0% ammonia) were supplied by Sigma-Aldrich. Methanol (MeOH; 99.8%) and n-hexane (95%) were purchased from Panreac. Gaseous Carbon dioxide (CO<sub>2</sub>, 99.95%) was supplied by Carburos Metálicos S.A. These reactants are necessary to obtain microparticles of silica aerogel.

Microparticles of silica aerogel were synthesized using a batch supercritical equipment as drying method as reported in a previous work [28]. They were prepared by a sol-gel reaction, using TMOS as precursor and methanol as solvent. While with this procedure it is common to prepare large gel monoliths, in this work the gel was synthetized as microparticles, in order to reduce the possible heat and mass transfer resistances that could be caused by larger aerogel monoliths. To do this, the sol-gel reaction media were dispersed in hexane under mechanical stirring, in order to obtain small droplets of TMOS in methanol dispersed within the hexane continuous phase. After 10 minutes of mechanical stirring of this mixture, an aqueous solution of NH<sub>4</sub>OH was added as condensation catalyst, which induced the gelation of TMOS. The molar ratio used was the following: 1 mol TMOS: 4.4 mol MeOH: 3.3 mol H<sub>2</sub>O: 4.5 mol hexane: 0.08 mol NH<sub>4</sub>OH. Gel particles were then kept in a closed vessel immersed in methanol for an aging period for at least 4 days. During this time, the methanol was renewed 2-3 times in order to remove the water content of the solution. After this time, microparticles of alcogel immersed in methanol were dried using supercritical  $CO_2$  at 110 bar and 40°C, employing the supercritical drying apparatus described in a previous work [28]. For this, the system was slowly pressurized and depressurized at a rate of 3 bar/min in order to avoid the cracking of the alcogel/aerogel and mechanical stresses that could damage the structural properties of the final product. Saturated  $CO_2$  in the system was renewed four times in order to obtain completely dried particles. Dried silica aerogel microparticles had a porous structure with surface area of 723 m<sup>2</sup>/g and a pore volume of 1.35 cm<sup>3</sup>/g (mean pore size =7.5nm) [29]. Obtained SiO<sub>2</sub> aerogel particles were collected and stored in a closed vial at room temperature until used for MBH–SiO<sub>2</sub> composite preparation.

# 2.2 The MBH-SiO<sub>2</sub> preparation and characterization

All handling and preparation of the samples took place in a MBRAUN Unilab glove box which continuously purified argon atmosphere where oxygen and moisture values were kept below 1ppm.

Commercial  $\gamma$ -MBH and silica aerogel particles were treated via a pressure-temperature method. First, SiO<sub>2</sub> aerogel were heated at 5°C/min in a alumina boat placed in a tubular oven at 110°C in the Ar glove-box for 2 hours in order to remove water traces adsorbed by the aerogel from the air. Subsequently, MBH and silica aerogel particles were mixed in a mortar with a mass ratio of 1:1. Around 200 mg of the mixture were introduced in a high pressure stainless steel sample holder under Ar atmosphere in the glove box and then transferred to the Sievert type apparatus (PCT-Pro 2000, Setaram and Hy-Energy), whose experimental set-up is schematically shown in figure 1. The sample was treated at 120 bar of hydrogen and 200°C for 3 hour in order to obtain a composite in which after phase transformation [6] the irreversible  $\beta$ -phase of MBH, can be coated by SiO<sub>2</sub>.

## (FIGURE 1)

Kinetic measurements and the study of reversibility were done in the same Sievert type apparatus. The prepared MBH–SiO<sub>2</sub> composite was dehydrogenated and rehydrogenated for 3 cycles in the same high pressure cell. Hydrogen desorption was tested at 300 and 400 °C both in static vacuum while rehydrogenation was made at 390 °C under 110 bar H<sub>2</sub>. The same conditions were used for bulk  $\gamma$ -MBH in order to compare its reversibility and cyclability with those of the prepared composites.

The gases evolved from the sample during dehydrogenation were analyzed in a residual gas analyzer (RGA Pro, Setaram & Hy-Energy) connected to the manometric instrument.

Coupled manometric-calorimetric measurements were done by connecting the Sievert apparatus to a high pressure differential scanning calorimeter (DSC, Sensys Setaram) in order to study the influence of silica aerogel in the composite system. The calorimeter cell was loaded with around 19 mg of the sample and heated up to 460°C at a rate of 5°C/min in static vacuum. Argon was

used as carrier gas at 10 mL/min. Calisto was used as software for data acquisition and processing.

Crystallinity of the different samples was examined using an ex-situ X-ray powder diffractometer (model D5005 Bruker). The measuring conditions were CuK $\alpha$  radiation,  $\lambda$ =1.54060 Å, 2 $\theta$  angle ranging from 5° to 90° with a scan rate of 10 s/step and a step size of 0.020°. A Bruker-dome was used in order to avoid the contact with air and perform measurements under Argon (gas in the glove-box) and room temperature.

Particle morphology was observed by Scanning Electron Microscopy (SEM) using EVO-MA10-HR (Zeiss, Germany) with energy-dispersive microprobe analyzer INCA Energy 350X Max from Oxford instruments. A special home-made sample holder was used in order to avoid the contact of the powders with atmospheric oxygen and humidity: in particular, the samples were fixed on Al stubs by C-tapes in the glove box and closed in the sample holder. Rough vacuum was created in it by a vacuum pump before its extraction from the glove box. Only after reaching high vacuum in the SEM, the sample holder was open and thanks to the 3D movements of the SEM stage and the samples could be analyzed.

#### **3.** Results and discussion

#### 3.1 XRD and SEM analyses

XRD profiles showed that crystalline  $\Upsilon$ -MBH and amorphous SiO<sub>2</sub> (Fig. 2*c*) were converted into an amorphous mixture after infiltration treatment (Fig. 2*b*). The same amorphous spectrum was obtained after several cycles. This fact suggests that in the prepared composites MBH was amorphous or infiltrated into the pores of SiO<sub>2</sub> aerogel.

# (FIGURE 2)

For the hand-made mixture of MBH and  $SiO_2$  aerogel Rietveld refinement was performed similar to the method proposed in [31]. Obtained results suggest that crystalline powder has particle size around 138 nm and a cell parameter of cubic structure was estimated of 1.5 nm.

In Fig. 3, the SEM image of the prepared MBH–SiO<sub>2</sub> composite is shown. SEM-EDX analysis suggested that MBH and silica were mixed uniformly and their homogeneity and contact of both components was higher compared to that of their hand-made mixture. At these conditions of pressure and temperature, it was difficult to conclude if MBH was infiltrated into the pores of the silica aerogel. More pressure would be necessary in order to achieve this, or alternatively another method of infiltration such as wet impregnation should be used. A similar morphology of the composite was observed after 3<sup>rd</sup> desorption, as it is observed in Fig. 3*b*, which means that silica coating avoided the aggregation of MBH during the performance of

cycles. In comparison, the powder of bulk MBH was sintered into a single aggregate after three dehydrogenation-hydrogenation cycles (Fig. 4). In Fig. 4*b* some single grains of the compacted pellet obtained after cycling were observed. The experimental results suggest that used  $SiO_2$  aerogel worked as nanoscaffold avoiding aggregation of MBH during hydrogen cycling.

# (FIGURE 3 AND 4)

# 3.2 Coupled calorimetric-manometric measurements.

DSC results obtained with bulk  $\gamma$ -MBH are shown in Fig. 5 (separately in Fig. S1). Obtained results are in good agreement with those previously reported in [6][13][32][8]. Small temperature differences between the results obtained in this work and literature results can be due to different applied conditions of pressure and heating rate during the measurement [32]. According to Soloveichik et al. [6], Li et al. [11] and Saldan [8], the first two endothermic peaks (162 and 194°C) are related to the phase transformation of MBH. Then, in the range of 300-450°C, MBH was decomposed in 3 different steps (D1-D3). During the first two steps (T=307°C and T=361°C, respectively), MBH was decomposed into MgH<sub>2</sub> (crystallized during E1) and amorphous boron. After this two-step reaction, MgH<sub>2</sub> was decomposed into Mg in the third step (T= 384°C) [33][13]. Since MBH–SiO<sub>2</sub> composites preparation was done at 200 °C, the used MBH should be converted to its  $\beta$  modification.

Coupled DSC-manometric measurements were carried out for both bulk  $\gamma$ -MBH and the prepared MBH–SiO<sub>2</sub> composite (Fig. 5). Onset of decomposition for the composite was shifted to lower temperature on approximately 60°C compare to bulk MBH. A similar result was obtained in a previous work [34], in which TiO<sub>2</sub> was used as additive. In this case, the dehydriding temperature was lowered of 50°C. Therefore, in this work silica could also act as additive improving the breakage of B-H bond from MBH which is the limitation of the decomposition process [21], similarly to those reported in [29].

Moreover, Fig. 5 showed that silica changes the mechanism of the reaction. The three decomposition steps from bulk MBH were converted into only one broad endothermic peak and one higher temperature shoulder of the prepared MBH-SiO<sub>2</sub> composite, a typical behavior of confined hydrides [20]. This fact could be related to the intimal contact between silica and MBH. Similar result was observed in the case of MBH-0.5LiH composite [24], only one broad peak located at 210°C, involving complicated interactions which may lead to formation of metastable phases. Also in [35] hydrogen was evolved in only one hydrogen step after the formation of t MBH-ethylendiamine composite.

Since mass of MBH in the prepared composite was 50%, the final hydrogen release was  $\sim$ 6.6 wt% H<sub>2</sub> that was two times lower than for the bulk MBH (almost 12.0 wt% H<sub>2</sub>). The small difference in the final amount that is released till 450 °C could be due to a small instrumental

error during the preparation of the MBH–SiO<sub>2</sub> sample or surface oxidation of MBH before the measurement. Otherwise it could be due to an effect of the SiO<sub>2</sub> matrix. This would mean that silica not only lowered the decomposition temperature but also its presence makes MBH to release more hydrogen. As it is known, MBH has a theoretical content of hydrogen of 14.8 wt%, so higher conditions of temperature (>500°C) are necessary in order to release all of it.

# (FIGURE 5)

# 3.3 Hydrogen desorption at 300°C and 400°C

Figure 6 shows the kinetics curves of hydrogen desorption for the first desorption cycle at 300°C and 400°C for bulk  $\gamma$ -MBH and the prepared MBH–SiO<sub>2</sub> composite.

Hydrogen release kinetics at 300°C was improved in the sample in which silica was present. Different results were obtained respect to the work of Al-Kukhun et al. [17]. In this work, no changes in hydrogen release kinetics from MBH at 300°C were observed after addition of SiO<sub>2</sub>. This fact confirmed our SEM observation that a better contact between MBH and SiO<sub>2</sub> was found in the prepared composite, which was not obtained with the simple mixing employed by Al-Kukhun et al. [17], therefore the used aerogel might be considered as nanoscaffold for MBH Alternatively, it could be also due to some different properties of the silica used in this work and in the work of Al-Kukhun et al. [17], such as acidity, which could be responsible of the weakening of B-H bond and the kinetic improvement during dehydrogenation [18].

Regarding the kinetics at the lowest temperature, it took less than 4 hours to complete desorption from MBH in the composite ( $\sim$ 5 wt% H<sub>2</sub>), whereas around 8 hours were necessary to release the same normalized amount of hydrogen ( $\sim$ 10 wt% H<sub>2</sub>) from bulk MBH. This means that dehydrogenation time was reduced by a factor of 2. Similar results were observed in [10] using TiF<sub>3</sub> and ScCl<sub>3</sub> as additives ball milled with MBH.

At 400°C, the release kinetics was similar in both samples. In the case of the composite, the kinetic was shifted to the left because borohydride was decomposed in a shorter time, as observed in the calorimetric measurements discussed in the previous section. Moreover, a different pathway was followed during the dehydrogenation as it was also reported in DSC measurements. Three different steps were distinguished in bulk MBH whereas only one was seen in the prepared composite.

The different mass of hydrogen that was released taking into account that  $MBH:SiO_2$  sample was prepared in a ratio 50:50 can be explained by the same reasons reported in couple DSC-manometric measurements: a different degree of oxidation, or small errors during the preparation of samples.

#### (FIGURE 6)

#### 3.4 Reversible hydrogen sorption

Figure 7 and 8 show the dehydrogenations at 300°C and 400°C, respectively for several cycles for bulk MBH and the prepared MBH-SiO<sub>2</sub> composite, performing the rehydrogenation at the same conditions of pressure and temperature of 110 bar and 390°C.

In the case that desorption was performed at 300°C (Fig. 7), both samples were successfully rehydrogenated at milder conditions than those reported in previous works. 3.1wt% H<sub>2</sub> was released from bulk MBH after rehydrogenation. A similar amount was reported in [6], which could be associated to the rehydrogenation of MgH<sub>2</sub>, after rehydrogenation of completely decomposed MBH (MgB<sub>2</sub>). However, in this work products after desorption at 300°C (incomplete desorption) were partially rehydrogenated.

In the prepared composite, a lower amount of  $H_2$  was released (2.6 wt%  $H_2$  normalized for the amount of MBH in the sample). This could be due to the formation of more intermediates than in bulk MBH, which were formed during desorption and were more difficult to rehydrogenate, as it was also observed in [10]. Another possibility could be the formation of  $B(OH)_x$  compounds due to the presence of surface hydroxyl groups on silica aerogel particles which might be the reason of the surface deactivation. Thus, the formation of this irreversible compound would explain the lower amount of  $H_2$  released in the second cycle in MBH-SiO<sub>2</sub> composite.

Since RGA for the both samples confirmed mainly hydrogen gas (diborane was suppressed), existence of stable solid borane products might be the preliminary problem of reversible hydrogen sorption in MBH. More analyses could be necessary in order to clarify the different dehydrogenation-rehydrogenation mechanism that was followed in each sample.

Regarding the kinetics, it was observed that the second desorption was slower than the first one for both samples. A sluggish kinetic for both samples suggests the existence of a kinetic barrier that suppresses the fast development of the reaction. The addition of a catalyst could be a solution in order to enhance it.

#### (FIGURE 7)

A different result was obtained in the case that the samples were rehydrogenated after dehydrogenation at 400°C (Fig. 8). After 1-st desorption both samples showed 42 % of reversible hydrogen sorption that resulted in 5.4 wt% H<sub>2</sub> for bulk MBH and 2.9 wt% H<sub>2</sub> for the prepared MBH–SiO<sub>2</sub> composite in practice.

This is the first time that more than 3 wt% has been rehydrogenated at moderate conditions, not only in bulk MBH but also preserving the reversibility in MBH-SiO<sub>2</sub> composite. A similar amount was rehydrogenated by Li et al [11] and [12], but applying a pressure of 400 bar, whereas in this work pressure was reduced to 110 bar. Desorption curves during 2-nd and 3-rd cycles were almost identical for the both sample that suggest a good cyclability. It is visible that complete hydrogen release for both samples was very similar with the only difference of lower kinetics for the composite suggesting activated or deactivated (which is our case) surface of MBH.

After three dehydrogenation-hydrogenation cycles at 400 °C, bulk MBH powder was sintered and converted into a single compacted pellet with the shape of the reactor used, in contrast to the fine powder morphology that was preserved in the case of the prepared MBH–SiO<sub>2</sub> composite. As it was seen in SEM analyses, SiO<sub>2</sub> avoided the aggregation which could be very disadvantageous for the cyclability of the borohydride.

Most probably partial reversible hydrogen sorption at 400 °C might be explained by irreversible products formed during dehydrogenation of both bulk MBH and the prepared MBH–SiO<sub>2</sub> composite.

# (FIGURE 8)

In order to analyze phase composition for the commercial bulk MBH before and after cycling, XRD measurements were performed for MBH as received and after second dehydrogenation at 400 °C (Fig. 9). As it was expected, after dehydrogenation diffraction profile revealed the presence of Mg (74.9%) and some MgO (25.1%) that could be formed due to the presence of some oxygen during the applied measurement or the presence of nanocrystalline MgO already in the commercial MBH. In [15], MgO was also discussed which could form a shell around the reversible  $Mg_xB_yH_z$  phases preventing the full rehydrogenation and therefore might be responsible for the reduced reversibility. In the case that some intermediates were present, they would be amorphous as in the case of pure boron. In the prepared MBH-SiO<sub>2</sub> composite, XRD analyses did not reveal any crystalline phase in the spectra after thermal treatment. Some additional methods to analyze amorphous species during dehydrogenation-rehydrogenation would be helpful to clarify the difference in the reaction mechanism between bulk MBH and the prepared MBH–SiO<sub>2</sub> composite.

# (FIGURE 9)

# 4. Conclusions

Microparticles of silica aerogel were proposed as host to infiltrate MBH using a thermal treatment in hydrogen at 200°C and 120 bar  $H_2$  for 3h. According to the volume of pores of

microparticles of  $SiO_2$  aerogel calculations, it would be possible to infiltrate up to 50 wt% of MBH.

However, as this method was not satisfactory to confine MBH in the pores of the support, less amount of silica could be used in a future work. As a result, a composite in which silica behaves as additive and a storage material with more total  $H_2$  content could be obtained. Wet impregnation is also proposed as future technique to infiltrate MBH in the pores of silica selecting the correct solvent in order to avoid the decomposition of the hydride.

The temperature of the decomposition of bulk MBH was decreased almost 60°C and the mechanism of the decomposition was changed in the prepared composite. Calorimetric measurements showed that the three steps of decomposition of bulk MBH were converted into only one in range of 240-400°C. This result indicates an intimal contact between MBH and silica, which acts as an additive changing the decomposition route, was successfully acquired with this method. Additionally, their kinetics of the first dehydrogenation at 300 °C was two times faster compared to that of the bulk MBH.

Moreover, for the first time MBH has been demonstrated to be partially reversible at moderate conditions of pressure and temperature (110bar H<sub>2</sub> and 390°C). Almost 6wt % H<sub>2</sub> was rehydrogenated when hydrogen desorption was performed at 400°C.

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