

Consequences of MgO activation procedures on its catalytic performance for acetone self-condensation

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ABSTRACT

A new procedure for increasing surface area and basicity of MgO has been developed in this work, leading to a tailored catalyst for the self-condensation of acetone. The preparation method, consisting on the hydration/dehydration of periclase, was optimized varying the activation procedures. The effect of the precursor (magnesium oxide or magnesium carbonate) and the thermal treatment on the morphological properties and the basic sites distribution was characterised. In particular, CO₂ calorimetry data attributed the increase of basic sites mainly to the increase in surface area. The material was tested for the base-catalyzed self-condensation of acetone, obtaining high conversions and selectivities towards the products of most industrial value, the isophorones. The selectivity reached to trimers at 723 K was almost 50 %, and towards isophorones, of more than 30 %.

Keywords: *basic materials; brucite-periclase transitions; CO₂ adsorption; hydromagnesite; acetone condensation; biomass upgrading*

1. INTRODUCTION

Basecatalysis plays a key role in the upgrading of biomass-derived platform molecules. Several base-catalysed reactions, such as ketonic or aldol condensations, lead to concurrent carbon chain growth and oxygen removal of oxygenates [1]. Aldol condensation consists of the condensation of molecules with two carbonyl groups yielding hydroxyl-ketones or unsaturated ketones [2], and it is a key reaction for biomass feedstock upgrading [3]. Among the reactants that can undergo this reaction, an increasing attention is being paid to acetone. In fact, although acetone is nowadays mainly produced from fossil stocks, new technologies based on chemical and biological processes are emerging: biomass pyrolysis [4] or the ABE fermentation of agricultural wastes [5]. Thus, acetone self-condensation becomes in an interesting reaction for obtaining chemicals by itself (mainly isophorones) or in combination with hydrogenation steps (methyl-isobutyl ketone, MIBK).

Gas phase self-condensation of acetone yields to isophorones (cyclic trimers) as the most valuable product, with applications in paints, pesticides, adhesives, resins, etc., whereas liquid phase condensations (usually performed in presence of H_2) are mainly devoted to manufacturing methyl-isobutyl ketone (MIBK). Different basic catalysts have been proposed for this condensation, such as niobium oxides [6], Mg-Al mixed oxides [7,8], Mg-Zr mixed oxides and Mg-Zr oxides dispersed on carbonaceous materials to increase the exposed surface area [9] and even MgO, non-promoted and promoted with alkaline or alkaline-earth metal ions [10]. Among all these catalysts, bulk MgO is the cheapest alternative and very attractive due to its preparation easiness. However, magnesium oxide presents an important drawback in comparison to other common catalysts, its low surface area. Furthermore, the selectivity towards isophorones is

reduced in comparison to, for example, Li/MgO [10]. This effect was attributed to the low concentration of high-strength basic sites in the non-promoted catalyst.

Commercial MgO has low surface areas (typically, around 20 m²/g) and also low concentration of basic sites (around 0.4 mmol/g). Consequently, many methodologies for the synthesis of MgO with high surface area and higher concentration of exposed basic sites have been developed. Bhagiyalakshmi et al. [11] reported a method for synthesizing mesoporous magnesium oxide from a commercial MgO that enhances the specific surface area of the starting material up to 250 m²/g. In addition, the CO₂ adsorption capacity, an indicator of the surface basicity, increases from 0.45 to 2.3 mmol/g at 373 K, being this value the highest ever reported for magnesium oxide. The drawback of the aforementioned preparation method stems from its complexity, since it requires the synthesis of SBA-15 and CMK-3, which involves steps as pyrolysis at 1173 K and the use of acids such as sulfuric or hydrofluoric. Other synthesis method proposed is the sol-gel [12,13], which involves a multi-step decomposition process of a precursor in the presence of an expensive template. Surface areas in the range of 250-650 m²/g have been reached [14], although the template should be removed in subsequent steps. In an attempt to avoid the use of templates and toxic solvents, Bian et al. [15] described a procedure to synthesize mesoporous MgO with a nanocrystalline framework via the thermal decomposition of magnesium acetate, obtaining surface areas around 120-136 m²/g. Another important synthetic route for the preparation of high surface area MgO involves thermal treatment of Mg(OH)₂, obtained either by precipitation or by MgO hydration, leading to surface areas ranging 100-300 m²/g [14]. This method is remarkable for its simplicity and reduced cost, which makes the MgO so formed a plausible inexpensive catalyst. Furthermore, in a recent work, Bartley et al.

[16] have obtained a high surface area MgO by thermal procedures with different morphologies and surface properties depending on the precursor and thermal conditions.

In the present work, the thermal treatment method will be considered according to its proven compromise between surface area enhancement and environmental benignity. Two precursors were used for the preparation of Mg(OH)₂, namely, commercial MgO and MgCO₃. Their thermal decomposition (temperature and time) into the final MgO was optimized taking into account the CO₂ adsorption capacity, as measurement of both their surface basicity and potential as CO₂ adsorbents. Morphological aspects of the catalysts were characterized by temperature-programmed XRD and N₂ physisorption. Since Mg(OH)₂ dehydration is believed to be a key step, transitions brucite-periclase were analyzed in detail by means of XRD scans recorded at increasing temperatures, as well as a function of time at the decomposition temperature. The number and strength of surface basic sites on the catalysts were studied by microcalorimetry and FTIR. Finally, the gas phase acetone condensation was used as model reaction of basic catalyst. It is shown that a simple catalyst with an appropriate morphology and high concentration of basic sites can be obtained.

2. MATERIALS AND METHODS

2. 1. Materials preparation

Increased surface area magnesium oxides were prepared from two different precursors, namely: commercial magnesium oxide (Panreac) and magnesium carbonate (Fluka).

Magnesium carbonate was firstly decomposed into magnesium oxide by thermal treatment in a tubular (23 mm i.d.) fixed bed reactor from room temperature to 973 K at

a rate of 5 K/min and then maintained the final temperature for 3 h under synthetic air flow (in order to ensure the absence of CO₂ and water). Both magnesium oxides - commercial precursor and obtained from decomposition of magnesium carbonate- were converted into magnesium hydroxide by slow addition of distilled water at room temperature under stirring, in the ratio of 10 mL of water per gram of magnesium oxide, according to the procedure outlined in [10]. The temperature was subsequently raised to 353 K and stirring was maintained for 4 h. Finally, the precipitate was dried overnight in an oven at 358 K. Magnesium oxide was re-synthesized by thermal decomposition of magnesium hydroxide under air flow in two steps: sample was first treated at 623 K for 5 h and then at 773 K during 8 h. The temperature programme for the decomposition of Mg(OH)₂ was optimized targeting the largest CO₂ uptake. For that, analogous analyses were performed using a thermogravimetry-differential scanning calorimetry (TG-DSC) instrument (Setaram, Sensys).

Samples considered in the present work were designed as follows: MgO(I) for the commercial magnesium oxide, MgO(II) for the magnesium oxide obtained from decomposition of the magnesium carbonate, and hsMgO(I) and hsMgO(II) for the samples resulting after the hydration and thermal treatment from MgO(I) and MgO(II), respectively (Scheme 1).

2. 2. Materials characterization

The textural properties of the samples were determined by N₂ adsorption-desorption isotherms at 77 K in a Micromeritics ASAP 2020. Prior to the analysis, the samples were outgassed at vacuum conditions (< 10⁻³ kPa) at 473 K for 4 h. The surface area (S_{BET}) was calculated according to the Brunauer-Emmett-Teller (BET) method, whereas

the pore size distributions and total pore volumes (V_p), corresponding to the cumulative pore volumes between 170 and 30000 nm, were determined from the desorption branch using the Barrett-Joyner-Halenda (BJH) method. Analysis by scanning electron microscopy (SEM) was done using a JeolJSM-6100 microscope; the samples were deposited on a standard aluminium SEM holder and gold-coated.

The crystalline structure of the samples was studied by XRD using a Philips X'Pert Pro powder diffractometer equipped with a $\text{CuK}\alpha$ radiation source (0.15418 nm) and operating in a 2θ range of 5-85° at a scanning rate of 0.02°/s. The X-ray tube voltage and current were set at 45 kV and 40 mA, respectively. Additional XRD experiments were carried out in a Seifert XRD 3000 diffractometer, which is equipped with a temperature controlled chamber, in order to follow the brucite-periclase transition. Magnesium hydroxide samples were subjected to a temperature programme from 298 up to 773 K at a heating rate of 5 K/min, while 25 scans were recorded. Once determined the transition temperature, the evolution of the structure with time at that constant temperature was studied by recording 30 consecutive scans on fresh hydroxide samples. The equipment settings were analogous to the above described.

2. 3. CO₂ adsorption

In order to quantitatively establish the CO₂ adsorption capacity, and consequently the total concentration of basic sites, calorimetric experiments were carried out in a Tian-Calvet microcalorimeter (C80, Setaram) linked to a volumetric line. The adsorption of CO₂ took place by repeatedly introducing small doses of gas onto the initially outgassed solid until a final equilibrium pressure of 6.7 kPa. Before adsorption, about 80 mg of sample were outgassed overnight at 623 K under vacuum (10^{-6} kPa). The equilibrium

pressure was measured after each dose by a Barocel Capacitance Manometer (Datametrix). The adsorption temperature was maintained at a constant value.

In order to gain further understanding on the strength of the surface basic sites, the chemical structure of the chemisorbed CO₂ was characterised by Fourier Transform Infrared Spectroscopy (FT-IR). Data were obtained using a Bruker Vector 22 FT-IR spectrophotometer (DTGS detector), in the 4000-400 cm⁻¹ range, with a resolution of 2 cm⁻¹ and using 100 scans. An inverted T-shaped Pyrex cell containing the sample pellet was used. The two ends of the short arm of the T were fitted with CaF₂ windows. The self-supporting wafer (20 mg, 18 mm diameter) was first pre-treated in situ at 573 K in an oxygen flow overnight in order to remove surface substances, then evacuated at the same temperature for 2 h and finally exposed to CO₂ (1 kPa) at room temperature for 5 min. Spectra were taken at room temperature after CO₂ adsorption and sequential evacuation for 20 min at room temperature, 323, 373, 473, 573, and 673 K. The background spectrum was measured before adsorption and used to correct the others spectra by absorbance subtraction.

2.4. Reaction studies

The reaction was performed in a 0.4 cm i.d. U-shaped fixed bed quartz reactor placed in a PID-controlled electric furnace. The catalyst (150 mg, 50-80 µm) was placed over a plug of quartz wool and a thermocouple was placed inside the catalyst bed. A gas flow of 0.05 L•min⁻¹ (s.t.p.) of helium- acetone mixture (3.2 vol% acetone) was fed. Acetone was injected as a liquid by a syringe pump in a He flow and vaporized in situ. Reactor outgoing gases were analyzed on-line by gas chromatography using a HP6890 GC, equipped with a FID detector and using a CP-Sil 5 CB capillary column as stationary phase. Product identification was confirmed by analysis in a

GC-MS (Shimadzu QP-2010), using the same column and conditions that in the gas chromatograph.

Conversions were calculated from the acetone concentration at the reactor inlet and outlet, whereas selectivities were calculated as the ratio between the concentration of each reaction product and the sum of the concentration of all the reaction products. Carbon balance was checked by comparing the total amount of carbon atoms at the reactor inlet and outlet.

3. RESULTS AND DISCUSSION

3.1. Materials characterisation

3.1.1. Crystallographic characterisation

X-Ray diffraction patterns of the samples, Fig. 1, showed the typical diffractograms of periclase (JCPDS 45-946), with peaks at $\sim 37^\circ$, 43° , 62° , 75° and 79° , which correspond to diffraction by planes (111), (200), (220), (311) and (222), respectively. Phases corresponding to the precursors were not detected. Sample MgO(I), the commercial magnesium oxide, presented peaks considerably sharper and more intense, indicating a markedly more developed crystallinity, which is in agreement with the considerable lower specific surface area exhibited by this sample. The crystallite size of the magnesium oxides calculated from the plane (200) using the Debye-Scherrer equation[17] are included in Table 1.

Brucite-periclase transitions were studied by temperature-programmed XRD, being the diffractograms recorded at different temperatures depicted in Fig. 2. The diffractograms recorded at 298 K correspond to the magnesium hydroxides obtained from the parent magnesium oxides. In the case of magnesium hydroxide resulting from MgO(I) (Fig. 2a), the peaks at $\sim 19^\circ$, 33° , 38° , 51° , 59° , 62° , 68° , 72° and 81° are characteristic of

brucite phase (JCPDS 7-239) and are related to planes (001), (100), (101), (102), (110), (111), (103), (201) and (202), respectively. Nevertheless, in the diffraction pattern corresponding to the sample obtained from hydration of MgO(II) (Fig. 2b), the presence of a second phase, hydromagnesite ($\text{Mg}_5(\text{CO}_3)_4(\text{OH})_2 \cdot 4\text{H}_2\text{O}$) is evidenced. This phase is unambiguously identified according to the peaks at $\sim 10^\circ$, 14° , 15° , 20° , 21° , 23° , 25° , 27° , 28° , 31° , 33° , 36° , 41° , 42° and 45° , which correspond to diffraction by planes (100), (-110), (011), (020), (-102), (002), (012), (-221), (211), (310), (-321), (230), (023), (113) and (-423), respectively (JCPDS 25-513). This phase is supposed to remain from magnesium carbonate used as precursor. From Fig. 2b, hydromagnesite vanished above 566 K, giving rise to a sole phase of brucite. In both cases, the increase in temperature led to a transition from brucite to periclase, which was patent above 608 K. With the aim of studying the kinetic underlying the phase transition, additional sets of scans were recorded at 608 K at increasing times. A first order kinetics was established for the transformation of brucite into periclase from the decrease with time of the peaks at 38° , assumed proportional to the concentration of brucite phase in the sample. The kinetic data are summarized in Fig. 3, where the natural logarithm of the ratio between the area under the peak at a given time, and this area at initial time is plotted versus time. The kinetic constant, obtained from the slope of the line, for sample I ($k_{\text{I}} = 6.8 \cdot 10^{-4} \text{ s}^{-1}$) was found to be more than twice the value of the constant for sample II ($k_{\text{II}} = 3.0 \cdot 10^{-4} \text{ s}^{-1}$), which is consistent with the fact that the complete transition was reached in half of the time for sample I. Thus, dehydroxylation is favoured in the pure magnesium oxide, whereas the carbonate remaining in the MgO(II) sample hinders the process, decreasing in this way its transformation velocity. The higher transformation rate observed for sample MgO (I)

also lead to the formation of metastable species and larger amount of crystalline defects [19], usually considered as responsible of basic sites.

3.1.2. Morphological characterisation

Adsorption isotherms for the magnesium oxides considered in the present work are shown in Fig. 4, whereas the textural properties of the samples are summarized in Table 1. With the exception of MgO(I), which exhibits an isotherm type II, the other isotherms correspond to the type IV (mesoporous solids), according to the IUPAC classification. The hysteresis loops are rather narrow and, in the case of hsMgO(I) and hsMgO(II), vertically oriented; whereas for MgO(II), it remains nearly horizontal and parallel over a wide range of p/p^0 , suggesting a change toward a slit-shaped porosity upon activation. In the case of MgO(I), the slight hysteresis loop could correspond to condensation of adsorbate in interparticle macropores.

Comparing the samples obtained after hydration and thermal treatment with their respective precursors, an increase in the specific surface area can be appreciated, as well as in the pore volume, being especially significant in the case of the commercial sample, where the treatment leads to a development of the BET surface area from 26 to 221 m^2/g and of the pore volume, reaching 1.15 cm^3/g . Evacuation of water formed as a product of $\text{Mg}(\text{OH})_2$ decomposition could be the responsible of these expansion in the structure.

SEM images of the samples under study are shown in Fig. 5. It is observed that MgO(I) and MgO(II) exhibit a compact structure, especially in the case of MgO(I). On the contrary, hsMgO samples present a more spongy morphology. All these facts are in agreement with surface area measurements, since hsMgO(I) presents the highest S_{BET} .

This high surface area of hsMgO(I) sample can be also explained according to its crystallographic transformation. It was reported that in the formation of periclase, by dehydroxylation of brucite, structural oxygen vacancies are formed by the removal of water [19]. This generation of structural defects becomes more important when the process is faster. Accordingly, hsMgO(I) sample should have a higher amount of morphological irregularities and therefore a larger surface area of .

3.2. Surface chemistry

3.2.1. Calorimetric studies of CO₂ adsorption

Equilibrium adsorption isotherms at 0- 6.7 kPa and 323 K were recorded by microcalorimetry, Fig. 6. The shape of these curves suggests the presence of combined chemisorption and physisorption mechanisms. It can be noticed that a larger uptake of CO₂ was reached by sample hsMgO(I) -1.2 mmol/g (52.8 mg/g)-, followed by hsMgO(II) -0.82 mmol/g (36.1 mg/g)-, confirming the suitability of the modification method. These values of capacity of adsorption are similar to Mg-Al mixed oxides used also in aldol condensation reactions [20], and considerable higher than zirconia catalysts doped with sodium, used in the ethanol coupling to 1-butanol reaction [21]. Considering the adsorption capacities in surface area basis (5.4 $\mu\text{mol}/\text{m}^2$ for hsMgO (I) vs. 9.4 for hsMgO(II)), it is clear that the higher global concentration of basic sites in sample hsMgO (I) is mainly caused by the increase of the surface area.

Adsorption microcalorimetry is a very useful technique for the characterization of basic properties of solids, not only the basic sites concentration but also their strength distribution [22]. From the distribution of basic sites (Fig. 7), heterogeneity of

the strength of the basic sites observed. These sites can be separated in three groups according to their strength: strong ($Q > 150$ kJ/mol), medium (120 kJ/mol $< Q < 150$ kJ/mol) and weak ($Q < 90$ kJ/mol)[23,24]. Fig. 7 also shows the distribution of the strength of the basic sites. It is noticed that MgO(I) and hsMgO(I) do not present the strongest sites, contrary to the samples obtained from magnesium carbonate. The fast transformation of brucite into periclase would hinder the formation of the strongest basic sites. These observations are in agreement with the kinetic of the CO₂ desorption studied for hsMgO(I) and hsMgO(II) by TPD profiles obtained by thermogravimetry [18]. Deconvolution of the curve revealed two adsorption sites for hsMgO(I), with activation energies of desorption of 56 and 111 kJ/mol; and three for hsMgO(II), with activation energies of desorption of 62, 85 and 142 kJ/mol [18]. The adsorption sites with activation energies lower than 150 kJ/mol were already observed in magnesium derived oxides, more concretely Mg-Al and Mg-Fe mixed oxides [20,25].

3.2.2. Spectroscopic characterisation of the adsorbed species (FTIR)

In order to get further information about the basic sites of the magnesium oxides, infrared spectroscopic studies of the surface of the samples after CO₂ adsorption and desorption at temperatures up to 673 K were performed (Fig. S1). Analysis of the spectra provides useful information on the nature of the species formed upon CO₂ adsorption, complementing the conclusions drawn from microcalorimetry about the strength and relative amount of basic sites.

FTIR spectra in the 900-1900 cm⁻¹ region of the samples after CO₂ adsorption at room temperature, and subsequent desorption under vacuum at temperatures up to 673 K were recorded. Computer decomposition of the spectra, evaluation of the resistance to

evacuation and the spectral parameter $\Delta\nu_3$ have been used for the assignment of the species modes. The splitting, $\Delta\nu_3$, can be considered as a measure of the strength of the base sites: the lower the splitting, the stronger the basic site [12]. For $\Delta\nu_3$ values of about 100, 300 and 400 cm^{-1} are generally proposed for unidentate, chelating bidentate and bridging bidentate structures, respectively [26]. Unidentate species are formed on oxygen ions showing the lowest coordination number (O^{2-} , strong base sites), whereas chelating and bridging bidentate carbonates require the participation of an adjacent cationic site ($\text{M}^{n+}\text{-O}^{2-}$ pairs, medium strength base sites) [27] (Scheme 2).

Table 2 summarizes the assigned bands for each species in the four samples after CO_2 adsorption at room temperature, as well as the area of the decomposed peaks corresponding to each species, which is proportional to the concentration of each type of basic site present in the samples, and the $\Delta\nu_3$ values. Bands at around 1685, 1460 and 1220 cm^{-1} can be assigned to asymmetric O-C-O stretching vibration ($\text{asym}\nu_{\text{O-C-O}}$), symmetric O-C-O stretching vibration ($\text{sym}\nu_{\text{O-C-O}}$) and C-OH bending mode (δ_{COH}), respectively, of bicarbonate. Pairs of bands at 1633-1640 ($\text{asym}\nu_{\text{O-C-O}}$)/1381-1386 ($\text{sym}\nu_{\text{O-C-O}}$), 1590-1603 ($\text{asym}\nu_{\text{O-C-O}}$)/1406-1409 ($\text{sym}\nu_{\text{O-C-O}}$) and 1515-1541 ($\text{asym}\nu_{\text{O-C-O}}$)/1422-1432 ($\text{sym}\nu_{\text{O-C-O}}$) cm^{-1} , with $\Delta\nu_3$ between 254, 184 and 85-119 cm^{-1} , respectively, may correspond to two different unidentate structures. The first value of $\Delta\nu_3$ is higher than the above reported for unidentate species, nevertheless the similar thermal stability displayed by the three pairs of bands suggests that they could be originated by analogous species.

Bands around 1685 and 1660 belong to the asymmetric $\nu_{\text{O-C-O}}$ vibrations of chelating bidentate carbonates, whereas the bands around 1310 and 1350 are attributed to their $\text{sym}\nu_{\text{O-C-O}}$ modes. Finally, bands around 1730 and 1705 correspond to the asymmetric $\nu_{\text{O-C-O}}$.

C-O vibrations of bridging bidentate carbonates, while the bands around 1220 and 1280 are attributed to their sym $\nu_{\text{O-C-O}}$ modes. $\Delta\nu_3$ of 312-379 and 423-507 displayed by chelating and bridging bidentate carbonates, respectively, are in relative good agreement with the above proposed.

Regarding the thermal stability of the adsorbed species, bridging bidentate carbonates and bicarbonates are formed on the weakest basic sites, since they disappear above 323 and 373 K, respectively. On the other hand, chelating bidentate and unidentate carbonates are still present at 573 K. Nevertheless, while for the samples MgO(I) and hsMgO(I) no peaks were detected in the last scan, for the samples MgO(II) and hsMgO(II) slight evidences of remaining unidentate carbonates are appreciated at 673 K, which indicates the presence of stronger sites in these samples compared with MgO(I) and hsMgO(I). Therefore, unidentate carbonates correspond to the strongest basic sites found by microcalorimetry, chelating bidentate corresponds to medium strength sites, and bridging bidentate and bicarbonates are the weakest ones. The unidentate carbonates are responsible of the irreversibility of the CO₂ adsorption process [20], but the role of these sites in the catalytic reaction is still unclear.

3.3. Reaction studies

The gas phase condensation of acetone was performed over the four catalysts at temperatures between 373 and 723 K. The presence of diffusional limitations was ruled out based on the Thiele modulus modified by Weisz [28]. In a previous work [8], it was checked the negligible conversion at these conditions in absence of catalyst. According to the reaction pathway published in the literature [6,29], the condensation reaction of acetone leads to diacetone alcohol, which after a dehydration reaction forms either

isomesityl oxide or mesityl oxide. The formation of these C₆ compounds requires the presence of basic sites [30]. Further condensation of these C₆ molecules with acetone form phorones (C₉), which rearrange to yield cyclic C₉ compounds called isophorones (the most valuable product).

The results of conversion for the four catalysts are shown in Fig. 8, whereas conversion, carbon balance and selectivities for the different products at 523 and 723 K are shown in Table 3. It should be noted that the formation of heavier reaction products, which can lead to catalyst deactivation in experiments performed at larger reaction times, lead to poorer carbon balance closures in the experiments performed at the highest temperature.

Acetone conversion increases with the temperature, reaching more than 37 % of conversion at 723 K for the hsMgO(I) sample. Concerning the selectivities, at temperatures until 573 K, C₆ compounds were the main products, whereas at higher temperatures the trimers increase their concentration. The highest selectivity towards isophorones (IP) were 32.8 %, and for mesitylene (M), 20.7 % at 723 K for hsMgO(I). These values are higher than those obtained previously under similar conditions: 13.9 (IP) and 3.2 (M) % at 723 K, in the case of Mg-Al oxides [8]; and 18 (IP) and 10 (M) % at 575 K for Mg-Zr oxides [9]. What is more, comparing MgO results with those obtained for Mg-Zr oxides dispersed on a high surface area graphite, it is observed that selectivities towards C₉ compounds are slightly higher (53.8 for MgO versus 42 % for Mg-Zr supported catalyst). Although the conversion is lower (32.8 versus 54 %) [9], MgO catalyst is much cheaper.

The conversion obtained for the studied catalysts can be directly correlated to the surface area of the material, increasing the conversion with the surface area, and also to the concentration of basic sites (increasing surface areas imply higher concentration of

active sites). This effect was already observed in the MPV reaction over MgO oxides [16]. It is remarkable that these relationships are not linear, but two marked trends are observed depending on the starting material of the catalysts, observing two similar trends for the correlation between surface area and basic sites with the conversion obtained at 723 K (Fig. 9 a and b). This analogy between MgO(I)/hsMgO(I) and MgO(II)/hsMgO(II) can be related both to the absence of the strongest basic sites in the MgO(I)-based samples, but also to the similar ratio between weak and medium strength basic sites: MgO(I) and hsMgO(I), 0.34 and 0.40, respectively; whereas for MgO(II) and hsMgO(II), 1.5 and 0.9, respectively.

In order to study the influence of morphology and surface chemistry on catalyst performance, results obtained can be analyzed as function of temperature. At low temperatures (up to 523 K), only C6 are formed in relevant concentrations, except for the hsMgO(I). This catalyst exhibits the highest activity and C9 products are already present. For the other catalysts, diacetone alcohol follows the behaviour typical of primary products, which decompose with the increase of temperature in other compounds. Concerning the mesityl oxide and isomesityl oxide, the first one is the main product. The diacetone alcohol formation requires medium strength basic sites (aldol condensation reaction), whereas the further dehydration needs both acid-base pairs, thus add to medium, also weak basic sites favour this reaction [9]. At this point, there is a remarkable difference with the behaviour of Mg-Zr mixed oxides, the formation of diacetone alcohol as non-reactive side product being observed.

At higher temperatures, lower selectivities towards C6 compounds are observed. At 723 K, the concentration of C6 compounds is enriched in isomesityl oxide. This result can be attributed to the presence of two different active sites for the diacetone dehydration, yielding either mesityl oxide or isomesityl oxide. Only the former is able to follow the

reaction toward trimers [9]. The samples MgO(II) and hsMgO(II), with lower concentration of weak basic sites, have also lower concentration of acid-base pairs, thus being limited in this step.

Concerning the formation of trimers, the first compound is the linear trimer, only detected over the hsMgO(I), which reacts to form either isophorones or mesitylene.

Isophorones, the main trimer, are formed by intramolecular aldol condensation of phorones, requiring strong basic sites. At 623 K (not shown), isophorones selectivity is higher for hsMgO(II), 16.1 %, than hsMgO(I), 15.4 %, in agreement with the strongest basicity of hsMgO(II). In fact, hsMgO(II) shows the highest concentration of strong basic sites and, according to FT-IR results, unidentate carbonates remain at the highest temperature studied. However, at 723 K, null presence of either isophorones or mesitylene is observed for hsMgO(II). Both MgO samples exhibit similar selectivity towards isophores (26.1 %), whereas hsMgO(I) exhibits the highest selectivity (32.8 %). Likewise, it is observed that the carbon balance at 723 K decreases for all catalysts, and this trend is more notorious in the case of hsMgO(II). Thus, further reactions not detected could occur on the surface of this catalyst consuming the C₉ compounds. These condensation products could be adsorbed on the strongest basic sites (by analogy with the adsorption of CO₂), hindering further reaction and limiting the formation of mesityl oxide.

4. CONCLUSIONS

The effect of the preparation procedure of magnesium oxide (precursor and hydrothermal treatment) has been studied for the condensation of acetone. Calorimetry

measurements, as well as FT-IR analysis, were used to understand the modifications of the surface chemistry.

The thermal decomposition procedure generates defects in the surface that are responsible for its high surface area ($221 \text{ m}^2/\text{g}$). and important concentration of surface basic sites. Furthermore, it was shown that high surface area MgO prepared by this methodology is an effective catalyst for the aldol condensation of acetone, reaching conversion similar to mixed oxides and selectivities towards the isophorones, even higher than for these materials.

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Table 1. Textural properties of the different magnesium oxides tested in this work.

Sample	S_{BET} (m^2/g)	$V_{\text{p, mesopores}}$ (cm^3/g)	L_{200} (nm)
MgO(I)	26	-	56.9
MgO(II)	70	0.189	7.2
hsMgO(I)	221	1.147	11.0
hsMgO(II)	87	0.530	8.7

S_{BET} : surface area obtained by nitrogen adsorption-desorption isotherms by the Brunauer–Emmett–Teller theory.

V_{p} : pore volume corresponding to the cumulative pore volumes between 170 and 30000 nm, determined from the desorption branch using the Barrett-Joyner-Halenda (BJH) method

L_{200} : crystallite size of the oxides calculated from the plane (200) of MgO using the Debye-Scherrer equation.

Table 2.Characteristic bands of the species formed upon CO₂ adsorption on the magnesium oxides

	bb	bb	h/cb	cb	u	u	u	h	u	u	u	cb	cb	bb	h/bb
MgO(I)		1706 (4)	1685 (4) [379]	1662 (3) [313]	1635 (6) [254]	1594 (9) [184]	1517 (9) [85]		1432 (5)	1409 (6)	1382 (2)	1349 (4)	1306 (8)		1220 (1)
MgO(II)	1726 (10) [507]	1706 (16) [417]	1686 (18) [375]	1665 (13) [316]	1640 (32) [254]	1603 (31)	1541 (41) [119]	1458 (9)	1422 (22)		1386 (23)	1349 (25)	1311 (18)	1289 (21)	1220 (10)
hsMgO(I)	1731 (7) [513]	1704 (11) [427]	1683 (13) [377]	1662 (10) [314]	1633 (30) [250]	1591 (18) [184]	1538 (14) [109]	1457 (5)	1429 (5)	1406 (4)	1383 (12)	1347 (12)	1306 (15)	1277 (6)	1218 (5)
hsMgO(II)	1727 (10) [507]	1706 (7) [423]	1685 (20) [374]	1661 (15) [312]	1635 (23) [254]	1590 (42) [184]	1515 (25) [87]	1456 (13)	1428 (13)	1406 (13)	1381 (22)	1349 (10)	1311 (20)	1283 (11)	1220 (8)

u (unidentate carbonates), *cb* (chelating bidentate carbonates), *bb* (bridged bidentate carbonates), and *h* (bicarbonates).

In bold(cm⁻¹):Infrared bands after CO₂ adsorption at room temperature

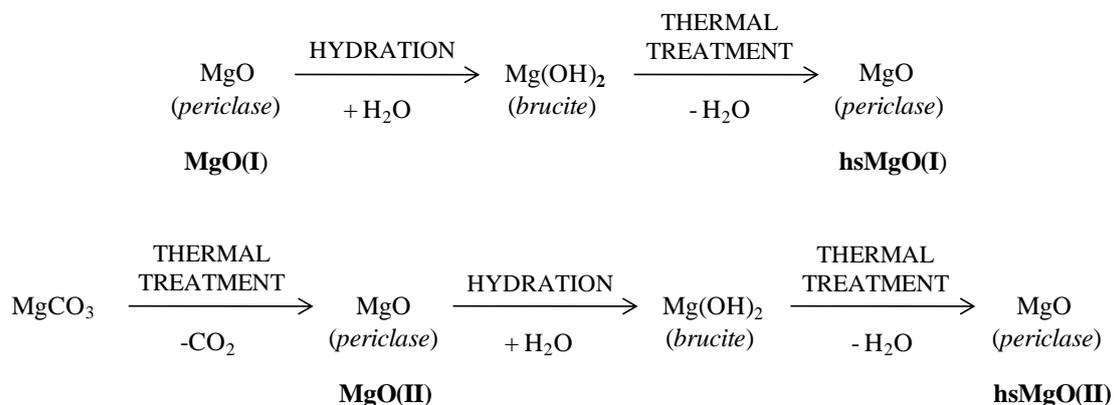
In brackets:area of the decomposed peaks

In square brackets: $\Delta\nu_3$ values (cm⁻¹)

Table 3. Evolution of selectivity of acetone condensation towards the reactions products at two temperatures: diacetone alcohol (DAA), isomesityl oxide (IMO), mesityl oxide (MO), phorone (P), mesitylene (M) and isophorone (IP)

Catalyst	Conversion (%) / carbon balance (%)		Selectivity (%)											
			Diacetone alcohol		Isomesityl oxide		Mesityl oxide		Phorone		Mesitylene		Isophorone	
	523 K	723 K	523 K	723 K	523 K	723 K	523 K	723 K	523 K	723 K	523 K	723 K	523 K	723 K
MgO (I)	20.0/88.9	22.7/95.6	5.1	0.7	15.1	58.7	79.7	3.2	0	0	0.1	11.3	0	26.1
MgO (II)	17.8/88.3	19.7/92.9	5.1	0.7	15.1	58.7	79.8	3.3	0	0	0	11.2	0	26.1
hsMgO (I)	33.0/72.9	37.0/85.2	0	0.2	7.2	9.3	49.4	33.8	10.9	3.2	10.6	20.7	21.9	32.8
hsMgO (II)	25.6/72.9	28.9/81.6	14.6	0.0	0	28.7	85.4	71.3	0	0	0	0	0	0

Scheme 1. Activation steps and notation (**bold**) used in this work for the different MgO used in this work



Scheme 2. Species formed upon CO₂ adsorption

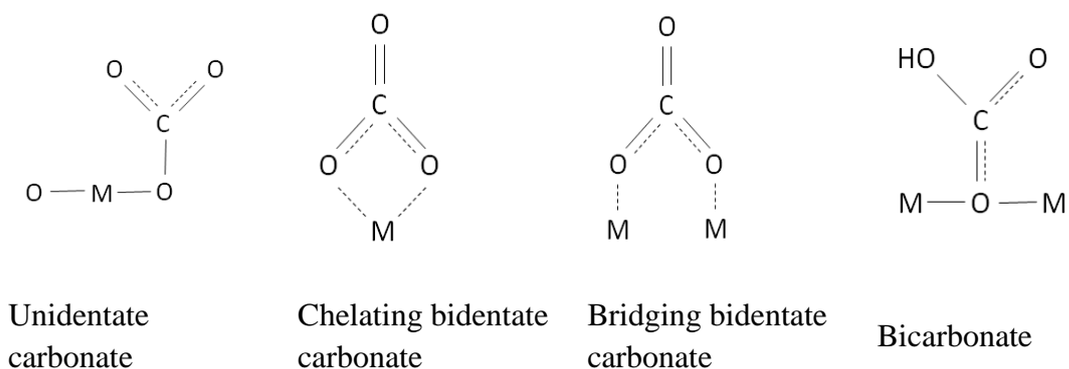


FIGURE CAPTIONS:

Fig. 1. XRD patterns corresponding to the magnesium oxides: MgO(I), MgO(II), hsMgO(I) and hsMgO(II)

Fig. 2. XRD patterns corresponding to the transition brucite-periclase at increasing temperature on magnesium hydroxides obtained from MgO(I) (a) and MgO(II) (b)

Fig. 3. Kinetic data corresponding to the transition brucite-periclase (A/A_0 : ratio of the area under the peak at a given time to the area under the peak at time zero)

Fig. 4. N_2 adsorption-desorption isotherms at 77 K for the MgO samples studied in this work.

Fig. 5. SEM images of MgO(I) (a), MgO(II) (b), hsMgO(I) (c), and hsMgO(II) (d)

Fig. 6. CO_2 adsorption isotherms at 323 K and 6.7 kPa, measured by microcalorimetry for: MgO(I) (\blacklozenge), MgO(II) (\blacksquare), hsMgO(I) (\square), and hsMgO(II) (\circ)

Fig. 7. Differential heats of CO_2 adsorption vs. coverage at 323 K on: MgO(I) (\blacklozenge), MgO(II) (\blacksquare), hsMgO(I) (\square), and hsMgO(II) (\circ). Distribution of interaction strengths of carbon dioxide adsorption on the samples: $210 > Q > 150$ kJ/mol (black); $150 > Q > 90$ kJ/mol (grey); $90 > Q > 30$ kJ/mol (white).

Fig. 8. Activity for the acetone condensation over the different MgO catalysts: MgO(I) (\blacklozenge), MgO(II) (\blacksquare), hsMgO(I) (\square), and hsMgO(II) (\circ)

Fig. 9. Influence of surface area (a) and total concentration of basic sites (b) (measured by calorimetry) on the conversion of the studied catalysts at 723 K

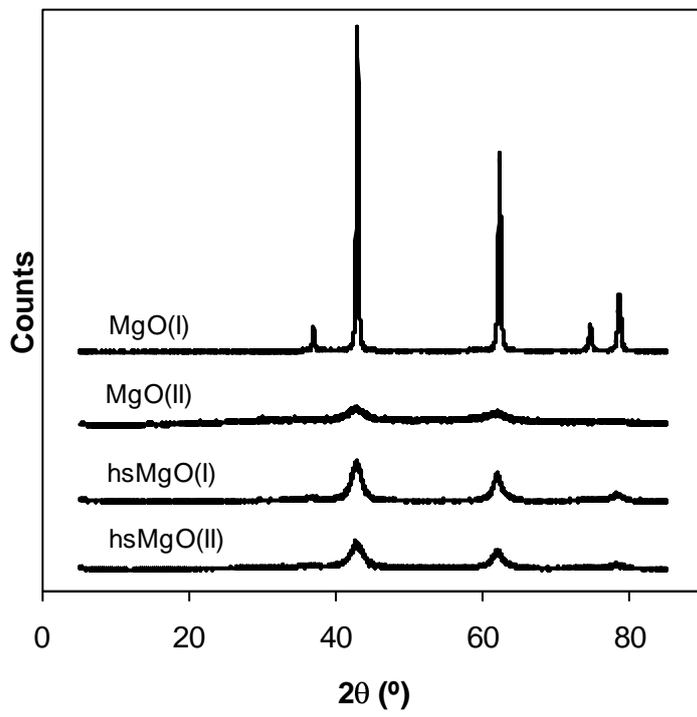
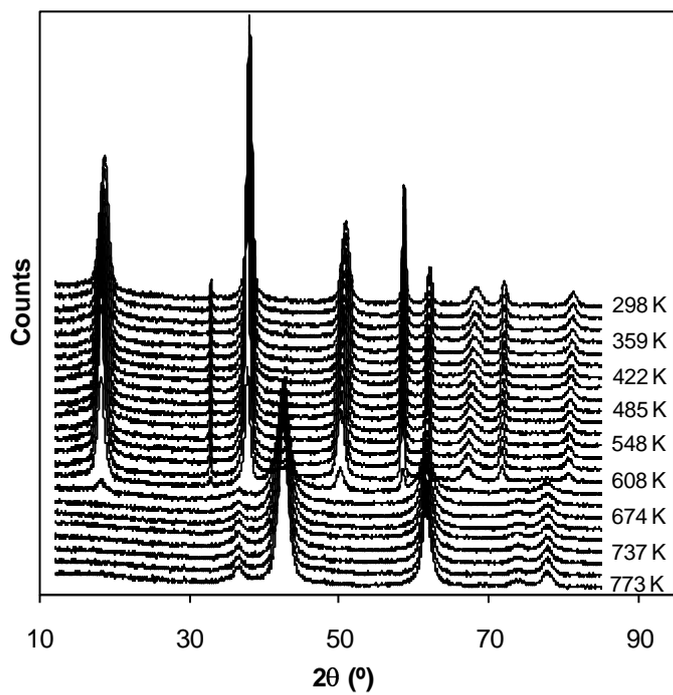


Fig. 1

(a)



(b)

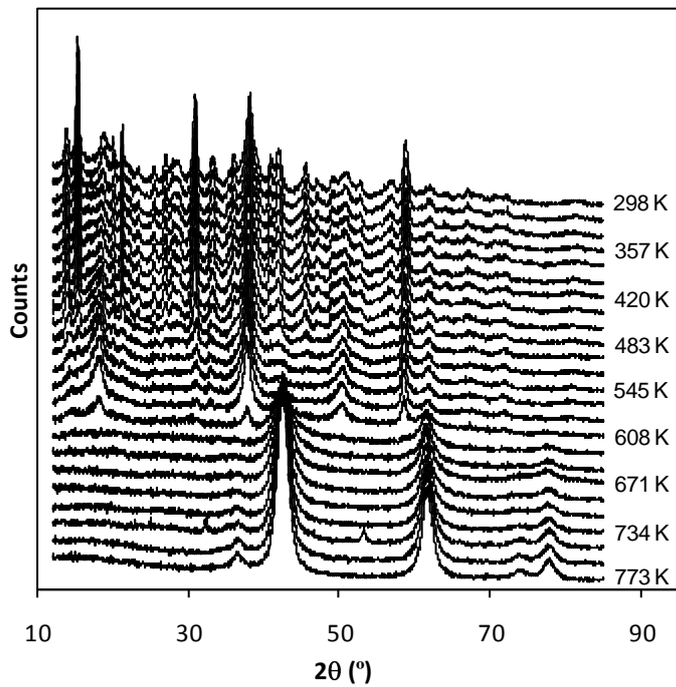


Fig. 2

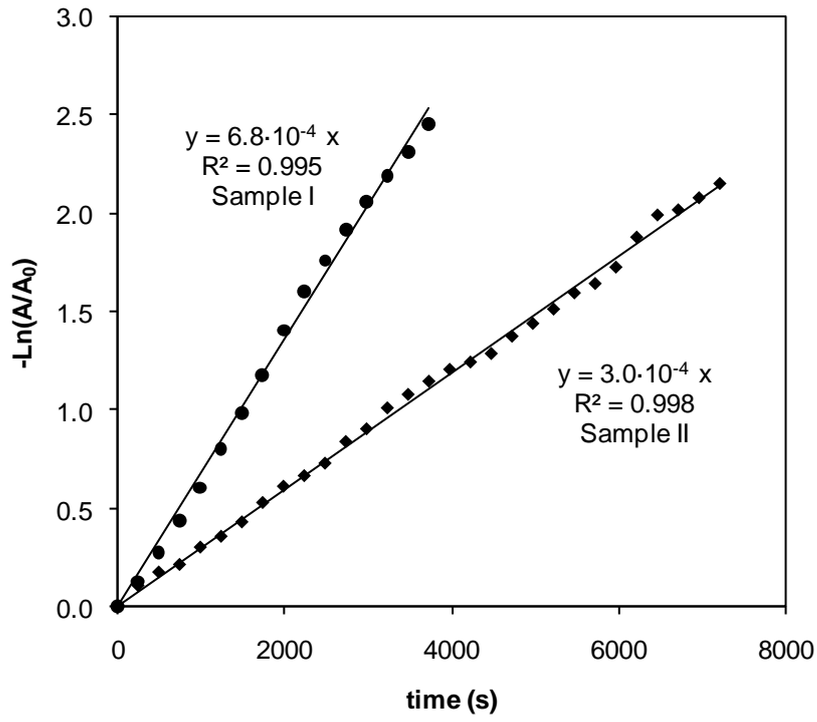


Fig. 3

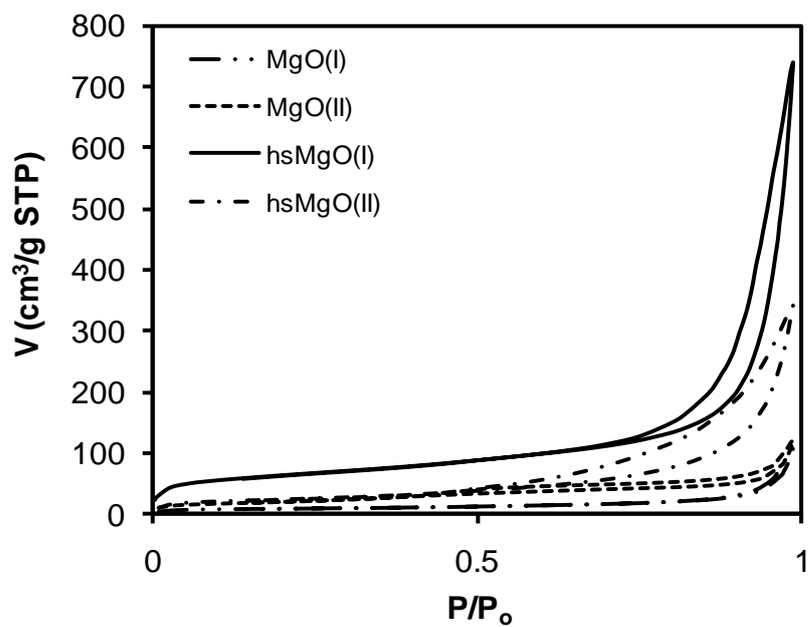


Fig. 4

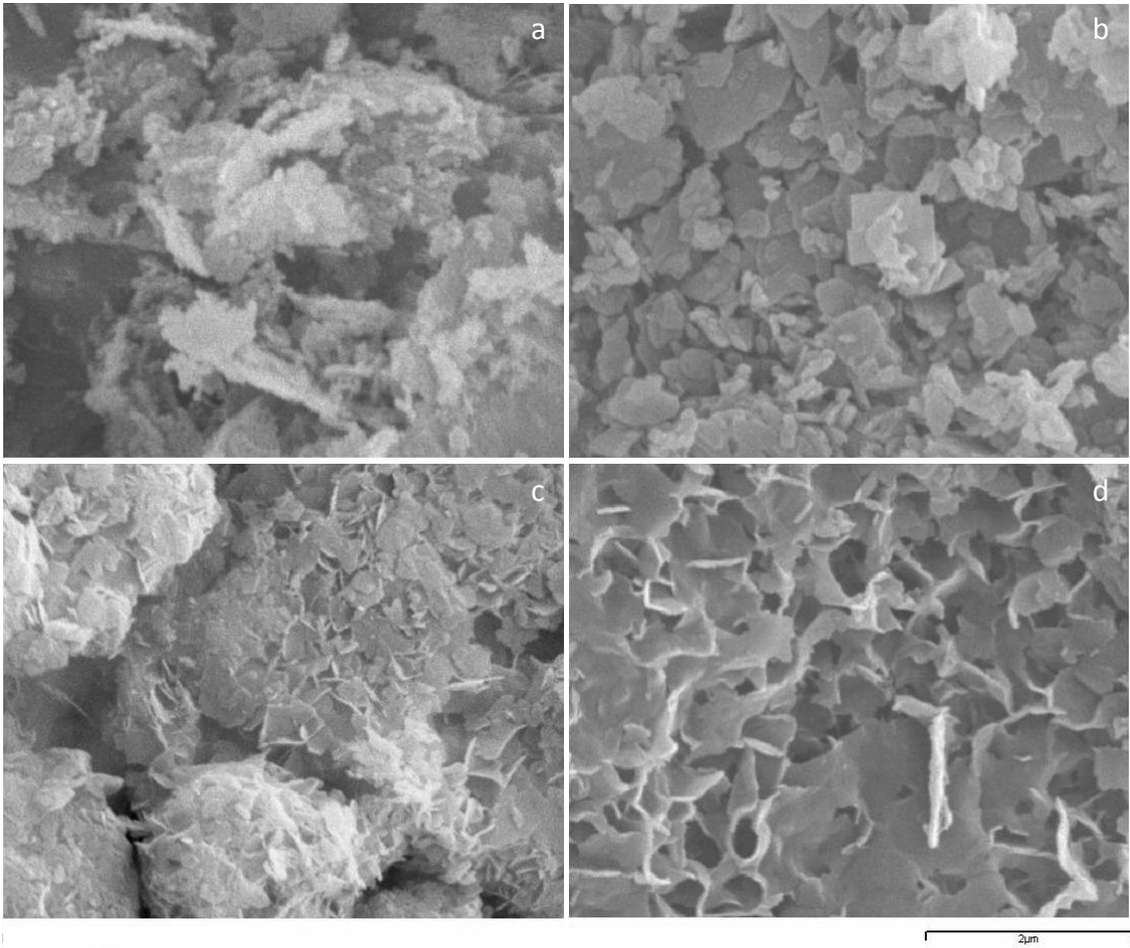


Fig. 5

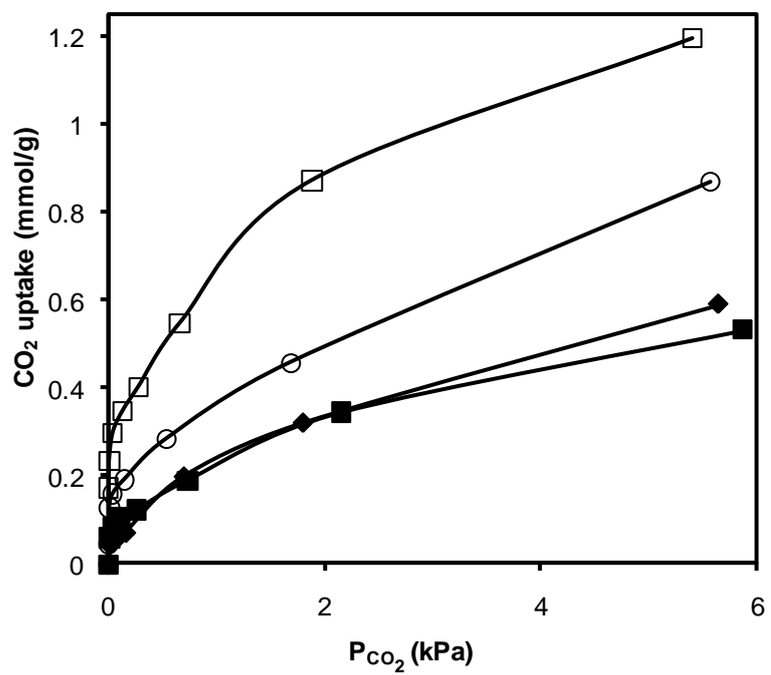


Fig. 6

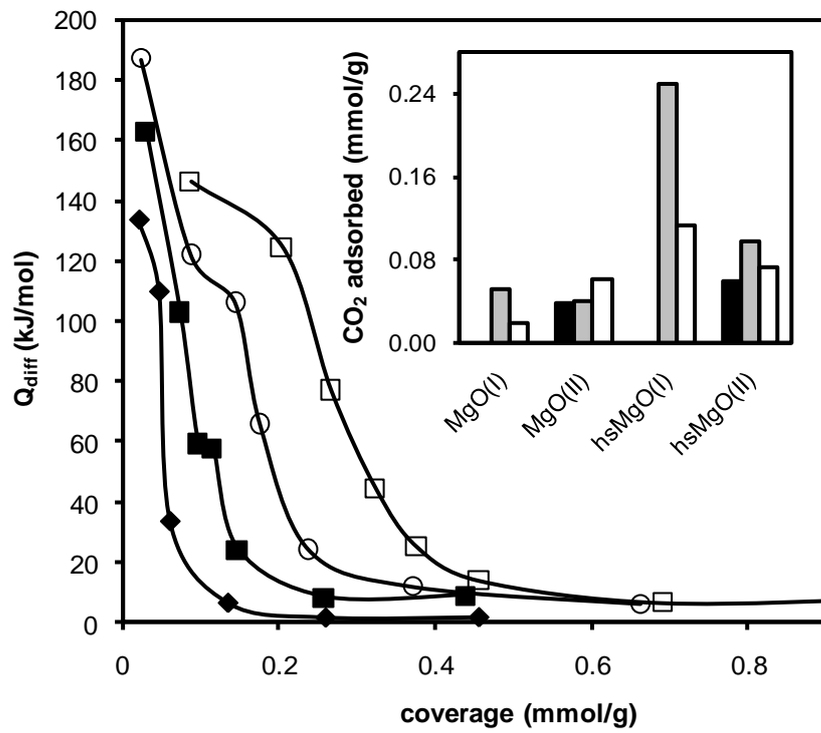


Fig. 7

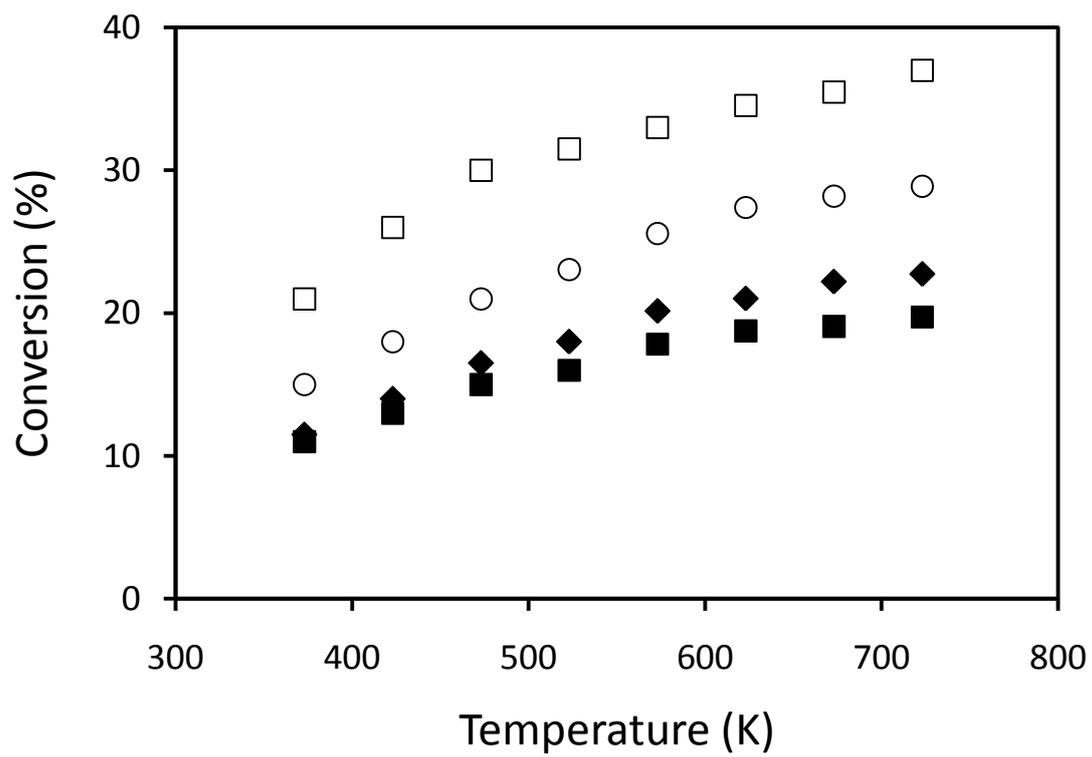


Fig. 8

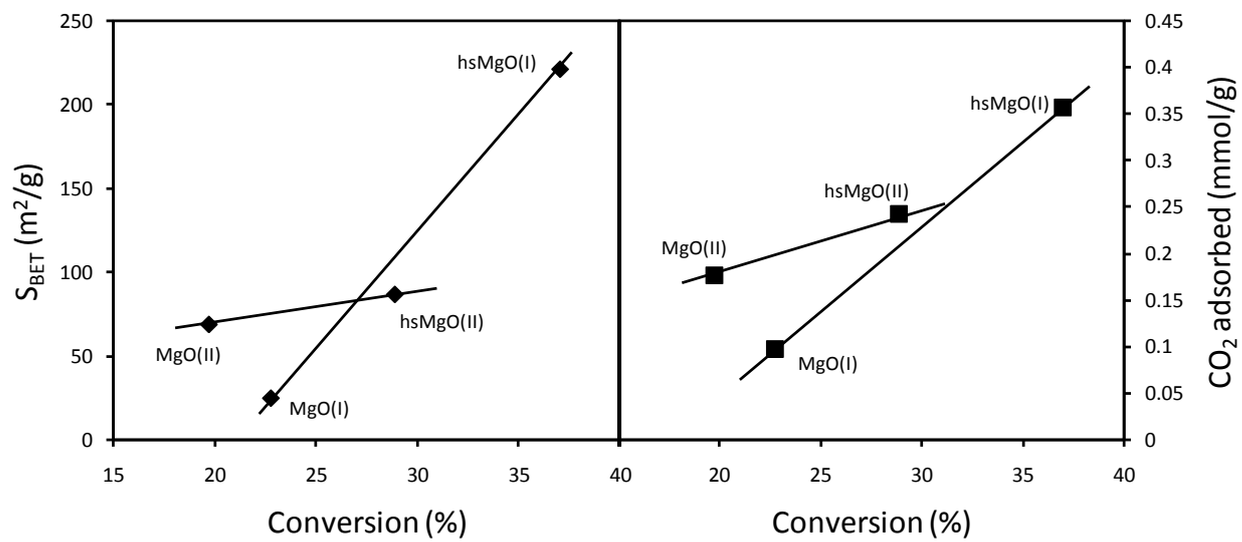


Fig. 9