Nonlinear spin control by terahertz-driven anisotropy fields

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Future information technologies, such as ultrafast data recording, quantum computation or spintronics, 1 call for ever faster spin control by light¹⁻¹⁶. Intense terahertz pulses can couple to spins on the intrinsic 2 energy scale of magnetic excitations^{5,11}. Here, we explore a novel electric dipole-mediated mechanism of 3 4 nonlinear terahertz-spin coupling which is much stronger than the linear Zeeman coupling to the terahertz magnetic field^{5,10}. Using the prototypical antiferromagnet thulium orthoferrite (TmFeO₃), we 5 demonstrate that resonant terahertz pumping of electronic orbital transitions modifies the magnetic 6 anisotropy for the ordered Fe³⁺ spins and triggers large-amplitude coherent spin oscillations. This 7 mechanism is inherently nonlinear, it can be tailored by spectral shaping of the terahertz waveforms, 8 and its efficiency outperforms the Zeeman torque by an order of magnitude. Since orbital states govern 9 10 the magnetic anisotropy in all transition metal oxides, the demonstrated control scheme is expected to be applicable to many magnetic materials. 11

Ultrafast magnetization control has become a key incentive of modern photonics, with a broad variety of successful concepts emerging at fast pace. Examples include light-induced spin reorientation in canted antiferromagnets³, the vectorial control of magnetization by light⁶, photoinduced antiferromagnet-ferromagnet phase transitions⁹, optical modification of the exchange energy^{4,14}, and driving spin precessions via nonlinear magneto-phononic coupling^{7,16}. Despite this remarkable progress, the lion share of the photon energy in all known concepts using visible and near-infrared light is idle with respect to the light-spin interaction, and avoiding dissipation of large excess energies requires special care.

In contrast, intense electromagnetic pulses at terahertz (1 THz = 10^{12} Hz) frequencies may interface spin 19 dynamics directly on their intrinsic energy scales^{5,11}. The magnetic field component of few-cycle THz pulses 20 has been used to coherently control magnons in the electronic ground state by direct Zeeman interaction^{5,11}. 21 22 Since magnetic dipole coupling is typically weak, however, THz-driven spin excitation has been confined to the linear response regime. Massive nonlinearities, such as THz-driven phase transitions^{17,18} and THz 23 lightwave electronics¹⁹⁻²², in turn, have been realized by all-electric coupling to the charge degree of freedom. 24 Apart from pioneering work on electromagnons in multiferroic TbMnO₃ (Ref. 13), efficient ways to exploit 25 the THz electric field for the control of magnetic order have been missing. 26

Here, we introduce a conceptually new universal route to control magnetism by THz electric fields. The strength and direction of the magnetic anisotropy in practically all materials is determined by the coupling of electronic orbital states to ordered spins. Therefore, an ultrashort electric field pulse can change the orbital state of electrons abruptly, leading to a sudden modification of the magnetic anisotropy. In our proof-ofconcept experiment, we exploit intense, phase-locked THz pulses to achieve such an abrupt change of the magnetic anisotropy, which in turn triggers magnon oscillations with large amplitudes that scale quadratically with the THz field strength.

Non-thermal pumping of orbital transitions in the optical range is known to induce a nonlinear spin-charge coupling on ultrashort timescales^{23,24}. In the THz spectral range, this concept can be applied to any material in which selected low-energy electronic transitions change the magnetic anisotropy, for example in oxides containing both 3d and 4f ions (e.g. orthoferrites, manganites, garnets and ferroborates) and in 3d-compounds 38 like hematite α -Fe₂O₃. However, despite the anticipated strong impact of the preparation of non-thermal 39 orbital states on the anisotropy field, terahertz spin control exploiting orbital transitions has remained largely 40 unexplored^{25,26}.

Figure 1 illustrates the fundamental idea of our experiment for the case of the prototypical antiferromagnet 41 TmFeO₃. This material crystallizes in a distorted perovskite structure (Fig. 1a). The four iron spins (blue 42 43 arrows) per unit cell occupy two antiferromagnetically coupled sublattices, whose spin orientations are mutually canted by the Dzyaloshinskii-Moriya interaction²⁷. The ${}^{3}H_{6}$ ground state of the paramagnetic rare-44 earth Tm³⁺ ions is fully split by the crystal field into a series of singlets with a characteristic energy spacing of 45 $\sim 1 - 10$ meV (Ref. 28). The angular momenta of these states are coupled with the Fe³⁺ spins by exchange and 46 dipolar interactions, which set the magnetic anisotropy. Thermal population of the singlet states within the ${}^{3}H_{6}$ 47 multiplet changes the magnetic anisotropy as a function of temperature, leading to spin reorientation phase 48 transitions²⁹: In the Γ_2 phase ($T < T_1 = 80$ K), the antiferromagnetic vector **G** is aligned along the 49 crystallographic z-axis, whereas it lies along the x-axis in the Γ_4 phase ($T > T_2 = 90$ K). For $T_1 < T < T_2$ (Γ_{24} 50 51 phase), G rotates continuously in the (xz)-plane (see Fig. 1b, Eq. (6) in Methods, Supplementary Fig. 1 and Supplementary Movie 1). The spin dynamics supports two eigenmodes, the quasi-ferromagnetic (q-FM) and 52 53 the quasi-antiferromagnetic (q-AFM) one. Instead of thermal activation, resonant pumping of electronic 54 transitions between orbital states of the rare-earth ions by THz pulses may be expected to abruptly modify the 55 magnetic anisotropy to trigger coherent magnon oscillations (Fig. 1c and Supplementary Movie 2).

We excite a 60-µm-thick window of TmFeO₃ by intense few-cycle THz transients generated by tilted-pulse-56 front optical rectification of near-infrared laser pulses¹¹. The THz peak field can be tuned up to $B_{\text{THz}} = 0.3 \text{ T}$ 57 without changing the waveform (Fig. 2a). The spectral content lies between 0.1 and 2 THz, covering both 58 magnon modes of Fe³⁺ spins and several transitions of the Tm³⁺ ground state multiplet²⁸ (Fig. 2b). Excitation 59 of phonons can be excluded as they feature frequencies above 3 THz^{28,30}. The induced ultrafast magnon 60 dynamics is revealed by tracking the polarization rotation imprinted by the Faraday effect and magnetic linear 61 dichroism on co-propagating near-infrared 30-fs probe pulses (Fig. 2c, Supplementary Figs. 2 and 3). Both the 62 q-FM and the q-AFM modes are excited (see Supplementary Fig. 4), and their frequencies exhibit 63

64 characteristic temperature dependences (Fig. 2d). Close to the phase transitions, we observe a dramatic 65 softening of the q-FM mode down to a frequency of 50 GHz, in agreement with theory²⁸.

Next, we systematically vary B_{THz} while keeping the sample in the Γ_{24} transition phase (T = 84.5 K), where 66 pumping of the rare-earth states should have maximum impact on the magnetic anisotropy. Figure 3a shows 67 the dynamic polarization rotation as a function of the delay time t between the THz pump and the optical 68 probe. Each transient is normalized by the corresponding THz peak field (see Supplementary Fig. 5 for a 69 70 quantitative analysis of the magnetization deflection angles). The signal exhibits an oscillatory behaviour with 71 two quasi-monochromatic components at frequencies of 0.1 THz and 0.8 THz corresponding to the q-FM and 72 the q-AFM mode, respectively (Fig. 3b). Most remarkably, the relative strength of the q-FM mode grows with increasing THz peak fields. Figure 3c summarizes the amplitude of both modes as a function of B_{THz} . The q-73 74 AFM mode scales linearly with the THz driving field as expected for the linear Zeeman interaction. In 75 contrast, the q-FM mode shows a distinctly nonlinear increase, which is well fit by a superposition of linear and quadratic functions of the peak field. The nonlinearity vanishes when the crystal leaves the Γ_{24} phase 76 (Fig. 3d). 77

One can show (see "Model for purely magnetic interaction" in Methods and Supplementary Fig. 6), that the 78 Zeeman torque exerted by the THz magnetic field on the Fe³⁺ spins cannot explain the nonlinear excitation of 79 80 the q-FM mode. In contrast, the THz electric field can influence the magnetic system: The point group of the 81 orthoferrites allows for an anisotropic energy term that scales quadratically with the electric field. Due to this 82 term, the THz driving field may change the magnetic anisotropy (see "The role of the THz electric field" in 83 Methods). Importantly, the link between the THz electric field and the spins is not restricted to a certain 84 microscopic mechanism. In our specific experiment, the THz electric field resonantly excites electronic transitions between the singlet states of the ${}^{3}H_{6}$ multiplet of the Tm³⁺ ions (Ref. 28 and Supplementary Fig. 7). 85 The concomitant non-thermal occupation change alters the magnetic anisotropy. 86

The fact that the nonlinear excitation is achieved most efficiently in the vicinity of the magnetic phase transition temperatures, where the static magnetic anisotropy is effectively zero, indicates that the excitation of the q-FM magnon is caused by the transient anisotropy. Indeed, our model shows that anisotropy changes 90 generated by the electric field of the THz pulse can drive large-angle excitations of the magnetic lattice, which 91 behaves "soft" at these temperatures (see 'The role of the THz electric field' in Methods). Figure 3d depicts 92 the temperature dependence of the modelled anisotropy torque (black curve) together with the deviation of the 93 q-FM amplitude from a linear scaling with the THz field (red data points). Our theory traces the experimental 94 data very well.

95 To put our interpretation to an ultimate test, we repeat the experiments with spectrally filtered THz pulses that selectively excite either the q-FM mode or the electronic transitions in the Tm^{3+} ions. Indeed, we find a linear 96 97 dependence of the magnon amplitude on the THz field if the q-FM mode is excited only (Fig. 4a). A comple-98 mentary THz spectrum (inset of Fig. 4b) that cannot couple by Zeeman interaction, resonantly prepares nonthermal orbital states of the Tm³⁺ ions, leading to a quadratic scaling (Fig. 4b). These results prove that the 99 nonlinear spin excitation is mediated by the rare-earth ions, which exert a strong effective torque on the Fe³⁺ 100 101 spin system. From our measurements, we extract that the strength of the anisotropy-mediated spin excitation is 102 8 times as large as the Zeeman interaction for THz pulses featuring peak magnetic fields of 0.3 T (see 103 Methods section). With the latter, peak-to-peak magnetization deflection angles of 2.6° are reached 104 (Supplementary Fig. 5). Neglecting saturation effects, we estimate that up-scaling the THz electric field to 105 ~ 3 MV/cm may suffice for non-thermal switching of the magnetization direction by 90° via the nonlinear 106 anisotropy torque (see Supplementary Fig. 8). Finally, we note that a similar nonlinear scaling of the magnon 107 amplitude with the THz field occurs in dysprosium orthoferrite ($DyFeO_3$) (see Supplementary Fig. 9), another 108 magnetic reference system, underlining the broad applicability of the new concept of a THz-induced 109 anisotropy torque.

In conclusion, we demonstrated a novel interface between THz fields and the spin system of an antiferromagnet which exploits *electric-dipole* transitions coupled to the *magnetic* degrees of freedom of electrons. In this way, we realized the first nonlinear excitation of the amplitude of spin oscillations using THz pulses. The spectral sensitivity of the effect and its high efficiency compared to the Zeeman excitation open an unprecedented doorway to further raise the amplitude of THz-driven spin deflection using pulse-shaping and coherent control. Throughout the broad class of rare-earth-transition metal compounds yet-predicted field thresholds⁸ for THz induced magnetic switching may be reduced by an order of magnitude. Our work exploits a new, general concept of electric field control of magnetic excitations by creating hidden states of matter which involve the spin degree of freedom. In the same spirit, one may now investigate the role of other lowenergy elementary excitations, such as excitons or phonons¹⁶, which could change the orbital wavefunctions of nearby atoms and lead to the creation of magnons by a related mechanism. Finally, the new principle of a symmetry-breaking preparation of low-energy non-thermal states may open unforeseen applications in future spin-based devices.

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182 Methods

Sample. A single crystal of TmFeO₃ (grown by a floating zone melting technique) with a thickness of ~ 60 μ m and lateral dimensions of ~ 5 mm was used in the experiments. The plate was cut perpendicularly to one of the crystal's optical axes which lies in the (*yz*)-plane at an angle of 51° with respect to the *z*-axis. This orientation allows for THz excitation of both magnon modes at all temperatures. A constant magnetic field of 0.1 Tesla saturates the magnetization of the sample.

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189 Setup. The detailed experimental setup is shown in Supplementary Fig. 10: A low-noise titanium:sapphire 190 laser amplifier (centre wavelength, 800 nm; pulse energy, 5.5 mJ; repetition rate, 3 kHz; pulse duration, 30 fs) 191 is used to generate intense few-cycle THz fields by tilted-pulse-front optical rectification in a cryogenically 192 cooled LiNbO₃ crystal. A pair of wire-grid polarizers and different THz spectral filters (see insets to Fig. 4 for 193 their transmission characteristics) allow for adjusting the peak fields and the spectral shape of the THz pulses. 194 A small portion of the laser pulses (pulse duration, 30 fs; pulse energy, ~ 10 nJ) is sent through a mechanical 195 delay line and is used as a polarization probe. The THz and the probe pulses are collinearly focused onto the 196 TmFeO₃ sample, which is mounted in a helium cryostat for temperature control. The THz-induced rotation of 197 the linear polarization of the probe pulses is analysed by polarization optics consisting of a half-wave plate, a 198 Wollaston polarizer and a pair of balanced silicon photodiodes. A lock-in amplifier is used to record the diode 199 signals as a function of the time delay between the THz and the probe pulses.

200

201 **Theoretical formalism**.

202 *Model for purely magnetic THz-spin interaction*. We describe the dynamics of the quasi-ferromagnetic mode 203 of the weak ferromagnet TmFeO₃ using the Lagrangian *L* and Rayleigh *R* functions of the angle θ of the 204 normalized antiferromagnetic vector **G** with respect to the crystal axis *x* (see Fig. 1a), and its time derivative 205 $\dot{\theta}$ in a form^{31,32}

206
$$L = \frac{M_{\rm Fe}}{2\gamma^2 H_{\rm E}} \dot{\theta}^2 - \frac{M_{\rm Fe}}{\gamma H_{\rm E}} B_y \dot{\theta} - W(\theta), \qquad (1)$$

207
$$R = \frac{\alpha M_{\rm Fe}}{2\gamma} \dot{\theta}^2.$$
 (2)

Here, $M_{\rm Fe}$ is the magnetization of the single Fe³⁺ sublattice, $H_{\rm E}$ is the effective field of the *d*-*d* exchange, α is the Gilbert damping parameter and $W(\theta)$ is the free energy. For TmFeO₃ subjected to the magnetic field **B** of the THz pulse, one has

211
$$W(\theta) = K_1 \sin^2 \theta + K_2 \sin^4 \theta - \frac{H_D}{H_E} M_{Fe} (B_z \cos \theta - B_x \sin \theta), \qquad (3)$$

212 where H_D is the Dzyaloshinskii field, K_2 is a constant parameter and $K_1 = 2K_2 \frac{T - T_2}{T_1 - T_2}$ with $T_1 \approx 80$ K,

213 $T_2 \approx 90$ K. Equation (3) accounts for the small spin canting angle $\varepsilon = \frac{H_D}{H_E}$ which is a result of

Dzyaloshinskii-Moriya interaction. In Eq. (3), the term containing the magnetic field arises from Zeeman coupling. The equation of motion reads

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$$\frac{d}{dt}\left(\frac{\partial L}{\partial \dot{\theta}}\right) - \frac{\partial L}{\partial \theta} + \frac{\partial R}{\partial \dot{\theta}} = 0, \qquad (4)$$

217 which, in the case of the functions (1) - (3), can be written as

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218
$$\ddot{\theta} + \omega_{\rm E} \alpha \dot{\theta} + \omega_{\rm E} \omega_{\rm A} w (\theta, T) = \omega_{\rm E} \gamma \dot{B}_{\rm y} - \frac{H_{\rm D}}{H_{\rm E}} \omega_{\rm E} \gamma B_{\rm z} \sin \theta - \frac{H_{\rm D}}{H_{\rm E}} \omega_{\rm E} \gamma B_{\rm x} \cos \theta , \qquad (5)$$

219 where γ is the gyromagnetic ratio, $\omega_{\rm E} = \gamma H_{\rm E}$, $\omega_{\rm A} = \gamma \frac{K_2}{M_{\rm Fe}}$ and $w(\theta, T) = \sin \theta \cos \theta \left(\frac{T - T_2}{T_1 - T_2} + \sin^2 \theta \right)$.

Equation (5) has the form of a generalized sine-Gordon equation and is nonlinear with respect to θ . Assuming $\ddot{\theta} = \dot{\theta} = 0$ in Eq. (5) one can find the equilibrium orientation θ_0 of the antiferromagnetic vector for a given temperature as

$$\theta_{0} = \begin{cases} \frac{\pi}{2}, T < T_{1} \\ \arcsin\left(\frac{T - T_{2}}{T_{1} - T_{2}}\right)^{\frac{1}{2}}, & T_{1} < T < T_{2} \\ 0, T > T_{2}. \end{cases}$$
(6)

224 The temperature dependence $\theta_0(T)$ is shown in Fig. 1b and Supplementary Fig. 1. We numerically find the 225 solution of Eq. (5) taking the time trace of the magnetic field B(t) from the experiment (see Fig. 2a) and assume standard initial conditions $\theta(t=0) = \theta_0$, $\dot{\theta}(t=0) = 0$. For calculations we take $M_{\text{Fe}} = 1000$ emu cm⁻³, 226 $H_{\rm E} = 2 \times 10^7$ Oe, $H_{\rm D} = 2 \times 10^5$ Oe and $\omega_{\rm E} \alpha \approx 0.05$ ps⁻¹ (Ref. 28). The resulting time evolution of the angle $\theta(t)$ is 227 228 shown in Supplementary Fig. 6 for different peak amplitudes of the driving magnetic field B_{THz} polarized 229 along the x-axis, as in the experiment. While we confirmed the possibility of excitation of the quasi-230 ferromagnetic mode via the Zeeman mechanism, we did not find any deviation from the linear relation between the maximum amplitude of θ and B_{THz} below 0.3 Tesla for any temperature and orientation of the 231 field in the (xz)-plane. Our analysis shows that the Zeeman interaction of the Fe^{3+} sublattices with the THz 232 233 magnetic field cannot lead to the nonlinear excitation observed in the experiment.

The role of the terahertz electric field. To account for the observed nonlinear interaction between the Fe^{3+} spins and the THz pulses we write the free energy term quadratic with respect to the THz electric field **E** and the antiferromagnetic vector **G** as

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$$W_{\rm int} = \sum_{i,k,l,m} g_{iklm} G_l G_m E_i E_k , \qquad (7)$$

which is allowed in centrosymmetric TmFeO₃. Here, g_{iklm} are the components of the nonlinear magnetoelectric susceptibility tensor \hat{g} . The symmetry point group D_{2h}^{16} of TmFeO₃ dictates the form of the tensor \hat{g} which reads (in the Voigt notation)

241
$$\hat{\mathbf{g}} = \begin{pmatrix} g_{11} & g_{12} & g_{13} & 0 & 0 & 0 \\ g_{12} & g_{22} & g_{23} & 0 & 0 & 0 \\ g_{13} & g_{23} & g_{33} & 0 & 0 & 0 \\ 0 & 0 & 0 & g_{4} & 0 & 0 \\ 0 & 0 & 0 & 0 & g_{5} & 0 \\ 0 & 0 & 0 & 0 & 0 & g_{6} \end{pmatrix}$$
 (8)

Taking into account that the antiferromagnetic vector in TmFeO₃ lies in the (*xz*)-plane, such that $G = (\cos \theta, 0, \sin \theta)$, we get

244
$$W_{\text{int}} = \cos^2 \theta \Big(g_{11} E_x^2 + g_{12} E_y^2 + g_{13} E_z^2 \Big) + \sin^2 \theta \Big(g_{12} E_x^2 + g_{22} E_y^2 + g_{33} E_z^2 \Big) + \frac{1}{2} g_5 \sin 2\theta E_x E_z$$
(9)

which can be rearranged as

246
$$W_{\rm int} = f(\mathbf{E}) + \sin^2 \theta \left(\chi_x E_x^2 + \chi_y E_y^2 + \chi_z E_z^2 \right) + \frac{1}{2} g_5 \sin 2\theta E_x E_z, \tag{10}$$

247 where $\chi_x = g_{12} - g_{11}$, $\chi_y = g_{22} - g_{12}$, $\chi_z = g_{33} - g_{13}$. The function $f(\mathbf{E}) = \left(g_{11}E_x^2 + g_{12}E_y^2 + g_{13}E_z^2\right)$ does not 248 depend on θ and can be omitted.

Importantly, the interaction term in Eq. (10) can be present in any crystal with D_{2h}^{16} point group regardless of the exact microscopic origin (electronic excitations, phonons, excitons, etc.) of coupling between the antiferromagnetic vector and the electric field. In our experiment the THz magnetic field is linearly polarized along the *x*-axis and therefore $E_x = 0$. Thus, the interaction energy (10) reduces to

253
$$W_{\text{int}} = (\chi_y E_y^2 + \chi_z E_z^2) \sin^2 \theta$$
(11)

and can be seen as a modulation of the anisotropy energy $W_A(\theta) = K_1 \sin^2 \theta + K_2 \sin^4 \theta$. The THz electric field changes the anisotropy parameter K_1 in Eq. (3) by $\Delta K_1 = \chi_y E_y^2 + \chi_z E_z^2$.

The coupling described by Eq. (11) generates a torque acting on spins which results in an additional term on the right-hand side of the generalized sine-Gordon equation (7) as

258
$$\ddot{\theta} + \omega_{\rm E} \alpha \dot{\theta} + \omega_{\rm E} \omega_{\rm A} w(\theta, T) = -\omega_{\rm E} \omega_{\rm A} \sin \theta \cos \theta \, a(t), \tag{12}$$

with 259

$$a(t) = \chi_y E_y^2(t) + \chi_z E_z^2(t)$$
⁽¹³⁾

being proportional to the THz intensity time trace. In Eq. (12) we do not include the additive Zeeman torque considered above to isolate the effect of the THz-induced transient anisotropy. If the duration of the THz pulse and the relaxation time of the excited Tm^{3+} states is much shorter than the period of the quasi-ferromagnetic mode one can show that the generated torque acts as an instantaneous excitation described by the initial conditions

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$$\theta(t=0) = \theta_0, \quad \dot{\theta}(t=0) = -\omega_{\rm E}\omega_{\rm A}\sin\theta_0\cos\theta_0\int_0^{\tau_{\rm THz}} dt, \qquad (14)$$

where τ_{THz} is the duration of the THz pulse. The transient anisotropy torque depends on the temperature as $\sin \theta_0(T)\cos \theta_0(T)$ which is shown in Fig. 3d. It is non-zero only in the temperature interval corresponding to the intermediate magnetic phase Γ_{24} . Obviously, the torque is quadratic with respect to the THz field **E** (Eq. (13) and (14)). The Zeeman torque and the transient anisotropy torque are additive, but scale differently with the THz peak field.

Finally we note that our phenomenological theory correctly reproduces the spin dynamics on timescale longer than the duration of the THz-driven perturbation of the anisotropy fields. To describe the strongly nonequilibrium state during the interaction of the medium with the intense THz pulse, one may need to account for the THz carrier wave within a time-dependent many-body theory.

Estimation of the strength of the Zeeman torque. Applying a THz low-pass filter with a nominal cut-off frequency of 0.3 THz allows us to selectively excite the q-FM mode (see inset to Fig. 4a). In this measurement, the q-FM amplitude amounts to $A_z = 8\%$ of the one induced by the unfiltered THz transients (Fig. 4a). Taking into account the filter transmission of $T_F = \sim 75\%$ at the q-FM resonance frequency, we conclude that the anisotropy induced torque is $\frac{1-A_z/T_F}{A_z/T_F} \approx 8$ times as large as the Zeeman interaction.

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285 Acknowledgements

286 The authors thank M. Furthmeier for technical assistance, R.V. Pisarev for providing the samples, and T. L. Cocker for stimulating discussions. Continuous support by Th. Rasing is acknowledged. S.B., M.H. and 287 R.H. were supported by the European Research Council through ERC Grant No. 305003 288 289 (QUANTUMsubCYCLE) and Deutsche Forschungsgemeinschaft (DFG) through collaborative research 290 center SFB 689. A.V.K., R.V.M and A.K.Z. were supported by the European Community Seventh Framework 291 Programme FP7-NMP-2011-SMALL-281043 (FEMTOSPIN), the European Research Council ERC Grant 292 agreement No. 257280 (Femtomagnetism), the Foundation for Fundamental Research on Matter (FOM) as 293 well as the Netherlands Organization for Scientific Research (NWO) and program "Leading Scientist" of the 294 Russian Ministry of Education and Science (14.z50.31.0034). T. K. thanks the Deutsche 295 Forschungsgemeinschaft and the European Research Council for support through priority program SPP 1538 296 and the ERC Grant No. 681917 (TERAMAG), respectively.

297 Author contributions

S.B., A.V.K., R.H. and R.V.M. conceived the study, carried out the experiments and analysed the data. A.K.Z.
and R.V.M. developed the theoretical model. S.B., M.H., A.V.K., R.H., R.V.M. wrote the manuscript. All
authors discussed the results.

301 Additional Information

302 Supplementary information is available in the online version of the paper. Reprints and permissions 303 information is available online at www.nature.com/reprints. Correspondence should be addressed to R.H. 304 (rupert.huber@physik.uni-regensburg.de) or R.V.M. (R.Mikhaylovskiy@science.ru.nl).

305 **Competing financial interests**

306 The authors declare no competing financial interests.

307 Figure legends

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- **Figure 1** | **Principle of spin control by a THz induced anisotropy torque. a,** Spin and lattice structure of TmFeO₃
- 310 arrows) form two antiferromagnetically coupled sublattices M_1 and M_2 , which are mutually canted by the

shown in the Γ_{24} phase. Green/blue spheres: Tm³⁺/Fe³⁺ ions. Oxygen atoms are not shown for clarity. The iron spins (blue

- 311 Dzyaloshinskii-Moriya interaction, giving rise to a weak ferromagnetic moment $\mathbf{F} = \mathbf{M}_1 + \mathbf{M}_2$. In the Γ_{24} phase (i.e. for
- 312 80 K < T < 90 K), the antiferromagnetic vector $\mathbf{G} = \mathbf{M}_1 \mathbf{M}_2$ encloses a finite angle $0^\circ < \theta_0 < 90^\circ$ with the x-axis. **b**,
- 313 Spin reorientation phase transitions: In the Γ_2 phase (i.e. T < 80 K) the antiferromagnetic vector **G** is aligned along the
- 314 crystallographic *z*-axis, whereas it lies along the *x*-axis above T = 90 K (Γ_4 phase). In the Γ_{24} phase, **G** rotates 315 continuously in the (*xz*)-plane (see also panel **a**). **c**, The crystal field splits the ground state ${}^{3}H_{6}$ of the rare-earth Tm³⁺ ions 316 into several energy levels with an energy spacing of $\sim 1 - 10$ meV (schematic level scheme). The corresponding orbital 317 wavefunctions set the magnetic anisotropy for the iron spins in thermal equilibrium (upper panel). Lower panel: Ultrafast 318 transitions between these energy levels resonantly induced by THz pulses should exert an abrupt torque on the spins and
- act as an efficient trigger for coherent spin dynamics. The small canting angle is not shown for clarity.
- Figure 2 | Overview of the experiment. **a**, THz transients used to excite magnon/Tm³⁺ resonances in TmFeO₃. **b**, Amplitude spectrum of the waveform shown in panel **a**. Arrows mark the frequencies of the magnon/Tm³⁺ resonances. **c**, Schematic of the experiment: THz-pump (red) and near infrared probe pulses (NIR, blue) are collinearly focused onto the TmFeO₃ sample with variable time delay *t*. Using a $\lambda/2$ plate, a Wollaston prism, and two balanced photodiodes, THz-induced magnetic dynamics in TmFeO₃ are measured by polarization rotation of the probe pulses. **d**, Resonance frequencies of the q-FM (red circles) and q-AFM (blue triangles) modes in dependence on the sample temperature *T*. Black curves are guides to the eye.
- Figure 3 | Nonlinear THz-magnon interaction. a, Normalized magnon traces for various THz excitation strengths B_{THz} : 327 328 Whereas quasi-monochromatic oscillations are found for the lowest THz field, a low-frequency oscillation is 329 superimposed onto the dynamics for higher pump fields. b, Amplitude spectra of the time domain data shown in panel a 330 allow for the identification of the q-FM and q-AFM modes at 100 and 830 GHz, respectively. c, Scaling of the 331 amplitudes from panel b: The q-AFM mode (blue triangles) scales linearly with the THz field strength, whereas the q-332 FM mode (red circles) shows a quadratic dependence on the latter. Error bars denote the standard deviation interval for 333 the THz amplitude, arising from uncertainties in the THz spot size and the repeatability of the polarizer angle. 334 d, Deviation of the experimental field-scaling of the q-FM mode with the THz field strength from a linear behaviour (red 335 data points) and THz-induced anisotropy torque exerted on the spins (black curve) as computed by our model for various temperatures at the magnetic phase transitions. The nonlinear behaviour vanishes outside the Γ_{24} phase. Error bars take 336 337 account of the uncertainty in the extracted nonlinearity owing to noise in the measurements.

Figure 4 | **Control of THz-induced nonlinear torque by spectral shaping: a,** Low pass filtering the pristine THz spectrum (grey shaded curve in inset) results in the black shaded spectrum (inset) featuring a dominant maximum at the q-FM resonance frequency. Main graph: Linear scaling of the q-FM amplitudes obtained by Zeeman excitation with the constrained THz spectrum (red circles). The q-AFM mode (blue triangles) is suppressed since the THz amplitude at the corresponding frequencies is strongly reduced by the low pass filter. **b**, The Zeeman type excitation of the q-FM mode is

- 343 switched off by bandpass filtering (centre frequency 1.2 THz) of the THz pulse (black/grey: filtered/pristine spectrum).
- 344 Main graph: Tm³⁺ excitation triggers q-FM oscillations, whose amplitudes scale quadratically with the THz fields (red
- 345 circles). Additionally, q-AFM magnons (amplitude: blue triangles) are excited by the Zeeman interaction. All curves are
- normalized to the maximum amplitude of the q-FM mode shown in Fig. 3c. Dotted/solid lines are linear/quadratic fits.
- 347 Error bars are retrieved as in Fig. 3c.