Effect of Strontium on Structure and Superficial Area of La$_2$O$_3$

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ABSTRACT

Recently, lanthanum oxide doped by group IIA elements may strongly influence solid state reaction. A series of samples, noted LaSrX (where X = % atomic of strontium) have been prepared by hydrolysis, in neutral medium from La$_2$O$_3$ and SrCO$_3$. These samples were calcined under air at 450°C and 1150°C then characterized by specific surface area (BET), X-ray diffraction, scanning electron microscopy (SEM), thermogravimetry (TG) and differential thermal analysis (DTA). Obtained results show that after calcinations at: 1) 450°C, the addition of strontium is without effect on surface of La$_2$O$_3$ and the XRD analysis revealed no interaction between lanthanum oxide and strontium carbonate; 2) 1150°C, the sintering of the samples is very important, reduction of 70% of surfaces compared to the samples calcined at 450°C; XRD results show that LaSrX are formed principally by two oxides: SrLa$_2$O$_4$ and La$_2$O$_3$, which is confirmed by SEM method. The reaction between La$_2$O$_3$ and SrCO$_3$ showed three endothermic weight losses; elimination of water, a partial de-hydroxylation of La(OH)$_3$ and formation of La$_2$O$_2$CO$_3$ and La$_2$(CO$_3$)$_3$.

Keywords: La$_2$O$_3$; Strontium; Calcination; Surface; Structure; TG-DTA

1. Introduction

The literatures [1-5] show that the rare earth oxides (REO) are very interesting in the conversion of greenhouse gases; particularly the valorisation of methane by direct oxidation or dry reforming and the reduction of nitrogen oxides. The ceria and lanthanum oxides are the most used and studied in catalysis. The literatures show that these oxides suffer significant damage under the reactions conditions: lower activities, reduction in surfaces… Recently we showed that the barium stabilizes La$_2$O$_3$ by its interaction and insertion [6].

We are interested in the effect of the elements of the group IIA on the REO; in this paper we reported the results of a study on the effect of strontium on La$_2$O$_3$.

2. Experimental

2.1. Preparation of the Samples

The samples are obtained according to the method used previously [6,7] in our laboratory. These samples are noted LaSrX, X (= 0, 5, 10, 15 and 20) indicates the atomic percentage of strontium (Table 1). These samples are then calcined under air (3 L/h) at 450°C and 1150°C during 3 hours.

2.2. Characterizations of the Samples

2.2.1. Surfaces Specific Measurements (BET)

Measurements of surfaces samples are carried out, by adsorption of N$_2$ at 77 K, on 1 gram of powder using Micrometrics ASAP 2000 apparatus.

2.2.2. Analysis by X-Rays Diffraction (XRD)

The apparatus used is a standard diffractometer Philips PW 1800 with cylindrical chamber of 36 mm. The tension of this apparatus is of 40 kV, the current is around 30 mA. The radiations are the K$_α$ of copper: K$_{α1}$ (1.554056 Å) and K$_{α2}$ (1.54439 Å); these can be easy separated because the ratio of their intensities is about 0.500. A nickel filter is used to eliminate the radiation K$_β$ from copper. The spectra obtained are indexed according to JCPDS file (Joint Committee one Powder Diffraction Standards).

Table 1. Composition of LaSrX samples.

<table>
<thead>
<tr>
<th>% Sr(at)</th>
<th>0</th>
<th>5</th>
<th>10</th>
<th>15</th>
<th>20</th>
</tr>
</thead>
<tbody>
<tr>
<td>La$_2$O$_3$</td>
<td>LaSr0</td>
<td>LaSr5</td>
<td>LaSr10</td>
<td>LaSr15</td>
<td>LaSr20</td>
</tr>
</tbody>
</table>

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2.2.3. Scanning Electron Microscopy (SEM)
A JEOL JSM-35 scanning electron microscope (SEM) was employed to study the particle morphology. Prior to SEM observations, samples were deposited from ethanolic solution on to holey-carbon copper grids.

2.2.4. Thermogravimetric Analysis (TG and DTA)
The thermobalance used is a SETARAM TG-DTA92-16 including an electric balance with continuous signal, a furnace equipped with a regulator-programmer of temperature and a recorder. Measurement is made under oxidizing atmosphere (air: 3 L·h⁻¹) with a speed of heating of 10°C·min⁻¹ from ambient to 1200°C.

3. Results and Discussion
3.1. Effects of Strontium on Surfaces of La₂O₃
The calcinations at 450°C showed that the strontium addition is without effect on surfaces of lanthanum oxide (Figure 1). Surfaces of the LaSrX samples are practically the algebraic sums “Equation 1” of surfaces of La₂O₃ and SrCO₃. This result is in favour of the segregation of lanthanum (III) and strontium carbonate.

\[ S = (1 - X) \cdot S_{LaO} + X \cdot S_{SrCO_3} \]  

Nevertheless, one notes a light surface enrichment of strontium for the samples LaSr5 and LaSr10, starting from X = 20 this tendency is reversed.

At 1150°C, the addition of 5 and 10% of strontium make slightly increase the surface of lanthanum oxide. This increase is 3.37% and 9.16% respectively for LaSr5 and LaSr10. Beyond 15% no effect on surface of La₂O₃ is detected. Reported on Figure 1, the measurements of surfaces show a surface enrichment with lanthanum for the samples LaSr5 and LaSr15 and particularly for LaSr10.

3.2. Characterization of the Samples by XRD and Microscopy SEM
3.2.1. Samples Calcined at 450°C
The XRD patterns of LaSrX calcined at 450°C are reported on Figure 2. The analysis of these spectra shows that the samples are mixtures of composed containing lanthanum (carbonates, oxides, oxy-hydroxides, hydroxides and oxy-carbonates) and of strontium carbonate. La(OH)₃ is in a majority [6] and crystallizes according to a hexagonal system pertaining to the space group P6₃mmc with parameters a = 4.06 Å and c = 6.43 Å, in accordance with JCPDS files. These results showed no interaction between lanthanum oxide and strontium carbonate. Sure enough the comparison between the spectra of the LaSrX samples and that of LaSr0 indicates a whole of lines at 2θ = 25°, 26°, 37° and 44° (Figure 2) specific to SrCO₃.

The intensity of these lines increases with X, the addition of strontium. These results are in agreement with those of the literature [8]. After calcinations at 450°C, one observes partial or total de-hydroxylation of the hydroxide of lanthanum La(OH)₃ and probably according to following reactions:

\[ \text{La(OH)₃} \rightarrow \text{LaO(OH)} + \text{H}_2\text{O} \]  
\[ 2 \text{LaO(OH)} \rightarrow \text{La}_2\text{O}_3 + \text{H}_2\text{O} \]  
\[ 4\text{LaO(OH)} + \left[ \text{CO}_2, \text{CO}_3^{2-} \right] \rightarrow \text{La}_2\text{O}_2\text{CO}_3 + \text{La}_2\left(\text{CO}_3\right)_3 + 2\text{H}_2\text{O} \]  

The formation of lanthanum oxy-carbonates is very frequent when La₂O₃ is used in catalysis [9-11]. Indeed
La$_2$O$_3$ activates CO$_2$ in the form of La$_2$O$_2$CO$_3$ during the reactions using CO$_2$ [12,13]. XRD patterns of the samples show clearly the crystalline phase of La$_2$O$_2$CO$_3$ (Figure 2) this one is promoted by the presence of strontium.

3.2.2. Samples Calcined at 1150°C

The analysis of XRD patterns reported in Figure 3, shows that samples are formed principally by two oxides: a mixed compounds with structure of SrLa$_2$O$_4$ and La$_2$O$_3$ [6]; the La$_{4}$Sr$_{2}$O$_{9}$, La$_{8}$SrO$_{7}$ and La$_{2}$Sr$_{2}$O$_{5}$ appears after treatment at 1500°C [14].

The interaction and the insertion of strontium in La$_2$O$_3$ are confirmed by the absence of the position of SrO on the spectra of the LaSrX samples. The literature [2] shows that the strontium oxide is not detected by XRD after calcinations between 800°C and 950°C. This result are confirmed by scanning electron microscopy, the SEM of sample LaSr15 calcined at 1150°C (Figure 4), shows porous phase with two homogenous regions probably for SrLa$_2$O$_4$ and La$_2$O$_3$.

3.3. Thermogravimetric Analysis

Thermogravimetric measurements (TG and DTA), of samples from ambient to 1000°C, are reported on Figures 5(a)-(f).

3.3.1. TG and DTA of the Precursors Alone

The TG curve, represented on the Figure 5(a), indicates three losses of mass for LaSr100. The first change of mass of about 0.53%, which intervenes with 100°C, corresponds at the beginning of the water of hydratation. One second loss of mass of 15% starts as of 480°C to finish with 620°C approximately. In this temperature range, it has formation of SrO$_2$ according to the “Reaction (5)”; the decomposition of SrCO$_3$ in SrO and CO$_2$ usually proceeds towards 1300°C [13]

$$\text{SrCO}_3 \rightarrow \text{SrO} + \text{CO} \quad (5)$$

DTA curves show, in this range of temperature, three peaks of endothermic reactions. The first two fine peaks, towards 480°C - 540°C, correspond to the phase shift of SrCO$_3$ [15]; the third towards 600°C is due to the reaction (5). Indeed, the TG shows that the ratio

$$\frac{\Delta m_{\text{measured}}}{\Delta m_{\text{calculated}}}=0.8$$

is close to the stoichiometry of CO.

The third change which takes place towards 900°C shows a loss of mass of 0.62% which could be allotted to a sub-
The formation of La$_2$O$_3$ is obtained after two stages “Reactions (7) and (8)” of the de-hydroxylation of La(OH)$_3$. Between 350°C and 400°C a loss of mass, about 7.76%, is attributed to the appearance of lanthanum oxy-hydroxides, and that of La$_2$O$_3$ worms 500°C. Finally between 650°C and 800°C, a loss of mass of 3.18% could be due to the de-carboxylation of La$_2$O$_2$CO$_3$.

For LaSr0, the Figure 5(b) shows four losses of mass corresponding to four reactions.

The first, below 300°C indicates the drainage of water.

The formation of La$_2$O$_3$ is obtained after two stages “Reactions (7) and (8)” of the de-hydroxylation of La(OH)$_3$. Between 350°C and 400°C a loss of mass, about 7.76%, is attributed to the appearance of lanthanum oxy-hydroxides, and that of La$_2$O$_3$ worms 500°C. Finally between 650°C and 800°C, a loss of mass of 3.18% could be due to the de-carboxylation of La$_2$O$_2$CO$_3$. 

Figure 5. (a): TG and DTA of LaSr100; (b): TG and DTA of LaSr0; (c): TG and DTA of LaSr5; (d): TG and DTA of LaSr10; (e): TG and DTA of LaSr15; (f): TG and DTA of LaSr20.
according to the “Reaction (9)”. This compound comes from the easy carbonation of lanthanum oxide and of the tri-hydroxide La(OH)₃ with the free air [8].

\[
\begin{align*}
\text{La(OH)}_3 & \rightarrow \text{LaO(OH)} + \text{H}_2\text{O} & (7) \\
2\text{LaO(OH)} & \rightarrow \text{La}_2\text{O}_3 + \text{H}_2\text{O} & (8) \\
\text{La}_2\text{O}_2\text{CO}_3 & \rightarrow \text{La}_2\text{O}_3 + \text{CO}_2 & (9)
\end{align*}
\]

3.3.2. TG and DTA of LaSrX Samples

Measurements TG and DTA of the samples LaSrX (X = 5, 10, 15 and 20) are reported on the Figures 5(c)-(f). The TG shows four stages of losses of mass. First is due to the removal of water towards 100°C. The last one towards 900°C is very weak and does not exceed the 0.25% for the four samples. As for the two other significant changes of mass one notes two different behaviours. Indeed, the DTA shows two simple peaks 380°C and the other towards 560°C for the samples LaSr5 and LaSr10; on the other hand for LaSr15 and LaSr20 the second peak presents two distinct shoulders. For the first couple of samples it had two reactions and four reactions for the second couple of samples.

- Samples LaSr5 and LaSr10

Towards 560°C, the change of mass is neither representative of La(OH)₃ (Figure 5(b)) or that of SrCO₃ (Figure 5(a)).

La(OH)₃ exit of the de-hydroxylation of La(OH)₃ reacts with SrCO₃ to form a mixed oxide according to “Reaction (10)”.

\[
2\text{LaO(OH)} + \text{SrCO}_3 \rightarrow \text{SrLa}_2\text{O}_4 + \text{CO}_2 + \text{H}_2\text{O} & (10)
\]

- Samples LaSr15 and LaSr20

Three peaks DTA, represented on the Figures 5(e) and (f), which makes the superposition of three pic, it can be to explain by the two phases shifts of strontium carbonate and by the departure of CO coming from the decomposition of SrCO₃ according to the “Reaction (5)”. Nevertheless, the formation of the mixed compound SrLa₂O₄ Reaction (10) is not excluded. This reaction would be supported by an acid catalysis of the ions La³⁺. The quantity of this mixed oxide would be weak and equivalent to that obtained during the treatment of LaSr5 and LaSr10.

Lastly, it should be noted that contrary to the samples LaSr5 and LaSr10, quantity of water which is eliminated from the samples LaSr15 and LaSr20 towards 100°C is relatively more significant. This water accounts for 1.5% (Δm/m) of the treated mass.

4. Conclusions

A series of samples, LaSrX where X is the atomic percentage in strontium was prepared by hydrolysis of La₂O₃ and SrCO₃ in neutral medium. These samples was calcined under air at 450°C and 1150°C, then characterized by BET specific surface area, XRD analysis and SEM microscopy. The reactivity of lanthanum oxide and SrCO₃ was followed by TG and DTA.

After calcination at 450°C, the addition of strontium is without effect on surfaces of lanthanum oxide. The XRD patterns shows data for crystalline phases of La₂O₃, La(OH)₃, LaO(OH) and La₂O₂CO₃ and of strontium carbonate. There are no interaction between lanthanum oxide and strontium carbonate.

After calcinations at 1150°C, the addition of 5% and 10% of strontium make slightly increase the specific surface of lanthanum oxide, at this temperature the sintering is important, 70% of values of the surfaces are decreases compared to those calcined at 450°C. The XRD spectra shows that LaSrX samples are formed by two oxides: a mixed compounds with structure of SrLa₂O₄ and La₂O₃. This result are confirmed by SEM micrography.

The TG curves of pure La₂O₃ and SrCO₃ indicate that, SrCO₃ decompose at 620°C approximately and the weight losses of La₂O₃ result for the drainage of water and the partial de-hydroxylation of La(OH)₃ with formation of La(OH). The reactivity of LaSrX showed three endothermic weight losses; elimination of water, a partial de-hydroxylation of La(OH)₃ and formation of La₂O₂CO₃ and La₂(CO₃)₃. In the range of temperature of 500°C - 580°C the DTA indicates, a first allotropic transition from the SrO₂ and its decomposition on CO and SrO₂ for LaSr15 and LaSr20.

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