Low-Temperature Formation of $\alpha$-Alumina from Polyhydroxoaluminum–Lactic Acid Composite Gels

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1. Introduction

Alpha-alumina (corundum), which is a thermodynamically stable polymorph, is widely used in applications including high-strength materials, electronics and catalytic supports. Recently, due to interest in anisotropically shaped oxide ceramics for metal-ceramic composites and polymer matrix composites, the synthesis of $\alpha$-alumina powders having controlled shape and size tailored to a specific application has attracted much attention.1,2

In general, $\alpha$-alumina is produced by the calcination of various alumina precursors, including aluminum hydroxides such as bayerite, gibbsite and boehmite, and aluminum salts such as sulfate, nitrate, chloride and organic acid salts. Typically, the precursors first transform into metastable transition aluminas such as $\delta$, $\gamma$, $\eta$, $\theta$, $\kappa$- and $\chi$-phase aluminas, depending on the starting species, temperature and ambient atmosphere. The transition aluminas then transform into $\alpha$-alumina upon further heating, and thus $\alpha$-alumina is difficult to synthesize at temperatures below $1100^\circ$C. It is difficult to obtain $\alpha$-alumina having fine primary particles because of the sintering at such high temperatures.3,4 To prevent particle growth, the low-temperature formation of $\alpha$-alumina has also been attempted using various methods, such as sol–gel,5-12 solution,13-17 vapor-phase,18 mechanical/thermal routes19-26 and the introduction of seeds.27-31 However, most of these approaches still require temperatures above $800^\circ$C to form $\alpha$-alumina.

Diaspore ($\alpha$-AIOOH) dehydrates to form $\alpha$-alumina directly at $\sim450^\circ$C.32 This unique transformation is attributable to the structural similarities between diaspore and corundum; both have a hexagonal close-packed anion sublattice. However, natural high-purity diaspore occurs in limited quantities in nature, and its synthesis requires severe hydrothermal conditions ($\sim300^\circ$C, $>30$ MPa). Diaspore-like precursors, which also transform into $\alpha$-alumina at temperatures as low as $450^\circ$C, have been synthesized under moderate conditions.5-12 Brand and co-workers5-8 obtained a mixture of $\alpha$-alumina ($\sim40$ mass%) and amorphous phase by the heat-treating of precursors prepared from the Al–O–H–Cl system at 500–800°C under a rapid flow ($\sim50$ L/h) of air or nitrogen. Kamiya et al.9 synthesized precursors by the sol–gel method using aluminum sec-butoxide. The precursors transformed into $\alpha$-alumina in static air at $500^\circ$C or below, and the fraction of $\alpha$-alumina was $\sim30$ mass% at $600^\circ$C. They assumed that the diasporalike structure was partially built during the sol–gel reaction. Kamiya et al. also reported the effects of seeding the gel with corundum particles; the fraction of $\alpha$-alumina formed from the seeded gel at $600^\circ$C was $40$ mass% or more.10 Smith et al.12 examined the effects of diaspor-seeding on the crystallization of gels prepared from aluminum sec-butoxide and reported a complete transformation to $\alpha$-alumina at $700^\circ$C by $4.1$ mass% seeding.

Concentrated polyhydroxoaluminum (hereafter PHA) solutions prepared by the Al metal dissolution method have such features as a high OH/Al ratio and a high content of the highly polymerized Keggin $\mathrm{Al}_13$ ion ($[\mathrm{Al}_{13}\mathrm{O}_{40}(\mathrm{OH})_{12}(\mathrm{H}_2\mathrm{O})_{24}]^{7-}$) species.25 PHA gels obtained by drying PHA solutions can be transformed into transition aluminas by heat-treatment.26 The preparation of transition aluminas with controlled polymorph and pore properties has been attempted by using PHA composite gels containing organic additives such as polyethylene glycol and quaternary ammonium ions.27-31 The interaction of additives with PHA ions to modulate the gel structure was key in controlling the polymorph and pore properties.

In the present study, lactic acid (hereafter LA) was used as an additive for the preparation of PHA composite gels as alu-
mina precursors. The small LA molecule contains both –OH and –COOH groups. The addition of LA may modulate the PHA gel structure through the interaction of OH groups in PHA ions with the functional groups of LA. The phase transition of the PHA–LA composite gels was investigated. We found that part of the composite gels transformed into α-alumina at low temperatures of around 500°C. Thus, the aims of this study were (1) to clarify the effects of LA content on the thermal transformation of PHA–LA composite gels and (2) to investigate the effects of heat-treatment conditions on the low-temperature formation of α-alumina from PHA–LA composite gels. The results revealed that the low-temperature formation of α-alumina occurred only when the LA content and the heat-treatment conditions were optimized.

2. Experimental

A PHA solution containing 23.4 mass% Al₂O₃ and having an OH/Al ratio of 2.51 was prepared by dissolving Al metal in an HCl solution, as previously described. LA (Wako Pure Chemical Industries) was added to this starting solution and stirred for 30 min to prepare the PHA–LA mixed solutions. The amount of LA was varied from 0 to 50 mass%, based on the mass of Al₂O₃ obtained from the PHA solution. The PHA–LA mixtures were then held at 60°C for 2 d to prepare the PHA–LA composite gels.

The PHA–LA composite gels were heat-treated at 400–1000°C for 3 h in air, and were brought to the heat-treatment temperature at a heating rate of 1.5°C/min. Heating rates of 1.0, 3.0 and 5.0°C/min were also used. The composite gels and their heat-treated products were ground using an agate mortar and pestle, and passed through a 325 mesh (45 μm) sieve prior to characterization.

Powder XRD patterns of the composite gels and their heat-treated products were recorded using Cu Kα radiation with a monochromator. The mass fraction of α-alumina in the heat-treated products was estimated from the relative intensity of the (104) reflection of α-alumina with respect to the (111) reflection of CaF₂ added as an internal standard. The FT-IR spectra of the samples were obtained by means of the KBr tablet method using a spectrometer in the range of 4000–450 cm⁻¹. ²⁷Al-MAS NMR spectra were recorded using a spectrometer at 78.2287 MHz with an MAS frequency of 10.3 kHz. The standard for the ²⁷Al chemical shifts was an aqueous solution of Al₂(SO₄)₃. The morphology of the heat-treated products was observed by means of SEM.

3. Results and discussion

3.1 PHA–LA composite gels

Figure 1 shows XRD patterns of PHA–LA composite gels of varying LA content. The PHA gel and PHA–LA composite gels had a broad reflection peak at approximately 2θ=8° (Cu Kα), ascribable to a loosely stacked structure consisting of highly polymerized PHA ions. The diffraction angle of the peak shifted slightly to the lower angle side when the LA content of the PHA–LA composite gels was increased up to 40 mass%, while additives such as polyethylene glycol and quaternary ammonium ions modulated the composite gel structure so that the peak at approximately 2θ=8° shifted clearly to lower angles with increasing the additive amount. In addition, the halo pattern, which appeared around 2θ=35°, was slightly changed in shape with the addition of LA. These suggest that LA has a slight effect on the modulation of the loosely stacked structure observed in the composite gels.

Figure 2 shows the IR spectra of typical PHA–LA compo-
effect on the modulation of the PHA gel structure or the fluctuation of the coordination number of Al ions.

3.2 Thermal decomposition of PHA–LA composite gels

Figure 3 shows the XRD patterns of the products obtained by heating PHA–LA composite gels at 400–1000°C for 3 h (heating rate to target temperature: 1.5°C/min). Various polymorphs of alumina were crystallized from the composite gels depending on the LA content and heating temperature. In the PHA gel (Fig. 3(A)), γ-alumina formed at 500°C, gradually increasing in amount as the heat-treatment temperature increased. A weak reflection of χ-alumina with a characteristic peak at 42.5° (Cu Kα) appeared above 700°C. Transformation into α-alumina occurred at 1000°C. This crystallization behavior of the PHA gel matched that reported previously.3,25 On the other hand, the PHA–LA5 composite gel partially crystallized into χ-alumina at temperatures as low as 500°C, following which transition aluminas formed from the residual amorphous phase above 600°C with increasing temperature (Fig. 3B). This unusual crystallization behavior, namely, the formation of α-alumina at considerably low temperatures, resembles the thermal decomposition route of natural diasporite (α-AlOOH)C,3,24 A similar phenomenon has been reported by several researchers in the case of sol–gel-derived precursors from AlCl3·6H2O or aluminum alkoxide as a starting material.3,12 In the present work, a small amount of LA (3–10 mass%) was all that was required for the low-temperature formation of α-alumina to occur. The peaks assignable to α-alumina became sharper at 1000°C, accompanied by a rapid decrease in the intensity of the peaks assignable to transition aluminas and also by the disappearance of the halo at 20°–40°. In contrast to composite gels containing ≤10 mass% LA, PHA–LA20% gels retained their amorphous state up to 700°C and transformed into γ-alumina above 800°C (Fig. 3C). A small amount of α-alumina was formed by heating the PHA–LA20% composite gel at 1000°C, due to the transformation of γ-alumina. The transformation behavior of composite gels containing >20 mass% LA was almost the same as that of the PHA–LA20% composite gel. This suggests that the addition of LA to PHA solution is effective for controlling the polymorph of the resultant aluminas, in particular, for producing α-alumina at low temperatures.

Figure 4 shows the mass fraction of α-alumina obtained at different temperatures. As mentioned above, α-alumina was crystallized from PHA–LA composite gels containing 3–10 mass% LA at around 500°C. The fraction of α-alumina increased with LA content up to 5 mass% and then decreased with further increase in LA content (≥8 mass%). On the basis of these results, PHA–LA5% is judged to be the best precursor under the present conditions, with respect to the low-temperature formation of α-alumina.

LA molecules in composite gels with low LA contents interact with PHA ions through their functional groups, combining the clustered domains of PHA ions. However, the addition of excessive amounts of LA leads to the isolation of these clustered domains, allowing the formation of a continuous LA matrix. The counterbalance between these two effects may explain why the PHA–LA5% composite gel yields the maximal α-alumina fraction.

Diaspore, one of the polymorphs of AlOOH, transforms into α-alumina at low temperatures (450°C).24 Kamiya and co-workers3,30 reported that alumina gels prepared under carefully controlled conditions transformed into α-alumina at
500°C or lower, and concluded that direct transformation into α-alumina at such low temperatures was probably caused by the partial formation of a diaspore-like structure during the sol-gel process.8) Hence, it is plausible that a similar process occurs in the PHA-LA system during the heat-treatment process.

The mass fraction of α-alumina increased in two steps: in the low-temperature range of 500 to 600°C and the high-temperature range of 900 to 1000°C. A similar stepwise increase in α-alumina fraction was reported by Kamiya and co-workers.9,10) The first increase in the α-alumina fraction results from the low-temperature formation of α-alumina directly from the precursor gels; the second is due to the transformation from transition aluminas, where the low-temperature α-alumina acts as a seed. In addition, residual amorphous alumina also may contribute to the second increase through the intermediate formation of transition alumina. The fraction of α-alumina remained almost constant between 700 and 900°C. This shows that the addition of LA (3–10 mass%) affects the crystallization of α-alumina, principally below 700°C.

3.3 Effects of heat-treatment conditions on the formation of α-alumina at low temperatures

To determine the effect of heating rate on the low-temperature formation of α-alumina, the PHA-LA5% composite gel was heat-treated under different heating rates. Figure 5 shows the XRD patterns. The formation of α-alumina was promoted by heating at 1.0°C/min, to a greater extent than at 1.5°C/min (refer to Fig. 3(B)). In the case of products obtained at faster heating rates (≥3.0°C/min), α-alumina was not crystallized in the low-temperature range (~500°C), but was rather formed at 1000°C by the transformation of the transition aluminas. Figure 6 shows the α-alumina fraction of the products as a function of the heating temperature. Clearly, the fraction of α-alumina obtained from the PHA-LA5% composite gel increased as the heating rate was reduced. This shows that α-alumina forms at a low temperature (i.e., at 500°C) only when the LA-poor gels (3–10 mass% LA) are heat-treated under low heating rates.

Low heating rates promote the formation of the diaspore-like structure because the time spent at its formation temperature is prolonged. Thus, the larger the diaspore-like region, the higher the amount of resultant low-temperature α-alumina. The fraction of α-alumina remained almost constant between 700 and 900°C regardless of the heating rate, as shown in Fig. 6. This means that the heat-treatment in this temperature range may not directly contribute to the formation of α-alumina. Thus, the formation of the diaspore-like region presumably occurs below 600°C.

The α-alumina fractions at 600–900°C in the present study ranged between 10 and 40 mass%, depending on the LA content and heating rate. Brand and co-workers5–8 achieved ~40 mass% α-alumina at ~800°C from the Al-O-H-Cl system using the precursors obtained from AlCl₃•6H₂O. Kamiya and co-workers8,10 reported ~20 and 40% α-alumina fractions from aluminum sec-butoxide-derived precursors without and with corundum-seeding, respectively, although the precursors and heat-treatment conditions differed from those of the present study. The difference in the fraction of low-temperature α-alumina may partly result from the ability to form the diaspore-like region in the precursors. Although the formation mechanism of the diaspore-like structure in the present composite gels is a problem that remains to be solved, the further enhancement of the low-temperature α-alumina fraction can be expected by increasing the diaspore-like structure in the composite gels through the improvement of the sol-gel and heat-treatment process.

3.4 NMR study and SEM observation of the products

To elucidate the formation mechanism of the low-temperature α-alumina, the coordination number of Al in the products obtained by heating the PHA-LA composite gels under the heating rate of 1.0°C/min at 500°C for 30 min was measured by means of 27Al-MAS NMR spectroscopy. Figure 7 shows the NMR spectra. All the samples gave peaks at around 65 ppm and 5 ppm, which were assignable to 4-coordinated Al (Al⁴⁺) and 6-coordinated Al (Al⁶⁺), respectively. 6) Between these peaks, a peak assignable to 5-coordinated Al (Al⁵⁺) whose intensity depended on the LA content, appeared at around 35 ppm. The heat-treatment products from the PHA gel gave an Al⁴⁺ peak and a weak Al⁶⁺ peak. The intensity of the Al⁴⁺ peak increased with the addition of LA, most markedly in the case of PHA-LA5%. As with α-alumina, the diaspore is composed of only Al⁴⁺. Al⁴⁺ changes easily into Al⁶⁺ by
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Upon heat-treating at low temperatures, the PHA–LA5% composite gel yielded products that contained uniformly distributed mesopores (~3.4 nm in diameter), which is characteristic of PHA gel-derived transition aluminas. This reflects the formation of AlIV in the product obtained by heating a AlCl3·6H2O-derived gel at 300°C and the subsequent change into AlVI at 550°C, accompanied by the decrease in intensity of the AlIV and AlV peaks. The AlV observed at 450–500°C in the present study is an intermediate configuration in the coordination change from AlIV to AlVI. The AlV formed in the amorphous products is most likely to change into AlVI during heat-treatment. AlVI may be partly related to the formation of the diaspore-like structure, resulting in the crystallization of α-alumina at low temperatures. Unlike the PHA–LA5% composite gel, the AlIV peak in the PHA–LA20% sample persisted even after heat-treatment at 700°C. Thus, the addition of a large amount of LA also led to the formation of AlIV below 500°C, but the AlVI in the system was sufficiently stable to suppress crystallization even at 700°C, plausibly because of the coexistence of thermally decomposed LA residues.

Figure 9 shows the SEM photographs of the products obtained by heating the PHA–LA5% composite gel at different temperatures for 3 h. The heating rate was 1.5°C/min. The heat-treatment products, which included α-alumina transformed in the low-temperature range, exhibited a heterogeneous texture, consisting of two different morphology areas; one of fine and uniform particles (Figs. 9(A), (C)), and the other of large aggregates of plate-like crystallites less than approximately 200 nm in size (Figs. 9(B), (D)). The former predominated at temperatures below ~800°C, while the latter increased as the heat-treatment temperature increased and/or the heating rate reduced. These results indicate that the former fine particles correspond to transition aluminas, while the latter plate-like crystallites correspond to α-alumina. In the case of products having a low α-alumina fraction, α-alumina was only partially observed in the heat-treatment products. Thus, the microstructure supports the mechanism of the low-temperature crystallization of α-alumina from PHA–LA composite gels via partial formation of a diaspore-like region. The frequency in the appearance of the plate-like regions corresponded to the formation of α-alumina, and the plate-like crystallites exhibited a more clear-cut morphology at 1000°C, retaining their dimensions of ~200 nm.

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the texture and morphology shown in Figs. 9(A) and (C). However, the formation of α-alumina markedly reduced the specific surface areas of the products, for example, to 24.0 and 6.4 m²/g at 800 and 1000°C, respectively. Unlike the PHA–LA5% composite gel, the products from composite gels containing ≥20 mass% LA had a porous texture consisting of fine, uniform particles (figure not shown), giving sharp and monomodal pore size distributions at around 3.4 nm, and had relatively large specific surface areas (∼120 m²/g) at 800–900°C. This indicates that the addition of larger amounts of LA (∼20 mass%) leads to the formation of uniform mesopores among γ-alumina crystallites upon heating. The formation of mesopores from composite gels containing ≥20 mass% LA probably resulted from the highly dispersed clustered domains of PHA ions in the LA matrix. Indeed, the composite gels containing larger amounts of LA gave a broader reflection with slightly larger d-spacing in the XRD patterns shown in Fig. 1. Thus, composite gels containing large amounts of LA are also interesting from the standpoint of ceramic processing using PHA solutions.

Recently, we reported the fabrication of alumina ceramics with high mechanical strength by pulse electric current sintering (PECS) using PHA gel-derived alumina powders consisting of transition aluminas and seeding α-alumina. 32–34 The α-alumina-containing alumina powders obtained from PHA–LA composite gels in the current study are also promising as starting powders for PECS.

4. Conclusion

Low-temperature formation of α-alumina was attempted using composite gels prepared from polyhydroxolalumnum (PHA) solution and lactic acid (LA). Effects of LA content and heat-treatment conditions on the low-temperature formation of α-alumina were investigated. The results can be summarized as follows:

1. PHA–LA composite gels containing 3–10 mass% LA transformed partially into α-alumina at around 500°C under heat-treatment at a heating rate of 1.5°C/min, while the PHA gel and PHA–LA composite gels containing ≥20 mass% LA transformed into α-alumina at temperatures above 1000°C. The fraction of α-alumina formed from the PHA–LA5% composite gel was ∼25 mass% at 600–900°C.

2. When the heating rate to the target temperature dropped to 1.0°C/min, the low-temperature formation of α-alumina was promoted and the fraction of α-alumina at 600–900°C reached ∼40 mass%. On the other hand, an increase in the heating rate prevented the direct low-temperature formation of α-alumina. This can be explained by the partial formation of a diaspore-like structure in the composite gels during heat-treatment. Lowering of the heating rate promotes the formation of the diaspore-like structure, which can transform into α-alumina at low temperatures, owing to the prolonged stay in the formation temperature range.

3. 27Al-MAS NMR revealed that the heat-treatment products from the PHA–LA5% composite gel contained a large amount of 5-coordinated Al, which changes into 6-coordinated Al with increasing temperature. The resultant 6-coordinated Al seems to be key to the formation of the diaspore-like region, which induces the low-temperature formation of α-alumina in the PHA–LA system.

References