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High- and Low-Energy Photoemission Study of Strongly Correlated Au-Ga-Ce Quasicrystal Approximants: Localized 4*f* Nature and Disorder Effects

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We have investigated the electronic structures of Ce-based 1/1 quasicrystal approximants $Au_{59.2}Ga_{25.7}Ce_{15.1}$ and $Au_{60.3}Ga_{26.1}Ce_{13.6}$ by hard X-ray photoemission (HAXPES) and high-resolution photoemission spectroscopy. The localized Ce 4*f* electronic states are revealed for both Au-Ga-Ce approximants. Moreover, disorders in the compounds notably affect their electronic states, which has been detected by the core-level HAXPES. Valence-band photoemission spectra show the slight spectral difference depending on the composition ratio, which can be explained by a rigid-band-like shift.

1. Introduction

Rare-earth-based compounds with partially filled 4f subshell show many intriguing quantum phenomena, such as magnetic ordering,^{1,2)} unconventional superconductivity,^{3,4)} heavy-Fermionic state^{5–8)} (rather itinerant 4f electronic state with enormously enhanced effective mass), and non-Fermiliquid behavior.^{9,10)} The strongly correlated 4f electronic states in the compounds are determined by the hybridizations between the originally localized 4f orbitals and itinerant conduction bands (*c*-*f* hybridization), which play the important roles for these quantum phenomena. Therefore, revealing the 4f electronic states is one of the fundamental issues for understanding of the mechanisms of the quantum phenomena and further developments of the functional rare-earth-based compounds.

Recently, fascinating electronic states have been reported for rare-earth-based quasicrystals (QCs) and approximants (ACs). Here, QC is a metallic alloy having an aperiodic structure with unconventional rotational symmetry forbidden to conventional crystals, and AC is a periodical metallic alloy with a similar local structure to that of QC. The unique non-Fermi-liquid behavior, which is robust against hydrostatic pressure in contrast to that in the crystalline compounds, has been observed in Au-Al-Yb QC,¹¹⁾ whereas the superconductivity has been found in Au-Ge-Yb AC.¹²⁾ Moreover, ferromagnetic transition has been observed in Au-Ga-(Gd,Tb) QCs.¹³⁾ Consequently, the 4*f* electronic states in QCs and

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Fig. 1. (Color online) (a) Shell structure of the Tsai-type cluster for Au_{60.3}Ga_{26.1}Ce_{13.6} (Refs.17, 18). Chemical disorder of Au/Ga ions at the M4 and M5 sites and positional disorder due to the orientationally disordered tetrahedron at the M7 site are noticed. (b) Same as (a) but for Au_{59.2}Ga_{25.7}Ce_{15.1} in which a Ce ion occupies the cluster center (Ce2 site) with 58.1 % occupancy, in addition to the similar chemical and positional disorders observed in (a).

ACs have attracted much attention.

Conventional crystalline Ce-based compounds exhibit the characteristic electronic states as mentioned above. Thus, the physical properties of Ce-based ACs have also been investigated so far. For Ag-In-Ce,¹⁴ Au-Ge-Ce,¹⁵ and Au-Al-Ce ACs,¹⁶ spin-glass transition has been observed. On the other hand, the presence of Kondo effect has been pointed out for

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Ag-In-Ce and Au-Al-Ce ACs.^{14,16)} Meanwhile, new Ce-based Tsai-type Au-Ga-Ce 1/1 ACs have been discovered.¹⁷⁾ The Au-Ga-Ce ACs have wide range of composition ratio with single-phase region and show the spin-glass transition for all composition ratio. In Au_{60.3}Ga_{26.1}Ce_{13.6}, the Ce ions are occupied only in the icosahedron sites (Figure 1(a)).^{17,18)} On the other hand, the injection of Ce ions into the cluster center has been confirmed in Au_{59.2}Ga_{25.7}Ce_{15.1} where the Ce ions are occupied in both icosahedron sites and cluster center sites (Figure 1(b)).^{17,18)} It should be noted that the chemical disorder by non-magnetic Au/Ga atom mixing, which is indispensable for obtaining the wide range of composition ratio, would be seen in these systems. In (Au_{0.6975}Ga_{0.3025})_{100-y}Ce_y, the electrical resistivity of Au_{60,3}Ga_{26,1}Ce_{13,6} hardly depends on temperature, but slight $-\log T$ -like temperature dependence gradually develops with increasing y.¹⁷⁾ This $-\log T$ like dependence is prominent for the end composition ratio Au_{59,2}Ga_{25,7}Ce_{15,1}, implying the possibility of Kondo effect in Au_{59.2}Ga_{25.7}Ce_{15.1}. In order to clarify the origins of such macroscopic properties, the element-selective electronic structure should be clarified.

To reveal the Ce 4f electronic states in the Au-Ga-Ce ACs, we have performed core-level and valence-band hard Xray photoemission spectroscopy (HAXPES). Note that HAX-PES has an advantage in probing the bulk electronic states compared with conventional photoemission spectroscopy at relatively low-energy excitations.^{19,20)} We have revealed the highly localized 4f electronic states in the Au-Ga-Ce ACs. Moreover, we have observed unexpected spectral broadening in the Ce 3d core-level HAXPES spectra of Au-Ga-Ce ACs. From the comparison of the spectra between the Au-Ga-Ce ACs and Tsai-type AC Cd₆Ce^{21,22)} (more accurate stoichiometry composition ratio as Cd₃₇Ce₆, but hereafter denoted as Cd_6Ce for simplicity), we have revealed that the observed broadening is mainly caused by the effect of the disorders. On the other hand, it is known that a so called "psudogap"like structure is often seen in the density of states near the Fermi level (E_F) for QCs and ACs.^{23–25)} We have also performed valence-band HAXPES measurement and the highresolution photoemission spectroscopy (PES). We have observed the "pseudogap"-like structure and the compositionratio dependence of the spectra.

2. Experimental details

Polycrystalline samples of Au-Ga-Ce ACs Au_{59.2}Ga_{25.7}Ce_{15.1} and Au_{60.3}Ga_{26.1}Ce_{13.6} were prepared by the arc-melting method.¹⁷⁾ The HAXPES measurements of Au-Ga-Ce ACs were performed at BL19LXU in SPring-8 with an MBS A1-HE hemispherical analyzer.²⁶⁾ We also carried out the Ce 3d core-level HAXPES of single-crystalline AC Cd₆Ce to discuss the effect of the disorders. The photon energy for HAXPES was set to 7.9 keV. The overall energy resolution was set to 500 meV for the Ce 3d core-level measurements and 200 meV for the Au 4f core-level and valence-band measurements. For the valence-band HAXPES of Au_{59.2}Ga_{25.7}Ce_{15.1}, we changed the linear polarization of the incident hard X-ray from the horizontal to vertical directions. The degree of linear polarization (P_L) was estimated as -0.94, which corresponds to the linear polarization components along the horizontal and vertical directions of 3 and 97 %, respectively. The high-energy-resolution PES

measurements of Au-Ga-Ce ACs were performed using a SCIENTA SES-2002 hemispherical analyzer with an MBS T-1 discharge lamp.^{27,28)} The overall energy resolution was set to 11 meV with the Xe I resonance line (8.4 eV). The temperature was set to 7 K for HAXPES and 11 K for high-resolution PES. The clean surfaces were obtained by *in-situ* fracturing at the measuring temperatures for all measurements.

3. Results and discussions

3.1 Localized Ce 4f electronic states

Figure 2(a) shows the Ce 3d core-level HAXPES spectra of Au_{59,2}Ga_{25.7}Ce_{15.1} and Au_{60,3}Ga_{26.1}Ce_{13.6}. Both spectra are mutually similar each other, where main peaks are seen at 884 and 902 eV in the spectra. The former (latter) is ascribed to the Ce $3d_{5/2}$ ($3d_{3/2}$) excitations, mainly corresponding to the $3d^94f^1$ final states. Slight shoulder structures are seen at 878 and 896 eV, about 6 eV lower than the main peaks. These are predominantly due to the $3d^94f^2$ final-states contributions, which originate from the *c*-*f* hybridizations between the Ce 4f orbitals and the valence bands as seen for many Ce compounds. The nearly identical spectral shape between these compounds indicates that the composition ratio dependence of 4f electronic states are negligible in the Au-Ga-Ce ACs.

For comparison, the Ce 3d core-level HAXPES spectra of CeRu₂Ge₂ and CeRu₂Si₂²⁹⁾ are also shown in Figure 2(a). Note that the Ce 4f states in CeRu₂Ge₂ are localized and magnetically ordered at low temperatures,²⁾ while CeRu₂Si₂ is known as a typical heavy Fermion system with the specific heat coefficient $\gamma \sim 350 \text{ mJ/(K^2mol)}$ in which the 4f electrons have more or less itinerant nature due to the Kondo effects.^{6–8)} The shoulder structures at 878 and 896 eV ascribed to the $3d^94f^2$ final states are clearly seen in both spectra of the crystalline compounds. The $3d^94f^2$ final-state contributions are definitely weaker for AC Au_{59,2}Ga_{25,7}Ce_{15,1} and Au_{60,3}Ga_{26,1}Ce_{13,6} than for CeRu₂Ge₂ and CeRu₂Si₂. In addition, a small but clear peak at ~914 eV seen in the spectrum of the heavy Fermion system CeRu₂Si₂ seems to be absent in the spectra of the Au-Ga-Ce ACs as well as that of CeRu₂Ge₂. The peak at ~914 eV is due to the $3d^94f^0$ final state, which reflects the effect of itinerant behavior of the 4f electrons. Instead, a broad hump structure is observed around 916 eV in the spectra of Au_{59,2}Ga_{25,7}Ce_{15,1} and Au_{60,3}Ga_{26,1}Ce_{13,6} with splitting energy of 13.3 eV from the Ce $3d_{3/2}$ main peak. As shown in Figure 2(b), the similar hump structures are also seen in the Ga 2p core-level HAXPES spectra of Au_{59.2}Ga_{15.7}Ce_{15.1} at 13.3 eV higher than the main peaks. These spectral weight are independent on element to be excited, being ascribed to the plasmon satellite. Therefore, we conclude that the $3d^94f^0$ final-state contribution is absent for the Au-Ga-Ce ACs. These spectral features, the relatively weaker $3d^94f^2$ contributions than even those for CeRu₂Ge₂ and the negligible $3d^94f^0$ spectral weight for the Au-Ga-Ce ACs, indicate that the 4f electrons are highly localized in the Au-Ga-Ce ACs. Although possible itinerant 4f nature due to the Kondo effect could be originally expected for Au_{59.2}Ga_{25.7}Ce_{15.1}, the localized 4f electronic states in both Au-Ga-Ce ACs have been revealed by the Ce 3d core-level HAXPES.



Fig. 2. (Color online) (a) Ce 3*d* core-level HAXPES spectra of $Au_{59,2}Ga_{25,7}Ce_{15,1}$, $Au_{60,3}Ga_{26,1}Ce_{13,6}$, Cd_6Ce , $CeRu_2Ge_2$, and $CeRu_2Ge_2$,²⁹⁾ where the linear background has been subtracted from the raw spectra of $Au_{59,2}Ga_{25,7}Ce_{15,1}$, $Au_{60,3}Ga_{26,1}Ce_{13,6}$, and Cd_6Ce . The black dashed lines indicate the binding energies corresponding to the $3d^94f^0$ and $3d^94f^2$ final-state contributions. The black markers indicate the plasmon satellite structures, which can be judged from the other core-level spectra of the other elements in (b) and (c). (b) Ga 2*p* core-level HAXPES spectrum of $Au_{59,2}Ga_{25,7}Ce_{15,1}$. The black dashed line shows the plasmon satellite located at the binding energy 13.3 eV higher than the Ga $2p_{3/2}$ main peak. (c) Cd 3*d* core-level HAXPES spectrum of Cd₆Ce, in which the plasmon satellites are observed at the binding energies 9.8 eV higher than the main peaks.

3.2 Effect of the disorders

It is known that the two main peaks at 884 and 902 eV in the Ce 3d core-level photoemission spectra are formed by the ionic-like on-site $3d^94f^1$ final-state multiplet structure which can become unclear when the c-f hybridizations become stronger.^{30–32)} As discussed above, the 4f states are localized for both Au-Ga-Ce ACs. However, the $3d^94f^1$ finalstate multiplet structures seem to be smeared out in the spectra of the Au-Ga-Ce ACs. In addition, the two main peaks are much broader than those for CeRu₂Si₂ and CeRu₂Ge₂. To clarify the origins of the peak broadening for the Au-Ga-Ce ACs, we have measured the Ce 3d core-level HAXPES spectrum of Tsai-type AC Cd₆Ce with no chemical and almost negligible positional disorders, $^{21,22)}$ as shown in Figure 2(a). The clear $3d^94f^1$ multiplet structures are seen in the Ce 3dmain peaks (clearer in the $3d_{3/2}$ peaks centered at 902 eV) for AC Cd₆Ce. The Ce 4f electronic states are found to be localized also for AC Cd₆Ce, of which the $3d^94f^2$ final-state spectral weight is weaker than that for CeRu₂Ge₂ and the $3d^94f^0$ peaks are negligible. Broad peaks centered at 894 and 912 eV in the Ce 3d core-level spectra of AC Cd₆Ce, being 9.8 eV higher than the main $3d^94f^1$ peaks, are ascribed to the plasmon excitations which are also observed in the Cd 3dcore-level spectrum as shown in Figure 2(c). From our finding about the localized nature of the 4f states for all ACs surveyed here and the fact that the chemical and positional disorders (their details are explained later) are seen for the Au-Ga-Ce ACs in contrast to AC Cd_6Ce ,^{21,22)} we conclude that the remarkable broadening of the Ce 3d core-level photoemission spectra of the Au-Ga-Ce ACs originates (at least, partly) from the disorders. Note that the Ce 3d main peaks of Au-Ga-Ce ACs are also broader than those of polycrystalline CePdSn with localized Ce 4f system,³³⁾ where the multiplet structures are seen in the Ce $3d_{3/2}$ peak for CePdSn.³⁴⁾ Thus, the peak broadening seen for the Au-Ga-Ce ACs is not simply explained by the effect of their polycrystalline structures.

We focus on the two main peaks in the Ce 3d spectra of Au-Ga-Ce ACs and AC Cd₆Ce corresponding to the $3d^94f^1$ contributions. Figure 3(a) shows the Ce $3d_{3/2}$ and Ce $3d_{5/2}$ peaks of Au-Ga-Ce ACs and AC Cd₆Ce. In AC Cd₆Ce, the $3d^94f^1$ final-state multiplet structures are clearly observed as mentioned above. In order to estimate how much the Ce 3dcore-level HAXPES spectra of the Au-Ga-Ce ACs are broadened beyond the life-time and instrumental broadenings, we have broadened the Ce 3d spectrum of AC Cd₆Ce to roughly fit the spectra of the Au-Ga-Ce ACs. Figure 3(b) shows the comparison of the spectra after the broadening by the Gaussian with the full width at half maximum (FWHM) of 2.2 eV for the spectrum of AC Cd₆Ce. This value is much larger than that of the instrumental broadening of 0.5 eV, reflecting the effect of the disorders in the Au-Ga-Ce ACs as well as their polycrystalline structures. In order to further reveal the effect on the other sites, we have measured the Au 4fcore-level HAXPES spectra of $Au_{59.2}Ga_{25.7}Ce_{15.1}$ and the Au metal evaporated on the sample surface, as shown in Figure 3(c). The Au 4f peaks of Au_{59,2}Ga_{25,7}Ce_{15,1} are shifted by



Fig. 3. (Color online) (a) Enlarged view of the Ce 3*d* core-level photoemission spectra of AC Au_{59.2}Ga_{25.7}Ce_{15.1}, Au_{60.3}Ga_{26.1}Ce_{13.6}, and Cd₆Ce focused on the main peaks. The vertical black lines indicate the $3d^94f^1$ final-state multiplet components for Cd₆Ce. (b) Comparison of the Ce $3d_{5/2}$ main peaks after subtracting the Shirley-type backgrounds among the ACs. For Cd₆Ce, broadened spectra by a Gaussian with the full width at half maximum of 2.2 eV to roughly fit the widths of the spectra of the Au-Ga-Ce ACs. (c) Au 4f core-level HAXPES spectra of Au_{59.2}Ga_{25.7}Ce_{15.1} and Au metal. (d) Comparison of the Au $4f_{7/2}$ HAXPES spectra between AC Au_{59.2}Ga_{25.7}Ce_{15.1} and Au metal where the spectrum of Au_{59.2}Ga_{25.7}Ce_{15.1} is shifted by -0.49 eV.

0.49 eV to higher binding energy relative to those of the polycrystalline Au metal. To discuss the difference of the peak width, the Au 4*f* spectrum of Au_{59,2}Ga_{25,7}Ce_{15,1} is shifted by -0.49 eV and compared with that of the Au metal as shown in Figure 3(d). We have confirmed that the Au 4*f* peaks of Au_{59,2}Ga_{25,7}Ce_{15,1} is broader than that of the Au metal as seen in the comparison of the Ce 3*d* spectra between the Au-Ga-Ce and Cd-Ce ACs. However, the broadening of the Ce 3*d* peaks is larger than that of the Au 4*f* peaks in Au_{59,2}Ga_{25,7}Ce_{15,1} for which the broadening with FWHM of 0.24 eV (not shown in the figure) has been estimated. These results indicate that the disorders affect the overall electronic states of the Au-Ga-Ce ACs although the effect on the Au sites is relatively smaller than that on the Ce sites.

The core-level peak broadening observed for the Ce and Au sites is ascribed to inhomogeneity of the core-level energy depending on site. Namely, the binding energy on each site would be deviated from the mean binding energy. Since the observed core-level photoemission spectra are formed by the superposition of the spectra from all sites, the deviation of the binding energy reflects the broadening effects. In general, the shift of the core-level binding energy ΔE can be given by the following formula:³⁵⁾

$$\Delta E = \Delta \mu + K \Delta Q + \Delta V_M - \Delta E_R \tag{1}$$

where $\Delta \mu$ denotes the change in the chemical potential, K and ΔQ stand for the Coulomb coupling constant between the valence and core electrons, and the change in the number of valence electrons on the atom considered, respectively, ΔV_M represents the change in Madelung potential, and ΔE_R denotes the change in the extra-atomic relaxation energy which is important for metallic atoms/ions.³⁶⁾ It should be noted that the effect of the polycrystalline structure is not included in Eq. (1). The first term $\Delta \mu$ can be ruled out from the origins of the shift leading to the broadening since we discuss the site dependence within the same compound. The positional and chemical disorders have been observed on M4, M5, and M7 sites of the Au-Ga-Ce ACs (Figure 1).¹⁷⁾ From these disorders, the local environment and Madelung potential are different at each Ce site in the Au-Ga-Ce ACs. It should be noted that the mutually similar Ce 3d core-level HAXPES spectra of Au-Ga-Ce ACs indicate that the effect of Ce injection into the icosahedron sites is minor compared with the effect of the other disorders or this effect is too small to detect within our experimental conditions. Meanwhile, the effects of the second and fourth terms ($K\Delta Q$ and ΔE_R) would be minor in the Ce 3d spectra, which is concluded from the comparison of the spectra with that of AC Cd_6Ce where the Ce 4f states are found to be localized with the Ce³⁺ configuration in all Ce-based ACs discussed above. Therefore, we can conclude that the broadening of the Ce 3*d* core-level photoemission spectra by the disorders is ascribed to ΔV_M . On the other hand, the relatively less broadening of the Au 4*f* core-level spectra (FWHM of 0.24 eV) would be due to a compensation by the extra-atomic relaxation (ΔE_R) yielded by the screening of the conduction electrons in the Au sites.

3.3 Valence-band photoemission spectroscopy

The polarization-dependent valence-band HAXPES is useful to reveal the orbital contribution in the valence-band electronic states.³⁷⁾ Figure 4(a) shows the polarization-dependent valence-band HAXPES spectra of Au_{59,2}Ga_{25,7}Ce_{15,1}. In both p- and s-polarization configurations, prominent spectral weights are observed in the binding energy region between 3 and 8 eV. In order to estimate the orbital contributions for this structure, the photoelectron intensity ratio (I_s/I_p) for the s-polarization (I_s) to the p-polarization (I_p) is also shown in Figure 4(a) whereas the calculated I_s/I_p values and photoionization cross-sections (relative to those for the Au 5d ortibals) for the orbitals forming the valence bands are listed in Table 1.^{38–41)} The experimental I_s/I_p closes to the calculated Au 5d value of I_s/I_p between 3 and 8 eV. This result as well as the fact that the Au 5d contribution is much predominant at the photoelectron kinetic energy (hv) of ~8 keV means that the pronounced structures between 3 and 8 eV originate from the Au 5d bands. Moreover, the experimental I_s/I_p in the vicinity of E_F is much larger than calculated value for s orbitals and comparable to that found in the Au metal,³⁷⁾ which indicates that the Au 5d orbital contributes to the density of states even around E_F .

Figure 4(b) shows the valence-band HAXPES spectra of Au_{59.2}Ga_{25.7}Ce_{15.1} and Au_{60.3}Ga_{26.1}Ce_{13.6} in the *p*polarization configuration, which are compared with the HAXPES spectrum of the Au metal evaporated on the sample surface where the evaporated Au layer is much thicker than the photoelectron probing depth. The spectral shape of both Au-Ga-Ce ACs is mutually similar including the pronounced structures between 3 and 8 eV. Therefore, the Au 5d contributions are also dominant in the valence-band HAX-PES spectrum of Au_{60.3}Ga_{26.1}Ce_{13.6}. The observed main Au 5d bands between 3 and 8 eV in the Au-Ga-Ce ACs are narrower than those of the Au metal where the top of the main Au 5d bands is located at \sim 2 eV. Actually, the surface color of the ACs is silver-like, which is consistent with the spectral feature with the top of the 5d valence bands are located at \sim 3 eV as shown in Fig. 4(b). A slight difference depending on the composition ratio is observed in the valence-band HAXPES spectra. The inset of Fig. 4(b) shows an enlarged view of the spectra around the pronounced structures in 7 eV for Au_{59.2}Ga_{25.7}Ce_{15.1} and Au_{60.3}Ga_{26.1}Ce_{13.6}. The structure of Au_{60,3}Ga_{26,1}Ce_{13,6} is shifted by about 60 meV to higher binding energy relative to that of Au_{59,2}Ga_{25,7}Ce_{15,1}. The similar slight shift is also observed around 4 eV (not shown in the figure).

To further investigate the composition-ratio dependence of the electronic structure, we have performed the high-resolution PES at $h\nu = 8.4$ eV. Figure 5(a) shows the high-resolution PES spectra of Au_{59.2}Ga_{25.7}Ce_{15.1} and Au_{60.3}Ga_{26.1}Ce_{13.6} near E_F . In both spectra, a slight reduction of the spectral weight toward E_F is observed whereas no peak structure is found. These reductions are reported as the



Fig. 4. (Color online) (a) Linear polarization dependence of the valenceband HAXPES spectra (bottom) and the photoelectron intensity ratio I_s/I_p (top) of Au_{59.2}Ga_{25.7}Ce_{15.1} after subtracting the Shirley-type backgrounds. The spectral weights at different polarization configurations have been normalized by the photon flux. (b) Valence-band HAXPES spectra of Au_{59.2}Ga_{25.7}Ce_{15.1}, Au_{60.3}Ga_{26.1}Ce_{13.6}, and Au metal in the *p*-polarization configuration. The inset shows the enlarged view of the spectra around 7 eV.

"pseudogap" in the previous PES studies.^{23–25)} The slightness of the reduction at E_F compared with that for the other ACs and QCs^{23–25)} could be caused by the disorders observed for the Au-Ga-Ce ACs when the spectral reduction at E_F is assumed to be intrinsic for QCs and ACs with less disorders. Meanwhile, the binding energy at which the spectral intensity starts to decrease is slightly different between the compounds. The spectral weight starts to decrease (toward E_F) at the binding energy of 0.28±0.03 eV in Au_{59,2}Ga_{25,7}Ce_{15,1}, which is defined here as $\Delta_{15,1}$ and has been estimated from the differential of the smoothed spectrum (not shown here). For Au_{60,1}Ga_{26,1}Ce_{13,6}, the binding energy $\Delta_{13,6}$ at which the intensity starts to decrease has been obtained as 0.35±0.03 eV.

Table I. Calculated photoelectron intensity ratio I_s/I_p and relative photoionization cross sections per electron σ for Au 5*d*, Au 6*s*, Ga 4*s*, Ga 4*p*, Ce 4*f*, and Ce 6*s* orbitals at the kinetic energy of 8 keV.^{38–41}

Orbital	Au 5 <i>d</i>	Au 6 <i>s</i>	Ga 4s	Ga 4 <i>p</i>	Ce 4f	Ce 6s
I_s/I_p	0.293	0.026	0.011	0.469	0.853	0.006
σ	1	0.39	0.43	0.06	0.03	0.16



Fig. 5. (Color online) (a) High-resolution valence-band PES spectra of $Au_{59,2}Ga_{25,7}Ce_{15,1}$ and $Au_{60,3}Ga_{26,1}Ce_{13,6}$ near E_F . The black lines indicate the binding energy at which the spectral intensity starts to decrease. (b) Schematic image of a rigid-band-like shift in the Au-Ga-Ce ACs.

It should be pointed out that the energy difference between $\Delta_{15.1}$ and $\Delta_{13.6}$ is comparable to the slight shift of the main Au 5*d* bands seen in the HAXPES spectra in Fig.4(b). In order to discuss the composition-ratio dependence, the number of valence electrons n_e is shown here. Note that the definition of n_e is the number of electrons which form the valence bands, *i.e.*, Au: $5d^{10},6s^1$, Ga: $4s^2,4p$,¹ and Ce: $6s^2,5d^1$. The n_e for both composition ratio is calculated as follows:

$$Au_{59,2}Ga_{25,7}Ce_{15,1}: n_e$$

$$= 11 \times 0.592 + 3 \times 0.257 + 3 \times 0.151 = 7.74$$
(2)

 $Au_{60.3}Ga_{26.1}Ce_{13.6}: n_e$

$$= 11 \times 0.603 + 3 \times 0.261 + 3 \times 0.136 = 7.82$$
 (3)

From Equations (2) and (3), n_e of Au_{60.3}Ga_{26.1}Ce_{13.6} is larger than that of Au_{59.2}Ga_{25.7}Ce_{15.1}. When it is assumed that a rigid-band-like shift arises, E_F of Au_{60.3}Ga_{26.1}Ce_{13.6} is shifted to the unoccupied side relative to that of Au_{59,2}Ga_{25,7}Ce_{15,1}. Figure 5(b) shows the schematic image of such a rigid-bandlike shift in Au-Ga-Ce ACs. $\Delta_{13.6}$ should be larger than $\Delta_{15.1}$ in the rigid-band-like shift when we consider the difference in n_e shown in Equations (2) and (3), which is consistent with the result of high-resolution PES. Moreover, the shift of Au 5d bands in the valence-band HAXPES spectra is also explained by the rigid-band-like shift. Thus, the composition-ratio dependence found in the valence-band PES is explained by the rigid-band-like shift. Namely, the additional Ce injection into the tetrahedron (M7) sites hardly affects the substantial electronic structure for the Au-Ga-Ce ACs. From our study, we also conclude that the slight $-\log T$ -like behavior in the resistivity of Au_{59,2}Ga_{25,7}Ce_{15,1}¹⁷⁾ does not originate from the Kondo effect although the mechanisms are unclear at present.

4. Summary

In summary, we have performed the HAXPES and high-resolution PES of AC Au_{59,2}Ga_{25.7}Ce_{15.1} and Au_{60,3}Ga_{26.1}Ce_{13.6}. The Ce 4*f* states are found to be highly localized in both Au-Ga-Ce ACs. The overall electronic structure is mutually similar irrespective of the Ce injection in the tetrahedron sites. We have also observed the unexpected broadening in the $3d^94f^1$ main peaks in the Ce 3*d* core-level photoemission spectra of both Au-Ga-Ce ACs. From the comparison of the spectra of AC Cd₆Ce, we conclude that this broadening is caused by the effect of the disorders in the Au-Ga-Ce ACs. The composition-ratio (substitution-ratio) dependence of the electronic state is found as the rigid-band-like shift.

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