Gallium oxide hydroxide (GaOOH$\cdot$H$_2$O) single crystals were synthesized in aqueous solutions by using two different precipitation techniques: homogeneous decomposition of urea and forced hydrolysis in pure water. Precipitation of crystals started at exactly the same pH value (i.e., 2.05 at 85°C) in both cases. The morphology of crystals turned out to be quite different (zeppelin-like with urea, rodlike without urea) in each of the above methods.Calcination of these gallium oxide hydroxide crystals in air at temperatures $\approx 500^\circ$C transformed them into Ga$_2$O$_3$. Characterization of the samples was performed by X-ray diffractometry, scanning electron microscopy, thermogravimetry/differential thermal analysis, Fourier transform infrared spectroscopy, and ICP, carbon, and nitrogen analyses.

I. Introduction

Gallium oxide hydroxide (GaOOH$\cdot$H$_2$O) single crystals were synthesized in aqueous solutions by using two different precipitation techniques: homogeneous decomposition of urea and forced hydrolysis in pure water. Precipitation of crystals started at exactly the same pH value (i.e., 2.05 at 85°C) in both cases. The morphology of crystals turned out to be quite different (zeppelin-like with urea, rodlike without urea) in each of the above methods. Calcination of these gallium oxide hydroxide crystals in air at temperatures $\approx 500^\circ$C transformed them into Ga$_2$O$_3$. Characterization of the samples was performed by X-ray diffractometry, scanning electron microscopy, thermogravimetry/differential thermal analysis, Fourier transform infrared spectroscopy, and ICP, carbon, and nitrogen analyses.

The high-temperature structure$^{4,5}$ of Ga$_2$O$_3$ (β form) is monoclinic (space group $\text{C}2/m$), with the lattice parameters $a = 12.214$, $b = 3.037$, $c = 5.798$ Å, $\beta = 103.83^{\circ}$. β-Ga$_2$O$_3$ is only one of five well-known forms$^{7-5}$ of gallium oxide: α-, γ-, δ-, and ε-Ga$_2$O$_3$, and all of these polymorphs are converted to β-Ga$_2$O$_3$ at $T > 870^\circ$C.$^{10}$ Growth of the single crystals$^{11}$ of β-Ga$_2$O$_3$ has been studied by Okada et al.$^{12}$ using the tin-flux method, and Ga$_2$O$_3$ nanowires were prepared by Zhang et al.$^{13}$ and Choi et al.$^{14}$

Manufacture of n-type semiconducting Ga$_2$O$_3$ thin films on electrically insulating substrates has been studied by Meixner et al.$^{15}$ which are considered and developed for the detection of reducing gases. It was also reported that different gas sensitivities can be tailored by the appropriate setting of the operating temperature, and at temperatures in excess of 900°C, the sensors may even be operated as O$_2$ sensors.$^{16-19}$ On the other hand, Haneda et al. reported$^{20,21}$ that the Ga$_2$O$_3$–Al$_2$O$_3$ ceramic catalysts prepared by using the sol–gel method showed superior activity (as compared with those of either pure Al$_2$O$_3$ or Ga$_2$O$_3$) for the selective reduction of NO with propane in the presence of H$_2$O and SO$_2$. It should be noted here that during the preparation of ceramics, Haneda et al.$^{20,21}$ mentioned that the pure Ga$_2$O$_3$ catalysts of their study were prepared “from gallium hydroxide obtained by adding urea to an aqueous solution of gallium(III) nitrate and stirring at 90°C for 10 h, followed by calcination of the recovered precipitates at 600°C for 5 h in an air atmosphere.”

Ga$_2$O$_3$ has recently been used in the synthesis of solid electrolytes of superior (as compared with Y-stabilized zirconia) ionic conductivity, i.e., La$_{0.8}$Sr$_{0.2}$Ga$_{0.8}$Mg$_{0.2}$O$_{2.8}$ (LSGM)$^{22,23}$ also beyond its significance for the semiconductor, optoelectronic, and catalysis technologies. We have encountered the incidental formation of gallium oxide hydroxide (Ga(OH)$_3$) in aqueous solutions throughout the course of our recent studies, which were mainly focused on the wet-chemical synthesis of powders of doped LaGaO$_3$ fuel cell ceramics.$^{24,25}$

The first experimental study on the transformation of gallium oxide polymorphs and hydroxides was performed by Laubengayer et al.$^{26}$ in 1939 over the temperature range of 110° to 1000°C. These authors synthesized α-Ga(OH)$_3$ from a gel obtained through the hydrolysis of gallium nitrate or gallium chloride, after calcination at temperatures between 110° and 300°C. It was claimed$^{27}$ that, in this temperature range, Ga(OH)$_3$ transforms in a sluggish manner to α-Ga(OH)$_3$ (β form) after exposure a 0.114 M aqueous solution of GaCl$_3$ to ultrasonic irradiation for 6 h, by inserting the titanium horn of an ultrasonic finger into this solution. The same authors repeated the above procedure for InCl$_3$ solutions (0.68M), but this time$^{27}$ to form rodlike In(OH)$_3$ particles. The researchers concluded in both of these studies$^{36,37}$ that the sonochemical reaction provided by the inserted tip of the ultrasonic finger led to the sonohydrolysis of the gallium chloride or indium chloride solutions.

Matijevic et al.$^{38}$ prepared uniform colloidal indium hydroxide (i.e., In(OH)$_3$) particles by using either the technique of “homogeneous precipitation with urea”$^{29-30}$ or forced hydrolysis of aqueous solutions at elevated temperatures in the absence of any additives. The authors have systematically varied the indium salt

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N. J. Dudney—contributing editor

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concentration, the addition of sodium sulfate and urea, the solution pH, the temperature, and the aging time to see the effect of these parameters on the composition, size, and shape of the resulting particles. Matijevic et al.51 have also studied the preparation of In(OH)₃ particles by peptization of precipitates obtained in NaOH–InCl₃ solutions of varying concentrations.

In this paper, we used the precipitation methods of (a) the homogeneous decomposition of urea,39–50,52–54 and (b) forced hydrolysis (without urea, at 90°C)38 of gallium nitrate solutions to form the GaO(OH) crystals. The product powders are fully characterized (as identified via, for example, phases present, precipitate morphology, infrared (IR) behavior, and thermogravimetric and chemical analyses).51

II. Experimental Procedure

Ga(NO₃)₃·0.4H₂O (99.999% pure, Sigma-Aldrich Chemie GmbH, Steinheim, Germany) was used as the only gallium source throughout this study. A 0.3704M (Ga³⁺) stock solution was first prepared by dissolving an appropriate amount of gallium nitrate in deionized water. Urea used was also reagent-grade (>99% pure, Merck, Darmstadt, Germany). Synthesis experiments were performed by two different precipitation techniques.

(a) Homogeneous Decomposition of Urea: A 6.7 mL aliquot (i.e., 0.0025 mol of Ga³⁺) of the gallium nitrate stock solution was placed into a 150 mL capacity glass beaker, followed by the addition of 90 mL of deionized water. The solution was mixed at room temperature for 15 min, and 1.756 g of urea (i.e., 0.0292 mol) was then dissolved in it. The resultant clear solution was again mixed at room temperature for another 15 min, followed by heating on a hot plate to 90°C in about 45 min. The beaker was tightly covered with Parafilm®, before heating, to avoid excessive evaporation. The solution was finally aged at a constant temperature of 90(±1)°C for 95 min. The molar ratio of urea to Ga³⁺ was kept constant at a value of 11.68 throughout this study. The above procedure was also repeated by placing the same solution (containing urea) into a 100 mL capacity screw-capped glass bottle at room temperature, followed by inserting it into a microprocessor-controlled oven for heating and aging (95 min), which was preheated to 90°C, to produce the same powders.

(b) Forced Hydrolysis: A 6.7 mL aliquot (i.e., 0.0025 mol of Ga³⁺) of the gallium nitrate stock solution was placed into a 150 mL capacity glass beaker, followed by the addition of 90 mL of deionized water. This solution was mixed at room temperature for 15 min, followed by heating on an hot plate to 90°C in about 45 min. The beaker was covered with Parafilm® before the start of heating to reduce evaporation, and the solution was aged for 95 min at a constant temperature of 90(±1)°C. A portion of the same solution was also forced to undergo hydrolysis in a 100 mL capacity screw-capped glass bottle, placed into an oven at 90°C, as described previously.

The formed precipitates (in each of the above-described procedures) after 95 min of aging at a constant temperature of 90°C, were finally separated from their mother liquors by centrifugal filtration (at 8000 rpm), followed by washing four times with 2-propanol (>99% pure, Merck). Washed precipitates were dried in an oven at 90°C, overnight. Dried powders were first finely ground by hand using an agate mortar and pestle, and finally calcined isothermally (as loose powders in clean alumina boats) in a stagnant-air-atmosphere box furnace over a temperature range of 250° to 1200°C. Calcined powders were recovered from the Al₂O₃ boats by gently tapping them onto a clean piece of paper. Each calcination bath of powders was heated to the specified temperature at a rate of 5°C/min, annealed at this temperature for 6 h, and then furnace-cooled to room temperature.

The phase constitution of the powders was analyzed, as a function of temperature (on isothermally calcined samples), using a powder X-ray diffractometer (Model D-5000, Siemens GmbH, Karlsruhe, Germany) with monochromated CuKα radiation (40 kV, 30 mA). For determination of the lattice parameters, a scan rate of 0.9° 2θ/min (over the range of 10–80° 2θ) was used; a silicon external standard, along with the Appleman Least Squares Refinement Program,55 was used to correct for systematic errors.

The pyrolysis of accurately weighed 224.8-mg portions of ground powders was monitored via simultaneous thermogravimetric and differential thermal analysis (TG/DTA) (Model STA501, Bähr GmbH, Bremen, Germany) in an air atmosphere at a heating rate of 5°C/min up to 1040°C. Fourier transform infrared (FTIR) spectroscopy analyses of the samples were performed (Model IFS 66, Bruker GmbH, Karlsruhe, Germany) after they were mixed (1 wt%) with dry KBr to form pellets that were used in these analyses. The residual carbon and nitrogen contents of the uncalcined and calcined powders were determined using a combustion-IR absorption method (Model CS-800, Eltra GmbH, Neuss, Germany). ICP-AES analyses were performed to test the contamination of Ga(OH) or Ga₂O₃ powders by aluminum, after calcination of the samples in alumina boats at elevated temperatures (Model JY-70 Plus, Jobin Yvon S.A., Longjumeau, France). The powder morphologies were monitored using both field-emission scanning electron microscopy (FESEM) (Model DSM 982-Gemini, Zeiss GmbH, Oberkochen, Germany) and scanning electron microscopy (SEM) (Model Stereoscan-200, Cambridge Instruments, Ltd., Cambridge, U.K.).

III. Results and Discussion

(1) Homogeneous Decomposition of Urea

It is known39–50,52–54 that the formation of a solid precursor in aqueous precipitation experiments, which relies on the homogeneous decomposition of urea, is governed by the controlled generation of hydroxide and carbonate ions. The kinetics of the decomposition of urea in water appears to depend on the urea concentration, as well as other important variables, such as temperature and the presence and concentration of cations. The present study, however, excludes the effects of the above-mentioned variables on precipitate morphology and composition, since the molar ratio of urea to Ga³⁺, as well as the solution temperature, has not been varied.

The decomposition of urea and the progress of gallium oxide hydroxide precipitation was followed as a function of aging time and pH, and a typical plot of pH versus reaction time is given in Fig. 1. The trace of pH variation with time for 0.026M Ga³⁺ solutions (labeled “w/o urea,” i.e., the “forced hydrolysis” experiments) is also included in Fig. 1 for the purpose of direct comparison. The solution temperature has been raised from 23°C to 90°C in about 45 min, and the insert in Fig. 1 displays the typical variation of temperature with time in both kinds of gallium oxide hydroxide precipitation experiments.

It is interesting to note that precipitation started (observed by the onset of a slight turbidity in the solutions) when the temperature reached 85°C, regardless of the presence of urea in the solutions. The initial pH value of the 0.026M Ga³⁺ solutions (with or without urea) was in the range of 2.75 to 2.80 at 23°C, and the precipitation started in both kinds of solutions when the pH dropped down exactly to 2.05. This means that urea had no effect at all in “starting” the precipitation in gallium nitrate solutions of this study. On the other hand, the rise in pH (when t > 80 min) observed in urea-containing gallium nitrate solutions, as seen in Fig. 1, is characteristic of the decomposition of urea, and it corresponds to the start of dissociation of NH₄OH (which is an initial byproduct of urea decomposition together with H₂CO₃) into NH³⁺ and OH⁻. The detailed sequence of chemical reactions describing the decomposition of urea in aqueous media has previously been documented.47,56

The extremely small amounts of precipitates formed just at the precipitation-start point (i.e., pH 2.05, T = 85°C) have been immediately recovered from the urea-containing solutions by quenching the beaker of that solution in an ice-water bath (0°C), followed by centrifugal filtration at 10000 rpm, and 2-propanol washing (5×) and drying at 90°C. The unique morphologies of these precipitates are depicted by the FESEM micrographs given in Figs. 2(a) and (b). These little (200 to 500 nm long) “zeppelins”
of \( \alpha \)-GaO(OH) were found to be crystalline with an orthorhombic unit cell \( (a = 4.5606(3), b = 9.7975(4), c = 2.9731(2) \text{ Å}, V = 132.85 \text{ Å}^3) \). The granular surface texture of the zeppelins is thought to be due to the local dehydration under long electron-beam exposure.\(^{53}\) The zeppelins also had a pronounced elongation in the (001) crystallographic direction. On the other hand, the precipitates recovered from the same solution (with urea) after 95 min of aging (at 90 \(^\circ\)C, pH 6.10) in the mother liquor still had the zeppelin morphology (see the SEM micrograph of Fig. 2(c)), but their lengths were found to increase to about 1 to 2 \( \mu \text{m} \). The bigger zeppelins had also the same orthorhombic unit cell as given above.

Figure 2(d), on the other hand, shows the morphology of \( \alpha \)-GaO(OH) precipitates collected from their mother liquors following 24 h of aging at 90\(^\circ\)C (pH 7.8). The zeppelin morphology disappeared by a mechanism of dissolution–precipitation and the spherical particles formed were found to have an average particle size of 50–60 nm. A similar dissolution–precipitation process was previously observed by Ramanathan et al.\(^{57}\) in the precipitation of crystalline boehmite \( (\text{AlO(OH)}) \) during the aluminum nitrate–urea reactions taking place at around 100\(^\circ\)C for prolonged aging times. This unique morphology (zeppelins or ellipsoidal acicular particles) observed by us this time for gallium oxyhydroxide crystals, which were only grown out of urea-containing solutions, has also been encountered by several other previous researchers\(^{55,52,53,58–60}\) working on the aqueous synthesis of \( \alpha \)-FeOOH (i.e., goethite) in the presence of urea. These researchers concluded that the morphology of goethite crystals was chiefly a function of the pH of the system\(^{61}\) and showed morphologic variations along the pH scale. However, it is quite interesting to mention here that a similar ellipsoidal particle morphology was detected by Lu et al.\(^{62}\) during the hydrothermal synthesis of zinc hydroxide powders from zinc nitrate solutions titrated with \( \text{NH}_4\text{OH} \).

(2) Forced Hydrolysis

Gallium oxide hydroxide precipitates formed in aqueous solutions (without urea), on the other hand, were only recovered from their mother liquors at the end of 95 min of aging at 90 \( \pm \) 1\(^\circ\)C and at the final pH value of 1.87, and they possessed a unique rodlike morphology as shown in the SEM micrograph of Fig. 3(a), with an average rod length of about 3 \( \mu \text{m} \). These rods were also crystalline. Formation of rodlike GaO(OH) is considered to proceed according to the following reaction:

\[
\text{Ga}^{3+} + 2\text{H}_2\text{O} \rightarrow \text{GaO(OH)} + 3\text{H}^+
\]

and the solution pH is slightly decreased with aging (Fig. 1).

A tentative XRD pattern (which is of better crystallographic quality than that of ICDD PDF 06-180 assigned to \( \alpha \)-GaOOH) has been created for these crystals and is given in Table I. The lattice parameters of these orthorhombic crystals were determined to be \( a = 4.5545(2), b = 9.8007(4), c = 2.9738(1) \text{ Å}, Z = 4, V = 132.74 \text{ Å}^3 \). The powder XRD data for these precipitates (without urea) are given by the bottom trace (labeled 90\(^\circ\)C) of Fig. 4. On the other hand, the little and big zeppelins formed in the urea-containing precipitation solutions also had the same XRD trace, with the same relative intensity values. The crystal structure presented by both kinds of gallium oxide hydroxide crystals was found to be analogous to the structure of diaspore, i.e., \( \alpha \)-AlOOH\(^{27,28}\) and the space group of the diaspore structure is regarded to be \( Pbnm \) (62). Goethite and groutite (i.e., \( \alpha \)-MnOOH\(^{31}\)) are also structurally similar to diaspore. However, the GaO(OH) crystals that we synthesized certainly did not have the \( Pbnm \) space group; instead they belonged to either one of the following two space groups: \( Pna2_1 \) (28) or \( Pnma \) (51). The reflections for 201, 011, 401, and 601 planes are forbidden for the space group of \( Pbnm \).

It should be noted that gallium nitrate solutions (without urea) are not stable. We observed several times that even the 0.37\( M \) stock solutions underwent \textit{in situ} hydrolysis (by forming tiny white precipitates at the bottom of the glass containers) in about 3 months, while they were just being stored in a fume hood kept at 21 \( \pm \) 1\(^\circ\)C. The simple way that we performed the forced-hydrolysis experiments during the course of this study also showed that only heating dilute gallium nitrate solutions resulted easily in the formation of monodispersed GaO(OH) crystals.
The recent work of Avivi et al. describes the ultrasonification of \( \text{GaCl}_3 \) solutions for 6 h, and mentions the strong necessity of "ultrasonic hydrolysis for the sonochemical formation" of scroll-like cylindrical or tubular gallium oxide hydroxide particles. It should be remembered that the insertion of an ultrasonic finger into a solution, and operating it (at 100 W/cm\(^2\)) in that solution for a long time like 6 h would certainly cause a lot of heating of that solution. What these authors actually observed at the end of 6 h might simply have been the forced hydrolysis of a gallium solution at an elevated temperature, in strong contrast to their sonochemical hypotheses about the nucleation of nanosized particles with a unique rodlike morphology between the interface of the thin liquid layer and the collapsing air bubbles.

\( \text{(3) Morphology of GaO(OH) Crystals} \)

Formation of zeppelin-like (or spindlelike) or rodlike particles under different conditions was reported by previous researchers in synthesizing \( \alpha-\text{FeO(OH)} \), via hydrolysis, from \( \text{Fe(NO}_3\text{)}_3 \), \( \text{FeSO}_4 \), or \( \text{FeCl}_3 \) solutions. In some of these studies, hydrolysis of the inorganic solutions was typically achieved by the addition of a strong base, such as NaOH. Goni-Elizalde et al. on the other hand, studied the homogeneous precipitation of \( \alpha-\text{FeO(OH)} \) spindles (or zeppelins) in iron(III) sulfate solutions hydrolyzed by urea.

\( \alpha-\text{GaO(OH)} \) crystals consist of double chains of edge-shared octahedra, where \( \text{Ga}^{3+} \) ions are surrounded by six oxygens and the \( c \)-axis \((001)\) lies parallel to the longitudinal direction of these zeppelins, strongly analogous to the case of \( \alpha-\text{FeO(OH)} \), while the vertical and horizontal axes are both \( \{110\} \). This implies that the preferred growth direction along the width of the zeppelins should be along the \( b \)-axis \((010)\), and the same direction along the thickness should be along the \( a \)-axis \((001)\) of the unit cell. Since each particle is a single crystal, the zeppelin-shaped cross section of the particle has a crystallographic acute angle of about 50°. \( \alpha-\text{FeO(OH)} \) and \( \alpha-\text{GaO(OH)} \) have exactly the same diaspore-like crystal structure, and for this reason, it is not surprising at all to see the precipitation of same zeppelins or spindles in both cases of oxyhydroxides. The growth of \( \alpha-\text{GaO(OH)} \) crystals is being realized by the continuous linking of the \( \text{Ga}^{3+} \text{O}_x \) chains along the \( c \)-axis. Kurokawa et al. reported the transition from rodlike to spindlelike (i.e., zeppelin) morphology in \( \alpha-\text{FeO(OH)} \) with the addition of \( \text{OH}^- \) ions (via NaOH) into the solution, which supports the findings of the present study. Decomposition of urea (into \( \text{NH}_2\text{OH} \) and \( \text{H}_2\text{CO}_3 \)) occurs at around 80°C and it is followed by dissociation of the reaction products to yield \( \text{OH}^- \) and \( \text{CO}_3^{2-} \) ions. The gradual but uniform release of \( \text{OH}^- \) ions into the mother liquor results in the nucleation (Figs. 2(a) and (b)) and growth (Fig. 2(c)) of uniformly sized and zeppelin-shaped crystals. Apparently,
the OH\textsuperscript{−} ions, supplied by the homogeneous decomposition of urea, allow nucleation and growth to take place also along the 110 and 010 directions of the formed zeppelins. Structural OH\textsuperscript{−} ions found along the faces of the diaspore (or goethite) unit cell might also contribute to the growth of the crystals along their width parallel to the 010 direction. However, in the case of pure water (i.e., in the absence of excess OH\textsuperscript{−} ions) α-GaO(OH) rodlike crystals are observed to grow only along the longitudinal c-axis. The exact role of CO\textsubscript{3}\textsuperscript{2−} ions in this mechanism is not yet clear.

(4) Calcination of Gallium Oxide Hydroxide

The zeppelin morphology of GaO(OH) crystals, which were initially formed in the presence of urea, drastically changed following calcination for 6 h in an air atmosphere at 750° and 1000°C, as shown in the SEM micrographs of Figs. 2(e) and (f), respectively. The aggregates of fused spheres, which totally replaced the initial GaO(OH) zeppelins, especially as seen in Fig. 2(f), had average radii of about 150 nm. Calcination at temperatures below 750°C (i.e., 250° and 500°C), on the other hand, did not cause any significant change in the zeppelin morphology. The calcination behavior of GaO(OH) crystals grown in pure water (without urea) can be followed by the micrographs of Fig. 3. The SEM micrograph given in Fig. 3(b) displays the morphology of gallium oxide hydroxide crystals after calcination at 250°C for 6 h. These crystals still had the same α-GaO(OH) XRD pattern as with those of shown in Fig. 3(a), and had the same morphology, as well. The FESEM micrographs of Figs. 3(c) through (f) show the surface morphology of rodlike crystals after calcination for 6 h in an air atmosphere at 500°, 750°, 1000°, and 1200°C, respectively. The rods preserved their initial shapes (in contrast to zeppelin-like crystals, which formed in urea-containing solutions) at all calcination temperatures, but they became nanoporous on their visible surfaces with increasing calcination temperature. The formation of these nanosized pores is due to the elimination of water from constitutional OH groups, and this process starts at temperatures above 300°C. Finally, as seen from the micrograph of Fig. 3(f), after calcination as a loose powder compact at 1200°C for 6 h, the rodlike crystals fused to each other at their mutual point of contact. ICP-AES analysis performed on all of the calcined powders did not show any contamination by Al, which might have arisen by the use of Al\textsubscript{2}O\textsubscript{3} boats during calcination.

The powder XRD data of precipitated GaO(OH) crystals, as a function of calcination temperature, is given in Fig. 4. The XRD traces were found to be the same for both kinds (i.e., with urea and without urea) of powders synthesized in this study. α-GaO(OH) was still the only phase after calcination at 250°C for 6 h. Powders
calcined at 500°C for 6 h, on the other hand, displayed the rhombohedral polymorph of Ga2O3. This phase was identified by ICDD PDF 06-0503. The only phase identified in the XRD spectra of 750°C, 1000°C, and 1200°C was the monoclinic polymorph of Ga2O3. This phase was identified by ICDD PDF 41-1103 and 76-0573. With an increase in the calcination temperature from 750°C to 1200°C, the intensities of the XRD peaks have increased considerably.

Knowing that the commercially available Ga2O3 powders (e.g., Sigma-Aldrich) possess the monoclinic crystal structure, and that it is almost impossible to convert them to the rhombohedral polymorph by heating, for instance, at 500°C even for prolonged times, the synthesis method described here can also be regarded as a simple way of manufacturing the rhombohedral Ga2O3 powders of uniform morphology.

Table I. Powder XRD Pattern of Gallium Oxide Hydroxide

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The results of the TG/DTA analyses of GaO(OH) crystals produced by using both kinds of synthesis procedures of this study are summarized in Fig. 5. It should be mentioned here that the DTA trace for the crystals produced in the presence of urea has been lowered by about 4 μV, during plotting, to avoid overlapping with the other DTA trace. All other traces are plotted as they were obtained.

Gallium oxide hydroxide produced by “forced hydrolysis” displayed a total weight loss of 13.01% (from 50°C to 110°C, 0.38%; from 110°C to 250°C, 1.26%; from 250°C to 350°C, 2.47%; from 350°C to 440°C, 8.86%; and from 440°C to 700°C, 0.04%), while the samples produced in the presence of urea showed a total weight loss of 13.38% (from 50°C to 110°C, 2.85%; from 110°C to 250°C, 2.75%; from 250°C to 350°C, 2.75%; from 350°C to 440°C, 4.50%; and from 440°C to 700°C, 0.53%). It can immediately be stated at this point that the formula of the freshly precipitated crystals was not simply GaO(OH). If they indeed were stoichiometric GaO(OH), then the total weight loss must have been in the vicinity of 8.8%.

The TG/DTA traces also indicate a significant difference between the dehydration nature of “urea-” and “no-urea-precipitated” crystals. The total dehydration of urea-precipitated samples took place in three differentiated steps (while the dehydration of samples produced by forced hydrolysis took place in two steps). In the first step (between 50°C and 110°C), the elimination of physical, as well as surface adsorbed, water was achieved, which was also accompanied by an endothermic DTA peak. The urea-precipitated little zeppelins were therefore shown (by the TG analysis alone) to adsorb about 7 times more surface water (2.85% weight loss in the first step) than rodlike crystals (0.41% weight loss). The second step of dehydration of the urea-precipitated samples was identified by a unique exothermic event (which was not observed at all in the forced-hydrolysis samples) at above 250°C. This exothermic event may be attributed to the release of surface bound hydroxyl groups. The granular surface texture of the zeppelins of Figs. 2(a) and (b) was believed to result (after long electron beam exposure) from such dehydroxylation. The broad endothermic peaks (for both samples) appearing between 290°C and 400°C were due to the elimination of water from constitutional OH groups and conversion to Ga2O3. The exothermic event observed in both samples over the temperature range of 250°C to 350°C was attributed to the release of surface bound hydroxyl groups. The granular surface texture of the zeppelins of Figs. 2(a) and (b) was believed to result (after long electron beam exposure) from such dehydroxylation. The broad endothermic peaks (for both samples) appearing between 290°C and 400°C were due to the elimination of water from constitutional OH groups and conversion to Ga2O3. The exothermic event observed in both samples over the temperature range of Fig. 4. XRD spectra of samples of this study as a function of temperature (90°C and 250°C, orthorhombic α-GaOOH·H2O; 500°C, rhombohedral Ga2O3; 750°C, 1000°C, and 1200°C, monoclinic Ga2O3).
635° to 670°C was due to the crystallization of monoclinic \( \beta \)-Ga\(_2\)O\(_3\). Apparently, for “urea-precipitated” powders the observation of a second exothermic shoulder at around 700° to 780°C was due to the slower and stepwise nature of crystallization taking place in these samples.

For the GaO(OH) crystals prepared from either urea-containing solutions or pure water solutions, representative infrared spectra are given in Fig. 6. The spectra labeled 5 through 8 were observed to be exactly the same for both types of samples. (However, the IR traces given as 5 to 8, in Fig. 6, are the spectra of forced-hydrolysis samples.) The IR spectra of 90°- and 250°C-heated samples (traces which are labeled 1 through 4) reveal a broad H–O–H stretching band at around 3400 cm\(^{-1}\) and an OH bending at 1630 cm\(^{-1}\). The bands at 2344 and 1384 cm\(^{-1}\) are due to adsorbed atmospheric CO\(_2\), which resulted from the preparation and processing of FTIR samples in the ambient atmosphere. On the other hand, the bands at 2036 and 1942 cm\(^{-1}\), together with the bands at 1026 and 952 cm\(^{-1}\), are assigned to constitutional Ga–OH bending bands, and their overtones, respectively, in close agreement with Sato et al.\(^{26}\)

The group of bands (including those at 690 and 645 cm\(^{-1}\)) appearing within the range of 1200–600 cm\(^{-1}\) are characteristic M–OH deformation bands seen in hydroxo complexes.\(^{68}\) With increasing calcination temperature, all IR bands attributed to anion vibrations disappear, and gallium oxide absorption bands start to be visible beginning with trace 5 (i.e., sample calcined at 500°C) of Fig. 6. FTIR analysis showed that structural CO\(_3^{2-}\) or NO\(_3^-\)
ions were not present (as supported by the quantitative C and N analyses on those samples) in either type of crystal in this study.

Relying on the combined results of the TG/DTA, FTIR, and C and N analyses, the chemical formula of the rodlike crystals produced (after 90°C drying) by the method of forced hydrolysis was estimated to be GaOOH·0.5H₂O. On the other hand, the formula of the crystals obtained (following drying at 90°C) by the technique of homogeneous decomposition of urea was determined to be GaOOH·0.11H₂O. The coefficients in front of the crystal waters in these formulas may vary on prolonged exposure of those to the atmosphere.

IV. Conclusions

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Acknowledgments

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(4) Carbon and nitrogen analyses, as well as the FTIR results, show that the formed crystals (following drying at 90°C) do not contain any structural CO₃²⁻ and NO₃⁻ ions.

June 2002

Synthesis of Gallium Oxide Hydroxide Crystals

1429


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