Electronic structure of layered $1T$-TaSe$_2$ in commensurate charge-density-wave phase studied by angle-resolved photoemission spectroscopy

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We present a detailed angle-resolved photoemission study of the electronic structure of layered $1T$-TaSe$_2$ in the commensurate charge-density-wave (CDW) phase. A considerable reduction in the spectral weight of a quasiparticle band centered at the binding energy of about 0.25 eV below the Fermi level is observed in the momentum space ranging from the end of the first surface Brillouin zone to the second surface Brillouin zone. Moreover, no crossings of the Fermi level are visible in the whole Brillouin zone, meaning that the Fermi level lies in a pseudogap created by the tails of two overlapping Hubbard subbands. Our results indicate that not only the electron-phonon coupling, which is responsible for the formation of the CDW, but also the subsequent electron correlation effects in the Ta 5$d$ band play an important role for the establishment of electronic structure of $1T$-TaSe$_2$ in the commensurate CDW phase.

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The layered transition metal dichalcogenides $1T$-TaS$_2$ and $1T$-TaSe$_2$ have attracted much attention because of their quasi two dimensionality (2D) and, consequently, their unique physical properties leading to formation of the charge-density-wave (CDW). Although these two materials have the same CdI$_2$-type crystal structure and similar CDW superstructure, they show dramatically different physical properties. In particular, it was revealed in $1T$-TaS$_2$ that the metal-to-insulator transition from the quasicommensurate (metallic) CDW to the commensurate (insulating) CDW phase occurring at 180 K is closely related to a Mott localization induced by the electron correlation effects in the Ta 5$d$ band. Very recently, Pillo et al. studied the electronic structure of $1T$-TaS$_2$ in the quasicommensurate (metallic) CDW phase in detail using angle-resolved photoemission spectroscopy (ARPES). They pointed out the importance of the electron correlation effects in the quasicommensurate (metallic) CDW phase. On the other hand, $1T$-TaSe$_2$ undergoes an incommensurate CDW to commensurate CDW transition at about 430 K without drastic change in the electronic conductivity in the whole commensurate CDW phase, and exhibits a metallic behavior. Because of the larger interaction among the layers due to a larger charge transfer between Ta 5$d$ and Se 4$p$ orbitals, $1T$-TaSe$_2$ has a three-dimensional (3D) character in the electronic structure stronger than $1T$-TaS$_2$. From this viewpoint, one may say that the Ta 5$d$ electrons in $1T$-TaSe$_2$ may be less susceptible to localization.

To elucidate these physical properties, it is essential to understand the role of the electron-phonon coupling, which is responsible for the occurrence of the CDW and the subsequent electron correlation effects in the Ta 5$d$ band, in the electronic structure. Although the importance of the electron correlation effects in $1T$-TaS$_2$ has been recognized enough by virtue of various experimental techniques, such as scanning tunneling spectroscopy (STS) and ARPES, direct information about the occurrence of the CDW, that is, the crucial evidence for the nesting behavior on the 2D Fermi surface (FS) has yet to be reported. The influence of the electron correlation effect for $1T$-TaSe$_2$ in the CDW phase is expected to be smaller than that for $1T$-TaS$_2$ in the commensurate (metallic) CDW phase, because of the larger interaction between Ta 5$d$ and Se 4$p$ orbitals, as mentioned above. In other words, it is thought that the role of the electron-phonon coupling to the electron correlation effects for the electronic structure of $1T$-TaSe$_2$ becomes more important compared with that for $1T$-TaS$_2$. To investigate the interplay between the electron-phonon coupling and the electron correlation effects of the Ta 5$d$ band on the electronic structure of $1T$-TaSe$_2$ in the commensurate CDW phase, we performed a detailed ARPES study.

ARPES measurements were carried out at BL-1C of the Photon Factory (KEK, Tsukuba) using an electron spectrometer mounted on a two-axis goniometer (VG ARUPS10). The sample goniometer used here provides independent polar, azimuth, and tilt rotation of the sample. All ARPES spectra were taken at the photon energy ($h \nu$) of 40 eV. The radiation was linearly polarized in the horizontal plane of incidence. The samples were mounted vertically and only photoelectrons emitted from the plane defined by the light beam and the surface normal were observed. The emission angle of the photoelectron measured from the surface normal
samples were cleaved angular resolution and the energy resolution were 6 CdI\textsubscript{2}-type structure and SBZ correspond to parallel momenta of rotating the samples to the surface normal. Single crystals of 2 polariza-
tion effects of the light reported previously,\textsuperscript{17} we ro-
ted the samples in the CDW phase.\textsuperscript{5,11,19–21} This means that the overall shape of the electron wave functions along the \(\overline{\Gamma M}\) direction is governed by the Fourier compo-
nents of the crystal potential of the undistorted 1T lattice, rather than the CDW-related contribution.\textsuperscript{21} In contrast, such an exchange behavior of the spectral weight is not seen along the \(MK\) direction, only band A appears around the \(M\) point, as shown in Figs. 1(b) and 1(d).

In order to get more detailed information about the actual behavior of these QP bands near the \(MK\) direction, we measured azimuth dispersion spectra through the first and the second SBZ for the different parallel momenta, as shown in Fig. 2. The data start near the \(\overline{\Gamma M}\) direction and end near the next \(\overline{\Gamma M}\) direction through the \(\overline{KM}\) lines.\textsuperscript{22} Each azi-
muth scan has been carried out along one of the arcs shown in the bottom right sketch of Fig. 2. The parallel momentum on the top right of each dispersion spectra corresponds to the radius of the azimuth scan. In Fig. 2, the azimuth spectra show no significant polarization effects of the light along the \(KM\) and \(KM\overline{K}\) lines\textsuperscript{17} because \(\theta_y\) was kept constant (45\textdegree) during the measurement. This facilitates the following dis-
cussion. In these azimuth dispersion spectra, one observes two bands A and B, but not band C, which is shown around the \(\Gamma\) point in Fig. 1. The bands A and B show significant modulation with parallel momentum. At \(k_z = 0.64\ \text{Å}^{-1}\), the band B dominates the spectral shape. The band A appears in the azimuth spectra at \(k_z = 0.74\ \text{Å}^{-1}\). As we go from \(k_z = 0.74\) to 0.95 \(\text{Å}^{-1}\), the spectral weight of the band A increases and that of the band B decreases gradually. Then, the band B becomes faint at \(k_z = 1.05\ \text{Å}^{-1}\). The considerable reduction in the spectral weight of the band B is shown in the momentum region from \(k_z = 1.05\) to 1.24 \(\text{Å}^{-1}\). Finally, the band B reappears in the azimuth spectra at \(k_z = 1.34\ \text{Å}^{-1}\) and \(k_z = 1.44\ \text{Å}^{-1}\).

In order to corroborate the reduction in the spectral weight of the band B, we showed (parallel) momentum dis-
tributions of the spectral weight at the binding energy \((E_B)\) of (a) 100, (b) 200, (c) 400, (d) 600, (e) 800, and (f) 1000 meV in Fig. 3. The topology of the Fermi surface, or the momentum distribution of the spectral weight at \(E_B = 0\ \text{meV}\) (not shown) is almost consistent with that at \(E_B = 100\ \text{meV}\). White and black corresponds to high and low spectral weight, respectively. The momentum distributions of the spectral weight at \(E_B = 100\) and 400 meV are in-
duced by band B and those at \(E_B = 600\) and 1000 meV are induced by band A. For the sake of convenience, the distributions of the high spectral weight at each binding en-

**FIG. 1.** EDC spectra of 1T-TaSe\(_2\) along the (a) \(\overline{\Gamma M}\), (b) \(MK\), and (c) \(\overline{K}\) directions, taken with \(h\nu = 40\ \text{eV}\) at RT, and (d) the empirical band dispersion visualized by use of the second derivatives of the EDC spectra. The empirical bands are displayed in the linear gray scale plots, where the black regions denote the negative peak of the second derivatives. The notation in the parenthesis are empirical band dispersion visualized by use of the second derivatives. The EDC spectra at the ~G point denote normal emission and the ~M and ~K points in the SBZ correspond to parallel momenta of \(k_z = 1.05\) and 1.21 \(\text{Å}^{-1}\), respectively. At first sight, we find three distinct quasiparticle (QP) bands centered at the binding energy of about 0.9, 0.25, and 0.1 eV, denoted by A, B, and C, respectively in the EDC spectra. These QP bands can be explained by a gross distortion of the band structure due to the formation of the CDW superstructure.\textsuperscript{19} Also, an exchange of the spectral weight between these QP bands along the \(\overline{\Gamma M}\) di-
rection is shown in Fig. 1(a). As a result of the exchange, it is shown in Fig. 1(d) that the center-of-gravity of the spectral weight disperses downward in energy along the \(\overline{\Gamma M}\) di-
rection. The empirical dispersive behavior along the \(\overline{\Gamma M}\) direction is predicted by theoretical calculations in the normal state without the formation of the CDW,\textsuperscript{9,10} which is very similar to that of 1T-TaSe\(_2\) in the CDW phase.\textsuperscript{5,11,19–21} This means that the overall shape of the electron wave functions along the \(\overline{\Gamma M}\) direction is governed by the Fourier compo-
nents of the crystal potential of the undistorted 1T lattice, rather than the CDW-related contribution.\textsuperscript{21} In contrast, such an exchange behavior of the spectral weight is not seen along the \(MK\) direction.
energy are shown by means of an arbitrary ellipse centered at the $M^\|$ point in Fig. 3. The extension of the ellipses from the bands A and B reduces gradually with increasing binding energy. This behavior is qualitatively consistent with the prediction of theoretical calculations in the normal state without the formation of the CDW, showing an electron pocket around the $M^\|$ point. 9,10,12 As shown in Figs. 3 (a) through (f), a considerable reduction in the spectral weight of the band B is shown in the momentum space for the area between the end of the first SBZ and the second SBZ. Very recently, Pillo et al. observed similar behavior with respect to the band B for 1$T\text{-TaSe}_2$ in the quasicommensurate CDW and commensurate CDW phases, showing that a considerable reduction in the spectral weight of the band B occurs around the $M^\|$ point. 5

Based on a theoretical picture where the nesting of the Fermi surface plays a key role in the formation of the CDW, it is considered that the observed reduction in the spectral weight of band B is ascribed to the CDW-induced energy gap. From previous tunneling studies for 1$T\text{-TaSe}_2$, it was reported that the values of CDW-induced energy gap $2\Delta_{\text{CDW}}$ are about 0.3 (Ref. 23) and 0.5 eV. 24 The size of these energy gaps is almost consistent with the binding energy of the band B (0.25 eV). However, there are two questions about this nesting scenario. First, the locations in the momentum space where the drastic reduction in the spectral weight of the band B occurs are different for 1$T\text{-TaSe}_2$ and 1$T\text{-TaS}_2$. This should be explained in terms of the difference in the dimensionality. 8–11 The electronic structure of 1$T\text{-TaS}_2$ is independent of the momentum normal to the layers ($k_z$) due to negligibly small charge transfer between Ta 5$d$ and S 3$p$ orbitals,11 resulting in the highly 2D character. 9,10 On the other hand, the electronic structure of 1$T\text{-TaSe}_2$ depends on $k_z$ because of a large charge transfer between Ta 5$d$ and Se 4$p$ orbitals. 8–10 In order to confirm the $k_z$ effects on the nesting of the FS for 1$T\text{-TaSe}_2$, further ARPES experiments are required using various photon energies. The other question is that the size of the energy gap is inconsistent with that predicted by mean-field (MF) theory. Using the binding energy of the band B for $\Delta_{\text{CDW}}$ (0.25 eV) and the transition temperature ($T_{\text{CDW}}$) of 600 K used in Ref. 23, $2\Delta_{\text{CDW}}/k_B T_{\text{CDW}}$ is estimated to be 9.7, indicating strong coupling in the CDW phase, where $k_B$ is Boltzmann’s constant.

The strong coupling behavior is supposed to be caused by the electron correlation effects in the Ta 5$d$ band. 23 For 1$T\text{-TaS}_2$ in the quasicommensurate (metallic) CDW phase, previous ARPES studies showed that no crossings of the Fermi level are visible in the complete Brillouin zone, result-
ing in a pseudogap of the FS. It was concluded that the pseudogap behavior is induced by the electron correlation effects in the Ta 5d band. Our ARPES spectra for 1T-TaSe₂ in the commensurate CDW phase also shows no crossings of the Fermi level, as shown in Figs. 1 and 2. For $h\nu=40$ eV used here, the $\Gamma M$ line in the first SBZ almost corresponds to the AL line in the bulk BZ, assuming a work function of 4.5 eV and an inner potential of 12.2 eV. Along the AL line, band calculations predict one-particle Ta 5d band crossing the Fermi level. This means that the lack of bands crossing the Fermi level is not due to the 3D character of the electronic structure of 1T-TaSe₂, but due to the electron correlation effects in the Ta 5d band, resulting in a pseudogap at the Fermi level created by the tails of two overlapping Hubbard subbands.

In our previous ARPES study, we concluded that 1T-TaSe₂ has a 3D electronic structure from the normal emission spectra results along the $\Gamma A$ direction. From the previous results and the present ARPES results, one may say that the electronic structure of 1T-TaSe₂ in the commensurate CDW phase can be described by the interplay of the electron correlation effects in the Ta 5d band and the large hybridization effects between Ta 5d and Se 4p orbitals which cause the observed 3D character.

We examined in detail the electronic band structure of 1T-TaSe₂ in the commensurate CDW phase by means of ARPES. In particular, a considerable reduction in the spectral weight of the QP band centered at the binding energy of about 0.25 eV was shown in the momentum space ranging from the end of the first SBZ to the second SBZ. Moreover, no crossings of the Fermi level were shown in the complete Brillouin zone, which means that the Fermi level lies in a pseudogap created by the tails of two overlapping Hubbard subbands. Our results indicate that not only electron-phonon coupling, which is responsible for the formation of the CDW, but also the effects of subsequent electron correlation in the Ta 5d band play an important role for the establishment of electronic structure of 1T-TaSe₂ in the commensurate CDW phase.

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16 The sample goniometer used here is constructed for motorized, computer-controlled data acquisition and can be cooled with LHe down to 12.5 K (R-Dec Co. Ltd., i GONIO LT); Y. Aiura et al., Rev. Sci. Instrum. 74, 3177 (2003).
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