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| 1 | Symmetry-Breaking Interlayer Dzyaloshinskii-Moriya Interactions in |
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| 2 | Synthetic Antiferromagnets |
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| 17 | The magnetic interfacial Dzyaloshinskii-Moriya interaction (DMI) in multi-layered thin |
| 18 | films can lead to chiral spin states, of paramount importance for future spintronic |
| 19 | technologies ^{1,2} . Interfacial DMI typically manifests as an intralayer interaction, mediated |
| 20 | via a paramagnetic heavy metal in systems lacking inversion symmetry ³ . Here we show |
| 21 | that, by designing synthetic antiferromagnets with canted magnetisation states ^{4,5} , it is also |
| 22 | possible to observe direct evidence of the interfacial interlayer-DMI at room temperature. |
| 23 | The interlayer-DMI breaks the symmetry of the magnetic reversal process via the |
| 24 | emergence of non-collinear spin states, which results in chiral exchange-biased hysteresis |
| 25 | loops. The spin chiral interlayer interactions reported here are expected to manifest in a |
| 26 | range of multi-layered thin film systems, opening up as yet unexplored avenues for the |
| 27 | development and exploitation of chiral effects in magnetic heterostructures ^{6–8} . |
| 28 | |

33 The interfacial Dzyaloshinskii-Moriya interaction (DMI) is an antisymmetric exchange interaction emerging in systems lacking inversion symmetry that promotes chiral coupling 34 between spins^{1,3}. In ferromagnets (FM), this gives rise to topological spin textures such as 35 skyrmions and chiral domain walls, with outstanding properties to store, transport and process 36 magnetic information ⁹⁻¹². Interfacial DMI in an ultra-thin FM layer describes the coupling of 37 spins S_i and S_i , mediated by a paramagnetic (PM) heavy metal atom l in a neighbouring layer 38 (left sketch in **Fig. 1a**), as described by the three-site Lévy-Fert model ¹³. The DMI energy per 39 atom pair is expressed as $E_{DMI} = D_{ii}$ ($S_i \times S_i$), where D_{ii} is the Moriva vector, whose direction 40 is dictated by symmetry rules ¹⁴. This interaction favours one sense of rotation of spins in the 41 same FM layer, *i.e.* it is a chiral *intralayer* interaction. 42

Together with the vast research in FM systems, DMI can potentially play an important role in the 43 emergent field of antiferromanetic (AF) spintronics⁸. In particular, the existence of a non-44 negligible interlayer DMI between neighbouring FM layers separated by a spacer has been 45 recently predicted ¹⁵. Similarly to intralayer-DMI, an interlayer-DMI will lead to the chiral 46 coupling of spins of different FM layers via PM atoms located in an interlayer between both FMs 47 (right sketch in Fig. 1a). However, due to the rapid decrease of the DMI interaction with distance 48 and the need for the correct crystallographic symmetry, this effect has not been experimentally 49 observed ¹⁶. Here, we report the experimental observation of a room-temperature chiral exchange 50 bias in SAF bilayers due to the interlayer-DMI, opening an unexplored route for the study and 51 manipulation of chiral spin interlayer interactions in multi-layered spintronic systems. 52

53 To obtain experimental evidence of the presence of the interlayer-DMI in synthetic antiferromagnets (SAFs), we have designed magnetic bilayers such as those depicted in Fig. 1b, 54 formed by two ultra-thin magnetic layers made of Co and CoFeB, with a heavy metal (Pt) on 55 both sides of the two layers providing perpendicular magnetic anisotropy (PMA) and acting as a 56 57 source of interfacial DMI. A Ru spacer couples both layers antiferromagnetically via Ruderman-Kittel-Kasuya-Yosida (RKKY) interactions. The Pt layers also tune the magnitude of the 58 59 effective RKKY coupling. The SAF is magnetically asymmetric: the bottom Co layer is significantly thinner than its spin reorientation transition (SRT), *i.e.* it is magnetically hard, with 60 its magnetisation strongly out-of-plane (z-direction). On the contrary, the top CoFeB layer is 61 slightly thicker than its SRT thickness, with a shape anisotropy moderately larger than its PMA 62 (Methods). Thus, the CoFeB layer is a soft magnetic layer which, because of the competition 63

- between its low in-plane anisotropy and the AF coupling with the out-of-plane Co layer, presents canted magnetisation configurations, *i.e.* it has a non-negligible magnetisation component along both in-plane and z directions ^{4,5}. Furthermore, the application of an in-plane magnetic field during growth breaks the symmetry during deposition (**Supplementary**), providing a moderate in-plane anisotropy along the field direction, referred to as the x-direction in the manuscript.
- To estimate the interlayer-DMI strength, the three-site model ¹³ is applied to our system, 69 represented as three layers arranged in an hexagonal close-packed (hcp) stacking, with two 70 71 magnetic atom layers separated by a distance t_{IL} from each other by one layer of non-magnetic atoms (Fig. 1c). The microscopic intralayer and interlayer DMI vectors D_{ii} are analytically 72 calculated ¹³ considering only next nearest neighbour FM and nearest neighbour PM atoms 73 (Methods). Fig. 1c shows the six non-zero resulting $D_{ii}^{(Co/Pt/CoFeB)}$ vectors corresponding to the 74 bonds connecting the central bottom Co spin *i* and the six outer CoFeB spins *j* of the top 75 hexagon. From these calculations, the interlayer-DMI strength $|D_{ii}^{(Co/Pt/CoFeB)}|$ is $\approx 0.02-0.03V_{I}$, 76 where V_1 is the so-called spin-orbit parameter of the material defining the magnitude of the D_{ii} 77 vectors ^{13,15}. For FM/Pt interfaces, $V_1^{(FM/Pt)} \approx 6.4$ meV/atom ¹³, of the same order of magnitude 78 as the direct exchange interaction of Co, $J^{(Co) 17}$. Hence, $|D_{ij}^{(Co/Pt/CoFeB)}| \approx 0.1-0.2$ meV/atom, 79 about one order of magnitude smaller than typical values for the intra-layer DMI¹⁸. The small 80 value of the interlayer-DMI in our samples is mostly due to the relatively large total interlayer 81 82 thickness of our samples (Pt/Ru/Pt \approx 2 nm) and the decrease of DMI with distance, as described by the three-site model (Methods). 83
- 84 We illustrate the effect of this interaction in the magnetic configuration of a bilayer SAF by depicting the ground state in Fig. 1d, for interlayer-DMI as the only (intra- or inter- layer) 85 exchange coupling interaction considered (direct exchange coupling, intra-layer DMI and RKKY 86 are excluded), and for large in-plane CoFeB and out-of-plane Co anisotropies. A strong 87 interlayer-DMI with positive $D_{ij}^{(Co/Pt/CoFeB)}$ results in an anticlockwise rotation between Co and 88 CoFeB spins along the z-direction -from bottom to top- for spins in the same row, and clockwise 89 for spins in adjacent rows. This creates an alternating configuration of spins in both top and 90 91 bottom layers along the x-direction, as illustrated in Fig. 1e, where the extended top view of the resulting hexagonal lattice is shown. 92

93 The presence of interlayer-DMI has been experimentally investigated under the following vector magnetic fields: First, a strong unipolar -either positive or negative- (~0.4 T) B_z field is applied, 94 saturating both layers. This field is then set to zero, leading to a canted CoFeB layer at 95 remanence. This initialisation is followed by a moderate bipolar oscillating in-plane field (-30 96 mT $< B_x < 30$ mT), applied while measuring the reversal of the CoFeB layer. Figs. 2a-d shows 97 experiments for one of the samples under investigation following this field sequence, where both 98 M_z (polar MOKE) and M_x (longitudinal MOKE) components of the magnetisation are probed as 99 a function of B_x (Methods). Importantly, the hysteresis loops associated to the CoFeB layer 100 reversal are shifted by $B_{bias} \approx \pm 1.1$ mT for the two possible Co orientations. 101

To complement experimental results, we have performed MC simulations (see Fig. 1c) using the 102 atomistic model described in Methods. The complex polycrystalline and amorphous 103 104 crystallographic structure of the sputtered layers, added to unknown spin-orbit parameters, 105 makes it challenging to estimate the DMI values of the samples. Moreover, V_1 will have different values for Co/Pt, Pt/CoFeB and Co/Pt/CoFeB interfaces. To incorporate realistic values in the 106 107 simulations, we have compared sets of $M_2(B_2)$ experimental results for a wide range of thicknesses with MC simulations (see Supplementary). This allows us estimate V_1 for the 108 different interfaces and associate an effective CoFeB thickness t for each sample, given by the 109 $|V_1^{(Pt/CoFeB)}/V_1^{(Co/Pt)}|$ ratio. These estimated spin-orbit parameters are then used in subsequent MC 110 simulations (Figs. 2e-h) that replicate the experimental minor loops described before. A good 111 112 qualitative agreement between experiments and simulations is observed, with simulations reproducing both the shape of the experimental loops and the chiral bias effect. Furthermore, a 113 good quantitative agreement is also found between experiments and simulations when estimating 114 the effective strength of the interlayer-DMI (Methods). We therefore conclude that the chiral 115 bias effect described here constitutes a fingerprint of the interlayer-DMI. Other indirect exchange 116 interactions such as the biquadratic interlayer coupling ¹⁹ cannot account for the chiral nature of 117 the observed effect. Furthermore, intra-layer DMI effects leading to asymmetric magnetic 118 hysteresis processes have been only observed in laterally-patterned nanomagnets, and require the 119 simultaneous application of orthogonal magnetic fields ^{20,21}, in contast to our experiments. 120

We have studied the dependence of the chiral B_{bias} magnitude as a function of CoFeB thickness (left and bottom axes in **Fig. 3a**) for the range 1.5 - 2.4 nm. This function rises sharply after the nominal SRT CoFeB thickness, peaking at 1.7 nm, and dropping to negligible values for 124 thicknesses above 2.2 nm, when the CoFeB becomes strongly in plane. The regime where nonzero B_{bias} is observed corresponds to the thickness range where the CoFeB magnetisation 125 becomes canted ⁴, as illustrated by the further right axis, where the function $sin2\theta$, as obtained 126 from macrospin MC simulations (Methods), presents non-zero values. θ is the effective 127 macrospin canting angle of the CoFeB (see Fig. 1b). In addition, the function plotted in nearer-128 right and top axes is the normalised $|B_{bias}|$ extracted from MC atomistic simulations as a function 129 of the effective CoFeB thickness t, showing an excellent agreement with experiments. Fig. 3b 130 displays the characteristic spin configurations of the system, obtained from atomistic 131 simulations, for the thickness ranges: t < 1.6 nm, (AP), 1.6 nm < t < 2.2 nm (CANT) and t > 2.2132 nm (PERP). The AP and PERP are standard spin configurations, whereas the spin state for the 133 CANT regime is explained below. No bias is observed for the AP and PERP regimes due to a net 134 zero $E_{DMI}^{(Co/Pt/CoFeB)}$ in both cases (Methods). A measureable B_{bias} is only present for the CANT 135 regime, where a small effective CoFeB anisotropy is expected to promote the emergence of 136 effects ruled by small energy contributions, such as the interlayer-DMI. 137

138 To understand in detail the CANT regime and its role in the chiral bias, Fig. 4 includes results from MC simulations for a SAF within this thickness regime. Fig. 4a shows snapshots during the 139 reversal process of the CoFeB layer at different B_x values, for Co pointing along the +z direction. 140 Overall, the magnetisation process follows the same mechanism previously reported for this type 141 of samples⁴, result of the competing energies present in the system: The soft layer (CoFeB) 142 reverses back and forth under B_x , while the hard (Co) layer remains unchanged because of its 143 high PMA. The AF RKKY promotes an antiparallel orientation of CoFeB and Co, leading to a 144 peak in M_z during CoFeB reversal (Fig. 2g). The AF RKKY also results in an incomplete in-145 plane saturation of CoFeB at the maximum B_r applied (Fig. 2h). In addition, the intralayer-DMI 146 promotes a chiral clockwise spin rotation -from left to right- across the CoFeB layer. To satisfy 147 this requirement, the magnetisation reverses via the propagation of domain walls with clockwise 148 149 chirality. To achieve the same wall chirality for both branches of the hysteresis loop and keep an antiparallel alignment with Co, a domain wall is nucleated at opposite edges of the simulated 150 area for either branch. However, none of these contributions is able to create a biased switching 151 in extended structures and under B_x only ²¹, requiring an additional symmetry-breaking 152 mechanism. The reversal process will be in reality strongly influenced by defects and 153 inhomogeneities of the layers ²², and driven by domains of very small sizes for thicknesses 154

around the SRT ²³, making their direct observation using magneto-optical methods as those used here very challenging ⁵. Despite these, the macroscopic bias observed experimentally indicates that a clear reversal asymmetry for both branches is present.

Complementing these results, Figs. 2i-j show the evolution of $E_{DMI}^{(Co/Pt/CoFeB)}$ during CoFeB 158 reversal, for the two possible z-directions of Co. Whereas standard magnetic energy terms are 159 symmetric under inversion of B_x , this is not the case for $E_{DMI}^{(Co/Pt/CoFeB)}$, which presents two 160 plateaus at moderate B_x values and a biased switching. An asterisk in those graphs marks the 161 state of the system that is energetically more favourable from an interlayer-DMI point of view, 162 which is depicted in the insets of Figs. 2f-h. These sketches show the spin configuration for top 163 CoFeB and bottom Co layers, where green (red) interconnecting lines indicate the pair bonds 164 where the interlayer-DMI is energetically favourable (unfavourable) for that spin configuration 165 (compare with Fig. 1d). The figure also indicates how unfavourable bonds cause canting of the 166 CoFeB spins (red arrows) which become more antiparallel to Co because of the strong AF 167 **RKKY** interaction. 168

MC simulations evidence the emergence of this type of noncollinear magnetisation states, as a 169 170 result of the competition between interlayer-DMI and RKKY coupling (Fig. 4b). Magnetisation amplitude changes of up to 15% for S_x , with a period corresponding to a few atomic lattice 171 constants, are observed in simulations, with this behavior dependent on the $|V_1^{(Pt/CoFeB)}|/|V_1^{(Co/Pt)}|$ 172 ratio (not shown here). The relevance of noncollinear magnetic phases for symmetry breaking 173 has already been pointed out ²⁴. Here, simulations indicate how this CoFeB magnetisation 174 modulation is different for either branch, due to the different configuration of -energetically 175 satisfied and unsatisfied- interlayer-DMI bonds for either branch (Fig. 4(c,d)). The subtle 176 symmetry breaking mechanism responsible for the chiral exchange bias is thus the result of these 177 two effects acting together: the emergence of noncollinear spin states during reversal, combined 178 with this asymmetric bond profile. This magnetisation modulation asymmetry also manifests as 179 other small asymmetric features in these loops. For instance, the M_z peak reaches larger values 180 for one of the two branches (Figs. 2e, g), revealing spin modulations of larger amplitude, due to 181 the interlayer-DMI competing less efficiently with the RKKY interaction. 182

In conclusion, we report a room temperature chiral exchange bias in ultra-thin asymmetric synthetic antiferromagnetic bilayers caused by the presence of DMI across the interlayer. The emergence of noncollinear spin modulation, subject to different interlayer-DMI profiles during magnetic reversal, is behind this symmetry breaking. Whereas the interlayer-DMI would appear to be too weak to significantly change the intralayer magnetic ordering, due to the competition with a strong direct exchange and intralayer-DMI contributions, it can however be effective in competition with RKKY coupling, co-defining the interlayer ordering.

The canted SAFs studied here have been specifically designed to probe the presence of the 190 191 interlayer-DMI, which manifests as a macroscopic chiral exchange bias. However, we expect 192 symmetry-breaking effects induced by this interaction to play an important role in other ultrathin SAFs away from the SRT and with more standard magnetic configurations. Specifically, 193 provided that the symmetry arguments exposed here are fulfilled, the interlayer-DMI will 194 become important in these systems when the magnetic reversal becomes dominated by areas 195 with a low effective anisotropy, such as defects and layer inhomogeneities²⁵. Moreover, larger 196 net interlayer-DMI energies than the one reported here are expected in other multilayered 197 systems formed by ultra-thin interlayers (see Methods), including *e.g.* the use of spacer materials 198 that simultaneously present both RKKY and DMI interactions²⁶. This interaction will also be of 199 particular importance in magnetic systems with large antisymmetric/symmetric exchange 200 interaction ratios²⁷. The realisation of systems integrating interlayer magnetic chiral interactions 201 paves the way for the creation and manipulation of unprecendented magnetic effects in synthetic 202 antiferromagnets, e.g. the introduction of indirect -via the magnetic state of a neighbouring 203 layer- control of asymmetric effects in the motion of domain walls²⁸ and spin waves²⁹ of a 204 magnetic system. It is also of great relevance towards the development of future three-205 dimensional spintronic systems ^{6,7}. 206

207 Data availability:

All data associated to this publication is available via Enlighten, the University of Glasgow public repository.

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280 Methods

Synthetic antiferromagnets: The samples studied here were grown by DC magnetron sputtering, 281 under a base pressure of 7×10^{-8} mbar and a growth pressure of 8×10^{-3} mbar. The structure is 282 Ta(4.0 nm)/Pt(10.0 nm)/Co(1 nm)/Pt(0.5 nm)/Ru(1.0 nm)/Pt(0.5 nm)/Co₆₀Fe₂₀B₂₀(1.6-2.4 283 nm)/Pt(2.0 nm)/Ta(2.0 nm). The Ru spacer provides antiferromagnetic coupling between the two 284 ferromagnetic films via RKKY interaction ³⁰. Although only one Pt layer at the interlayer is in 285 principle needed to observe interlayer-DMI, a symmetric interlayer with two Pt interfaces was 286 used to improve the PMA of the FM layers, as well as to provide fine tuning over the RKKY 287 coupling between the two³¹. The surface PMA of the Co and CoFeB layers, determined by 288 growing single layers with analogous structure, are 1.2 mJ/m^2 and 0.7 mJ/m^2 . This corresponds 289 to SRT thicknesses of ≈ 1.95 nm for Co and 1.55 nm for CoFeB, when the shape anisotropy 290 balances the surface PMA, *i.e.* when the effective anisotropy $K_{eff} = 2K_s/t - 0.5 \mu_0 M_s^2 = 0$. RKKY 291 coupling is created by a 1 nm Ru layer, which corresponds to the first AF peak, and tuned by the 292 Pt on both sides. For 0.5 nm of Pt, this corresponds to an AF surface energy $J_{RKKY}^{(Co/Ru/CoFeB)}$ of -293

294 0.08 mJ/m^2 . A magnetic field of $\approx 100 \text{ mT}$ is applied during the sputtering process, resulting in a 295 moderate in-plane anisotropy for the CoFeB layer along the field direction ³², measured to be up 296 to $1.8 \times 10^3 \text{ J/m}^3$. A chiral bias effect has also been observed in another similar set of samples 297 (Supplementary).

Atomistic Three-Site Model: The interlayer-DMI effect is modelled using a FM₁/PM/FM₂ atomistic trilayer with hcp stacking. The z-position of each atomic plane corresponds effectively to the middle point of each layer ³³. Each magnetic layer is represented by a single monolayer of Heisenberg spins S_i and S_j at atomic positions R_i and R_j . The spin-orbit parameters for the bottom and top layers are defined as $V_1^{(Co/Pt)}$ and $V_1^{(Pt/CoFeB)}$, respectively, and between layers as $V_1^{(Co/Pt/CoFeB)}$. Microscopic DMI vectors describing the interaction between spins S_i and S_j as mediated by impurity l within and in-between layers are obtained using the three-site model ¹³:

305
$$\boldsymbol{D}_{ijl}(\boldsymbol{R}_{li}, \boldsymbol{R}_{lj}, \boldsymbol{R}_{ij}) = -V_1 \frac{\sin(k_F(R_{li}+R_{lj}+R_{ij})+(\pi/10)Z_d)(\boldsymbol{R}_{li}\cdot\boldsymbol{R}_{lj})(\boldsymbol{R}_{li}\times\boldsymbol{R}_{lj})}{|R_{li}|^3|R_{lj}|^3R_{ij}},$$
(1)

where R_{ii} , R_{ij} are the distance vectors from the impurity *l* to the corresponding FM atom sites *i* and *j*, and R_{ij} the distance vector between these FM sites. The parameter $V_1 = \frac{135\pi}{32} \frac{\lambda_d \Gamma^2}{E_F^2 k_F^3} sin\left(\frac{\pi}{10} Z_d\right)$ refers to the material specific quantity defining the DMI strength. Hereby, k_F and E_F are the Fermi wave vector and energy respectively, λ_d is the spin-orbit coupling parameter, Γ the interaction parameter between the localised spins and the spins of conduction electrons, and Z_d the number of d-electrons.

An effective DMI vector describing the interaction between a given *ij* atomic pair can be calculated by performing a sum over all nearest neighbour PM impurities $l^{13,15}$:

314
$$\boldsymbol{D}_{ij}^{eff} = \sum_{l} \boldsymbol{D}_{ijl}(\boldsymbol{R}_{li}, \boldsymbol{R}_{lj}, \boldsymbol{R}_{ij})$$
(2)

The total DMI energy between two magnetic layers is then given by

316
$$E_{DMI} = \sum_{ij} \boldsymbol{D}_{ij}^{eff} \cdot (\boldsymbol{S}_i \times \boldsymbol{S}_j)$$
(3)

317 where this time, next-nearest neighbour *ij* pairs are considered in the calculations.

If we evaluate equation (3) for the three thickness regimes discussed in **Fig. 3b**, we find that the AP configuration obviously leads to zero interlayer-DMI, due to both layers forming 180° , resulting in $S_i \times S_j = 0$ for all pairs. The net interlayer-DMI is also zero in the PERP configuration, despite Co and CoFeB spins forming 90°. In that case, equation (3) becomes

322 $E_{DMI} = (\mathbf{S}_i \times \mathbf{S}_j) \cdot \sum_{ij} \mathbf{D}_{ij}^{eff} = 0$ (4)

since $(S_i \times S_j)$ is the same for each pair, and the total sum of $D_{ij}^{eff}=0$ for an hexagonal lattice. However, noncolinear spin configuration in the soft CoFeB layer (CANT configuration) will result in a non-zero DMI energy as describes by eq (3).

The arguments presented here for hcp stacking can be also extended to other crystallographic structures. A net non-zero $D_{ij}^{(Co/Pt/CoFeB)}$ vector is obtained, for instance, for distorted or disordered cubic phases ¹⁵. **Supplementary** includes additional information about the atomistic model.

- 330 *Magnetometry measurements:* The samples were investigated using focused magneto-optical 331 Kerr effect, with a 3.5 mW laser Gaussian spot of FWHM \approx 5 µm and wavelength = 635 nm. To 332 probe both M_z and M_x components of the samples, two different setups were used, with either 333 normal or 45° incidence geometries. Optical analyser and quarter-wave plate angles were tuned 334 to detect either Polar or Longitudinal Kerr signals, respectively.
- Analogous (bulk) vibrating sample magnetometer (VSM) measurements with two sets of perpendicular pick-up coils and Kerr control experiments complement these measurements (**Supplementary**).
- 338 *Monte Carlo atomistic simulations:* As a complement to analytical calculations, we perform 339 atomistic MC simulations using the model Hamiltonian
- 340

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$$H = -\sum_{ij} J^{(Co)}(\boldsymbol{S}_{i} \cdot \boldsymbol{S}_{j}) - \sum_{ij} J^{(CoFeB)}(\boldsymbol{S}_{i} \cdot \boldsymbol{S}_{j}) - \sum_{ij} J^{RKKY}_{ij}(\boldsymbol{S}_{i} \cdot \boldsymbol{S}_{j}) - K^{(Co)}_{i} \sum_{i} (S^{z}_{i})^{2} - K^{(CoFeB)}_{i} \sum_{i} (S^{z}_{i})^{2} - \sum_{ij} \boldsymbol{D}^{eff}_{ij}(\boldsymbol{S}_{i} \times \boldsymbol{S}_{j})$$

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(5)

based on experiments and typical parameters of Co-based alloys³⁴: A strong PMA is included for the bottom Co layer, $K_i^{(Co)} \approx K_z^{(Co)} \approx 0.7 J^{(Co)}$, and the top CoFeB layer is close to the SRT, $K_z^{(CoFeB)} \approx 0$. Additionally, we introduce an additional in-plane anisotropy in this layer to mimic experiments: $K_x^{(CoFeB)} \approx 0.4 J^{(CoFeB)}$. The FM intralayer exchange interaction for both layers is set as $J^{(CoFeB)}/J^{(Co)} = 0.5$ and the AF RKKY coupling between both layers as $J_{ij}^{RKKY} = -0.1 J^{(Co)}$.

Samples with lateral dimensions of up to $50a \times 50a$ on an hcp lattice with periodic and open boundaries have been considered. Calculations have been performed for a wide of temperatures $kT = 0.05 \cdot 0.1 J^{(Co)}$. The Monte Carlo simulations are used to calculate magnetisation curves, comparing them with experimental data. This allows us to access to the atomic-scale configuration during magnetisation reversal. For calculations of the $M_x(B_x)$ curves, out-of-plane Co and in-plane CoFeB magnetic orientations were used as the initial configuration, whereas

- fully saturated out-of-plane states were used as the initial state for $M_z(B_z)$ loops (Supplementary). In the simulations, 10^5 MC initial steps were used first to reach magnetic equilibrium. After those, the magnetisation curves were recorded by sweeping over the lattice at every MC step, updating the orientations of the spins following single-spin Metropolis dynamics. At every field, the system was again thermalised for 10^5 steps, then the averaging was performed.
- Monte Carlo macrospin simulations: Macrospin MC simulations were carried out to determine the effective canting angle of the CoFeB layer as a function of its thickness, for the SAF under investigation. PMA, in-plane shape anisotropy and RKKKY AF coupling were considered (*i.e.* neither type of DMI is included). The parameters used were extracted from experiments. PMA: $K_s^{(Co)} = 1.2 \text{ mJ/m}^2$, $K_s^{(CoFeB)} = 0.7 \text{ mJ/m}^2$. In-plane volume anisotropy: $K_v^{(CoFeB)} = 1.8 \times 10^2 \text{ J/m}^3$. Spontaneous magnetisation: $M_s^{(Co)} = 1.4 \times 10^6 \text{ A/m}$, $M_s^{(CoFeB)} = 1.2 \times 10^6 \text{ A/m}$. $J_{RKKY}^{(Co/Ru/CoFeB)} = 366$ -0.08 mJ/m².
- Estimation of the interlayer-DMI from the magnitude of the bias field: Since the interlayer-DMI 367 is considered as the only symmetry-breaking source in the system, $|B_{bias}|$ can be identified with 368 the effective strength of the interlayer-DMI. Hence, e.g. a bias of 1 mT for the 2.1 nm thick 369 CoFeB corresponds to an effective energy of 10^{-4} meV/atom, given by $E_{DM}^{(Co/Pt/CoFeB)} = m|B_{bias}|$ 370 $\approx 2 \mu_B$ /atom $\times 1 \text{ mT} \approx 10^{-4} \text{ meV/atom}$, with m the magnetic atomic moment, expressed in units 371 of the Bohr magneton μ_{B} . This compares with the bias energy extracted from simulations for 372 hcp stacking with interlayer distance $t_{IL}=2a\sqrt{2/3}=0.4$ nm for a lattice constant a=0.25 nm 373 (Figs. 2(i,j)) in the main manuscript): $E_{DM}^{(Co/Pt/CoFeB)} = mB_{bias} \sim 0.001 J^{(Co)} \approx 2 \times 10^{-2} \text{ meV/atom},$ 374 if $J^{(Co)} \approx 20$ meV/atom is considered ^{17,35}. If we consider instead $t_{IL} = 8 a\sqrt{2/3}$, ≈ 2 nm as in 375 experiments, an interlayer- DMI energy of 5×10^{-4} meV/bond is obtained. This is in rather good 376 377 agreement with the experimental $|B_{bias}|$, despite the difference between the complex experimental system investigated, comprising polycrystalline/amorphous sputtered samples and rough 378 379 interfaces, in contrast with the model, where a perfect crystalline hcp structure has been 380 considered.
- 381 *Raw data and Monte Carlo codes*

382 All metadata for this publication is available via the following link: http://dx.doi.org/10.5525/gla.researchdata.787. The atomistic and macrospin Monte Carlo codes 383 used for this study are available from the corresponding authors on reasonable request. 384

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401 **Competing interests:**

402 Authors declare no competing interests.

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Fig. 1. Interlayer-Dzyaloshinskii-Moriya Interaction (DMI) investigations in canted 408 synthetic antiferromagnets (SAFs). (a) Left. Intralayer-DMI coupling between spins *i* and *j* of 409 the same ferromagnetic (FM) layer via a paramagnetic (PM) atom; the figure depicts this type of 410 coupling for the top FM layer only. **Right.** Interlayer-DMI coupling (right) between spins of two 411 neighbouring FM layers separated by a spacer, mediated by PM atoms. (b) Schematic of the 412 magnetic state at remanence of the SAFs studied in a macrospin approximation: two ultra-thin 413 CoFeB (top) and Co (bottom) layers with Pt at the interfaces, separated by Ru to create 414 antiferromagnetic coupling between both FM layers via Ruderman-Kittel-Kasuva-Yosida 415 (RKKY) interaction. The two FM layers have different proximities to their corresponding spin-416 reorientation-transition, with Co remaining out-of-plane and CoFeB becoming canted with 417 respect to the substrate plane. θ is the (polar) effective macrospin canting angle of this layer. (c) 418 **D**_{ii}^(Co/Pt/CoFeB) interlayer-DMI vectors (green) calculated via the 3-sites model for a 419 Co(i)/Pt(l)/CoFeB(i) trilayer with hexagonal close-packed (hcp) structure. The distance between 420 magnetic atoms is the interlayer thickness (t_{II}) . The *j* letter denotes one of the seven next-nearest 421 neighbours of the i central bottom spin, with l the corresponding PM atom for this bond included 422 in the calculations. The $D_{ii}^{(Co/Pt/CoFeB)}$ vector corresponding to the interaction between both 423 central atoms at top and bottom hexagons equals zero when computed across the three nearest 424 neighbour impurities. (d) Ground state spin configuration based solely on the interlayer-DMI, for 425 426 a hexagonal-closed-packed trilayer with in-plane top and out-of-plane bottom magnetisations (no FM direct or AF RKKY exchange is considered). All green bonds connecting the middle Co to 427 the outer CoFeB spins are interlayer-DMI energetically favourable. (e) Extended top view of the 428 hexagonal lattice for the same ground state as in (d), x is the direction of the CoFeB in-plane 429 anisotropy. 430



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Fig. 2. Chiral exchange bias due to the interlayer-DMI. (a-d) Chiral exchange bias observed 432 during the reversal of the canted CoFeB layer, for a sample with CoFeB thickness = 2.1 nm. The 433 magnetisation components M_z (a,c) and M_x (b,d) of the CoFeB film are measured by Kerr effect 434 under B_x magnetic fields, after negative (a,b) and positive (c,d) initial saturating orthogonal B_z 435 fields that define the magnetic state of the Co layer for the rest of the field sequence. This 436 magnetic field sequence is thus a minor loop used to probe the reversal of the canted free layer, 437 while the out-of-plane layer remains fixed along the z-direction. The bias effect, obtained from 438 the switching field (M_x) and peaks (M_z) , is marked by a red dashed line. The insets show the most 439 favourable state of the two under moderately high B_x fields, based on the interlayer-DMI; 440 red/green lines denote interlayer-DMI energetically unfavourable/favourable bonds connecting *j* 441 top outer spins to the central *i* bottom spin. Canted spins promoted by the RKKY interaction and 442 443 an unfavourable interlayer-DMI are colored in red, in contrast to blue spins, where the two interactions promote instead an in-plane spin configuration. (e-h) Monte Carlo atomistic simulations reproducing the experiments, with $V_1^{(Pt/CoFeB)}/V_1^{(Co/Pt)} = 1.7$, corresponding to a 444 445 CoFeB thickness t = 2.1 nm. (i, j) Evolution of the interlayer-DMI energy $E_{DMI}^{(Co/Pt/CoFeB)}$ during 446 the hysteresis loops; an asterisk marks the states sketched in the inset of (f) and (h). Both mB_x 447 and interlayer-DMI energies are normalised with respect to $J^{(Co)}$, the direct intralayer exchange 448 energy, with *m* the magnetic moment of the system. 449





451 Fig. 3. Bias field dependence with CoFeB thickness. (a) Left, bottom axes (black squares and dashed line) are experiments, showing a peak around the spin reorientation transition (SRT). 452 Nearer-right, top axes (blue triangles and dash-dot line) show computed normalised bias from 453 atomistic MC simulations, with t the effective CoFeB thickness, corresponding to an interval of 454 $|V_1^{(Pt/CoFeB)}/V_1^{(Co/Pt)}|$ between -1.7 and +1.9 (Supplementary). The same behaviour is evidenced 455 for experiments and simulations. Further-right, bottom axes (red stars and dotted line) plots the 456 effective degree of canting of the CoFeB layer (when it is neither in-plane nor out-of-plane) as a 457 function of its thickness, parametrised as $sin2\theta$, as extracted from macrospin MC simulations; 458 only anisotropies and RKKY coupling interactions are considered. The magnitude of the bias 459 correlates well with the magnetisation effective degree of canting of the CoFeB layer, revealing 460 that a low competing effective anisotropy is necessary to observe a bias effect. (b) Schematics of 461 the three types of spin configurations: antiparallel (AP), canted (CANT), and perpendicular 462 (PERP) across the SRT. A non-zero net interlayer-DMI is only present for the CANT regime. 463 Red spins in the CANT state are those more favourable to become out-of-plane, due to an 464 energentically unfavourable interlayer-DMI and the effect of the RKKY interaction. 465

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470 Fig. 4. Emergence of spin modulations. (a) Snapshots of Monte Carlo simulations at remanence for a SAF with an effective CoFeB thickness = 2.1 nm and Co pointing upwards. 471 Figs. 2(g,h) are the corresponding hysteresis loops. Forward (top) and backward (bottom) 472 branches of the B_x hysteresis loop are included. Top spins in red and blue indicate the value of 473 $M_{\rm r}$ for the top CoFeB layer during reversal. The grey bottom spins represent the Co layer along 474 +z. The reversal process is asymmetric for both loop branches and occurs at different magnetic 475 fields, resulting in a biased hysteresis loop. (b) Three components of the magnetisation as a 476 function of the atomic spin number across the dashed line in the inset, for $B_x = 0$ and starting 477 from negative fields. Periodic changes in the amplitude of the three components in the 478 simulations reveal the presence of spin modulations in the CoFeB layer. Different periods for the 479 three components are observed due to their anharmonic character. (c,d) Top extended view of the 480 hexagonal lattice, with bottom Co spins colored in grey and CoFeB top spins in blue. 481 E_{DM} (Co/Pt/CoFeB) = 0 for both spin configurations. However, a different number and symmetry of 482 favourable (green) and unfavourable (red) interlayer-DMI bonds is obtained for (b) and (c), 483 breaking the symmetry of the system. This leads to a chiral bias when spin modulations become 484 present during the switching of the CoFeB layer. 485

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