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# Angle-Resolved and Resonant Photoemission Spectroscopy Study of the Fermi Surface Reconstruction in the Charge Density Wave Systems of CeTe<sub>2</sub> and PrTe<sub>2</sub>

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The electronic structures of a charge-density-wave (CDW) system  $RTe_2$  (R=Ce, Pr) have been investigated by employing angle-resolved photoemission spectroscopy (ARPES) and the first-principles band structure method. The R 4f hybridization peak ( $4f^nc^{m-1}$ ) in the R 4f PES spectrum is located deeper in PrTe<sub>2</sub> than in CeTe<sub>2</sub> and R 4f spectral intensity near  $E_F$  is much weaker in PrTe<sub>2</sub> than in CeTe<sub>2</sub> and R 4f spectral intensity near  $E_F$  is much weaker in PrTe<sub>2</sub> than in CeTe<sub>2</sub>, implying the importance of the hybridization between Ce 4f and Te(1) 5p electrons. For both CeTe<sub>2</sub> and PrTe<sub>2</sub>, the metallic states crossing the Fermi level ( $E_F$ ) are observed below the CDW transition temperature, indicating the existence of the partially ungapped Fermi surfaces (FSs). The zigzag features having the four-fold rotational symmetry are observed near the X point in the FS of CeTe<sub>2</sub>, but not in the FS of PrTe<sub>2</sub>. The tight-binding model calculations show that the zigzag FS features in CeTe<sub>2</sub> can be described as the CDW-induced FS reconstruction due to the  $4 \times 4$  CDW supercell structure. The effect of the linear dichroism is observed in ARPES, suggesting that the  $E_F$ -crossing states have mainly the in-plane orbital character. The photon-energy maps for the near- $E_F$  states exhibit the straight vertical dispersions for both CeTe<sub>2</sub> and PrTe<sub>2</sub>, demonstrating the dominant two-dimensional character in  $RTe_2$  (R=Ce, Pr).

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# I. INTRODUCTION

The charge-density-wave (CDW) formation is one of the most interesting phenomena in solid state physics.<sup>1–3</sup> Transition-metal (T) dichalcogenides of  $TX_2$  (X=S, Se) are the well-known CDW systems,<sup>4</sup> where T d electrons play an important role in determining the CDW phenomenon. The less-studied  $RTe_2$  (R: rare-earth ion) belongs to the *p*-electron CDW system, having a high CDW transition temperature of  $T_{CDW} \sim 1000 \text{ K}$ . <sup>5–8</sup> R Te<sub>2</sub> crystallizes in the quasi-two dimensional layered Cu<sub>2</sub>Sb-type tetragonal structure with two types of Te sites: Te(1)and Te(2). Te(1) atoms form the planar square sheets, which are sandwiched along the c axis by the corrugated double layers of R and Te(2) atoms.<sup>6,9</sup> Due to the underlying R-Te(2) layer, the two-dimensional (2D) unit cell of a Te(1) sheet is doubled  $(\sqrt{2} \times \sqrt{2})$ , resulting in a reduced Brillouin Zone (BZ) to a half of that of the Te(1) square lattice. Thereby the bands are folded into the reduced BZ to produce the shadow bands. In reality, the periodicity of the potential energy in the Te(1) layer depends on the interaction strength between Te(1) and R-Te(2) layers. Hence, the electronic structures of Te(1) sheets would keep the 2D nature of the planar square lattice if the interlayer interaction is small, while they would exhibit the three-dimensional (3D)-like nature if the interlayer interaction becomes strong.

In contrast to the triangular  $TX_2$  (X=S, Se) CDW systems,<sup>4</sup> the CDW states of  $RTe_2$  are not well explored yet. The ionic configuration of  $RTe_2$  is considered to be  $R^{3+}Te(2)^{2-}Te(1)^{1-}$ , which was confirmed experimentally for R=La, Ce.<sup>10</sup> So hole carriers are produced in Te(1) sheets. Then the square net of Te(1) are easily distorted by the Peierls-like mechanism<sup>11</sup> due to the partial filling. Band-structure calculations indicate that the CDW instability occurs due to the nesting between the Fermi surfaces (FSs) in Te(1) square sheets in the *ab* plane,<sup>5,9,12,13</sup> which was supported by experiment.<sup>6</sup> The 2 × 2 supercell structure in the form of a double herringbone pattern<sup>6</sup> was reported to be a candidate CDW structure for CeTe<sub>2</sub>, but the real CDW structure of CeTe<sub>2</sub> has not been confirmed either experimentally or theoretically yet. Further, some electronic structure calculations indicated that the 3D nature is stronger in CeTe<sub>2</sub> than in LaTe<sub>2</sub>.<sup>12-14</sup>

Angle-resolved photoemission spectroscopy (ARPES) is a very good experimental tool for studying the FS topology and the electronic structures of the CDW systems.<sup>4</sup> Despite the extensive ARPES studies for the similar CDW systems of  $RTe_3$ ,<sup>8,15,16</sup> there are not many ARPES reports on  $RTe_2$ .<sup>8,13,15,17,18</sup> Shin *et. al.*<sup>8</sup> reported that the FS topology of CeTe<sub>2</sub> in the  $k_x$ - $k_y$  plane is different from that of LaTe<sub>2</sub>,<sup>15</sup> and that the magnitude of the CDW gap  $\Delta_g$  varies around the FS with the minimum value of  $\geq 600$  meV, which is much larger than that found in another ARPES study ( $\Delta_g \approx 100$  meV).<sup>13,17</sup> Ito *et. al.*<sup>18</sup> argued the observed intensity modulation in the FS along the  $k_z$  axis to support the strong 3D nature in CeTe<sub>2</sub>. In contrast, the 3D nature is considered to be negligible in LaTe<sub>2</sub>.<sup>12,15</sup>

Hence the following issues remain to be resolved for  $R\text{Te}_2$ : (i) What are the modulated structures after the

CDW distortion? (ii) How different are the FS topology and the CDW states depending on R element? Namely, what is the role of R ions in the CDW distortion? (iii) How large is the effect of the band foldings on the FS of Te(1) sheets, arising from the interaction between Te(1)and R-Te(2) layers and the CDW supercell formation? We have resolved these questions by performing careful ARPES measurements for high-quality stoichiometric single crystals of  $RTe_2$  (R=Ce, Pr), and first-principles electronic structure calculations. CeTe<sub>2</sub> and PrTe<sub>2</sub> have very similar lattice constants in agreement with the Lanthan de contraction: a = 4.47 Å and c = 9.11 Å for  $CeTe_2$ ,<sup>12</sup> and a = 4.44 Å and c = 9.06 Å for  $PrTe_2$ .<sup>8</sup>.

### п. EXPERIMENTAL AND CALCULATIONAL DETAILS

High-quality  $RTe_2$  (R=Ce, Pr) single crystals were grown by using the self-fluxed Bridgeman method.<sup>19</sup> The quality and the orientation of the single crystal were checked by Laue patterns. ARPES measurements were carried out at the MERLIN 4.0.3. beamline at the Advanced Light Source (ALS). The ARPES endstation is equipped with a Scienta R8000 electron energy analyzer and a low temperature 6-axis sample manipulator cooled with an open-cycle He flow cryostat. The sample temperature can be cooled down to 6 K, but we kept the samples at  $\sim 35 K$  to improve their electrical conductivity. Single crystalline samples were cleaved in situ and measured under the pressure better than  $6 \times 10^{-11}$ Torr. The Fermi level  $(E_F)$  and the instrumental resolution of the system were determined from the Fermi edge spectrum of an evaporated Au metal. The photon energy  $(h\nu)$  corresponding to the  $\Gamma$  plane was determined from the photon energy scans for the Fermi-edge states at the normal emission.<sup>20</sup> For both  $CeTe_2$  and  $PrTe_2$ ,  $h\nu \approx 108 - 110 \text{ eV}$  turned out to be close to the  $\Gamma$  plane within the experimental uncertainty (see Fig. 8).

For band calculations, we have employed the *ab-initio* full-potential linearized augmented (FLAPW) band calculations implemented in the Wien2k.<sup>21</sup> The generalized gradient approximation (GGA) was utilized for the exchange-correlation interaction, and the spin-orbit interaction was taken into account in the second variation manner.

### III. **RESULTS AND DISCUSSION**

Figures 1(a) and (b) show the optical microscopy images of the cleaves surfaces of  $RTe_2$  (R=Ce, Pr) single crystals, taken after ARPES measurements. These optical images show that the ARPES data presented in this paper were obtained from the flat region because the beam size employed in ARPES measurements is less than  $\sim 40 \ \mu m$ . Figure 1(c) shows the angle-integrated valenceband resonant photoemission spectroscopy (RPES) spec2



FIG. 1: (Color online) Optical Microscopy Images and Angleintegrated PES Spectra. (a), (b) Optical microscopy images of the cleaves surfaces of  $RTe_2$  (R=Ce, Pr) single crystals, taken after ARPES measurements. (c) Angle-integrated valence-band  $R 4d \rightarrow 4f$  resonant photoemission spectroscopy (RPES) spectra of  $CeTe_2$ . Those obtained at the Ce 4f on-resonance energy ( $h\nu$ =121 eV: red) and off the resonance energy ( $h\nu$ =110 eV: blue) are compared. (d) Comparison of the angle-integrated  $R \ 4f$  on-resonance valence-band PES spectra of  $RTe_2$  (R=Ce, Pr). They are measured at the Ce 4f resonance energy ( $h\nu$ =121 eV) and Pr 4f resonance energy ( $h\nu$ =124 eV), respectively. (e) Comparison of the Te 4d core-level PES spectra of  $RTe_2$  (R=Ce, Pr), measured at  $h\nu = 110$  eV. All the PES data were obtained at T=35 K.

tra of CeTe<sub>2</sub>, obtained at the Ce  $4d \rightarrow 4f$  on-resonance  $(h\nu=121 \text{ eV})$  and off the 4f resonance  $(h\nu=110 \text{ eV})$ , respectively. The resonance enhancement of the Ce 4f (between  $E_F$  and  $\sim -5 \text{ eV}$ ) and 5p (between  $\sim -18 \text{ eV}$  and  $\sim -22$  eV) states is observed clearly. A similar resonance behavior was observed in  $PrTe_2$  (not shown in this paper).

Figure 1(d) compares the angle-integrated valenceband RPES spectra of  $RTe_2$  (R=Ce, Pr), obtained at the  $R 4d \rightarrow 4f$  on-resonance energies. This figure reveals that both Ce and Pr 4f PES spectra exhibit the double-peak structures, one near  $E_F$  and the other well below  $E_F$ . The *R* 4*f* peaks well below  $E_F$  correspond to the trivalent  $R \ 4f^n c^m \rightarrow 4f^{n-1}c^m$  transition (c: conduction-band electron, m=3). The 4f peaks near  $E_{\rm F}$  are known to arise from the hybridization between R 4f and conduction-band electrons, corresponding to the  $4f^n c^{m-1}$  final states.<sup>24–26</sup> From R=Ce to Pr, the peak positions of both the  $4f^{n-1}c^m$  and  $4f^nc^{m-1}$  final states shift to higher BE's. The position of the near- $E_{\rm F}$  hybridization peak  $(4f^n c^{m-1})$  in  $R={\rm Pr}$  (BE ~ 1.5 eV) is deeper than in R=Ce (BE ~ 0.8 eV), and its relative intensity is much weaker than in R=Ce, resulting in the much weaker 4f spectral intensity near  $E_{\rm F}$ . Such trends are similar to those of metallic rare-earth transition-metal compounds.<sup>27,28</sup> Since the metallic electronic states near  $E_F$  in the normal phase (non-CDW





FIG. 2: (Color online) The FS and CE maps of  $RTe_2$  (R=Ce, Pr). (a) The constant initial-state energy (CE) maps of the ARPES intensity in the momentum space at increasing binding energies (BE's) for CeTe<sub>2</sub> from BE=0 eV (E<sub>F</sub>) to BE  $\sim 0.5 \text{ eV}$  (BE= $|E_i|$ ). Dotted lines denote the BZ. ARPES data were measured at T=35 K and with  $h\nu = 110$  eV. (b) Similarly for PrTe<sub>2</sub>. (c) The FS map of CeTe<sub>2</sub> over the wider BZ region. (d) Similarly for PrTe<sub>2</sub>. In (c) and (d),  $\frac{\pi}{a}(0,0)$ and  $\frac{\pi}{a}(2,0)$  correspond to the two  $\Gamma$  points in the first and second BZs, respectively. (e) A schematic sketch of the zigzag features near X points, drawn on top of the measured FS for CeTe<sub>2</sub>. The ring at the center with an arrow represents the four-fold rotational symmetry of zigzag features.

phase) play an important role in the CDW distortion, the observed trends from CeTe<sub>2</sub> to PrTe<sub>2</sub> indicate that the contribution from Ce 4f electrons to the CDW formation is stronger than that from Pr 4f electrons. Note, however, that the R 4f spectral intensity at E<sub>F</sub> is almost negligible for both R=Ce and Pr. Hence, even if the contribution of R 4f electrons to the CDW exists, it would be indirect, *i. e.*, it will contribute through the R4f-Te(1) 5p hybridization. Indeed this effect is observed in this work, which will be discussed under Fig. 2(c).

Figure 1(e) compares the Te 4d core-level PES spectra of  $RTe_2$  (R=Ce, Pr). Their line shapes are similar to each other except that the low-BE shoulder is more pronounced in PrTe<sub>2</sub> than in CeTe<sub>2</sub>. Their lineshapes indicate that there are several components, consistent with the presence of two different Te sites of Te(1) and Te(2) as well as the *bulk* and *surface* components for each site. We have found that the the lineshapes of the Te 4d corelevel spectra of  $RTe_2$  do not change much for different spots in the cleaved surfaces.

Figure 2 shows the FS and constant initial-state energy (CE) maps of  $RTe_2$  (R=Ce, Pr). These maps correspond to the constant bind ing[-energy (BE) slices of the angle maps between  $E_F$  ( $E_F \equiv 0 \text{ eV}$ ) and BE=0.5 eV ( $BE=|E_i|$ , where  $E_i$  denotes the initial-state energy). All the FS

FIG. 3: (Color online) The CDW distorted unit cells and the corresponding FSs for CeTe<sub>2</sub>. (a) The CDW supercell structure composed of a candidate  $2 \times 2$  unit cell (yellow-colored cell) in the Te(1) sheet. Dumbells represent the dimerized Te(1) atoms in the *ab* plane. (b) Similarly for a candidate  $4 \times 4$  unit cell (yellow-colored cell) in the Te(1) sheet. (c) and (d) show the calculated FSs using the tight-binding models for the CDW supercell structures shown in (a) and (b), respectively. Those FSs are unfolded into the first BZ of the non-CDW phase (depicted as square boxes in (c) and (d)).

and CE maps in this figure were obtained by integrating the spectral intensity of  $E_i \pm 40$  meV for each  $E_i$ . This figure shows that the FS map of PrTe<sub>2</sub> is close to that of an ideal Te(1) square net, while that of CeTe<sub>2</sub> has the extra features in addition to that of the ideal Te(1) square net, which will be described in detail in Fig. 2(c). Nevertheless, the shapes, sizes, and energy-dependent behavior of the CE maps of CeTe<sub>2</sub> and PrTe<sub>2</sub> are similar to each other, exhibiting the four-fold rotational symmetry. The energy-dependent behavior is consistent with that of the calculated CE maps in Fig. 2 of Ref.<sup>[13]</sup>.

The FSs of  $RTe_2$  (R=Ce, Pr), measured over the wide angle regions, are shown in Fig. 2(c) and (d). The CDW gaps are found to open only partially in some part of the FS, which is consistent with the semimetallic nature of RTe<sub>2</sub>.<sup>17,29,30</sup> The overall similarity of the FS/CE maps of CeTe<sub>2</sub> and PrTe<sub>2</sub>, as well as those of LaTe<sub>2</sub>,<sup>15</sup> implies that the contribution of the R 4f (R=Ce, Pr) electrons to the states at  $E_F$  is minor. Nevertheless, Figs. 2(c) and (d) show the crucial differences between  $CeTe_2$  and PrTe<sub>2</sub>. The zigzag shape features are observed near the X points in the FS map of CeTe<sub>2</sub>, while these features are absent in PrTe<sub>2</sub>.<sup>31</sup> To help to identify the zigzag features in  $CeTe_2$ , a schematic sketch of the zigzag features near X points is shown on top of the measured FS for  $CeTe_2$ in Fig. 2(e). This finding suggests that the origin of these extra features in CeTe<sub>2</sub> is the hybridization between Ce 4f and Te(1) 5p states because they are not observed in  $PrTe_2$ . We also think that these X-point features in CeTe<sub>2</sub> arise from the CDW-induced FS reconstruction because the calculated FS maps for the non-CDW phases



FIG. 4: (Color online) MDCs for  $R\text{Te}_2$  (R=Ce, Pr) along the (100) direction. (a) The MDC stack for CeTe<sub>2</sub> for cuts parallel to (100), obtained at T=35 K and  $h\nu = 110$  eV. In the MDC stack, each MDC curve is shifted vertically by a constant. The green arrows on the MDC curve represent the  $\vec{k}$ -points, where the states cross  $E_{\rm F}$  near the X point. The inset shows the FS and the white dotted lines on the FS represent the three symmetry lines in the MDC cuts. (b) Similarly for PrTe<sub>2</sub>.

do not reveal these zigzag features near X (see Fig. 7).

The zigzag structures near X show the four-fold symmetry, but not the mirror symmetry. These features in CeTe<sub>2</sub> are reminiscent of those in LaTe<sub>2</sub>.<sup>15</sup> But the zigzag features in LaTe<sub>2</sub> were detected for the CE map with deeper BE of ~ 100 - 160 meV. As mentioned above, the zigzag features seem to arise from the CDW supercell structure in Te(1) sheets, which would yield the additional band foldings. Note, however, that this kind of zigzag features in the FS cannot be obtained for a double herringbone-type  $2 \times 2$  supercell structure of CeTe<sub>2</sub>.<sup>6</sup>

Figure 3 shows the reported  $2 \times 2$  and the assumed  $4 \times 4$  supercell structures for CeTe<sub>2</sub>, and the corresponding FSs that are obtained from the tight-binding model calculations.<sup>32</sup> In these calculations, the effect of the Ce 4 f orbitals was taken into account in the effective Te-Te hopping parameters of the tight-binding model Hamiltonian. The FS for the  $2 \times 2$  structure in Fig. 3(a) does not exhibit the zigzag features near X. In contrast, new features are observed near X in the FS for the  $4 \times 4$ structure in Fig. 3(b), which are analogous to those in the measured FS. These findings indicate that the zigzag structures near X are not compatible with the reported  $2 \times 2$  supercell structure<sup>6</sup> and that the CDW states in  $CeTe_2$  might be formed as the  $4 \times 4$  supercell structure rather than the  $2 \times 2$  structure. Note that the zigzag features in the calculated FS for the  $4 \times 4$  supercell structure result from the special combination of the Peierls distortions of the crossing 1D Te chains in the 2D Te square net, as shown in Fig. 3(b). Further experimental and theoretical studies are demanded to clarify this issue.



FIG. 5: (Color online) Near-E<sub>F</sub> EDCs for  $RTe_2$  (R=Ce, Pr). (a) EDC stacks for CeTe<sub>2</sub> for a cut through the X point, *i.e.*, the EDC stacks along the M-X-M direction, obtained at T=35 K and  $h\nu = 110$  eV. The solid lines, superposed on the EDCs, provide the guidelines for the EDC lineshapes. The EDC in blue corresponds to that at X. The inset at the top shows the FS of CeTe<sub>2</sub>. A and B denote the  $k_y$  points where the top and bottom EDCs are obtained. The EDCs that have stronger intensities than others are denoted in red color. (b) Similarly for PrTe<sub>2</sub>.

Figure 4 shows the momentum distribution curves (MDCs) for the cuts parallel to the (100) direction in the FS maps of  $RTe_2$  (R=Ce, Pr), as shown in the insets. In the MDC stacks, each MDC curve is shifted vertically by a constant. These MDCs correspond to the intensity profiles of the states at  $E_F$  versus  $k_x$  for a fixed  $k_y$  value. Hence the peaks in MDCs represent the states that cross  $E_{\rm F}$  at the corresponding  $\vec{k}$ -points. The MDC curves, depicted by thick red lines, represent those along M-X-M, X- $\Gamma$ -X, and M-X-M, respectively. The MDC peaks in  $CeTe_2$  reveal the near-X features very well, as discussed in Fig. 2(c). In order to identify the peak positions due to the zigzag shape near X in CeTe<sub>2</sub>, green arrows are added on the MDC along M-X-M (bottom cut) of  $CeTe_2$ . In contrast, no such MDC peaks are observed in  $PrTe_2$  around X points. The differences in the MDCs between CeTe<sub>2</sub> and PrTe<sub>2</sub> suggest that the hybridization between Ce 4f and Te(1) 5p states plays an important role in determining the CDW formation in  $CeTe_2$  and gives rise to the extra zigzag-shape FS near X in CeTe<sub>2</sub>.

The opening of the  $\vec{k}$ -dependent CDW gaps in the FS would be observed in the energy distribution curves (EDC's) as the vanishing spectral intensity near  $E_F$ . Figure 5 shows the near- $E_F$  EDCs for the *M*-*X*-*M* cut in the FS maps of *R*Te<sub>2</sub> (*R*=Ce, Pr). In the EDC stacks, each EDC curve is shifted vertically by a constant. Due to the very low spectral intensity near  $E_F$ in *R*Te<sub>2</sub>, the Fermi-level crossings of the near- $E_F$  metallic states are not observed clearly in the EDCs for *R*Te<sub>2</sub> (*R*=Ce, Pr). Nevertheless, the intensity modulations are



FIG. 6: (Color online) Linear Dichroism (LD) in ARPES for PrTe<sub>2</sub> along  $M - \Gamma - M$ . (a)-(c) The measured ARPES image plots of PrTe<sub>2</sub>, along  $M - \Gamma - M$  (in the 1st BZ), obtained with the linear horizontal (LH) and linear vertical (LV) polarizations, and the difference between the LH and LV ARPES, respectively, obtained at T=35 K and  $h\nu = 110$  eV. (d)-(f) The ARPES image plots near E<sub>F</sub>, corresponding to the wideenergy-range ARPES of (a)-(c), respectively. Here (c) and (f) represent the LD of the ARPES. The schematic drawings on the left show the configurations for LH and LV measurements.

very distinct between CeTe<sub>2</sub> and PrTe<sub>2</sub>. The evidence for the E<sub>F</sub>-crossings in CeTe<sub>2</sub>, corresponding to the zigzag FS features around X, is provided by the stronger intensity modulation in the EDCs for CeTe<sub>2</sub> than for PrTe<sub>2</sub>. To identify the stronger intensity modulation in CeTe<sub>2</sub> clearly, the EDCs having strong intensities in Fig. 5(a) are denoted in red color. In contrast to PrTe<sub>2</sub>, the strong intensity modulation in CeTe<sub>2</sub> yields the finite intensity at E<sub>F</sub> at several  $\vec{k}$  points, indicating no gap at the corresponding  $\vec{k}$  points. Figure 5 also reveals that the magnitude of the gap varies continuously along M-X-M (along the cut through the zigzag-shaped FS). Hence the EDCs shown in Fig. 5 support the findings of both the MDCs (Fig. 4) and the measured FSs (Fig. 2).

Figure 6 shows the measured ARPES image plots of  $\Pr Te_2$  along M- $\Gamma$ -M (in the 1st BZ), obtained with two different polarizations of LH (linear horizontal) and LV (linear vertical) polarizations. The schematic drawings on the left show the configurations for LH and LV measurements. In the LH configuration, the electric field vector  $\vec{E}$  of the incident photons is nearly perpendicular to the sample surface, while  $\vec{E}$  is parallel to the sample surface in the LV configuration. The upper and lower layers show the wide-energy-range and near- $E_F$  ARPES image plots, respectively. Figures 6(c) and (f) show the differences between LH and LV ARPES data, defined as linear dichroism (LD): LD=LH-LV.

The ARPES data for PrTe<sub>2</sub> in Fig. 6 reveal several things. (i) Many dispersive bands are observed clearly in ARPES. The large-energy-scale band structures of PrTe<sub>2</sub> in ARPES are very similar to those of CeTe<sub>2</sub>.<sup>13</sup> (ii) The



FIG. 7: (Color online) Calculated Band Structures of CeTe<sub>2</sub> and PrTe<sub>2</sub>. (a) The calculated band structures of CeTe<sub>2</sub>, where the Ce 4f electrons are treated as core electrons. The different colors denote the different character of the electronic states, such as Ce 4f, Ce 5d, Te(1) 5p, and Te(2) 5p states. The sizes of the circles for each line represent the contribution of each electronic character at different  $\vec{k}$  points. (b) Similarly for PrTe<sub>2</sub>. (c) The calculated FS of CeTe<sub>2</sub> in the tetragonal Brillouin Zone (BZ) of the non-CDW phase.  $\Gamma$ , X, and M represent  $\vec{k}$ =(0, 0, 0),  $\frac{2\pi}{a}(1/2, 0, 0)$ , and  $\frac{2\pi}{a}(1/2, 1/2, 0)$ , respectively. (d) Similarly for PrTe<sub>2</sub>. (e) Comparison of the calculated FSs of CeTe<sub>2</sub> and PrTe<sub>2</sub> in the  $\Gamma$  plane. Red and blue colors denote CeTe<sub>2</sub> and PrTe<sub>2</sub>, respectively, and violet color appears when red and blue colors overlap each other.

effect of LD, arising from the different polarizations is observed clearly. The states at  $\sim 1 \text{ eV}$  BE exhibit the stronger intensity with the LH polarization than with the LV polarization, while those at  $\sim 5 \text{ eV}$  BE exhibit the opposite trend, implying that the orbitals at  $\sim 1 \text{ eV}$ BE and at  $\sim 5 \text{ eV}$  are ordered mainly along the c axis and in the *ab* plane, respectively. On the other hand, the LD of the  $E_{\rm F}$ -crossing states is opposite to the LD of the  $\sim 1 \text{ eV BE}$  states. The E<sub>F</sub>-crossing bands exhibit the stronger intensity with the LV polarization than with the LH polarization, implying that the E<sub>F</sub>-crossing orbitals are ordered mainly in the ab plane than along the c axis. (iii) The band crossing through  $E_F$  is clearly observed in the near- $E_F$  ARPES image plots. These states are expected to produce the FSs at the corresponding  $\vec{k}_F$ values.

The E<sub>F</sub>-crossing positions, observed in ARPES, agree very well with those in the calculated bands for the non-CDW phase of RTe<sub>2</sub> (see Figs. 7), which have mainly Te(1) 5*p* character.<sup>12</sup> In the near-E<sub>F</sub> ARPES for CeTe<sub>2</sub>,<sup>13</sup> two crossing bands near  $\Gamma$  (~ 0.2  $\Gamma M$ ) and M (~ 0.7  $\Gamma M$ ) were observed, which produced the inner bright FS and outer dim FS, respectively, along  $\Gamma$ -M in the first BZ. The much stronger spectral weight of the former band (~ 0.2  $\Gamma M$ ), as compared to the latter band (~ 0.7  $\Gamma M$ ), was explained by the main (former) versus the shadow (latter) band nature. In contrast, in the near- $E_F$  ARPES for PrTe<sub>2</sub>, only the band near  $\Gamma$  (~ 0.2  $\Gamma M$ ) is observed in Fig. 6(d)-(f). This feature is consistent with the almost vanishing outer FS of PrTe<sub>2</sub> along  $\Gamma$ -M in Fig. 2(d), reflecting that the 3D interaction between Te(1) and R-Te(2) layers is weak.

Figures 7(a) and (b) show the calculated band structures of  $RTe_2$  (R=Ce, Pr), where R 4f electrons are treated as core electrons. Here, the different colors denote the different character of the electronic states, such as  $5p_{x+y}$  and  $5p_z$  states of Te(1) and Te(2), and R 5d. These figures reveal that the calculated band structures of  $CeTe_2$  and  $PrTe_2$  are very similar to each other, as expected. This is because Ce and Pr f electrons are treated as core electrons and so the R f electrons do not contribute to the electronic structures near  $E_F$ . As shown experimentally in Fig. 1(d), the Ce 4f states are expected to contribute more to the near- $E_F$  electronic structures than the  $\Pr 4f$  states. For  $\Pr Te_2$ , a very good agreement is found in the overall band structures between ARPES (Fig. 6) and calculations (Fig. 7). Especially, the calculational results of the dominant  $5p_{x+y}$  character in the ab plane near  $E_F$  and at ~ 5 eV BE are quite consistent with the observed LD in Fig. 6(c).

Figures 7(c) and (d) show the calculated FSs of  $RTe_2$  (R=Ce, Pr) in the tetragonal BZ of the non-CDW phase. Note that the calculated FSs for CeTe<sub>2</sub> and PrTe<sub>2</sub> are very similar to each other, as shown more clearly in Fig. 7(e). No extra features are observed near the X points in the calculated FSs of CeTe<sub>2</sub>, contradictory to experiment (see Fig. 2(c)). Such discrepancy in CeTe<sub>2</sub> is expected to arise from the CDW distortion because the FSs in Fig. 7 are calculated for the non-CDW phases. The similarity between the measured FS and the calculated FS for PrTe<sub>2</sub> implies that the metallic states remain even after the CDW transition with the remnant ungapped FS and that the FS reconstruction due to the CDW transition is quite weak.

Figure 8 shows the measured and calculated FSs in the  $k_x$ - $k_z$  plane for RTe<sub>2</sub> (R=Ce, Pr). Figure 8(a) shows the measured Fermi-edge  $h\nu$  ( $k_z$ ) maps between 80-120eV with the horizontal direction along (100). So these maps show the FSs in the  $k_x - k_z$  plane. The  $k_x - k_z$   $(h\nu)$ map for  $CeTe_2$  is obtained for the BZ of MXM-ARA, and that for  $PrTe_2$  is for the BZ of  $X\Gamma X$ -RZR. For CeTe<sub>2</sub>, the *MXM*-*ARA* cut is chosen because the zigzagtype features, present around X in the measured FS for  $CeTe_2$  (Fig. 2(c)), are expected to be observed along M-X-M. In the  $h\nu$  maps for both CeTe<sub>2</sub> and PrTe<sub>2</sub>, the E<sub>F</sub>crossing states exhibit straight vertical dispersions along  $k_z$ . These features indicate that they originate from the 2D-like states, and that the interaction between Te(1)and R-Te(2) layers does not have a crucial contribution to the electronic states responsible for the CDW formation in  $RTe_2$ . The bright feature is observed in  $CeTe_2$ around X for  $h\nu \sim 110 - 115$  eV, which reflects the existence of several E<sub>F</sub>-crossing states due to the zigzag-type



FIG. 8: (Color online) Photon Energy Map. (a) The Fermiedge state  $h\nu$ -map for CeTe<sub>2</sub> for  $h\nu$  between 80 – 120 eV, obtained at T=35 K. Horizontal  $k_x$  direction is parallel to (100) and cuts through M(A)-X(R)-M(A) along  $k_z$  ( $h\nu$ ). (b) Similarly for PrTe<sub>2</sub> for  $k_x$  through X(R)- $\Gamma$ (Z)-X(R). (c) The calculated FS for the  $k_x$ - $k_z$  BZ in CeTe<sub>2</sub>. (d) Similarly for PrTe<sub>2</sub>.

FS features around the X point. The calculated FS of CeTe<sub>2</sub> in Fig. 8(c) also shows the straight vertical dispersions along  $k_z$ . In the calculated FS of PrTe<sub>2</sub> (Fig. 8(d)), however, there is an extra FS centered at  $\Gamma$ , which is not detected in the measured FS (Fig. 8(b)). The existence of this FS remains to be checked more carefully.

## IV. CONCLUSIONS

The overall shapes and sizes of the measured FSs of  $RTe_2$  (R=Ce, Pr) are similar to each other and they agree well with the calculated FSs for the normal-phase  $RTe_2$ . However, the zigzag-shaped features, having the four-fold rotational symmetry, are observed near the X points in the FS map of CeTe<sub>2</sub> only, but not in PrTe<sub>2</sub>, suggesting that these features are due to the hybridization between Ce 4f and Te(1) 5p electrons. According to the tightbinding model calculations, these zigzag FS features in CeTe<sub>2</sub> reflect the CDW-induced FS reconstruction due to the  $4 \times 4$  supercell structure. The partially ungapped FSs, measured below the CDW transition, reflect that the metallic states remain even below the CDW transition for both R=Ce and R=Pr. The effect of the LD is observed in ARPES. The  $E_{\rm F}$ -crossing states are stronger with the LV polarization than with the LH polarization, suggesting that the  $E_{\rm F}$ -crossing states have mainly the *ab*-plane orbital character.

We have found the following answers to the questions addressed in the beginning: (i) The CDW modulation of Te(1) sheets is large for CeTe<sub>2</sub>, but negligible for PrTe<sub>2</sub>. The CDW modulated structure for CeTe<sub>2</sub> is likely to be the  $4 \times 4$  structure (see Fig. 3(b)), but not compatible with the previously reported  $2 \times 2$  double herringbone structure. (ii) The R 4f states in RTe<sub>2</sub> do not contribute directly to the CDW formation, but the R 4f-Te 5p hybridization is much weaker in PrTe<sub>2</sub> than in CeTe<sub>2</sub>, which is reflected as the much weaker 4f spectral intensity near  $E_F$  in PrTe<sub>2</sub>. (iii) The opposite effects of the LD are observed between the  $E_F$ -crossing states and the states at ~ 1 eV BE, indicating that the  $E_F$ -crossing orbitals are ordered mainly in the *ab* plane. (iv) The  $\vec{k}$ -dependent energy gap  $\Delta_g(\vec{k})$  was observed. (v) The straight vertical dispersions along  $k_z$  are observed in the  $h\nu$  maps for the  $E_F$ -crossing states along  $k_z$  for both CeTe<sub>2</sub> and PrTe<sub>2</sub>. These features demonstrate the dominant 2D character for the states near  $E_F$ .

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- $^{31}$  The polarization of the incident light is not the origin of

the zig-zag features near X points in  $CeTe_2$  because these features are also observed when the sample is rotated azimuthally by  $45^{\circ}$ .

 $^{32}$  Kyoo Kim, et al., unpublished (2014).