

## Surface Area, Crystal Morphology and Characterization of Transition Alumina Powders from a New Gibbsite Precursor

Antonio Carlos Vieira Coelho<sup>a\*</sup>, Helena de Souza Santos<sup>b</sup>, Pedro Kuniiko Kiyohara<sup>b</sup>,

Kelly Nanci Pinto Marcos<sup>a</sup>, Pêrsio de Souza Santos<sup>a</sup>

<sup>a</sup>Laboratório de Matérias-Primas Particuladas e Sólidos Não Metálicos, Departamento de Engenharia Metalúrgica e de Materiais, Escola Politécnica, Universidade de São Paulo – USP, Av. Prof. Luciano Gualberto, travessa 3, 380, 05508-970 São Paulo - SP, Brazil

<sup>b</sup>Laboratório de Microscopia Eletrônica – LME, Departamento de Física Geral, Instituto de Física – IF, Universidade de São Paulo – USP, CP 66318, 05315-970

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A new procedure was used to prepare a microcrystalline powder constituted by thin euhedral hexagonal gibbsite plates, 0.2 to 0.6  $\mu\text{m}$  in diameter and 32 nm thick. The powder, fired between 200 and 1000  $^{\circ}\text{C}$ , produced chi and kappa transition aluminas. Alpha-alumina is formed from 1000  $^{\circ}\text{C}$  and recrystallized up to 1500  $^{\circ}\text{C}$ . At 1000  $^{\circ}\text{C}$ , kappa- and alpha-alumina coexisted, but kappa-alumina could only be characterized by SAED. The details of the internal organization of the transition alumina pseudomorphs were clearly observable in TEM due to the great thinness of the I-gibbsite plates. The specific surface area varied from pristine I-gibbsite (24.9  $\text{m}^2\cdot\text{g}^{-1}$ ) to chi- and kappa transition aluminas (25.4  $\text{m}^2\cdot\text{g}^{-1}$ ) at 1000  $^{\circ}\text{C}$  to alpha-alumina (4.0  $\text{m}^2\cdot\text{g}^{-1}$ ) at 1500  $^{\circ}\text{C}$ . The maximum value of specific surface area is 347  $\text{m}^2\cdot\text{g}^{-1}$  in chi-alumina powder at 300  $^{\circ}\text{C}$ , a difference from Bayer gibbsite, in which the chi-alumina highest surface area is 370  $\text{m}^2\cdot\text{g}^{-1}$  at 400  $^{\circ}\text{C}$ .

**Keywords:** gibbsite, transition aluminas, active aluminas, alpha-alumina

### 1. Introduction

Synthetic gibbsite is the most important alumina chemical, either as an aluminum trihydroxide or as precursor for alumina powders for advanced and traditional ceramics, catalysts and catalyst carriers, adsorbents, aluminum, paper, rubber, plastics and other chemical process industries. A great industrial value of gibbsite is due to fact that it is the only aluminum hydroxide that, by thermal processing, is able to produce all transition aluminas (from the chi and gamma series) and alpha-alumina.

Usually, gibbsite, natural or synthetic crystals, are of micrometric size; their structure is triclinic, but common shapes are tabular with pseudo-hexagonal profiles and prisms of hexagonal basis<sup>1,2</sup>. By transmission electron microscopy (TEM), many of the prisms appear as rectangles if they stand sideways on the TEM support.

McBride et al.<sup>3</sup> presented TEM of hexagonal platy microcrystals of gibbsite prepared by Gastuche and Herbillon's procedure<sup>4</sup> of aging of gelatinous amorphous aluminum hydroxide precipitate in water, at pH 4.6, for one month at room temperature; the aluminum hydroxide was prepared by the reaction between aluminum chloride and sodium hydroxide solutions. Hsu<sup>5</sup>, reviewing the literature on aluminum hydroxides, reproduced TEM of hexagonal plates of microcrystals of gibbsite, from the aging of aluminum hydroxide, in water containing sodium and perchlorate ions, at room temperature<sup>6</sup>.

McZura et al.<sup>7</sup> and Misra<sup>8</sup> described, by TEM, very uniform hexagonal platy microcrystals, about 1  $\mu\text{m}$  in diameter, produced by a patented process from an "organic-free aluminate liquor" for use as coating pigment for paper<sup>9</sup>, in ALCOA Hydral Series. Sweegers et al.<sup>10</sup> reviewed the morphology of gibbsite crystals grown from pure and impure aqueous sodium aluminate solutions and Adamo et al.<sup>11</sup> in tartrate solutions.

Pseudo-hexagonal platy gibbsite microcrystals are normal components of Brazilian bauxites<sup>12</sup> and high alumina gibbsite clays<sup>13,14</sup>.

Brown et al.<sup>15</sup> were the first to demonstrate that the thermal decomposition of gibbsite crystals may follow dual transformations at normal pressure; the first leads to the chi-alumina series and the second to the boehmite (AlOOH) or gamma series. Both transformations are pseudomorphic and topotactic in relation to gibbsite axis<sup>16</sup>. Both series produce several alumina structures, the so-called transition aluminas and both end, at 1100  $^{\circ}\text{C}$ , in alpha-alumina. In consequence, the great value of gibbsite is because it is the only of all aluminum hydroxides that may produce all transition aluminas. This property makes synthetic gibbsite the most important alumina chemical, either as a crystalline aluminum hydroxide or as an alumina precursor. The so-called Bayer gibbsite is produced as an intermediate in Bayer process for aluminum production<sup>8</sup>. It is used as a precursor for production of alumina powders for several chemical processes. So, the thermal transformation of gibbsite in the two series was extensively studied, specially the development of high surface areas, pore structures and texture of the transition aluminas. Wefers and Bell<sup>17</sup>; Wefers and Misra<sup>18</sup> reviewed and summarized the most important results from these studies. Wefers<sup>19</sup> reviewed the industrial and laboratorial processes for producing crystalline gibbsite.

Bayer gibbsite particles, as in ALCOA C-30, when examined by scanning electron microscopy (SEM), are roughly spherical agglomerates of tabular and prismatic single crystals; the agglomerates diameter vary between 40 and 200  $\mu\text{m}$ ; each crystal, tabular or prismatic, has a diameter between 5 and 10  $\mu\text{m}$ <sup>7,8</sup>. The majority of the thermal decomposition studies of gibbsite has been conducted on Bayer crystal agglomerates.

\*e-mail: acvcoelh@usp.br

Wefers and Misra<sup>18</sup> presented TEM data on the texture of the crystals of the chi-alumina series; the micrographs are from chi- and kappa-alumina pseudomorphs from prismatic or acicular single crystals of Bayer gibbsite, observed sideways: the original gibbsite crystal appears as rectangular and the transformation into chi-alumina is described as an exfoliation of lamellae. Suzuki<sup>20</sup> and later Wefers and Bell<sup>17</sup>, published micrographs of thin hexagonal platy chi-alumina pseudomorphs observed parallel to the c-axis: the texture of the hexagonal plate is described as a “network of submicroscopic cracks and devices in the heated crystal”, this texture, later, has been described as “sponge-like” or as being constituted by “jigsaw” particles.

Pandolfelli et al.<sup>21</sup> described the alterations by calcination of the physical, morphological and rheological properties of ALCOA C-30 Bayer gibbsite.

The aluminum hydroxides which crystallizes in water in which no substance is dissolved is the trihydroxide bayerite; if Na<sup>+</sup> or K<sup>+</sup> ions are present, gibbsite is formed<sup>22</sup>. Amalgamated aluminum reacts with water producing amorphous aluminum hydroxide which crystallizes in bayerite; if linear alcohols are added several aluminum hydroxides may crystallize, depending on the alcohol<sup>23</sup>.

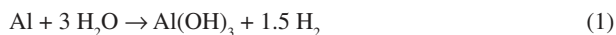
Recently, Souza Santos et al.<sup>24</sup> studied the reaction between aluminum powder and water, using iodine as an “activator”<sup>25</sup> in place of mercury. Euhedral, very thin, regular hexagonal plates were formed in the narrow temperature range of 50-60 °C, having 0.2 to 0.6 μm diameter by TEM; a few crystals may have diameters in the nanometric range (6-7 nm). The smaller size and thinness of these gibbsite hexagonal platelets could allow a detailed study by TEM of the variation with of the pseudomorphs texture increasing temperature in the series: gibbsite → chi-alumina → kappa-alumina → alpha-alumina.

The purpose of this paper is to characterize the differences from Bayer gibbsite crystals which are observed in the specific surface area, internal structure and morphology of the microcrystals of the transition aluminas of the chi-alumina series formed from the gibbsite crystals by the new method after heated between 200 and 1500 °C, aiming to obtain aluminas with new or improved properties.

## 2. Materials and Methods

### 2.1. Gibbsite preparation by the “iodine method”<sup>24</sup>

ALCOA 123 uncoated Al-powder was used (100% minus n° 325 ABNT sieve; Al-99.7%; Si-O-0.15%; Fe-0.17%). In a round bottom two liters three necked Pyrex flask, in the following sequence were placed: 500 mL of water; 27.0 g of Al powder and 50 mL of an iodine solution (5.0 g of iodine crystals dissolved in 50 mL 95% ethanol). The system was stirred one hour for homogenization and heated to 60 ± 1 °C by thermostat controlled heating mantle; a water cooled condenser was fitted to the flask. It took a minimum of two weeks for the aluminum react completely. The condenser was taken out in the last four hours to assure that if any remaining iodine was eliminated. The white dispersion presented strong dityndallism by stirring, an indication of anisodiametric shape of the particles. The white dispersion is centrifuged in polyethylene tubes in a Servall centrifuge at 3500 rpm for 30 minutes. The white precipitate was washed four times with its original volume of 95% ethanol. The white precipitate of aluminum hydroxide gibbsite (called “I-gibbsite” in this paper) was dried at 60 °C for 24 hours and the white powder was ready for characterization and other tests; the yield in Al(OH)<sub>3</sub> was 86-87%, based in the equation:



### 2.2. Thermal treatment

Three to four grams of the unpressed dry white powder were program fired on a platinum foil in an electric furnace EDG, São

Carlos, SP, for 3 hours at the maximum temperature, from 200 °C up to 1500 °C, with 100 °C intervals; natural cooling was used. After cooling, the fired powder was characterized by X ray diffraction, transmission electron microscopy, selected area electron diffraction (SAED), elemental microanalysis and for measurement of the specific surface area.

### 2.3. Methods

The Differential Thermal Analysis (DTA) was conducted in an equipment by B.P. Engenharia, Campinas, SP, Brazil, up to 1100 °C in open air.

The X ray diffraction (XRD) was conducted in a Philips X-pert MPD model equipment, operating at 40 kV and 40 mA, between 1° (2θ) and 90° (2θ). All lines of the XDR curve of the aluminum hydroxide powder prepared at 60 °C correspond to gibbsite lines listed in ICDD file n° 7-324 and ref.<sup>18</sup>; the more intense line is the 4.85 Å gibbsite reflection. No other Al-hydroxide is detected. The transition aluminas were characterized from ICDD files and ref.<sup>18</sup>.

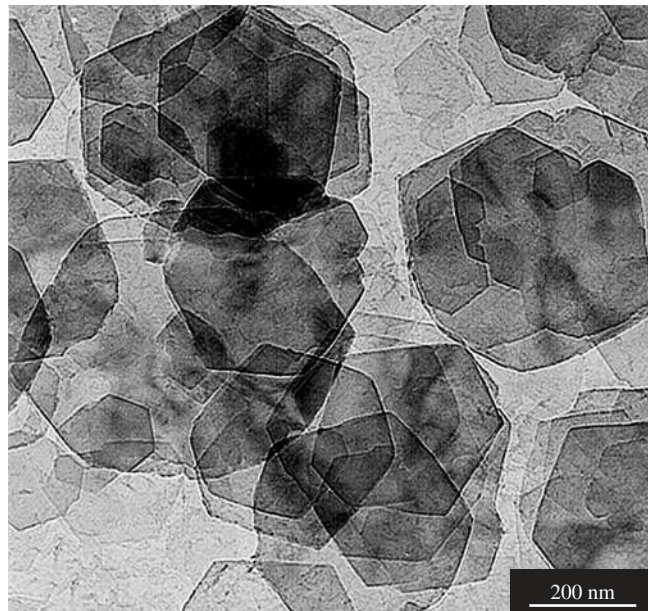
The dried powder was dispersed in distilled water and prepared in the conventional manner for transmission electron microscopy (TEM). A Philips CM-200 equipment, operating at 200 kV was used. The same microscope was used as a diffraction camera to characterize the crystalline alumina phases structures by SAED and for elemental microanalysis by electron diffraction spectroscopy (EDS).

The specific surface area (in m<sup>2</sup>.g<sup>-1</sup>) of the fired powders was measured by the BET-nitrogen adsorption method using a Micromeritics ASAP 2010 model. The adsorption data were taken using 99.999% pure nitrogen gas (Air Liquid), at the liquid nitrogen temperature (77 K). The experimental data were taken in the 0.01-0.30 range of partial pressures.

## 3. Results and Discussion

### 3.1. Gibbsite characterization

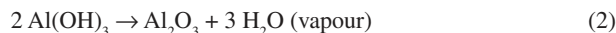
Figure 1 is a transmission electron micrograph of some well-formed gibbsite platy crystals. Their diameters are in the 0.2 μm to 0.6 μm interval. They have hexagonal profiles, various thickness, the



**Figure 1.** Transmission electron micrograph of some well-formed I-gibbsite platy crystals (LME-IF-USP).

majority being very thin. The SAED confirms the gibbsite structure characterized by the XRD curve of Figure 2. EDS elemental microanalysis indicates the absence of iodine in the microcrystals that were examined. The hexagonal platy crystals are similar to those used for paper coating<sup>9</sup> and are completely different from the Bayer gibbsite agglomerates<sup>7,8</sup>.

The DTA curve of Figure 3a shows an intense endothermic peak starting at 250 °C and with maximum intensity at 300-320 °C. No endothermic peak is observed at 525-550 °C, an indication of absence of boehmite, either as an impurity or formed by partial dehydroxilation. The endothermic peak corresponds to the complete dehydroxilation of the gibbsite crystal, according to the equation:



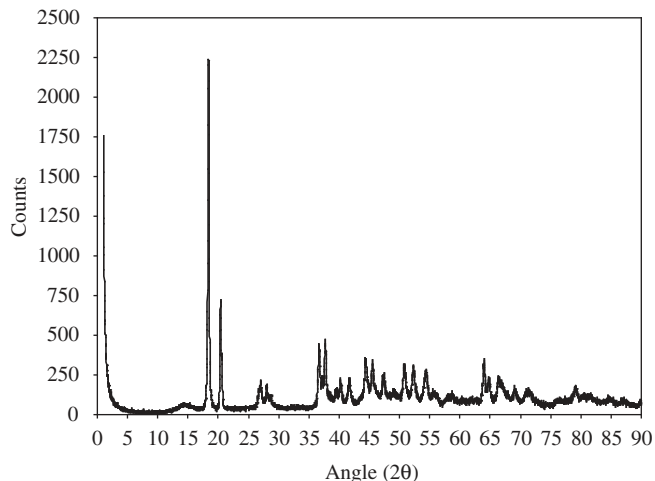
That reaction occurs when the gibbsite crystals are fine grained and there is free escape of the water vapour. In these conditions, the transition chi-alumina series is formed, starting at 273 °C, according to Wefers and Misra<sup>18</sup>.

### 3.2. Aluminas characterization by XRD

Figure 4, adapted from ref.<sup>18</sup>, shows the Bayer gibbsite to the chi-alumina series transformation by increasing temperatures. The following temperature differences can be observed in the transition temperatures of the I-gibbsite microcrystals prepared by iodine activation of aluminum powders from Bayer gibbsite: a) chi-alumina starts to form at 200 °C (273 °C in ref.<sup>18</sup>); b) chi-alumina remains up to 900 °C, while 753 °C is the upper limit, according to ref.<sup>18</sup>; c) kappa-alumina exists in a shorter temperature interval (900 to 1000 °C), instead of 753 to 1020 °C<sup>18</sup>; and d) alpha-alumina crystallization starts at 1000 °C, because its three strongest lines are observed (instead 1020 °C in ref.<sup>18</sup>); no kappa-alumina line is observed in the 1000 °C XRD curve; sintering, recrystallization and growth of the alpha-alumina microcrystals continues up to 1500 °C. Figure 5 presents some XRD curves from I-gibbsite fired between 200 and 1500 °C.

### 3.3. Specific surface area

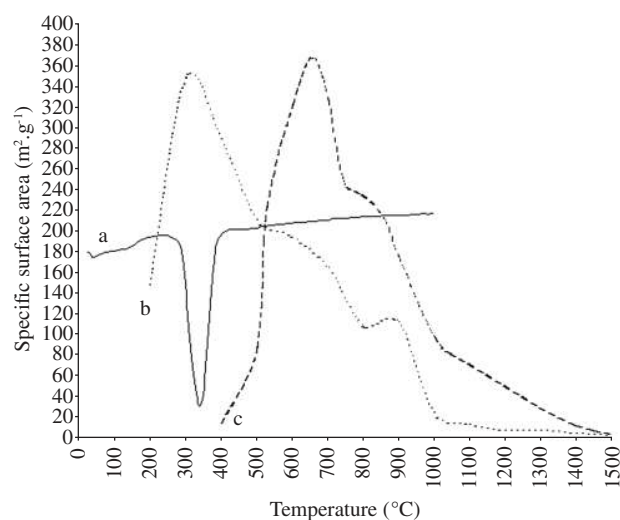
The value of the specific surface area of the pristine I-gibbsite crystals is 29 m<sup>2</sup>.g<sup>-1</sup> (Figure 3), which is bigger than from Hydral aluminum hydroxide, having the same crystal shape (the values are 6-8 m<sup>2</sup>.g<sup>-1</sup> and 12 m<sup>2</sup>.g<sup>-1</sup> for the two types<sup>9</sup>). The variation of the specific surface area of the I-gibbsite powder with firing temperatures



**Figure 2.** XRD curve of well-formed gibbsite platy crystals showed in Figure 1, confirming the gibbsite structure.

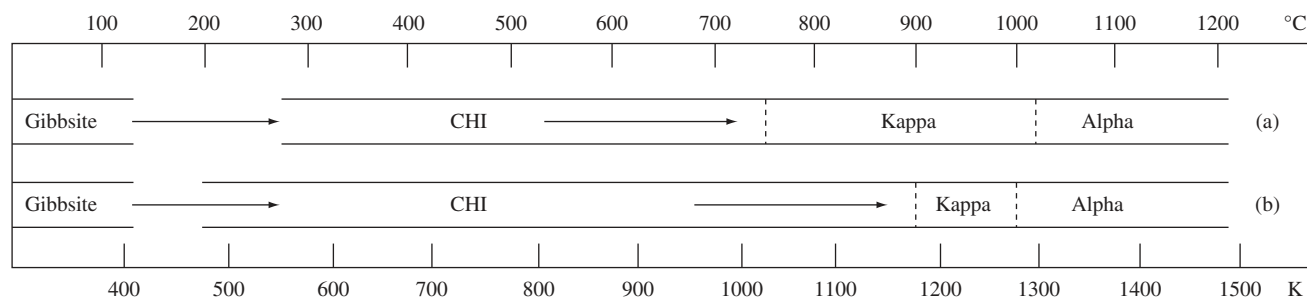
between 200 and 1500 °C is shown in Figure 3b in comparison with Bayer gibbsite data from literature (Figure 3c)<sup>18</sup>. It can be observed that the maximum value observed for the fired powders of I-gibbsite is obtained at 300 °C, the same temperature of the endothermic dehydroxilation peak of the gibbsite to chi-alumina transformation shown in the DTA curve (Figure 3a).

The maximum value is 347 m<sup>2</sup>.g<sup>-1</sup> from a sample fired at 300 °C, which is close to the maximum value from of the gibbsite form literature - Figure 3c, 370 m<sup>2</sup>.g<sup>-1</sup> at 650 °C. Therefore, either at 300 °C (I-gibbsite) or at 650 °C (Bayer gibbsite), both have chi-alumina structure; so, the most probable cause of the difference for achieving the temperature for producing maximum surface area is the smaller internal texture of the chi-alumina crystals from I-gibbsite. According to Oberlander<sup>26</sup>, activated aluminas having specific surface areas higher than 100 m<sup>2</sup>.g<sup>-1</sup>, are considered "high surface area aluminas". So, high surface area alumina can be produced firing I-gibbsite in the 200-900 °C range.

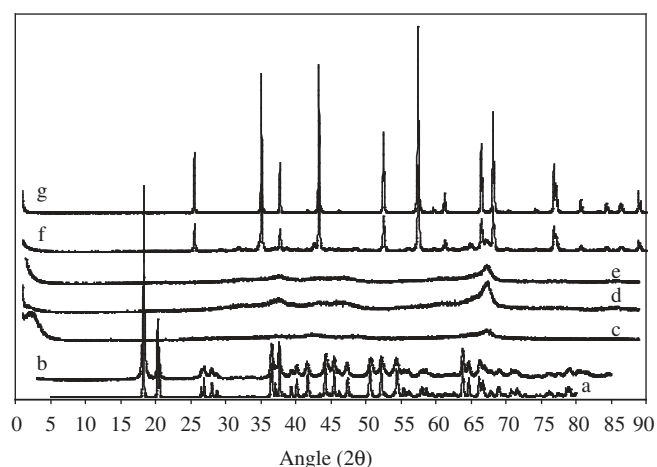


Temperature (°C)	BET area (m <sup>2</sup> .g <sup>-1</sup> )
25	29.4
200	148
300	347
400	289
500	211
600	195
700	165
800	110
900	113
1000	25.4
1100	14.4
1200	8.2
1300	8.3
1400	5.9
1500	4.0

**Figure 3.** a) DTA curve of I-gibbsite; b) Curve representing specific surface area data of I-gibbsite after thermal activation; c) Curve representing specific surface area data of a Bayer gibbsite after thermal activation; and d) Specific surface area (BET) data of I-gibbsite after thermal activation.



**Figure 4.** Thermal transformation series: a) Bayer gibbsite: gibbsite  $\text{Al}(\text{OH})_3 \rightarrow$  chi-alumina  $\rightarrow$  kappa-alumina  $\rightarrow$  alpha-alumina (adapted from ref. <sup>18</sup>); and b) I-gibbsite: same transformation series, but in different temperatures.



**Figure 5.** XRD curves from I-gibbsite fired between 300 and 1500 °C: a) Bayer gibbsite; b) pristine I-gibbsite; c) I-gibbsite fired at 300 °C; d) 800 °C; e) 900 °C; f) 1000 °C; and g) 1500 °C.

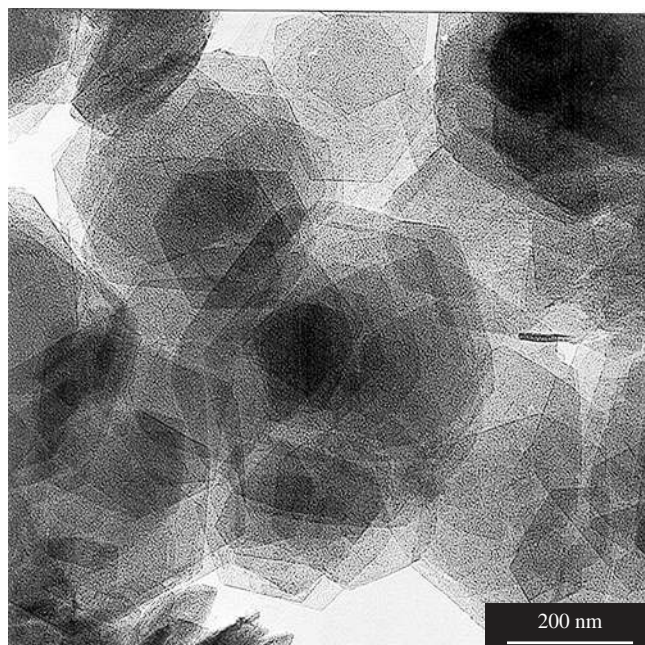
### 3.4. Electron microscopy and SAED

The thickness of I-gibbsite crystal can be calculated, assuming the microcrystal as a prism of regular hexagonal basis; the gibbsite density is  $2.42 \text{ g/cm}^3$ ,  $0.5 \mu\text{m}$  diameter and specific surface area of  $29.4 \text{ m}^2 \cdot \text{g}^{-1}$  (Figure 3d): the calculated thickness is 32 nm, which corresponds to 66 gibbsite (0.485 nm) layers.

The fired samples, after being characterized by XRD, were examined by TEM to compare the internal morphology changes of the aluminas pseudomorphs with the pristine I-gibbsite hexagonal platy microcrystals of Figure 1. SAED was used to characterize the crystalline structure of single microscopical crystals.

Very early, at 200 °C, but better at 300 °C, SAED characterized gibbsite in many crystals, but several chi-alumina hexagonal pseudomorphs were also characterized; so, several gibbsite microcrystals had dehydroxilated into chi-alumina and the original shapes of the crystals was maintained, forming the chi-alumina pseudomorphs. Figure 6 shows 300 °C chi-alumina pseudomorphs with an internal fine granularity; the majority of the grains are chi-alumina crystals; the surface of the plates is not smooth anymore; that stage constitute the initial nucleation stage of the chi-alumina series characterized by the appearance of the granularity.

At 300 °C, but better at 500/600 °C, inside the thinnest hexagonal plates (pseudomorphs), an arrangement can be observed of the granules into an hexagonal patterns, giving the appearance of a mosaic structure to the pseudomorphs. The arrangement is still visible at 700 °C, but it is not easily observable in thicker plates or at higher temperatures due to the increased particle size. Each granule is a microcrystal of chi-



**Figure 6.** TEM of chi- $\text{Al}_2\text{O}_3$  from I-gibbsite platy crystals fired at 300 °C (LME-IF-USP).

alumina. It can be concluded that the maximum value of the specific surface area occurs at 300 °C, when the chi-alumina granules inside the pseudomorphs (Figure 6) have the smallest size and the gibbsite to chi-alumina transition or reaction is complete; increasing temperature, increases the sizes of the chi-alumina nuclei and decreases the specific surface area of the transition alumina.

The earlier transmission electron micrographs of chi-alumina pseudomorphs from fired I-gibbsite hexagonal plates showed only an internal porosity and no regular arrangement of the grains; even the more recent papers by Levin and Brandon<sup>27</sup> and by Souza Santos et al.<sup>28</sup> do not show that hexagonal arrangement of the chi-alumina microcrystals shown in Figure 6.

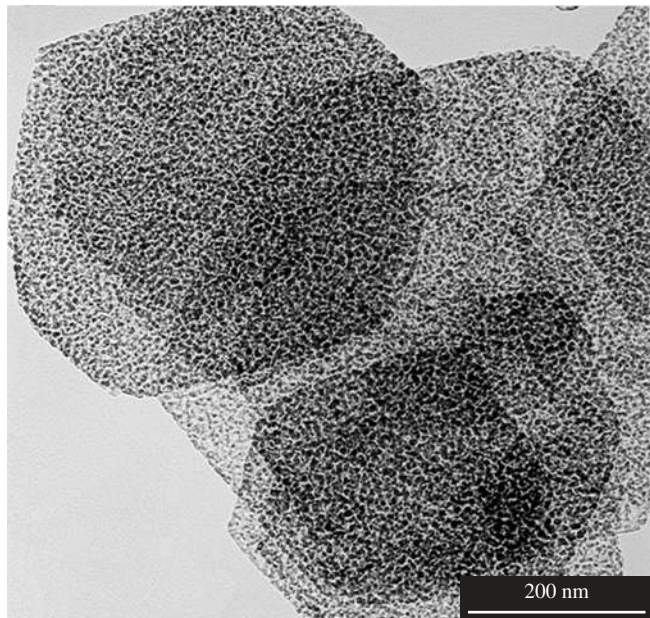
Figure 7 is a TEM of chi-alumina pseudomorphs fired at 700 °C, where the increased granularity in comparison with Figure 6 can be observed. It can clearly be observed that the coarse grains have lost the round shape shown in Figure 6 and acquired a sharp platy one; from this crystal growth results sharper and more intense SAE and X ray reflections.

At 800 °C, it is possible to observe that many platy granules inside the pseudomorphs have acquired hexagonal profiles; XRD characterizes only chi-alumina and SAED confirms that structure for the hexagonal grains. Increasing the temperature, XRD and SAED still

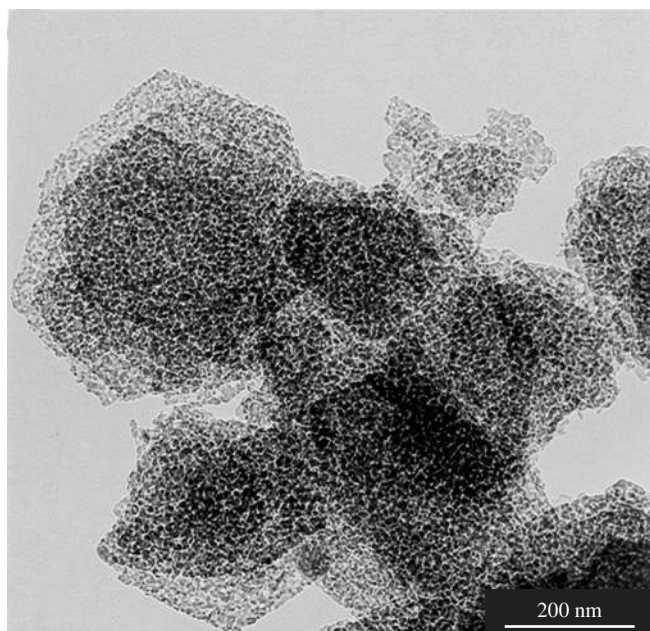
shows the chi-alumina structure. The granularity of the pseudomorphs becomes coarser with the increased temperatures.

But at 900 °C, all the granules have platy hexagonal shapes and XRD and SAED confirm kappa-alumina structure for the grains. The pseudomorphs profiles have not changed. Figure 8 is a TEM of kappa-alumina pseudomorphs fired at 900 °C.

At 1000 °C, only alpha and no kappa-alumina lines are detected, specially the characteristic 0.257 nm kappa-alumina reflection; however, their mixture is easily detected by SAED. The coexistence of kappa and alpha-alumina particles in mixture in the 1000 °C fired



**Figure 7.** TEM of chi- $\text{Al}_2\text{O}_3$  from I-gibbsite platy crystals fired at 700 °C (LME-IF-USP).



**Figure 8.** MET of kappa- $\text{Al}_2\text{O}_3$  from I-gibbsite platy crystals fired at 900 °C (LME-IF-USP).

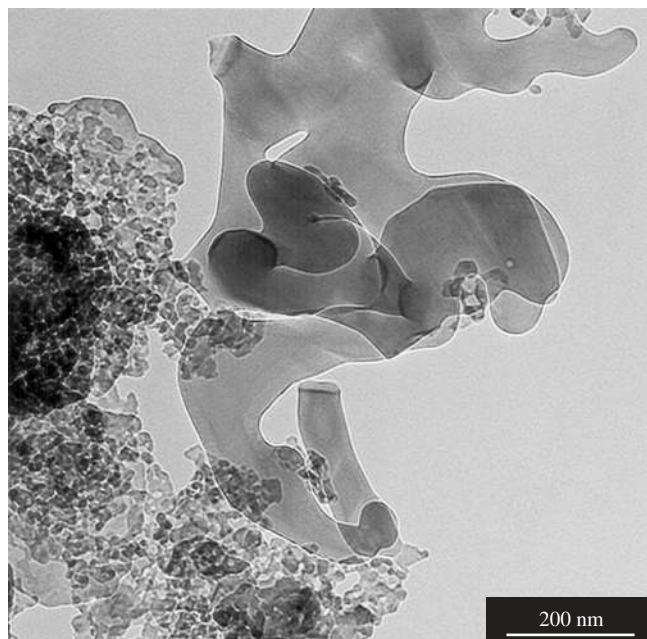
sample is clearly shown in transmission electron micrographs, as shown in Figure 9: a) the same pseudomorphs of Figure 9, constituted by small hexagonal platelets of kappa-alumina; b) electron dense lamellar crystals, many superposed, having smooth round irregular profiles, exhibiting Bragg contours and Moire fringes: they are sintered alpha-alumina crystals, as characterized by SAED. The kappa-alumina pseudomorphs are destroyed in the alpha-alumina crystallization. At 1100 °C, only electronically coalesced round profile alpha-alumina crystals are observed.

Lippens and De Boer<sup>29</sup> called attention to the work of De Boer et al.<sup>30</sup>: “gibbsite crystals heated at lower temperature (350 °C) formed slit-shaped pores which are separated by platelike particles parallel to each other and perpendicular to the c-axis of the original gibbsite crystal; at higher temperatures (not stated), however, rodlike particles are formed which are perpendicular to the c-axis of the original gibbsite crystal and parallel to each other. During sintering, even up to 1200 °C, the rodlike structure of the  $\text{Al}_2\text{O}_3$  is left undisturbed, in spite of the fact of the chi- $\text{Al}_2\text{O}_3$  is converted, via a number of intermediate stages, into alpha- $\text{Al}_2\text{O}_3$ ”.

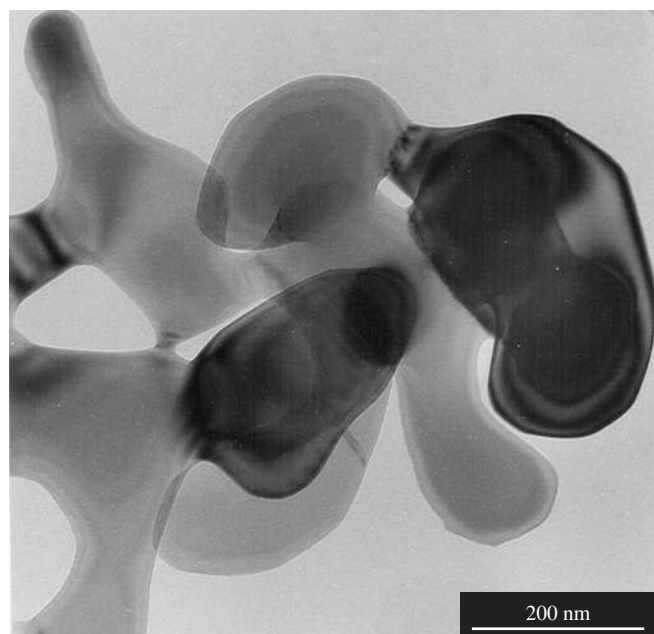
Wefers and Misra<sup>18</sup> (p. 56-59) presented TEM micrographs of chi, kappa and alpha-alumina from heated acicular gibbsite crystal: parallel the slit-shaped pores and platelike particles and perpendicular to the c-axis are observed, but no rodlike particle is mentioned from the micrographs.

In this paper, the original gibbsite crystals were observed perpendicular to the c-axis; so, the rodlike particles would appear inside the hexagon pseudomorph of chi-alumina and aligned radially. No rodlike particle was observed in the gibbsite crystals fired from 200 to 1100 °C: the chi and kappa-alumina microcrystals always appeared as hexagonal plates.

Up to 1500 °C, the alpha-alumina crystals present sintering, pore formation, growth and recrystallization into round particles and plates, many with some 120° angle; no hexagonal plate so as observed. Figure 10 is a TEM micrograph of alpha-alumina sample fired at 1500 °C.



**Figure 9.** MET of kappa- $\text{Al}_2\text{O}_3$  + alpha- $\text{Al}_2\text{O}_3$  from I-gibbsite platy crystals fired at 1000 °C (LME-IF-USP).



**Figure 10.** MET of  $\alpha\text{-Al}_2\text{O}_3$  from I-gibbsite platy crystals fired at 1500 °C (LME-IF-USP).

#### 4. Conclusions

Gibbsite microcrystals, prepared by a new procedure using the reaction between aluminum powder and water, having iodine as activator, produce powders constituted by thin pseudo-hexagonal plates, 0.2 to 0.6  $\mu\text{m}$  in diameter, with specific surface areas of 24-25  $\text{m}^2\cdot\text{g}^{-1}$ . The thermal transformation sequence of the I-gibbsite powder presents some differences from literature data on Bayer gibbsite; these differences are attributed to the small diameter and thinness of the I-gibbsite platy microcrystals. The I-gibbsite powder, fired between 200 and 1500 °C, produces the chi-alumina series up to alpha-alumina (corundum). The maximum value of 347  $\text{m}^2\cdot\text{g}^{-1}$  of specific surface area, is obtained at 300 °C; the powder has chi-alumina structure; 300 °C is the temperature of the endothermic dehydroxilation peak which indicates the end of the reaction I-gibbsite  $\rightarrow$  chi-alumina + water (vapour). The new observed stability temperature ranges of the alumina phases are: a) I-gibbsite (up to 200 °C); b) chi-alumina (200 to 900 °C); c) kappa-alumina (900 to 1000 °C); d) kappa-alumina + alpha-alumina, (at 1000 °C); and e) alpha-alumina (1000 to 1500 °C). Therefore, it is possible with this gibbsite to obtain chi-alumina until 900 °C, while with Bayer gibbsite the maximum temperature of stability is 673 °C.

The selected area diffraction has shown more sensitivity than the XRD powder method for characterizing kappa-alumina, not detected by the later method at 1000 °C due to the very small particle size of the microcrystals.

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