Quantum spin liquid (QSL) is a novel state of matter which refuses the conventional spin freezing even at 0 K. Experimentally searching for the structurally perfect candidates is a big challenge in condensed matter physics. Here we report the successful synthesis of a new spin-1/2 triangular antiferromagnet YbMgGaO$_4$, with $R3m$ symmetry. The compound with an ideal two-dimensional and spatial isotropic magnetic triangular-lattice has no site-mixing magnetic defects and no antisymmetric Dzyaloshinsky-Moriya (DM) interactions. No spin freezing down to 60 mK (despite $\theta_w \sim -4$ K), the power-law temperature dependence of heat capacity and nonzero susceptibility at low temperatures suggest that YbMgGaO$_4$ is a promising gapless ($\leq |\theta_w|/100$) QSL candidate. The residual spin entropy, which is accurately determined with a non-magnetic reference LuMgGaO$_4$, approaches zero ($< 0.6\%$). This indicates that the possible QSL ground state (GS) of the frustrated spin system has been experimentally achieved at the lowest measurement temperatures.

Low-spin geometrically frustrated systems in two-dimensional (2D) lattices have received significant interest in condensed-matter physics. The two most studied frustrated spin systems are spin-1/2 triangular and kagomé antiferromagnets, in which strong quantum fluctuations prevent spin freezing even at very low temperatures$^1$. With respect to theoretical studies, Anderson first proposed that the triangular Heisenberg antiferromagnet (THAF) has a resonating valence bond GS, which is a type of spin liquid$^{2,3}$. However, recent numerical studies have consistently indicated a long-range Néel GS for spin-1/2 THAF. The calculated order parameter is much smaller than the classical value, indicating that it is very close to a quantum critical point between magnetic ordered and disordered GSs$^{4,5}$. On the experimental side, “structurally perfect” triangular or kagomé antiferromagnets (AFs) are still extremely rare, although many spin-1/2 geometrically frustrated triangular and kagomé AFs have been proposed$^{6-12}$. Most of the existing candidates suffer from spatially anisotropic intralayer exchange interactions$^{8,9,12}$, site-mixing between magnetic and nonmagnetic ions$^{1,12-14}$, interlayer exchange interactions$^{8-10}$, and/or antisymmetric Dzyaloshinsky-Moriya (DM) interactions$^{15}$. These factors are critical to the GSs of the frustrated spin

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systems and are difficult to determine precisely. The complications caused by these factors make extraction of the intrinsic physics from real systems difficult.

In this paper, we report the successful synthesis of a new triangular antiferromagnet with effective spin-1/2, YbMgGaO₄. The aforementioned structural disadvantages are avoided in the new compound. First, it has spatially isotropic and perfect triangular layers with R₃₃m symmetry. Second, the number of magnetic defects is negligible because of the large chemical difference between the Kramers magnetic Yb³⁺ ions and the nonmagnetic ions. Third, the magnetic triangular layers are well magnetically separated by nonmagnetic double layers of Mg/GaO₅ triangular bipyramids, indicating an ideal two-dimensionality of the magnetic layers and a negligible interlayer exchange interaction. Finally, the antisymmetric DM interactions between first-, second- and third-neighbor spins are strictly excluded because inversion centers are located at any Yb³⁺ ion and at the half-way sites between them. Moreover, the nonmagnetic reference compound LuMgGaO₄ is also available for control experiments, such as precisely excluding the lattice heat capacities for YbMgGaO₄.

No magnetic ordering is observed at least down to 60 mK from both magnetization and heat capacity measurements, despite the obvious AF exchange interaction between nearest-neighbor spins (θₓ ≈ −4 K) suggesting that YbMgGaO₄ is a new QSL candidate. And the power-law temperature dependence of heat capacity and nonzero susceptibility further indicate that the excitation gap from the GS should be no more than ~ |θₓ|/100 at low temperatures. Almost zero residual spin entropies are observed under 0 to 9 T at T < 0.3 K. This experimentally indicates that the frustrated spin system extremely approaches a possible gapless QSL GS at low temperatures. Moreover, an anomalous susceptibility plateau is first observed at paramagnetic states (0.5 K), which imply an unusual field-induced quantum spin state.

Results and Discussion

YbMgGaO₄ and LuMgGaO₄ are members of the Ln³⁺M²⁺M′³⁺O₄ family (R₃₃m symmetry, a ≈ 3.4 Å and c ~ 25 Å), where Ln is Lu or Yb, and M and M’ are 3d transition metals. Spin frustration has been observed in the other members of this structural family, including YbCuGaO₄, LuCuGaO₄, LuCoGaO₄, LuZnFeO₄ and LuCuFeO₄. Unfortunately, the chemical disorder in these compounds, along with the geometrical frustration, resulted in spin-glass behavior. Compounds with magnetic ions occupying only the lanthanide sub-lattice (triangular layer), such as YbZnGaO₄, have not been reported thus far. Our experiments demonstrate that pure YbZnGaO₄ is difficult to synthesize in air because ZnO is volatile at T > 1200 °C. However, MgO remains stable at T < 1500 °C, as confirmed by the almost zero mass loss of this compound after heating. This thermal stability of MgO allows us to successfully synthesize pure YbMgGaO₄.

For accuracy, we employed two starting crystal structure models: YbFe₂O₄ (I) and LuCuGaO₄ or YbCoGaO₄ (II). In model I, Ln³⁺ locates at its ideal position (0, 0, 0). We fit the observed XRD intensities with reasonable refined parameters and small residuals in the Supplementary Information. In the case of the alternative model II, Ln³⁺ slightly deviates from its ideal position to (0, 0, z₁ₐ), where z₁ₐ ≈ 0.004. In this case, we also achieved a reasonable refinement using model II. However, an extra refinement parameter z₁ₐ was required. Even when the deviation exists, the maximum displacement along the c-axis is only approximately ± 0.004c (0.1 Å). This possible displacement may be due to Mg²⁺:Ga³⁺ disorder in the interleaved double layers (Fig. 1a) and may cause slight bond randomness. Other researchers have reported that z₁ₐ is ~ 0.009 in LnCuGaO₄, whereas z₁ₐ is ~ 0.006 in LuCoGaO₄ and z₁ₐ is ~ 0.005 in YbCoGaO₄. These results are consistent with the fact that Cu²⁺ is Jahn-Teller active, whereas Mg²⁺, Co²⁺ and Ga³⁺ are not. Notably, the strict R₃₃m symmetry is maintained in both cases; i.e., a triangular spin lattice remains spatially isotropic.

The crystal structure is shown in Fig. 1a. Triangular layers of magnetic YbO₆ octahedra, interleaved with double layers of nonmagnetic MgO₂ and GaO₂ triangular bipyramids, are ABC-stacked along the c-axis. Thus, the interlayer magnetic coupling can be negligible compared to the intralayer superexchange interaction between neighboring Yb³⁺, which is directly mediated by two parallel anions, O₁⁻. Site-mixing between magnetic and nonmagnetic ions is forbidden because the radius of Yb³⁺ is much larger than that of Mg²⁺ or Ga³⁺. This result is confirmed by the lack of observable narrow electron spin resonance (ESR) signals of isolated Yb³⁺ ions in YbMgGaO₄ (Supplementary Information). The spatially isotropic triangular layer of YbO₆ is shown in Fig. 1b. Because the inversion centers are located at halfway sites between neighboring Yb³⁺ ions (Fig. 1b) and Yb³⁺ sites (Fig. 1c), the DM interactions between the first-, second-, and third-neighbor Yb³⁺ vanish according to Moriya’s rules. Since Yb³⁺ is a Kramers ion, the low-T (≤ 30 K, see below) magnetism of YbMgGaO₄ should be dominated by the Kramers doublet GSs with effective spin-1/2.

The magnetizations are strongly suppressed with increasing Yb³⁺ concentration or effective exchange coupling (Fig. 2a,b); this result suggests an AF neighbor exchange energy. The AF coupling is confirmed by the negative Weiss temperatures, θₓ < 0 (Fig. 2c). The Curie-Weiss fit gives the Weiss temperature, θₓ = −4.11(2) K, and the averaged g-factor, 3.2(1), for YbMgGaO₄. At T ≪ |θₓ|, there is a slight deviation from the Curie-Weiss law (Fig. 2c), which corresponds to an upturn of susceptibility (Fig. 2d). A broad peak in bulk susceptibility well below |θₓ|, which was observed in lots of frustrated spin systems, is absent in our samples. The underlying reason why the broad peak is absent in YbMgGaO₄ is still unclear at present, considering the fact that impurity spins can be ignored and DM interactions...
are symmetrically forbidden in YbMgGaO$_4$ (Fig. 1b,c). To reveal the exact reason, further experimental and theoretical efforts based on high-quality single crystals$^{21}$ are highly required. In fact, the broad peak seems not to be a common feature of 2D frustrated spin systems. For instance, it is absent in the bulk susceptibility of herbertsmithite no matter whether the contribution from impurity spins is taken into account or not$^{22}$. The spin-1/2 antiferromagnetic Heisenberg model on the triangular lattice with a quenched randomness$^{23}$ ($\Delta > 0.7$) or on the kagome lattice with DM interactions$^{24}$ ($D_p \sim 0.3 J$) also gives no broad peak in uniform susceptibility well below $|\theta_w|$. Saturation behavior is observed in the MH curves below 10 T (Fig. 2b), whereas a linear dependence emerges above 10 T; this behavior is attributed to Van Vleck magnetism$^8$ and is related to field-induced electronic transitions. The saturation magnetizations are in agreement with the above averaged g-factor, $M_s = 1.600(2) \mu_B$/Yb$^{3+}$, gave/2, where the powder-averaged g-factor, $g_{\text{ave}} = 2/3 g_{\perp} + 1/3 g_{\parallel}$. In addition, a Van Vleck susceptibility (the fitted slope), $\chi_{vv} = 0.0122(1) \mu_B$/Yb$^{3+}$/T, is also obtained.

Experimentally determining the GS of YbMgGaO$_4$ is of particular interest and non-trivial. The susceptibilities measured under zero field cooling (ZFC) and FC show no observable differences at least down to 0.48 K (Fig. 2d). A complete magnetization loop ($-7$ T to $7$ T) was also measured at 0.5 K, and no hysteresis was observed (inset of Fig. 2d). The heat capacities measured under 0 to 9 T show no sharp $\lambda$-type peak down to 60 mK (Fig. 3a,b). These observations are incompatible with the long-range AF/ferromagnetic or short-range spin glass$^{25}$/spin ice$^{26}$ transition. It should be pointed out that the quasi-long-range Kosterliz-Thouless (K-T) transition also seems unlikely, since the above results are contrary to the typical thermodynamic changes accompanying the K-T transition$^{27,28}$. The results consistently suggest no spin freezing at least down to 60 mK, despite the AF exchange energy ($\Theta_w \sim -4$ K). In addition, YbMgGaO$_4$ is a good insulator with a room temperature resistance greater than 20 M$\Omega$. These results imply that the compound is a new QSL candidate.

The discrepancy between the experimental observations and numerical results$^{29}$ of the spin-1/2 triangular isotropic Heisenberg model, XY model or more general XXZ model is attributed to the following possibilities. First, the effective-1/2 exchange matrix between neighbor Yb$^{3+}$ may significantly deviate from the ideal Heisenberg, XY or XXZ model$^{30}$, and may cause much stronger spin frustration and fluctuation at low temperatures. And we would like to discuss this in detail in the further experimental and theoretical works basing on high-quality single crystals$^{31}$. Second, slight bond randomness may exist in YbMgGaO$_4$ according to the crystal structure model II, which may influence the GS. The bond randomness can prevent spin ordering in THAF according to the exact diagonalization (ED) study$^{32}$. Third, the next-nearest-neighbor and/or longer exchange interactions may stabilize a spin disordered GS against AF states$^{31,32}$. Finally, the multiple-spin$^{33}$/ring exchanges$^{34}$ can also prevent Néel long-range ordering in

![Figure 1](https://www.nature.com/scientificreports/)

**Figure 1.** (a) Polyhedral structure of YbMgGaO$_4$. The black dashed lines indicate the unit cell. (b) Top view of the triangular layer of YbO$_6$ octahedra. (c) Local crystal structure around the Yb$^{3+}$ Kramers ions.
a triangular lattice. Quantum fluctuations play an important role in the calculated ordered GS because of the geometrical frustration even in Heisenberg case\(^5\). This fact means that some seemingly small perturbations may be critical in the determination of the GS of the frustrated spin system at low temperatures.

The heat capacity is a highly sensitive probe for the low-energy excitations from the GS. The availability of a perfect nonmagnetic reference compound LuMgGaO\(_4\) enables an accurate exclusion of the lattice heat capacities without any fitting. Such advantage is absent in most of the existing QSL candidates. The measured heat capacities of LuMgGaO\(_4\) well follow the Debye law with a Debye temperature \(\approx 151\) K (Fig. 3a). Thus the exact magnetic heat capacities of YbMgGaO\(_4\) can be precisely extracted by directly subtracting the lattice contributions, i.e., the heat capacities of LuMgGaO\(_4\), from those of YbMgGaO\(_4\) (Fig. 3b). The heat capacity measurements were performed up to 30 K, at which the spin-1/2 entropy has been fully released as \(T \gg |\theta_w|\), and the Yb\(^{2+}\) ions remain in the Kramers doublet GSs with effective spin-1/2 as \(C_m(\sim 30\) K, 0 T) \(\approx 0\) (Fig. 3b). As the temperature decreases, the magnetic heat capacities of YbMgGaO\(_4\) exhibit a broad hump, whose position is almost field-independent (\(\mu_0 H \leq 2\) T), and shifts to a higher temperature with further increasing applied magnetic fields (\(\mu_0 H \geq 4\) T). The broad hump centered at 2.4 K (under 0 T) may suggest a crossover into a QSL state\(^35\). At \(T < 2\) K, the magnetic heat capacities well follow power-law temperature dependences down to the lowest measurement temperatures (Fig. 3b). The fitted power exponent, \(\gamma \approx 0.7\), approaches the theoretical value of 2/3 reported in the THAF spin liquid with ring exchanges\(^34\). It is another evidence that YbMgGaO\(_4\) is a new strongly correlated QSL candidate.

The released magnetic entropies from the lowest measurement temperature to 30 K are exactly \(\approx R\ln 2\) per mol Yb\(^{2+}\) in YbMgGaO\(_4\) (Fig. 3d). In other words, the residual spin entropy at 60 mK is almost zero. This means that the gapless QSL candidate has a disordered but macroscopically non-degenerate GS at low temperatures, and thus the third law of thermodynamics remains unviolated. Moreover, the near-zero upper limit of residual entropy (\(-0.6\%\)) at 60 mK almost excludes the possibility of magnetic transitions.
at lower temperatures and indicates that a possible QSL GS is experimentally achieved in YbMgGaO$_4$, since there are no sufficient entropies for spin symmetry breaking any more below 60 mK.

The neighbor exchange energy, $|\theta_w|$, is on the order of several Tesla. The magnetization of YbMgGaO$_4$ under a stable magnetic field at very low temperatures is critical and interesting. Unconventional quantum spin states, such as quantum magnetization plateaus $8^{-10}$, can be induced by applied fields. The careful magnetization measurements of YbMgGaO$_4$ at 0.5 K under 0 to 7 T, as well as at higher temperatures, are shown in Fig. 4a. The derived susceptibilities (dM/dH) show a clear and anomalous plateau under fields of 1.6 to 2.8 T at 0.5 K, and this plateau completely disappears at higher temperatures (1.9, 2.5 and 4.2 K), as shown in Fig. 4b. The unconventional magnetization behavior was confirmed by repeating the measurements several times. It is unlikely that the anomalous plateau originates from impurity spins. As aforementioned, the ESR measurements have determined a negligible amount (<0.04%) of impurity/isolated spins. This is further confirmed by the fact that no Schottky anomaly from isolated spins is observed in magnetic heat capacity of YbMgGaO$_4$ (Fig. 3b) down to 60 mK$^{12}$. The very low level of impurity/isolated spins also can be naturally explained by the large chemical difference between the magnetic Yb$^{3+}$ and the other nonmagnetic ions. The difference prevents the magnetic site-mixing and hence guarantees an extremely low level of impurity/isolated spins. This is one of the key advantages of the new compound. We have made a linear fit to the magnetization data at 0.5 K in the constant susceptibility range (form 1.6 to 2.8 T) (Supplemental material), supposing it is contributed by easy-saturated “impurity” spins and field-independent “intrinsic” susceptibility$^{12}$. The fitted intercept, $\sim 0.173(4) \mu_B$/Yb$^{3+}$ ($\sim$11% of the saturation magnetization), would give $\sim 11\%$ “impurity” spins, which is an unreasonably large amount and clearly contradicts with the ESR and heat capacity measurements. The magnetization curve at 0.5 K (Fig. 4a) is very similar to that of the triangular lattice compound C$_6$Eu along the c-axis at 4.2K$^{36}$. In the case of C$_6$Eu, this behavior has been attributed to traces of the 1/3 plateau$^{27}$. It should be pointed out that the fitted power exponent $\gamma$ also shows an unusual quasi plateau ($\gamma \sim 1.5$) in the similar magnetic-field range at extremely low temperatures (inset of Fig. 3c). The unusual $\chi$-plateau must be related to a field-induced unconventional quantum state at extremely low temperatures ($\leq |\theta_w|/10$).

The one-third quantum magnetization plateau, as well as higher-fraction quantum states, have been reported in Néel ordering phases ($T < T_N$) in spin-1/2 triangular-lattice antiferromagnets such as Ba$_2$Co$_3$Sb$_2$O$_9$ and Cs$_2$CuBr$_4$, and the Néel transitions are caused by the interlayer coupling$^{8^{-10}}$. To the best of our knowledge, YbMgGaO$_4$ is the first example that exhibits the abnormal magnetization behavior in

Figure 3. (a) Temperature dependences of total heat capacity under different magnetic fields in YbMgGaO$_4$ and LuMgGaO$_4$. The dashed curve denotes the Debye heat capacity. (b) Temperature dependences of magnetic heat capacity under different magnetic fields in YbMgGaO$_4$. The colored dash lines show the power law fits to the low-T magnetic heat capacities. (c) Magnetic heat capacity vs. 1/T in YbMgGaO$_4$. The red dash line shows the gapped spectral function fit to the low-T magnetic heat capacities under 0 T. Inset: fitted power exponent $\gamma$. (d) Temperature dependences of integral magnetic entropy under different magnetic fields in YbMgGaO$_4$. 

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
a paramagnetic phase in a 2D lattice. To fully understand the abnormal magnetization behavior, further theoretical and experimental studies are required.

Conclusion
A new triangular antiferromagnet with effective spin-1/2 and perfect $R3m$ symmetry, YbMgGaO$_4$, was successfully synthesized. The new compound overcomes some critical disadvantages in the existing materials, such as spatially anisotropic intralayer exchange interactions, magnetic defects, interlayer exchange coupling, and antisymmetric DM interactions. Despite the significant AF exchange coupling between neighbor spins ($\theta_w \approx -4 K$), no spin freezing was observed at least down to 60 mK, suggesting that YbMgGaO$_4$ is a new QSL candidate. The gapless low-energy excitations from the GS are evidenced by the low-$T$ power-law temperature dependence of heat capacity and nonzero susceptibility. Almost zero residual spin entropy suggests that the new compound extremely approaches a disordered and possible gapless QSL GS at low temperatures. An unconventional field-induced quantum state featuring an abnormal susceptibility plateau was observed for the first time, where the frustrated spin system was still paramagnetic at 0.5 K.

Methods
Yb$_x$Lu$_{1-x}$MgGaO$_4$ ($x = 1, 0.4, 0.16, 0.08$ and 0) white powder samples were synthesized from stoichiometric mixtures of Yb$_2$O$_3$ (99.99%), Lu$_2$O$_3$ (99.9%), MgO (99.99%) and Ga$_2$O$_3$ (99.999%). The mixtures were heated in air to 1450 °C for 4 days, with an intermediate grinding. The phase purities of all of the samples were confirmed by powder X-ray diffraction (XRD) (Bruker D8 Advance, 40 kV, 40 mA) before further study. High-intensity and monochromatic synchrotron XRD was used to more precisely determine the crystal structures of YbMgGaO$_4$ and LuMgGaO$_4$ at the diffraction station (4B9A) of the Beijing Synchrotron Radiation Facility (BSRF). The General Structure Analysis System (GSAS) program was used for Rietveld crystal structure refinements. No structural transitions were observed from the XRD patterns (Rigaku, 40 kV, 200 mA) at least to 12 K in YbMgGaO$_4$. Magnetization measurements at 0.48 to 300 K under 0 to 14 T were performed using a superconducting quantum interference device (SQUID) magnetometer (Quantum Design Magnetic Property Measurement System, MPMS) and a vibrating sample magnetometer (VSM) (Quantum Design Physical Property Measurement System, PPMS). The magnetizations were measured at 0.48 to 2 K using a He3 refrigeration system. The magnetizations of Yb$_x$Lu$_{1-x}$MgGaO$_4$ ($x = 1, 0.4, 0.16, 0.08$ and 0.04) were carefully corrected for the weak contribution ($<3\%)$ of the brass sample holder to the diamagnetic background. Heat capacity measurements between 3 K and 60 mK were performed using a dilution refrigeration system.

Figure 4. (a) Magnetic field dependence of magnetization at 0.5, 1.9, 2.5 and 4.2 K. The dashed line represents Van Vleck magnetism. (b) Magnetic field dependence of susceptibilities (dM/dH) at 0.5, 1.9, 2.5 and 4.2 K. The dashed lines show the interval where almost constant susceptibilities are observed.
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Author Contributions
Q.Z. conducted all of the experiments and analyses and wrote the paper. Y.L. synthesized the samples, collected synchrotron and laboratorial XRD patterns, performed structure refinements and magnetization and heat capacities and ESR measurements, analyzed all of the data, and prepared the figures and the draft. H.L. analyzed the magnetization data. L.L., Z.Y. and L.P. performed the magnetization measurements at 0.48 to 2 K. L.Z. collected low-temperature XRD patterns. Y.Z. performed ESR measurements. J.W. performed magnetization measurements. F.J. and Z.W. collected synchrotron XRD patterns. Z.Z. and S.L. performed the heat capacity measurements at 0.06 to 3 K.

Additional Information
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