

School of Natural Sciences and Mathematics

***Missing Magnetism in $Sr_4Ru_3O_{10}$: Indication
for Antisymmetric Exchange Interaction***

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Missing magnetism in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$: Indication for Antisymmetric Exchange Interaction

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Metamagnetism occurring inside a ferromagnetic phase is peculiar. Therefore, $\text{Sr}_4\text{Ru}_3\text{O}_{10}$, a $T_C = 105$ K ferromagnet, has attracted much attention in recent years, because it develops a pronounced metamagnetic anomaly below T_C for magnetic fields applied in the crystallographic ab -plane. The metamagnetic transition moves to higher fields for lower temperatures and splits into a double anomaly at critical fields $H_{c1} = 2.3$ T and $H_{c2} = 2.8$ T, respectively. Here, we report a detailed study of the different components of the magnetization vector as a function of temperature, applied magnetic field, and varying angle in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$. We discover for the first time a reduction of the magnetic moment in the plane of rotation at the metamagnetic transition. The anomaly shifts to higher fields by rotating the field from $H \perp c$ to $H \parallel c$. We compare our experimental findings with numerical simulations based on spin reorientation models taking into account magnetocrystalline anisotropy, Zeeman effect and antisymmetric exchange interactions. While Magnetocrystalline anisotropy combined with a Zeeman term are sufficient to explain a metamagnetic transition in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$, a Dzyaloshinskii-Moriya term is crucial to account for the reduction of the magnetic moment as observed in the experiments.

$\text{Sr}_4\text{Ru}_3\text{O}_{10}$ belongs to the Ruddlesden-Popper family of ruthenium oxide perovskites $\text{Sr}_{n+1}\text{Ru}_n\text{O}_{3n+1}$. This class of metallic compounds caught much attention in recent years due to its rich variety of ground states. Sr_2RuO_4 the $n = 1$ member, is discussed as an example of rare p -wave superconductivity¹. A quantum critical endpoint covered by a high entropy phase was found in the $n = 2$ layer system $\text{Sr}_3\text{Ru}_2\text{O}_7$ ². The compound $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ ($n = 3$) discussed here shows ferromagnetism below $T_C = 105$ K³. Neutron diffraction experiments in zero magnetic field reveal ordering of the Ru moments along the c -axis. No ferromagnetic (FM) or antiferromagnetic (AFM) correlations are observed in the ab -plane^{4,5}. $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ contains four inequivalent Ru sites with two different magnetic moments of $0.9 \mu_B$ and $1.5 \mu_B$ sitting on outer and inner RuO layers, respectively. The magnetic unit cell contains 8 of the smaller and 4 of the larger magnetic moments averaging to $1.1 \mu_B$ per Ru. The higher order ruthenate SrRuO_3 with $n = \infty$ also orders FM at a Curie temperature of 165 K^{6,7}. The $\text{Sr}_{n+1}\text{Ru}_n\text{O}_{3n+1}$ are strongly 2-dimensional electron systems with the trend to become more isotropic for higher n , because of their layered structure. Two-dimensionality is reflected in anisotropic transport properties as seen for $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ in the ratio of the electrical resistivity $\rho_c/\rho_{ab} \simeq 400$ ⁸ and confirmed by optical conductivity experiments⁹.

Metamagnetism is a phenomenon observed in magnetic materials, where hidden magnetism is suddenly uncovered by the application of an external field. The origin of metamagnetism can be spin flip transitions in antiferromagnets^{10–12}, but also changes of the band structure in itinerant electron systems. Latter scenarios are in the vast majority described on the basis of the well-known Stoner model¹³ and refined^{14–16}, to accommodate special cases, e.g. Fermi surface reconstruction^{17,18} or in the vicinity of a quantum critical point^{19,20}. In a general description, metamagnetism is a phase transition or crossover from a magnetically disordered or ordered state with small net magnetization to a field polarized (FP) or partially FP state. In the case of $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ the term metamagnetism refers to the sudden increase in the magnetization when the field applied in the (ab) in-plane of this layered compound exceeds 2 T. Magnetism is hidden only because the spontaneous moment is mainly

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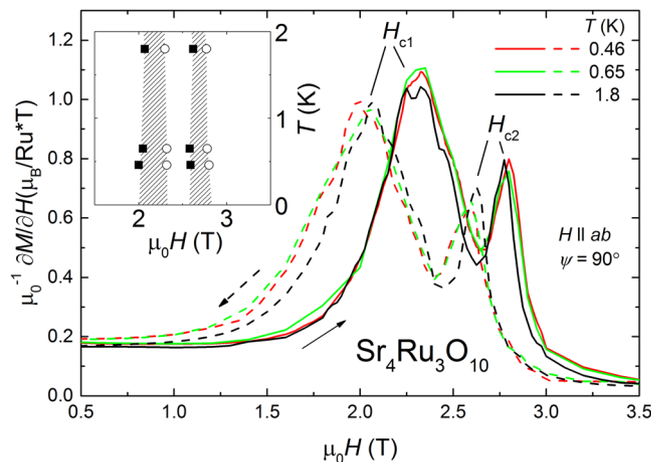


Figure 1. Field derivative of the magnetization $\partial M/\partial H$ at 1.8 K, 0.65 K and 0.46 K with clear anomalies at two critical fields $H_{c1,2}$. Solid lines show measurements during increasing field sweeps and dashed lines during decreasing field sweeps as labeled by arrows. The inset shows $H - T$ phase diagram close to the MM transition with regions of hysteresis marked as striped patterns.

aligned with the easy c -axis at smaller fields; nonetheless, we continue to refer to this as metamagnetic (MM) transition. Interestingly, the saturation magnetization above the MM transition ($H \perp c$) is about 10% smaller than the saturated moment for $H \parallel c$ ^{3,21}, which points to a more complex underlying scenario for the metamagnetism than just a simple spin flip transition. While the MM transition in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ was discovered from early on in flux grown single crystals^{3,21}, it took more than a decade to improve the crystal quality to a level to see a double step in the magnetization at the MM transition²². This strong dependence of physical properties on the crystal purity is a characteristic signature of strontium ruthenates $\text{Sr}_{n+1}\text{Ru}_n\text{O}_{3n+1}$ and was also observed in the sister compound $\text{Sr}_3\text{Ru}_2\text{O}_7$ ^{2,23–25}. The MM transition in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ develops below 68 K as a double-transition close to zero field and shifts gradually to about 2.5 T with temperatures down to 1.7 K²². Carleschi *et al.*²² speculate that the double transition originates either in the ordering of Ru magnetic moments on two inequivalent crystallographic sites or in the presence of two van Hove singularities in the density of states close to the Fermi level. A transport study based on electrical resistivity^{26,27} reveals steps in the magnetoresistance at various critical fields around H_c accompanied by pronounced hysteresis. Fobes *et al.*²⁷ interpret the transport data as domain movement of regions with high and low electronic spin polarization. Anomalous behavior at the MM transition was also observed in specific heat experiments²⁸ up to 9 T and in thermopower investigations²⁹. Neutron diffraction experiments up to 6 T reveal a change of lattice parameters at the critical field H_c ⁴. Field and pressure dependent Raman measurements³⁰ as well as a recent study of thermal expansion and magnetostriction³¹ confirm strong magnetoelastic coupling in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$.

This work aims to increase our understanding of MM phenomena in $4d$ oxides in general and the peculiar MM transition inside the ferromagnetic order of $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ in particular. We carry out magnetization measurements up to 7 T and down to lowest temperatures of 0.46 K and under rotational fields between the c -axis and the ab -plane as well as (ab) in-plane rotation. Our investigations include a detailed analysis of the behavior of the magnetization modulus M at the MM transition and its individual components M_{ab} and M_c simultaneously. In the following, we analyze and interpret our data in a localized picture, meaning the magnetic moments are mainly confined on the Ru^{4+} sites in the crystal structure of $\text{Sr}_4\text{Ru}_3\text{O}_{10}$. This scenario is supported by neutron diffraction experiments which have determined the spin and orbital momentum distribution in great detail^{4,5}. Our main discovery is the observation of a reduced measured moment at the MM transition caused by a spin component pointing out of the rotational plane which we assert can best be explained by significant anisotropic exchange interactions in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$.

Results

At first, we focus on the magnetization measured for $H \perp c$ at temperatures below 2 K. Figure 1 shows the susceptibility $\partial M/\partial H$ between 0.5 T and 3.5 T. We observe a clear double MM phase transition with a main anomaly at $H_{c1} = 2.3$ T and a second anomaly at $H_{c2} = 2.8$ T for increasing field as observed by Carleschi *et al.*²². Our new experimental data down to 0.46 K clarify that the transition neither sharpens to lower temperatures nor is there splitting into more distinct anomalies. The inset in Fig. 1 shows the $H - T$ phase diagram with near-vertical phase boundaries for $T \rightarrow 0$ at the MM transition. Both anomalies are shifted by -0.3 T for measurements in decreasing magnetic field. The size of the hysteretic region, marked as striped pattern, remains similar for all temperatures below 1.8 K.

The operation mode of the SQUID magnetometer allows the simultaneous collection of longitudinal M_{long} and transversal component M_{trans} of the sample magnetization in respect to the applied magnetic field \vec{H} as sketched in the inset of Fig. 2 (see Methods). Any magnetization component M_{perp} occurring perpendicular to the rotational plane is not recorded during the measurements. This geometry allows us to calculate the magnetization

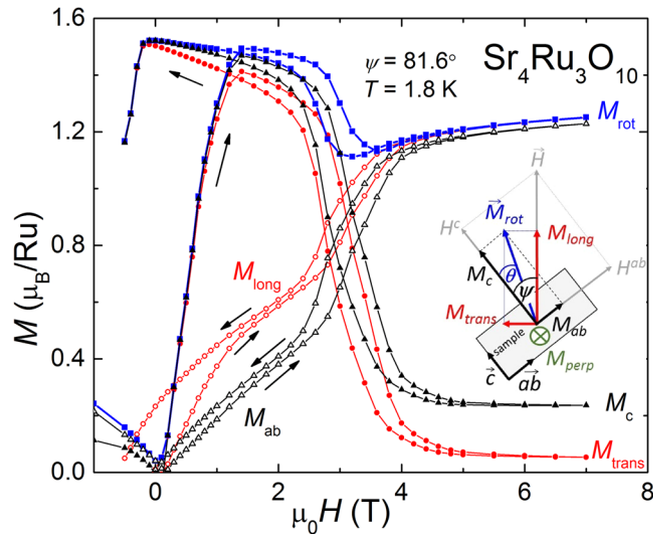


Figure 2. The inset shows a geometrical sketch of the sample $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ mounted inside the SQUID magnetometer. ψ is the rotation angle of the applied field \vec{H} and θ the angle of the magnetization \vec{M} , both in respect to the magnetic easy c -axis. M_{long} and M_{trans} are measured components of \vec{M} parallel and perpendicular to the applied field in the plane of rotation. The component M_{perp} parallel to the axis of rotation is not captured during the measurement. The main panels compares the different components of the magnetization M_{long} , M_{trans} , M_{ab} , M_c , and the modulus M_{rot} versus magnetic field H measured at 1.8 K for $\psi = 81.6^\circ$.

$M_{\text{rot}}^2 = M_{\text{long}}^2 + M_{\text{trans}}^2$ in the rotational plane. The knowledge of the rotation angle ψ and relation $\tan(\psi - \theta) = M_{\text{trans}}/M_{\text{long}}$ enables the determination of the angle of the magnetization θ with respect to the magnetic easy axis c in the rotational plane. We can now calculate $M_{\text{ab}} = M_{\text{rot}} \sin \theta$ and $M_c = M_{\text{rot}} \cos \theta$, the magnetization occurring in the plane of rotation. Note, we follow closely the notation of angles used for magnetic anisotropic materials. Figure 2 illustrates the different components of the magnetization for one particular measurement with $\psi = 81.6^\circ$ taken at 1.8 K. Prominent feature is the hysteresis loop around ± 1 T, caused by FM domain dynamics. The longitudinal magnetization M_{long} increases moderately in small fields and shows a sudden rise at $H_{c1,2} \simeq 3$ T at the MM transition. M_{trans} on the other hand consists mainly of the M_c component with a sudden decrease of the magnetization at the same critical fields $H_{c1,2}$. $M_{\text{trans}} \neq 0$ above $H_{c1,2}$ indicates incomplete field polarization meaning that \vec{M} is not perfectly aligned with \vec{H} . This observation points to the presence of magnetic anisotropy. The calculated magnetization modulus M_{rot} in the plane of rotation is depicted as black line in Fig. 2. We find a maximum moment of $1.5 \mu_B$ slightly higher than obtained in neutron experiments^{4,5}, but in good agreement with previous magnetization studies^{21,32}. Most peculiar is that M_{rot} drops suddenly below $1.2 \mu_B$ at the MM transition and only recovers partially to $1.2 \mu_B$ up to maximum applied field of 7 T. This missing component of the magnetic moment in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ was never recognized before. Furthermore, we observe strong hysteresis at the MM transition between up and down measurements as reported in previous investigations^{8,21,22,27,32}.

Geometrical effects can distort magnetic properties during magnetization experiments. To avoid this problem, we plot in Fig. 3 M_{ab} as a function of the field component in the ab -plane $H^{ab} = H \sin \psi$ to examine how $H_{c1,2}$ change with ψ . In contrast to previous results by Jo *et al.*³³ obtained by torque magnetometry, we observe a clear simultaneous increase of both critical fields $H_{c1,2}$ to higher values while rotating from $H \perp c$ to $H \parallel c$. In fact, $H_{c1,2}$ move out of the observable field range of $H \leq 7$ T for $\psi \lesssim 72^\circ$. A similar shift to higher critical fields was observed in measurements of the longitudinal magnetoresistance for currents $j \parallel c$ and $j \perp c$ as a function of rotating magnetic field as reported by Fobes *et al.*^{8,27}.

Figure 4 summarizes the critical fields $H_{c1,2}$ in the $H - \psi$ phase diagram for field up and down sweep measurements. The difference $H_{c1} - H_{c2}$ increases slightly with smaller ψ . As mentioned above, the double anomaly is accompanied by significant hysteresis. The inset of Fig. 4 shows the evolution of combined step size ΔM_{ab} of both MM transitions for decreasing ψ which were extracted from the curves in Fig. 3. It follows a quadratic fit function marked as solid line and extrapolates to zero step size at about 65° . The magnetization modulus M_{rot} recorded in the plane of rotation for ψ between 85.3° and 70.5° is plotted in Fig. 5. We only show field-down sweep measurements for clarity. Striking is the occurrence of a drop from about $1.5 \mu_B$ to below $1.2 \mu_B$ at the critical field of the main anomaly H_{c1} followed by a minimum and a small step at the second anomaly at H_{c2} . The described features are marked in Fig. 5 by arrows for the measurement at $\psi = 85.3^\circ$. The MM anomaly broadens and moves to higher fields for decreasing angles ψ .

So far, we only carried out magnetization measurements under rotational fields at 1.8 K and did not expand to higher temperatures. Therefore, we cannot conclude with certainty, how the “moment loss” changes with increasing temperature. We have shown, however, that H_{c1} and H_{c2} of the double step in M_{ab} are intimately connected to the missing moment and represent basically the same critical magnetic fields, where the magnetization drops and partially recovers. Please compare Figs 3 and 5. Carleschi *et al.* found out in a temperature study²² that the

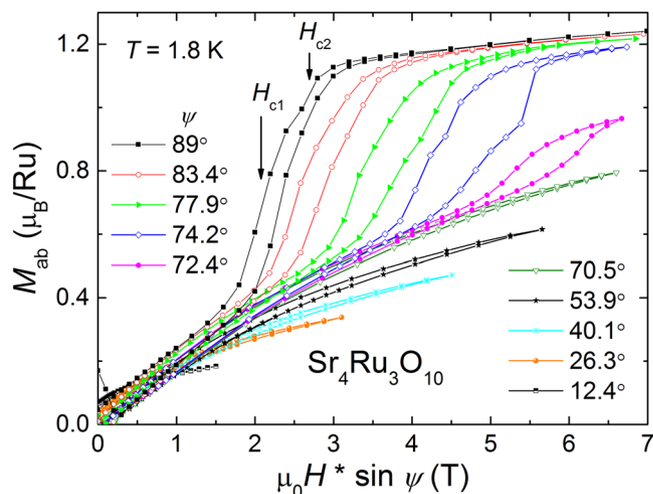


Figure 3. ab -plane magnetization M_{ab} as a function of $H^{ab} = H \sin \psi$ is shown for selected angles ψ between 89° and 12.6° measured at 1.8 K. A clear double step is observed at $H_{c1,2}$ as labeled by arrows for $\psi = 89^\circ$.

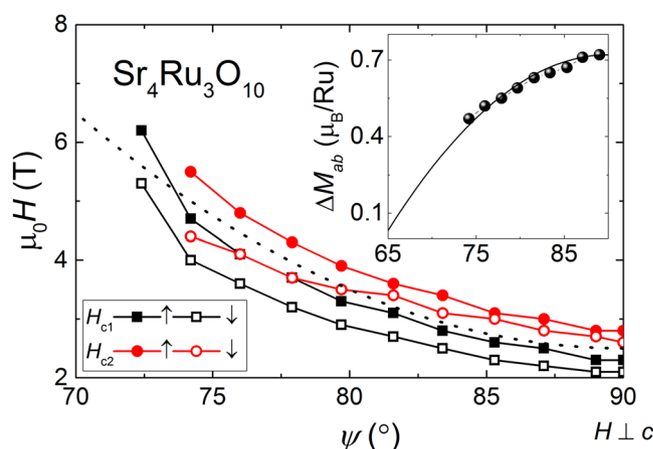


Figure 4. Angular ψ dependent shift of the double anomaly at the MM transition in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$. Solid points mark positions for increasing and empty points decreasing field sweeps. The dotted line is a quadratic fit to the data. The inset shows the reduction of the magnetization step ΔM_{ab} at the MM transition as a function of ψ including a quadratic extrapolation marked as solid line.

double transition occurs right below the ferromagnetic ordering temperature $T_C = 105$ K. This observation makes us believe that the effect of a missing moment occurs right below T_C as well with some temperature broadening.

Discussion

The “loss” of magnetic moment in the rotational plane can be explained either by partial AFM alignment or by a moment M_{perp} occurring perpendicular to the rotation (parallel to rotation axis of ψ). The first scenario can be excluded based on neutron experiments where no short or long range AFM coupling neither in zero nor in magnetic fields $H > H_{c1,2}$ was observed in the ab -plane^{4,5}. The second scenario is rather unexpected since magnetic moments tend to align with field and stay within the rotational plane, if no further coupling is present. We want to focus in our discussion on two mechanisms that potentially lead to a M_{perp} component in the magnetization. First one is based on general magnetocrystalline anisotropy in tetragonal symmetry, with an easy c -axis and 4-fold in-plane anisotropy. The second mechanism is antisymmetric exchange between spins, also called Dzyaloshinskii-Moriya (DM) interaction, causing a canting of the spins $\vec{S}_i \times \vec{S}_j$.

We have to have a closer look at the crystal structure of $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ in order to understand and model its magnetic anisotropy caused by spin-orbit coupling. $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ crystals consist of three layers of corner sharing RuO_6 octahedra separated by a double layer of Sr-O. Primary Bragg reflections in synchrotron experiments can be indexed assuming a tetragonal unit cell with space-group $I4/mmm$, but a more detailed analysis of secondary reflections reveals orthorhombic $Pbam$ symmetry³. The lower symmetry originates in c -axis rotation of the RuO_6 octahedra that are correlated between different layers, meaning $+11.2^\circ$ clockwise rotation for inner and -5.6° counterclockwise rotation for outer layers.

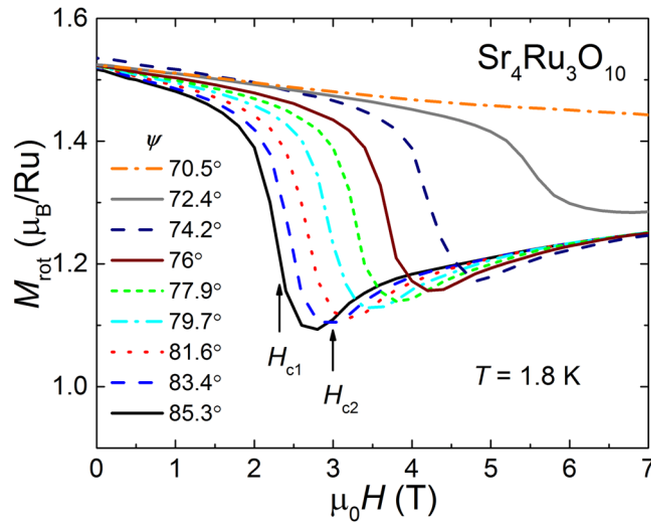


Figure 5. Magnetization modulus M_{rot} in the rotational plane versus magnetic field H for angles ψ between 85.3° and 70.5° in decreasing fields. The drop in M_{rot} coincides with H_{c1} and the step-like increase with H_{c2} as marked by arrows for the measurement at $\psi = 85.3^\circ$.

The free energy F accounting for magnetocrystalline anisotropy in a tetragonal lattice, can be modeled³⁴ by

$$F = F_0 + K_1 \sin^2 \theta + (K_2 + K_3 \sin(2\varphi)) \sin^4 \theta - F_Z. \quad (1)$$

F_0 is a constant background contribution independent of \vec{H} or \vec{M} . \vec{M} is expressed in polar coordinates (θ, φ) , with $\theta = 0$ along the crystal c -axis and $\varphi = 0$ defining the in-plane hard axis for $K_3 < 0$. Here, $K_1 > 0$ defines the easy direction and the Zeeman term F_Z can be written as

$$F_Z = MH(\sin \theta \sin \psi (\cos \omega \cos \varphi + \sin \omega \sin \varphi) + \cos \theta \cos \psi). \quad (2)$$

The applied field has the spherical coordinates (ψ, ω) , with $\omega = 0$ corresponding to field rotation from the c -axis to the in-plane hard direction and $\omega = \pi/4$, field rotation to the easy direction. We used numerical minimization of equation (1) to determine θ and φ as functions of the applied field. In the uniaxial case with $K_3 = 0$, the MM behavior in M_{ab} and M_c at the critical field H_c is reproduced by choosing correct parameters K_1 and K_2 (data not shown). However, uniaxial anisotropy is unable to reproduce any reduction of the magnetization in M_{rot} since the moment always stays in the rotational plane. Therefore, we considered in-plane anisotropy with $K_3 \neq 0$ in the next step.

Note, we do not know precisely the in-plane orientation of our sample. However, the rectangular shape suggests that ψ rotation axis is parallel to one of the principal axes such as [100] or [110]. For 4-fold tetragonal symmetry either one of them would be the intermediate or hard axis, respectively. We consider in the following a projection of \vec{H} onto the magnetic hard axis in the ab -plane with $\omega = 0$, because tilting of \vec{M} towards the hard axis forces the magnetic moments to align spontaneously toward either one of the intermediate axes, which are 45° apart from the hard axis. This spontaneous alignment $\pm 45^\circ$ is energetically degenerated and could lead to domain formation with an overall smaller net magnetization as observed in our measurements.

Figure 6 compares numerical results of $\psi = 78^\circ, 81^\circ, 84^\circ$ based on equation (1) with experimental data of the magnetization modulus M for $\psi = 77.9^\circ, 81.6^\circ, 83.4^\circ$. We are able to reproduce i) a critical field value of about 2.5 T that increases with smaller ψ , ii) a drop ΔM at $H_{c1,2}$ that is comparable in size with the experimental data, and iii) a gradual slope $M(H)$ for $H > H_{c1,2}$. The double feature at the MM transition is missing due to the simplicity of the model. We obtain anisotropy parameters $K_1 = 3.1$ K, $K_2 = 0.1$ K and $K_3 = -2.2$ K in Kelvin energy scale which convert to the following values 300 kJ/m³, 10 kJ/m³, and -210 kJ/m³, respectively, in units widely used in magnetic anisotropy tables. The 4th order parameter K_2 being more than 10 times smaller than K_1 implies that it is irrelevant for the description of the anisotropy in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$. For comparison, the 3d FM metal cobalt has anisotropy constants of $K_1 = 450$ kJ/m³ and $K_2 = 150$ kJ/m³, which are of similar size as K_1 in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ ³⁵.

Despite the reasonable agreement between experiment and model, it is necessary to check in a subsequent experiment our initial assumption of tilting the field towards the magnetic hard axis in the ab -plane. Therefore, we rotate the sample by $\omega = 45^\circ$ in the plane and measure again M at three different angles ψ as shown in Fig. 7. We anticipate the 45° change would bring the intermediate anisotropy axis into the rotation plane. Specifically, the magnetization would rotate toward the intermediate axis with the total magnetization remaining in the rotation plane and therefore no “loss” of magnetic moment effect. Surprisingly, the magnetization M_{rot} shows exactly the same behavior as for the $\omega = 0$ experiments within experimental uncertainty. Even if in both experiments $\omega = 0$ and 45° , the plane of ψ -rotation would not include exactly the principal axes, we would at least expect the observation of a reduced anomaly in M_{rot} at H_c . Based on our last finding, we exclude general magnetic anisotropy as sole cause for the reduction of moment at the MM transition in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$.

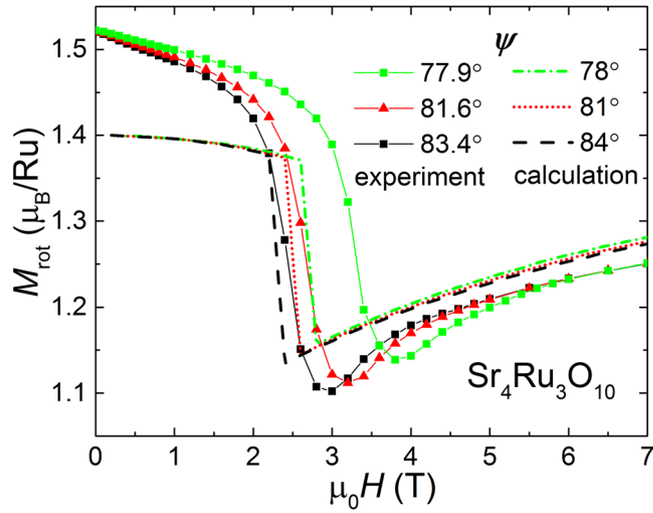


Figure 6. The experimental magnetization moduli M_{rot} for 3 different angles ψ (symbols) measured at 1.8 K in decreasing magnetic field are compared with numerical simulations (broken lines) of the general anisotropy model as described by Eq. (1).

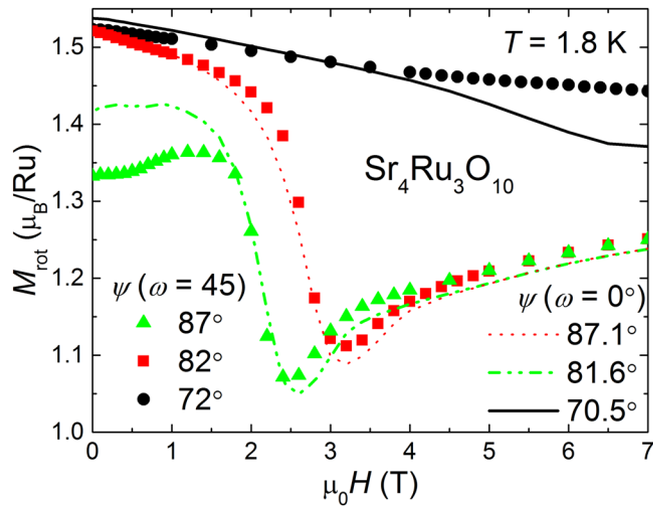


Figure 7. Decreasing field H measurements of the magnetization modulus M_{rot} taken at 1.8 K at three angles $\psi \simeq 87^\circ$, 82° , and 71° are shown for two different in-plane angles $\omega = 0$ and 45° . The almost identical results for 0 and 45° disable magnetocrystalline anisotropy being the cause for the loss of magnetic moment in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$.

Anisotropic exchange interactions caused by spin-orbit coupling under certain symmetry constraints were first considered by *Dzyaloshinsky and Moriya*^{36,37} to explain weak FM ordering inside an AFM phase in transition metal oxides. Recently, *Bellaiche et al.*³⁸ pointed out that tilting of oxygen octahedra in perovskites can be described by a rotation (pseudo) vector ν_i sitting at position i of spin S_i . The oxygen octahedra rotation leads to an energy reduction

$$\Delta E = K \sum_{ij} (\nu_i - \nu_j) \cdot (\vec{S}_i \times \vec{S}_j) \quad (3)$$

in analogy to DM antisymmetric exchange coupling. The summation is done over nearest-neighboring spins $\vec{S}_{i,j}$ and K is a constant. Consequently, we approximate a DM interaction between the in-plane component of the Ru center magnetization and that of the two nearest-neighbor Ru atoms, whose spins are assumed to remain parallel. This gives rise to an effective energy term

$$F_{DM} = -D \sin^2(\theta) \sin(2\varphi) \quad (4)$$

in replacement of the K_3 term of the magnetocrystalline anisotropy. Angle φ is interpreted as the angle between in-plane magnetic moments sitting on one inner and two outer layers with $\varphi = 0$ being the direction of in-plane magnetic field. The change in the dependence on angle φ between the magnetization vector and the c -axis

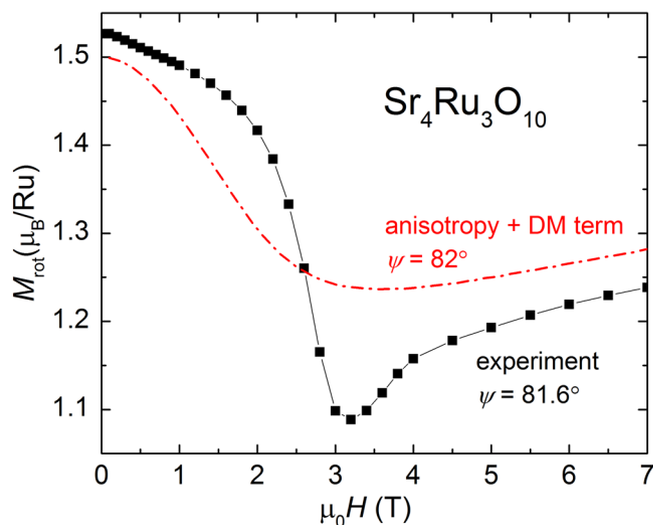


Figure 8. Experimental magnetization modulus M_{rot} (symbol) in the plane of rotation as a function of magnetic field $\mu_0 H$ up to 7 T and numerical results including tetragonal magnetocrystalline anisotropy and a DM like energy term (broken line) for $\psi \approx 82^\circ$.

prevents an accurate minimization of the total energy. Nonetheless, the approximate solution shown in Fig. 8 for $\psi = 82^\circ$ does produce a “missing” portion of the total magnetization, with the minimum moving to higher field with decreasing ψ . The parameter D is approximately 5.3 K and comparable to the energy scale of the magnetocrystalline anisotropy K_1 .

We want to point out that the opening of 2φ between the inner and outer magnetic moments can be seen as AFM order if it occurs periodically along the c -axis. Neutron scattering experiments on periodic holmium-yttrium superlattices³⁹ e.g., were able to distinguish between different types of AFM periodicity along $[00l]$ with increasing in-plane magnetic field, such as helical, helifan-shaped, fan-shaped and FM order. However, the particular crystal structure of $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ with three inequivalent layers of RuO_6 octahedra connected through a double layer of Sr-O doesn't give rise to additional periodicity, even if the orientation of the magnetic moments follows a repetitive pattern such as $-\varphi, +\varphi, -\varphi$ along c inside the triple layer. As mentioned before, the inner and outer Ru positions are crystallographic inequivalent and carry different magnetic moment sizes. Consistently, in-plane AFM ordering was excluded by neutron scattering studies⁴. Another experimental route to gain insight into the nature of the missing moment is the exploration of transversal magnetoresistance $\rho_c(H, T)$ as a function of temperature along the c -direction with in-plane magnetic field ($j \parallel c, j \perp H$). In case of a periodic opening 2φ of neighboring spins in different layers, we expect to observe coherent scattering and a decreasing magnetoresistance with lower temperature and in higher magnetic field. In case the missing magnetic moment is caused by domain formation of any sort, ρ_c should also go down with increasing magnetic field, but stay temperature independent because of scattering on domain walls.

In summary, the MM transition in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$ as a function of magnetic field and rotation angle between $H \perp c$ and $H \parallel c$ has been studied in great detail by magnetization measurements down to lowest temperatures in a SQUID magnetometer. Our experimental results reveal a reduced magnetic moment in the plane of rotation which was never recognized before. It is robust to (ab) in-plane rotation. We find furthermore that the double step at the MM transition is stable down to lowest temperatures of 0.46 K. Our experimental results are interpreted in a strict localized picture with magnetic moments of different sizes sitting on inner and outer RuO_6 layers in the crystal structure. We completed our study with numerical calculations based on energy minimization including Zeeman effect, magnetocrystalline anisotropy and antisymmetric exchange and compared them with our experimental data. We conclude that all three contributions are essential ingredients to understand the behavior of the magnetization and that a *Dzyaloshinsky - Moriya* like component is crucial to model a reduced magnetic moment in $\text{Sr}_4\text{Ru}_3\text{O}_{10}$.

Methods

$\text{Sr}_4\text{Ru}_3\text{O}_{10}$ single crystals were grown in an image furnace by a floating zone technique⁴⁰ and characterized by energy dispersive spectroscopy, scanning electron microscopy, electron backscattering diffraction and x-ray diffraction techniques.

Magnetization measurements in fields up to 7 T were carried out in a Quantum Design MPMS SQUID magnetometer equipped with a standard ^4He setup for measurements between 1.7 K and 100 K and in an iQuantum ^3He insert that fits inside the MPMS sample space for the temperature range 0.46 K to 2 K. Excellent agreement between the ^3He and ^4He data was observed in the temperature range of overlap. Angular dependent measurements at 1.8 K were obtained with a mechanical rotator mounted inside the MPMS magnetometer. Note that the MPMS operates with a pair of coils for signal detection mounted parallel (longitudinal coil) and perpendicular

(transversal coil) to the applied magnetic field. The rotator is aligned with the rotation axis normal to the plane defined by both SQUID coil axes.

The measured single crystal has a rectangular shape of (1.87×2.99) mm² in *ab* and 0.54 mm along the crystallographic *c*-direction. We considered demagnetization effects by approximating the sample shape with an ellipsoid⁴¹ and estimated small correction fields of -30 mT for $H \perp c$ and -140 mT for $H \parallel c$, which are negligible for the analysis and discussion of our magnetization results close to the metamagnetic transition.

References

- Ishida, K. *et al.* Spin-triplet superconductivity in Sr₂RuO₄ identified by ¹⁷O Knight shift. *Nature* **396**, 658–660 (1998).
- Grigera, S. A. *et al.* Magnetic field-tuned quantum criticality in the metallic ruthenate Sr₃Ru₂O₇. *Science* **294**, 329–332 (2001).
- Crawford, M. K. *et al.* Structure and magnetism of single crystal Sr₄Ru₃O₁₀: A ferromagnetic triple-layer ruthenate. *Physical Review B* **65**, 214412–1–5 (2002).
- Granata, V. *et al.* Neutron diffraction study of triple-layered Sr₄Ru₃O₁₀. *Journal of Physics: Condensed Matter* **25**, 056004 (2013).
- Granata, V. *et al.* Spin-orbital nature of the high-field magnetic state in the Sr₄Ru₃O₁₀. *Physical Review B* **93**, 115128–1–9 (2016).
- Callaghan, A., Moeller, C. W. & Ward, R. Magnetic interactions in ternary ruthenium oxides. *Inorganic Chemistry* **5**, 1572–1576 (1966).
- Kanbayasi, A. Magnetic properties of SrRuO₃ single crystal. *Journal of the Physical Society of Japan* **41**, 1876–1878, doi:10.1143/JPSJ.41.1876 (1976).
- Fobes, D. *et al.* Anisotropy of magnetoresistance in Sr₄Ru₃O₁₀: Evidence for an orbital-selective metamagnetic transition. *Physical Review B* **81**, 172402–1–4 (2010).
- Mirri, C. *et al.* Anisotropic optical conductivity of Sr₄Ru₃O₁₀. *Physical Review B* **85**, 235124–1–7 (2012).
- Flippen, R. B. Changes in magnetization of single crystal dysprosium, erbium, and holmium in high magnetic fields. *Journal of Applied Physics* **35**, 1047–1048, doi:10.1063/1.1713372 (1964).
- Held, K., Ulmke, M., Blümer, N. & Vollhardt, D. Correlated-electron theory of strongly anisotropic metamagnets. *Physical Review B* **56**, 14469–14480 (1997).
- Eundeok Mun, E. *et al.* Partially disordered antiferromagnetism and multiferroic behavior in a frustrated ising system CoCl₂-2S(NH₂)₂. *Physical Review B* **93**, 104407–1–8 (2016).
- Stoner, E. C. Collective electron ferromagnetism. *Proceedings of the Royal Society of London A* **165**, 372–414 (1938).
- Wohlfarth, E. P. & Rhodes, P. Collective electron metamagnetism. *Philosophical Magazine* **7**, 1817–1824, doi:10.1080/14786436208213848 (1962).
- Levitin, R. Z. & Markosyan, A. S. Itinerant metamagnetism. *Soviet Physics Uspekhi* **31**, 730 (1988).
- Sakakibara, T., Goto, T., Yoshimura, K. & Fukamichi, K. Itinerant electron metamagnetism and spin fluctuations in nearly ferromagnetic metals Y(Co_{1-x}Al_x)₂. *Journal of Physics: Condensed Matter* **2**, 3381 (1990).
- Honerkamp, C. Charge instabilities at the metamagnetic transition of itinerant electron systems. *Physical Review B* **72**, 115103–1–5 (2005).
- Deppe, M. *et al.* Pronounced first-order metamagnetic transition in the paramagnetic heavy-fermion system CeTiGe. *Physical Review B* **85**, 060401(R)–1–5 (2012).
- Binz, B. & Sigrist, M. Metamagnetism of itinerant electrons in multi-layer ruthenates. *Europhysics Letters* **65**, 816–822 (2004).
- Weickert, F., Brando, M., Steglich, F., Gegenwart, P. & Garst, M. Universal signatures of the metamagnetic quantum critical endpoint: Application to CeRu₂Si₂. *Physical Review B* **81**, 134438–1–8 (2010).
- Cao, G. *et al.* Competing ground states in triple-layered Sr₄Ru₃O₁₀: Verging on itinerant ferromagnetism with critical fluctuations. *Physical Review B* **68**, 174409–1–7 (2003).
- Carleschi, E. *et al.* Double metamagnetic transition in Sr₄Ru₃O₁₀. *Physical Review B* **90**, 205120–1–9 (2014).
- Perry, R. S. *et al.* Multiple first-order metamagnetic transition and quantum oscillations in ultra pure Sr₃Ru₂O₇. *Physical Review Letters* **92**, 166602–1–4 (2004).
- Grigera, S. A. *et al.* Disorder-sensitive phase formation linked to metamagnetic quantum criticality. *Science* **306**, 1154–1157 (2004).
- Gegenwart, P., Weickert, F., Garst, M., Perry, R. S. & Maeno, Y. Metamagnetic quantum criticality in Sr₃Ru₂O₇ studied by thermal expansion. *Physical Review Letters* **96**, 136402–1–4 (2006).
- Mao, Z. Q., Zhou, M., Hooper, J., Golub, V. & O'Connor, C. J. Phase separation in the itinerant metamagnetic transition of Sr₄Ru₃O₁₀. *Physical Review Letters* **96**, 077205–1–4 (2006).
- Fobes, D. *et al.* Phase diagram of the electronic states of trilayered ruthenate Sr₄Ru₃O₁₀. *Physical Review B* **75**, 094429–1–5 (2007).
- Cao, G., Chikara, S., Brill, J. W. & Schlottmann, P. Anomalous itinerant magnetism in single crystal Sr₄Ru₃O₁₀: A thermodynamic and transport investigation. *Physical Review B* **75**, 024429–1–6 (2007).
- Xu, Z. *et al.* Magnetic, electrical transport, and thermoelectric properties of Sr₄Ru₃O₁₀: Evidence for a field-induced electronic phase transition at low temperatures. *Physical Review B* **76**, 094405–1–6 (2007).
- Gupta, R., Kim, M., Barath, H., Cooper, S. L. & Cao, G. Field- and pressure-induced phases in Sr₄Ru₃O₁₀: A spectroscopic investigation. *Physical Review Letters* **96**, 067004–1–4 (2006).
- Schottenhamel, W. *et al.* Dilatometric study of the metamagnetic and ferromagnetic phases in the triple-layered Sr₄Ru₃O₁₀ system. *Phys. Rev. B* **94**, 155154, doi:10.1103/PhysRevB.94.155154 (2016).
- Zhou, M. *et al.* Electronic and magnetic properties of triple-layered ruthenate Sr₄Ru₃O₁₀ single crystals grown by a floating-zone method. *Materials Research Bulletin* **40**, 942–950 (2005).
- Jo, Y. J. *et al.* Orbital-dependent metamagnetic response in Sr₄Ru₃O₁₀. *Physical Review B* **75**, 094413–1–5 (2007).
- Horner, H. & Varma, C. M. Nature of spin-reorientation transitions. *Physical Review Letters* **20**, 845–846 (1968).
- Cullity, B. D. & Graham, C. D. *Introduction to Magnetic Materials*, 2 edn. (Wiley-IEEE Press, 2008).
- Dzyaloshinsky, I. A thermodynamic theory of weak ferromagnetism of antiferromagnetics. *Journal of Physics and Chemistry of Solids* **4**, 241–255, doi:10.1016/0022-3697(58)90076-3 (1958).
- Moriya, T. Anisotropic superexchange interaction and weak ferromagnetism. *Phys. Rev.* **120**, 91–98, doi:10.1103/PhysRev.120.91 (1960).
- Bellaiche, L., Gui, Z. & Kornev, I. A. A simple law governing coupled magnetic orders in perovskites. *J. Phys.: Condensed Matter* **24**, 312201–1–6 (2012).
- Moriya, T. Anisotropic superexchange interaction and weak ferromagnetism. *Journal of Physics: Condensed Matter* **11**, 6529–41 (1999).
- Fittipaldi, R., Sisti, D., Vecchione, A. & Pace, S. Crystal growth of a lamellar Sr₃Ru₂O₇-Sr₄Ru₃O₁₀ eutectic system. *Crystal Growth & Design* **7**, 2495–2499, doi:10.1021/cg070180p (2007).
- Osborn, J. A. Demagnetizing factors of the general ellipsoid. *Physical Review* **67**, 351–357 (1945).

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Author Contributions

F.W., A.M.S. and E.C. initiated the research. F.W. led the project. L.C. and B.M. conducted the experiments. F.W., M.B.S. and M.J. analyzed experimental data and performed numerical calculations. R.F., V.G. and A.V. grew and characterized single crystals. F.W., M.B.S. and M.J. wrote the manuscript. All authors discussed the results and commented on the manuscript.

Additional Information

Competing Interests: The authors declare that they have no competing interests.

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