

CHCRUS

This is the accepted manuscript made available via CHORUS. The article has been published as:

Strongly anisotropic antiferromagnetic coupling in math xmlns="http://www.w3.org/1998/Math/MathML">mrow>ms ub>mi>EuFe/mi>mn>2/mn>/msub>msub>mi>As/mi>mn >2/mn>/msub>/mrow>/math> revealed by stress detwinning

Joshua J. Sanchez, Gilberto Fabbris, Yongseong Choi, Yue Shi, Paul Malinowski, Shashi Pandey, Jian Liu, I. I. Mazin, Jong-Woo Kim, Philip Ryan, and Jiun-Haw Chu Phys. Rev. B **104**, 104413 — Published 10 September 2021 DOI: 10.1103/PhysRevB.104.104413

1	Strongly anisotropic antiferromagnetic coupling in
2	EuFe ₂ As ₂ revealed by stress detwinning
3	Authors: Joshua J Sanchez ¹ , Gilberto Fabbris ³ , Yongseong Choi ³ , Yue Shi ¹ , Paul Malinowski ¹ , Shashi
4	Pandey ² , Jian Liu ² , I.I. Mazin ⁵ , Jong-Woo Kim ³ , Philip Ryan ^{3,4} , Jiun-Haw Chu ^{1*}
5	Affiliation:
6	¹ Department of Physics, University of Washington, Seattle, Washington 98195, USA.
7	² Department of Physics and Astronomy, University of Tennessee, Knoxville, Tennessee 37996, USA.
8	³ Advanced Photon Source, Argonne National Laboratories, Lemont, Illinois 60439, USA.
9	⁴ School of Physical Sciences, Dublin City University, Dublin 9, Ireland.
10	⁵ Department of Physics and Astronomy and Quantum Science and Engineering Center, George Mason
11	University, Fairfax, VA 22030
12	*Correspondence to: jhchu@uw.edu (JH.C)
13	

15 Abstract:

16	Of all parent compounds of iron-based high-temperature superconductors, EuFe ₂ As ₂ exhibits by far the
17	largest magnetostructural coupling due to the sizable biquadratic interaction between Eu and Fe
18	moments. While the coupling between Eu antiferromagnetic order and Fe structural/antiferromagnetic
19	domains enables rapid field detwinning, this prevents simple magnetometry measurements from
20	extracting the critical fields of the Eu metamagnetic transition. Here we measure these critical fields by
21	combining x-ray magnetic circular dichroism spectroscopy with in-situ tunable uniaxial stress and
22	applied magnetic field. The combination of two tuning knobs allows us to separate the stress-
23	detwinning of structural domains from the field-induced reorientation of Eu moments. Intriguingly, we
24	find a spin-flip transition which can only result from a strongly anisotropic interaction between Eu
25	planes. We argue that this anisotropic exchange is a consequence of the strong anisotropy in the
26	magnetically ordered Fe layer, which presents a new form of higher-order coupling between Eu and Fe
27	magnetism.
28	
29	

Magnetism is the origin of a wide range of intriguing phenomena in iron-based superconductors, including electronic nematicity and high temperature superconductivity [1–5]. In contrast to the magnetism of the high- T_c cuprates, the stripe spin density wave (SDW) ground state breaks fourfold rotational symmetry and the spin dynamics are highly anisotropic [6–9]. Key to this highly anisotropic magnetism is a sizable biquadratic coupling that is not captured in a simple Heisenberg model and likely arises from the dual itinerant-localized character of the Fe magnetism [10,11]. It has also been suggested that this biquadratic term is necessary for the emergence of spin nematicity [12,13].

39 Very recently, it was shown that a similar biquadratic coupling plays a role in the unusually large 40 magnetoelastic coupling in EuFe₂As₂, where the introduction of a magnetic lanthanide element adds 41 another layer of complexity into the magnetism of iron-based superconductors [14]. In addition to the 42 SDW order, EuFe₂As₂ also hosts an A-type layered antiferromagnetic (AFM) order in the Eu layer, sharing the same easy axis with the Fe-SDW. Like in other iron pnictides, the SDW in EuFe₂As₂ creates 43 44 orthorhombic twin domains. In BaFe₂As₂, due to the strong coupling between the structural distortion 45 and the SDW, an applied field of order 25 T is able to fully detwin the structural domains [15]. Surprisingly, 46 EuFe₂As₂ can be fully detwinned with less than 1 T, and partial detwinning can persist even after the field 47 is turned off [16,17]. Magnetization, NMR and neutron diffraction data show that this structural 48 detwinning coincides with the reorientation of Eu moments towards the applied field direction, suggesting 49 that the Eu magnetism and the associated large magnetic moments are responsible for this drastic 50 reduction of detwinning field [16,18,19]. Nevertheless, as no single-ion anisotropy is present for the halffilled Eu 4f⁷ electrons and no dipolar coupling between the Fe and Eu layers is allowed by symmetry, it 51 52 remained an open question how Eu moments even sense the orthorhombic direction. Recently, Maiwald et al solved this mystery by considering a biguadratic coupling between Fe-SDW and Eu-AFM 53 moments [14]. The biquadratic coupling of the form $K(f_i \cdot e_j)^2$, where f_i and e_i represent the Fe and Eu 54 55 moments, respectively, provides an effective single-ion anisotropy that couples the Eu moment orientation with the Fe-SDW direction [14]. Therefore, while the Fe-Fe biquadratic coupling generates the
nematicity in the FeAs layer, the Eu-Fe biquadratic coupling provides a pathway for the Eu magnetism to
couple to the structural orthorhombicity and the underlying nematicity.

59 Here, we report the discovery of another consequence of the higher order Eu-Fe coupling in 60 $EuFe_2As_2 - a$ highly anisotropic Eu-Eu interplanar coupling. The degree of anisotropy of Eu-Eu interlayer exchange $\left(\frac{J_x - J_y}{J_x + J_y}\right)$ is about 75 times larger than the structural orthorhombicity, which can only be 61 understood by considering the influence of the Fe-SDW order. The anisotropy of Eu-Eu interlayer 62 63 exchange was overlooked previously because the field detwinning process masked the spin-flip nature of 64 Eu metamagnetic transition. We overcome this challenge by a direct measurement of the Eu 65 metamagnetic transition in a mechanically detwinned sample using a piezoelectric stress device, which 66 allows us to apply magnetic fields either parallel or perpendicular to the easy axis of Eu moments within 67 a single structural domain. Conventional magnetometry techniques are difficult to apply to a sample mounted to a strain device due to the added size and background magnetization contributed by the 68 69 device. We therefore employ x-ray magnetic circular dichroism (XMCD) on the Eu L₃ edge to measure the 70 Eu-specific in-plane magnetization induced by an applied magnetic field. We show that we are able to 71 strain the crystal into a monodomain which exhibits either a large jump in magnetization, consistent with 72 a spin flip transition, or has a perfectly linear magnetization from continuous canting of Eu moments (i.e., 73 we are able to turn the metamagnetic transition on and off). From the measurement of the critical field 74 for the Eu spin flip and the field dependence of the spin canting, we determine the energies of the Eu-Eu 75 and Eu-Fe coupling and discover that the Eu-Eu interaction itself is directionally dependent on the 76 orientation of Fe moments. We then confirm this by first principles calculations. The discovery of the 77 anisotropic Eu-Eu interplanar coupling also sheds light on the evolution of the Eu magnetism in doped

EuFe₂As₂ which we reevaluate in the discussion. Finally, simultaneous transport measurements suggest a
 unique Eu origin of the large magnetoresistance jump previously found in the system [17,20].

80 **RESULTS**:

81 Figure 1a shows the fully magnetically ordered unit cell at T=7 K and zero applied magnetic field. 82 Eu moments are aligned ferromagnetically within each plane and antiferromagnetically between planes. 83 The Fe spin density wave ordering results in a small structural orthorhombicity and the formation of 84 structural twin domains that are identical up to a 90-degree rotation. Within each domain, the Eu AFM 85 and Fe SDW easy axes are aligned with the longer orthorhombic a lattice constant [21,22]. We use a 86 geometry with the \hat{x} and \hat{y} axes aligned to the orthorhombic a and b directions, and tensile or 87 compressive stress applied along the \hat{x} direction detwins the sample toward the A monodomain (a lattice 88 vector along \hat{x}) or B monodomain (b lattice vector along \hat{x}), respectively (Fig.1b). Given the maximum 89 orthorhombicity of EuFe₂As₂ (0.28% at 2.5 K [22]) and the maximum strain of the sample device at 7 K 90 (0.3%), we are capable of nearly fully detwinning the sample (the minor domain is estimated to be 5% or 91 less of the sample volume, see Supplementary Figure 2 [23] and ref. [24]). Once in the A (B) monodomain state, the measured resistivity ρ_{xx} becomes sensitive to the anisotropic resistivity ρ_a (ρ_b) along the 92 93 orthorhombic a(b) direction. Magnetic field is applied perpendicular to the current/strain axis and at 10 94 degrees above the ab plane. Except for a change in the strain state, the sample is not reoriented in any 95 way during the experiment, ensuring identical effective fields and XMCD-illuminated sample volumes. 96 XMCD measures the induced Eu magnetic moment along the field direction, which is fixed parallel to the 97 incident x-ray direction in this study.

First, we address the effect of the applied magnetic field at T=30K, below the orthorhombic and SDW transitions ($T_{SDW} = 187K$), but above the Eu AFM ordering temperature ($T_N = 19.1K$). In the Eu²⁺ valence state, the $4f^7$ electrons have zero orbital angular momentum (L = 0), and as such are expected

101 to show an isotropic response to applied field. After detwinning to either the A or B monodomain, we 102 applied fields from 0 to 1 T and measured the XMCD and resistivity simultaneously in 0.02 T steps (XMCD 103 data were not collected for the B monodomain for 0.8 T-1 T at this temperature). We find an XMCD signal 104 that is indistinguishable between the two domains, suggesting that at this temperature and field range 105 the Eu-Fe interaction is negligible compared to the Eu paramagnetic coupling to applied field (Fig.2). We 106 note that compared to the data presented next, the XMCD values at 1 T and 30 K are roughly 3 times 107 smaller than the 1 T XMCD saturation value within the Eu AFM phase, consistent with a lower 108 susceptibility in the paramagnetic phase. As in BaFe₂As₂, the zero-field resistivity is considerably larger 109 along the orthorhombic b direction than the a direction [25–27]. Both ρ_a and ρ_b have a weak field 110 dependence at 30K. The inset to Fig. 2 shows the detwinned sample cooling through the Eu AFM 111 transition. As demonstrated previously in mechanically detwinned EuFe₂As₂, we see no change in the Eu 112 transition temperature between the tensile and compressive cooling data, nor do we see any additional resistivity anisotropy induced by the Eu AFM ordering [28]. Indeed, the resistivity anisotropy $\frac{\rho_b - \rho_a}{\rho_b + \rho_a}$ 113 114 0.084(2) is unchanged through this temperature range.

115 We now discuss results from within the Eu AFM phase at 7K. We applied field through a ± 1 T loop 116 to each detwinned state and measured the XMCD and resistivity simultaneously in 0.02 T steps (Fig.3). In 117 the A domain (Fig.3a), the Eu moments are initially aligned with the AFM easy axis transverse to the field 118 direction. The linear growth of XMCD signal with field indicates that Eu moments cant continuously to 119 align with the field, with no observable hysteresis. Conversely, in the B domain (Fig.3b) the easy axis is 120 along the field direction, and so for fields below 0.4 T the XMCD is nearly flat as no canting can occur. The 121 jump in XMCD from 0.4 T to 0.6 T and accompanying hysteresis is a clear sign of a metamagnetic spin flip 122 transition between the Eu AFM and FM states. The weak linear field dependence of magnetization in the 123 pre- and post-spin-flip field ranges are due to the out of plane magnetization induced by the small out of 124 plane field component, which is assumed to equally contribute to the A domain magnetization. We note

that the observation of spin flip transition contradicts the expectation from the spin Hamiltonian derived from ref [14], which predicts a spin flop transition. As will be shown in the discussion section, this contradiction can only be resolved by including an anisotropic interlayer exchange between Eu moments.

128 The resistivity is approximately linear in field for $|\mu_0 H| < 0.4T$ and $|\mu_0 H| > 0.6T$ for both domains. For $0.4T < |\mu_0 H| < 0.6T$, a large hysteretic drop occurs in the B domain resistivity, coinciding 129 130 with the jump in XMCD. A much smaller drop also occurs in the A domain, which also shows a small 131 hysteresis (Fig.3a, inset), and is likely due to a remnant B domain that was not fully detwinned and which 132 is not resolved in the XMCD measurement. For each domain, the resistivity returns to the initial zero-field 133 value after the field loop, indicating that there is no persistent field-detwinning in our setup as is found 134 reliably from previous studies of freestanding samples [16,17]. Further, given the field and current 135 orientation, the sample in either strain state would be expected to detwin towards the higher-resistivity 136 B domain above ~0.5T, and so the drop in resistivity for both domains suggests that our stress device is 137 indeed preventing field detwinning. This is also strong evidence for a purely Eu spin origin of the resistivity 138 jump.

139 Both the XMCD and resistivity data show excellent agreement between positive and negative field 140 values. In Figure 4a, we plot the average value over positive and negative field sweeps of the XMCD and 141 the magnetoresistance. While we were unable to apply enough field to fully saturate the A domain XMCD, 142 we can extrapolate the field dependencies of each domain to estimate the saturation field. Linear fits to the XMCD of the A and B domains at fields greater than 0.6 T are shown, with an intersection at $\mu_0 H_4^{sat}$ 143 144 = 1.17 T. Beyond this field, the Eu magnetic moment is expected to be fully saturated in each domain, as 145 seen in freestanding crystal magnetometry studies [14,17,18,29]. To more precisely determine the B domain spin flip field H_B^{flip} we use the value at the center of the field hysteresis. Figure 4b shows the 146 147 difference between the increasing and decreasing field values of the XMCD and resistivity. In the B domain, a sharp peak in both quantities occurs at $\mu_0 H_B^{flip} = 0.48T$. In principle, the lattice distortions induced by the detwinning strain could cause a change in the interaction strengths between Eu and Fe planes that could alter these critical fields. In Supplementary Figure 3 we present data for the same sample tuned to a nearly-zero strain state in which both A and B domains are present and show that H_A^{sat} and H_B^{flip} are essentially unchanged, demonstrating that the strain applied to detwin the sample does not appreciably affect the interplanar coupling strengths.

154

155 DISCUSSION

We can relate the spin flip and saturation fields H_{R}^{flip} and H_{A}^{sat} to the microscopic interactions in 156 157 the sample using a spin Hamiltonian (Supplementary Information section IV). We will start with a modified version of the spin Hamiltonian presented in ref. [14] and show that to explain the experimental results 158 159 an additional anisotropic exchange term is needed. We consider first a purely isotropic exchange energy 2J between Eu antiferromagnetic planes in the doubled unit cell of the fully ordered state ($Eu_2Fe_4As_4$). 160 From the symmetry of the magnetic ordering structure (Fig. 1a), the dipolar interactions of Eu and Fe 161 moments cancel each other and so do not contribute to the magnetic energy. The half-filled $4f^7$ orbital 162 of Eu^{2+} has zero orbital angular momentum (L = 0) and negligible single ion anisotropy, but the posited 163 164 biguadratic coupling between Eu and Fe moments creates an effective magnetocrystalline anisotropy. We 165 define a biquadratic coupling energy K between the 8 inequivalent Eu-Fe moment pairs, with a total Eu-Fe planar coupling energy 8K. Following the usual treatment of spin-flip and spin canting transitions we 166 167 determine the critical fields as

168
$$H_B^{flip} = \frac{2J}{M}$$

169
$$H_A^{sat} = \frac{4J + 16K}{M} = 2\left(H_B^{flip} + \frac{8K}{M}\right)$$

For the B domain, Eu and Fe moments are aligned both before and after the spin flip, and so the presence of the biquadratic coupling does not change the value of the spin flip field, but only serves to provide the necessary magnetocrystalline anisotropy to enforce a sharp spin flip transition. For the A domain, the biquadratic coupling provides an extra energy barrier that must be overcome by the field to reach the fully saturated canted state. From the measured values $\mu_0 H_A^{sat} = 1.17$ T and $\mu_0 H_B^{flip} = 0.48$ T, and using the expected Eu moment $M = 6.8 \mu_B$ [16,17], we obtain $J = 94.5 \mu eV$ and $8K = 41.3 \mu eV$. Thus, the Eu-Eu and Eu-Fe planar coupling energies are comparable, $\frac{J}{8K} = 2.3$.

177 At this point we are facing an apparent contradiction: we observe experimentally a sharp spin flip transition, but for $\frac{J}{8K} > 1$ the spin Hamiltonian would actually be expected to result in a spin flop 178 179 transition, as shown in Fig. 5(a). (See ref. [14] and Supplementary Information section IV for further 180 discussion). This discrepancy has important implications for correctly modelling the field detwinning 181 process, which has been based on the (never actually observed) spin flop transition. This apparent 182 contradiction indicates that an additional term is needed in the spin Hamiltonian. The simplest such term is a symmetric anisotropic exchange term $W(e_{i,x}e_{i+1,x} - e_{i,y}e_{i+1,y})$, where $e_{i,x}$ and $e_{i,y}$ are the x and y 183 components of the Eu moment in the *i* th layer, using a notation with Fe moments aligned 184 185 antiferromagnetically along \hat{x} . This term then increases (decreases) the interaction strength between Eu 186 moments when aligned parallel (perpendicular) to the Fe moments, and provides the additional 187 anisotropy needed to enforce a spin flip transition. Although other higher order terms could also be 188 introduced to the spin Hamiltonian, they generally lead to non-linear M-H curves which were not observed in the experiment. With this additional term, the criterion for a spin flip transition becomes $\frac{J}{2K+W} < 1$, 189 190 and the critical fields are:

191
$$H_B^{flip} = \frac{2(J+W)}{M}$$

192
$$H_A^{sat} = \frac{4J + 16K}{M} = 2\left(H_B^{flip} + \frac{8K - 2W}{M}\right).$$

The three parameters *J*, *K* and *W* cannot be uniquely determined by the two experimentally measured values. Therefore, additional constraint is needed. This constraint can be provided by the measurement of $H_{45^\circ}^{sat}$, i.e. the field required to saturate the magnetization when the field is aligned 45 degrees to the easy and hard axis, such that the field has an equivalent effect on both domains. It has the following expression:

$$H_{45^{\circ}}^{sat} = \frac{4J}{M}$$

199 As our strain+XMCD measurements have demonstrated the need for the W term, we now discuss the 200 extraction of I, K and W from magnetization measurements using a vibrating sample magnetometer 201 (VSM) on a second sample from the same growth batch. We stress that this extraction is not possible 202 without the confirmation of a spin-flip transition by XMCD measurement on a stress-detwinned sample. 203 The sample was encased in GE varnish so that the domain configuration is fixed to 50/50 and the field 204 detwinning is prohibited. The thin octagonal shape of the sample also ensures that field applied totally in-205 plane along the $\begin{bmatrix} 1 & 0 \end{bmatrix}_T$ and $\begin{bmatrix} 1 & 1 \end{bmatrix}_T$ directions have an identical (negligible) demagnetization factor (see 206 Methods). As shown in Figure 5b, at T=2 K for field applied along the $[1 \ 0 \ 0]_T$ direction (blue curve), i.e. 207 at 45° to both domain easy axes, the magnetization exhibits continuous spin canting toward saturation at $\mu_0 H_{45^\circ}^{sat} = 0.73$ T. For field applied along the $[1\ 1\ 0]_T$ direction (red curve) the *M* vs *H* perfectly overlaps 208 209 with the combination of the responses of A and B domains from the XMCD measurements (grey), confirming the absence of field detwinning. We extract $\mu_0 H_A^{sat} = 1.15 \text{ T}$ and $\mu_0 H_B^{flip} = 0.45 \text{ T}$ from 210 211 these spin flip and saturation fields, which is in good agreement with the XMCD-measured values. Using 212 these 3 measured values, we can uniquely solve for the interaction terms and find $J = 71.8 \, \mu eV$, W=

16.7 μeV and 8 K = 82.7 μeV , with a ratio $\frac{J}{8K+W} = 0.72 < 1$ satisfying the sharp spin flip criterion. Therefore, we find that the Eu-Eu interaction is much stronger for Eu moments aligned parallel ($J_x = J + W = 88.5 \ \mu eV$) compared to perpendicular ($J_y = J - W = 55.1 \ \mu eV$) to Fe moments. Further, the normalized difference of the anisotropic interplanar interaction, $\frac{J_x - J_y}{J_x + J_y} = 23.3\%$, is nearly two orders of magnitude greater than the corresponding normalized difference of in-plane lattice constants (the orthorhombicity), $\frac{a-b}{a+b} \sim 0.28\%$, which strongly implies the Fe-SDW origin of the anisotropy.

219 To gain more insight, we used density functional theory (DFT) to calculate the exchange coupling 220 between Eu layers as the difference between ferro- and antiferromagnetically stacked Eu layers, for Eu 221 moments parallel and perpendicular to the Fe moments. We used the standard VASP package [30,31], 222 and verified that the results were fully converged with respect to the Brillouin zone integration, plane 223 wave cutoff, and the number of bands included in the diagonalization. We also varied the effective 224 Hubbard repulsion parameter for Eu f orbitals, U-J, between 5 and 7 eV, which had little impact on the 225 result. We find that for parallel Eu-Fe moments, J_x =150-170 μeV , while for perpendicular Eu-Fe moments, $J_{\rm y}$ is essentially zero. While this result seems to considerably overestimate the exchange anisotropy (see 226 ref. [14], Supplementary Materials Section 1A, for a discussion of the difficulties of DFT calculation for 227 228 noncollinear Eu-Fe moments, which we assume contributes to this overestimation), it clearly shows that 229 DFT calculations also support a strongly anisotropic Eu interplanar interaction.

The anisotropic interaction between Eu planes can be thought to result from the anisotropic hopping of conduction electrons through the Fe plane. Considering a standard superexchange interaction, conduction electrons with spins polarized along the Eu direction will generally have a larger Eu-Fe hopping amplitude *t* when the Eu and Fe moments are parallel ($J_x = J + W$) rather than perpendicular ($J_y = J - W$), which generates a stronger antiferromagnetic interaction for collinear Eu-Fe moments. Further, the weak but finite *ferromagnetic* interaction between Eu planes is also mediated through the Fe layer, and 236 as the Fe moments have a much larger susceptibility perpendicular to their ordering direction, the 237 ferromagnetic interaction is stronger for perpendicular Eu and Fe moments, which weakens their overall 238 effective antiferromagnetic interaction (this can be considered an extreme case of the RKKY interaction). 239 In this sense, we can view this Eu interplanar interaction anisotropy (W) as a generalized Eu-Fe biquadratic 240 coupling independent of the previously investigated Eu-Fe biquadratic coupling (K), where the square of Fe moments $(f_{i,x}^2 - f_{i,y}^2)$ couples the Eu moments above and below the iron plane. We note that this 241 picture is reminiscent of the Fe-Fe biquadratic coupling within the FeAs plane that generates an effective 242 243 anisotropic in-plane exchange of Fe moments $J_{1a} - J_{1b}$ [11]. These types of interactions have been long 244 overlooked in the past, but are relevant to rare-earth and transition metal intermetallic systems with 245 multiple magnetic orders [32,33].

246 A complete determination of the spin Hamiltonian in EuFe₂As₂ also sheds light on the doping 247 dependence of Eu magnetic order, which has yet to be fully understood. As in other iron pnictides, 248 chemical doping in EuFe₂As₂ rapidly suppresses the Fe-SDW and stabilizes superconductivity [34–42]. In 249 contrast, the doping has only a weak effect on the magnetic ordering temperature in the Eu layer, but 250 causes a smooth evolution from an A-type AFM order to a c-axis canted AFM order (c-AFM), and finally to 251 a c-axis ferromagnetic order (FM) [21,22,41–47]. This doping dependence can be naturally understood as 252 the consequence of Eu moments lowering their energy by aligning with the Fe-SDW, with this energy 253 saving being gradually diminished as doping weakens the Fe-SDW. Further, in the parent compound both 254 the Fe-SDW and Eu-AFM are robust under moderate hydrostatic pressure even as superconductivity 255 develops [48], while in the underdoped case a pressure-induced transition from c-AFM to FM occurs only 256 after the SDW is nearly fully suppressed [49]. This suggests the SDW plays a role in both the orientation 257 and the interaction of Eu moments. Future doping dependence studies may provide more insight on how 258 the Eu interlayer interaction is influenced by the various orders in the FeAs plane, including by 259 superconductivity [50].

260 In conclusion, our sample environment allows us to approach the magnetic coupling and 261 magnetotransport properties of the EuFe₂As₂ system in an unprecedented fashion. Through mechanical 262 stress we can prevent field detwinning and gain access to the meta-magnetism and the associated 263 magnetotransport behavior of a monodomain sample. From measurements of the spin flip and moment 264 saturation fields we are able to determine the strengths of coupling between Eu and Fe planes and 265 discover the presence of an anisotropic exchange term in the spin Hamiltonian. We emphasize again that 266 in a freestanding crystal, the rapid field-detwinning has prevented any previous determination of the 267 anisotropic Eu interplanar interaction in this system. This new technique not only deepens our 268 understanding of the EuFe₂As₂ system, but can also be applied to a variety of systems to disentangle the 269 strongly coupled spin, orbit and lattice degree of freedom.

271 **METHODS**:

272 Sample Preparation

273 Single crystal samples of $EuFe_2As_2$ were grown from a tin flux as described elsewhere [43]. The sample 274 was cleaved from a large as-grown single crystal plate and cut along the tetragonal [1 1 0] direction into 275 a bar with dimensions 3.2 x 0.50 x 0.065 mm. These sample dimensions correspond to a demagnetization 276 factor of N=0.13 along the applied field direction [51], resulting in a small maximum demagnetization 277 field of only NM_{Eu}~0.005 T. A piezo-actuator uniaxial stress device (Razorbill Instruments, CS-100) was 278 used to provide in-situ stress in the B_{2g} symmetry channel, i.e. in the direction of the orthorhombic 279 distortion, such that the applied tensile or compressive stress detwins the sample to either the A or B 280 domain, respectively (Fig.1). The four-wire electrical contact geometry is illustrated in Figure 1, with wires 281 underneath the sample to not obstruct the x-ray fluorescence off the top surface of the crystal. 282 Measurements of the resistivity coefficient ρ_{xx} aligned along the stress axis were performed using a 283 standard 4-point measurement and an SR830 lock-in amplifier. A second sample from the same growth 284 batch was cut into a thin octagon with surface area 2.18 mm² and thickness 0.0165 mm, with negligible 285 in-plane demagnetization factor [51]. Magnetization was measured in a Quantum Design PPMS.

286 X-ray Magnetic Circular Dichroism

287 XMCD was measured at the Advanced Photon Source beamline 4-ID-D at Argonne National Laboratory. 288 We probed the Eu L₃ edge using x-rays of 6.97 keV, which measure the spin polarization of the Eu 5d band 289 due to the magnetic moment of the 4f orbital. Generally, the Eu L₃ edge XMCD signal can be taken as 290 proportional to the 4f moment magnetization; however, as the Eu 5d band has a significant hybridization 291 with the As 4p orbitals [52], which themselves hybridize with the Fe 3d orbitals, the exact value of the 292 XMCD is expected to have some dependence on Fe conduction effects. Nonetheless, we can use the sharp 293 changes in XMCD signal to mark the fields at which magnetic transitions and saturations occur at. A superconducting split coil magnet with a large bore was used to apply magnetic field. The sample temperature was controlled using He flow. XMCD was collected in fluorescence geometry by monitoring the Eu L_{α} line using a four element Vortex detector integrated with the Xspress module to enable a larger dynamical range. Circularly polarized x-rays were generated using a 180 microns thick diamond (111) phase plate. Data was corrected for self-absorption.

The XMCD spot size illuminates the whole sample width across the y direction and is roughly 100 microns wide along the x direction (between the transport wires) and probes a depth of about 5 microns. The beam is centered on the middle of the crystal where strain is most transmitted and homogenous. The transport wires are separated by about 1700 microns, and transport is sensitive to the whole bulk of the sample. While the resistivity and XMCD are not measuring exactly the same volume of crystal, the tight correlation between the two data sets suggests no major difference in crystal behavior between the two sampled volumes.

306 Magnetization measurement

307 The magnetization of the single crystal EuFe₂As₂ sample was measured by the vibrating sample 308 magnetometer option of a Quantum Design Dynacool. The sample was cut into a thin octagon so that field 309 applied along the $[1 \ 0 \ 0]_T$ and $[1 \ 1 \ 0]_T$ directions have the same (minimal) demagnetization factor. The 310 sample was encased in GE varnish, which is known to fix the domain configuration and prevent field 311 detwinning. In Figure 5, at T=2 K the field was applied and magnetization measured along the $[1 \ 0 \ 0]_T$ 312 direction (blue), i.e. at 45° to both domain easy axes, and the magnetization indicates a continuous spin canting towards a saturation at $\mu_0 H_{45^\circ}^{sat} = 0.73 \text{ T}$. Conversely, for field applied along $[1 \ 1 \ 0]_T$ the 313 magnetization appears to be a combination of both a B domain spin flip at $\mu_0 H_B^{flip} = 0.45$ T and an A 314 domain continuous spin canting which saturates at $\mu_0 H_A^{sat} = 1.15$ T, in strong agreement with the 315 316 detwinned XMCD data of Figure 4. To further demonstrate this, we normalize the XMCD data of Figure 4

by the B domain 1 T saturated value and average their values over the field range to simulate a 50/50 perfectly twinned sample. This data is plotted in grey in Figure 5b and is in very strong agreement with the octagonal sample, from which we conclude that the domain populations are indeed held fixed by the GE varnish.

321 DATA AVAILABILITY

The authors declare that all data supporting the findings of this study are available within the article or from the corresponding author upon reasonable request.

324 ACKNOWLEDGEMENTS

325 We thank Jannis Maiwald for useful discussion. This work was supported by NSF MRSEC at the University

of Washington (DMR-1719797) and the Air Force Office of Scientific Research Young Investigator

Program under Grant FA9550-17-1-0217. J.H.C. acknowledges the support of the David and Lucile

328 Packard Foundation, the Alfred P. Sloan foundation and the State of Washington funded Clean Energy

329 Institute. J.L. acknowledges support from the National Science Foundation under grant no. DMR-

1848269. The work performed at the Advanced Photon Source was supported by the US Department of

Energy, Office of Science, and Office of Basic Energy Sciences under Contract No. DE-AC02-06CH11357.

332 J.J.S. acknowledges the support by the U.S. Department of Energy, Office of Science, Office of Workforce

- 333 Development for Teachers and Scientists, Office of Science Graduate Student Research (SCGSR)
- program. The SCGSR program is administered by the Oak Ridge Institute for Science and Education

335 (ORISE) for the DOE. ORISE is managed by ORAU under contract number DE-SC0014664. All opinions

expressed in this paper are the author's and do not necessarily reflect the policies and views of DOE,

337 ORAU, or ORISE.

338 AUTHOR CONTRIBUTIONS

- 339 J.J.S. conceived the project and grew the samples. G.F., Y.C, J.-W.K., P.R. and J.J.S. performed the x-ray
- 340 measurements. Y.S. and P.M. performed transport and magnetization measurements on the freestanding
- samples. S.P. and J.L. contributed to the theoretical analysis. I.I.M. contributed to the theoretical analysis
- and performed DFT calculations. J.-H.C. supervised the research. J.J.S. and J.-H.C. wrote the manuscript
- 343 with the input from all the authors.

344 ADDITIONAL INFORMATION

345 **Competing interests:** The authors declare no competing financial interests.

346 **REFERENCES**

- 347 [1] P. Dai, J. Hu, and E. Dagotto, Nat. Phys. 8, 709 (2012).
- 348 [2] P. Dai, Rev. Mod. Phys. 87, 855 (2015).
- 349 [3] R. M. Fernandes, D. K. Pratt, W. Tian, J. Zarestky, A. Kreyssig, S. Nandi, M. G. Kim, A. Thaler, N. Ni,
- 350 P. C. Canfield, R. J. McQueeney, J. Schmalian, and A. I. Goldman, Phys. Rev. B **81**, 140501 (2010).
- 351 [4] F. Kretzschmar, T. Böhm, U. Karahasanović, B. Muschler, A. Baum, D. Jost, J. Schmalian, S.
- 352 Caprara, M. Grilli, C. Di Castro, J. G. Analytis, J. H. Chu, I. R. Fisher, and R. Hackl, Nat. Phys. **12**, 560
 353 (2016).
- M. D. Watson, P. Dudin, L. C. Rhodes, D. V. Evtushinsky, H. Iwasawa, S. Aswartham, S. Wurmehl,
 B. Büchner, M. Hoesch, and T. K. Kim, Npj Quantum Mater. 4, 1 (2019).
- J. Zhao, D. T. Adroja, D. X. Yao, R. Bewley, S. Li, X. F. Wang, G. Wu, X. H. Chen, J. Hu, and P. Dai,
 Nat. Phys. 5, 555 (2009).
- 358 [7] S. O. Diallo, V. P. Antropov, T. G. Perring, C. Broholm, J. J. Pulikkotil, N. Ni, S. L. Bud'Ko, P. C.
- 359 Canfield, A. Kreyssig, A. I. Goldman, and R. J. McQueeney, Phys. Rev. Lett. **102**, 187206 (2009).
- J. Pelliciari, K. Ishii, M. Dantz, X. Lu, D. E. Mcnally, V. N. Strocov, L. Xing, X. Wang, C. Jin, H. S.
 Jeevan, P. Gegenwart, and T. Schmitt, Phys. Rev. B **95**, 115152 (2017).
- 362 [9] X. Lu, R. Zhang, H. Luo, A. H. Nevidomskyy, Q. Si, and P. Dai, Science. **345**, 657 (2014).
- 363 [10] A. N. Yaresko, G. Q. Liu, V. N. Antonov, and O. K. Andersen, Phys. Rev. B **79**, 144421 (2009).
- 364 [11] A. L. Wysocki, K. D. Belashchenko, and V. P. Antropov, Nat. Phys. 7, 485 (2011).
- 365 [12] R. M. Fernandes, A. V. Chubukov, J. Knolle, I. Eremin, and J. Schmalian, Phys. Rev. B 85, (2012).

- J. K. Glasbrenner, I. I. Mazin, H. O. Jeschke, P. J. Hirschfeld, R. M. Fernandes, and R. Valentí, Nat.
 Phys. 11, 953 (2015).
- 368 [14] J. Maiwald, I. I. Mazin, and P. Gegenwart, Phys. Rev. X 8, 011011 (2018).
- 369 [15] J. P. C. Ruff, J. H. Chu, H. H. Kuo, R. K. Das, H. Nojiri, I. R. Fisher, and Z. Islam, Phys. Rev. Lett. 109,
 370 1 (2012).
- 371 [16] Y. Xiao, Y. Su, W. Schmidt, K. Schmalzl, C. M. N. Kumar, S. Price, T. Chatterji, R. Mittal, L. J. Chang,
 372 S. Nandi, N. Kumar, S. K. Dhar, A. Thamizhavel, and T. Brueckel, Phys. Rev. B 81, (2010).
- 373 [17] S. Zapf, C. Stingl, K. W. Post, J. Maiwald, N. Bach, I. Pietsch, D. Neubauer, A. Löhle, C. Clauss, S.
- Jiang, H. S. Jeevan, D. N. Basov, P. Gegenwart, and M. Dressel, Phys. Rev. Lett. **113**, 227001
 (2014).
- 376 [18] S. Jiang, Y. Luo, Z. Ren, Z. Zhu, C. Wang, X. Xu, Q. Tao, G. Cao, and Z. Xu, New J. Phys. **11**, 025007
 377 (2009).
- Q.-P. Ding, N. S. Sangeetha, W. R. Meier, M. Xu, S. L. Bud'ko, P. C. Canfield, D. C. Johnston, and Y.
 Furukawa, Phys. Rev. B **102**, (2020).
- 380 [20] Z. Xu, J. Pan, Z. Tao, R. Liu, and G. Tan, Chinese Phys. B **29**, 77402 (2020).
- 381 [21] J. Herrero-Martín, V. Scagnoli, C. Mazzoli, Y. Su, R. Mittal, Y. Xiao, T. Brueckel, N. Kumar, S. K.
- 382 Dhar, A. Thamizhavel, and L. Paolasini, Phys. Rev. B **80**, 134411 (2009).
- 383 [22] Y. Xiao, Y. Su, M. Meven, R. Mittal, C. M. N. Kumar, T. Chatterji, S. Price, J. Persson, N. Kumar, S.
- 384 K. Dhar, A. Thamizhavel, and T. Brueckel, Phys. Rev. B **80**, 174424 (2009).
- 385 [23] See Supplementary Materials.
- 386 [24] J. J. Sanchez, P. Malinowski, J. Mutch, J. Liu, J.-W. Kim, P. J. Ryan, and J.-H. Chu, ArXiv: 2006.09444

387 (2020).

- 388 [25] R. M. Fernandes, E. Abrahams, and J. Schmalian, Phys. Rev. Lett. 107, 217002 (2011).
- 389 [26] B. Valenzuela, E. Bascones, and M. J. Calderón, Phys. Rev. Lett. **105**, 207202 (2010).
- 390 [27] C. C. Chen, J. Maclejko, A. P. Sorini, B. Moritz, R. R. P. Singh, and T. P. Devereaux, Phys. Rev. B 82,
 391 100504 (2010).
- J. J. Ying, X. F. Wang, T. Wu, Z. J. Xiang, R. H. Liu, Y. J. Yan, A. F. Wang, M. Zhang, G. J. Ye, P.
 Cheng, J. P. Hu, and X. H. Chen, Phys. Rev. Lett. **107**, 067001 (2011).
- T. Terashima, N. Kurita, A. Kikkawa, H. S. Suzuki, T. Matsumoto, K. Murata, and S. Uji, J. Phys. Soc.
 Japan **79**, (2010).
- 396 [30] G. Kresse and J. Hafner, Phys. Rev. B 47, (1993).
- 397 [31] I. I. Mazin, M. D. Johannes, L. Boeri, K. Koepernik, and D. J. Singh, Phys. Rev. B 78, (2008).
- 398 [32] N. J. Ghimire, R. L. Dally, L. Poudel, D. C. Jones, D. Michel, N. Thapa Magar, M. Bleuel, M. A.
- 399 McGuire, J. S. Jiang, J. F. Mitchell, J. W. Lynn, and I. I. Mazin, Sci. Adv. 6, eabe2680 (2020).
- 400 [33] J. X. Yin, W. Ma, T. A. Cochran, X. Xu, S. S. Zhang, H. J. Tien, N. Shumiya, G. Cheng, K. Jiang, B. Lian,
- 401 Z. Song, G. Chang, I. Belopolski, D. Multer, M. Litskevich, Z. J. Cheng, X. P. Yang, B. Swidler, H.
- 402 Zhou, H. Lin, T. Neupert, Z. Wang, N. Yao, T. R. Chang, S. Jia, and M. Zahid Hasan, Nature 583, 533
 403 (2020).
- 404 [34] N. Kurita, M. Kimata, K. Kodama, A. Harada, M. Tomita, H. S. Suzuki, T. Matsumoto, K. Murata, S.
 405 Uji, and T. Terashima, Phys. Rev. B 83, 214513 (2011).
- 406 [35] W. Uhoya, G. Tsoi, Y. K. Vohra, M. A. McGuire, A. S. Sefat, B. C. Sales, D. Mandrus, and S. T. Weir,
 407 J. Phys. Condens. Matter 22, (2010).

408	[36]	Y. He, T. Wu, G. Wu, Q. J. Zheng, Y. Z. Liu, H. Chen, J. J. Ying, R. H. Liu, X. F. Wang, Y. L. Xie, Y. J
409		Yan, J. K. Dong, S. Y. Li, and X. H. Chen, J. Phys. Condens. Matter 22 , 235701 (2010).

410 [37] Anupam, P. L. Paulose, S. Ramakrishnan, and Z. Hossain, J. Phys. Condens. Matter 23 455702,
411 (2011).

- 412 [38] M. Nicklas, M. Kumar, E. Lengyel, W. Schnelle, and A. Leithe-Jasper, J. Phys. Conf. Ser. 273,
 413 (2011).
- 414 [39] G. H. Cao, W. H. Jiao, Y. K. Luo, Z. Ren, S. Jiang, and Z. A. Xu, J. Phys. Conf. Ser. **391**, 012123
 415 (2012).
- 416 [40] W. H. Jiao, Q. Tao, J. K. Bao, Y. L. Sun, C. M. Feng, Z. A. Xu, I. Nowik, I. Felner, and G. H. Cao, Epl
 417 95, 67007 (2011).
- 418 [41] U. B. Paramanik, P. L. Paulose, S. Ramakrishnan, A. K. Nigam, C. Geibel, and Z. Hossain,
 419 Supercond. Sci. Technol. 27, 075012 (2014).
- 420 [42] S. Nandi, W. T. Jin, Y. Xiao, Y. Su, S. Price, D. K. Shukla, J. Strempfer, H. S. Jeevan, P. Gegenwart,
 421 and T. Brückel, Phys. Rev. B 89, 014512 (2014).
- 422 [43] W. T. Jin, Y. Xiao, Z. Bukowski, Y. Su, S. Nandi, A. P. Sazonov, M. Meven, O. Zaharko, S. Demirdis,
 423 K. Nemkovski, K. Schmalzl, L. M. Tran, Z. Guguchia, E. Feng, Z. Fu, and T. Brückel, Phys. Rev. B 94,
 424 184513 (2016).
- 425 [44] S. Jiang, H. Xing, G. Xuan, Z. Ren, C. Wang, Z. A. Xu, and G. Cao, Phys. Rev. B **80**, 184514 (2009).
- 426 [45] A. Baumgartner, D. Neubauer, S. Zapf, A. V. Pronin, W. H. Jiao, G. H. Cao, and M. Dressel, Phys.
 427 Rev. B **95**, 174522 (2017).
- 428 [46] W. H. Jiao, Q. Tao, Z. Ren, Y. Liu, and G. H. Cao, Npj Quantum Mater. **2**, 50 (2017).

- 429 [47] V. H. Tran, T. A. Zaleski, Z. Bukowski, L. M. Tran, and A. J. Zaleski, Phys. Rev. B 85, 052502 (2012).
- 430 [48] W. T. Jin, Y. Xiao, S. Nandi, S. Price, Y. Su, K. Schmalzl, W. Schmidt, T. Chatterji, A. Thamizhavel,
 431 and T. Brückel, Phys. Rev. B 100, 14503 (2019).
- 432 [49] W. T. Jin, J. P. Sun, G. Z. Ye, Y. Xiao, Y. Su, K. Schmalzl, S. Nandi, Z. Bukowski, Z. Guguchia, E. Feng,
- 433 Z. Fu, and J. G. Cheng, Sci. Rep. **7**, 1 (2017).
- 434 [50] Z. Devizorova and A. Buzdin, Phys. Rev. B **100**, 104523 (2019).
- 435 [51] R. Prozorov and V. G. Kogan, Phys. Rev. Appl. 10, 014030 (2018).
- 436 [52] G. Adhikary, N. Sahadev, D. Biswas, R. Bindu, N. Kumar, A. Thamizhavel, S. K. Dhar, and K. Maiti, J.
 437 Phys. Condens. Matter 25, (2013).
- 438 [53] P. Malinowski, Q. Jiang, J. J. Sanchez, J. Mutch, Z. Liu, P. Went, J. Liu, P. J. Ryan, J.-W. Kim, and J.439 H. Chu, Nat. Phys. (2020).
- 440 [54] Y. Xiao, Y. Su, S. Nandi, S. Price, B. Schmitz, C. M. N. Kumar, R. Mittal, T. Chatterji, N. Kumar, S. K.
- 441 Dhar, A. Thamizhavel, and T. Brückel, Phys. Rev. B **85**, 094504 (2012).

442





Fig. 1 (a) EuFe₂As₂ unit cell at T=7 K and zero applied magnetic field. Both Fe and Eu antiferromagnetic 446 447 orders are stabilized with easy axes aligned with the longer a lattice constant of the orthorhombic unit 448 cell. (b) Uniaxial stress is applied along the \hat{x} direction aligned with the orthorhombic a/b orthorhombic 449 unit cell lattice directions such that tension (compression) detwins the sample to the A (B) domain 450 (orange/blue outline). Resistivity measurements along the stress axis measure ρ_a (ρ_b) aligned with the 451 a(b) lattice constant of the A (B) domain. A magnetic field is applied perpendicular to the strain axis at 452 10° above parallel from the a/b plane, causing a reorientation of Eu moments to align along the field 453 direction. XMCD is proportional to the Eu magnetization along the applied field direction. For simplicity, 454 we collapse the 4 Eu atoms and 8 Fe atoms of the doubled orthogonal unit cell into 2 Eu (blue arrows) 455 and 4 Fe (red arrows) effective moments.

456

457





Fig. 2 T = 30K single pass applied magnetic field sweep for the detwinned A and B domains (XMCD data was not collected for B domain 0.8T-1T). XMCD above the Eu AFM ordering temperature shows a nearly isotropic response to field. ρ_{xx} vs applied magnetic field reveals a minimal magnetoresistance. (Inset, left) ρ_{xx} vs temperature for the detwinned A and B monodomains reveal no additional anisotropy induced at the Eu AFM ordering temperature $T_{N,Eu} = 19.1K$. (Inset, right) The resistivity anisotropy $\eta = \frac{\rho_b - \rho_a}{\rho_b + \rho_a}$ (black).

466



Fig. 3 Field sweep at T=7 K in the fully magnetically ordered phase presented in Fig.1a. Applied field
ramped from 0 T to 1 T, -1 T and 0 T in each detwinned A (a) and B (b) monodomain states. Inset to
bottom panel of (a) shows the small magnetoresistance hysteresis visible near 0.5 T in the A domain.





479 Fig. 4 (a) Data from Fig.3a-b replotted as the average value of positive and negative field sweeps of 480 XMCD and the magnetoresistance against the absolute value of applied magnetic field for the 481 detwinned A (orange) and B (blue) monodomains. Linear fits (black lines) to the XMCD magnitude for $|\mu_0 H| = 0.6T$ to 1T indicate both values coincide at $|\mu_0 H_A^{sat}| \sim 1.17T$. The magnetoresistance for 482 483 positive and negative field are nearly identical and tightly overlap. (b) The difference in XMCD and ρ_{xx} for increasing and decreasing fields yields $\Delta XMCD = XMCD(H_{inc}) - XMCD(H_{dec})$ and $\Delta \rho_{xx} =$ 484 $\rho_{xx}(H_{inc}) - \rho_{xx}(H_{dec})$. For the B domain the peak values of $\Delta XMCD$ and $\Delta \rho_{xx}$ coincide at $\mu_0 H_B^{flip} =$ 485 0.48T. 486

487

488





Fig. 5 (a) The XMCD data of Fig.4a for the B domain normalized to the 1 T mean value. Black (magenta) line represents the T=0 K metamagnetic spin flip (spin flop) transition calculated using the anisotropic JKW (isotropic JK) model (see main text). **(b)** Octagon sample magnetization with field applied along the tetragonal $[1 \ 0 \ 0]_T$ (blue) and $[1 \ 1 \ 0]_T$ (red) directions. Critical fields marked by dashed lines. Magnetization normalized by 1 T saturation value along $[1 \ 0 \ 0]_T$. (grey) Average of the two domain

XMCD data of Figure 4a, normalized by the B domain saturated value at 1 T.

497

496

498