1. Introduction

Conventional Solid Oxide Fuel Cells (SOFCs) of operating temperature at around 900°C is widely studied because almost whole members are made from ceramics not utilizing rare metals such as platinum and fuels are able to be chosen from not only hydrogen but also hydrocarbons, carbon monoxide, and even carbon.1)–4) SOFCs have these usefulness, on the other hand, reliability and durability are problems because of its high operating temperature.5)–8)

Electrolyte thicknesses of electrolyte supported SOFCs were about 150 μm to 1 mm because it must have a certain level of mechanical strength for its self-standing.9) Yttria-stabilized zirconia (YSZ) was often used for the electrolyte owing to its high electrical conductivity and hardness. By the thickness of the electrolyte, the resistance of the electrolyte was large, for that reason, high operating temperature was necessary for SOFC system because YSZ presented the low oxide-ion conductivity and high activation energy.9)

Due to decrease the electrolyte resistance, SOFCs with thin film electrolyte layer have attracted attention over the years.10)–11) Operating temperature could be decreased at the range of intermediate temperature around 600°C by the thin film electrolyte. The low operating temperature could be utilized of low cost metallic interconnects and decrease the thermal stresses and reduce the reaction products.

High temperature at about 1400°C was necessary to be crystallization and sintering for YSZ. When the thin film was generated by the chemical vapor deposition and the Atomic Layer Deposition with the substrate heating, the substrate could not be applied such high temperature because of the durability of the metallic material of the devices. Therefore, the crystallization temperature lower than 1000°C was necessary for the YSZ precursors as the thin film electrolyte SOFCs.

Sintering at high temperature when manufacturing flat plate type or tube type SOFC may cause the problem of reaction with the electrolyte to the other parts. When producing SOFC with a thin film electrolyte, the residual stress between the electrode and the electrolyte also becomes a problem. These problems could be avoidable to sinter SOFCs at low temperatures. It is preferring that the precursor powder of the constituent material can be crystallized and sintered at a low temperature.

Homogeneous precipitation method is one of an advanced method for the conventional precipitation methods. This method is utilized for urea as reducing agent with heating, which generates ammonia. More homogeneous precipitation of hydroxide and/or oxide is obtained by the method because the neutralization reaction is occurred everywhere in the solution. However, synthesis of YSZ by
homogeneous precipitation method has not been reported much because it is in principle difficult that several cations cannot be simultaneously precipitated due to showing inherent degree of solubility for each cation.\(^{12-14}\)

We have been reported that the 8 mol% Y\(_2\)O\(_3\)/92 mol% ZrO\(_2\) (8YSZ) precursor by homogeneous precipitation method with microwave heating could crystallize at lower temperature in shorter time than that with the conventional heating by using autoclave reactor.\(^{12}\) However, detail of precursor and general versatility of this synthesis method for other yttria content YSZ are still unclear. Knowledge concerning general versatility of the present synthesis method can also be expected to contribute to the fields of thin film fuel cells, capacitors and all solid lithium batteries. In this study, the precursors of \(x\) mol% Y\(_2\)O\(_3\)/(100 - \(x\)) mol% ZrO\(_2\) (\(x = 3, 5, 8, 10, 13\)) were prepared by that method due to the investigation of the general versatility. In addition, the crystalline behavior for yttria contents was investigated by X-ray diffraction (XRD), Raman spectroscopy, and differential scanning calorimetry (DSC).

2. Experimental

YZS precursor powder were prepared by urea homogeneous precipitation method with microwave irradiation. Y\(_2\)O\(_3\) powder (Wako, reagent grade, 99.9%) was dissolved into nitric acid (Wako, special grade, 60%). ZrOCl\(_2\)
\(\bullet\)8H\(_2\)O powder (Wako, reagent grade, 99.0%) was dissolved into distilled water and then both solutions were mixed as the total cation concentration of 0.3M adjusted by deionized water and then urea was added into the solution. The solution in the autoclave reactor, containing the cation and urea in a 1:4 mole ratio, was heated by microwave irradiation (2.45 GHz, 500 W) for 7 min using by Shikoku Instrumentation CO., LTD., SMW-060. The autoclave reactor, PFADJ-0120-02, was made from perfluoroalkoxy alkanes by WINTEC. The water in the autoclave can be heated to 118.5 ± 2.5°C within 7 min by microwave irradiation, which was monitored by a heat label (MICRON Corp., 6R-99). The maximum internal pressure of the autoclave was less than 0.5 MPa. The ion ratios of Zirconium/Yttrium in the precursor solution were adjusted for 97/6, 19/2, 23/4, 9/2, 87/26 which were named as 3YSZ, 5YSZ, 8YSZ, 10YSZ and, 13YSZ, respectively. After microwave irradiation, white precipitation was obtained, then washed and centrifuged by distilled water at 4000 rpm then dried over 12 h at 85°C. Crystallization of the precipitation underwent at 500°C for 5 h in the static air. For comparison, specimens of conventional heating process were also carried out in the stainless autoclave. At 100°C for 5 h, no precipitation was obtained. At 120°C for 5 h, precipitate was obtained clearly. The precipitate crystallized in the same manner as the microwave heating.

Crystallization behavior and crystal structure of the precursors and the specimens were investigated by XRD Bruker Corp. D8 DISCOVER, Raman spectroscopy HORIBA Ltd. XploRA, and Fourier Transform Infrared Spectroscopy (FT-IR) Shimadzu Corp. FTIR-8400.

Calorimetric studies of crystallization were estimated by DSC Shimadzu Corp. DSC-50. DSC measurements were carried out until 500°C from room temperature. Heating and cooling rates were 10°C/min.

Particle size and morphology of the precursors and the specimens were measured by Field Emission Scanning Electron Microscope (FE-SEM) JEOL Ltd. JSM-7100F.

Sintering performances were evaluated from absolute density of the sintered pellets. The absolute density was bulk density calculated from the volume and weight of the pellets. The specimens after drying at 85°C was shaped into pellet of \(\phi\) 10 mm by uniaxial pressing under 20 MPa for the density evaluation. The evaluation was carried out at room temperature after heat treatments at 500 to 1200°C each 100°C.

3. Results and discussion

Figure 1(a) shows the SEM image of the 8YSZ precursor powder by the microwave heating. The shape of the precursor particle was a roundish cuboid, which was agglomerated. The primary 8YSZ precursor particle size was around 300 nm. Figure 2(a) shows the SEM image of the 8YSZ precursor powder by the conventional heating. The shape of the 8YSZ precursor by the conventional...
heating was more rotund than that by the microwave heating. The primary particle size was distributed from 0.2 to 1.5 μm. Figure 1(b) shows the SEM of the 8YSZ powder after heat treatment at 500°C. The wide particle distribution was observed that the width was from 10 nm to several microns. The shape and size for the conventional heating one was almost the same manner as shown in Fig. 2(b).

Figure 3 shows the XRD patterns of the 8YSZ precursor powder by microwave heating and after heat treatment at 500°C. No peak was observed for the precursor powder as shown in the Fig. 3(a). After heat treatment, the XRD pattern was indexed as a cubic fluorite-type structure as presented in the Fig. 3(b). Figure 4(a) shows the XRD patterns for the 8YSZ precursor powder by conventional heating. A broad peak at around 25° and a peak at 32.7° were observed from the precursor which was consisted with that by the microwave heating. Figure 4(b) shows the XRD patterns of the 8YSZ by conventional heating after heat treatment at 500°C. That was almost consistent with the microwave one.

Figure 5(a) presents the Raman spectra of the 8YSZ precursor by microwave heating and the heated specimen at 500°C. The Raman peaks for the precursor were observed at 374, 536, 717, 1051 cm⁻¹. The Raman peaks at 374 and 536 cm⁻¹ were originated from Zr-O vibration in the precursor. The peaks at 717 and 1051 cm⁻¹ were assigned to Zr-OH vibration, which suggested that the precursor consisted with tetrametric hydroxo-cation \([\text{Zr}_4(\text{OH})_8(\text{H}_2\text{O})_x]^4+\). After 500°C heat treatment, the Raman peak at 626 cm⁻¹ was appeared, shown in Fig. 5(b). This peak was originated from Zr–O vibration in the cubic structure. Moreover, extremely broad peaks at 154, 269, 330, 486 cm⁻¹ were observed for tetragonal phase of YSZ by Raman spectroscopy, which was not observed by XRD. In the precursor, it suggested that tetragonal crystals appeared because not all \(\text{Zr}^{4+}\) and \(\text{Y}^{3+}\)
bonded. Since it was cubic after crystallization, it was thought that $Y^{3+}$ was uniformly dispersed in the precursor and that they were mutually bonded after heat treatment.

The Raman spectra of the 8YSZ precursor by conventional heating and the heated specimen at 500°C are shown in Fig. 6. For the spectrum of the precursor, peaks were observed at the same wavenumber as that of the microwave heating. The precursor by conventional heating was also essentially same for the microwave heating one. However, for the precursor, peak intensity ratio of Zr–O vibration/Zr–OH vibration for conventional heating was lower than that of microwave heating. Figure 6(b) shows the Raman spectrum for the heated specimen at 500°C. The crystal structure after heating was identified as mixed structure both cubic and tetragonal.

Figure 7 shows FT-IR spectra for the as prepared 8YSZ precursor by microwave heating, after annealing at 350°C and 500°C in the static air. As prepared precursor showed the absorption bands assigned to vibration of Zr–O bond at about 480 and 644 cm$^{-1}$, respectively, and also at 1045 cm$^{-1}$ for C–OH bond. Bonds at around 3404 and 1623 cm$^{-1}$ were originated to the vibration from carbonate. After annealing at 500°C, the bonds for OH stretching, water hydration, C–OH and carbonate were disappeared and only appeared Zr–O bond which was assigned tetragonal ZrO$_2$.

In contrast, conventional heating YSZ after annealing at 500°C carbonate bonds still observed as shown in Fig. 8. Figure 9 shows the DSC curves of the specimens by microwave heating for each yttria content during the heating process.
ing. Figure 11 shows the crystallization temperatures estimated by the DSC curves. The crystallization temperatures increased with increasing yttria ratio and were lower for microwave heating than that for conventional heating.

We have been reported that the microwave technique can reduce the crystallization temperature. In this study, this decrease phenomenon was reappeared for 8 mol% yttria, moreover, that was also observed from 3 to 13 mol% yttria. It is interesting that an inclination is different between low yttria region from 3 to 8 mol% and high region of 10 to 13 mol%. In the range of 10 to 13 mol% of yttria, the crystallization temperatures were more decrease than that in the range of 3 to 8 mol%. We believe that this difference is caused by generation ratio of crystal structures both cubic and tetragonal.

Figure 12 presents the density of the pellet for the precursors by microwave heating and conventional heating for each yttria content. The density by microwave heating at room temperature was higher than that by conventional one. At 900 and 1000°C, the density was gently increased according to increase temperature. The density of microwave heating was higher than that of the conventional one in this range. This difference of the sinterability could be caused by microwave heating.

4. Conclusion

The YSZ precursors of x mol% Y₂O₃/(100 – x) mol% ZrO₂ (x = 3, 5, 8, 10, 13) could be successfully synthesized by homogeneous precipitation method with microwave heating. The precursor was found to be containing zirconium hydroxide by Raman spectroscopy. By the DSC measurements revealed that the crystallization temperatures for the microwave heating was smaller than those for the conventional heating. For high yttria content YSZ above 10 mol%, the crystallization temperatures were more decrease than those of the lower yttria content. For microwave heating, the crystallization temperature of the precursor was lower, and the density was higher than that of conventional heating. This advance could cause by microwave technique.

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