Magnetoresistance (MR) is the magnetic field–induced change of electrical resistance. The MR effect not only has wide applications in hard drivers and sensors but also is a long-standing scientific issue for complex interactions. Ferromagnetic/ferri-ferromagnetic oxides generally show negative MR due to the magnetic field–induced spin order. We report the unusually giant positive MR up to 17,200% (at 2 K and 7 T) in 12-nm Sr₂CrWO₆ thin films, which show metallic behavior with high carrier density of up to $2.26 \times 10^{28} \text{ m}^{-3}$ and high mobility of $5.66 \times 10^{4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. The possible mechanism is that the external magnetic field suppresses the long-range antiferromagnetic order to form short-range antiferromagnetic fluctuations, which enhance electronic scattering and lead to the giant positive MR. The high mobility may also have contributions to the positive MR. These results not only experimentally confirm that the giant positive MR can be realized in oxides but also open up new opportunities for developing and understanding the giant positive MR in oxides.

INTRODUCTION

The magnetoresistance (MR) effect of a material is the change in electrical resistance in response to an external magnetic field. Materials with large MR are a long-standing goal for research and still at the forefront of contemporary science because of their applications in sensors, memories, and spintronic devices and the fundamental physics of complex interactions. MR is generally defined as $\frac{\rho(H) - \rho(0)}{\rho(0)} \times 100\%$, where $\rho(H)$ and $\rho(0)$ are the resistivities with and without magnetic field, respectively. Accordingly, MR can be positive and negative. In general, positive MR occurs in normal metals because of the Lorentz force, but the magnitude is only about a few percent. Giant positive MR has been found in many types of materials, such as magnetic semiconductors (1, 2), small-gap or zero-gap semiconductors (3, 4), ferromagnetic alloys (5), topological insulators (6), and semimetals (7, 8). Many topological Dirac or Weyl semimetals also show extremely larger positive MR (9–13). Negative MR, on the other hand, specifically relates to the multilayer structure of adjacent ferromagnetic layers (metals or ABO₃-type perovskite oxides), known as giant MR (GMR) and colossal MR (CMR). In the GMR and CMR materials, the magnetic field reduces the local spin disorder and thus decreases resistivity and leads to negative GMR or CMR (14–18).

Ferromagnetic ABO₃-type perovskite oxide generally illustrates negative CMR (15–19), although, by artificial microstructure design, positive MR can be achieved in perovskite oxides, but the MR value is very small (20, 21). ABO₃-type perovskite oxides provide structural flexibility to tune functional properties. For example, by introducing B and B′ to the B site alternatively, that is, forming B site–ordered $\text{AB}_2\text{B}′\text{BO}_₅$-type double-perovskite ferrimagnetic oxides, which generally show extremely high Curie temperatures (17, 22), positive MR with a moderate value of up to ~265% can be achieved (23–27). However, at least two issues about the positive MR in double-perovskite ferrimagnetic oxides need further investigations. The first issue is that the reported phenomenon is delicately dependent on experimental parameters. In most cases, double-perovskite materials show negative MR (17, 28), and only a few studies report the positive MR (23–27) or even both the positive and negative MR (29, 30). These paradoxical phenomena are due to the existence and domination of long-range magnetic field–induced ferromagnetic background (15, 31), which tends to result in negative MR (14–18). The other issue is that all the experimentally observed positive MRs in double-perovskite oxides (23–27, 29, 30) are about three to four orders of magnitude lower than those in semimetals or semiconductors (1–13). Therefore, ferromagnetic/ferromagnetic oxides with giant positive MRs are still desirable.

Here, we report the robust giant positive MR up to 17,200% (at 2 K and 7 T) in half-metallic double-perovskite Sr₂CrWO₆ (SCWO) thin films fabricated under low pressure and discuss the mechanism of the GMR. SCWO is a typical $\text{A}_2\text{B}′\text{BO}_₅$-type double-perovskite oxide, where $\text{Cr}^{3+}$ and $\text{W}^{5+}$ occupy the B and B′ sites, respectively. The antiferromagnetic coupling between $\text{Cr}^{3+}$ (3d⁵, $S = 3/2$) and $\text{W}^{5+}$ (5d⁵, $S = 1/2$) leads to a macroscopic ferrimagnetic nature with a net magnetization of $2 \mu_B$/formula unit (f.u.), as will be shown in the following theoretical calculation. The corresponding Curie temperature ($T_c$) is ~460 K (28, 32). However, the synthesis of single-phase, high-quality SCWO is very difficult due to the parasitic phases (33, 34). Therefore, only several experimental works on SCWO have been reported (30, 32–34).

RESULTS

Figure 1 (A to C) shows the atomic force microscopy (AFM) surface morphologies of the SCWO films deposited for 1000, 2000, and 4000 laser pulses under a low pressure of $10^{-5}$ Pa. As can be seen, all films show very flat surfaces, and the surface roughness increases slightly as the thickness increases, indicating the high quality of these films. The root mean square surface roughness is 348.4, 502.7, and 1420 pm for the three films, respectively. From the typical cross-sectional transmission electron microscopy (TEM) morphology of the films deposited for 4000 pulses shown in Fig. 1D, the thicknesses of the films deposited for 1000, 2000, and 4000 pulses are 12, 24, and 48 nm, respectively, which are abbreviated as S-1, S-2, and S-3, respectively. The homogeneous cross-sectional TEM morphology and the sharp, coherent substrate-film interface (Fig. 1, D to F) indicate the high quality of the epitaxial films, which is further confirmed by the x-ray diffraction (XRD) measurements (Fig. S1). Both S-1 and S-2 only show diffraction
shoulders at the left of the SrTiO₃ (STO) diffraction peaks, indicating that the films are well strained to form a coherent interface with the (001) STO substrate surface. By noting that STO has a cubic structure with a lattice constant of \( a = 3.90 \) Å and that SCWO has a cubic structure with a lattice constant of \( a = 7.89 \) Å (28, 30), it is deduced that S-1 and S-2 should have a slight compressive strain. With increasing thickness, the strain tends to relax. Therefore, S-3 has an obvious independent diffraction peak.

The magnetization-magnetic field \((M-H)\) curves of all the samples under low pressure were measured at 10 K with a magnetic field of 2 T and plotted in Fig. 2. The maximum magnetization values are 1.4, 1.1, and 0.6 \( \mu_B/\text{f.u.} \) for the S-1, S-2, and S-3 thin films, respectively, which are smaller than the theoretical prediction of 2.0 \( \mu_B/\text{f.u.} \), as will be shown later. The decreasing magnetization in the double perovskite is usually due to the anti-site defects (ASDs) or the oxygen vacancy (28, 35). However, we believe that the concentration of ASD or the oxygen vacancy should not be very high because we find that the films have a very small resistivity, as we will discuss later. The magnetization values obtained here are closer to those reported by other studies on SCWO films (30, 34), and the magnetization increases monotonically as the temperature decreases in the temperature range between 10 and 300 K (fig. S2), which is attributed to the ferrimagnetic nature of SCWO (30, 32).

The temperature dependence of the resistivity \((\rho)\) was measured by method-1 (schematically shown in fig. S3A) with and without an external magnetic field (2 T), which is shown in Fig. 3 (A to C). Without magnetic field, all films show a metallic behavior in the whole temperature range (that is, the resistivity increases as the temperature increases). The measured resistivities at 2 and 300 K are \( 4.9 \times 10^{-11} \) and \( 9.5 \times 10^{-7} \) ohm·m, \( 4.5 \times 10^{-10} \) and \( 1.1 \times 10^{-6} \) ohm·m, and \( 9.2 \times 10^{-10} \) and \( 3.7 \times 10^{-6} \) ohm·m for the S-1, S-2, and S-3 thin films, respectively. The resistivity of the SCWO thin films is comparable to that of a good metal and much smaller than that reported by others on SCWO epitaxial films or bulk ceramics (30, 32–34). The residual resistance ratios \((\text{RRR})\) between 300 and 2 K are 19,387, 2444, and 4111 for the S-1, S-2, and S-3 thin films, respectively. The extremely high RRR value again confirms the good quality of these films. To extract the characteristics of the carriers and carrier density, we measured the temperature-dependent Hall effect. The corresponding calculated temperature-dependent carrier density and mobility of all samples are plotted in Fig. 4 (A and B), respectively (the inset of Fig. 4B presents the Hall coefficient). At 2 K, the n-type carrier densities were calculated to be 2.25 \( \times 10^{28} \) m⁻³, 1.18 \( \times 10^{28} \) m⁻³, and 3.82 \( \times 10^{27} \) m⁻³ for the S-1, S-2, and S-3 thin films, respectively. The carrier density shows almost negligible temperature dependence. The mobilities, on the other hand, were 5.66 \( \times 10^{4} \) cm² V⁻¹ s⁻¹ for the S-1, S-2, and S-3 thin films, respectively, at 2 K. For each film, the mobility decreases as the temperature increases.
It is necessary to emphasize that low pressure during deposition is important to obtain high-quality samples. As a comparison, two SCWO films were also deposited on the (001) STO substrate for 2000 pulses under oxygen pressures of 0.5 and 1.5 Pa. Although the XRD patterns indicate that the films are well oriented along the $c$-axis (fig. S4), the AFM and TEM morphologies show that the films deposited under relatively higher oxygen pressures have rougher surfaces (fig. S5). The temperature-dependent resistance of the SCWO films fabricated under an oxygen pressure of 0.5 Pa is plotted in fig. S6. The figure shows a semiconductor-like behavior in the whole temperature range, which is due to the bad quality of the film. The SCWO films fabricated under an oxygen pressure of 1.5 Pa show similar behavior in the higher temperature range, but the resistance is too high to be measured in the lower temperature range. Here, we will focus on the films fabricated under low oxygen pressure ($10^{-5}$ Pa).

However, the low pressure during sample fabrication could induce the oxygen vacancies in both the SCWO films and the STO substrate. The oxygen vacancies have different effects on the SCWO films and the STO substrate. For the metallic SCWO films, they can increase the scattering of the electrons and the resistivity of the materials, which can even possibly lead to the semiconductor behavior due to the defective band that crosses the Fermi level ($36–42$). However, as shown...
above, the resistivity of the three films is very small, which is comparable to that of the best metals. Therefore, we believe that the oxygen vacancy has no predominant effect on the transport property in these thin films.

The oxygen vacancies, on the other hand, have a totally different effect on the insulating STO substrate. The oxygen vacancies can introduce carriers in the insulator or semiconductor and increase the electric conductivities. It is well known that the oxygen-deficient STO behaves like metals (43, 44). Therefore, it is interesting to know the impact of the low pressure–induced oxygen vacancies on the STO substrate and how the oxygen vacancies affect the transport and MR results. To investigate these influences, the bare STO substrate was heated at the same temperature (830°C) and pressure (10^5 Pa), and then the temperature-dependent resistance was measured as shown in fig. S3. The resistance was first measured by method-1, which is schematically shown in fig. S3A, and the result is shown in fig. S3B. It is found that the heated bare STO substrate shows metallic behavior due to the oxygen vacancies, which is consistent with other reports (34, 43, 44). However, two important features should be emphasized. The first feature is that the resistance of the heated bare STO is at least two orders of magnitude higher than that of the SCWO film (S-2) that is measured with the same method. With a simple model, the resistances of the STO substrate and the SCWO film are in parallel connection; therefore, the significantly smaller resistance of the SCWO film should be dominant for the macroscopic resistance (that is, the resistance of the STO can be neglected). The second feature is that the heated bare STO shows almost zero MR in the whole temperature range (fig. S3C), consistent with a previous report (44). Because our SCWO films show the giant positive MR at low temperatures (fig. S3D and Fig. 3), as will be discussed later, the heated STO has almost no influence on the macroscopic MR. For further confirmation, the resistivity and MR of the S-2 films with the same geometry size were measured by using the wire bonding techniques, as schematically shown in fig. S3A (labeled as method-2); the result is shown in fig. S3D. In this case, the contributions of both the substrate and the film to macroscopic transport properties are measured together. As can be seen from fig. S3 (B and D), good metallic behavior and giant positive MR are still observable. The results are similar to those measured in the SCWO films but different from those in the heated bare STO substrate. On the basis of these discussions, it may be reasonable to conclude that the contribution from oxygen vacancy of the heated STO substrates to the macroscopic transport property and MR should be negligible.

Here, it is also necessary to briefly discuss the effect of strain on magnetic property. In general, strain in ultrathin metallic perovskite oxide films can induce metal-insulator transition; the critical thickness is less than 10 units (for example, <4.0 nm) (41, 45). However, the thicknesses of our films are 12, 24, and 48 nm, indicating that these SCWO films are quasi–two-dimensional or even three-dimensional, that is, the strain cannot have determinant contributions to the transport nature of these films.

To understand the basic physical properties of SCWO, we have calculated its band structure and density of states (DOS) by the OpenMX code (46) within the framework of the density functional theory in the generalized gradient approximation (47). From the results shown in Fig. 5, we can see that SCWO is a perfect half-metal, where one spin component shows the metallic properties and the other has a large band gap of about 2.3 eV. The half-metallicity nature of SCWO is commonly found in double-perovskite materials, such as Sr₃FeMoO₈ (17). Note that the SCWO is definitely not a semimetal or a zero-gap semiconductor from the band structure. Besides, we find that the total net spin moment is 2.0 μB/f.u. in our calculation, which is the same as the theoretical expectation. Because of the ASD or oxygen vacancies in SCWO, the observed magnetization in our experiments is smaller than that of calculated results mentioned above.

As shown in Fig. 3 (A to D), when an external 2-T magnetic field perpendicular to the thin film is applied, the resistivity of each sample tends to increase when the temperature is lower than 20 K and increases sharply when the temperature is lower than 10 K. This increased resistivity leads to a giant positive MR of 1960%, 245%, and 190% at 2 K for SCWO films S-1, S-2, and S-3, respectively. It is found that the positive MR has negligible dependence on the external magnetic field direction. As evidenced by the temperature-dependent resistivity of S-2 between 2 and 300 K with the applied magnetic field parallel to the thin film (but still perpendicular to the electric current) (fig. S7), the positive MR value is about 235% at 2 K, which is very close to that in Fig. 3B.

Figure 6 shows the field-dependent MR of the S-1, S-2, and S-3 thin films at 2, 4, 10, and 20 K. The maximum positive MR values at 2 K and 7 T are 17,200%, 2060%, and 1185% for S-1, S-2, and S-3, respectively. The MR shows no saturation trace even up to 14 T, as confirmed by the field-dependent MR of the S-2 films (fig. S8). To the best of our knowledge, this positive MR is the largest value ever observed in ferromagnetic/ferrimagnetic perovskite oxides and is two orders larger than that reported by others on half-metallic double-perovskite oxide thin films or bulks (23–27, 29, 30).

**DISCUSSION**

We will first analyze the mechanism of the giant positive MR in SCWO thin films. In our opinion, the mechanism for the giant positive MR here might be a very novel one. We therefore can only provide the following discussion. Recently, it is theoretically proposed that giant positive MR could be observed in the antiferromagnetic metallic double perovskite (31). An applied field could suppress the long-range antiferromagnetic ordering, leading to a state with short-range antiferromagnetic correlations and strong electronic scattering, which induces the giant positive MR. According to the above discussion, our SCWO thin films show very good quality and metallicity, which implies a long-range antiferromagnetic ordering (ferrimagnetic) and high spin ordering degree. Therefore, the SCWO could possibly be a materialization of the...
not require the semimetallicity and can be extended to other types of materials. It is obvious that the SCWO is not a semimetal according to Fig. 5, but the carrier mobility is also very high, as shown in Fig. 4B. Therefore, the high mobility in the SCWO thin films could also be the possible origin of the giant and nonsaturating positive MR. Especially, it is observed that when the thickness of the films increases, the conductivity and the mobility decrease because of the increase of ASD, leading to the decrease of MR, which is possibly due to the decrease of the mobility.

However, it seems that we cannot distinguish the above two mechanisms based on the present work; thus, more future experimental works are needed. For these studies, high-quality samples with metallic nature and low ASD might be the key factor because ASD plays a detrimental role in suppressing conductivity and, therefore, mobility (49, 50).

Finally, we need to exclude the possible contributions of nanosized Cr, CrO2, or Cr2O3 clusters, which might be formed during sample fabrication or processing, to macroscopic transport properties. According to literatures, metallic Cr or CrO2 shows negative MR (14, 51), whereas insulating Cr2O3 cannot lead to the observed metallicity and thus large positive MR. Both are in contrast to our experimental results. Therefore, it is reasonable that Cr or CrO3 is not responsible for the observed giant positive MR. Metal Cr is not detectable because the Cr cations have mixed valences of +2 and +3 at room temperature and at a low temperature of 14 K (52), as shown in Fig. S10A. In addition, the x-ray photoelectron spectroscopy (XPS) measurements confirm that there is no W metal (fig. S10B). The asymmetric peaks centered around 35.4 and 37.6 eV should be the sum of two peaks. According to the simulation, the peaks around 35.0 and 37.2 eV are attributed to W5+, whereas those around 35.7 and 37.8 eV are attributed to W6+ (53). This means the W cations have mixed valence states of +5 and +6. One of the possible reason for the formation of W6+ is that Ar+ ions sputter on the surface, disrupt the lattice, alter bonding and electronic configuration, leading to the decrease of MR, which is possibly due to the decrease of the mobility.

In conclusion, unusually giant positive MR up to 17,200% is observed in half-metallic ferrimagnetic SCWO films deposited under low pressure. This finding is possibly attributed to the fact that external magnetic field suppresses the long-range antiferromagnetic coupling between Cr and W cations, leading to short-range spin fluctuations. This field-induced spin disorder enhances the electronic scattering and thus results in positive MR. At the same time, high mobility may also contribute to the giant positive MR. In addition, ASD can suppress the positive MR by forming local antiferromagnetic regions, which destroy the metallic nature of SCWO. We believe that this work not only sets a record for the giant positive MR in magnetic perovskite oxides but also stimulates further investigation and understanding of the MR materials.
**MATERIALS AND METHODS**

Sample preparation

SCWO thin films were prepared on a (001) STO substrate by pulsed laser deposition using a 248-nm KrF excimer laser. A stoichiometric SCWO ceramic was used as the target. Before each deposition, the chamber was evacuated to a base pressure of 10⁻⁵ Pa. During each deposition, the substrate temperature was 830°C, the laser repetition rate was 2 Hz, the laser energy density was 1.2 J/cm², and the pressure was 10⁻⁵ Pa. After each deposition, the films were in situ annealed in the chamber for 10 min, cooled to 400°C at a rate of -10°C/min, and then naturally cooled to room temperature.

**Structural and property measurements**

The structures of the films were investigated by XRD (Rigaku Ultima III), AFM (Cypher), and TEM (FEI Tecnai F20). The cross-sectional sample was prepared by a focused ion beam (FEI Strata 235). The transport properties of the SCWO films fabricated with an oxygen pressure of 0.5 Pa were measured using a physical property measurement system (PPMS-9, Quantum Design). The complex impedance measurements were performed using an ac bridge (ZItester, Autolab). The magnetoresistance measurements were performed using a SQUID magnetometer (Magnex,model 3C). The magnetic properties were measured using a superconducting quantum interference device (MPMS XL-7, Quantum Design) and a physical property measurement system (PPMS-9 and PPMS-16, Quantum Design).

**First-principles calculations**

The band structure and DOS of double-perovskite SCWO were calculated by the OpenMX code [46] within the framework of the density functional theory in the generalized gradient approximation [47]. SCWO has a cubic structure with the space group of Fm3m, and its experimental lattice constant (a = 7.89 Å) was used in our calculation. The bases used in the OpenMX code are s⁺p⁺d⁺³, s⁺p⁺d⁺², and s⁺p⁺d⁺ for Sr, Cr, W, and O elements, respectively. The energy cutoff for real-space integration is 300 rydbers, and a k-points mesh of 15 × 15 × 15 was used for the Brillouin zone integration. To include the effects of the on-site Coulomb energy, we used the effective Hubbard U of 2.13 eV for Cr [55].

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/11/e1701473/DC1.

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