Giant Magnetoresistance in an all-oxide spacerless junction

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We report the fabrication of an oxide-specific type of magnetoresistive junction, which is a ferromagnetic bilayer. Both electrodes are high spin-polarization oxides:agnetite (Fe₃O₄) and manganite (La₀.₇Sr₀.₃MnO₃). Negligible magnetic coupling between both ferromagnetic electrodes is realised, which allows to obtain parallel and antiparallel magnetic configurations of the electrodes when sweeping the applied magnetic field. The structure exhibits negative giant magnetoresistance (GMR) at low temperatures. This negative MR shows that both electrodes stay spin-polarized at the interface and have opposite spin polarizations, i.e. the Fe₃O₄ layer has a negative spin polarization at low temperature. Maximum GMR (-5 %) is obtained at 55 K.

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I. INTRODUCTION

Highly spin-polarized ferromagnetic oxides, such as CrO₂, Fe₃O₄, La₀.₇Sr₀.₃MnO₃ (LSMO) have been the focus of recent fundamental and technological studies in the field of spin electronics. Using these materials, various devices, such as giant magnetoresistance (GMR) junctions [1] and tunnel magnetoresistance (TMR) junctions [2] have been fabricated and studied. To fabricate a magnetoresistive device based on a junction, usually two ferromagnetic layers are separated by a thin non magnetic spacer (TMR).

In the usual case of transition metal electrodes, the thickness of the spacer is chosen in order to control the spin-dependent transport mechanism at the interface: metallic spacer (GMR) or insulating spacer (TMR).

In the usual case of transition metal electrodes, the thickness of the spacer is chosen in order to magnetically decouple the magnetic layers (i.e. thickness larger than a few atomic planes to break the direct coupling exchange path, and to prevent indirect coupling such as the RKKY one). The transport across the spacer must conserve the spin information, thus the spacer thickness must be kept thinner than a few mean free paths (current-in-plane cip-GMR) or spin diffusion lengths (current-perpendicular-to-plane cpp-GMR) or a few 1/kF (tunnel probability in TMR junction).

A spacer is not necessary if it is possible to weaken the magnetic coupling between both electrodes. Such devices have already been proposed: ballistic junctions [3] or break junctions [4]. These junctions were designed to break the exchange coupling between two transition metals. To achieve this, both electrodes have to be mechanically separated, which is a difficult step, source of non reproducibilities, and sensitive to parasitic phenomena such as magnetostriction [5]. In this paper we propose a simple solid state structure, adapted to collective fabrication.

In oxides, magnetic coupling is due to indirect exchange (3d ion - oxygen - 3d ion) and it is very sensitive to the atomic details of such a bond. For example it is possible to weaken the coupling by changing the bond angle (manganite’s Tₘₐₜ varies as a function of the Mn-O-Mn bond angle [4]). Thus, tuning the interfacial magnetic coupling is achievable in oxides.

The spin-polarization of a material is positive if the majority spin at the Fermi level is parallel to the magnetization and negative if the minority spin at the Fermi level is parallel to the magnetization. Half metallic ferromagnets have a spin polarization of 100 % (only one spin direction is present at the Fermi level). Magnetite (Fe₃O₄) stands out as a predicted half metallic ferromagnet (ferrimagnet in fact) with negative spin polarization [6] and a remarkably high Curie temperature (Tₘₐₜ) of 858 K. La₀.₇Sr₀.₃MnO₃ is predicted to have 100 % positive polarization [8] with Tₘₐₜ of 350 K. A junction between two such half metallic ferromagnetic compounds would in theory behave as an ideal magnetic-field-controlled switch with 100 % negative magnetoresistance (MR).

As mentioned above, recent efforts have been made to fabricate Fe₃O₄/La₀.₇Sr₀.₃MnO₃ junctions, where I is SrTiO₃, CoCr₂O₄ [9, 10], but a Fe₃O₄/La₀.₇Sr₀.₃MnO₃ junction without a spacer has never been proposed.

II. EXPERIMENTAL DETAILS

The Fe₃O₄/La₀.₇Sr₀.₃MnO₃ bilayers were grown on (001)-oriented SrTiO₃ (STO) substrates using pulsed laser deposition. First, the LSMO layer was grown at 1023 K under 40 Pa of O₂, then the Fe₃O₄ layer was grown at 623 K under 5.10⁻⁴ Pa of O₂. The thickness of LSMO was 50 nm whereas the Fe₃O₄ was grown with two different thicknesses, 15 nm and 50 nm, estimated in situ by optical reflectometry. Prior to the deposition,
the substrate was heated in oxygen up to the deposition temperature.

To study the magnetotransport properties of the junction, 50 nm Au was deposited upon the 15/50 nm Fe\textsubscript{3}O\textsubscript{4}/LSMO/STO structure at room temperature using the sputtering technique and subsequently, junctions of 500 × 500 μm and 140 × 140 μm were fabricated by photolithography and Ar ion etching process. All transport measurements were carried out with ccpp geometry and applied magnetic field parallel to the plane.

X-ray diffraction (XRD) study was carried out to examine the structural properties of the bilayers. Despite the 1 % lattice mismatch between LSMO and STO, the LSMO growth is pseudomorphic up to a critical thickness (100 nm) larger than the thickness chosen for these bilayers. The LSMO film is epitaxially-strained (0.18° FWHM rocking-curve) and a large epitaxially-induced magneto-elastic anisotropy is present. The Fe\textsubscript{3}O\textsubscript{4} film on LSMO is textured with multiple orientations (diffraction peaks corresponding to [001] and [011] directions, but not [311]), whereas Fe\textsubscript{3}O\textsubscript{4} films on SrTiO\textsubscript{3} deposited under similar conditions, were grown textured along the [001] direction with 1° FWHM rocking curve. The details of the deposition of films of LSMO and Fe\textsubscript{3}O\textsubscript{4} on STO have been reported elsewhere.

### III. RESULTS

To check for magnetic coupling between both oxide layers, M(H) hysteresis loops of the unpatterned 50/50 nm bilayer structure were measured in the temperature range 10-350 K and up to 3 Tesla using a VSM and a SQUID magnetometer.

Fig. 1 shows the typical hysteresis loop from an unpatterned bilayer structure at 50 K, with the magnetic field applied along the substrate [110] axis, which is the easy direction of the La\textsubscript{0.7}Sr\textsubscript{0.3}MnO\textsubscript{3} layer. The hysteresis loop clearly shows two distinct coercive fields (about 5 mT and 100 mT), which correspond to LSMO and Fe\textsubscript{3}O\textsubscript{4}, respectively. The temperature dependence of Fe\textsubscript{3}O\textsubscript{4} coercivity shows the Verwey transition at 110 K (T\textsubscript{V} = 122 K in single crystals).

To study the magnetic coupling between these layers, minor loops of the softer layer (LSMO) were measured at 10 K. No shift of the loops was detected. Thus no exchange bias field larger than 3 mT exists. The large coercivity difference between layers and squareness of the LSMO hysteresis loop create well-defined parallel and antiparallel magnetic configurations.

As far as transport is concerned, we have measured I(V) characteristics from 5 to 300 K. They are linear up to the point where heating effects come into play. The evolution of the resistance with temperature (Fig. 2) can be divided into 3 regimes. The high temperature regime, above 90 K, exhibits the well-known resistance and CMR (colossal magnetoresistance) of manganites, and can thus be attributed to the LSMO electrode, which dominates the transport at these temperatures due to geometrical reasons. Between 30 and 90 K, the transport is thermally activated, due to the increasing Fe\textsubscript{3}O\textsubscript{4} dominance when temperature decreases. Below 30 K, a plateau is observed, which is surprising in a Fe\textsubscript{3}O\textsubscript{4}-dominated regime. This remains to be investigated, but could be explained by the onset of a hot electron transport mechanism due to the high electric field (40 kV/cm), such as the one observed in Fe\textsubscript{3}O\textsubscript{4}.

At high field (1 to 6 T range), the junction shows a negative magnetoresistance. This high field MR is large below 40 K (over -1 %/T, consistent with Fe\textsubscript{3}O\textsubscript{4} thin films), smaller between 40 and 100 K, and increases to high values (-2 %/T) at high temperature (LSMO CMR). In the intermediate regime, though, this negative slope is only visible above 2 T. Under that field, Fe\textsubscript{3}O\textsubscript{4} is poorly saturated, and GMR dominates (see below), giving a positive slope.

In the intermediate temperature range, the magnetic field dependent transport measurement shows a characteristic inverse GMR behavior (Fig. 3, the applied field is swept as the arrows indicate). The magnetic fields at which the junction resistance changes the most abruptly correspond to the coercivities of both oxide layers and the junction resistance is lower when the magnetizations of both layers are antiparallel to each other. The GMR was measured at ± 100 μA constant current as a function of temperature (Fig. 2). R\textsubscript{1↑↓} is measured at 20 mT and R\textsubscript{1↑↑} at 400 mT. This magnetoresistance diminishes in absolute value both above 90 K and below 30 K. The maximum GMR of -5.2 % is found at 55 K.

![FIG. 1: Hysteresis loops of unpatterned bilayer film at 50 K](image-url)
FIG. 2: Resistance and GMR of a 140×140 µm junction measured with a current of 100 µA

FIG. 3: R(H) magnetotransport measurement of a 140×140 µm junction for T = 55 K

IV. DISCUSSION

The two keypoints which have to be discussed are the origin of the magnetic decoupling of the La0.7Sr0.3MnO3 and Fe3O4 electrodes and the transport mechanism responsible for the large magnetoresistance. The ferromagnetic coupling mechanism is double exchange (Mn^{3+}-O-Mn^{4+} bonds) in the case of LSMO, and is superexchange (Fe^{3+}(A)-O-Fe^{2+/3+}(B) bonds) as well as double exchange (Fe^{3+}(B)-O-Fe^{2+}(B) bonds) in the case of Fe3O4. To magnetically decouple the two layers, these nearest-neighbour mechanisms have to be weakened. The interface between LSMO and Fe3O4 is a structurally disordered layer due to the 6.7 % lattice mismatch, which prevents heteroepitaxy. In oxides, due to the localised character of electrons, weakening the exchange is much easier than in transition metals where electrons are more delocalised (RKKY coupling range can reach a few nanometers). So one disordered layer due to the lattice mismatch between a perovskite and a spinel ferromagnet is enough to reduce the exchange coupling and to decouple both layers. This is a general statement since the lattice mismatch between a spinel and a perovskite structure will always be a few %.

We claim that our structure is a bilayer. However the presence of an intermixed layer between the electrodes has to be ruled out to support this claim. In our system any intermixing layer would be made of Fe and Mn ions and therefore it would be magnetic. Since transport in conducting ferromagnetic oxides is based on a nearest-neighbour hopping mechanism, any magnetic layer depolarizes the current. Thus, the characteristic type of magnetoresistance we can measure rules out the presence of an intermixing layer. Furthermore since the Fe3O4 layer is deposited at low temperature (623 K), the spinel/perovskite interface is expected to be stable. We have recently conducted a TEM study on the SrTiO3 / Fe3O4 interface, which showed that the perovskite / spinel mismatch can be accommodated through a regular array of dislocations [14] and confirms that no intermixing takes place.

As the MR is small (-5 %), spin disorder in the interfacial plane, leading to a partial depolarisation, cannot be ruled out. However, this cannot be called a distinct magnetic layer, it is better characterized as an interfacial disorder.

The two electrodes are in direct electrical contact, so the nature of the transport mechanism is related to the presence or absence of an electrical barrier. The cpp transport characteristics are ohmic, and the R(T) exhibits no regime that could be interpreted as a tunnel transport.

Since a significant MR has been measured and the characteristic fields of this MR are the coercive fields of both electrodes a spin-coherent mechanism has to be proposed. Through the interface, transport could still be based on a hopping mechanism. Interface disorder only reduces hopping integrals (also impacting magnetism by
reducing the double-exchange coupling). Thus the MR mechanism is closer to the cpp-GMR than to the TMR mechanism and the only significant interfacial resistance is the GMR itself.

The value of the magnetoresistance is difficult to interpret in a quantitative manner, given that the resistance of the interface is not dominant compared to that of the electrodes. Thus the -5.2 % GMR ratio is not intrinsic, and could be enhanced through an optimized junction pattern.

It is also worth noting that it is difficult to obtain a real parallel or antiparallel state with Fe₃O₄. Because of structural defects present in all Fe₃O₄ thin films (antiphase boundaries), the remanence is less than 80 %, and full saturation is not achieved even at large fields. Lastly, the decay of the GMR at high temperature can be explained by the decrease of LSMO polarisation well below Tₐ, as is known from other studies ([15]). Nonetheless, our measurements evidence negative spin polarization of Fe₃O₄, as predicted, down to the interface with LSMO.

For these reasons, we propose that the mechanism responsible for the exchange weakening and the large MR in our bilayer is fundamentally different from the TMR and spacer-assisted cpp-GMR mechanisms reported in other studies. It is purely interfacial (i.e. spacerless) and specific to oxides.

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