Electronic bandstructure of ReS$_2$ by high resolution angle resolved photoemission spectroscopy

James L. Webb,* Lewis S. Hart, and Daniel Wolverson
Centre for Nanoscience and Nanotechnology, Department of Physics,
University of Bath, Bath BA2 7AY, United Kingdom

Chaoyu Chen, Jose Avila, and Maria C. Asensio†
Synchrotron SOLEIL, Saint Aubin, and Université Paris-Saclay, BP 48 91192 Gif-sur-Yvette, France

(Dated: September 6, 2017)

The rhenium-based transition metal dichalcogenides (TMDs) are atypical of the TMD family due to their highly anisotropic crystalline structure and are recognized as promising materials for two dimensional heterostructure devices. The nature of the band gap (direct or indirect) for bulk, few and single layer forms of ReS$_2$ is of particular interest, due to its comparatively weak inter-planar interaction. However, the degree of inter-layer interaction and the question of whether a transition from indirect to direct gap is observed on reducing thickness (as in other TMDs) are controversial. We present a direct determination of the valence band structure of bulk ReS$_2$ using high resolution angle resolved photoemission spectroscopy (ARPES). We find a clear in-plane anisotropy due to the presence of chains of Re atoms, with a strongly directional effective mass which is larger in the direction orthogonal to the Re chains (2.2 $m_e$) than along them (1.6 $m_e$). An appreciable inter-plane interaction results in an experimentally-measured difference of $\approx$100-200 meV between the valence band maxima at the Z point (0,0,1) and the Γ point (0,0,0) of the three-dimensional Brillouin zone. This leads to a direct gap at Z and a close-lying but larger gap at Γ, implying that bulk ReS$_2$ is marginally indirect. This may account for recent conflicting transport and photoluminescence measurements and the resulting uncertainty about the nature of the band gap in this material.

I. INTRODUCTION

The transition metal dichalcogenides (TMDs) are a class of material that can form thin sheets down to a single monolayer, analogous to to graphene but consisting of compound semiconducting materials rather than a single atomic species. It is these semiconducting properties that have attracted considerable recent interest with regard to fabricating and controlling new devices from single atomic species. It is these semiconducting properties of compound semiconducting materials rather than a single monolayer, analogous to graphene but consisting of thin sheets down to a few layers, that have attracted considerable recent interest with regard to fabricating and controlling new devices from single atomic species. It is these semiconducting properties of compound semiconducting materials rather than a single monolayer, analogous to graphene but consisting of thin sheets down to a few layers, that have attracted considerable recent interest with regard to fabricating and controlling new devices from single atomic species.

Of these materials ReS$_2$ is of particular interest due to several properties. Its optical [6, 7] and electrical transport properties are highly anisotropic, in particular with higher mobility in certain in-plane crystallographic directions as determined by electrical measurements using bulk-like flakes, [8, 9] making it an interesting material for the fabrication of FETs and polarization-sensitive photodetectors [8, 10, 11]. In this context, ReS$_2$ has the advantage of being stable under ambient conditions, unlike some similar 2D materials [12]. Next, spin-orbit coupling is important in ReS$_2$ because the presence of inversion symmetry means that spin-orbit splitting in unperturbed layers of any thickness is zero and may be manipulated via doping or gating. Finally, unlike many other TMD materials [13, 14] it has been proposed there is no transition from indirect to direct band gap with reduced thickness, meaning that the distinction between mono- and few-layer structures is not crucial for device concepts[15]. If true, this could be advantageous in terms of building efficient optical devices.

Figure 1(a) shows a schematic model of the atomic structure of ReS$_2$ with the lattice vector directions $\mathbf{a}$, $\mathbf{b}$ and $\mathbf{c}$ indicated. Here $\mathbf{a}$ lies in the direction of the Re chains and $\mathbf{b}$ at about 120° to them. This structure has been confirmed experimentally by scanning probe microscopy studies [16], indirectly by Raman spectroscopy [15, 17] and by single crystal x-ray diffraction by Lamfers et al. [18] with a triclinic structure and in-plane lattice parameters $a=6.352$ and $b=6.446$ Å. The question of whether the unit cell contains one or two layers stacked along the out-of-plane $c$ axis has been resolved in favor of a single layer, giving four formula units per unit cell[19] with $c = 6.403$ Å to 6.461 Å[20].

In performing the ARPES measurements we effectively probe the projection of the $k$-points in the 3D Brillouin zone (BZ) of the material onto a quasi-2D flat plane ($k_z=$constant; the value of $k_z$ is determined by the choice of excitation photon energy). This is exemplified in Fig. 1(b) for points in the full 3D BZ in a plane passing through the Γ point (shaded yellow), which are projected onto the measurement plane to produce a quasi-2D Brillouin zone in $k_x$, $k_y$. This quasi-BZ is shown at the bottom of Fig. 1(b) and is an irregular hexagon centered on $\bar{\Gamma}$. Here we show the projection of the lattice vectors $\mathbf{a}$ and $\mathbf{b}$ onto the measurement plane as $\mathbf{a}^\perp$ and $\mathbf{b}^\perp$. If the bulk material were thinned to a 2D mono-

* j.l.webb@bath.ac.uk
† maria-carmen.asensio@synchrotron-soleil.fr
FIG. 1. The atomic structure of ReS$_2$: (a) Re atoms in blue and S in yellow with the unit cell shown by the vectors $\text{a}$, $\text{b}$, $\text{c}$. Chains of Re atoms run along the in-plane vector taken here as $\text{a}$, with the second in-plane vector $\text{b}$ at about 120° to the chain direction. (b) The conventional triclinic Brillouin zone of ReS$_2$, showing the reciprocal lattice vectors $\text{a}^*$, $\text{b}^*$, and $\text{c}^*$, with the latter passing through the labeled $\Gamma$ and $Z$ points along the direction $\Lambda$. Two planes which pass through $\Gamma$ (the $\Gamma$ plane) and $Z$ (the $Z$ plane) and other high symmetry points are shaded; in our ARPES experiments, we effectively measure a projection of the 3D Brillouin zone onto the plane $k_2=$constant. This is shown schematically at the bottom of (b) by the irregular hexagon which is the projection of the $\Gamma$ plane. $\text{a}^*$ and $\text{b}^*$ represent the projections of the reciprocal lattice vectors in the $k_2=$constant plane, $\overrightarrow{\text{K}}, \overrightarrow{\text{M}}$ are the $k_2=$constant projections of their respective high symmetry points, and and $\overrightarrow{\text{X}}$ is the projection of the general point $\Lambda$

layer, this quasi-BZ would ultimately represent a good approximation to the 2D BZ as used in other work, for example Tongay et al. [15] to calculate the properties of monolayer ReS$_2$. We label directions $\overrightarrow{\text{M}}$ and $\overrightarrow{\text{K}}$ following this and other work and in order to show the direction of measurement of the ARPES data. It is necessary to distinguish between directions $\overrightarrow{\text{M}}_1..\overrightarrow{\text{M}}_3$ and $\overrightarrow{\text{K}}_1..\overrightarrow{\text{K}}_3$ since, in this triclinic material, none of these are related by symmetry. The directions $\text{b}^*$ and $\text{b}^*$ are orthogonal to the real space vector $\text{a}$, so that directions $\overrightarrow{\text{M}}_1$ and $\overrightarrow{\text{K}}_2$ are exactly orthogonal and approximately parallel, respectively, to the rhombic chains of ReS$_2$

Although electrical transport and optical absorption measurements have given some indirect insight into the ReS$_2$ bandstructure, along with ab initio calculations of the material properties [15, 21, 22], it is evident that gap size as well as the direct or indirect character of the gap is particularly sensitive to the details of any computational model (for instance, the choice of pseudopotential, and whether spin-orbit coupling is included). Consequently, the direct, accurate determination of the electronic bandstructure is necessary and timely. This is a task for which angle resolved photoemission spectroscopy (ARPES) is ideally suited. However, two primary difficulties exist in performing these measurements: they must be performed on a clean surface, ideally a crystal cleaved under ultra-high vacuum conditions, and the crystal facets of the material must be larger than the spot size of the X-ray beam in order to obtain clear, monocrystalline data. Recent advances in TMD handling and advances towards micro and nano-ARPES systems with beam sizes on the 100 nm scale mean these problems can now be overcome.

In this work we present the main results of our direct determination of the valence bandstructure of bulk ReS$_2$ using nano-ARPES, mapping the photoemission intensity in directions along and perpendicular to the Re atomic chains of the material and measuring also constant binding energy contours throughout large portions of the full 3D Brillouin zone. Our findings show the effect of the in-plane anisotropy on the electronic dispersions and the van der Waals interactions between planes, revealing the electronic dispersion perpendicular to the electronic band gap.

**METHODS**

Samples were commercially grown via Bridgman single crystal growth by 2D Semiconductors USA and were confirmed 99.9995% pure using secondary ion mass spectrometry. We performed prior studies on the crystals using Raman spectroscopy in order to confirm their phase and high crystal quality[23].

Nano-ARPES measurements were performed using the k-microscope at the ANTARES beamline at the Soleil synchrotron, Paris, with a spot size of 100 nm. At a photon energy of 100 eV, this beamline has an angular resolution of $\sim 0.2^\circ$ and an energy resolution of $\sim 10$ meV. The advantage of the nano-ARPES technique for the study of ReS$_2$ is that the X-ray beam spot size is smaller than the size of the crystallites (generally a few microns to tens of microns, as determined by optical microscopy), ensuring that the measured dispersion is obtained only from a single facet, which is important due to the high degree of direction-dependent anisotropy in the band structure. The samples were prepared by cleaving in-situ under UHV conditions (pressure $< 1 \times 10^{-10}$ mbar). A nano-positioning system was used to locate clean and flat areas of the sample with maximum ARPES intensity with measurements performed at photon energies from 95-180 eV. The sample was rear-cooled using liquid nitrogen to approximately 100 K in order to reduce thermal noise.
DFT calculations were performed using the Quantum Espresso package [24] to perform structural relaxation and obtain total energy and bandstructure simulations. We focus here on results obtained using a non-relativistic Perdew Burke Ernzerhof (PBE) generalised gradient approximation (GGA) exchange-correlation functional[25] but we also explored the use of a fully-relativistic Perdew Zunger (PZ) local density approximation (LDA) functional[26] with projector augmented wave (PAW) pseudopotentials generated by QE and PSLibrary46. The GGA and LDA results are compared to the experimental data in the Supplemental material, Figures S1 and S2 respectively [27]. The valence of Re was taken as 15 (configuration 5s²5p⁶5d⁵6s²). Kinetic energy cutoffs were 70 Ry (816 eV) and Monkhorst-Pack k-point meshes of 12 × 12 × 12 were used with a single 12-atom unit cell.

III. RESULTS

First, we consider how the observed constant energy maps in the \( k_z \) =constant plane reflect the crystal symmetry. Fig. 2 shows views of the crystal structure looking along the \( a \) and parallel to the \( c \) directions respectively. In Fig. 2(c), we show a constant energy surface plot of the ARPES signal intensity probing a set of binding energies \( E_b \) moving downwards in energy from the Fermi energy \( E_F \) and recorded by illuminating the samples with photons of energy \( h\nu = 100 \) eV. This excitation energy was chosen since it gives the optimum transmission of the zone plate used to focus the excitation beam. As we shall see below, this excitation energy means that the plane probed intersects the \( c^* \) axis at a point lying on the line \( \Gamma-Z \). Following the notation of Fig. 1(b), we label a general point of this type as \( \Lambda \); this point is the origin \( (k_x = k_y = 0) \) of the 2D quasi-Brillouin zone, and thus is the origin of the binding energy contour plots for a given excitation energy.

Using a gold sample \textit{in situ} in the ARPES system, the Fermi edge of its density of states was precisely determined; this experimental Fermi energy is shared by the ReS₂ sample as they have a common ground potential. We find that the Fermi level of the semiconductor sample is 1.5 eV above the valence band maximum, putting a lower bound on the single-particle band gap of 1.5 eV (unoccupied bands are not recorded by ARPES, so the conduction band minimum was not recorded here). This value is close to the lowest-lying excitonic band gap of \( E_{\text{ex1}}^\text{c} = 1.55 \) eV recorded at the same temperature.
FIG. 3. (a) shows the constant energy surface in the valence band at 1.5 eV below the Fermi energy $E_F$ as shown in Fig. 2(c) with the labels of the high symmetry points projected onto the $(k_x,k_y)$ plane. Panels (b) and (c) show high resolution ARPES plots recorded using an excitation photon energy of $h\nu = 100$ eV along the six inequivalent $K'\bar{\Lambda}K$ and $M'\bar{\Lambda}M$ directions.

(100 K) [28], showing the $n$-type character of our material. The doping state of ReS$_2$ is dependent on the details of the crystal growth with $p$-type material also being possible depending on the vapor transport method used[6, 29]. Consequently, the ARPES constant energy plots of Fig. 2(c) can be labeled at this local maximum of the valence band with binding energies of 1.5 eV, 1.7 eV and 1.9 eV below the Fermi level.

The plots of Fig. 2(c) clearly indicate a “wavy” shape of the contours related to the in-plane chains of Re atoms sketched in Fig. 2(b). This implies a marked difference between the dispersion in the direction along the Re chains and that perpendicular to them, with a more abrupt drop in the valence band energy along the chains. Such anisotropy directly affects the effective mass and hence the mobility in the two perpendicular directions. This aspect will be studied in detail below, where we obtain representative effective masses in both directions from our ARPES data. However, it is important to note that this in-plane anisotropy is not exclusive to these two directions. Fig. 3 indicates that the band structure along different $\overline{M'-\Lambda-M}$ directions is dissimilar, confirming that the in-plane anisotropy is not restricted to the directions along and perpendicular to the chains of Re atoms. The same remark can be made about the electronic dispersions along the $K'\bar{\Lambda}K$ directions, which are also not equivalent, even though the tops of the bands in all directions are centered at the surface $\bar{\Lambda}$ projection, as shown in Fig. 3(a). We distinguish in Fig. 3 between points either side of $\bar{\Lambda}$ (e.g, $M'$ and $M''$) because there is inversion symmetry only through the true $\Gamma$ point and so these pairs are not related by real reciprocal lattice vectors. Indeed, it can be seen in some of the ARPES plots of Fig. 3(b) and (c) that the dispersion is not exactly symmetrical about $\bar{\Lambda}$ for this reason.

Thus far, we have measured the bands only at an arbitrary point along $c^*$ (corresponding to the excitation energy of 100 eV) and this limits our analysis. Consequently, it is important to probe the limits of the 3D Brillouin zone, at the $\Gamma$ and $Z$ points. This can be achieved in ARPES by varying the excitation photon energy; the systematic study of the energy dependence of the ARPES signals allows the mapping of the whole ReS$_2$ 3D BZ and a precise analysis then provides the exact location of the...
FIG. 4. (a) 3D ARPES intensity plots along the $k_z$, $k_y$ versus binding energy $E_B$. The $k_z$ periodicity shows that at photon energies of 131 eV and 111 eV, the ARPES mapping probe planes containing the Γ and the Z symmetry points, respectively. From this experimental determination we have estimated the inner potential $V_{in} = 16 \pm 2$ eV. (b) Second derivatives of ARPES intensity taken at Z and Γ along the $K_2$ and $M_1$ directions. In both cases, we see a sharper peak at Z, compared to the flatter bands in Γ, particularly in the $M_1$ direction perpendicular to the Re chains. We see a difference in energy between the top of the valence band at Z and that at Γ of approximately 150-200 meV.
electron mobility was found to be about three times larger along the rhenium chains compared to the direction perpendicular to them [8]. In the Supplemental material, Figure S4, we give the conduction band effective masses derived from DFT calculations, which reflect this anisotropy [27]. The valence band masses given above would also lead to a similar degree of anisotropy in the hole mobility even if all other parameters determining the phonon-limited mobility (deformation potential constant, elastic modulus) were isotropic. However, no electrical measurements of hole mobility in ReS$_2$ are yet available for comparison.

To aid in interpretation of the data, we performed simulations of the present data using DFT calculations using parameters detailed in the Method section; some results are shown in Figure 6. We obtain qualitative agreement with the experimental data in both the $\overline{K}_2$ and $\overline{M}_1$ directions, with the calculations matching the uppermost experimental bands best for $\overline{K}_2$-$Z$ and worst for $\overline{M}_1$-$\Gamma$. To facilitate comparison of these simulations to experiment, figures S1 and S2 of the Supplemental material show the simulations superimposed on the ARPES data for calculations using both the PBE functional (as shown in Figure 6; Fig. S1) and a fully relativistic LDA functional (Fig. S2) [27].

Importantly, in both cases we replicate the anisotropy observed in experiment with respect to the Re chain direction. As an example, we focus on the dispersion along $\overline{K}_2$-$\Gamma$-$\overline{K}_2$, Fig. 6(a), where the highest energy valence band has a strongly peaked and approximately parabolic form whilst the next valence band down in energy has a distinctive double-peak structure; this is exactly as found in experiment, as shown in Figs. 4(b) and 5(b). In these simulations, we did not take into account the slight curvature of the plane in $k$-space probed in ARPES that arises from the non-conservation of $k_z$ [32]; for these rather high excitation energies, we expect that this does not introduce a significant error but it may affect the comparison to experiment particularly for bands deeper in the valence band.

Based on these simulations of the experimental data we used DFT to calculate the full band structure of the material including the lowest-lying conduction band states. Here, both PBE (GGA) and fully relativistic PZ (LDA) functionals were used in order to estimate the effects of spin-orbit coupling due to the high atomic number of rhenium. The results are qualitatively similar apart from the well-known underestimation of the band gap in the LDA, and so we focus here on the PBE (GGA) functional as used to produce the results shown in Fig. 6.
FIG. 6. Calculations of the valence bands measured by ARPES in Figure 4 along the directions: a) $K'_{2}-Z-K_{2}$; b) $K'_{2}-\Gamma-K_{2}$; c) $M'_{1}-Z-M_{1}$; d) $M'_{1}-\Gamma-M_{1}$ by DFT using a PBE functional (parameters given in method), showing the same anisotropy with respect to the Re chain direction.

Figure 7(a) shows the calculated energy of the highest energy VB and lowest energy CB states, with Fig. 7(b) showing the absolute CB minimum (CBM) and VB maximum (VBM) energies (dashed) and the local CBM and VBM values moving along the direction $\Gamma$ to $Z$, that is, as a function of $k_z$. We performed a full 3D calculation of the entire Brillouin zone in $k_x$, $k_y$ and $k_z$ to obtain this data. As noted above, conduction band effective masses obtained from parabolic fits to the DFT data at the Z point are given for the $k_x$ and $k_y$ directions in the Supplemental material (Fig. S4) [27]. From our calculations, we also obtained the electronic density of states for bulk ReS$_2$ (Supplemental material, Figure S3) which reproduces earlier calculations well [37, 38].

At $Z$ we obtain an estimate of the zero-temperature direct band gap as 1.525 eV. We note that a broad range of experimental band gaps exist from previous work from around 1.6eV [15] down to around 1.3eV[16, 28] (or lower in defect-rich materials or if defect states are probed[39]). In our calculations, the band gap value obtained by the GGA is the most reliable and is in good agreement with experimental values[28]. Interestingly, towards $\Gamma$ we find a transition to an indirect band gap. The conduction band minimum remains at $(k_x,k_y)=0$ (bottom left panel of Fig. 7a). We find the gap narrows at around $k_z=0.18$ to lower energy (1.49 eV) than at $Z$, as indicated by the arrow on Fig. 7(b). Moving from $Z$ to $\Gamma$ and calculating at $(k_x,k_y)=0$ (solid line on Fig. 7) we observe the valence band maximum to decrease by 125 meV for PBE (and 210meV for PZ). This is very similar to the 100-200 meV drop in the valence band maximum from $Z$ to $\Gamma$ observed experimentally and can be seen in Fig. 4.

This behavior may account for some of the uncertainty in the literature regarding the direct or indirect gap nature of bulk ReS$_2$, with other groups reporting electrical transport properties characteristic of indirect behavior in bulk samples [28, 40] or PL peaks within this range close
to the direct transition[7], whilst one group found a direct gap via electron energy loss spectroscopy of 1.42 eV at room temperature [22], consistent with the direct gap at 100 K of ~1.5 eV that we find [41]. Unlike here, previous calculations have often not considered the full bulk BZ and have instead focused on 2D or quasi-2D simulations of ReS$_2$ where calculations have predicted the gap to be direct at the 2D Γ point [15] in the monolayer form. As yet, only one brief study of ARPES measurements has been reported on a 2D monolayer of ReS$_2$ [11], and an important next step in this field will be to investigate few-layer structures in more detail. Use of a hybrid functional may also enhance the quality of the calculations. To confirm fully the nature of the band gap in the material it is necessary to measure the lowest states of the conduction band (potentially through alkali metal doping of the material). During the preparation of this paper, two more reports of ARPES studies of ReS$_2$ in bulk and thin-layer forms appeared on arxiv (both now published) which confirm the three-dimensional dispersion of the valence band structure [42] and, for the bulk material, indicate via Rb-doping that the band extrema are indeed located at Z as proposed above [43].

IV. CONCLUSIONS

In conclusion, we have performed ARPES measurements on the transition metal dichalcogenide ReS$_2$ in its bulk form. The anisotropic curvature of the valence band in directions perpendicular and parallel to the Re atomic chains can account for the reported anisotropic electrical properties (effective mass and mobility) of the material. We find the valence band maximum at $Z$ to be higher in energy than that at Γ and we obtain good agreement with DFT calculations of the band structure. Our calculations for the electronic bands over the whole bulk Brillouin zone predict a direct bandgap at $Z$ and a wider gap towards Γ with a small shift of 100-200 meV in the valence band maximum between these two points. This may account for uncertainty in the literature as to the direct or indirect nature of the band gap in bulk ReS$_2$. Future work to measure the conduction band directly (for example by doping to move the Fermi level into the conduction band) is required to investigate this more fully and also to map out the band structure of two-dimensional monolayer ReS$_2$.

ACKNOWLEDGMENTS

This work was supported by the Centre for Graphene Science of the Universities of Bath and Exeter and by the EPSRC (UK) under grants EP/G036101, EP/M022188 and EP/P004830; LSH is supported by the Bath/Bristol Centre for Doctoral Training in Condensed Matter Physics, grant EP/L015544. We thank the SOLEIL synchrotron for the provision of beam time; work at SOLEIL was supported by EPSRC grant EP/P004830/1. Computational work was performed on the University of Bath’s High Performance Computing Facility. We thank Dr Philip King of St Andrews University for useful discussions and for communicating his data [43] prior to publication. M.C.A., J.A. and C.C. thank Young-Hee Lee and Matthias Bantzill for enlightening exchanges. The Synchrotron SOLEIL is supported by the Centre National de la Recherche Scientifique (CNRS). Data created during this research are freely available from the University of Bath data archive at DOI:10.15125/BATH-00331.

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