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By combining bulk sensitive soft-X-ray angular-resolved photoemission spectroscopy and first-principles calculations we explored the bulk electron states of WTe$_2$, a candidate type-II Weyl semimetal featuring a large non-saturating magnetoresistance. Despite the layered geometry suggesting a two-dimensional electronic structure, we directly observe a three-dimensional electronic dispersion. We report a band dispersion in the reciprocal direction perpendicular to the layers, implying that electrons can also travel coherently when crossing from one layer to the other. The measured Fermi surface is characterized by two well-separated electron and hole pockets at either side of the Γ point, differently from previous more surface sensitive ARPES experiments that additionally found a significant quasiparticle weight at the zone center. Moreover, we observe a significant sensitivity of the bulk electronic structure of WTe$_2$ around the Fermi level to electronic correlations and renormalizations due to self-energy effects, previously neglected in first-principles descriptions.

**Introduction** - The observation of unconventional transport properties in WTe$_2$ [1], such as the large non-saturating magnetoresistance with values among the highest ever reported, prompted experiments and theory to address the electronic structure of this semimetallic transition metal dichalcogenides (TMD) [2–6]. WTe$_2$ consists of layers of transition metal (TM) atoms sandwiched between two layers of chalcogen atoms, similarly to other TMDs such as MoS$_2$ and MoSe$_2$. Because of the layered structure, TMDs have commonly been considered as quasi-two-dimensional solids. The easiness of exfoliation down to a single layer makes them appealing for nanoscale electronic applications. WTe$_2$ has also been theoretically described, in a recent paper, as the prototypical system to host a new topological state of matter called type-II Weyl semimetal [7]. At odds with standard type-I Weyl semimetals showing a point-like Fermi surface, type-II Weyl excitations arise at the contact between hole and electron pockets. Theoretical predictions were immediately followed by several surface sensitive angle-resolved photoemission (ARPES) studies claiming evidence of topological Fermi arcs [8–10].

Our previous investigation by surface sensitive ARPES, spin-resolved ARPES and DFT calculations, gave clear hints on the non-purely two-dimensional (2D) electron states of WTe$_2$ and suggested interlayer, i.e. $k_z$ perpendicular ($k_z$) dispersion and cross-layer compensation of electrons and holes [11]. However, a direct inspection of the electronic properties by means of bulk sensitive soft-X-ray ARPES technique, and more accurate calculations are needed in order to prove the three-dimensional (3D) character of the bulk electronic structure. By measuring ARPES with photon energy $h\nu$ in the range 400-800 eV, one probes the electron states averaging on several layers and therefore reducing the weight of the surface specific features that otherwise dominate the spectra when excitations energies in the VUV-range are employed. Furthermore, the increase of photoelectron mean free path in the soft-X-ray energy range results in a high intrinsic $k_z$ resolution of the ARPES experiment [12], essential to explore 3D effects in electronic band structure.

WTe$_2$ displays an unprecedentedly large non-saturating magnetoresistance even at magnetic fields $B$ as high as 60 Tesla [1]. A large orbital magnetoresistance is expected in semimetals, so that WTe$_2$ shares this intriguing feature with bismuth and graphite [13], all showing small concentrations of very mobile hole and electron carriers. Differently from bismuth and graphite, however, the magnetoresistance in WTe$_2$ exactly follows a $B^2$ dependence typical of an electron-hole compensated semimetal [14]. Carrier compensation, in turn, is only
FIG. 1: (Color online) a) View of the WTe$_2$ crystal structure. W and Te atoms are shown in orange and blue, respectively. b) Relative bulk and (001) surface Brillouin zones. c) $k_x$-$k_z$ Fermi surface ($k_y = 0$) taken in the 400-800 eV range of excitation energies. The value of the inner potential $V_0 = -6.5$ eV is used to compute the $k_z$ periodicity over many Brillouin zones (see rectangles), assumed here to be $2\pi/c \sim 0.45 \text{ Å}^{-1}$, see below for a discussion. d-f) $E$ vs $k$ band dispersions along the blue dashed lines in c). Arrows are guides to the eyes to highlight features from hole and electron pockets in a way consistent with c). g) Zoom of the $k_x$-$k_z$ Fermi surface around the $\Gamma_{30}$ point (red rectangle in c), and h-i) calculated Fermi surface within the LDA and LDA+U ($U = 2$ eV) approximations.

Results and Discussion – In Fig. 1c) we report the $k_z$ evolution of the Fermi surface spanning about 9 Brillouin zones in the out-of-plane reciprocal direction and about 3 in-plane Brillouin zones along the W chain direction $k_x$. Our measurements unambiguously unveil a clear continuous $k_z$ dispersion of the electronic states at the Fermi level, definitely proving that WTe$_2$ has a 3D bulk electronic structure despite the layered geometry common to all TMDs [12], and despite the preferential direction for electronic dispersion given by the zigzag TM chains. These results provide a direct spectroscopic validation of the early proposals based on quantum oscillations experiments and temperature dependent magnetoresistance measurements [5, 6].

A closer look at the $k_z$ evolution of the electronic states around $k_z = 0$ would suggest the out-of-plane periodicity be $\sim 0.9 \text{ Å}^{-1}$ (two rectangles along $k_z$ in Fig. 1c), as resulting from a unit cell containing only one WTe$_2$ layer, and contrary to the experimental structure consisting of two layers, as depicted in Fig. 1a) [17]. It is not unusual that the experimentally measured periodicity of a system does not match that of the structural unit cell. Common examples are given by nonsymmorphic systems such as CrO$_2$, graphite, and 2H-WSe$_2$ [18–20], for which selection rules restrict the final state symmetry. WTe$_2$ crystallizes in the nonsymmorphic space group $Pmn\bar{1}$. The nonsymmorphicity ensures a symmetry protected band degeneracy at the Brillouin zone point $Z = (0, 0, \pi/c)$ (see Fig. 1a in the Supp. Mat. [21]), despite the fractional translation invariance along the $k_z$ direction is broken by the non-equivalence of WTe$_2$ monolayers. To model this double periodicity in band structure calculations, we have projected the effective band structure of WTe$_2$ onto the irreducible representations of a Brillouin zone compatible with the one extracted from our experiments, with a necessary condition. It is equally mandatory that the carrier mobility does not depend on the applied magnetic field, a feature met by WTe$_2$ [1] but not for example by pure bismuth [15]. The bulk electronic structure of WTe$_2$ has been so far only investigated by means of transport measurements [5, 6]. The behavior of the resistivity under an external magnetic field is hard to reconcile with the picture of a layered solid: when the magnetic field is applied parallel to the layers, unexpected quantum oscillations were observed, suggesting that electrons may travel in a coherent way also across the weakly bonded layers [5]. Moreover, recent spin-ARPES data indicated both in-plane and out-of-plane spin polarization of the electron states below the Fermi level, deviating from what expected from spin-orbit interaction (SOI) in a non-interacting 2D layered system [11], and the balance between the hole and electron states was shown to be fully established only if cooperation of several layers, i.e. bulk 3D character, was included [11]. The above results, albeit indirectly, suggest that the electronic structure of WTe$_2$ has an intrinsic 3D character, unexpectedly for a TMD. However, up to now, no conclusive spectroscopic evidence of three-dimensionality was reported.
as laser excited ARPES [16] is the lack of any evident effects, enhancing selectively electron and hole contributions in subsequent Brillouin zones.

Quantum oscillation measurements. c) Calculated $k_x$-$k_y$ bulk Fermi surface within the LDA+U approximation for $U = 2$ eV.

The measured electron pocket is characterized by a bow-like $k_z$ dispersion (Fig. 1g), in agreement with the calculated Fermi surface based on the local density approximation (LDA) plus an on-site Hubbard $U$ of 2 eV (compare with cyan areas in Fig. 1i). However, discrepancies arise when comparing the hole pocket dispersion. In the measured Fermi surface, hole pockets seem to disperse all over the $k_z$ extension of the Brillouin zone, while first-principles calculations give disconnected pockets (purple areas in Fig. 1i). However, it is worth to note that the calculated $k_x$-$k_y$ Fermi surface (Fig. 2c), nicely reproduces the features and the extensions of the measured Fermi surface. This improves over standard (i.e. without U) LDA calculations (see Fig. 1h, discussions in Refs. [2, 5, 11], and discussion below), suggesting that electronic correlation could play a significant role. In fact, previous theoretical studies have demonstrated that LDA is capable of providing an overall good description of the electronic structure of WTe$_2$, especially the coexistence of electron and hole features and the onset of topological surface states, but at a quantitative level, significant discrepancies with experiment remain. In particular, LDA tends to overestimate the dimensions of the Fermi surface along the W chains direction (see Fig. 1h), positioning the minimum of the electron pocket at momentum values larger than ARPES [11]. This, together with the difficulty in resolving tiny features dispersing around the Fermi level, has limited so far the understanding of surface and bulk contributions on the Fermi surface. Moreover, since calculations predict Weyl points to be above the Fermi level, a direct comparison with experiment requires an accurate treatment of the unoccupied states.

Despite the fairly delocalized character of 5d W or-
orbitals, the inclusion of a moderate U in LDA leads to a sizeable modification of the electronic states in the proximity of the Fermi level, as shown in Fig. 3. The general trend as a function of increasing U (see Fig. 4 of the Supplement [21] for more values) is the small shift of the electron pocket toward lower momentum values and the sizeable modification of the hole pocket. It is interesting to note how for U = 2 eV, the value that gives a nice comparison between measured and calculated Fermi surfaces as shown in Fig. 2, the two pockets almost linearly cross at ~50 meV above the Fermi level along the k_x direction. A further increase of the U (Fig. 4 of Supp. Mat. [21]) causes the reopening of the gap between these pockets. If this correlation driven trend leads to a change of the topological properties of WTe_2 it would deserve a proper investigation. What we highlight here is that for U values slightly smaller than 2 eV, bands linearly cross at finite k_y, symmetric with respect the k_x axis, giving rise to type-I Weyl points, as shown in Fig. 3c). Such Weyl points, having opposite chirality, are connected by Fermi arcs when projected onto the surface Brillouin zone (Fig. 5 of Supp. Mat. [21]). At the critical value U_c = 1.98 eV these Weyl points touch on the k_x axis and annihilate. The appearance of a type-I Weyl point has been suggested as the fingerprint of topological transitions in noncentrosymmetric topological insulators [29].

Despite the approximate way in which DFT+U treats the electron-electron interaction i.e. within a energy independent Hartree approximation, the improved description of the Fermi surface both in the k_x-k_y plane and in the k_z direction, as well as of the optical conductivity of WTe_2 (Fig. 6 of Supp. Mat. [21]), indicates that the electronic states experience a non-negligible degree of correlation.

To fully endorse this thesis, we perform GW calculations in which band renormalization is induced by truly many-body self-energy effects. The resulting band structure, obtained within a relativistic framework, is shown in Fig. 3d). It displays significant changes with respect to the underlying DFT (dotted-dashed lines) and DFT+U ones: the position of the electron pockets shifts closer to the Γ-point, in better agreement with ARPES experiments, but in contrast to DFT(+U), GW finds a density of empty states at Γ right above the Fermi level, which leads to a strong renormalization of the hole pocket; finally, the quasiparticle bands exhibit a larger SOI induced splitting, which could indicate a strong coupling between relativistic effects and electronic correlations, a novel quantum phenomenon recently observed in other heavy materials subjected to Lifshitz-type instabilities [30]. Remarkably, our GW results may explain the observed large spin polarization and complex spin texture recently reported in Ref. [11], hardly reconcilable with standard DFT calculations.

**Concluding remarks** – In this Letter, we address the bulk electronic properties of WTe_2 by complementary bulk-sensitive electron spectroscopy and theoretical methods. Since the prediction of topological surface states in WTe_2 owing to a topological nature and its classification as type-II Weyl semimetal, the spectroscopic study of the bulk electronic structure of WTe_2 was missing. Our soft-X-ray ARPES measurements, by means of an unprecedentedly high intrinsic definition of k_z and a large range of its variation in the Fermi surface mapping, definitely demonstrate a 3D character of the electronic states. These results prove that layered materials as TMDs may host electrons moving from layer to layer in a coherent way, in agreement with the quantum oscillation transport results [5]. Moreover, our theoretical investigation shed light on the role of electronic correlations and self-energy effects on those electronic states, dispersing around the Fermi level, that play a relevant role in the transport properties of WTe_2.

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See Supplemental Material at http://xxxx.xxxx for experimental and computational details, as well as additional ARPES spectra and calculations. Supplemental Material includes Refs. [17, 34–42].

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Spectra in Fig. 1d, e) and f) refer to photon energies $h\nu = 425$ eV and 620 eV, such that Fermi surfaces in the first in-plane Brillouin zone ($k_z \sim 0$) look different (see $k_x = 10.8 \, \text{Å}^{-1}$ and $k_z = 13.0 \, \text{Å}^{-1}$ in Fig. 1c). Spectra for $h\nu = 425$ eV, $570$ eV ($k_z = 12.5 \, \text{Å}^{-1}$) and $668$ eV ($k_z = 13.4 \, \text{Å}^{-1}$) are similar.

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