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I. INTRODUCTION

The conversion of missed heat generated during industrial processes into electricity based on thermoelectricity is of particular interest and is becoming more and more crucial in the current environmental context. Direct conversion of thermal energy into electrical energy requires the use of thermoelectric (TE) modules. These modules are composed of TE material pairs which are connected in series, electrically and in parallel, thermally. Each couple is made of two legs of n- and p-type semiconductors. Applying a temperature gradient results in the carriers’ mobility from the hot to the cold sides and the generation of a voltage through the Seebeck effect. The efficiency of the TE module is dependent on the TE properties of the leg materials which are determined from conductivity, and more commonly by the dimensionless figure of merit, $Z$ ($Z = \alpha^2\sigma/k$, where $\alpha$ is the Seebeck coefficient, $\sigma$ the electrical conductivity, and $k$ the thermal conductivity), and more commonly by the dimensionless figure of merit, $ZT$ (where $T$ is the temperature). Skutterudite compounds have been reported to be promising TE materials for applications in the medium temperature range from 400 to 600 °C. Doped or filled CoSb$_3$ and CeFe$_4$Sb$_{12}$-based skutterudites have been shown to be good candidates for n- and p-type legs of TE devices, respectively. ZT values as high as 0.85 were obtained for CeFe$_4$Sb$_{12}$ and CoSb$_3$ compounds at about 400 °C and as high as 1.9 for doped skutterudites at 550 °C. However, the performance of TE devices is strongly dependent on the connection and adherence between TE legs with metal electrodes and on the quality of the interface. One of the key issues, particularly for the hot-side TE material/metal electrode joint. Maintaining at high temperature can influence the performance of the device drastically by elemental interdiffusion and interfacial reactions which raise the interface electrical resistivity and decrease its mechanical properties.

Consequently, a diffusion barrier layer is required to avoid any interfacial reaction between the thermoelectric substrate and the hot-side electrode. Investigations have been carried out to develop joining of CoSb$_3$-based skutterudite couples with metal electrodes. Fan et al. studied the effectiveness of a Ti foil in joining a CoSb$_3$ substrate to a Mo electrode by spark plasma sintering at 580 °C under a pressure of 40 MPa in vacuum. They observed the formation of an intermediate TiSb layer at the CoSb$_3$/Ti boundary and of a Ti–Mo solid solution layer with graded composition at the Ti/Mo interface. They assumed that this Ti–Mo graded layer may contribute to the bonding between Ti and Mo and that the interfacial resistance in the CoSb$_3$/Ti/Mo is minimized. They showed that the thickness and the composition of the intermediate layer remained unchanged after annealing at 500 °C during 1000 h under vacuum. Zhao et al. investigated the interfacial reactions
in CoSb3/Ti/Mo–Cu joints after thermal aging in the 500–600 °C temperature range from 0 to 30 days in vacuum. They showed the formation of an intermetallic compound layer composed of TiCoSb, TiSb2, and TiSb at the interface. The thickness of this layer increases with aging due to the diffusion of Ti and Sb. This results in a decrease of the shear strength with the formation of cracks at the interface. Chen et al. studied the interfacial reactions in Ni/CoSb3 couples after annealing at 450 °C during 2–24 h. They observed interfacial reactions with the formation of Ni5Sb4/(Co,Ni)3Sb interlayers. Chen et al. investigated Co, Ni, and Ti materials as potential barrier layers for CoSb3/Ag–Cu couples at 450 and 600 °C for 1–44 days in vacuum. They observed interfacial reactions at both temperatures even if they are reduced with the introduction of barrier layers. They reported the formation of CoSb3 and CoSb, Ni5Sb4 and (Co,Ni)Sb, and TiSb, TiSb2 and TiCoSb phases for Co, Ni, and Ti barriers. They concluded that Co is a good barrier at 450 °C with a good adhesion to both CoSb3 and Ag–Cu and relatively low interfacial reaction rates.

While diffusion barriers have been largely investigated for n-type CoSb3 skutterudites, only few results were reported for the p-type skutterudite, CeFe4Co4−xSb12, which is the most promising family. Gu et al. studied the interfacial stability of the p-type Ti/CoFe2Sb2/Sb12 joints in comparison with the n-type Ti/Yb52Co4Sb12 after aging in vacuum at 550 °C from 1 to 30 days. They showed that interdiffusion results in the formation of intermetallic compound layers at the interface. The formation of a bilayer (TiSb2, Ti–Sb) is observed for the p-type one. Moreover, the growth rate of the interfacial layers in the p-type skutterudite joint is lower than in the n-type one. The authors concluded that Ce filling and Fe doping slow down the decomposition of the p-type joint and that Ti is a promising barrier layer material for the p-type CeFe4Co4−xSb12 skutterudite.

Previous studies reported the interest in using TaN against Cu diffusion for microelectronics where TaN was selected due to its high thermal stability and electrical conductivity. More recently, Hsu et al. showed that TaN is a good diffusion barrier for thermoelectric applications up to 200 °C. In a previous work, we studied the efficiency of a TaN interlayer in CeFe5Sb12/Cu after aging at 600 °C. It was found that CeFeSb12 and Cu chemically react and form CeCu2 and CuSb phases. Using 1.5 μm thick TaN thin films as the diffusion barrier showed its efficiency against Ce diffusion but does not inhibit Sb diffusion from the skutterudite substrate to the copper surface film. The formation of a CuSb phase indicates that interfacial reactions induce the decomposition of the skutterudite material and the decrease of the figure of merit.

The current paper focuses on the thermal stability and efficiency of a multilayer, TaN-Ta-TaN, as a diffusion barrier for CeFe5Sb12/Cu in comparison with single TaN layer in the temperature range 400–600 °C.

II. MATERIAL AND METHODS

CeFe5Sb12 skutterudite ingots were synthesized by direct reaction of stoichiometric amounts of Ce, Fe, and Sb elements as previously described. CeFe5Sb12 ingots were cut into 3 mm thick pieces followed by mechanical polishing and then ultrasonic cleaning in ethanol. The TaN 900 nm thick single layer and TaN-Ta-TaN, each 300 nm, multilayer thin films were deposited onto the skutterudite substrate by radio frequency (RF) sputtering as reported in previous papers. A Cu layer, about 1 μm thick, was subsequently sputtered on these barriers. The stability of the multilayer barrier in comparison to the single barrier was investigated after 6 h annealing under vacuum (3 · 10−4 Pa) in the temperature range of 400–600 °C. The interfacial reactions and the phase formation were studied using X-ray diffraction (XRD) and transmission electron microscopy (TEM) associated to energy dispersive X-ray spectroscopy (EDS). A 0–20 diffractometer (Philips Xpert Pro MRD) with a Cu target X-ray tube working at 40 kV and 25 mA was used for crystallography and phase analysis. The microscope was operated at an accelerating voltage of 200 kV. STEM-EDS analyses were obtained with a spot size of 0.7 nm and with acquisition times from 200 to 740 s on an Oxford Instrument X-MaxN 100 TLE EDX detector. TEM specimens were thinned in cross section using the focused ion beam (FIB) technique with a Ga ion beam and attached to the molybdenum sample holder.

III. RESULTS AND DISCUSSION

TEN observations of the CeFe5Sb12/barrier/Cu are reported in Fig. 1. The as-deposited TaN and TaN-Ta-TaN films present a columnar growth perpendicular to the substrate surface. This columnar growth is commonly observed in sputtered films considering the process parameters used as predicted by the model of Movchan and Demchishin extended by Thornton. The columnar grain structure with intercolumn voids and density-deficient grain boundaries is characteristic of sputtered films. The mean grain size of 4 and 6 nm was determined from X-ray diffraction analyses, respectively, for Ta and TaN layers using the Scherrer formula.

A. Study of the TaN single layer barrier

The role of the single TaN layer in elemental interdiffusion was investigated first. Figure 2(i) shows the XRD patterns of the
CeFe₄Sb₁₂/TaN/Cu sample in the as-deposited state and after annealing. In the as-deposited state, the characteristic peaks of CeFe₄Sb₁₂, TaN, and Cu phases are observed. The filled CeFe₄Sb₁₂ phase crystallizes in the cubic structure of the Im-3 space group, where Fe atoms are located at the center of distorted octahedral formed by Sb and Ce atoms. The tantalum nitride thin film is observed to be the δ-TaN phase with an NaCl-type structure and the copper film crystallizes in the polycrystalline fcc-Cu phase. After annealing at 400 °C, the presence of the Sb phase was confirmed by XRD and TEM investigation (Fig. 3). This new Sb phase is formed in the Cu layer. The formation of Sb-rich phases in Cu confirms diffusion of this element through the columnar TaN layer to the Cu layer. Sb is well known as very mobile specie depending on temperature. Cu₂Sb and FeSb₂ phases are formed after annealing in the 500–600 °C temperature range [Fig. 2(i)]. The Cu₂Sb phase results from the diffusion of Sb and the reaction with Cu. Parallel to this, depletion in Sb of CeFe₄Sb₁₂ produces a partial decomposition of the skutterudite and the formation of FeSb₂.

These results point out that a TaN single layer, 900 nm thick, does not appear to be effective against Sb diffusion for the temperature above 400 °C. However, this single layer limits Ce diffusion, inhibiting the subsequent formation of the Cu₂Sb and CeCu₂ intermetallic phases.

B. Study of the TaN-TaN multilayer barrier

The presence of characteristic peaks of Ta, TaN, Cu and CeFe₄Sb₁₂ is remarked in as deposited TaN-TaN multilayer films [Fig. 2(ii)]. After annealing at 400 °C for 6 h, the XRD pattern does not show the presence of the Sb phase as previously observed in the case of the TaN single layer barrier. These results are confirmed also by TEM observations which do not show the presence of any new phase in the Cu substrate and by the Sb concentration profile (Fig. 4). Figure 5 shows the EDS mapping of Sb and Cu elements for the single and multilayer barriers. These elemental images show the presence of Sb in the Cu layer in the case of the sample with the TaN monolayer after...
FIG. 3. (a) Cross section STEM micrograph of CeFe$_4$Sb$_{12}$/TaN/Cu and (b) Sb EDS profile after 6 h annealing at 400 °C (arrows: Sb-rich phases).

FIG. 4. (a) Cross section STEM micrograph of CeFe$_4$Sb$_{12}$/TaN-TaN/Cu and (b) Sb EDS profile after 6 h annealing at 400 °C.
treatment at 400 °C for 6 h and confirm the diffusion of this element through TaN. On the other hand, no Sb is found in the case of sample with the TaN-Ta-TaN multilayer after a treatment of 400 °C for 6 h, which put in evidence the effectiveness of the chosen multilayer structure. However, the XRD spectra showed the formation of the secondary phases, Cu$_3$Sb and FeSb$_2$, in the multilayer sample above 500 °C as observed also for the monolayer sample.

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<td>Sb</td>
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**FIG. 5.** STEM-EDS mapping of Sb (L$_{α_1}$) and Cu (L$_{α_1-2}$) after 6 h annealing at 400 °C (a) CeFe$_4$Sb$_{12}$/TaN/Cu and (b) CeFe$_4$Sb$_{12}$/TaN-Ta-TaN/Cu.

**FIG. 6.** Schematic illustration of single and multilayer barrier samples cross sections after annealing at 400 °C and 500 to 600 °C.
C. Discussion: TaN-Ta-TaN multilayer vs TaN single layer barriers

The efficiency of the multilayer barrier in comparison with the single layer barrier is schematically reported in Fig. 6. At 400°C, the performance of the TaN-Ta-TaN multilayer is evidenced as the Sb phase formation is not observed in the Cu layer. In the case of the single TaN layer, the Sb diffused to the Cu layer and formed Cu3Sb precipitates, which suggests in parallel a chemical decomposition of the skutterudite. The efficiency of TaN single layers vs TaN-Ta-TaN multilayers as diffusion barriers in the CMOS technology process was investigated previously, where copper is used for electrical interconnection.52 It has been reported that the efficiency of the TaN thin film as the diffusion barrier depends on its microstructure.5,7,53 In the reactive sputtering process, TaN layers are grown with a characteristic columnar structure perpendicular to the substrate, associated with intercolumn voids.34,55 The diffusion mechanism may be assumed to be simultaneously governed by the copper migration through the TaN-intercolumn voids and grain boundaries. In our previous study, we showed that a TaN/Ta/TaN multilayer is more efficient against copper diffusion than a TaN single layer.52 The copper diffusion is stopped at the Ta/TaN interface, where a discontinuity in the columnar growth is observed. In the present study, we show that the critical point is the diffusion of Sb from the skutterudite material to the copper surface layer. The formation of Sb-rich phases in the copper layer is observed when a TaN single layer is used as a diffusion barrier. In the case of the TaN-Ta/TaN multilayer barrier, Sb diffusion is not observed and seems to be impeded by the multilayer stacking. Based on our previous study, the Sb diffusion mechanism may be assumed to be mainly governed by its migration through the TaN-intercolumn voids. The multilayer structure promotes the barrier efficiency that is assumed to be due to the discontinuity of the columnar growth and the creation of additional layer interfaces. These results are supported by Sb concentration profiles and mappings which are clear evidence for the higher efficiency of the multilayer barrier.

At temperatures higher than 500°C, the Sb diffusion is not prevented neither by the single layer nor the multilayer barriers, which results in the formation of FeSb2 and Cu3Sb precipitates. Degradation of the thermoelectric properties of the skutterudite and of the skutterudite/copper electrical contact is, therefore, expected.

IV. CONCLUSION

The efficiency of TaN single layer and TaN-Ta-TaN multilayer diffusion barriers in CeFe3Sb2/Cu joining has been studied after annealing at 400°C and 500–600°C for 6 h in secondary vacuum. Regarding the single TaN layer, Sb diffusion from the skutterudite substrate to the copper layer is observed at 400°C. Further elemental diffusion is evidenced for higher temperatures, from 500 to 600°C, with the additional formation of FeSb2 and Cu3Sb precipitates. For the TaN-Ta-TaN barrier, the microstructural study shows a similar behavior as for the TaN barrier in the 500–600°C temperature range. However, this multilayer structure is interestingly an efficient diffusion barrier for the temperature below 500°C. The diffusion of Sb to the Cu surface layer is then inhibited which points out the role of sublayer interfaces in the diffusion mechanisms.

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