Origin of First-Order-Type Electronic and Structural Transitions in IrTe$_2$

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We have explored the origin of unusual first-order-type electronic and structural transitions in IrTe$_2$, based on the first-principles total energy density functional theory analysis. We have clarified that the structural transition occurs through the interplay among the charge density wavelike lattice modulation with $q_{1/5}=(1/5,0,1/5)$, in-plane dimer ordering, and the uniform lattice deformation. The Ir-Ir dimer formation via a molecular-orbital version of the Jahn-Teller distortion in the Ir-Ir zigzag stripe is found to play the most important role in producing the charge disproportionation state. Angle-resolved photoemission spectroscopy reveals the characteristic features of structural transition, which are in good agreement with the density functional theory bands obtained by the band-unfolding technique.

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Much attention has been recently paid to IrTe$_2$ since Yang et al. [1] and Pyon et al. [2] discovered superconductivity (SC) in Pd- and Pt-doped IrTe$_2$. Aside from the SC in doped IrTe$_2$, the parent compound IrTe$_2$ itself exhibits quite intriguing electronic and structural properties, which have not yet been fully comprehended. IrTe$_2$ crystallizes in a hexagonal structure of CdI$_2$ type ($P\bar{3}m1$) at room temperature ($T$). Upon cooling, IrTe$_2$ undergoes a structural transition around $T_\text{s} \approx 280$ K, exhibiting a sudden jump in the resistivity with a large hysteretic feature of the first-order type. It was argued that this behavior stems from the partial gap opening due to a charge density wave (CDW) transition with $q_{1/5}=(1/5,0,1/5)$ [1,3]. With Pd doping, the CDW is suppressed and the SC emerges with $T_C$ up to 3 K at 4% Pd doping. The interplay between the CDW and the SC results in a domelike phase diagram with respect to the Pd doping ratio.

The structural transition in IrTe$_2$, however, looks different from a standard CDW transition. Namely, the system has the first-order-type transition to a commensurate structure and exhibits heavy reconstruction of the electronic structure over a large energy window [4]. The phonon-softening instability signifying the structural transition was not obtained in the phonon dispersion for hexagonal IrTe$_2$ [5]. Furthermore, unlike other transition-metal dichalcogenide systems that have the SC ground state, for which the pressure suppresses the CDW structural transition but enhances the SC transition, the opposite trend was observed for IrTe$_2$ [6]. Hence, several different scenarios have been suggested as the origin of the structural transition in IrTe$_2$, such as the charge or orbital density wave [1], an orbital-induced Peierls instability [7,8], a crystal field of Te $p$ orbital [4,5,9], or an anionic depolymerization transition [10].

Using single-crystal x-ray diffraction, Pascut et al. [11] refined the low-$T$ crystal structure of IrTe$_2$ as a triclinic structure ($P\bar{1}$), as shown in Fig. 1(a). They argued that Ir dimer formation occurs during the structural phase transition, and the resulting energy gain plays a crucial role in

![FIG. 1 (color online). (a) High-$T$ HEX unit cell (blue), and low-$T$ 5X unit cell (black) of IrTe$_2$. The violet (grey) spheres are Ir (Te) atoms, yellow octahedra represent IrTe$_2$ octahedra, and red rods indicate Ir$_3$-Ir$_3$ dimers. (b) The near-$E_F$ Ir wave function of $d_{z^2}$ character forms antibonding with Te wave functions located along diagonal directions. The bright (dark) red triangle describes the Te lattice above (below) the Ir layer. x, y, and z are local coordinates of Ir, approximately pointing toward Te sites. x–y lies in the Ir plane. (c) The orbital-projected Fermi surface (FS) of the high-$T$ phase of IrTe$_2$. The color of the FS indicates the weight contribution from Ir $d_{z^2}$, which shows the clear 1D nature (to plot orbital-projected FS, we used the MOLISO package [12]).](image-url)
the structural transition. In fact, IrTe$_2$ is a rare system that exhibits charge ordering or disproportionation together with structural modulation in the metallic state. This property suggests that IrTe$_2$ at low $T$ is on the verge of a localized and extended electronic system.

In this Letter, we have explored the microscopic origin of the first-order-type electronic and structural transitions in IrTe$_2$. As described above, there has yet been no consensus on the microscopic mechanism elucidating the first-order structural transition in IrTe$_2$. Most previous studies concentrated on the question of which atom, Ir or Te, is responsible for the structural transition. However, results of density functional theory (DFT) and x-ray absorption spectroscopy indicate that the Ir-Te hybridization is strong enough to yield covalent states [13]. In this situation, separating out the contributions of Ir and Te is not possible; they would contribute together to the relevant physics in IrTe$_2$. We have found that the first-order-type electronic and structural transitions occur through the interplay among the CDW-like lattice modulation with $q_{1/5}$ (hereafter 1/5 lattice modulation), in-plane dimer ordering, and the uniform lattice deformation. We have also analyzed Angle-resolved photoemission spectroscopy (ARPES) data for the low-$T$ phase of IrTe$_2$, employing the band-unfolding technique for the first time, and demonstrated that our structural model is quite consistent with the ARPES band dispersions, Fermi surfaces, and spectral weights.

Electronic structures within the DFT were obtained by using the full-potential linearized augmented plane wave band method implemented in WIEN2K code [14]. For the structural relaxations, both VASP (pseudo-potential projected augmented wave code) [15] and WIEN2K were employed. We optimized the hexagonal structure with the inclusion of the spin-orbit coupling, utilizing three exchange-correlation potentials: local density approximation (LDA) and generalized gradient approximation (GGA) in the Perdew-Burke-Ernzenhof (PBE) and PBEsol (revised PBE for solid) scheme [16]. Since PBEsol gives the optimized volume closest to the experimental one, we chose the PBEsol results throughout this Letter [17].

Figure 1(c) shows the DFT FS of the high-$T$ phase of IrTe$_2$. The color indicates the weight contribution from the Ir $d_{z^2(x-y)}$ orbital character, the wave function form of which is depicted in Fig. 1(b). This orbital-projected FS manifests that the FS has a quasi-one-dimensional (1D) nature along three different diagonal directions. The 1D nature of the FS is more clearly seen in the tight binding (TB) model analysis in the Supplemental Material [18]. This 1D nature, together with saddle-point scattering, produces the FS nesting and the corresponding susceptibility peak along the diagonal direction [1]. Even though the susceptibility peak itself is not fully sufficient to explain the first-order structural transition of IrTe$_2$, it is to be an important ingredient for the 1/5 modulation observed for the low-$T$ phase of IrTe$_2$. Hereafter, we will refer to high- and low-$T$ phases of IrTe$_2$ as HEX and 5X, respectively.

We have first examined the energetics between the HEX and 5X phases of IrTe$_2$. The relaxed 5X structure was obtained starting from the experimental 5X structure refined by Pascut et al. [11]. To compare total energies under the same condition, we considered a supercell of the high-$T$ phase of IrTe$_2$ (HEX5), which is described with the same group symmetry as for 5X IrTe$_2$. It is expected that the structural transition occurs through the uniform lattice deformation (changes of lattice constants and angles between them) and the modulation of internal degrees of freedom (the 1/5 lattice modulation and the Ir dimerization). Therefore, to separate out the effects of the lattice deformation and the modulation, we considered artificial structural phase space in between the HEX5 and 5X structures, which was obtained by linearly interpolating the lattice structures and the internal parameters (atomic positions) of HEX5 and 5X phases independently. Total energy calculations were performed on this simplified structural phase space by using WIEN2K.

The energy contour in Fig. 2(a) shows that there are two energy minima in the structural phase space, which correspond to HEX5 and 5X structures. The energy of the 5X phase is lower than that of the HEX5 by 6.25 meV per formula unit. The double-well structure with a saddle-shaped barrier in between the two minima indicates an existence of the first-order-type structural transition. As shown in Fig. 2(b), the height of energy barrier is about 11 meV per formula unit. It has been suggested that the

![FIG. 2 (color online). (a) Energy contour of IrTe$_2$ in the structural phase space. A and B stand for the structural parameters of HEX5 and 5X, respectively. The horizontal and vertical axes represent the in-plane dimerization coupled to the 1/5 lattice modulation and the uniform lattice deformation, respectively, which are linearly interpolated between structural parameters of A and B on 13 $\times$ 13 mesh points [18,26]. Energy zero corresponds to the energy of the HEX5 structure. C represents a saddle point where the energy barrier is the lowest. (b) Energy profile along the path connecting A and B directly. The 5X structure is seen to be lower in energy than the HEX5 by 6.25 meV per formula unit. (c) A schematic mechanical model describing the interplay between the Ir$_3$-Ir$_3$ dimerization ($x$) and the uniform lattice deformation ($y$), which are coupled through Te$_1$ distortion ($z$) [18].]
in-plane Ir dimerization plays a crucial role in stabilizing the 5X structure [11,22–25]. Figure 2(a), however, reveals that just the Ir dimer formation, which corresponds to the increment along the horizontal axis from A, increases the energy solely monotonically. Indeed, it reveals that not only the in-plane Ir dimerization, but also the lattice deformation, is an essential ingredient for the structural transition to 5X. This explains why there is no softening feature in the phonon dispersion for HEX IrTe, which does not take into account the uniform lattice deformation [5].

The Ir dimerization pushes away nearby Te atoms; this brings about the lattice deformation. This mechanism can be schematically described by a simple mechanical model in Fig. 2(c), which contains the attractive interaction between Ir dimers and the additional harmonic interactions describing the lattice deformation. It shows that, as x decreases with dimerization, Te is pushed away from Ir, and so y increases due to the spring z. This model yields the double-well energy minima properly, as shown in the Supplemental Material [18]. It is thus deduced that the interplay among the 1/5 lattice modulation, the in-plane Ir dimerization, and the uniform lattice deformation is the origin of the first-order structural transition in IrTe2. Pascut et al. [11] performed similar energy calculations for the HEX5 and 5X phases, but they obtained lower energy for HEX5 than for 5X, as opposed to what is shown in Fig. 2(b). Presumably, their unphysical result comes from the less precise description of structural parameters, through the use of the different exchange-correlation functional.

It is worth noting that the direction of the 1/5 lattice modulation and the direction of the in-plane dimer formation do not coincide. Moreover, there are two substructures that depend on the Ir3-Ir3 dimer ordering patterns along the a direction (see Fig. 3). We expect that this feature arises from the cooperative Jahn-Teller (JT) type distortion that occurs with the 1/5 lattice modulation. In Fig. 3, we constructed an artificial structure with no dimer ordering (no-DO), lattice constants, and atomic positions that are identical to those of HEX5 except for the Ir3 and Te1 positions. In fact, the no-DO phase results from the 1/5 lattice modulation. As shown in Fig. 3(e), in no-DO, Ir3-Ir3 dimer ordering is not realized yet, but the Ir3-Ir3 distance becomes uniformly shorter and the Te1 atoms moved farther from the Ir layer, forming a Ir3 zigzag chain along the a direction. In HEX5, the DOSs for three t2g orbitals are degenerate, as shown in Fig. 3(d). However, as the 1/5 lattice modulation occurs, the degeneracy of t2g orbitals is lifted, separating out the dxy orbital, which has weaker hopping in the in-plane zigzag chain.

Notice that the charge transfer to the Te1p orbital occurs in this process [7,11,13], resulting in the increase of hole carriers in the degenerate antibonding Ir3-Ir3 dxy/dyz molecular orbitals. These two degenerate orbitals, which belong to different molecules [as depicted in Fig. 3(e)] couple to two modulations in the Ir3 zigzag chain, leading to a specific dimer ordering, as shown in Fig. 3(f). This is a molecular-orbital version of the JT effect in the zigzag stripe, composed of Ir3 and Te1 atoms. As a consequence of the dimer ordering, Te1 moves away from the Ir3-Ir3 center; thereby, Ir3 dimerization induces the uniform lattice deformation, as described in Fig. 2(c), giving rise to the first-order structural transition. Though we describe the processes in sequence, they would occur simultaneously.

There is experimental evidence for charge disproportionation or ordering of Ir (3+ /4+) in IrTe2 [7,10,28,29]. To inspect the charge disproportionation property in the 5X phase of IrTe2, we counted the number of Ir d (t2g) electrons inside the muffin-tin sphere. Table I shows that Ir3 has fewer electrons than Ir1 and Ir2, by about 0.5 electrons, reflecting that the valence state of Ir3 is close to 4+, while those of Ir1 and Ir2 are close to 3+. Table I also presents the core-level shift of Ir 4f levels in the 5X phase with respect to those in the HEX phase. The 4f level in Ir3 is deeper than those in Ir1 and Ir2 by about 0.4 eV, which indicates less screening of nuclear potential due to fewer valence electrons in Ir3 than in others. These values are consistent with experimental findings by Qian et al. [9].

![Figure 3](image)

**Figure 3** (color online). (a)–(c) Partial densities of states (DOSs) for HEX5, no-DO, and 5X structures, where no-DO corresponds to an artificial structure without dimer ordering, as depicted in (e). (d)–(f) In-plane structures and corresponding schematic DOSs for HEX5, no-DO, and 5X. The yellow shade represents the compressed region due to the 1/5 lattice modulation. Large (small) circles denote Ir (Te) atoms. The red and white small circles are Te atoms located above and below the Ir layer. Te atoms in the yellow shade are Te1. Two DOS peaks in blue and green correspond to bonding and antibonding states of dxy and dyz. Note that dxy and dyz orbitals participate in two different dimer orderings: one specific dimer ordering (red thick lines) is shown in (f), which is associated with the dxy orbital.

| Table I. Ir 5d t2g orbital occupation n and Ir 4f1/2 core-level shift ΔE in the 5X phase with respect to that in the HEX phase of IrTe2. |
|-----------------|-----------------|-----------------|
| **Ir type**     | **n [Ir 5d t2g]** | **ΔE [Ir 4f] (eV)** |
| Ir1             | 4.49            | 0.040           |
| Ir2             | 4.48            | -0.057          |
| Ir3             | 3.97            | -0.432          |
The properties in Table I show a strong indication of the charge disproportionation feature in the 5X phase of IrTe₂, which indeed arises from the Ir₃-Ir₃ dimerization.

To examine how the structural transition and the dimer formation are reflected in the electronic structure of the 5X structure, we compared the DFT band dispersions with ARPES data in Fig. 4 [18]. For this purpose, we have utilized the band-unfolding technique [30–33], which takes into account the Bloch phase factor between 5X and HEX cells, and maps the wave function in the 5X cell to that in the HEX cell. The relation between the HEX and the 5X Brillouin zone is shown in Figs. 4(b) and 4(c) [18]. Figure 4(a) presents Ir-projected band dispersions of 5X that are unfolded into a hexagonal in-plane BZ along the k path in Fig. 4(c).

The photoionization cross section of Ir 5d is 11 times stronger than Te 5p at 75 eV [34], therefore, we considered Ir-projected bands in Fig. 4(a) [18]. It is seen that the band splittings appear at some k points near EF. For example, there appears an abrupt change or splitting in the near-EF band along K1-Γ, which is quite different from that along K2-Γ. Another noticeable feature is the appearance of flat bands around −1.4 eV near Γ (see along K2-Γ-M3). According to our TB model analysis along the in-plane Ir chain, this unique dispersion comes from the on-site energy term of dimerized Ir₃, which is different from those of other Ir atoms; this happens due to the charge transfer or the crystal field effect (see Supplemental Material [18]).

In Fig. 4(d), unfolded 5X FSs are plotted in the HEX BZ for various k色泽. For kₓ = 0, one of the lobes in outer FS almost disappears; this is due to the gap opening in this part of the FS of 5X. Note that this gap opens not by the 1/5 lattice modulation, but by the lifting of orbital degeneracy due to the JT distortion [see Fig. 3(f)]. In actual ARPES data, however, the gap opening at one of the outer FS lobes is not clearly observed. Instead, the reduction of ARPES intensity is observed at all the outer FS regimes for 5X with respect to those for HEX; this is because of the existence of three types of domains, which yields the domain-averaged ARPES intensity [18].

Figures 4(e) and 4(f) provide the comparison of DFT band structures and ARPES data. In Fig. 4(e), ARPES data measured at T = 290 K are compared with DFT bands of HEX, while in Fig. 4(f), those at T = 260 K are compared with DFT unfolded bands of 5X. Quite good agreements in the spectral weights as well as the dispersions are revealed at both T's, which indeed verifies the formation of Ir₃ dimer ordering upon cooling. Furthermore, it supports the reliability of the structural data employed in this study [11].

Finally, it is worthwhile to examine the doping and pressure effects on the structural and SC transitions in IrTe₂ based on the present model. We have found that, under positive pressure, the peak structure in the susceptibility χ(q) of HEX IrTe₂ becomes enhanced at q₁/5, while under negative pressure, that becomes reduced. The reduction in the peak structure of χ(q) is also obtained when considering the carrier doping of Pd and Pt and associated structural relaxations. Even though more detailed study is needed, this feature provides qualitative clarification of the intriguing pressure effects on the structural and SC transitions in IrTe₂ [6], and also the doping-induced SC transition in doped IrTe₂ [2,35]. The c/a ratio, as well as the internal structural parameters, will strongly depend on the pressure. In fact, the angle between atoms in the Ir-Te-Te-Ir chain is found to change sensitively with variation of the pressure. The 1D quality in IrTe₂, an essential ingredient of the structural transition, is then to be altered, and, accordingly, the SC transition is induced.

In conclusion, we have demonstrated that there is an energy barrier in between the high-T (HEX5) and low-T (5X) phases of IrTe₂, which explains the first-order-type electronic and structural transitions in IrTe₂. This feature explains the limitation of linear response-based calculations, such as phonon calculation, in describing the structural transition in IrTe₂. We have clarified that the structural transition in IrTe₂ originates from the interplay among the CDW-like lattice modulation with q₁/5 = (1/5, 0, 1/5), the in-plane Ir₃-Ir₃ dimer formation, and the uniform lattice deformation, all of which are
coupled through the molecular-orbital version of the JT distortion. With Ir$_3$-Ir$_3$ dimerization, the charge disproportionation state is realized. Quite good agreement between the unfolded DFT bands and FSs and the ARPES data for both HEX and 5X phases of IrTe$_2$ supports our analysis of the structural transition, especially the formation of Ir$_3$-Ir$_3$ dimerized states.

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