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Understanding the Role of Mono and Ternary Alkali Metal Salts on CO₂ Uptake of MgO Sorbents

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Abstract: CO₂ uptake by MgO-based sorbents at intermediate temperatures is attractive for preand post-combustion CO₂ capture applications. However, besides the high CO₂ uptake potential of these materials (1.1 g CO₂ g^{-1} sorbent), in practice, the realistic CO₂ capture is far from that of the theorical values. In this work, the sol-gel method was used to synthetize unsupported and supported MgO sorbents (10% Ca⁻ or 10% Ce⁻ support, mol) that were impregnated with different fractions (15, 25, and 35; % mol) of a NaNO3 single salt or a ternary alkali salt (NaNO3, LiNO3 and KNO₃ (18/30/52; % mol)). To understand the role of alkali metal salts (AMSs) in the MgO sorbents' performance, the working and decomposition temperature ranges of AMS under different atmospheres (CO_2 and air) were evaluated. The findings show that the CO_2 uptake temperature range and maximum uptake (20-500 °C, CO₂ atmosphere) of sorbents are correlated. The cyclic CO₂ uptake of the most promising sorbents was tested along five carbonation-calcination cycles. For the first and fifth cycles, respectively, the 15 (Na, K, Li)-MgO sorbents showed the highest carrying capacity, i.e., 460–330 mg CO $_2$ g $^{-1}$ sorbent, while for the 15 (Na, K, Li)-MgO-Ca sorbents, it was $375-275 \text{ mg CO}_2 \text{ g}^{-1}$. However, after the first cycle, the carbonation occurred faster for the 15 (Na, K, Li)-MgO-Ca sorbents, meaning that it can be a path to overpassing carbonation kinetics limitations of the MgO sorbent, making it viable for industrial applications.

Keywords: MgO-based sorbents; alkali metal salts; intermediate-temperature carbon capture

1. Introduction

MgO is an intermediate-temperature CO₂ sorbent (200–400 °C) that gained relevance in the scientific community during the last decade [1,2]. This sorbent is suitable for both post- and pre-combustion CO₂ capture technologies. Most of the studies in the literature have been performed for post-combustion conditions [3,4], but the application in sorptionenhanced water gas shift (SEWGS) reactions [5,6], i.e., under pre-combustion conditions, is also relevant since it increases the H₂ production and purity.

 CO_2 capture is based on the following reversible chemical reaction: MgO (s) + CO_2 (g) \rightleftharpoons MgCO₃ (s), and since one mole of MgO can absorb one mole of CO_2 , its theoretical CO_2 capture capacity is high (1.1 g CO_2 g⁻¹ MgO) [7]. The calcination and carbonation reactions are both dependent on temperature, and its equilibrium is also related to the CO_2 partial pressure according to Equation (1) [8]:

$$\Gamma_{\rm eq} = \frac{13636}{\ln\left(\frac{[\rm bar]}{P_{\rm CO_2}}\right) + 20.01} - 273.15 \tag{1}$$

Calcination is an endothermic reaction (Δ H 25 °C = +116.9 kJ mol⁻¹) that, ideally, should be performed in an atmosphere rich in CO₂ (>450 °C) to allow for a high CO₂ concentration at the exit of the calciner with an efficient compression and storage for the



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Copyright: © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). utilization of the captured CO_2 . Some advantages of this sorbent, which is available in nature as MgCO₃ or MgO, include its characteristics of being nontoxic and noncorrosive and its availability at a relatively low cost. Moreover, as mentioned above, its regeneration can occur at around 450–500 °C; then, the energy consumption during this step is reduced in comparison with that of CaO or Li₄SiO₄ sorbents' regeneration that needs temperatures higher than 700 °C [5].

In addition to the interesting advantages of MgO as a CO₂ sorbent, MgO has a very low experimental CO₂ uptake capacity of 11–20 mg CO₂ g⁻¹ and very poor experimental sorption kinetics at a very favorable temperature of 200 °C from a thermodynamic point of view [9]. Some explanations have been proposed to justify the low performance of MgO [7]: (i) the low surface area does not allow for sufficient exposure of its basic sites for CO₂ carbonation; (ii) the low porosity can obstruct the CO₂ diffusion through the pores and delay the carbonation equilibrium; (iii) the MgO has a volume expansion of 2.49 due to the formation of MgCO₃, while the formation of dense layers adjacent to basic active sites of the MgO sorbent can inhibit the CO₂ carbonation; (iv) the MgO has an intrinsically high lattice enthalpy.

For an upscale application of MgO sorbents, the kinetic and carrying capacity limitations need to be overcome. In this sense, different paths to enhance the MgO sorbents performance, based on their dependence on intrinsic and extrinsic factors, have been assessed. To upgrade the internal properties of sorbents, the effects of distinct factors and their synergies have been evaluated: different MgO precursors, the synthesis of mesoporous MgO, dispersion on inert supports, and doping with alkali metal salts (AMSs). Factors like the CO_2 concentration, pressure, presence of water, and presence of gas impurities are extrinsic factors that can also influence CO_2 uptake by sorbents.

The most common magnesium precursors include inorganic salts, organometallic salts, and natural minerals. In general, the organometallic salts of magnesium are better precursors than the other two types in producing efficient MgO sorbents. This is mainly because the former usually have a larger molecular weight, which results in MgO with a better pore size and porous structure [2]. However, the organometallic precursors are expensive. On the contrary, natural Mg-containing minerals are abundant in nature and, thus, are costeffective. For instance, magnesium oxalate (MgC_2O_4) is a better precursor than $MgCO_3$ because it generates MgO sorbents with better performance [3]. Guo et al. [3] obtained a MgO sorbent derived from magnesium oxalate dihydrate ($MgC_2O_4.2H_2O$) with the highest CO_2 capture capacity of 194 mg CO_2 g⁻¹ sorbent. This sorbent is characterized by excellent textural properties, a uniform surface, and abundant basic sites. Hanif et al. [10] synthesized mesoporous MgO with a BET surface area higher than 350 m² g⁻¹ using the ammoniumhydroxide-assisted precipitation of Mg(OH)₂ from a Mg(NO₃)₂ aqueous solution followed by the thermal degradation of the precipitated $Mg(OH)_2$ in vacuum. This sample presented a CO_2 capture capacity of 75 mg CO_2 g⁻¹ sorbent at 300 °C. The synthesis of mesoporous MgO is also an effective way to improve sorbents' CO_2 capture capacity. When the CO_2 reacts with MgO, it forms a covering layer of the thermodynamically stable MgCO₃ on the unreacted sorbents' surface, which delays the CO₂ molecules' diffusion through the product layer. Thus, the production of an MgO sorbent with a high surface area and porosity should facilitate the CO_2 diffusion, enhancing the capture capacity and kinetics reactivity of the sorbent [2]. The dispersion of MgO on a porous support also increases the surface area availability of wellexposed basic active sites, which in turn improves the sorbent's capture capacity. Ideally, a good support must present a stable porous structure, which should enhance the sorbent performance and reduce its sintering after several carbonation-calcination cycles, contributing to the stability of the sorbent's CO_2 uptake capacity.

Among the approaches to improve the CO_2 uptake properties of MgO, the doping with alkali metal salts is the most widely recognized promising approach. The aim of the most recent experimental works is to improve the CO_2 uptake capacity of these materials up to 0.7–0.8 g CO_2 g⁻¹ sorbent [11]. The main categories of alkali doping described in the

literature [2] are the alkali carbonate doping, alkali nitrate/nitrite doping and binary or ternary alkali doping.

A US patent [12] described that alkali metal carbonate-doped MgO sorbents prepared by coprecipitation registered CO₂ capture capacities of 48–570 mg CO₂ g⁻¹ sorbent at the temperature range of 350 to 400 °C and under 70% dry CO₂. Moreover, it was observed that the performance of the doped sorbent was influenced by the doping precursor. For instance, the MgO sorbents doped with Na₂CO₃ showed better performance than when doped with Li₂CO₃ or K₂CO₃.

Zhang et al. [13] doped MgO sorbents with NaNO₃. The authors achieve a good CO_2 carbonation kinetics and a MgO conversion of 75% against only 2% for an undoped MgO, both at 330 °C and ambient pressure. It was stated that molten NaNO₃ decreases the dissociation energy of Mg-O ionic bonds in bulk MgO; thus, the molten NaNO₃ acts as a phase transfer catalyst (PTC) between bulk MgO and CO_2 molecules which, in turn, facilitates the carbonation reaction. In addition, molten alkali metal nitrates prevent the formation of a rigid, CO_2 impermeable, and monodentate carbonate layer on the surface of MgO as it occurs with bare MgO and promote the rapid generation of carbonate ions to allow a high rate of CO_2 uptake [2].

Concerning to binary or ternary alkali doping, very promising results are found in the literature. Lee et al. [14] studied the Na₂CO₃/NaNO₃-doped MgO sorbent that maintained a CO₂ sorption capacity of 153 mg CO₂ g⁻¹ sorbent after seven cycles. The authors related this performance with the interaction established between MgO, CO₂ and the molten NaNO₃ that results in the formation of the double salt Na₂Mg(CO₃)₂. Identical results were registered for other cases of binary doping, such as K₂CO₃/KNO₃-doped MgO sorbents. Zhang et al. [15] provided a detailed explanation for the enhancement of the CO₂ uptake capacity of MgO sorbents doped with the binary Na₂CO₃/NaNO₃ system. The mechanism is stated to be similar to that of the single alkali nitrate. The bulk MgO dissolves in the molten salt because Mg-O ionic bonds are easy to break. Na₂CO₃ also dissolves in the same liquid medium because carbonate salts have good solubility in molten salts. Hence, ion pairs of Mg²⁺, O²⁻ and CO₃²⁻ are formed and react with the CO₂ molecules, generating the double salt (Na₂Mg(CO₃)₂). The same mechanism is appropriate to binary alkali nitrate/carbonate doping, which forms the CaMg(CO₃)₂ double salt [2].

The binary doping with alkali nitrate/nitrite is also an interesting matter of study. Zhao et al. [16] compared the CO₂ sorption capacities of the single NaNO₃ and of the binary NaNO₃/NaNO₂-doped MgO sorbents. The latter showed higher CO₂ sorption capacity than the former. This new evidence was explained by the reduction in the melting temperature of the eutectic mixture. While single NaNO₃ and NaNO₂ present a theoretical melting point of 308 °C and 271 °C, respectively, the eutectic mixture of NaNO₃/NaNO₂ exhibits a melting temperature of 185 °C. Thus, the eutectic mixture facilitates the carbonation process by providing a molten phase that works like a liquid channel. Ternary doping with NaNO₃, LiNO₃ and KNO₃ (18/30/52; % mol) registered a sharper reduction in the eutectic mixture's melting point (120 °C) and an enhanced CO₂ sorption performance [2].

Summarizing, the doping with AMS improves the cyclic CO_2 sorption capacity of MgO sorbents when compared to that of commercial MgO, of mesoporous MgO and of MgO obtained from different precursors. Nevertheless, most of the AMS-doped MgO sorbents exhibit a CO_2 uptake capacity of less than 50% of the maximum theoretical capacity. Table 1 shows different studies found in the literature using unsupported and supported MgO-based sorbents doped with AMS and the corresponding CO_2 uptake capacity after multicyclic tests.

As mentioned above, the existence of a molten state generated by the melting of alkali metal salts is essential to increase the sorbent's CO₂ uptake and to improve the kinetic of MgO carbonation. In this work, the role of the NaNO₃ single salt and the amount of a ternary alkali salt (NaNO₃, LiNO₃ and KNO₃ (18/30/52; % mol)) on the CO₂ uptake temperature range, as well as on the CO₂ maximum uptake, will be evaluated for unsupported and supported sorbents. To evaluate the calcination atmosphere effect on

the working temperature range of the alkali salts, its decomposition was studied under air and CO_2 atmosphere. As innovative work in relation to the experiments shown in Table 1, the sorbent calcination will be performed under a CO_2 atmosphere to obtain a concentrated CO_2 stream useful for storage or utilization, which is more interesting for industrial applications.

Table 1. CO₂ uptake and experimental conditions of unsupported and supported MgO-based sorbents impregnated with AMS.

MgO-Based Sorbent	Temperature (°C)/Atmo	Number of	CO_2 Uptake (mg CO_2 g ⁻¹ Sorbent)		Ref.	
-	Carbonation	Calcination	Cycles	Initial	Final	
10 NaNO ₃ _MgO	300/10% CO ₂ /180	300/He/180	5	390	210	[17]
10 NaNO ₃ _MgO	300/CO ₂ /30	$450/N_2/30$	20	140	100	[18]
10 (Li, Na, K)NO ₃ _MgO	300/CO ₂ /60	$450/N_2/15$	5	480	410	[11]
15 (Li, Na, K)NO ₃ _MgO	300/CO ₂ /60	$350/N_2/30$	5	440	350	[19]
15 (Li, Na, K)NO ₃ _MgO	300/CO ₂ /60	$350/N_2/30$	12	440	310	[19]
	Supported: x	AMS—MgO—Y supp	ort			
20 (Li, Na, K)NO ₃ _MgO_5 Ca	300/CO ₂ /30	$425/N_2/10$	5	396	330	[20]
12 (Li, Na, K)NO ₃ _MgO_10 Ce	$300/CO_2/60$	425/N ₂ /15	5	340	180	[21]
17 (Li, Na, K)NO ₃ _MgO_10 Ce	300/CO ₂ /60	$425/N_2/15$	5	420	330	[21]

x—molar fraction (%) of AMS, Y—molar fraction (%) of support.

2. Experimental Procedure

2.1. Materials

All the MgO-based sorbents were prepared with magnesium nitrate hexahydrate $(Mg(NO_3)_2 \cdot 6H_2O)$, with an assay of 98–102% (Sigma-Aldrich, St. Louis, MO, USA), and for the supported materials were used calcium nitrate tetrahydrate (Ca(NO_3)_2 \cdot 4H_2O) or cerium nitrate hexahydrate (Ce(NO_3)_3 \cdot 6H_2O), both with an assay \geq 99% (Sigma-Aldrich, St. Louis, MO, USA). The citric acid monohydrate (C₆H₈O₇ · H₂O) with an assay of 99.5–102%, was used as a chelating agent (Panreac, Castelar del Vallés, Barcelona, Spain).

2.2. MgO-Based Sorbents Synthesis and Doping with Alkali Metal Salts

The Mg(NO₃)₂·6H₂O was dissolved in distilled water with a molar ratio of citric acid and distilled water to magnesium of 1:1 and 120:1, respectively. For the supported MgO sorbents, the same procedure was used but adding Ca(NO₃)₂·4H₂O or Ce(NO₃)₃·6H₂O.

As shown in Figure 1, each solution was continuously stirred at 80 °C for 6 h. Afterward, the wet gel was dried in the oven at 120 °C for 14 h, milled, and the sample obtained was calcined in the muffle by ramping the temperature to 500 °C at a rate of 2 °C min⁻¹ plus 2 h more at 500 °C.



Figure 1. Synthesis of MgO based sorbents by the sol-gel method.

The synthesized MgO-based sorbents were doped with mono AMS (NaNO₃) and with a ternary AMS mixture (NaNO₃, LiNO₃ and KNO₃ (18/30/52; mol.%)) [11,22] using the wet-impregnation method. Prior to the impregnation, the AMS was dried in the oven at 120 °C for 24 h. First, the MgO-based sorbent and AMS have been dissolved in 20 mL of distillate water, using magnetic stirring for 1 h at room temperature. The obtained aqueous slurry was dried in the oven at 120 °C for 14 h. Afterward, the dried sample was placed in the muffle for calcination using a heating rate of 2 °C min⁻¹ until it reached 450 °C, plus 4 h at 450 °C. Table 2 summarizes all the prepared AMS-MgO-based sorbents and the different ratios of AMS in each sorbent. The following nomenclature was used for the doped sorbent samples: x (Na or Na-Li-K)-MgO-Z, where x corresponds to the molar fraction (%) of AMS, and Z corresponds to the Ca or Ce support.

Table 2. Summary of the molar composition of unsupported and supported MgO sorbents.

		6	Alkali Metal Salts						
Sorbent Precursor	Magnesium (mol, %)	Support (mol, %)	NaNO ₃			(Na,Li,K)NO ₃			
			(mol, %)	Sample ID	(mol, %)	Sample ID			
			15	15 Na-MgO	15	15 (Na,K,Li)-MgO			
MgO-SG ¹	100	0	25	25 Na-MgO	25	25 (Na,K,Li)-MgO			
-			35	35 Na-MgO	35	35 (Na,K,Li)-MgO			
MgO-Ca-SG	90	10		-	15	15 (Na,K,Li)-MgŌ-Ca			
MgO-Ce-SG	90	10			15	15 (Na,K,Li)-MgO-Ce			

¹ SG: sol gel.

2.3. Characterization Methods

A D8 Advance diffractometer (Bruker AXS GmbH, Karlsruhe, Germany) using Cu K α ($\lambda = 0.15406$) radiation and a Ni filter was used to identify the crystalline phases of the synthetized sorbents before and after the cyclic experiments on a thermogravimetric analyzer (Setsys Evolution TGA, Setaram Instruments, Caluire, France). The equipment was operated at 30 mA and 40 kV, scanned within the 2 θ range of 5–80°, and had a step size (scanning duration) of 0.03° (0.5 s). The crystallography open database (COD) was used to identify the crystalline phases. The MgO average crystallite size of the sorbents was estimated using Scherrer's equation (D = K λ /bcos θ), based on the XRD data, where D is the crystallite size (nm), b corresponds to the full width at half maximum (FWHM) of the XRD peak considered, λ is the wavelength (0.15406 nm), θ is the Bragg angle (degree) and K is Scherrer's constant (K = 0.9, assuming that particles are spherical).

The specific surface area (S_{BET}) of the synthetized sorbents based on the BET method and the total pore volume (V_p), were determined by the N₂ sorption technique at -196 °C, (Quantachrome Instruments, Model autosorb IQ, Graz, Austria) at a relative pressure (p/p₀) of 0.95. The degasification procedure was performed in two steps: the first one at 90 °C for 1 h and the second one at 350 °C for 5 h.

2.4. Thermal Decomposition of Mono and Ternary Alkali Nitrates Salts

The thermal decomposition of the molten salts was assessed for the pure NaNO₃ and for the ternary mixture of 52% (mol) KNO₃, 18% (mol) NaNO₃ and 30% (mol) LiNO₃ by thermogravimetric analysis with air and pure CO₂ atmospheres between 25 and 1000 °C at a heating rate of 10 °C min⁻¹.

2.5. CO₂ Uptake TGA Studies

2.5.1. Effect of Mono and Ternary AMS Molar Fraction and Inert Support Addition

The effect of mono or ternary alkali metal salts doping, as well as the best molar fraction of AMS-MgO were evaluated in a thermogravimetric analyzer (TGA) system TG-DSC Setsys Evo 16. The TGA studies were carried out with MgO-based sorbents for the range of temperature from 20 to 500 °C and a temperature ramp of 10 °C min⁻¹. During the experiments, a constant flow of 80 mL min⁻¹ of CO₂ and ~10 mg of sample were used.

The best two mono and ternary AMS samples, i.e., with higher CO_2 uptake capacity, were identified and selected to proceed to the tests with five carbonation–calcination cycles in the TGA as described below. Then, using the same procedure, the effect of the addition of an inert support to the MgO sorbent was tested with Ca and Ce additions (10%, mol), considering the most promising AMS doping and the respective molar fraction.

2.5.2. Carbonation–Calcination Cycles

A TGA system TG-DSC Setsys Evo 16 was used to carry out the studies for the assessment of both the cyclic stability and the CO₂ uptake of the supported and unsupported MgO sorbents. These tests consisted of five carbonation–calcination cycles with carbonation and calcination temperatures of 300 °C (60 min) and 445 °C (60 min), respectively, using a constant flow of 80 mL min⁻¹ of CO₂, a ramp temperature of 10 °C min⁻¹ and ~10 mg of sample. A blank experiment was performed to correct apparatus buoyancy effects like atmosphere density changes with temperature. The unsupported sample showing the best performance was identified and tested through 12 carbonation–calcination cycles using the same experimental conditions.

The TGA mass variation allows determining the mass of CO_2 captured by the sorbent during the carbonation step (Equation (2)), and subsequently, the carrying capacity and the MgO conversion to carbonate can be obtained using Equations (3), (4) and (5), respectively.

$$m_{CO_2 captured} = m_{carbonation} - m_{calcination}$$
(2)

where $m_{CO2captured}$ is the mass of CO₂ captured in each cycle, $m_{carbonation}$ is the mass of the sample after the carbonation reaction, and $m_{calcination}$ is the mass of the sample after the calcination reaction. The carrying capacity (CC) of the sorbent is defined as the mass of CO₂ captured per mass unit of the sorbent (mg CO₂ g⁻¹ sorbent).

$$CC = m_{CO_2 captured} / m_{sorbent}$$
(3)

where $m_{sorbent}$ is the mass of the sorbent. The theoretical carrying capacity (CC_{theor.}) is calculated with Equation (4),

$$CC_{theor.} = M_{CO_2} / M_{MgO} = 1.09$$
 (4)

where M_{CO_2} and M_{MgO} represent the molar mass of CO_2 and MgO (g mol⁻¹), respectively. The MgO conversion of the sorbent is defined as the percentage of the CC when compared to the CC_{theor}.

$$MgO_{conversion} = CC / \left(CC_{theor.} \times w_{MgO_{sorbent}} \right) \times 100(\%)$$
(5)

where $w_{MgO_{sorbent}}$ represents the nominal mass fraction of the MgO (%, wt.) in the sorbent.

3. Results

3.1. Properties of MgO-Based Sorbents

The crystalline structures of the MgO-SG sorbent impregnated with 15, 25 and 35% of either NaNO₃ (Figure 2a) or a ternary mixture of KNO₃, NaNO₃ and LiNO₃ (Figure 2b) were analyzed by powder XRD. All the XRD-obtained patterns show the characteristic peaks of the MgO (42.9°, 62.2°, 78.5°) pattern and a main peak located at ~29.4° that is referred to the NaNO₃ pattern. For the case of the sorbents impregnated with the ternary alkali metals salts, the main peak of KNO₃ (~23.6°) was also observed, and the second main peak should be overlaid by the NaNO₃ peak (29.4°). The LiNO₃ was not identified, which can be justified by its high dispersion on the sorbent sample, leading to the Li concentration below the instrumental detection limit [23] or its partial decomposition to Li₂O₂ or LiO₂ whose main peaks are around 32.9° and 33.5°. Wang et al. [24] studied a similar ternary system, LiNO₃ (25.9 mol.%), NaNO₃ (20.06 mol.%) and KNO₃ (54.1 mol.%), and concluded

that unlike the sodium and potassium nitrate phases, lithium nitrate is an unstable phase in the ternary system that starts to decompose to Li_2O_2 and LiO_2 at temperatures higher than 440 °C.



Figure 2. XRD patterns of MgO-SG sorbent doped with 15, 25 and 35 (mol, %) of: (**a**) NaNO₃, (**b**) ternary mixture of KNO₃, NaNO₃ and LiNO₃.

The MgO peaks are significantly sharper in the impregnated samples, suggesting that the impregnation influences the average size of MgO crystallite. It is usually expected that the sorbents with smaller MgO crystallites have higher surface areas and an enhanced reactivity; thus, the effect of impregnation on the growth of MgO crystallites is evaluated and correlated with the sorbent's performance along the carbonation–calcination cycles in Section 3.3. For instance, the average crystallite size of the MgO-SG sample is 10 nm, while the 15 Na-MgO presents a crystallite with 15 nm, and the MgO-SG impregnated with 15% of (Na, Li, K)NO₃ achieves a crystallite size of 26 nm. Such results indicate that the impregnation with alkali metal nitrates boosts the growth of the MgO crystals, which is more pronounced in the case of the impregnation does not seem to influence significantly the growth of the MgO crystallite. The MgO-SG sorbent doped with NaNO₃ shows an average crystallite size of ca. 15 nm and the MgO-SG sorbent doped with (Na, Li, K)NO₃ of ca. 28 nm regardless the molar percentage of AMS impregnation (Table 3).

AMS	Average Crystallite Size (nm) of MgO-SG Undoped and Doped with AMS (%, mol)						
	0	15	25	35			
NaNO ₃	10	15	15	16			
(Na, Li, K) NO ₃	10 –	26	28	29			

Table 3. Average crystallite size of MgO-SG undoped and doped with 15, 25 and 35% (mol) of AMS.

The supported sorbents, MgO-Ca-SG and MgO-Ce-SG, were doped with 15% of a ternary mixture of (Na, Li, K)NO₃. Figure 3 shows the X-ray diffraction peaks and Table 4 compares the MgO crystallite size for the unsupported and supported sorbents before and after the AMS doping.

The 15 (Na, Li, K)-MgO, 15 (Na, Li, K)-MgO-Ca and 15 (Na, Li, K)-MgO-Ce sorbents exhibit the characteristic peaks of the MgO pattern (42.9° , 62.2° , 78.5°). Moreover, the 15 (Na, Li, K)-MgO-Ca and 15 (Na, Li, K)-MgO-Ce also present the main peaks of the XRD pattern of CaCO₃ (29.4°) and CeO₂ (28.5°), respectively. Regarding the nitrate salts used in the impregnation, as justified above, the LiNO₃ was not detected by XRD. The main peaks of the KNO₃ pattern (23.5° and 29.4°) were identified for the 15 (Na, Li, K)-MgO and of the

15 (Na, Li, K)-MgO-Ca as well as the main peaks of the NaNO₃ pattern (29.4°, 31.9°, 38.9°, 47.9°). However, for the 15 (Na, Li, K)-MgO-Ce sorbent, as the characteristic peak of the CeO₂ XRD pattern is very wide, it is impossible to identify the peak at 29.4° of the KNO₃ and NaNO₃ patterns. Still, a small peak at 27° of the KNO₃ pattern is exhibited as well as the peaks at 31.9°, 38.9°, 47.9° related to the NaNO₃ pattern.



Figure 3. XRD patterns of MgO-SG, MgO-SG-Ca and MgO-SG-Ce before and after doping with 15(Na, K, Li)NO₃.

Table 4. Average crystallite size (nm) of unsupported and supported MgO-SG undoped and doped with 15% of (Na, K, Li)NO₃.

Unsupported and Supported MgO-SG Sorbents	Average Crystallite Size (nm) of Sorbents Undoped and Doped with 15% of (Na, K, Li)NO ₃				
	0	15			
MgO-SG	10	26			
MgO-SG-Ca	12	30			
MgO-SG-Ce	12	27			

Table 4 shows that the AMS doping influenced the crystallite size of supported sorbents, causing it to increase but in a similar way for both supports.

The specific surface area (S_{BET}) and the pore size distribution of MgO-SG, MgO-SG-Ca and MgO-SG-Ce before and after doping with 15 of (Na, K, Li)NO₃ was evaluated (Figure 4a,b) by the N₂ sorption technique.

The MgO-SG-Ce shows the highest S_{BET} (230 m² g⁻¹) followed by MgO-SG (187 m² g⁻¹) and the MgO-Ca-SG (144 m² g⁻¹). However, after doping, all the sorbents show a severe reduction in their S_{BET} , namely, 83%, 88% and 87% for 15 (Na, Li, K)-MgO, 15 (Na, Li, K)-MgO-Ca and 15 (Na, Li, K)-MgO-Ce sorbents, respectively. This evidence is in line with the previous conclusions from the XRD characterization, in which we observed an increase in the MgO crystallite size, indicating a reduced surface area available for the MgO sorbent to uptake CO₂.

Dal Pozzo et al. [11] reported a S_{BET} of 22 m² g⁻¹ for a MgO sorbent promoted with 10% of (Na, Li, K)NO₃, which is smaller than that of 15 (Na, Li, K)-MgO and 15 (Na, Li, K)-MgO-Ce sorbents but higher than that of 15 (Na, Li, K)-MgO-Ca. In addition, all the analyzed sorbents of the set also registered a decrease in the total pore volume after AMS doping. This is probably due to its partial occupation by the ternary mixture [17], which should affect the CO₂ uptake due to the higher occurrence of pores' blocking.



Figure 4. Textural properties of MgO-SG, MgO-SG-Ca and MgO-SG-Ce before and after doping with 15(Na, K, Li)NO₃: (**a**) specific surface area (S_{BET}) and total pore volume (V_p), (**b**) pore size distribution (PSD) estimated by the BJH desorption.

The PSD of the MgO-SG and MgO-SG-Ce sorbents (Figure 4b) indicates a predominance of the presence of mesopores (2–50 nm). There is also a small percentage of macropores (>50 nm). The MgO-SG-Ca consists mainly of a macroporous sorbent, although the MgO-SG-Ca also exhibits a small share of mesopores. After doping with 15(Na, K, Li)NO₃, all the sorbents are predominantly made up of macropores (>50 nm), but in case of 15 (Na, Li, K)-MgO-Ce, a small share of mesopores is still present.

3.2. Thermal Decomposition of Mono and Ternary Alkali Nitrates Salts

The mass loss (%), the first derivative of the mass change (DTG) and the heat flow profiles with temperature were obtained in the TGA, between 25 and 1000 °C (heating rate: 10 °C min^{-1}) under 100% of air or 100% of CO₂ flow, for pure NaNO₃ and a ternary mixture of 52 mol.% KNO₃, 18 mol.% NaNO₃ and 30 mol.% LiNO₃ (Figure 5a–c). The knowledge about the mono and ternary alkali nitrates thermal decomposition under different atmospheres contributes to the understanding of the AMS stability on the doped sorbents.



Figure 5. Thermogravimetric analysis: (**a**) thermal decomposition in mass loss (%), (**b**) first derivative of mass change and (**c**) heat flow profiles of NaNO₃ and (Na, Li, K)NO₃ with temperature.

Figure 5a shows a mass loss until 200 °C related with the release of the adsorbed water, since these salts are very hygroscopic [25]. It stands out that all the samples remain almost stable up to a temperature of around 600 °C, where an abrupt loss of mass is observed regardless of the atmosphere. The atmosphere affects the extent of the salt's decomposition: under pure air atmosphere, higher decomposition temperatures (~800 °C) are achieved when compared to those achieved for the case of 100% CO₂ atmosphere (~715 °C) [26]. Due to the formation of alkali carbonates with higher molar mass than the corresponding oxides, i.e., Na₂O, K₂O and Li₂O/Li₂O₂, the observed mass loss is lower under a CO₂ atmosphere.

The alkali carbonates are very stable at high temperature, especially under CO₂ atmosphere; i.e., Na₂CO₃ and K₂CO₃ start to decompose slowly above 900 °C [27] and, in the case of Li₂CO₃, the decomposition starts at 737 °C, but only at temperatures near 900 °C is the decomposition significant [26].

The decomposition temperature of each sample can be defined as the upper stability limit of a mixture (T3): that is, the maximum temperature at which it loses 3% of the initial weight [25]. The definition of the decomposition temperature allowed to set the working temperature range with the melting point and the T3 being the extremes of that interval [25]. In the present study, since we identified the presence of water in all the samples, the initial weight was replaced by the weight at 200 °C. The decomposition temperature range was also determined. The results obtained are summarized in Table 5.

Table 5. Thermal decomposition of mono and ternary alkali salts under air and CO₂ atmosphere: working range and decomposition range (°C).

Sample	Atmosphere	T3 (°C)	Melting Point (°C)	Working Range (°C)	Decomposition Range (°C)
NaNO ₂	Air	620	308 [11]	308–620	620-814
	CO ₂	581		308–581	581–731
(Na. Li. K)NO2	Air	575	120 [28]	120–575	575-818
CO_2 595		[]	120–595	595-736	

The results show that the ternary mixture offers a larger temperature window to operate when compared with the single salt, NaNO₃. This advantage relies mostly on that already described in the literature: the addition of the LiNO₃, which significantly decreases the melting temperature of the mixture with relation to the sodium nitrate salt, that is, 120 °C [28] against 308 °C [11], and not in the T3, which do not vary as much between both cases. Nevertheless, in air atmosphere, the eutectic mixture presents a T3 lower than the single salt, which is associated with the presence of Li that prematurely decomposes as stated by Bauer [28], but it does not happen under the CO₂ flow, evidencing that the atmosphere type is also a relevant parameter.

The heat flow TG curves presented in Figure 5c exhibit several up and downward peaks, whether an endothermic or exothermic heat event occurs, respectively, which allow properly identifying that the temperature occurs at each event. The analysis of both the heat flow profile (Figure 5c) and corresponding DTG curves (Figure 5b) reveals the existence of corresponding peaks in the same temperature intervals for all the samples (T3 \geq 575 °C) regardless of the atmosphere considered. In addition, it can be observed that the NaNO₃ melting (endothermic reaction) occurs around 310 °C (Figure 5c), which agrees with the literature [28]. In the case of ternary salt, it was difficult to identify its melting. Due to the water vaporization, which was confirmed by the mass loss in both salts, only a small peak around 129 °C was visible for (Na, Li, K)NO₃ salt, meaning that probably the water vaporization is masking the endothermic reaction associated with the ternary salt melting.

3.3. CO₂ Uptake TGA Studies

3.3.1. Assessment of the Effect of Mono and Ternary AMS Molar Fraction

Temperature-programmed carbonation–calcination followed by TGA studies were carried out with samples of MgO sorbent undoped and doped with different molar fractions of mono or ternary AMS to assess the effect of the promotors ratio (AMS/MgO) on the CO₂ uptake (Figure 6a,b).

Undoped MgO-SG sorbent immediately captures CO_2 in the beginning of the TGA test, achieving its maximum CO_2 uptake at 50 °C, and gradually releases it with the increasing of the temperature from 75 to 325 °C. The corresponding CO_2 desorption peak is significantly wider than those of the MgO-doped samples, which is probably due to the type of surface of each sorbent and, in turn, with the type of species being formed during

the CO_2 uptake. In contrast, the extent of CO_2 uptake is insignificant for all the doped MgO-SG sorbents at low temperatures until a certain value of temperature is reached, where it starts to uptake CO_2 significantly. According to the literature, in the case of the undoped MgO-SG sorbent, the CO_2 molecules were quickly adsorbed at the surface of the MgO particles to form an unidentate carbonate layer impermeable for gaseous reactants. On the other hand, in nitrate-promoted MgO, the molten layer dissolves the CO_2 . Since the diffusivities of the carbonate ions and the oxygen ions in MgCO₃ layers are higher in the absence of the unidentate layers, the formation of $MgCO_3$ occurs faster [11]. This enhancement is justified by the melting of the nitrates, since the CO₂ solubility increases in the phase transition from the solid to liquid state of the nitrates. In this way, the AMS impregnation is proved to increase considerably the CO₂ uptake by the MgO sorbents. The mentioned temperature was determined as the inflection point of each CO_2 uptake curve, that is, where the corresponding second derivate changes its signal. The different threshold temperatures were obtained for each sorbent (Supplementary Materials, Figure S1), and the results are summarized in Table 6. The sharp increase in the CO_2 uptake by the doped sorbents, with the temperature, can be attributed to the physical state of the corresponding AMS [29], ranging from solid to melting. At low temperatures, the CO₂ uptake is hindered due to the formation of a solid layer of nitrates that covers the surface of the sorbent. Only with the increasing temperature until close to the melting point of the AMS, this covering solid layer starts to melt. It is called the pre-melting phenomenon, where an interfacial liquidlike film of the partially disordered surface of the promoted sorbent is formed at temperatures below the corresponding melting point, enhancing the kinetic of the CO_2 uptake [29]. For instance, the melting point of the single salt NaNO₃ is 308 $^{\circ}$ C, but it suffers a solid-state transition from an ordered to a disordered rhombohedral structure at a lower temperature of 275 °C that, in turn, is believed to potentiate its pre-melting [11]. The present experiments indicate that this transition occurs at even lower temperatures (234–240 °C). In addition, the MgO-SG sorbent doped with single NaNO₃ appears to have a higher threshold temperature than the corresponding MgO-SG sorbent doped with the ternary mixture. Hence, it is suggested that the eutectic ternary mixture lowers the CO₂ uptake temperature. One study in the literature [11] describes a CO_2 uptake temperature of 160 °C using the ternary eutectic mixture of $(Na, Li, K)NO_3$. In this work, it was possible to reduce the temperature to around 175 °C. The difference between the inflection temperature and the temperature that corresponds to the maximum CO_2 uptake (mg CO_2 g⁻¹ sorbent) achieved during the temperature-programmed experiment suggests that at low temperatures, the accelerating effect of the addition of alkali metal nitrates on the CO₂ uptake by MgO is reduced, which is justified by the thermodynamic equilibrium of MgO/MgCO₃ [11].

The maximum instantaneous rate of change in temperature with respect to CO_2 uptake was obtained based on the upward peak of the first derivative of each TG curve of the tested sorbents (Supplementary Materials, Figure S1). The results indicate that the maximum CO_2 uptake rate temperature does not significantly vary with the AMS impregnation percentage for both sorbents (Table 6). The samples of MgO sorbent impregnated with the ternary mixture show lower maximum conversion temperatures than those impregnated with the single salt. As the former presents lower inflection temperatures, it is also possible to associate them with a larger CO_2 uptake temperature range, which extends the carbonation region. Moreover, the highest CO₂ uptake of 292 mg CO₂ g^{-1} sorbent was obtained for the MgO-SG sorbent doped with 35% of NaNO₃, which, in turn, corresponds to the highest percentage of NaNO3. This result suggests that the sorbents with higher percentages of NaNO₃ might favor the Mg^{2+} ions diffusion on the molten NaNO₃ [11]. For the ternary mixture, the highest CO₂ uptake was 239 mg CO₂ g^{-1} sorbent for the MgO-SG doped with 15% of AMS and decreased with the increasing AMS molar fraction. This suggests that a certain thickness of the molten layer covering the surface of the sorbent hinders the CO₂ uptake by increasing the mass transfer resistance. For the synthesized undoped MgO sorbent, there was found to be a maximum CO_2 uptake of 36 mg CO_2 g⁻¹ sorbent, which is better than the value found in the literature (19 mg CO_2 g⁻¹ sorbent) [11].



Figure 6. Profile of CO₂ uptake of undoped MgO-SG sorbent and doped with 15%, 25% and 35% of (a) NaNO₃ or (b) (Na, Li, K)NO₃ performed under a 100% CO₂ atmosphere.

Table 6. Properties of MgO sorbent undoped and doped with different molar fractions of mono or ternary AMS: maximum instantaneous rate of change in temperature with respect to CO₂ uptake (°C), inflection temperature (°C), maximum CO₂ uptake temperature (°C), CO₂ uptake range (°C) and maximum CO₂ uptake (mg CO₂ g⁻¹ sorbent).

	Undoped	Doped							
MgO-Based Sorbent		NaNO ₃	(mol, %)	(Na, Li, K)NO3 (mol, %)					
		15	25	35	15	25	35		
Maximum instantaneous rate of change in temperature with respect to CO_2 uptake (°C)	-	310	304	310	322	319	Peak not defined		
Inflection temperature (°C)	-	234	237	240	175	184	Not detected		
Maximum CO ₂ uptake temperature (°C)	-	399	396	395	391	388	388		
CO ₂ uptake temperature range (°C)	20–50	234–399	237–396	240–395	175–391	184–388	-		
Maximum CO_2 uptake (mg $CO_2 g^{-1}$ sorbent)	36	165	231	292	239	161	139		

Figure 7a,b shows the variation of the MgO crystallite size (CS) of the sorbents before and after the TG tests. Summarizing, the AMS impregnation contributes to the growth of the MgO crystallites with a higher increase in the MgO crystallite size when the impregnation is carried out using the ternary mixture of KNO₃, NaNO₃ and LiNO₃ than the single salt NaNO₃. The variation of the MgO crystallite size $(\Delta CS = CS_{dopedsorb} - CS_{undopedsorb}/CS_{undopedsorb} \times 100)$ of the sorbent doped with NaNO₃ varies between 50 and 60%, but it increases for values between 160 and 200% for the sorbent with higher (Na, Li, K)NO₃ content. Conversely, the samples impregnated with NaNO₃ have a greater increase in the crystallite size before and after the TGA tests, i.e., between 50 and 60%, and 7% to 15%, for NaNO₃ and (Na, Li, K)NO₃, respectively. The AMS molar percentage of the impregnation does not affect the crystallite size regardless of using the single salt or the ternary mixture.

The CO₂ uptake and the cyclic stability of the most promising doped sorbents identified above for both AMS doping, i.e., 25% and 35% for NaNO₃ and 15% and 25% of (Na, Li, K)NO₃, were tested in a TGA by performing five carbonation–calcination cycles. Figure 8a,b show the cyclic CO₂ carbonation–calcination profiles obtained in the TGA apparatus and Figure 9a,b show the respective carrying capacity and MgO conversion obtained for the MgO-SG sorbent impregnated with the single salt NaNO₃ (25 and 35%, mol) and with the ternary mixture of (Na, Li, K)NO₃ (15 and 25%, mol).



Figure 7. Crystallite size of undoped MgO-SG sorbent and doped with 15%, 25% and 35% of (**a**) NaNO₃ or (**b**) (Na, Li, K)NO₃, before (blue, violet, dark pink and light pink bars) and after (green bars) TG test between 20 and 500 °C.



Figure 8. Profile of carbonation (300 °C, 60 min) and calcination (445 °C, 60 min) tests carried out along 5 cycles in a TG (80 mL min⁻¹ of CO₂) for (**a**) 25% and 35% of NaNO₃ or (**b**) 15% and 25% (Na, Li, K)NO₃ impregnated MgO-SG sorbent.



Figure 9. Carrying capacity (mg CO₂ g⁻¹ sorbent) and MgO conversion (%) of (**a**) 25% and 35% of NaNO₃ or (**b**) 15% and 25% (Na, Li, K)NO₃ impregnated sorbents achieved along 5 carbonation–calcination cycles.

Both MgO-SG sorbents impregnated with 35% of NaNO₃ and with 25% of NaNO₃ exhibit a sharp decay on the CO₂ uptake in the 2nd cycle followed by a gradual reduction until the 5th cycle. The 25 Na-MgO registered an MgO conversion of 55% in the 1st cycle, of 35% in the 2nd cycle and of only 7% in the last cycle. The 35 Na-MgO achieved even higher reductions with an MgO conversion of 78% in the 1st cycle, 10% in the 2nd cycle and only 2% in the 5th cycle. A qualitative comparison of Figure 8a,b immediately reveals

that the MgO-SG samples impregnated with the ternary mixture of (Na, Li, K)NO₃ have a better performance than those impregnated with the single salt NaNO₃. Regardless of the molar percentage of ternary impregnation used, the two tested samples exhibited consistent peaks after five carbonation–calcination cycles, showing good CO₂ uptakes and cyclic stability. The 15 (Na, Li, K)-MgO reached a MgO conversion of 58% and of 42% in the 1st and 5th cycles, respectively. Meanwhile, the 25 (Na, Li, K)-MgO presented a MgO conversion of 64% in the 1st cycle and of 37% in the last cycle. Although the 25 (Na, Li, K)-MgO has the highest MgO conversion in the 1st cycle, it shows the lowest MgO conversion in the last cycle. Hence, between all the studied sorbents, the 15 (Na, Li, K)-MgO sorbent was considered the most promising since the MgO conversion values remained stable over five carbonation–calcination cycles. Subsequently, a TG test was carried out with 12 carbonation–calcination cycles using the 15 (Na, Li, K)-MgO sorbent (Figure 10a). Figure 10b plots the carrying capacity and the MgO conversion of the cyclic CO₂ carbonation–calcination test of the 15 (Na, Li, K)-MgO sorbent.



Figure 10. MgO-SG sorbent impregnated with 15% of (Na, Li, K)NO₃: (a) profile of carbonation (300 °C, 60 min) and calcination (445 °C, 60 min) and (b) carrying capacity (mg CO₂ g⁻¹ sorbent) and MgO conversion (%) for tests carried out along 12 cycles in a TG (80 mL min⁻¹ of CO₂).

The literature [11] reports a study with a MgO sorbent doped with 10% of (Na, Li, K)NO₃ that registered carrying capacities of 480 and 410 mg CO₂ g⁻¹ sorbent for the 1st and 5th cycles, respectively. Another article describes the carrying capacities of 440 and 352 mg CO₂ g⁻¹ sorbent for the 1st and 5th cycles, respectively, of a MgO sorbent impregnated with 15% of the same ternary mixture [19]. Since the 15 (Na, Li, K)-MgO sorbent synthesized in this work presents a carrying capacity between 460 and 330 mg CO₂ g⁻¹ sorbent from the 1st to the 5th cycle, it is clear that these results are in agreement with the results found in the literature. Dal Pozzo et al. [11] justify the MgO-based sorbents' deactivation along cycles with the nitrate's segregation. As shown in Figure 5a,b, the NaNO₃ and (Na, Li, K)NO₃ decomposition only starts above 600 °C, so these sorbents remain stable under the temperatures used for the above carbonation–calcination tests. However, since the AMS melting starts at lower temperatures (Table 5 and Figure 5c), the segregation of AMS can occur as it was demonstrated by SEM images for NaNO₃ [11] and in our previous work (unpublished), where the segregation of K particles was also demonstrated.

Due to the AMS segregation, the dispersion of MgO in the melted phase of the nitrates will be spoiled, and the CO₂ carrying capacity decreases. Apparently, under the used conditions, since the CO₂ uptake during the 1st carbonation is more effective and faster (the CO₂ uptake temperature range is lower) for the NaNO₃ salt, the nitrates segregation is more pronounced, and the sorbent's deactivation after the 1st cycle is abrupt.

In general, the 15 (Na, Li, K)-MgO sorbent exhibited a stable carbonation–calcination profile over the 12 cycles. Its MgO conversion dropped from 58% to 34% from the 1st to the 2nd cycle, but it gradually increased up to 48% until the 6th cycle, from where it progressively decreased until the last cycle, reaching a final MgO conversion of 33%. The decrease in MgO conversion is more significant after the 1st cycle, which can be correlated

with the fast initial MgO-CO₂ carbonation reaction, increasing the MgCO₃ content in the sorbent and contributing to the alkali's segregation [11]. Indeed, due to the segregation, the alkali will be less dispersed on the MgO particles, and its role as a phase transfer catalyst between bulk MgO and CO₂ molecules decreases alongside the CO₂ capture capacity. Thus, between the 2nd and the 6th cycles, the carbonation reaction occurred slowly, and more time was required for the carbonation stabilization as confirmed by the narrow peaks shown in Figure 10a and Figure S2 in the Supplementary Materials. However, it seems that there is an arrangement involving the alkalis and the MgO that allows the sorbent to recover some CO_2 carrying capacity. After the 7th cycle, the carbonation reached the stabilization faster (evidenced in Figure 10a and Figure S2 in the Supplementary Materials), meaning that the alkalis were still enhancing the kinetic stage of carbonation reaction, but the sorbent started to lose its capture capacity. Harada et al. [19] reported carrying capacities of 440 and 310 mg CO_2 g⁻¹ sorbent between the 1st and the 12th cycle for a MgO sorbent doped with 15% of (Na, Li, K)NO₃. In line with the literature, the 15 (Na, Li, K)-MgO synthesized in this work achieved a carrying capacity of 460 mg CO_2 g⁻¹ sorbent in the 1st cycle and 262 mg CO_2 g⁻¹ sorbent in the last cycle.

3.3.2. Assessment of the Effect of Inert Support Addition

Temperature-programmed carbonation–calcination studies followed by TGA were carried out on samples of supported MgO sorbents (Ca or Ce-based) undoped and doped with 15% of (Na, Li, K)NO₃ to assess the effect of the support on CO₂ uptake (Figure 11a,b).



Figure 11. Profile of CO₂ uptake of supported MgO sorbents undoped and doped with 15% of (Na, Li, K)NO₃: (**a**) Ca-based support or (**b**) Ce-based support.

Undoped MgO-SG-Ca and MgO-SG-Ce sorbents have their maximum CO2 uptake of 73 mg CO₂ g⁻¹ sorbent and 86 mg CO₂ g⁻¹ sorbent, respectively, at the beginning of the TGA test at 25 °C, and we observed a gradual CO₂ release with the increase in temperature until ~400 °C. Both supported sorbents exhibit CO₂ calcination peaks significantly wider than those of the corresponding unsupported AMS sorbents (Figure 6). Conversely, the CO₂ uptake of the 15 (Na, Li, K)-MgO-Ca and of the 15 (Na, Li, K)-MgO-Ce sorbents is negligible at low temperatures until it reaches the respective inflection temperature. This parameter was obtained as it was explained in Section 3.3.1, and the respective plots can be found in the Supplementary Materials, Figure S3. The maximum instantaneous rate of change in temperature with respect to CO_2 uptake (°C) was also determined, as it was explained above. Both parameters are summarized in Table 7. The promoted MgO-SG-Ce sorbent showed a higher inflection temperature than the promoted MgO-SG-Ca, meaning that the latter starts the CO_2 uptake sooner. As expected, the AMS doping enhanced considerably the CO₂ uptake of both MgO sorbents, 15 (Na, Li, K)-MgO-Ca and 15 (Na, Li, K)-MgO-Ce sorbents achieving a maximum CO_2 uptake of 299 mg CO_2 g⁻¹ sorbent and 252 mg CO₂ g^{-1} sorbent at 384 °C and 393 °C, respectively. Summarizing, undoped MgO-SG-Ce exhibited higher CO₂ uptake than MgO-SG-Ca, but AMS doping was more

effective for the latter, with 15 (Na, Li, K)-MgO-Ca exhibiting higher CO₂ uptake than 15 (Na, Li, K)-MgO-Ce. Moreover, 15 (Na, Li, K)-MgO-Ca also provides a more extensive carbonation than 15 (Na, Li, K)-MgO-Ce, since the former presents a wider carbonation temperature range than the latter.

Table 7. Properties of unsupported and supported MgO sorbents (Ca or Ce-based support) undoped and doped with 15% of (Na, Li, K)NO₃: maximum instantaneous rate of change in temperature with respect to CO₂ uptake (°C), inflection temperature (°C), maximum CO₂ uptake temperature (°C), CO₂ uptake range (°C) and maximum CO₂ uptake (mg CO₂ g⁻¹ sorbent).

	MgO-SG *	MgO-Ca-SG		MgO-Ce-SG				
MgO-Based Sorbent	% (Na, Li, K)NO ₃							
	15	0	15	0	15			
Maximum instantaneous rate of change in temperature with respect to CO_2 uptake (°C)	322	-	308	-	294			
Inflection temperature (°C)	175	-	153	-	210			
Maximum CO_2 uptake temperature (°C)	391	25	384	25	393			
CO_2 uptake temperature range (°C) Maximum CO_2 uptake (mg CO_2 g ⁻¹ sorbent)	175–391 239	- 73	153–384 299	- 86	210–393 252			

* Values from Table 6 for comparison.

Papalas et al. [20] report a CO₂ uptake of 317 mg CO₂ g^{-1} sorbent for an MgO sorbent promoted with 5% (mol) of CaCO₃ and impregnated with 20% of ternary mixture (Na, Li, K)NO₃ at 300 $^{\circ}$ C for 30 min under an atmosphere consisting of 30% of CO₂, which is in line with the CO₂ uptake obtained in the present work for the 15 (Na, Li, K)-MgO-Ca sorbent, despite the difference between the CO_2 atmospheres. Jin et al. [21] synthesized a supported sorbent with CeO_2 (10% mol) impregnated with a mixture of $LiNO_3$:NaNO₃:Na₂CO₃:K₂CO₃ = 0.2:0.76:0.04:0.5 (17% mol) MgO sorbent that achieved a CO₂ uptake of 450 mg CO₂ g⁻¹ sorbent at 325 °C for 120 min under 100% CO₂, which is almost double the CO₂ uptake obtained for the 15 (Na, Li, K)-MgO-Ce. It is important to note that in the above example a temperature plateau of 120 min (325 °C) was applied, while the present study does not include a temperature plateau. The 15 (Na, Li, K)-MgO-Ca sorbent starts to uptake CO_2 at a lower temperature (153 °C) than the 15 (Na, Li, K)-MgO-Ce (210 °C). However, the last one reaches the maximum instantaneous rate of change in temperature with respect to CO₂ uptake at a lower temperature (294 vs. 308 °C). The higher inflection temperature of 15 (Na, Li, K)-MgO-Ce was unexpected, since the same alkali ternary mixture was used. An explanation can be assigned to the high dispersion of both the CeO_2 and AMS on the sorbent. As it can be observed in the XRD patterns of the samples supported with Ce (Figure 3), broader peaks are observed, meaning there is a high dispersion of these compounds in the sorbent, which probably delays the molten effect of the ternary alkali salts mixture on the CO₂ uptake. Yu et al. [30] synthesized CeO₂-MgO and MgO sorbents and found that despite the similarity of their initial surface areas, the doped sorbent had an increased CO_2 uptake, which was justified by changes in the pore structures and by the increase in the basicity of the MgO phase induced by the addition of CeO₂. In the present study, the corresponding values of S_{BET} are also similar $(31 \text{ vs. } 32 \text{ m}^2 \text{ g}^{-1})$, and comparatively with the unsupported sorbent, the maximum CO₂ uptake of 15 (Na, Li, K)-MgO-Ce (239 vs. 252 mg CO₂ g^{-1} sorbent) was slightly improved. It must be noted that no temperature plateau was considered during this experiment. It is important to note that as shown in Table 7 and in accordance with the literature [2], the addition of both Ca and Ce-based supports improved the MgO sorbents' performance.

Figure 12a,b shows the variation of the MgO crystallite size of the undoped and doped sorbents with the Ca- or Ce- based support before and after the TG tests.

Both undoped MgO-SG-Ca and MgO-SG-Ce sorbents maintained the MgO crystallite size after undergoing the TGA test. The same behavior was already observed for the unsupported sorbent (Figure 7). For 15 (Na, Li, K)-MgO-Ca and 15 (Na, Li, K)-MgO-Ce, the MgO crystallites size increased 150% and 125% (Table 4), which was mainly due to the

AMS impregnation procedure, while after the TGA experiment, the increase was only 10% and 22%, respectively. Hence, the AMS impregnation with 15% (Na, Li, K)NO₃ affected identically the supported MgO-Ca-SG and MgO-Ce-SG sorbents.



Figure 12. Average crystallite size of supported MgO sorbents undoped and doped with 15% of (Na, Li, K)NO₃ before (yellow, blue, orange and mustard bars) and after (green bars) TG test between 20 and 500 °C: (a) Ca-based support or (b) Ce-based support.

The performance of supported sorbents along the cyclic carbonation–calcination cycles was assessed in TGA tests (Figure 13a,b).



Figure 13. MgO-SG supported sorbents (Ca- or Ce- based support) impregnated with 15% of (Na, Li, K)NO₃: (a) profile of carbonation (300 °C, 60 min) and calcination (445 °C, 60 min) and (b) carrying capacity (mg CO₂ g⁻¹ sorbent) and MgO conversion (%) for tests carried out along 12 cycles in a TG (80 mL min⁻¹ of CO₂).

The 15 (Na, Li, K)-MgO-Ca sorbent exhibited a more promising carbonation–calcination profile along the five cycles than the 15 (Na, Li, K)-MgO-Ce sorbent, since a higher CO₂ carrying capacity was achieved over cycles. However, both supported sorbents present quite stable peaks until the end of the test. Quantitively, the MgO-SG-Ca impregnated with 15% of (Na, Li, K)NO₃ obtained a carrying capacity of 375 mg CO₂ g⁻¹ sorbent in the 1st cycle and of 275 mg CO₂ g⁻¹ sorbent in the 5th cycle. Papalas et al. [20] reports a Ca-based supported MgO promoted with 20% of ternary mixture (Na, Li, K)NO₃ that varied its carrying capacity from 396 to 330 mg CO₂ g⁻¹ sorbent for the same number of cycles, which is in agreement with the obtained results. Jin et al. [21] report a Ce-based supported MgO sorbent promoted with 12% of ternary mixture of (Na, Li, K)NO₃ with carrying capacities of 340 and 180 mg CO₂ g⁻¹ sorbent between the 1st and the 5th cycles.

The same sorbent but with an impregnation molar percentage of 17% managed to achieve 420 mg CO₂ g⁻¹ sorbent in the 1st cycle and 330 mg CO₂ g⁻¹ sorbent in the 5th cycle. The 15 (Na, Li, K)-MgO-Ce sorbent presents an impregnation molar percentage of 15%, which is in between those mentioned in the literature, and it achieved a carrying capacity of 375 mg CO₂ g^{-1} sorbent, which is within the same range. In fact, the unsupported 15(Na, K, Li)-MgO sorbent performs better than the supported sorbents, since a higher carrying capacity was achieved for this sorbent, i.e., 460 and 330 mg CO_2 g⁻¹ sorbent for the 1st and 5th cycle, respectively. Nevertheless, after the 1st carbonation-calcination cycle, the 15 (Na, K, Li)-MgO-Ca is the unique one that maintains a similar carbonation rate profile along the cycles. The carbonation profiles (Figures 8 and 13a) show that the 1st carbonation rate is similar for all the sorbents, but after the 2nd cycle, the carbonation slope decreases, meaning that the carbonation rate is delayed, except for (Na, K, Li)-MgO-Ca. The carbonation reaction kinetics is one of the MgO-based sorbent's drawbacks, and the faster carbonation of 15 (Na, K, Li)-MgO-Ca is a huge advantage, which can contribute to a comparative upscale of MgO sorbents. Similar results were found by Cui et al. [31]; the authors performed kinetic studies with Ca and Mg precursors and verified that the activation enthalpy of CO₂ sorption for doped AMS-MgO sorbent was lower than for the undoped AMS-MgO, indicating that the CaCO₃ decreases the activation energy of CO₂ sorption on MgO. The CaCO₃ should act as a spacer among MgO crystallites, enhancing the AMS dispersion on the MgO surface area and making the carbonation reaction easier.

The positive effect of the calcium support was already evidenced by the lower inflection temperature of the 15 (Na, K, Li)-MgO-Ca sorbent relatively to 15 (Na, K, Li)-MgO (153 vs. 175 °C, Table 7) and the maximum CO₂ uptake obtained when no temperature plateau was applied (299 vs. 239 mg CO₂ g⁻¹ sorbent, Table 7), respectively.

4. Conclusions

This study evaluated the role of a NaNO₃ salt and a ternary mixture (NaNO₃, LiNO₃) and KNO_3 (18/30/52; % mol.)) molar fraction (15, 25 and 35%) on the MgO carbonation temperature range and on the maximum CO_2 uptake temperature. Regarding the carbonation temperature range, the ternary mixture offers a larger temperature window of operation when compared to that of the single salt $NaNO_3$, which is justified by the lower melting temperature of the ternary mixture (120 vs. 308 °C). Also, it is observed that the increase in the percentage of the NaNO₃ single salt between 15 and 35% allows enhancing the CO_2 uptake by the sorbent, but for the case of the ternary mixture sorbents, the opposite behavior is observed. This should be explained by the lower melting temperature of the ternary mixture that promotes the enhancement of the AMS dispersion on the MgO surface at the selected carbonation temperature (300 °C). Probably, the increase in the thickness of the molten layer covering the surface of the sorbent beyond a certain limit hinders the CO_2 uptake by increasing the mass transfer resistance, which justifies the enhanced results obtained with 15% of the ternary mixture. Despite the lower melting temperature of the ternary mixture, the inflection temperature only starts around 175 °C, suggesting that at low temperatures, the accelerating effect of the addition of the ternary alkali metal nitrates on the CO₂ uptake by MgO is reduced, which is justified by the thermodynamic equilibrium of MgO/MgCO₃.

Regarding the supported sorbents (Ca and Ce- based), both show a lower carrying capacity along the carbonation–calcination cycles than the unsupported sorbent. However, the 15 (Na, K, Li)-MgO-Ca sorbent shows a stable slope for the carbonation, while for the other sorbents, it decreases after the 1st cycle, meaning that the maximum carbonation rate was delayed. Thus, the carbonation performs faster for the 15 (Na, K, Li)-MgO-Ca sorbent, overcoming some kinetic limitations, which is considered as a huge advantage, and it can contribute for a sooner upscale of MgO sorbents for industrial CO₂ capture applications.

Supplementary Materials: The following supporting information can be downloaded at https://www.mdpi.com/article/10.3390/ma16247539/s1, Figure S1: Illustration of 1st and 2nd derivative of MgO sorbent doped with different molar fractions of mono or ternary alkali metal salts; Figure S2: Carbonation profile of MgO-SG sorbent impregnated with 15% of (Na, Li, K)NO₃; Figure S3: Illustration of 1st and 2nd derivative of supported MgO sorbent (Ca- and Ce- support) doped with different molar fractions of mono or ternary alkali metal salts.

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