Influence of Ca substitution by Mg on the Ca$_3$Co$_4$O$_9$ performances

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Ca$_{1-x}$Mg$_x$Co$_4$O$_9$ polycrystalline thermoelectric ceramics with small amounts of Mg have been synthesized by the classical solid state method. Microstructural characterizations have shown that all the Mg has been incorporated into the Ca$_3$Co$_4$O$_9$ structure and no Mg-based secondary phases have been identified. Apparent density measurements have shown that samples do not modify their density until 0.05 Mg content, decreasing for higher contents. Electrical resistivity decreases and Seebeck coefficient slightly raises when Mg content increases until 0.05 Mg addition. The improvement in both parameters leads to higher power factor values than the usually obtained in samples prepared by the conventional solid state routes and close to those obtained in textured materials.

Influencia de la sustitucion de Ca por Mg en las prestaciones del compuesto Ca$_3$Co$_4$O$_9$

Se han preparado cerámicas termoeléctricas policristalinas de Ca$_{1-x}$Mg$_x$Co$_4$O$_9$ con pequeñas cantidades de Mg, utilizando el método clásico de estado sólido. La caracterización microestructural ha mostrado que el Mg se ha incorporado en la estructura del Ca$_3$Co$_4$O$_9$ y que no se han formado fases secundarias con el Mg. Las medidas de densidad aparente han mostrado que las muestras prácticamente poseen la misma densidad hasta un contenido de 0.05 Mg, disminuyendo para contenidos mayores. La resistividad eléctrica disminuye y el coeficiente Seebeck aumenta cuando el contenido de Mg se incrementa hasta valores de 0.05. La mejora de ambos parámetros conduce a valores del factor de potencia mayores que los obtenidos en muestras preparadas por la vía convencional de estado sólido, alcanzando valores próximos a los que se encuentran en materiales texturados.

Palabras clave: Dopeado; Sinterizado; Cobaltitas; Propiedades eléctricas; Poder termoeléctrico.

1. INTRODUCTION

Thermoelectric (TE) materials are able to transform a temperature gradient to electrical power directly, without moving parts, due to the Seebeck effect. This intrinsic property has focused attention on these materials as they can be applied in waste heat recovery devices [1] or solar thermoelectric generators [2]. Furthermore, they can also be used as heating/refrigeration devices due to the Peltier effect [3]. The conversion efficiency of these materials is usually quantified by the dimensionless figure-of-merit ZT, $S^2/\rho k$ (in which the electrical part $S^2/\rho$ is also called power factor, PF), where $S$ is the Seebeck coefficient (or thermopower), $\rho$ the electrical resistivity, $k$ the thermal conductivity, and $T$ is the absolute temperature [4].

Taking into account the above expression for the figure-of-merit, high performance materials should possess a large Seebeck coefficient together with low electrical resistivity and thermal conductivity. The low electrical resistivity is necessary to minimize Joule heating, while the low thermal conductivity helps to maintain a large temperature gradient between the hot and cold sides in the thermoelectric device, increasing the developed voltage.

Nowadays the industrially applied TE devices are based on intermetallic materials, such as Bi$_2$Te$_3$ or CoSb$_3$, with high ZT values at relatively low temperatures. However, they cannot be applied in devices working at high temperatures under air due to their degradation and/or heavy metals evaporation. As a consequence, their ability to harvest energy from waste heat sources is strongly limited by their working temperatures. These limitations in temperature were overcome in 1997 by the discovery of relatively high thermoelectrical properties in a Co-based oxide ceramic, the Na$_2$Co$_3$O$_9$ [5]. From its discovery, many works have been performed on different cobaltite ceramic families as promising thermoelectric materials for high temperature applications, as the Bi-M-Co-O-based (M: alkaline earth) materials [6-8]. At the moment, research is mainly focused on Bi-free ceramic materials with high thermoelectric performances and higher thermal stability, such as Ca$_3$Co$_4$O$_9$ with promising thermoelectric properties [9,10].

Crystallographic studies performed on those Co-based materials have demonstrated that they present monoclinic
crystal structure which is, in turn, composed of two different layers. These layers show an alternate stacking of a common conductive CdI₂-type hexagonal CoO₂ layer with a two-dimensional triangular lattice, and a block layer composed of insulating rock-salt-type (RS) layers. The two sublattices (RS block and CdI₂-type CoO₂ layer) possess common a- and c-axis lattice parameters but different b-axis length, causing a misfit along the b-direction [11,12]. Furthermore, the high crystallographic anisotropy of these materials is reflected in their preferential growth habit along the ab plane (coincident with the conducting plane), which leads to the formation of plate-like grains during the sintering process. This preferential growth habit can be exploited to preferentially align the grains using physical, mechanical and/or chemical processes. Such processes allow the alignment of the conducting planes producing macroscopic properties comparable to the obtained on single crystals, by decreasing electrical resistivity. Numerous methods have been reported to be efficient in producing well aligned bulk materials, in these or in similar anisotropic systems, such as hot uniaxial pressing [13], spark plasma sintering (SPS) [14], microwave texturing [15], laser floating zone melting (LFZ) [16,17], electrically assisted laser floating zone (EALFZ) [18,19], templated grain growth (TGG) [20], etc. The main advantage of these techniques is the production of materials with very low electrical resistivity values due to the preferential alignment of conducting planes with the current flow direction. On the other hand, they possess some drawbacks, as the high price of the spark plasma device and the relatively long processing time of the hot uniaxial pressing or the TGG. In the case of the processes involving samples melting, it has been reported that the electrical properties are strongly dependent on the growth speed [15,21-23]. All these drawbacks have, as a consequence, an important influence on the industrial interest for the materials processed by these techniques.

It has also been reported that the Seebeck coefficient values are governed by the incommensurability ratio and/or the charge of the RS block layer between the CoO₂ ones [24]. This effect provides the basis for the modification of thermoelectric properties of a given material via chemical substitutions, as Sb⁺⁺ for Ca⁺⁺ in Ca₂Co₃O₇ [25], or Pb⁺⁺ for Bi⁺⁺ in Bi₂SrCo₂O₆ or Bi₂CaCo₂O₆ [26-28]. Other improvements can be produced by metallic Ag additions where an increase on the materials performances is produced by raising the electrical connectivity between the grains [29,30], or substituting Co³⁺⁺ by other transition metal, as Cu⁺⁺ or Cr⁺⁺ in Ca₃Co₄O₉ [31,32].

The aim of this work is to study the effect of small amounts of Mg⁺⁺ for Ca⁺⁺ isovalent substitution on the microstructure and high temperature thermoelectric properties of Ca₃₋ₓMgₓCo₄O₉ prepared by the simple classical solid state synthesis route. The structural and microstructural modifications produced by the Mg addition will be related with the changes on the thermoelectric performances.

2. EXPERIMENTAL

Ca₃₋ₓMgₓCo₄O₉ polycrystalline ceramic materials, with x = 0.00, 0.01, 0.03, 0.05, and 0.07, were prepared by the conventional solid state route using CaCO₃ (Panreac, 98 + %), MgO (Panreac, 98 + %), and Co₃O₄ (Aldrich, 98 + %) commercial powders as starting materials. They were weighed in the appropriate proportions, well mixed and ball milled for 30 minutes at 300 rpm, in acetone media, in an agate ball mill. The obtained slurry has been heated under infrared radiation during about 30 minutes to produce the acetone evaporation, leaving a homogeneous powder mixture. This dry mixture was then manually milled in order to avoid the presence of agglomerates in the next steps. After milling, the homogeneous mixture of fine powders was thermally treated twice at 750 and 800 ºC for 12h under air, with an intermediate manual milling in order to assure the total decomposition of calcium carbonate, as reported previously [10], which improves the mixture reactivity during the sintering processes. After the thermal treatments, the powders were uniaxially pressed at 400 MPa for 1 minute in order to obtain green ceramic parallelepips (−3 mm × 2.5 mm × 14 mm), with an adequate size for their thermoelectric characterization. These green ceramics were subsequently sintered at 910 ºC for 24 h, typical temperatures used in this kind of compounds [31].

Powder X-ray diffraction (XRD) patterns have been systematically recorded in order to identify the different phases in the thermoelectric materials after sintering procedure. Data have been collected at room temperature, with 2θ ranging between 5 and 60 degrees, using a Rigaku D/ max-B X-ray powder diffractometer working with Cu Kα radiation. Apparent density measurements have been performed by measuring the dimensions and weighting the samples for three times to avoid errors. At least four sintered samples of each composition were used to obtain the mean apparent density values and compared with the theoretical density (4.677 g/cm³) [33].

Microstructural observations were performed on the surface of samples, using a Field Emission Scanning Electron Microscope (FESEM, Carl Zeiss Merlin) fitted with an energy dispersive spectrometry (EDS) analyzer. Several micrographs of the samples have been used to analyze the different phases composition and their distribution. Electrical resistivity and Seebeck coefficient were simultaneously determined by the standard dc four-probe technique in a LSR-3 measurement system (Linseis GmbH), in the steady state mode and at temperatures ranging from 50 to 800 ºC under He atmosphere. From the resistivity values, the activation energy and the relative carrier concentration for each sample have been estimated. Moreover, with the electrical resistivity and thermopower data, the power factor has been calculated in order to determine the samples performances. These properties have been compared with the results obtained in the undoped samples and with those reported in the literature at low temperatures (∼50 °C), where oxygen diffusion is negligible, to avoid the influence of the atmosphere on the compared values.

3. RESULTS AND DISCUSSION

Powder XRD patterns for the different Ca₃₋ₓMgₓCo₄O₉ samples are displayed in Fig. 1 (from 5 to 40° for clarity). From these data, it is clear that all the samples have very similar diffraction patterns. In these plots, the peak marked by 9 in Fig. 1e belongs to the (111) diffraction plane of Si, used as internal reference. In Fig. 1a, corresponding to the undoped samples, major peaks have been associated with the thermoelectric Ca₃Co₄O₉ phase, indicated by its reflection
planes, in agreement with previously reported data [34]. When Mg is added to the samples, the peaks corresponding to the Ca₃Co₂O₆ secondary phase (shown by * in Fig 1e) [34] increase their intensity, indicating the raise of this phase contents. On the other hand, careful observation of these diagrams shows that there is no evidence of other secondary phases, especially Mg-containing ones, which is a clear indication that Mg has entered into the Ca₃Co₄O₉ and/or Ca₃Co₂O₆ phases.

Fig. 2 shows SEM micrographs performed on the surface of some representative samples corresponding to the 0.01 (a) and 0.05 (b) Mg substituted ones. In the figure, it can be clearly seen that all samples are composed by randomly oriented plate-like grains with similar sizes. When observing the samples in more detail, it has been found that major phase is the thermoelectric Ca₃Co₄O₉ one (grey contrast, #1) which can be observed as well defined plate-like grains. Ca₃Co₂O₆ secondary phase (#2) can be also seen in all samples as a very slightly darker grey contrast, easily distinguishable by their shape (as illustrated with Fig. 2c), but in a much lower proportion than the main

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<tr>
<th>Nominal composition</th>
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<tr>
<td>Ca₃₋ₓMgₓCo₄O₉</td>
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Figure 1. Powder X-ray diffraction patterns obtained for the Ca₃₋ₓMgₓCo₄O₉ samples; x = 0.00 (a); 0.01 (b); 0.03 (c); 0.05 (d); and 0.07 (e). The diffraction planes indicate the Ca₃Co₄O₉ phase and those marked by * the Ca₃Co₂O₆ one. The # symbol denotes the (111) plane of Si used as internal reference.

Figure 2. Surface SEM micrographs of Ca₃₋ₓMgₓCo₄O₉ samples for x = 0.01 (a) and 0.05 (b) showing the randomly oriented plate-like grains. The two different contrasts have been associated with: 1) Ca₃Co₄O₉ and 2) Ca₃Co₂O₆ phases. Close view (c) of one Mg-substituted sample showing with an arrow the Ca₃Co₂O₆ phase.
phase. These observations are in agreement with the above discussed XRD data. On the other hand, EDS analysis has shown that Mg incorporation is produced in the $\text{Ca}_3\text{Co}_4\text{O}_9$ crystal structure by substituting Ca (see Table I).

Other interesting feature which can be observed in Fig. 2 is that the samples are not fully dense, presenting porosity which can be observed in the micrographs as black contrast. This is due to the relatively low temperature stability of the thermoelectric phase (maximum stability temperature $\approx 926^\circ\text{C}$), compared with the minimum temperature to produce a liquid phase ($\approx 1350^\circ\text{C}$) [34]. The great difference between both temperatures leads to a very slow densification process at the sintering temperatures ($\approx 910^\circ\text{C}$), explaining the porosity obtained in these samples.

When observing the samples at higher magnification, some small differences can be detected in those with the highest Mg addition (0.07 Mg). This is illustrated in Fig. 3 where a high magnification micrograph performed on the surface of this sample is presented. As it is evident in the micrograph, a new type of grains is appearing (indicated by #3). These grains possess a shape which is different to the observed for the phases #1 and #2, and they can be seen as more or less triangular grains. Furthermore, EDS analysis has shown that their composition corresponds to a Mg-rich and Ca-deficient phase (Table I). In any case, this phase is in a very small proportion which explains the reason why it has not been detected with the XRD technique.

In order to evaluate the samples density, apparent density measurements have been performed on at least 4 samples for each composition. They were measured and weighed for three times to minimize the errors. The calculated apparent densities, together with their standard error, are represented in Fig. 4 vs. Mg contents. In the figure, it is clear that samples density is maintained approximately constant until 0.05 Mg-doping, and significantly decreasing for the 0.07 Mg containing sample. Moreover, the standard error is increased for these samples, indicating that the values dispersion is higher than for the other samples. These differences on the apparent density imply that they are $\approx 75\%$ of the theoretical one for all the samples except for the 0.07 Mg-doped one which is decreased to $\approx 70\%$. These density variations for the low doped samples are small enough to assume that density values practically have no effect on the electrical behaviour of the samples except in the case of the highest Mg-doped samples.

Fig. 5 shows the variation of the Seebeck coefficient with the temperature, as a function of the Mg doping. In the plot, it can be clearly seen that the sign of the thermopower is positive for the entire measured temperature range, which confirms a conduction mechanism mainly governed by holes. The values of the Seebeck coefficient increase with temperature, with similar behaviour for all the samples. The main differences between samples are observed at high temperatures, as the room temperature values are approximately the same for all samples.
samples, independently of Mg content. This measured value (around 135 $\mu V/K$) is slightly higher than those reported elsewhere for Mg-doped samples ($\sim 125 \mu V/K$) at 200 °C [35]. At high temperatures, undoped samples and 0.01Mg ones show the same values in all the measured temperature range due to the very small amount of Mg, which is not enough to modify Seebeck coefficient values. The effect of Mg can only be seen in samples with 0.03 and 0.05 Mg which slightly increase the Seebeck values at high temperatures, compared with the lower doped samples. This behaviour is in agreement with previously published results which show that the decrease on the rock salt layer size promotes higher Seebeck values [24]. Finally, for the highest Mg content samples the Seebeck coefficient decreases, due to the higher content of Ca$_3$Co$_2$O$_6$ secondary phase and the formation of isolated Ca$_{3-}$Mg$_{0.15}$Co$_4$O$_9$ grains which probably do not contribute to the measured Seebeck coefficient values. The maximum measured Seebeck coefficient value ($\sim 210 \mu V/K$) at 800 °C corresponds to the 0.03 Mg-doped samples higher than the obtained for the undoped ones. Moreover, 5 values at 625 °C ($\sim 195 \mu V/K$) are also higher than the best values obtained for undoped Ca$_3$Co$_2$O$_6$ samples sintered and textured by spark plasma sintering (170-175 $\mu V/K$) at the same temperature [36]. The relatively similar values obtained for the undoped and the Mg-doped samples indicate that Mg doping does not affect, in a great extent, the Ca$_3$Co$_2$O$_6$ conduction band [32].

The temperature dependence of electrical resistivity, as a function of the Mg content, is shown in Fig. 6. As can be clearly seen, the $\rho$ ($T$) curves show a decrease of the measured resistivity from 0.00 to 0.05 Mg substitution. Higher Mg contents increases resistivity, compared with the other Mg-doped samples. These curves reflect a slope change at about 400 °C, from semiconducting-like ($dp/dT < 0$) to metallic-like ($dp/dT > 0$) one. All the Mg-substituted samples present lower resistivity values than the undoped ones. The lowest measured room temperature resistivity values ($\sim 15 \Omega \cdot cm$ for the 0.05 Mg-substituted samples) is around the best values obtained for Ca$_3$Co$_2$O$_6$ samples sintered and textured by spark plasma sintering (15-18 $\Omega \cdot cm$) [36] and about 30 % lower than the undoped samples prepared in this work.

In order to explain the resistivity behaviour of these samples, it should be considered that the temperature dependence of the electrical conductivity, in the semiconductor behaviour zone, can be described as:

$$\sigma_T = \exp\left(\frac{-E}{k_B T}\right)$$

where $E$, $k_B$, and $T$ are the activation energy, Boltzmann constant, and absolute temperature, respectively. The activation energy values were obtained from the log ($\sigma_T$) versus $1/T$ plot as the curve fit slope in the different samples below $T_1$, as illustrated in Fig. 7. $T_1$ is defined as the temperature where the behaviour of the samples changes from semiconducting-like to metallic-like one. The calculated values have been found to be around 40 meV in all cases, confirming that Mg addition does not modify appreciably the Ca$_3$Co$_2$O$_6$ conduction band. On the other hand, two differences between doped and undoped samples can be pointed out from this graph. Firstly, Mg addition shift the $T_1$ to lower temperature values, indicating a change to metallic-like behaviour at lower temperature due to the fact that Mg doping increases the metallic characteristics of the compound (see $T_1^*$ and $T_1^*$ in Fig. 7, corresponding to the undoped and Mg-doped samples, respectively). Secondly, the carrier concentration is increased when Mg is added to the samples, as the log ($\sigma_T$) versus $1/T$ curves appear at higher values, until 0.05 Mg addition, decreasing for higher Mg contents. Under the assumption that these changes in conductivity are mainly due to variations on the hole concentration, the relative increase,
compared with the undoped samples, has been estimated and displayed in Fig. 8. From this figure, it is clear that the raise in Mg contents increase the hole carrier concentration until a maximum of around 45 % for the samples with 0.05 Mg substitution. Further Mg addition decreases the conductivity due to an increase of defects in the crystal structure reflected in lower carrier mobility, avoiding the effect of a higher carrier concentration.

In order to evaluate the thermoelectric performances of these materials, the power factor has been calculated. The temperature dependence of the power factor, calculated from the data represented in Figs. 5 and 6, is plotted in Fig. 9. When considering PF values at around 50 ºC (~ room temperature), it can be clearly seen that all the Mg-doped samples possess higher PF values than the undoped ones. The maximum increase is obtained for the 0.05 Mg-doped samples (~ 45 % higher than the undoped ones). The highest PF value obtained at 800 ºC (around 0.28 mW/K•m) for the 0.05 Mg-doped samples is relatively close to the values obtained in textured samples produced by spark plasma sintering (~ 0.32 mW/K•m) [36]. This 0.28 mW/K•m maximum value is a very important result when taking into account that the Mg substituted samples prepared for this work possess randomly oriented grains and, as a consequence, they should show much lower thermoelectric performances than the textured materials.

4. CONCLUSIONS

This paper demonstrates that Mg can substitute Ca in Ca$_{x}$Mg$_{y}$Co$_{4}$O$_{9}$ in small proportions (x ≤ 0.05) without appreciably modifying the crystal structure and improving the thermoelectric properties. Further Mg addition slightly diminishes thermoelectric performances. The optimal Mg for Ca substitution has been determined using the values of the power factor at 50 and 800 ºC, which is maximum for the 0.05 Mg-doped samples with values around 0.13 and 0.28 mW/K•m, respectively, which are about 45 and 55 % higher than the obtained for the undoped samples at these temperatures. Moreover, the value at 800 ºC is very close to the obtained in textured samples produced by spark plasma sintering under pressure. This value is very important for samples which possess randomly oriented grains.

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