Formation of Cerium-Filled Skutterudite Thermoelectric Materials
Sintered from Gas-Atomized Powder*1


1Japan Ultra-high Temperature Materials Research Institute, Ube 755-0001, Japan
2Materials Research Laboratory, Kobe Steel, LTD., Kobe 651-2271, Japan
3Department of Electronics and Computer Science, Science University of Tokyo in Yamaguchi, Onoda 756-0884, Japan

A process for forming the Ce-filled skutterudite thermoelectric materials sintered from gas-atomized powder has been investigated. The rapidly cooled particles obtained by the Ar gas-atomizing method consist of the phases of CeFe3CoSb12 (skutterudite), FeSb2, FeSb and Sb. The differential scanning calorimetry curve indicates that sintering proceeded through the two exothermic reactions: one due to the oxidation of Sb at low temperatures and the other due to the skutterudite phase formation from three phases of FeSb2, FeSb and Sb at high temperatures. On the basis of these results, sintering conditions were best tuned to reduce the oxidation of Sb while ensuring the skutterudite phase formation. Consequently, it becomes possible to increase the Seebeck coefficient, and to decrease the thermal conductivity in the temperature range of 300–850 K. We consider the achievement of favorable results to be mainly attributed to a decrease in the Sb phase remaining in sintered materials.

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1. Introduction

CoSb3-based skutterudite compounds are known as excellent thermoelectric materials.1) CeFe3CoSb12, one of “filled-skutterudite”, which is derived from CoSb3 by partially replacing the Co atoms with Fe atoms and filling cavities in the crystal lattice of CoSb3 with Ce atoms, has received particular attention because of its high thermoelectric efficiency, especially at high temperatures.2)

These materials have been conventionally prepared by the powder metallurgy technique, and the powder as a starting material is generally obtained by the method of both melting and grinding and/or mechanical milling of the raw materials.3, 4) However, these methods seem to be suitable only on a laboratory-scale but not on a commercial-scale. Several attempts have been proposed such as the application of a gas-atomizing method to powder production. The thermoelectric properties of the materials sintered from the powder have been reported.5) In our previous investigation,5) it was pointed out that a combined process of gas-atomizing and sintering is a promising on a production scale. However the sintering condition was not optimized at that time, and the characterization of the atomized powder particles and clarification of the formation process of these materials are required to establish the optimum sintering condition.

In order to appraise the newly adjusted sintering conditions, the characterization of gas-atomized powder, the process of forming the skutterudite phase of CeFe3CoSb12, and the thermoelectric properties of the materials sintered under varying conditions have been investigated in the present study. The difference between the properties of laboratory-scale materials and atomizing-featured ones is also discussed.

2. Experimental Procedure

Two kinds of powders were produced by gas-atomization. Chemical compositions of those materials in weighing are listed in Table 1. CoSb3 was atomized for comparison. In this table, the value of Sb is normalized to 3 in the case of CoSb3, and the total amount of Fe, Co and Sb is also normalized to 16 for CeFe3CoSb12.

Authors pointed that controlling the chemical compositions of atomized powder close to stoichiometry might not be the best way to achieve better thermoelectric properties only in the combination process of gas-atomizing and sintering.6) A slight amount of excess Sb over stoichiometry is rather effective for reducing the forming of Ce-rich portions on grain boundaries of compounds.6) In present study, therefore Sb was weighted with a slightly rich composition than that in stoichiometry for the reason described above and in order to compensate for the vaporizing loss during melting.

Table 1 Chemical compositions of specimens.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Ce</th>
<th>Fe</th>
<th>Co</th>
<th>Sb</th>
</tr>
</thead>
<tbody>
<tr>
<td>CoSb3</td>
<td></td>
<td></td>
<td>0.94</td>
<td>3</td>
</tr>
<tr>
<td>Weighed</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Analyzed</td>
<td></td>
<td></td>
<td>0.81</td>
<td>3</td>
</tr>
<tr>
<td>CeFe3CoSb12</td>
<td>0.90</td>
<td>2.74</td>
<td>0.92</td>
<td>12.34</td>
</tr>
<tr>
<td>Weighed</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Analyzed</td>
<td>0.94</td>
<td>2.74</td>
<td>0.92</td>
<td>12.34</td>
</tr>
</tbody>
</table>
Gas-atomization was carried out in the following manner. Ce ingots with a purity of 99.95%, Fe ingots with a purity of 99.99%, Co flakes with a purity of 99.9% and Sb ingots with purity of over 99.999% were used as raw materials. The weighed metals were melted in a graphite crucible with a high-frequency induction furnace. Melting took place in vacuum below 473 K, and in a pure Ar (99.999%) atmosphere above 473 K. In this experiment, Fe, Co and Sb were melted initially, and then Ce was added to the liquid metal mixture of Fe–Co–Sb at 1473 K to avoid the formation of CeSb with a high melting point. The liquid materials were injected through a nozzle capillary by a surge pressure of 0.18 MPa, and then a fast Ar gas jet with a pressure of 4.9 MPa blew the liquid stream off. In this manner, rapidly cooled and spherical particles were obtained. Fine particles of less than 100 μm in diameter were selected in an Ar atmosphere for sintering and analyses.

Chemical compositions were analyzed by using inductively coupled plasma spectrometry (ICP). Optical microscopy (OM), an electron probe microanalyzer (EPMA) and X-ray diffraction (XRD) were used to identify the phases of the atomized powder. Differential scanning calorimetry (DSC) was performed from 300 K to a maximum temperature of 973 K to detect a phase transformation during sintering. Measurements of DSC were done at a heating rate of 0.67 K/s utilizing a four-probe method utilizing a DC current was used for measurements of \( \sigma \). The thermal diffusion coefficient \( \kappa \) was chosen as 3 to 5 K under ordinary conditions. \( \alpha \) and \( \omega \) were obtained as the quotient of \( \nu \) over \( \delta T \). A standard four-probe method utilizing a DC current was used for measurements of \( \sigma \). The thermal diffusivity coefficient \( \lambda \) and the specific heat \( c \) of the sample of 10 mm in diameter and 2 mm in thickness were measured by the laser-flash method at room temperature and in the range from 473 to 873 K at intervals of 100 K. The thermal conductivity \( \kappa \) was calculated from the formula \( \kappa = \lambda c \rho \), where \( \rho \) is density.

3. Results and Discussion

3.1 Characterization of atomized powder and skutterudite formation

The analyzed chemical compositions of gas-atomized powders are shown in Table 1. As explained below, as-atomized powder particles consist of plural phases. These analyzed compositions, therefore, indicate averaged values.

Figure 1(a) represents the DSC curve of CoSb\(_3\) from 300 to 873 K. An exothermic peak is observed in the temperature range from 650 to 800 K. Uchida et al. reported that as-atomized CoSb\(_3\) powder particles consisted of CoSb, CoSb\(_2\) and CeSb. The CoSb\(_3\) skutterudite phase formation from these three phases took place through the temperature elevation to 873 K during sintering. Since the powder particles used in this study involve the same phases as the three phases above, it is considered that the exothermic peak corresponds to the CoSb\(_3\) skutterudite formation reaction. In comparison with CoSb\(_3\), the DSC curve of CeFe\(_3\)CoSb\(_{12}\) is slightly more complex, as shown in Fig. 1(b). There is a small exothermic reaction (peak-I), which seems to be induced by a group of two or more reactions, in the temperature range of 530 to 650 K. After the peak-I, another exothermic reaction (peak-II) occurs from 680 to 950 K. This peak most likely corresponds to the CeFe\(_3\)CoSb\(_{12}\) skutterudite formation reaction from the case for CoSb\(_3\). The reaction however continues over a broader temperature range than that of CoSb\(_3\), and furthermore the end of the reaction shifts to a higher temperature than the endothermic peak due to melting of Sb.

In order to identify the phase transformation associated with these exothermic peaks, XRD was measured at the temperatures of 300 K, 523 K (just below the onset of the peak-I), 573 K (the center of the peak-I), 673 K (just above the end of the peak-I), 773 K and 873 K. Line profiles obtained at the respective temperatures are shown in Fig. 2. As is obvious from Fig. 2, some peaks are weakened and disappear during heating, and a single skutterudite phase is eventually formed. Apparently, there are several peaks, relative intensity ratios of which are almost kept constant during heating, such as the peaks indicated by solid circles in Fig. 2.

Judging from these results for the case of CeFe\(_3\)CoSb\(_{12}\), we tend to believe that a part of the skutterudite phase al-
ready exists in as-atomized particles and its formation from the remaining phases is accelerated during sintering, which is clearly different from the case of CoSb$_3$. Therefore, the peaks of the XRD profile for the as-atomized powder can be consistently identified as consisting of four phases of skutterudite, FeSb$_2$, FeSb and Sb, as shown in Fig. 3. Since the lattice parameter of the skutterudite phase is definitely expanded by filling of Ce, peaks shift to lower angles than those of CoSb$_3$. FeSb$_2$ represents in the FeSb$_2$ type structure (FeSb$_2$, CoSb$_2$ and CeSb$_2$) and FeSb represents in the CoSb type structure (FeSb and CoSb). Both groups give rise to diffraction patterns similar to each other and cannot be distinguished from the measured spectrum.

Figure 4 represents the temperature dependence of intensities of the three specific peaks (marked with open arrows in Fig. 3) normalized with respect to the highest peak of the skutterudite phase (marked with a solid arrow). All intensities exhibit a plateau up to 673 K with subsequent monotonic decrease with increasing temperature. This result indicates that the peak-I on the DSC curve has nothing to do with the skutterudite formation, while the peak-II should correspond to the skutterudite formation.

The microstructures of heat-treated powders obtained in the several stages during DSC measurement and that of as-atomized particles were investigated. Figure 5(a) shows an optical micrograph of as-atomized particles. From the results of EPMA, it is determined that the gray areas represent a mixture of skutterudite, FeSb$_2$ and FeSb, and the white portions correspond to Sb. Black and acicular structures were identified as the Ce-oxide. As compared with the as-atomized powder, the microstructures of the powders heated up to 523 K, 573 K and 673 K are similar to that of the as-atomized powder. Although segmentations of Ce oxides are observed in some parts of the powder heated to 773 K, the micro-structural elements are similar to those of the as-atomized powder as shown in Fig. 5(b). No essential change could be detected in the XRD profiles in the temperature range up to the end of the peak-I. Since the peak-I accompanies any micrographic or structural change, it will be conjectured that the peak-I corresponds to the oxidation of elements involved. We carried out the DSC measurement for Sb fine powder of less than 100 µm in diameter under the same condition as that for the atomized powders. An exothermic reaction was observed as shown in Fig. 1(c). From the results of XRD for the powder after DSC measurement, this reaction was judged as the oxidation of Sb. In comparison between this reaction and the peak-I, heat flow of each reaction are approximately similar, whereas the temperature range are slightly different. Therefore, it is highly likely that the peak-I corresponds to the oxidation reaction of pure Sb remaining in the atomized particles.
3.2 Sintering and thermoelectric properties

As discussed in Section 3.1, the sintering process of gas-atomized CeFe$_3$CoSb$_{12}$ powder involves the stages of the oxidation of Sb and the formation of the skutterudite phase. From this point of view, the sintering condition was elucidated in comparison with that of Uchida et al. In their work, the experiments had been done in an Ar gas atmosphere at the temperature of 873 K in order to avoid the vaporization of Sb. In the present study, the sintering was carried out under vacuum to minimize the oxidation of Sb up to the temperature of 923 K to ensure the complete formation of the skutterudite phase.

Sintered compounds were produced by holding at 873 K or 923 K in vacuum. Another sintering was also carried out at 873 K in an Ar atmosphere, the condition used by Uchida et al. for comparison. Other parameters such as the heating rate, holding time and compression were kept constant for all specimens.

Figure 6 shows the temperature dependence of the Seebeck coefficient, $\alpha$ under different sintering conditions. The value of $\alpha$ is enhanced in the whole temperature range by changing the atmosphere from Ar to vacuum and by a rise of the sintering temperature from 873 K to 923 K. It is apparent that the formation of the skutterudite is enhanced as a result of the reduction of the oxidation of Sb, and also the volume fraction of the residual Sb is reduced by holding at a higher temperature in vacuum. Figure 7 shows optical micrographs of sintered materials in vacuum at 873 and 923 K. Residual Sb (white portion) is clearly seen along boundaries of atomized particles in the case of 873 K. However, the amount of the residual Sb decreased drastically when the specimen was held at 923 K due to evacuation of melted Sb to the outside of the mold during sintering. We consider the phenomena described above to contribute to a decrease in the carrier concentration and, hence, an increase in $\alpha$.

Since the amount of Sb, which serves as a good conductor both of electricity and heat, is reduced, both $\sigma$ and $\kappa$ decreased in the following order; 873 K in Ar, 873 K in vacuum and 923 K in vacuum as shown in Figs. 8 and 9.

A dimensionless figure of merit ($ZT$) calculated from $\alpha$, $\sigma$ and $\kappa$ is shown as a function of temperature in Fig. 10. In all specimens, $ZT$ values increase monotonously as the temperature increases until the maximum value is achieved at around 700 K, and then start to decrease as the temperature further increases. The elaboration of sintering conditions resulted in a significant enhancement in $ZT$. The maximum $ZT$ value of 0.63 for the specimen sintered at 923 K in vacuum is obtained at 700 K, which is 0.2 greater than that obtained at the same temperature for the specimen sintered at 873 K in an Ar atmosphere.

The maximum $ZT$ value obtained in this study is lower than the value of 0.9 at 700 K reported by Fleurial et al. More specifically, at 700 K, the value of $\alpha$ in this study is slightly lower than that of their report. $\sigma$ and $\kappa$ in this study are larger than each value of their report. The difference of these three thermoelectric properties between the two studies could be explained mainly by the still existing residual Sb, as pointed out by Uchida et al. Therefore, if the more residual Sb were pushed out from a compound during sintering, the smaller the difference would become. The optimum combination of the holding time and the pressure at high temperatures during sintering must be found out.
Reduction in the thermal conductivity by the filling of Ce ("rattling") may not function effectively in our specimens. As mentioned above, Ce-oxides already exist in atomized powder particles, and remain even after sintering. Since Ce is added under the assumption that the amount of Ce would be enough to fill the cavities in this study, the generation of Ce-oxide causes the shortage of Ce-filling into cavities. And it is thought that this is another reason why the thermal conductivity has not been sufficiently improved.

Kitagawa et al.\textsuperscript{9)} presented the relationship between the XRD intensity ratio of (211) to (310) and the Ce-filling fraction ($f$) in the melt-spun Ce$_x$Fe$_{8-x}$Co$_x$Sb$_{24}$ ribbon. Meisner et al.\textsuperscript{10)} reported the relationship between the Ce-filling fraction ($y$) and lattice constants of Ce$_y$Fe$_{4-x}$Co$_x$Sb$_{12}$ with varying $y$. The value of $f$ or $y$ for the specimen showing the best $ZT$ value in this study (sintered at 923 K in vacuum) was estimated using both results. The value of $f = 1.56$ was derived from the intensity ratio of $I_{211}/I_{310} = 0.011$ in the former case, and the value of $1.56$ is equivalent to $y = 0.78$ in the latter notation, namely, Ce$_y$Fe$_{4-x}$Co$_x$Sb$_{12}$. While the lattice constant turned out to be as 0.9076 nm, and it yielded the value of $y = 0.46$ in the latter case.

The Ce-filling fraction is affected by the value of $x$, the substitution number of Fe by Co, and $x$ is different between for our specimen and for theirs. Therefore it is not clear whether the Ce-filling fractions estimated above for our specimen are reasonable or not. Anyway, reducing the oxidation of Ce in atomizing process is required in order to enhance the
Ce-filling fraction to 1.0.

It is reported that the fine oxide particles or layers reduce the thermal conductivity and improve the thermoelectric properties.\textsuperscript{11,12} Although the permissible size of them for the effect has not been clarified, since the size of the Ce-oxides observed in this study is rather large up to several micrometers in the long axis (see Fig. 7), they might exert negative effects on the thermoelectric properties. Ce-oxides are thermodynamically very stable. For example, the standard free energy of formation ($\Delta G^0$) for Ce$_2$O$_3$ is $-1676.9$ KJ/mol and $-983$ KJ/mol, respectively.\textsuperscript{13} Since the standard free energy of formation for H$_2$O is $-219.0$ KJ/mol,\textsuperscript{13} Ce-oxides could not be reduced by heat-treatment even in a hydrogen atmosphere. From the above, controlling the oxidation of Ce as low as possible during atomization is strongly desired to achieve excellent thermoelectric properties. And keeping as small size of Ce-oxides as possible might be also effective. The former will become possible by using some kind of Ce alloy (e.g. Ce–Fe) instead of pure Ce, the latter by accelerating cooling speed in the injection (solidification) process of gas-atomization.

4. Conclusion

The formation of Ce-filled skutterudite thermoelectric materials sintered from gas-atomized powder was investigated. The conclusions obtained in this work are as follows;

(1) A gas-atomized powder particle of CeFe$_3$CoSb$_{12}$ consists of a skutterudite phase, FeSb$_2$, FeSb and Sb.

(2) The oxidation of Sb and the formation of the skutterudite phase take place during sintering in the written order.

(3) Adjusting of the sintering conditions to minimize the oxidation of Sb and to ensure the formation of skutterudite causes increases in the Seebeck coefficients and decreases in the electrical conductivities and the thermal conductivity simultaneously. Such results are attained mainly by decreasing the residual Sb in sintered materials.

(4) The highest value 0.63 of $ZT$ is obtained at 700 K for the material sintered at 923 K in vacuum.

(5) Controlling the oxidation of Ce as low as possible is necessary in order to enhance Ce-filling fraction.

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