Magnetic Weyl semimetals are a newly discovered class of topological materials that may serve as a platform for exotic phenomena, such as axion insulators or the quantum anomalous Hall effect. Here, we use angle-resolved photoelectron spectroscopy and ab initio calculations to discover Weyl cones in CoS₂, a ferromagnet with pyrite structure that has been long studied as a candidate for half-metallicity, which makes it an attractive material for spintronic devices. We directly observe the topologically non-trivial surface states that link the Weyl nodes, which will influence the performance of CoS₂ as a spin injector by modifying its spin polarization at interfaces. In addition, we directly observe a minority-spin bulk electron pocket in the corner of the Brillouin zone, which proves that CoS₂ cannot be a true half-metal.

**INTRODUCTION**

Since the experimental discovery of topological insulators and topological semimetals, there has been a substantial effort to functionalize topological materials for spintronic applications, most prominently in the family of Bismuth chalcogenides (1–4). However, because the field of spintronics predates the first prediction of topological insulators (5, 6) and topological semimetals (7), topological phenomena in many well-studied spintronic materials may have been overlooked during initial investigations of their electronic properties. Here, we reveal that CoS₂, a material that has been long studied because of its itinerant ferromagnetism and potential for half-metallicity, actually hosts Weyl fermions and Fermi arc surface states in its band structure close to the Fermi level $E_F$, as well as topological nodal lines below the Fermi level. CoS₂ is, therefore, a rare example of the recently discovered class of experimentally verified magnetic topological metals (8–10), which have been proposed to realize new mechanisms of spin-to-charge conversion (11), and are of broader interest for fundamental science, as a platform to realize axion insulators (7), the intrinsic anomalous Hall effect (12), or the anomalous fractional quantum Hall effect (13).

Beyond the discovery of its topological properties, we also clarify the question of whether CoS₂ is a true half-metal. Half-metallic ferromagnets, i.e., materials that are metallic in one spin channel but gapped in the other, have been considered as promising materials for spintronic devices, where they could act as sources of highly spin-polarized charge carriers. There has been a longstanding debate on whether CoS₂ or the related alloys Co₁−ₓFeₓS₂ are true half-metals, which has important implications for materials and device engineering. Calculations within the local spin density approximation (LSDA) (14, 15) [see Fig. 1 (A and B)] conclude that CoS₂ hosts a minority-spin electron pocket at the R point that leads to a peak in the minority-spin density of states (DOS), which would make CoS₂ a minority-spin conductor. This scenario is supported by magnetotransport experiments that suggest a sign flip in the spin polarization upon hole doping with iron that may turn CoS₂ into a half-metal (i.e., vanishing minority-spin DOS at $E_F$) due to the depopulation of a minority-spin electron pocket (16–20). On the other hand, calculations based on the generalized gradient approximation (GGA) (15, 21) [see Fig. 1 (B and C)] conclude that half-metallicity is already obtained in the undoped compound, and quantum oscillation experiments were unable to detect the putative minority-spin electron pocket (21). To resolve the debate about the half-metallicity of CoS₂, a direct spectroscopic band structure investigation would be clearly desirable, also because the spin polarization measured with transport probes can be reduced by surface defects (22) and can therefore not entirely rule out intrinsic half-metallicity. It should be noted that finite temperature effects can also reduce the spin polarization of nominally half-metallic ferromagnets (23).

Besides the bulk band structure, electronic surface states also influence the spin polarization at the Fermi level, which becomes particularly important for heterostructure interfaces in spintronic devices where half-metals could function as spin injectors. One important class of surface states are topical Fermi arcs in Weyl semimetals, which are considered to be particularly robust against passivation due to the protection by topological bulk invariants. Our LSDA calculations of CoS₂ shown in Fig. 1 (A and B) predict Weyl nodes close to the Fermi level on the $k_x = 0$ and $k_y = 0$ planes...
parallel to the magnetization direction (see Fig. 1D), which should give rise to topological Fermi arc surface states. In addition, we find topological nodal lines a few hundreds of milli–electron volts below the Fermi level, which are protected by a \( \{ ½ 0 ½ \} \) glide operation (also see the Supplementary Materials). Since the concept of topological semimetals and Fermi arc surface states was not yet established at the time, previous theoretical studies of CoS\(_2\) overlooked these important features in its band structure, which can have a decisive influence on spin transport properties.

Previous angle-resolved photoelectron spectroscopy (ARPES) experiments (24–26) on CoS\(_2\) were performed with photon energies between \( h\nu = 20 \) to 120 eV where the inelastic mean free path of the photoelectrons can be expected to be close to its minimum \(-0.5\) to \(-1\) nm, resulting in very surface sensitive probes. This implies that photoelectrons can be expected to be close to its minimum \(-0.5\) to \(-1\) eV around \(-0.65\) eV by the exchange splitting. The magnitude of the exchange splitting extracted from the energy distribution curve at the R point (Fig. 2D) is \( \Delta E = 0.60\) eV (3). Our LSDA calculations of the band dispersion shown in Fig. 2E are in good qualitative agreement with the experimental data and indicate that the observed electron pocket is of minority-spin character, which implies that CoS\(_2\) is not a true half-metal. However, the experimentally observed electron pocket is of minority-spin character and therefore affect the spin polarization at a heterostructure interface.

### RESULTS

Our samples were synthesized using chemical vapor transport. We studied their elemental composition with core-level spectroscopy and energy-dispersive x-ray spectroscopy, which confirmed the expected stoichiometry (see Fig. 1E). Powder x-ray diffraction confirmed the previously reported cubic space group 205 and a lattice constant of \( a = 5.5287\) (5) (see Materials and Methods and the Supplementary Materials for more details about synthesis and characterization). We measured a Curie temperature of \( T_c = 124\) K and a saturation magnetization of \( 0.91 \mu_B/\text{Co} \) at 9 T (which becomes linear at 0.2 T and 0.89 \( \mu_B \), in good agreement with the literature values (28) (see Fig. 1F for magnetization curves). The coercivity in our data is small, about 10 to 15 Oe, which agrees also with the literature values (16).

When cleaving CoS\(_2\) for ARPES experiments, we obtained two distinct cleavage planes with the surface normal pointing along the (111) and (100) directions. We performed photon energy–dependent ARPES measurements with soft x-ray photons (\( h\nu = 350 \) to 800 eV) to locate the high-symmetry planes along the \( k_z \) direction normal to the sample surface (see the Supplementary Materials). The band structure in the \( k_z = \pi \) plane containing the R point is for the (111) surface displayed in Fig. 2 (see fig. S9 for a definition of the high-symmetry planes). We can clearly identify circular Fermi surface pockets at the R point in the corner of the Brillouin zone in Fig. 2A. Our calculated Fermi surface (Fig. 2B; using the LSDA) is in good qualitative agreement with the experimental data, confirming the existence of Fermi surface pockets at the R point. When inspecting the experimental band dispersion along the R–X–R direction (Fig. 2C), we see that the circular pockets at the R point are electron-like and are related to another parabolic band with a minimum at around \(-0.65\) eV by the exchange splitting. The magnitude of the exchange splitting extracted from the energy distribution curve at the R point (Fig. 2D) is \( \Delta E = 0.60\) eV (3). Our LSDA calculations of the band dispersion shown in Fig. 2E are in good qualitative agreement with the experimental data and indicate that the observed electron pocket is of minority-spin character, which implies that CoS\(_2\) is not a true half-metal. However, the experimentally observed exchange splitting is \(~250\) meV smaller than in the LSDA calculations, such that the majority spin bands are located closer to the Fermi level in the experiment than expected from the calculations. The data measured on the (100) surface also show electron pockets at the R point and are displayed in the Supplementary Materials. To search for the topological nodal line and Weyl nodes in CoS\(_2\), we also probed the bulk band structure in the \( k_z = 0 \) plane, containing the \( \Gamma \) point, as illustrated in Fig. 3. Figure 3 (A to D) displays the experimental and calculated Fermi surfaces for the (111) and (100) cleavage planes, which are in good qualitative agreement. Figure 3E shows the band dispersion along the M–\( \Gamma \) direction (black arrow in
Fig. 2. Bulk band structure of the $k_z=\pi$ plane measured on the (111) surface. (A) Experimental Fermi surface was measured on the (111) cleavage plane with photon energy $h\nu = 602$ eV and linear-vertical polarization, integrated over 50 meV below the Fermi energy. The red arrow indicates an electron pocket located at the $R$ point. The black arrow indicates the position of the line cut shown in (C). (B) Calculated Fermi surface spectral function $A(k,E_F)$ for the same plane as shown in (A) obtained with the LSDA. (C) Line cut along the $R-X-R$ direction as shown in (A); red arrows indicate the electron pockets at the $R$ point. (D) Energy distribution curves for the two $R$ points shown in (E). The black arrow indicates the magnitude of the exchange splitting of $\Delta E = 0.60(3)$ eV. (E) Calculated band structure obtained with LSDA; red arrows indicate minority-spin electron pocket.

Fig. 3A), measured on the (111) surface (black arrow in Fig. 3A), and the $\Gamma-M$ direction, measured on the (100) surface (black arrow in Fig. 3C). Note that the observable bands along these two directions are very different, possibly due to matrix element effects. The line cut obtained from the (111) surface shows a V-shaped feature centered at the M point and a quasi-parabolic band centered at the $\Gamma$ point. In contrast, the dispersion obtained from the (100) surface shows a single band dispersing in the opposite direction from the quasi-parabolic band. To enhance the contrast of our data, we also show the corresponding second derivative spectrum in Fig. 3F. The combined band dispersion from both surfaces is illustrated in Fig. 3G, which displays the peak positions from a fit of the momentum distribution curves. By comparison with the calculated band dispersion shown in Fig. 3H, we can see that the band crossing between the blue and red bands [from the (111) surface and (100) surface, respectively] is part of a topological line node, while the blue bands form a Weyl cone. The Weyl point that corresponds to the Weyl cone is shown in Fig. 3l, which displays the calculated band dispersion along the $M^*-\Gamma$ direction, where $M^*$ is a point that is slightly displaced from $M$ (0, 0.5, 0.4581) in $k$ space. This is identical to the $M$ point within the experimental uncertainty. Since we cannot observe the band top of the blue bands along the M-\Gamma direction in our experimental data, we conclude that the Weyl point must be located slightly above the Fermi level.

Fermi arc surface states are a hallmark of Weyl points in topological semimetals. Therefore, the Weyl points in CoS$_2$ must be accompanied by Fermi arc surface states that are connecting the projections of the Weyl points in the surface Brillouin zone. We used surface-sensitive VUV-ARPES to investigate the surface electronic structure of the (100) surface in CoS$_2$, the results of which are displayed in Fig. 4. Figure 4A shows the experimentally obtained Fermi surface, which was measured on a strongly tilted crystal plane. The photon energy dependence of the Fermi surface maps (see the Supplementary Materials) indicates that all Fermi surface pockets measured by VUV-ARPES are surface states and that the signal from bulk states is mostly suppressed in this photon energy range. The positions of the projections of the Weyl point in the (100) surface Brillouin zone are shown in Fig. 1D. From our ab initio calculations, we expect that there should be two Weyl cone projections with total Chern numbers $C = \pm 2$ in the vicinity of the $X$ point of the surface Brillouin zone, which originate from the Weyl cones in the vicinity of the $M$ point of the bulk Brillouin zone that we have experimentally observed in Fig 3 (E to G). These two Weyl points have to be connected by two Fermi arcs, which can form a circular Fermi surface (see Fig. 4B). This relationship between Weyl cone projections and topological surface states is precisely what we observe in our experimental data in Fig. 4A, which shows small circular Fermi surface pocket centered at the $X$ point. A corresponding line cut of the experimental data along the $X-\Gamma-X$ path is shown in Fig. 4C. It displays a surface state band crossing the Fermi level that connects two hole-like pockets that are located at the $X$ points at opposite ends of the Brillouin zone. Our ab initio calculations of the surface electronic structure displayed in Fig. 4D confirm that this surface state is a Fermi arc that connects two Weyl points. Note that the renormalized energy scale of the Fermi arcs in the calculation compared to the experiment is expected owing to the reduced exchange splitting, which we already observed for the bulk band structure. Therefore, the Fermi arcs cross the Fermi level in the experiment, while they are located below the Fermi level in the calculation. Figure 4 (E to G) shows that the same line cut is measured at different photon energies, which shows that the Fermi arc dispersion is independent of the $k_z$ momentum, which proves its two-dimensional nature. Since the Fermi arcs are derived from majority-spin bulk bands, they have majority-spin character (see the Supplementary Materials for calculations of their spin polarization). This will influence the spin polarization of electrons at the Fermi level at heterostructure interfaces where CoS$_2$ can act as a spin injector.

DISCUSSION

One may wonder whether the appearance of the conical bulk-band pockets connected by surface states could just be a coincidence, and whether the circular surface state pocket observed here may actually be just a trivial surface state. However, such a coincidence would require an unrealistic degree of fine-tuning, because there are no symmetry constraints that should pin a trivial surface state to the projections of the conical bulk bands. The only plausible explanation is, therefore, that the surface states mentioned above are Fermi arcs, and the observed linear bulk bands are part of a Weyl cone. Although we cannot directly observe the Weyl point since it is located...
in the unoccupied part of the DOS, the overall agreement between the experimental data and LSDA calculations in conjunction with experimentally observed momentum space alignment between surface states and bulk bands provides sufficient evidence to conclude that CoS₂ is a magnetic Weyl semimetal.

Because of the relatively low effective mass of the minority-spin electron pocket at the R point, the minority-spin DOS at the Fermi level is large, and some calculations suggested that CoS₂ is a minority-spin conductor (18). Our discovery of the majority-spin Fermi arc surface states implies that the total interface spin polarization of a heterostructure involving CoS₂ will be reduced compared to the bulk value. Compared to trivial dangling bond surface states, Fermi arc surface states are more robust against attempts of passivation, because they are protected by the topological invariants (the Chern numbers) of the bulk Weyl points; hence, engineering of the interface potential cannot easily remove the Fermi arcs. On the other hand, hole doping with iron has been suggested to transform CoS₂ from a minority-spin to a majority-spin conductor and ultimately to a full half-metal by depopulation of the minority-spin bulk electron pocket (18). Such a transition will be facilitated by the majority-spin Fermi arcs at interfaces in heterostructures since they compensate the DOS of the minority-spin electron pocket. Therefore, CoS₂ provides a prime example of a spintronic material whose performance is affected by its topologically nontrivial band structure. Note that nonmagnetic Weyl semimetals have already been successfully implemented for applications in spin-to-charge conversion and magnetic switching (29, 30), and theoretically, proposals have highlighted the importance of topological Fermi arcs for interfacial spin accumulation (31). We therefore expect that our results will inform the design of future spintronic prototypes that use magnetic Weyl-semimetals, for instance, for spin-to-charge conversion (11). There may be magnetic domains on the sample surface that are smaller than the size of our beamspot (diameter of 50 to 70 µm), but on the basis of our ab initio calculations, we believe that the direction of the magnetization vector will have a negligible effect on the interpretation of the Fermi arc structure. This is because spin-orbit coupling is relatively small in CoS₂, and its inclusion has only a negligible effect on the band dispersion in our ab initio calculations (see the Supplementary Materials for further discussion). CoS₂ is also known as a good catalyst, e.g., for the hydrogen evolution reaction (32), and it has recently been speculated that Fermi arcs in Pt- and Pd-based chiral topological semimetals (33, 34) may play a role in catalysis due to their d-electron character and their robustness against hydrogen passivation (35, 36). Since the Fermi arcs in CoS₂ are also derived from bulk bands of d orbital character (see the Supplementary Materials), they may contribute to the catalytic performance of CoS₂.

**Fig. 3. Bulk band structure of the k_z = 0 plane.** (A) Experimental Fermi surface measured on the (111) cleavage plane with photon energy hν = 512 eV and linear-vertical polarization, integrated over 50 meV below the Fermi energy. The black arrow indicates the position of the line cut shown in (E). (B) Calculated Fermi surface spectral function for the same plane as shown in (A) obtained with LSDA. (C) Experimental Fermi surface spectral function δk(E_F) measured on the (100) cleavage plane with photon energy hν = 475 eV and linear-vertical polarization, integrated over 50 meV below the Fermi energy. The black arrow indicates the position of the line cut shown in (E). (D) Calculated Fermi surface for the same plane as shown in (C) obtained with LSDA. (E) Line cuts along the M–Γ direction from the (111) surface, as shown in (A), and the Γ–M direction from the (100) surface, as shown in (C). (F) Second-derivative spectrum of (E). (G) Result of the momentum distribution curve (MDC) fitting of the bands along the M–Γ–M direction, where blue circles originate from data of the (111) plane, and red circles originate from data of the (100) plane. (H) Calculated band dispersion along the M–Γ direction. (I) Band dispersion along the M*–Γ direction where M* (0,0,0.4581)Å⁻¹ is a point slightly displaced from the M point (see Fig. 1D), such that the k-path is passing through the Weyl point in the vicinity of M.
**MATERIALS AND METHODS**

**Sample growth**

Single crystals of CoS$_2$ were synthesized using chemical vapor transport. The elements cobalt (200 mg, 1 eq, 99.5%; Sigma-Aldrich) and sulfur (217 mg, 2 eq, 99.5%; Alfa Aesar) were mixed with 50 mg of iodine (99.999%; Sigma Aldrich) and sealed in an evacuated quartz glass ampoule. The ampoule was slowly (within 12 hours) heated to 1000°C and kept there for 140 hours. The growth was along a natural temperature gradient of a tube furnace. An increase in crystal size can be accomplished using an additional slow cooling step from 1000°C to 800°C in 100 hours.

**ARPES measurements**

SX-ARPES measurements were performed at the SX-ARPES endstation (37) of the ADRESS beamline (38) at the Swiss Light Source, Switzerland, with a SPECS analyzer with an angular resolution of 0.1°. The photon energy varied from 350 to 1000 eV, and the combined energy resolution ranged between 50 and 150 meV. The temperature during sample cleaving and measurements was below 20 K, and the pressure was lower than 1 × 10$^{-10}$ mbars. The increase of the photoelectron mean free path in the soft x-ray energy range results, by the Heisenberg uncertainty principle, in a higher $k_z$ resolution of the ARPES experiment compared to measurements at lower photon energies (39), which was critical to measure the bulk band structure of CoS$_2$.

VUV-ARPES measurements were performed at the high-resolution ARPES branch line of the beamline 105 at the Diamond Light Source, UK (40). Measurements at the high-resolution branch were performed with a Scienta R4000 analyzer, and a photon energy range between 90 and 130 eV, at a temperature below 20 K. Measurements in the VUV-ARPES regime are more surface sensitive than SX-ARPES and therefore most suitable to image the Fermi arcs in CoS$_2$.

**Ab initio calculations**

We used density functional theory as implemented in the Vienna Ab Initio Simulation Package. For the GGA calculations, the exchange correlation term is described according to the Perdew-Burke-Ernzerhof prescription together with projected augmented wave (PAW) pseudopotentials, while for the LSDA, the exchange correlation term is described according to the Dudarev simplified, rotationally invariant approach, together with PAW pseudopotentials. The kinetic energy cutoff was set to 400 eV. For the self-consistent calculation, a grid of 7 × 7 × 7 k points was used. For DOS calculation, a grid of 11 × 11 × 11 k points was used, with 1000 energy points. The spectral function of the Fermi surfaces shown in Figs. 2 and 3 was computed with the programs Wannier90 (41) and WannierTools (42). Using ab initio calculation of bands, we constructed maximally localized Wannier functions considering d orbitals for cobalt sites and p orbitals for sulfur sites using the Wannier90 software (41). We computed an effective tight-binding model that accurately reproduces the band structure of the bulk system. We then computed the surface spectrum by the method of iterative Green’s function as implemented in WannierTools (42) software for a slab of 50 unit cells.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/51/eabd5000/DC1

**REFERENCES AND NOTES**


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