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Fast carbon dioxide recycling by reaction with $\gamma$-Mg(BH$_4$)$_2$

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$\gamma$-Mg(BH$_4$)$_2$ was found to be a promising material for CO$_2$ recycling (mainly to formate and alkoxide-like compounds). CO$_2$ conversion occurs with unprecedented fast kinetics at 30°C and 1 bar. A multi-technique approach allowed to ascribe superior performances to its large specific surface area.

Porous complex hydrides are a new class of materials in the panorama of metal hydrides. Among them, gamma phase of Mg(BH$_4$)$_2$ possesses the interesting peculiarity to show a large permanent porosity, accounting for about the 33% of the material volume (see Fig. 1). Its structure is constituted by Mg$^{2+}$ ions in tetrahedral coordination, linked through a shared [BH$_4$]$^-$, giving rise to an highly 3D porous structure characterized by hexagonal overtures with a narrowest dimension of 5.95 Å (geometrical distance). Each hexagon, having a chair conformation, shares every Mg-BH$_4$-Mg side with a vicinal hexagon through a tetrahedral angle. The high surface of this material ($1160$ m$^2$ g$^{-1}$) makes it interesting as nanosponge for gas adsorption and its ability to adsorb large quantities of molecular hydrogen was recently reported.

Carbon dioxide is generally recognized as the most important greenhouse gas. The astonishing increase in CO$_2$ concentration in the terrestrial atmosphere in the past 200 years (from 270 to 399 ppmv, amounting to approximately 900 Gt CO$_2$) and, moreover, its monotonic trend are expected to have important fallout on the climate of the planet. In fact, a linear dependence of the terrestrial temperature on the CO$_2$ concentration has been experimentally determined, estimating an increase of $13.5$°C per 130 ppmv of CO$_2$, with important consequences for example on the sea level. It is then evident the necessity to employ selective sorbers that capture CO$_2$ in order to reduce the anthropogenic emissions, either by molecular adsorption or by dissociative chemisorptions. Among the first class of materials, it was demonstrated that magnesium cations have a larger affinity toward CO$_2$ with respect to other cations. Concerning dissociative chemisorption, magnesium oxide based materials have received strong attention because of the relatively low temperature of decomposition of the corresponding carbonates. The capture of CO$_2$ is only the first step in the carbon dioxide cycle: capture, storage and transportation and underground sequestration. Nevertheless, being the total amount of CO$_2$ released each year in the atmosphere of the order of gigatons, finding an effective way to reuse this molecule as valuable chemical reagent as possible alternative to its sequestration, is a key point in the global warming research. In this perspective, the necessity to identify different CO$_2$ derivative products is required. Unfortunately, the CO$_2$ chemical inertia makes this problem even more challenging.

Fig. 1. $\gamma$-Mg(BH$_4$)$_2$ structure: Mg atoms are represented as dark grey tetrahedra, B atoms as light grey polyhedra and H atoms as spheres. The Connolly surface obtained by using a probe with radius of 1.82 Å is represented.

The open structure of $\gamma$-Mg(BH$_4$)$_2$ and the large concentration of magnesium cations makes it an ideal material to be studied as CO$_2$ concentrator. In fact, the proximity to the cation of hydrogen atoms, negatively charged, gives rise to one Lewis basic-acid couple per Mg atom: these couples are well known to be the most suitable ones to enlarge the affinity of unreactive CO$_2$ toward surfaces. For example, late transition metal (Ni,Rh) hydrides instantaneously catalyse the transformation of CO$_2$ in formiate in the presence of borane. Nickel hydride with a diposphinite-based ligand is an high efficient catalyst for the reduction of CO$_2$ with catecholborane, and the hydrolysis of the...
resulting methoxyboryl species produces CH₃OH in good yield. However, the complex synthesis of these catalysts makes them not applicable to large scale. Methane and ethylene formation for reaction of CO₂ with amalates has been already reported; CO₂ reduction by BH₃NH₃, LiBH₄ and NaBH₄ was studied previously both theoretically and experimentally as a method to enhance the thermolytic dehydrogenation of borohydride compounds or as a cheap method to obtain boron-doped carbons or graphene oxide. It was also reported that sodium and lithium borohydrides react with CO₂ to give valuable chemical products as formic acid, acyloxyborohydrides and formamidoborohydride, depending on reaction conditions (temperature and solvents). The separation of the products from the reagents is possible by simply dissolving the material in a suitable solvent (e.g. water, although this procedure can be safely adopted only after a high degree of CO₂ conversion has been reached to avoid explosions due to the high exothermicity of unreacted borohydrides reaction with water). Nevertheless, the very low reaction rates at RT (1/90 molar ratio of reacted CO₂/BH₄ after 120 h), make unpractical its industrial application.

In this work, the potential of γ-Mg(BH₄)₂ in carbon dioxide recycling are explored for the first time. It is shown as the kinetics of reaction of CO₂ can be enhanced in γ-Mg(BH₄)₂ by several hundreds of time with respect to all previous explored metal hydrides. In order to quantify the CO₂ uptakes, CO₂ isotherms were obtained at 30°C by means of a microbalance, allowing also to investigate the kinetics of the reaction. The effect of the reaction on the structure of γ-Mg(BH₄)₂ were verified by X-Ray powder diffraction (XRPD) and nitrogen adsorption at 77 K whereas the reaction products were identified by means of Fourier Transformed Infrared in attenuated total reflectance spectroscopy (ATR-FTIR) in the medium infrared, coupled with Far-IR investigation and ¹³C CP-MAS NMR spectroscopy. Moreover, an in situ study of the sorption process was performed by FTIR spectroscopy in transmission, in order to get more insights into the reaction mechanism. All the measurements were replicated on the unporous polymorph α-Mg(BH₄)₂ (4 m² g⁻¹, see Table S1), in order to estimate the importance of the surface/microporosity on the kinetics of the CO₂ conversion.

The reaction kinetics were evaluated by following the CO₂ uptake as a function of time by means of microgravimetric measurements at 30°C and 1 bar, as shown in Fig. 2a. The γ-Mg(BH₄)₂ phase reaches the impressive value of 1:11 molar ratio of reacted CO₂ for [BH₄]¹⁻ unit after only 10 min. The effect of the surface area on the reaction kinetic is well evident by comparing the data obtained for γ-Mg(BH₄)₂ with those for α-Mg(BH₄)₂ (black and grey curve, respectively). The absence of highly reactive surface defects on both the Mg(BH₄)₂ surfaces has been confirmed by an infrared study of CO adsorbed at 100 K (results not shown). Fig. 2b compares the CO₂ isotherms obtained at 30°C for both γ and α phases of Mg(BH₄)₂. Each point of the isotherms was recorded after a fixed equilibration time of 4 h; without fixing the time, the equilibration procedure would protract further especially at low pressures. In fact, a plateau value of 12 mol/kg (34.5 mass%) was obtained for the γ phase at 1 bar only after 7 days (data not shown). This value is slightly larger than the highest value reported so far at RT and 1 bar for microporous materials (27.5 mass% in the metal organic frameworks Mg₂(dobdc) and HKUST-1), where CO₂ is molecularly adsorbed in the material pores. Nevertheless, this value is about one half of that reported for the most performing oxidative systems, although in that case the formation of carbonates and bicarbonates species requires to adopt very high temperatures (e.g. CaO/Al₂O₃, 68 mass% at 1 bar and 600°C). The uptake is completely irreversible, as demonstrated by the second isotherm, indicating that almost the whole material was reacted at the end of the first absorption isotherm. Also in this case, the key-role of the surface area is evident by comparison with the data collected on the α phase.

These results were confirmed by XRPD characterization. Fig. 3 shows the XRPD patterns of γ-Mg(BH₄)₂ before (γ) and after (γ') the reaction with CO₂. Most of the diffraction peaks present in the pattern of γ disappear in the pattern of γ', suggesting the complete collapse of the pores structure and the formation of an
amorphous phase. The few diffraction peaks still visible are due to a fraction of α phase present as impurity in the starting material (amounting to 16% in volume estimated by Rietveld refinement, see ESI). On the contrary, the crystalline structure of α-Mg(BH₄)₂ is almost unaffected (α') by the interaction with CO₂ as expected on the basis of the lower CO₂ uptake (see Fig. 2b). In both cases, after the reaction a broad peak is observed at around 2θ = 13.6°. It is interesting to notice that a signal in that region, although broader, as well as the formation of an amorphous phase was also observed after thermal dehydrogenation of γ-Mg(BH₄)₂. The structure collapse of γ-Mg(BH₄)₂ is confirmed also by nitrogen volumetry, showing a decrease of the surface area of more than two order of magnitude, after reaction with CO₂ (see Table S1 and Electronic Supplementary Information for further details and comments).

![Fig. 3. XRPD patterns of γ-Mg(BH₄)₂ and α-Mg(BH₄)₂ before (black curves) and after CO₂ interaction at 30°C and 1 bar for 3 days (grey curves).](image)

**13C CP-MAS NMR and ATR-IR spectroscopies allowed the identification of the new species formed upon CO₂ sorption.** Fig. 4a shows the ATR-IR spectra of the γ-Mg(BH₄)₂ material before and after CO₂ reaction. The IR spectrum of the starting material is quite simple and dominated by the absorption bands related to BH₄ units, at about 2270 cm⁻¹ (ν(BH₄)) and 1258-1126 cm⁻¹ (δ(BH₄)). After prolonged interaction with CO₂ the IR spectrum becomes more complex. The absorption bands characteristic of BH₄ decrease in intensity, although they do not disappear. This observation suggests that the reaction with CO₂ proceeds through a dehydrogenation of γ-Mg(BH₄)₂, but involves only a fraction of the BH moieties (in agreement with the CO₂ sorption measurements). Simultaneously, new IR absorption bands appear, providing an evidence that new chemical species are formed. Although a detailed assignment of all the absorption bands is not straightforward, some of them clearly reveal the formation of formate (-OOCCH₃ band at 1622 cm⁻¹ due to ν(C=O) mode) and methoxy species (-OCH₃ bands in the 3000-2800 cm⁻¹ region due to ν(CH₃) and around 1100 cm⁻¹ due to ν(C-O)). Finally, the broad absorption band centred around 1300 cm⁻¹ are assigned to ν(BO) mode. The Far-IR spectra of γ-Mg(BH₄)₂ and α-Mg(BH₄)₂ phases before (α and γ) and after (α' and γ') the reaction with CO₂ are shown in Fig. 5, compared with that of MgO (taken here as a reference for ν(Mg-O) vibrations). Both of them are characterized by two sets of absorption bands. The band centered around 425 cm⁻¹ (with a shoulder at 377 cm⁻¹) is due to ν(Mg-B) whereas the couple of bands centered around 215 cm⁻¹ are assigned to δ(B-Mg-B) vibrational modes. The two δ(B-Mg-B) bands have a different intensity ratio in the spectra of the two polymorphs. The Far-IR spectrum of α-Mg(BH₄)₂ does not change after reaction with CO₂ in agreement with all the results shown above. On the contrary, the spectrum of γ' is characterized by the almost total disappearance of the absorption bands due to δ(B-Mg-B) modes; the remaining couple of bands are due to the presence of a small fraction of the α phase, which is unaffected by the reaction with CO₂, in agreement with XRD data. Moreover, in the spectrum of γ' the ν(Mg-B) absorption band slightly shifts at lower frequency, accompanied by the appearance of a broad shoulder. All these observations, coupled with the fact that no hydroxyl species (3600 cm⁻¹ region) are formed, indicate that most of the reaction products are chemically bounded to B atoms. No evolution of gaseous reaction products (e.g. methane) was evidenced in these temperature and pressure conditions (see Fig. S5).
In order to elucidate the reaction mechanism, the reaction with CO₂ was monitored at room temperature by means of in situ FT-IR spectroscopy on samples in the form of self-supporting pellets. Fig. 4b) and c) show the background subtracted sequence of spectra collected at regular time intervals, in the two most relevant wavenumber regions; the spectra in the whole region are reported in the ESL. They demonstrate that formate (band at 1622 cm⁻¹) and methoxy (bands in the 3000-2800 cm⁻¹ region) species are formed simultaneously since the beginning of the reaction at the expenses of BH₄⁻ (bands at 2270 cm⁻¹), and no other reaction intermediates are detected. This is an important difference with respect to what reported for aminoborane reaction with CO₂, where an evolution of the products with time toward compounds with lower valence of carbon is observed.²⁰

Analogous information was obtained by NMR spectroscopy. ¹³C CP MAS spectrum (Fig. 6) is characterized by peaks at 170.7, 85.7, 50.2 and around 33-13 ppm, (33.3, 19.3 and 13.8 ppm), which are characteristics of C=O, OC-H, OCH₃ and aliphatic groups, respectively. This is in agreement with previous results on aminoborane.²⁰ Beside that, NMR allows also to quantify the relative abundance of the reaction products. Interestingly, it is found as the methoxy species (large peak at 50.2 ppm) are the most abundant followed by C=O groups.

In conclusions, the whole set of data discussed above demonstrate that the γ-Mg(BH₄)₂ material efficiently reacts with large amount of CO₂ converting it into valuable chemical products (among which formate and methoxy species). The reaction products can be easily recovered by simple dissolution of the material in water.²⁵ Fast kinetics of reaction were observed, unexpected on the basis of previous results reported for metal hydrides and ascribable to the large surface area of this material. This clearly indicates that this new class of porous metal hydrides are promising materials for CO₂ recycling. The mechanism involves a partial de-hydrogenation of the starting material. The maximum CO₂ uptake, estimated by very prolonged contact time (7 days at 1 bar and 30°C), is about 12 mol/kg. This value is lower than the theoretical one calculated by assuming a 1:1 reaction between CO₂ and BH₄⁻ (37.0 mol/kg, corresponding to 62 mass%), but it is very close to that expected by considering the completeness of the first step in the dehydrogenation of γ-Mg(BH₄)₂ (12.3 mol/kg, corresponding to 35 mass%).²² This first step is characterized by an almost complete reversibility.²³, ²⁴ In this respect, the results shown herein acquire interesting perspectives: if re-hydrogenable, γ-Mg(BH₄)₂ could be used as catalyst for CO₂ transformation into valuable organic products instead as reagent. Preliminary attempts in this direction were at the moment unsuccessful for both performing the reaction in a CO₂/H₂ atmosphere or by exposing the sample to 20 bar of H₂ at 250°C after the reaction, likely because of the higher CO₂ interaction energy with respect to H₂ and the large stability of the B-O bond. Nevertheless, the very fast kinetics associated to the reaction would provide to this hydride the peculiar characteristics to be able to couple in one step the separation (in post combustion systems) and recycling of CO₂ in a wide temperature range (RT < T <190°C) allowing to avoid the storage and transportation steps. The possibility to use γ-Mg(BH₄)₂ as co-reagents in other CO₂-based reactions would enlarge the number of obtainable products and will be the subject of future investigations.

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Notes and references

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²³ † Electronic Supplementary Information (ESI) available: samples and experimental methods details, in situ FTIR in transmission of CO₂ reaction with Mg(BH₄)₂ samples, Nitrogen adsorption measurements and Rietveld refinements results of the materials before and after reaction with CO₂, IR spectrum of the gas phase composition after CO₂ reaction with γ-Mg(BH₄)₂ for 3 days. See DOI: 10.1039/b000000x/

Graphical abstract

\(\gamma\)-Mg(BH\(_4\))\(_2\) was found to be a promising material for CO\(_2\) recycling (mainly to formate and alkoxide-like compounds) with very fast kinetics because of its very large surface area.