

Effect of aluminum precursor on physicochemical properties of Al₂O₃ by hydrolysis/precipitation method
Efecto del precursor de aluminio en las propiedades fisicoquímicas de la γ -Al₂O₃ por el método hidrólisis/precipitación

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Resumen

Este estudio reporta la síntesis de alúmina nano-fibrilar mesoporosa preparada por vía de hidrólisis-precipitación. Se realizó a partir de una solución acuosa de reactivo analítico (AR) de sulfato de aluminio y se comparó con un sulfato de aluminio grado técnico (TG) de baja pureza bajo condiciones similares usando amoníaco como agente precipitante. Las propiedades fisicoquímicas de estas muestras se estudiaron mediante diversas técnicas de caracterización, como el análisis térmico termogravimétrico y diferencial (TG/DTGA-DTA), difracción de rayos-X (DRX), Isotermas de adsorción-desorción N₂, Espectroscopía infrarroja (FTIR), tamaño de partícula y Microscopía Electrónica de Transmisión con análisis de Energía Dispersiva de rayos-X (TEM/EDAX). Los resultados de TG-DTA y XRD muestran una mayor estabilidad y una cristalinidad ligeramente mayor en la muestra de Al₂O₃-TG que Al₂O₃-AR. Los resultados de adsorción-desorción de N₂ muestran que ambos materiales tienen una alta área superficial de 311 m²/g para Al₂O₃-TG y 272 m²/g para Al₂O₃-AR exhibiendo características de materiales mesoporosos. Los resultados FTIR muestran un bajo porcentaje de grupos OH superficiales para Al₂O₃-TG, mostrando una acidez menor debido a la baja concentración de especies Al-OH (Al^{IV}). Las mediciones de TEM confirmaron que el tamaño de las fibras varió de 20 a 100 nm para Al₂O₃-TG y 20-80 nm para Al₂O₃-AR. El análisis EDAX muestra la presencia de 0.33% peso de Mg como impureza en Al₂O₃-TG. Se atribuye que esta cantidad es suficiente para generar defectos estructurales y disminuir ligeramente la acidez. Además, extendió la cadena fibrilar de la alúmina de Al₂O₃-TG.

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Abstract

This study reports the synthesis of mesoporous nano-fibrillar alumina prepared by hydrolysis-precipitation route. It was done from aqueous solution of aluminum sulfate analytical reagent (AR) and compared to aluminum sulfate technical grade (TG) of low purity under similar conditions using ammonia as the precipitating agent. The physicochemical properties of these samples were studied with the assistance of characterization techniques such as Thermogravimetric and differential thermal analysis (TG/DTGA–DTA), X-ray diffraction (XRD), N₂ adsorption-desorption isotherms, Fourier transform infrared (FTIR) spectroscopy, particles size and Transmission electron microscopy with energy-dispersive X-ray analysis (TEM/EDAX). The TG-DTA and XRD results show greater stability and a slightly greater crystallinity in Al₂O₃-TG sample than Al₂O₃-AR. N₂ adsorption-desorption results show for both materials greatly surface area of 311 m²/g for Al₂O₃-TG and 272 m²/g for Al₂O₃-AR exhibiting characteristics of mesoporous materials. The FTIR results show a low percentage of surface OH groups for Al₂O₃-TG showing a lower acidity due to the low concentration of Al-OH species (Al^{IV}). TEM measurements confirmed that fibers size ranged from 20 to 100 nm, for Al₂O₃-TG and 20-80 nm, and for Al₂O₃-AR. EDAX shows the presence of 0.33 % wt. of Mg as an impurity in Al₂O₃-TG. It is attributed that this amount is sufficient to generate structural defects and decrease acidity slightly. Likewise, it extended the fibrillar chain of the alumina.

Introducción

Alumina is one of the most widely used materials as support in heterogeneous catalysis, adsorbent, electronic, abrasives, reinforcement of ceramic composites for its high strength, corrosion resistance, low thermal conductivity, and good electrical insulation due to several attractive features (Glorias *et al.*, 2014; Yang, 2011, Jiratova *et al.*, 1986). It is inexpensive and available in many different qualities making it suitable in a wide range of different chemical reactions. Furthermore, its synthesis is simple and economic. Though mainly used as support, alumina has also some catalytic activities by its own, this due to its acidity (Ancheyta *et al.*, 2005). For specific applications, the surface area, the pore size, pore volume, thermal and mechanical stability of the support it has a large impact on the catalytic effect. Several Authors have emphasized that γ -Al₂O₃ with high pore volume may improve the adsorption performance if used as an adsorbent, and loading capacity if applied as a catalyst support, to minimize diffusion and transport influences (Xu

et al., 2016, Stanislaus *et al.*, 2002). The alumina is a material that still stands in limelight owing to its extensive application.

The used aluminum oxides exist in a variety of metastable structures which are usually called transitions alumina such as η , χ , κ , θ , λ , δ and γ , as well as the stable α - Al_2O_3 phase. Especially, the variety of γ -alumina is widely appreciated due to its excellent properties as surface chemical properties that can be either acidic or basic depending on the transition alumina structure and the degree of hydration and hydroxylation of the surface, which is a major attraction from catalytic/adsorption point of view (Sinfontes *et al.*, 2014; Zhang *et al.*, 2016).

Nowadays, there are many materials as aluminum sources to synthesis and obtain Al_2O_3 , for example $\text{Al}(\text{NO}_3)_3$, $\text{Al}_2(\text{SO}_4)_3$, $\text{NH}_4\text{Al}(\text{SO}_4)_2$, AlCl_3 , NaAlO_2 , among others. The mesoporous γ - Al_2O_3 synthesized with the above aluminum sources has excellent properties (Zhang *et al.*, 2016). However, these aluminum sources are expensive, because of that; it is difficult to realize the industrialization with these ways. However, aluminum sulfate with the presence of impurities can be attractive, because of its low cost. The presence of impurities can influence in alumina quality and consequently affect the oxide morphology and/or physicochemical properties. Interestingly, even a tiny amount of impurities in the Al_2O_3 structure, can exert a strong effect on acidity, including structural defects. On other hand, a small amount of impurity could be detrimental for the performance. However, not all impurities are harmful; there are impurities that may present advantages benefiting the reaction and/or application. Among the impurities contained in alumina powders we can find: Na, Si, Fe, Ca, Mg, etc. Moreover, depending upon the synthesis method, alumina can contain anion impurities from the precursors used in the synthesis as for xample Cl^- ions, NO_3^- or $(\text{SO}_4)^{2-}$, something which is accepted by most of the people working in the field (Mishra *et al.*, 2002).

Herein, we describe the synthesis and comparison of physic-chemicals features through several characterization techniques of two alumina materials prepared in similar conditions. One of them synthesized from an aqueous solution of aluminum sulfate analytical grade (Al_2O_3 -AR) and a second alumina synthesized from aluminum sulfate technical grade with presence of impurities (Al_2O_3 -TG), both alumina obtained by hydrolysis-precipitation route. We obtained materials with relatively high surface areas, with pore sizes and high pore volumes. Emphasis was put on the impurities of aluminum precursor and on the physic-chemical properties of the Al_2O_3 . This paper contributes to a basic understanding of the purity of the aluminum salt and how it

influences on textural characteristics of mesoporous materials with the presence of 0.33 % wt. of Mg. This fundamental knowledge could be exploited in the design of new materials.

Materials and Method

Materials

In the present study, the starting materials to obtain hydrated alumina (Al₂O₃) were Al₂(SO₄)₃•18H₂O with a purity of 95% (by weight, wt.) technical grade (TG), Alfa-Omega chemical, S. A. and an Al₂(SO₄)₃•18H₂O with a purity of 99.98% (wt.), analytical reagent (AR), CTR Scientific, S.A., deionized water (Karat S. A.) and anhydrous NH₃ gas (Praxair, 99.98% (wt.)) as precipitant. Then, the impurities of both materials of aluminum sulfate are described (Table 1). Table 1. Chemical composition wt. % by atomic adsorption of Al₂(SO₄)₃-TG, Alfa omega, S.A., and Al₂(SO₄)₃-AR, CTR Scientific, aluminium precursors.

Precursor	unsolvable	chlorides	Calcium	Magnesium	Potassium	Sodium	heavy metals	Iron
Al ₂ (SO ₄) ₃ TG	4±0.5%	ND	ND	0.33	ND	0.03	ND	0.03
Al ₂ (SO ₄) ₃ AR	0.003	<0.001	<0.0002	<0.0001	0.0002	<0.0075	<0.001	0.002%

**ND: Not Determined

Synthesis of pseudoboehmite

The flow chart of the preparation procedure is depicted in Figure 1. Al₂(SO₄)₃•18H₂O TG and/or AR were dissolved into deionized water. The Al₂(SO₄)₃•18H₂O technical grade was filtered for removal of insoluble impurities. Precursors were prepared by dropping saturated solution of aluminum sulfate into a solution of water and ammonia gas under rigorous magnetic stirring at temperature of 50-70 °C. The precipitated solution was filtered; the precipitated pseudoboehmite was rinsed with distilled water three times respectively, and then dried at 110 °C for 12 h. The phase change of pseudoboehmite to γ-Al₂O₃ was performed by thermal treatment at 500 °C for 7 h. The materials were labeled as follows: Al₂(SO₄)₃•18H₂O Technical Grade (Al₂O₃-TG) and Al₂(SO₄)₃•18H₂O Analytical Reagent (Al₂O₃-AR).

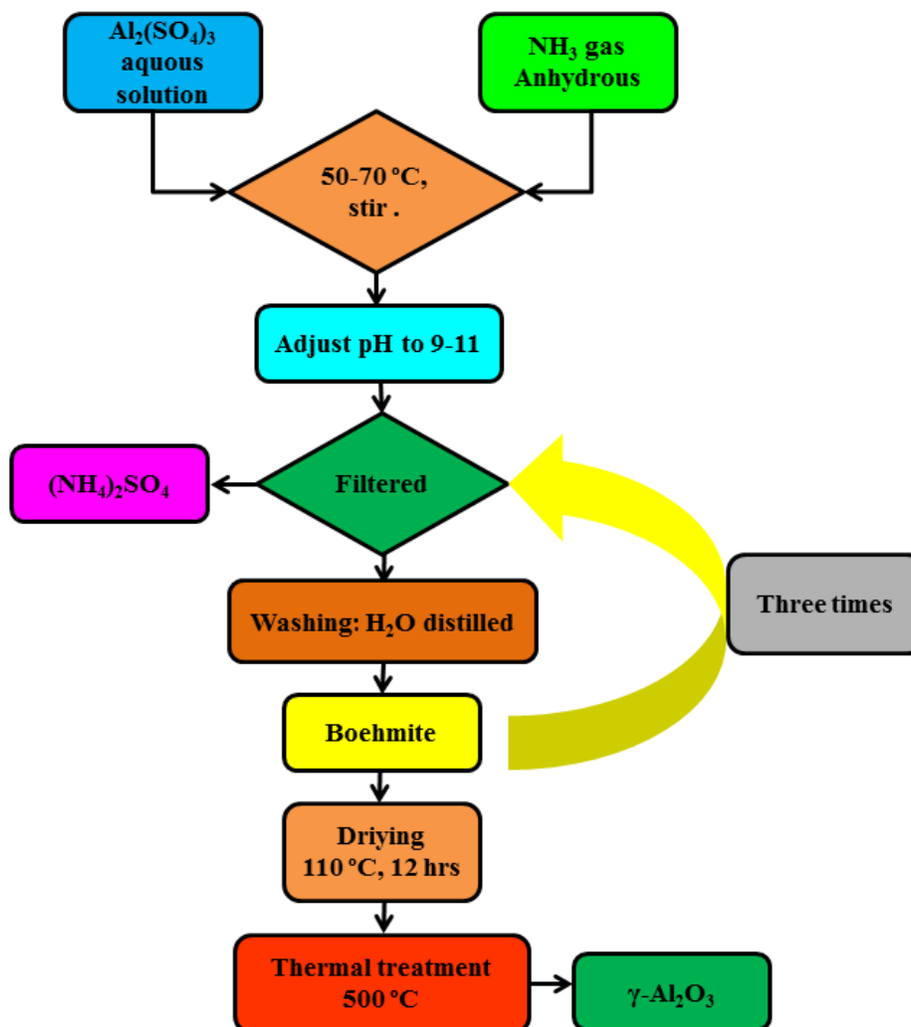


Figure 1. Flow chart of the preparation procedure of γ - Al_2O_3 .

Materials characterization

Thermal analysis TGA-DTA

The thermal stability of the samples was carried by Thermogravimetric and Differential thermal analysis (TGA/DTGA-DTA) was performed from 30 °C to 1000 °C in a Thermogravimetric analyzer SDT Q600 V20.5 Build 15 under air flow (100 mL / min) at a heating rate of 10 °C/min.

N₂ physisorption

The textural properties were determined by N₂ adsorption (Micromeritics, ASAP 2010). The samples were degassed at 200 °C for 3 h under vacuum. Nitrogen adsorption isotherms were measured at liquid N₂ temperature (77 K) and N₂ pressures ranging from 10⁻⁶ to 1.0 P/P₀. Surface

area was calculated according to Brunauer–Emmett–Teller (BET) method and the pore size distribution was obtained according to the Barret–Joyner–Halenda (BJH) method.

X-ray diffraction (XRD)

The crystalline properties of the samples were determined at room temperature by using X-ray powder diffraction. A Siemens D-500 diffractometer equipped with a Cu K α radiation anode was used for these measurements under the following conditions: sweep of 10-80° at an angle 2 θ , with applied voltage of 30 kV and current of 20 mA.

FT-IR analysis

The vibrational spectroscopy measurements were performed at room temperature by Fourier transform infrared (FTIR) spectroscopy on a Bruker Model 27 spectrometer through KBr (99%) disk method in a frequency range of 4000 - 400 cm⁻¹.

Transmission electron microscopy (TEM/EDAX)

The morphology was studied by transmission electron microscopy (TEM) using a Philips Tecnai F20 microscope operating with a 200 kV equipped with energy dispersive X-ray spectroscopy (EDAX) for the determination of the chemical composition.

Particle size

The particle size of the materials was measured by electroacoustic, the technique with a particle size analyzer (AcoustoSizer II, ESA; Colloidal Dynamics, USA) to pH neutral.

Results and discussion

TEM/energy-dispersive X-ray (EDAX) analysis

The surface morphology of Al₂O₃-TG and Al₂O₃-AR samples is studied by TEM and the corresponding micrographs are presented in Figure 2. It can be seen that the products obtained represent typical TEM images of Al₂O₃ powders. They exhibit disordered morphology nanofibrillar with a length of about 20-100 nm for the Al₂O₃-TG and 20-80 nm for Al₂O₃-AR. The products obtained in our experiment are similar as those reported in the literature (Zhang *et al.*, 2016). The development tendency of alumina is to be a more thermodynamically favored crystal

(Nuru *et al.*, 2015; Yu *et al.*, 2012). TEM images indicate alumina products have better dispersion performance for Al₂O₃-TG, which indicate alumina product obtained has good tendency to be a thermodynamic favored crystal due to presence of the Mg as impurity. Figure 2(c-d) shows the secondary particles size of the obtained products. The size varied from 0.45 μm to 0.85 μm for Al₂O₃-TG and from 0.35 μm to 0.65 μm for Al₂O₃-AR, observing a larger size for the sample Al₂O₃-TG, this is attributed to larger sizes of fibers that tend to form agglomerates or higher particles.

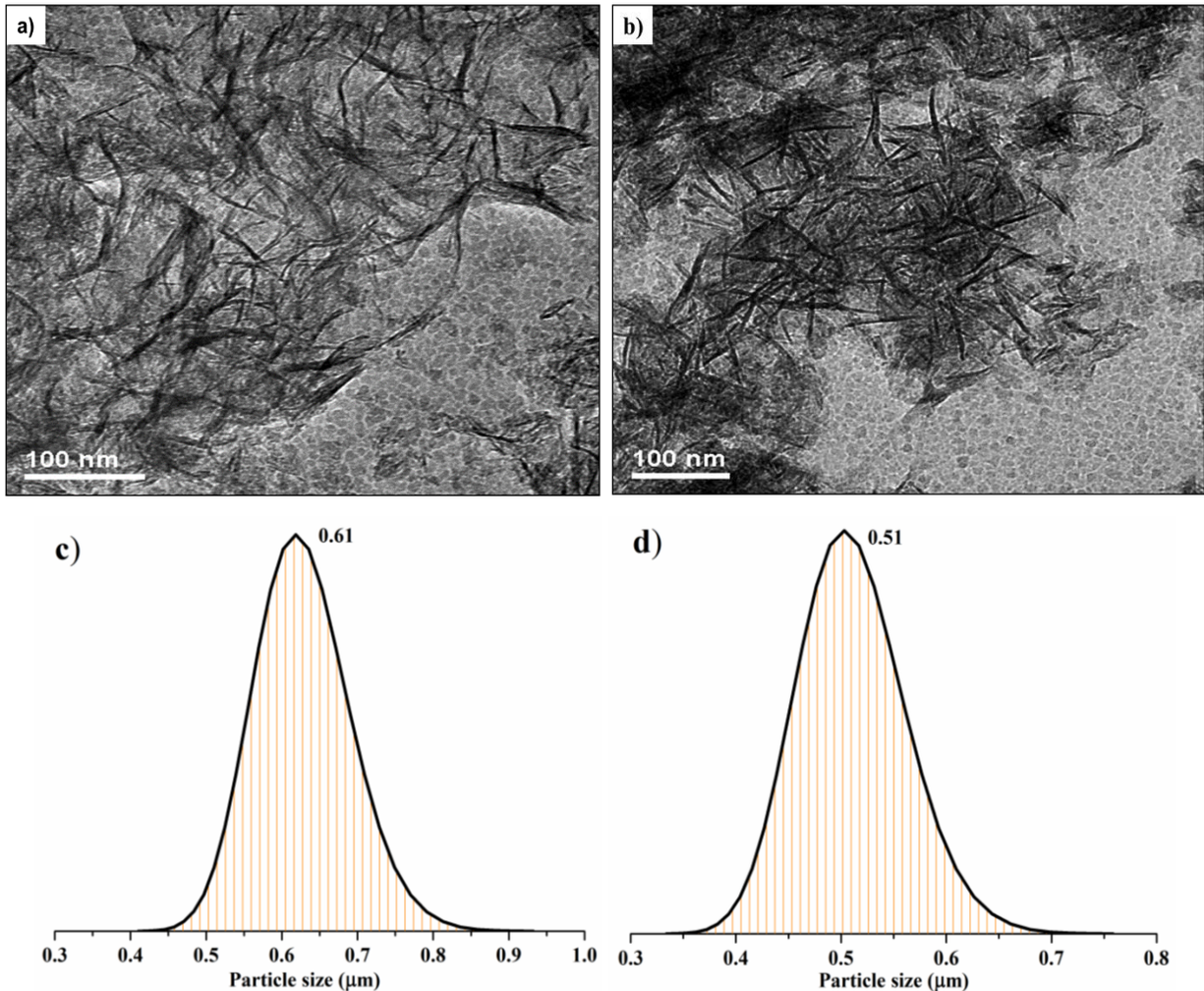


Figure 2. TEM images of a) Al₂O₃-TG, b) Al₂O₃-AR and particles size c) Al₂O₃-TG, d) Al₂O₃-AR

Energy dispersive X-ray (EDAX) data of both samples are provided in Table 2. The punctual analysis (EDAX) shows Al and O were detected on surfaces, respectively. The determination of

other present chemical elements as N and S are attributed to residues generated during the synthesis process. While than the presence of Mg is attributed as an impurity present in the low purity aluminum sulfate. The Mg was only possible to be observed since a micro-area analysis was performed. However, the assumption in this paper is that the trace element is homogeneously distributed on the matrix and the limit is expressed as an atomic fraction, this attributing that the Na and Fe were removed during the synthesis process.

Table 2. Average elemental analysis of Al₂O₃-TG and Al₂O₃-AR.

Elements: Atomic % by EDAX							
Sample	O K	Al K	C K	Mg K	N K	S K	Total
Al ₂ O ₃ -TG	57.96	15.58	24.28	0.20	2.84	0.07	100
Al ₂ O ₃ -AR	47.46	37.68	14.76	-	-	0.09	100

*(-): No detected

Thermogravimetric and differential thermal (TGA/DTGA–DTA) analysis

TGA/DTGA and DTA analysis of Al₂O₃-TG and Al₂O₃-AR are shown in Figure 3. Two endothermic peaks are observed in the DTA curve at 85 and 500 °C, the position of which varies with the type of the purity of aluminum salts that was used. The first peak is assigned to elimination of adsorbed water. At 500 °C pseudoboehmite decomposition to the γ -phase is produced; this involves dehydroxylation/dehydration [2AlO(OH) \rightarrow γ -Al₂O₃+H₂O]. The derivative of thermogravimetric analysis DTGA and DTA shows a peak centred at 85 °C indicating a thermal decomposition for Al₂O₃-TG. The weak broad DTGA and sharp DTA peaks around 500 °C imply a transformation phase. These facts are confirmed by the thermogravimetric study, which shows a complete weight loss (17%) at 500 °C, this is consistent with the reported in the literature (Urretavizcaya *et al.*, 1998). The Al₂O₃-AR shows two endothermic peaks, which are observed in the DTA curve at 60 and 500 °C. The first peak is assigned to elimination of adsorbed water. At 500 °C pseudoboehmite decomposition to the γ phase is produced. These facts are confirmed by the thermogravimetric study, which shows a complete weight loss (15%) at 500 °C.

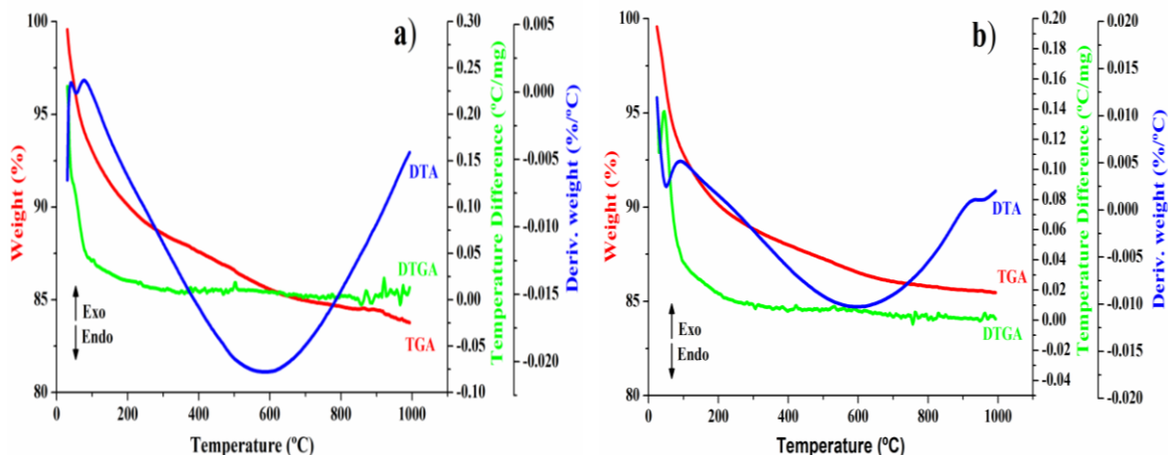


Figure 3. TGA/DTGA and DTA curves of a) Al₂O₃-TG and b) Al₂O₃-AR.

The endothermic peak observed in the DTA curve at 950 °C, Figure 3b, which is not accompanied by a weight loss (TGA), which is corroborated by DTGA, is attributed to θ -Al₂O₃ formation according to the reported in the literature (Jan *et al.*, 2014). Thermal transformations γ -Al₂O₃ → δ -Al₂O₃ → θ -Al₂O₃ → α -Al₂O₃ are not detected by DTA or TGA. In these transformations a small amount of energy is involved and no weight loss is produced. The temperature of phase transition is directly proportional to the grain size of the precursor. However, it is observed that Al₂O₃-TG exhibits greater thermal stability attributed to the presence of Mg. In fact several authors demonstrated that the presence MgO plays a structural role in the Al₂O₃ that alter its nature (Chen *et al.*, 2016).

Powder X-ray diffraction (XRD)

The high-angle XRD patterns of Al₂O₃-TG and Al₂O₃-AR samples are shown in figure 4. The X-ray diffraction pattern was performed of 10 – 80° at an angle 2 θ . The results of the XRD patterns wide angle shows seven weak diffraction peaks, those are observed at 2 θ = 19.2, 31.0, 36.6, 39.3, 46, 61.5, and 67°, which can be indexed as the (111), (220), (311), (222), (400), (511), and (440) reflections of γ -alumina according to JCPDS card: 100425 (Cheng *et al.*, 2006; Jan *et al.*, 2014). When the precursors are calcined at 500 °C, there are no obvious peaks of others phases in XRD pattern indicating that the products structure is amorphous.

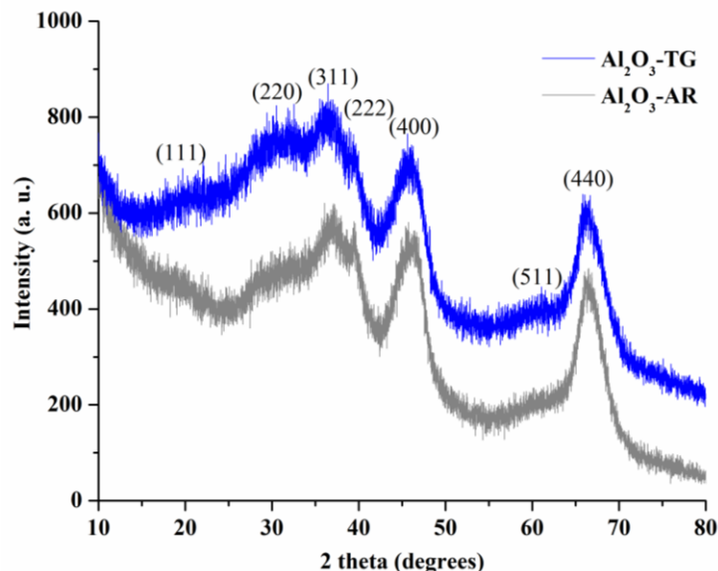


Figure 4. XRD patterns of a) Al₂O₃-TG and b) Al₂O₃-AR.

On other hand, a disorder in the structure can be observed. This disorder is reflected in the unusual broadening of some X-ray diffraction peaks for the Al₂O₃-TG material in the region $2\theta=20-48^\circ$. This is attributed to a strong faulting on the (111), (110), and (100) in the spinel-like structure of these materials, which occur as individual and/or interconnected defects on different plane families (Sakashita *et al.*, 2001). According to the literature, this non spinel model is more stable than spinel type structures (Wolverton *et al.*, 2000). The XRD results confirm the TG-DTA analyses indicating that the Al₂O₃-TG is more stable than Al₂O₃-AR. However, in this work the structural defect is attributed to the presence of Mg. This is due to that excess alumina introduced a substitution defect on the tetrahedral magnesium site and an aluminum vacancy or magnesium vacancy is formed to compensate the positive charge excess, as reported in the literature (Yuji *et al.*, 2006).

Adsorption/desorption analysis

The nitrogen adsorption isotherms and BJH pore size distribution of samples calcined at 500 °C are shown in Figure 5. It is seen that the two materials exhibit the shape of type-IV Isotherm curves according to the IUPAC classification (Sing *et al.*, 1985; Jan *et al.*, 2014), which is characteristic of mesoporous materials. These materials show irregular shape isotherms with hysteresis loop type E and indicate that the pores in the material have an inkwell-type shape. Existence of the hysteresis loops is ascribed to the capillary condensation of N₂ gas occurring in the pores (Zhang *et al.*, 2016).

Characterizations of powders obtained at 500 °C and values reported in the literature by different authors from different routes of synthesis were shown in Table 3. Surface area and pore volume values of the materials were higher for the sample Al₂O₃-TG than for Al₂O₃-AR. Due to that exhibit nanofiber with a higher length, as it is shown in TEM images, it is attributed to the presence of Mg in the structure of the Al₂O₃-TG. A predominance of mesopores that could be of interest for heterogeneous catalysis and the adsorption area is observed. In addition, the intercrossing degree and disorder among the fibers (interfibrillar porosity) as well as their length may cause the non-uniform pore size distributions.

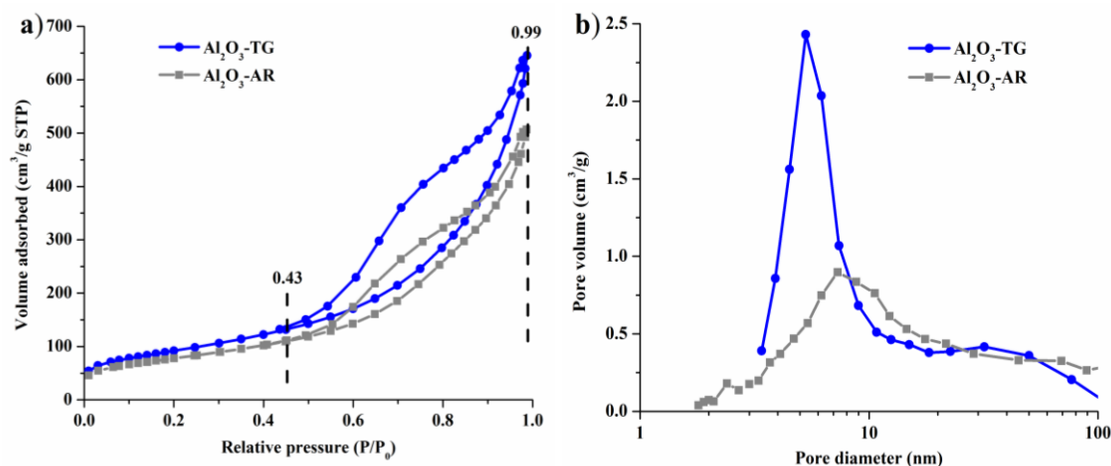


Figure 5. a) N₂ adsorption–desorption isotherm and b) pore size distribution curve of Al₂O₃-TG and Al₂O₃-AR.

Table 3. Results of the properties textural of samples obtained and reported by different synthesis routes.

Samples	SSA ^a	Vap ^b	Dap ^c	References
γ-Al ₂ O ₃ -TG (Hydrolysis-precipitation)	311	0.86	11.6	This work
γ-Al ₂ O ₃ -AR (Hydrolysis-precipitation)	272	0.76	10.8	This work
γ-Al ₂ O ₃ (hydrolysis method)	183	0.4	10	Sifontes <i>et al.</i> , 2014
γ-Al ₂ O ₃ (hydrothermal route)	248	0.30	4.3	Guzman <i>et al.</i> , 2005
γ-Al ₂ O ₃ (CONDEA commercial)	220	-	5.9	Del Angel <i>et al.</i> , 2005
γ-Al ₂ O ₃ (Template method)	328.9	0.81	12.0	Renuka <i>et al.</i> , 2016
γ-Al ₂ O ₃ (precipitation/digestión)	220	0.48	7.0	Potdar <i>et al.</i> , 2007
γ-Al ₂ O ₃ (dual-templating route)	320	0.60	7.0	Sandeep <i>et al.</i> , 2015
Mesoporuos Alumina (double hydrolysis method)	335.5	0.60	7.1	Benjing <i>et al.</i> , 2017
γ-Al ₂ O ₃ (sol–gel method)	370	0.52	6.0	Rudina <i>et al.</i> , 2012

γ -Al ₂ O ₃ (sol-gel hydrolysis method)	236	0.4	7.0	Wei <i>et al.</i> , 2016
γ -Al ₂ O ₃ commercial (J&K chemical)	331	-	5.8	Zongbo <i>et al.</i> , 2016
γ -Al ₂ O ₃ (Solid state Method)	220	0.49	6.1	Soodeh <i>et al.</i> , 2016
γ -Al ₂ O ₃ (coprecipitation method)	346.4	1.0	10.2	Zhang <i>et al.</i> , 2016
γ -Al ₂ O ₃ (Reverse precipitation method)	240	0.90	7.6	Wu <i>et al.</i> , 2012
γ -Al ₂ O ₃ (coprecipitation/sol-gel method)	264	0.30	5.1	Mohammed <i>et al.</i> , 2012

^a Specific surface area (m²/g)
^b Average pore volume (cm³/g)
^c Average pore diameter (nm)

The comparative values obtained and reported by different authors from several synthetic pathways are similar and in some cases higher. This textural behavior is typical of materials where condensation occurs mainly between particles. This is related to grain size. After the calcination there is a considerable increase in the values of specific areas caused by dehydroxylation leading to the formation of γ -Al₂O₃. It is seen that the synthesized material from an inexpensive aluminum precursor with impurity presence make larger nanofibers presenting very attractive textural properties obtained by a simple and economical synthesis method.

Fourier transforms infrared spectroscopy (FT-IR) analysis

The FTIR spectra for the different products obtained, is shown in Figure 6. The results of the synthesized materials are consistent with the reported in literature (Riad *et al.*, 2007). In the FT-IR patterns, the broad band at 3200–3700 cm⁻¹ its characteristic of -OH stretching vibration that is bonded to Al³⁺, and the band at 1638 cm⁻¹ corresponding to physisorbed water are observed. Meanwhile the peak at 1470 cm⁻¹ was also due to water deformation vibrations (Boumaza *et al.*, 2009). Band appearing at 1100 cm⁻¹ is typically for γ -alumina due to Al–O vibration mode (Xu *et al.*, 2017). The valley between 1000 cm⁻¹ and 435 cm⁻¹ confirms the γ -form, and the one at 873 cm⁻¹ is assigned to the bending vibrations of Al–O bond. Al–O–Al bond in the gamma phase of alumina generates the band at 670 cm⁻¹ (Cheng *et al.*, 2006). These results are consistent with the XRD analysis where the γ -phase was identified.

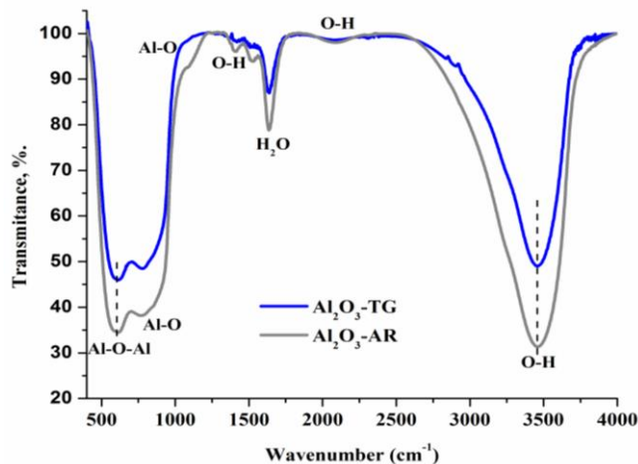


Figure 6. FT-IR spectra of Al_2O_3 -TG and Al_2O_3 -AR.

The peaks in the region of $500 - 750 \text{ cm}^{-1}$ are assigned to Al^{VI} , whereas the shoulder at 750 and the line at 890 cm^{-1} are assigned to Al^{IV} . Thus, $\gamma\text{-Al}_2\text{O}_3$ phase contains both tetrahedral and octahedral coordination (Renuka *et al.*, 2012). In this sense, it is possible to observe more intense vibrations and slightly more widened in the material Al_2O_3 -AR, reason why a greater content of the tetrahedral coordination assuming a greater acidity is attributed. Several Authors have observed that the presence of Mg decreases the weak Brønsted acidity and increases the basicity of the alumina (Zhang *et al.*, 2016). Then that product obtained of Al_2O_3 -TG presents lower content of tetrahedral coordination.

Effect of salt's purity

The results clearly show that the presence of Mg into the Al crystal lattice change the acid and basic properties of this oxide. It is interesting to verify that the values of the surface area (Table 3) change when comparing Al_2O_3 -TG with Al_2O_3 -AR. In recent years the use of different grades of aluminum salts have become important areas of research and development due to their importance as precursors for aluminum oxides. In some applications, such as advanced manufacturing alumina-based ceramics, and high-purity aluminium oxides compounds are required. Here tolerable concentrations of impurities are rather low and an analytical control became a limiting step of the technology (Koksal *et al.*, 2002). Other applications that required high-purity aluminum oxides are pharmaceuticals products, refractories, electronics, among others (Mishra, 2002). However, there are applications where the presence of impurities can benefits the application or reaction, as

mentioned earlier. This study determined that a small amount of impurity in precursor salt can modify the physical properties of alumina compared with conventional γ -alumina. In addition it, exhibit attractive properties for several applications as the process of adsorption and catalytic, in addition to a low cost.

Conclusion

The material of γ -Al₂O₃ nanofibrillar synthesized by hydrolysis-precipitation method presented physicochemical properties attract as possible catalytic support and adsorbent material. The results showed that 0.33 wt. % of Mg as impurity was sufficient to generate structural defects and decreased the acidity in the Al₂O₃-TG material. However, the presence of Mg extended the fibrillar chain of the alumina besides giving greater thermal stability and greater textural properties. Hence from precursors of aluminum with presence of impurities can be obtained attractive materials for several applications.

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