Persistent spin dynamics in the pressurized spin-liquid candidate YbMgGaO₄

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Single-crystal x-ray diffraction, density functional band structure calculations, and muon spin relaxation (μSR) are used to probe the pressure evolution of the triangular spin-liquid candidate YbMgGaO₄. The rhombohedral crystal structure is retained up to at least 10 GPa and shows a nearly uniform compression along both the in-plane and out-of-plane directions, whereas local distortions caused by the random distribution of Mg²⁺ and Ga³⁺ remain mostly unchanged. The μ SR data confirm persistent spin dynamics up to 2.6 GPa and down to 250 mK with no change in the muon relaxation rate. Longitudinal-field μ SR reveals power-law behavior of the spin-spin autocorrelation function, both at ambient pressure and upon compression.

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I. INTRODUCTION

Spin-liquid states in frustrated magnets are nowadays actively studied as hosts for unconventional excitations representing magnetic monopoles [1,2] and other exotic quasiparticles [3,4]. One relatively unexplored aspect in this field is the evolution of spin-liquid materials under pressure and the change in spin dynamics caused by tuning magnetic interactions via lattice compression.

Here, we focus on the spin-liquid candidate YbMgGaO₄ [5–7] that recently evolved as a unique triangular antiferromagnet with the robust threefold symmetry, persistent spin dynamics, and a broad continuum of (potentially fractionalized) magnetic excitations. The crystal structure of this compound features triangular layers of the pseudospin- $\frac{1}{2}$ Yb³⁺ ions that are well separated by slabs of nonmagnetic Mg²⁺ and Ga³⁺ [Fig. 1(a)]. Thermodynamic measurements [5,6] and muon spin relaxation (μ SR) [8] suggest the absence of magnetic order down to at least 50 mK. Weak spin freezing around 100 mK was indicated by ac-susceptibility data [9], although it involves only a tiny amount of the magnetic entropy [5] and leaves no signatures in either dc susceptibility [10] or μ SR [8].

Magnetic excitations of YbMgGaO₄ form a broad continuum that can be interpreted in terms of gapless spinons [11–14] or as arising from short-range valence bonds [10,15], the latter suggestion being particularly interesting, as it makes a direct link to Anderson's pioneering work at the outset of the spin-liquid research [16]. Continuum features were also

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reported for other Yb³⁺ triangular antiferromagnets [17–19], but they show a different distribution of the spectral weight and probably have a separate origin. Indeed, YbMgGaO₄ is known to be strongly affected by structural randomness that arises from the random distribution of Mg²⁺ and Ga³⁺ between the Yb³⁺ layers and modulates magnetic interactions via random crystal electric fields acting on Yb³⁺ [20,21]. No such randomness occurs in other Yb-based triangular antiferromagnets.

In the following, we explore the effect of hydrostatic pressure on the structure and magnetism of YbMgGaO₄, and juxtapose the behavior of this material with the evolution of other spin-liquid candidates upon compression. We show that in YbMgGaO₄, spins remain dynamic up to at least 2.6 GPa, and quantify associated structural changes for an eventual comparison across different classes of spin-liquid materials.

II. CRYSTAL STRUCTURE

A. Average structure

Room-temperature x-ray diffraction (XRD) data were collected at the ID15B beam line of the European Synchrotron Radiation Facility (ESRF) between ambient pressure and 10 GPa. A diamond-anvil cell loaded with He gas and a small single crystal of YbMgGaO₄ from the batch reported in Ref. [6] were used for the experiment.

Three positional parameters (z coordinates) and three independent atomic displacement parameters were refined for Mg/Ga, O1, and O2. The position of Yb was fixed at (0,0,0), with its thermal ellipsoid refined anisotropically to account for local displacements caused by the random distribution of Mg and Ga. Altogether, nine structural parameters were refined from about 120 symmetry-independent reflections collected at each pressure point (see Supplemental Material) [22]. Details

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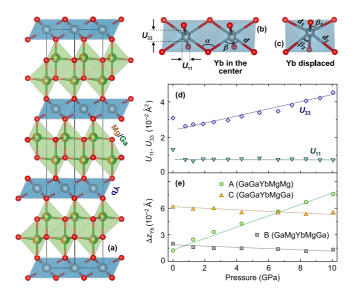


FIG. 1. (a) Crystal structure of YbMgGaO₄ with Mg²⁺ and Ga³⁺ randomly occupying the position in the trigonal bipyramids. (b) Local structural parameters, the Yb-O distance (*d*) and Yb-O-Yb (α) or O-Yb-O (β) angles with $\alpha = \beta$. (c) Distortion of the YbO₆ octahedron caused by an asymmetric distribution of the Mg and Ga atoms. (d) Experimental atomic displacement parameters of Yb. (e) Displacements of the Yb atoms obtained for different local configurations shown in Fig. 3.

of the data collection and structure refinement are given in Appendix A.

The $R\bar{3}m$ symmetry of YbMgGaO₄ keeps all Yb-O distances equal, but allows a trigonal distortion of the YbO₆ octahedra with the O-Yb-O angle β deviating from 90°. Incidentally, this angle is equal by symmetry to the Yb-O-Yb bridging angle α , which is responsible for superexchange interaction [Fig. 1(b)]. Two structural parameters, one distance and one angle, are thus sufficient to characterize both the local geometry of Yb³⁺ and the geometry of the nearest-neighbor exchange pathway within the average structure.

No changes in the crystal symmetry were observed in our experiment. The pressure evolution of the a and c lattice parameters demonstrates similar compressibility along the different crystallographic directions [Fig. 2(a)]. The fit with the second-order Birch-Murnaghan equation of state,

$$p(V) = \frac{3B_0}{2} \left[\left(\frac{V_0}{V} \right)^{\frac{7}{3}} - \left(\frac{V_0}{V} \right)^{\frac{5}{3}} \right],\tag{1}$$

using the EosFit routine [23], returned the bulk modulus of $B_0 = 142(2)$ GPa and the unit-cell volume of $V_0 = 253.12(7)$ Å³. A similar fit for individual lattice parameters using a^3 or c^3 in place of V yields $B_{0,a} = 151(3)$ GPa and $B_{0,c} = 130(2)$ GPa, suggesting a nearly isotropic compression of the structure. YbMgGaO₄ is more compressible than Yb₂Ti₂O₇ ($B_0 = 219$ GPa [24]) and other rare-earth pyrochlores that typically feature the bulk moduli in excess of 200 GPa [25].

The Yb-O distances shrink by about 0.6% at 2.6 GPa (the highest pressure of our μ SR experiment; see Sec. III) and by 1.7% at 10 GPa [Fig. 2(b)]. The $\alpha = \beta$ angle shows a weak

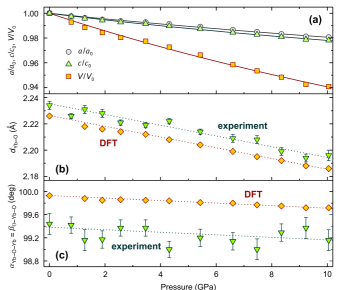


FIG. 2. Pressure-induced changes of the average structure determined using single-crystal XRD: (a) lattice parameters and unit-cell volume; (b) Yb-O distance d; (c) Yb-O-Yb/O-Yb-O angles ($\alpha = \beta$). The lines in (a) are fits with the second-order Murnaghan equation of state. (b) and (c) additionally show the geometrical parameters obtained from DFT structure relaxations detailed in Appendix B, and dotted lines are a guide for the eye.

downward trend only, with a larger error bar caused by the lower accuracy in the determination of oxygen position due to the low scattering power of oxygen. In order to confirm this downward trend, we relaxed the experimental structures using density functional theory (DFT) band structure calculations. As DFT cannot treat the Mg/Ga disorder explicitly, ordered structural models have to be used, as explained in Appendix B. This approximation leads to a constant offset between the DFT results and experiment. Nevertheless, not only the qualitative trends but also the slope are well reproduced (Fig. 2). We thus confirm that the Yb-O-Yb angle decreases under pressure. Compared to the ambient-pressure value, it changes by 0.07° at 2.6 GPa and by 0.2° at 10 GPa.

B. Local structure

Moderate changes of the average crystal structure are accompanied by a strong elongation of the Yb thermal ellipsoid. The in-plane displacements characterized by U_{11} are not affected by pressure, whereas the out-of-plane displacement component increases by 70% [Fig. 1(d)]. This out-of-plane displacement has been previously linked to the local distortions of the YbO₆ octahedra caused by the random (and, generally, asymmetric) distribution of the differently charged Mg^{2+} and Ga^{3+} ions around Yb³⁺ [20]. At first glance, the increase in U_{33} implies a strong enhancement of the structural randomness under pressure, but the actual situation is more complex.

We analyze the pressure evolution of the local structure by DFT relaxations for several ordered models of YbMgGaO₄. First, we consider the models of Ref. [20] that were previously used to interpret crystal-field excitations of Yb³⁺, and evaluate

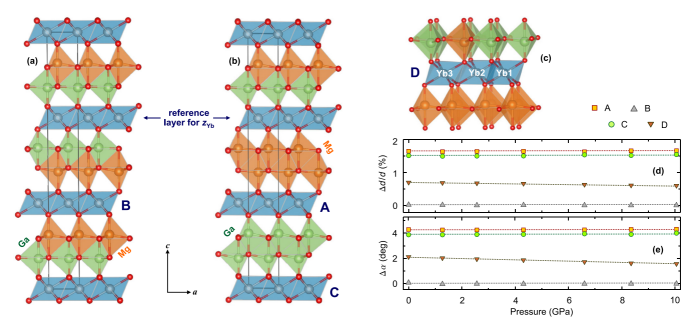


FIG. 3. Ordered models for evaluating pressure evolution of the local structure. (a),(b) Layer-by-layer ordering of Mg²⁺ and Ga³⁺ leading to the Yb layers A, B, and C with the same YbO₆ octahedra throughout each layer. (c) A more complex model with layer D containing nonequivalent YbO₆ octahedra. (d) Relative deformations of the octahedra obtained as $(d_1 - d_2)/(d_1 + d_2)$ for layers A, B, and C and as a difference between the average d for the Yb1 and Yb2 octahedra in layer D. (e) Angular deformation obtained as $\Delta \alpha = \Delta \beta = \beta_2 - \beta_1$ for layers A, B, and C and as the difference in the Yb1-Yb2 and Yb2-Yb3 superexchange angles for layer D.

 Δz_{Yb} as the displacements of the Yb atoms relative to each other. These displacements indicate the spread of the Yb electron density, which is gauged by the U_{33} parameter of the average structure.

Three scenarios visualized in Figs. 3(a) and 3(b) are considered, with the Yb³⁺ ions sandwiched between the following: (i) two Ga³⁺ ions on one side and two Mg²⁺ ions on the other side (octahedra A), (ii) one Ga³⁺ and one Mg²⁺ ion on each side (octahedra B), and (iii) two Ga³⁺ ions on one side and a combination of Ga³⁺ and Mg²⁺ on the other side (octahedra C). Pressure has a strong effect on the Yb position in the octahedra A and nearly no effect on the Yb atoms in B and C [Fig. 1(e)]. Surprisingly, this effect is a shift of the whole octahedron A along the c direction without any change in the octahedron itself. Using the geometrical parameters shown in Fig. 1(c), we determine that for octahedra A, the deformation expressed by $(d_2 - d_1)$ changes from 1.65% at 0 GPa to 1.66% at 10 GPa, whereas $(\beta_2 - \beta_1)$ evolves from 4.29° to 4.32° [Figs. 3(d) and 3(e)]. The changes in octahedra B and C are equally small. Since $\alpha = \beta$ in our material, the distribution of the Yb-O-Yb angles also does not change with pressure. Therefore, neither local randomness at each Yb site nor randomness of the exchange couplings should be affected by pressure.

We also consider a different scenario where two adjacent Yb atoms within the same layer have a different local environment. In layer D [Fig. 3(c)], Yb1 is surrounded by 3 Ga atoms on one side and 3 Mg atoms on the other side. In contrast, Yb2 and Yb3 are surrounded by 2 Ga and 4 Mg atoms each. These dissimilar local environments lead to different α_1 and α_2 for the Yb1-Yb2 and Yb2-Yb3 superexchange pathways and reflect the effect of structural randomness on nearestneighbor magnetic interactions in YbMgGaO₄. However, the

difference between these two pathways appears to be only weakly pressure-dependent and decreases upon compression [Fig. 3(e)].

Our results suggest that the visible enhancement of U_{33} under pressure is *not* related to the increased distortions around the Yb³⁺ ions. It rather indicates a change in the overall position of octahedra A that may be explained by the accumulation of different charges above (Ga³⁺) and below (Mg²⁺) the Yb layer. But the key result at this juncture is that such a change reflects a rearrangement within the Mg/Ga slabs and does not affect the structure of the magnetic layer itself. The YbO₆ octahedra undergo a uniform compression and simply keep the deformation that they had at ambient pressure. The randomness effect on the exchange couplings should be largely unchanged or even decrease under pressure.

III. SPIN DYNAMICS

The pressure evolution of spin dynamics is probed by muon spin relaxation (μ SR). The experiments were performed at the GPD and Dolly spectrometers at the Paul Scherrer Institute (PSI) at ambient pressure and at the GPD spectrometer under hydrostatic pressure [26] down to 250 mK on a polycrystalline sample of YbMgGaO₄. The data were collected in zero field (ZF) and in the longitudinal-field (LF) mode, where the magnetic field was applied parallel to the spin of the implanted muons.

A. ZF μSR

ZF μ SR time spectra measured at ambient pressure and at 2.6 GPa, the highest pressure of our experiment, are compared

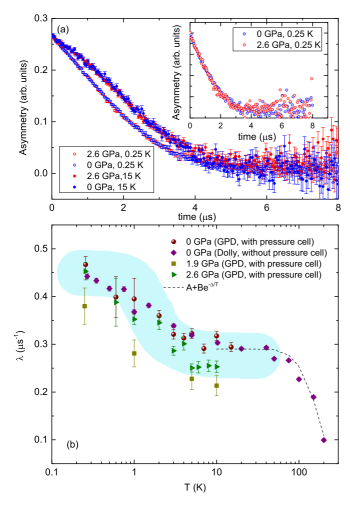


FIG. 4. (a) Zero-field μ SR spectra measured at 0 and 2.6 GPa at both 15 K and 250 mK. The inset shows the sample contribution after subtracting $f_{PC}A_{PC}$ of Eq. (2). (b) Temperature dependence of the zero-field muon relaxation rate measured at different pressures. The dashed line is the fit with the activated behavior above 40 K.

in Fig. 4. The similar behavior of the μ SR time spectra indicates no change of the magnetic ground state, and the absence of oscillations excludes pressure-induced magnetic ordering in YbMgGaO₄ up to at least 2.6 GPa. To estimate the temperature dependence of the relaxation rate λ , we fitted the ZF- μ SR time spectra by

$$A(t) = f_{PC}A_{PC} + (1 - f_{PC})e^{-\lambda t},$$
 (2)

where $A_{\rm PC}$ is the pressure-cell contribution described in Appendix C. The fraction of signal coming from the pressure cell was $f_{\rm PC}=0.7$ at 0 and 2.6 GPa, and $f_{\rm PC}=0.5$ at 1.9 GPa, where a different cell was used in an attempt to reduce the background.

The zero-field muon relaxation rate λ is temperature-independent between 4 and 40 K (Fig. 4). Its increase below 4 K indicates the onset of spin-spin correlations that fully develop around 0.8 K, where λ flattens out and remains temperature-independent upon further cooling. This temperature evolution is essentially similar to the ambient-pressure μ SR data reported in Ref. [8] and remains unchanged at

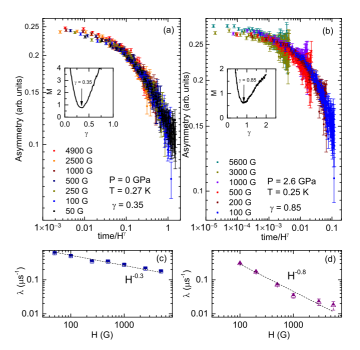


FIG. 5. (a),(b) Universal scaling of the LF- μ SR data measured at 0 GPa and 270 mK (left, $\gamma=0.35$) and 2.6 GPa and 250 mK (right, $\gamma=0.85$). The insets show the data mismatch function as a function of γ . (c),(d) Power-law behavior of the muon relaxation rate, $\lambda(H)\sim H^{-\gamma}$.

2.6 GPa (Fig. 4). Even absolute values of λ are the same as at ambient pressure within the error bar.

Above 40 K, λ shows a steady decreasing trend described by an activated behavior $\lambda = A + B e^{-\Delta/T}$, with $\Delta = 320 \pm 20$ K, which is reminiscent of the lowest crystal-field excitation energy of about 450 K [20]. This observation suggests that at high temperatures, the relaxation is governed by an Orbach process [27] involving the excited crystal-field doublets of Yb³⁺.

B. LF μ SR

LF measurements complement the data obtained in the zero field. Even a longitudinal field, which is more than 10 times higher than the local static field estimated from the low-temperature value of λ , does not decouple the muon relaxation, suggesting that the spins are dynamic in nature as has been seen at ambient pressure [8].

The LF data measured in different fields follow universal scaling initially proposed by Keren *et al.* [28] for systems with glassy dynamics. This scaling manifests itself in the power-law behavior of the muon relaxation rate $\lambda(H)$ extracted by fitting individual LF curves with a stretched exponential $e^{-[\lambda(H)t]^{\beta}}$ supplied with the time-independent and field-independent background, which originates from the sample holder and pressure cell. The ensuing values of λ follow $\lambda(H) \sim H^{-\gamma}$ with $\gamma \simeq 0.3$ at 0 GPa and 0.8 at 2.6 GPa (Fig. 5).

This behavior is confirmed by the scaling of muon asymmetry plotted against t/H^{γ} . To this end, the data points from all fields up to $t=6 \,\mu s$ [29] are arranged with increasing t/H^{γ} for every value of γ . An empirical data mismatch function is calculated by taking the difference between the

neighboring points and weighing them by the corresponding error bars. This mismatch function is defined as

$$M = \frac{1}{N} \sum_{i}^{N} \frac{(A_i - A_{i+1})^2}{(\delta_i - \delta_{i+1})^2},$$
 (3)

where N is the number of data points, and A_i and δ_i correspond to the asymmetry and error bar of the ith data point, respectively. The lowest value of the mismatch function is obtained at $\gamma = 0.35$ (0 GPa) and $\gamma = 0.85$ (2.6 GPa) that produce the universal scaling over at least three orders of magnitude in t/H^{γ} and show excellent agreement with the γ values from the analysis of $\lambda(H)$.

The scaling may not hold in the whole time and field range—e.g., at very short times comparable to the width of the muon pulse [28] or in very high fields that affect spin dynamics—but the scaling over three orders of magnitude (Fig. 5) is congruent with all earlier observations [30,31] and serves as a robust evidence of the power-law behavior $q(t) \sim t^{\gamma-1}$ of the dynamic spin-spin autocorrelation function $q(t) = \langle S_i(t) \cdot S_i(0) \rangle$. It indicates collective dynamics, as, for example, in spin-glass systems where γ increases toward 1.0 upon approaching the freezing temperature from above [28].

IV. DISCUSSION AND SUMMARY

YbMgGaO₄ is remarkably insensitive to pressure and thus different from other spin-liquid candidates. For example, herbertsmithite [32,33] becomes magnetically ordered at 2.5 GPa [34], whereas stoichiometric Yb₂Ti₂O₇ [35] shows pressureinduced magnetic order already at 0.1 GPa [36]. These materials have different compressibilities, with the bulk modulus of $B_0 = 70$ GPa in herbertsmithite [34] and $B_0 = 219$ GPa in Yb₂Ti₂O₇ [24]. YbMgGaO₄ shows an intermediate value of 142 GPa and is certainly comparable to other spin-liquid candidates as far as elastic properties are concerned. Its structural parameters are affected by pressure. The persistence of spin dynamics will then indicate that structural changes influence exchange couplings only weakly, whereas the material lies sufficiently far away from the phase boundary between the spin-liquid and magnetically ordered states. Alternatively, spin dynamics may not be caused by frustration that controls this phase boundary, and originate from structural randomness, which is not affected by pressure.

Triangular antiferromagnets develop 120° order or stripe order, depending on the extent of exchange anisotropy [37] and on the ratio J_2/J_1 of the second-neighbor to nearestneighbor exchange coupling [38,39]. The formation of a spin liquid is mostly controlled by J_2/J_1 , with the complete suppression of magnetic order between $J_2/J_1 \simeq 0.07$ and 0.15 [7,38,39]. Experimental estimates of this parameter vary in a broad range between 0.0 and 0.26 [7] and would place YbMgGaO₄ in any of the regions on the phase diagram: 120° order [6], stripe order [21,40], or the boundary between the latter and the spin-liquid phase [41].

External pressure of 2.6 GPa leads to a shortening of the Yb-O distance d by 0.013 Å and a reduction in the Yb-O-Yb angle α by 0.07° (Fig. 2). Using the superexchange theory of Ref. [42], we estimate that the change in the angle increases J_1 by 6 %, whereas the shortening of the Yb-O distance

increases J_1 by another 5 % assuming exponential dependence of Slater-Koster parameters on the interatomic distance. Overall, we expect that J_1 increases by 11%, which is similar in magnitude to pressure-induced changes in Cu-based quantum magnets [43,44]. No significant changes are expected for J_2 because long-range couplings are less sensitive to the structural geometry. In YbMgGaO₄, the Yb-O-O angles decrease by as little as 0.11° upon compression to 2.6 GPa. In contrast, in a Cu-based quantum magnet, even the 2° change of the respective angle reduces the coupling by only 20% [45].

We conclude that J_1 increases, J_2 is roughly unchanged, and the J_2/J_1 ratio of YbMgGaO₄ should be reduced under pressure. Assuming the parametrization of Ref. [41] with $J_2/J_1=0.18$, it would imply that pressure pushes the material into the spin-liquid region, in agreement with the persistence of spin dynamics observed experimentally. Alternatively, and perhaps even more likely, the lack of significant changes under pressure may indicate that the spin dynamics is triggered by structural randomness and thus unaffected by pressure.

The only change we observe is the increase in the γ parameter that describes scaling behavior in longitudinal fields. For a system with glassy dynamics, the increase in γ would reflect an evolution toward a frozen state [28] that was indeed proposed for YbMgGaO₄ below 100 mK (at ambient pressure) based on the ac-susceptibility data [9]. However, neither dc susceptibility [10] nor μ SR [8] support bulk spin freezing at ambient pressure. The scaling behavior may, in fact, be unrelated to glassy dynamics, and indeed it was also reported for a variety of systems with critical fluctuations influenced by structural disorder [30,31]. On the other hand, the very presence of this scaling serves as an additional argument for collective spin dynamics caused by the structural randomness, and the importance of this randomness for the spin-liquid behavior of YbMgGaO₄.

Altogether, hydrostatic pressure leads to a uniform compression of the YbMgGaO₄ structure with the reduction in the Yb-O distances and Yb-O-Yb angles, whereas local distortions of the YbO₆ octahedra and consequent randomness effects are nearly unchanged. Spin dynamics is not affected by pressure and appears to be collective yet influenced by the structural randomness. This puts YbMgGaO₄ into the group of materials where randomness effects can be integral to the spin-liquid formation, but collective spin dynamics is nevertheless observed.

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TABLE I. Details of the single-crystal XRD data collection and structure refinement at 10 GPa.

298
3.34272(13)
24.6033(7)
0.41114
$-4 \leqslant h \leqslant 3$
$-5 \leqslant k \leqslant 5$
$-39 \leqslant l \leqslant 38$
126
9
0.0249
0.0310

APPENDIX A: X-RAY DIFFRACTION

Membrane-driven LeToullec-type diamond-anvil cells (DACs) were used, equipped with Boehler-Almax anvils. Stainless steel was used as the gasket material and helium was loaded as the pressure-transmitting medium. Diffraction patterns were collected with a Mar555 flat-panel detector using steps of 0.5° oscillations over a total ω scan range of 76° about the vertical axis. The pressures were measured using the ruby fluorescence method. Lattice parameter determination and integration of the reflection intensities were performed using the CRYSALISPRO software [46]. Structures were refined using SHELXL [47] within the SHELXLE [48] graphical interface.

Details of the data collection and structure refinement at 10 GPa are listed in Table I. The refinements at other pressures were similar and can be found in the cif file provided as Supplemental Material [22]. We note that the ambient-pressure experiment was performed before loading the cell with the He gas. This may be the reason for the abrupt change in the Yb displacement parameters U_{11} and U_{33} between 0 and 0.78 GPa [Fig. 1(d)].

APPENDIX B: COMPUTATIONAL RESULTS

1. General methodology

Nonmagnetic DFT calculations were performed in the FPLO [49] and VASP [50] codes. Atomic positions were optimized until the energy minimum was reached, and residual forces dropped below 0.005 eV/Å. The typical k mesh included 64 points within the first Brillouin zone. Experimental

lattice parameters were used at each pressure, and only atomic positions were relaxed.

The results of structure optimization may be affected by the computational methodology. In Table II, we analyze the role of three factors: (i) band structure code, (ii) exchange-correlation potential, and (iii) treatment of the Yb 4f shell. The calculations are performed for the simplest R3m ordered model of YbMgGaO₄ described in Appendix B 2.

The choice of the band structure code affects the results of the relaxation. VASP calculations predict the too-long Yb-O distances and, consequently, the too-low angles. On the other hand, the relaxations within FPLO produce, irrespective of the methodology, the average distance $(d_1+d_2)/2$ and the average angle $(\beta_1+\beta_2)/2$ within, respectively, 0.01 Å and 0.5° from the experimental values. The larger deviations of the VASP results are probably related to the lower accuracy of the default pseudopotential for the Yb atoms, whereas FPLO does not rely on pseudopotentials and introduces no approximations to the crystal potential.

All choices mentioned here, including the VASP calculations, produce qualitatively similar results as a function of pressure. The trends obtained from DFT are thus robust. The results shown in the manuscript are obtained with FPLO, GGA functional, and Yb 4f states placed into the core, as this choice improved the convergence for larger unit cells.

2. Ordered structural models

As DFT cannot treat mixed sites, ordered structural models are used for calculations. We adopted three types of such ordered models:

- (1) The R3m model obtained from the parent $R\bar{3}m$ structure by splitting the mixed Mg/Ga site into two: one fully occupied by Ga and the other one fully occupied by Mg. In this case, each Yb layer is sandwiched between the Mg and Ga layers, leading to the off-center Yb displacement, two nonequivalent Yb-O distances $d_1 \neq d_2$, and two O-Yb-O angles $\beta_1 \neq \beta_2$. The R3m structure was used in Table II for testing the effect of band structure code, basis set, and exchange-correlation potential.
- (2) *P3m* models that preserve threefold symmetry and feature layer-by-layer ordering of Mg and Ga. We chose two of such ordered structures (Fig. 3) that correspond to the third and fourth structures from Fig. S5 of Ref. [20].

In the structure from Fig. 3(a), all Yb atoms develop a nearly symmetric local environment. This structure was used

TABLE II. Comparison of the relaxed geometrical parameters for the ambient-pressure crystal structure in the primitive cell with the R3m symmetry. Different band structure codes (FPLO vs VASP), exchange-correlation potentials (local density approximation (LDA) [51] vs generalized gradient approximation (GGA) [52]), and approaches to the treatment of the Yb 4f shell (valence vs core) have been used.

Code	FPLO	FPLO	FPLO	FPLO	VASP	Experiment
V_{xc} Yb 4 f	LDA valence	LDA core	GGA valence	GGA core	GGA core	
$\frac{d_1/d_2 (\text{Å})}{\beta_1/\beta_2 (\text{deg})}$	2.229/2.248 98.62/99.73	2.214/2.233 99.52/100.68	2.231/2.257 98.09/99.60	2.215/2.243 98.88/100.62	2.258/2.270 97.31/98.02	2.234(4) 99.44(13)

to evaluate the pressure dependence of the Yb-O distances and Yb-O-Yb angles shown in Fig. 2. In this case, the octahedra are still deformed, but $d_2 - d_1 < 0.005$ Å and $\beta_2 - \beta_1 < 0.2^{\circ}$ indicate a negligibly small deformation. The "symmetric" structure (left panel of Fig. 3) is also used to obtain the displacements (Δz_{Yb}) of Yb within octahedra B (GaMgYb-MgGa).

The structure from Fig. 3(b) covers the opposite scenario of the Yb atoms with a highly asymmetric local environment. It contains octahedra A (GaGaYbMgMg) and C (GaGaYb-MgGa) showing the largest values of $d_2 - d_1$ and $\beta_2 - \beta_1$. As explained in the main text, the thermal displacement parameter of the average structure, U_{33} , does not reflect deformations of individual octahedra and instead shows relative displacements of the Yb atoms with respect to each other. In the ideal structure, the Yb atoms should be at $z = 0, \frac{1}{3}, \frac{2}{3}$. To assess the effect of pressure on U_{33} , we compare the positions of the Yb atoms within the octahedra A and C to the remaining Yb atom at $z \simeq \frac{2}{3}$ that shows a rather symmetric local environment and can be used as reference. The values of Δz_{Yb} are obtained as $|z_{A,C}-z_{\text{ref}}-\frac{1}{3}|$, where z_{ref} is the z coordinate of the reference

(3) The P3m model with the fourfold $2\mathbf{a} \times 2\mathbf{b} \times \mathbf{c}$ supercell, where a, b, and c are lattice vectors of the parent $R\bar{3}m$ structure. This supercell allows one to construct configurations where adjacent Yb atoms have a different local environment. We focus on the fragment shown in Fig. 3(c), where Yb1-Yb2 and Yb2-Yb3 represent two nonequivalent superexchange pathways caused by the uneven distribution of Mg and Ga around the Yb layer.

APPENDIX C: μSR EXPERIMENT

μSR measurements were performed in double-walled MP35 pressure cells with Daphne oil 7373 as the pressuretransmitting medium. The pressure value was determined by measuring the superconducting transition of a small piece of

0.30 T = 270 mK= 0 GPa 0.20 Asymmetry

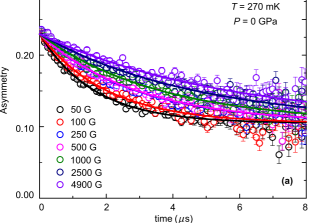


TABLE III. Details of the μ SR experiments.

P (GPa)	Spectrometer	Pressure cell	T (K)	Mode
0	Dolly	none	0.26-200	ZF/LF
0	GPD	MP35N + MP35N	0.25-10	ZF
1.9	GPD	MP35N + CuBe	0.25-10	ZF
2.6	GPD	MP35N + MP35N	0.25-10	ZF/LF

indium positioned next to the sample inside the pressure cell. Further experimental details are summarized in Table III.

Two different pressure cells were used. The 0 and 2.6 GPa data were collected in the double-walled MP35N+MP35Ntype cell [26], whereas the 1.9 GPa data were collected low-background double-walled separately in the MP35N+CuBe-type cell [53]. For both cells, we employed Eq. (2) and expressed the pressure-cell contribution as the Kubo-Toyabe depolarization function multiplied by an exponential damping,

$$A_{PC}(t) = \left[\frac{1}{3} + \frac{2}{3}\left(1 - \sigma_{PC}^2 t^2\right)e^{-\sigma_{PC}^2 t^2/2}\right] \exp(-\lambda_{PC} t).$$

In the case of the MP35N+MP35N-type cell, σ_{PC} = $0.29~\mu s^{-1}$ was temperature independent, whereas λ_{PC} remained constant down to 1 K and increased at lower temperatures, similar to Ref. [26]. The fraction of the signal coming from the pressure cell was $f_{PC} = 0.7$. In the case of the MP35N+CuBe-type cell, both $\sigma_{PC}=0.32~\mu s^{-1}$ and $\lambda_{PC} = 0.025 \ \mu s^{-1}$ are temperature independent, in agreement with those of Ref. [53], and $f_{PC} = 0.5$. The usage of different cells may be the reason for a slight offset between the 1.9 GPa data and 2.6 GPa data in Fig. 4.

In longitudinal fields above 100 G, the contribution of the pressure cell should be mostly decoupled, and the data include only a constant, field-independent background. Figure 6 shows the LF data fitted by the stretched exponentials to obtain $\lambda(H)$ shown in Fig. 5.

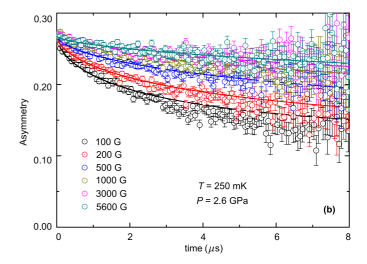


FIG. 6. Decoupling experiments (asymmetry as a function of time in different applied longitudinal fields) at (a) ambient pressure and 270 mK, and at (b) 2.6 GPa and 250 mK.

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