Giant Hall Effect in Nonmagnetic Granular Metal Films

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Nearly 3 orders of magnitude enhancement in the Hall coefficient is observed in Cu$_x$-(SiO$_2$)$_{1-x}$ granular films. This large enhancement of the Hall coefficient not only is significantly larger than the prediction of the classical percolation theory, but also occurs at a metal concentration identified to be the quantum percolation threshold. Measurements of the electron dephasing length and magnetoresistance, plus the TEM characterization of microstructures, yield a physical picture consistent with the mechanism of the local quantum interference effect.

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As a basic material constant, the Hall coefficient is generally indicative of the density and sign of the charge carriers. Thus, in granular metals, as the metal concentration decreases, the lower carrier density is expected to yield an enhanced Hall coefficient which peaks at the percolation threshold with a factor of ~30 for ~1 μm thick films [1,2]. Recently, however, it was found that in the magnetic (NiFe)-SiO$_2$, and Fe-SiO$_2$ granular films [2–5], the extraordinary Hall coefficient was enhanced by a factor of 10$^4$ when the metal volume fraction is close to $x = 0.53$ (the classical percolation threshold). An especially intriguing feature of this discovery is that, even after magnetic saturation, the ordinary Hall coefficient was still observed to increase by almost 3 orders of magnitude [4], suggesting a magnetic-independent mechanism could be operative.

In this Letter, we focus on the origin of the ordinary giant Hall effect (GHE) by studying the nonmagnetic Cu-SiO$_2$ granular system. We find the same 3 orders of magnitude enhancement in the Hall coefficient. By carrying out measurements on the electron dephasing length and the magnetoresistance (MR), and by characterizing our samples by transmission electron microscope (TEM) pictures, we find the Hall coefficient to peak at the quantum percolation threshold. Based on the picture that inhomogeneities (due to the small substructures) inside a dephasing length would necessarily cause local quantum interference and thereby modify the effective local properties, we show that all experimental data can be quantitatively accounted for within this simple framework. In particular, when the small substructures are suppressed through annealing, the GHE is shown to disappear, in agreement with the theoretical prediction [6].

Cu-SiO$_2$ granular films with different metal volume fractions were fabricated by using the cosputtering technique with a glass or Kapton substrate, at a temperature of 50°C. The base pressure of the chamber was kept below $2 \times 10^{-7}$ Torr. The films deposited on the glass were used for transport measurements and the films on Kapton were for composition determination. The metal volume fraction $x$ for all the films was obtained from energy-dispersive x-ray spectroscopy analysis. The dc resistance was measured by using the standard four-probe technique, and the Hall resistance was measured by using the five-contacts method.

More than 25 samples, with $0.4 < x < 0.81$ and a thickness of ~1 μm, have been fabricated and characterized. Longitudinal resistivity, $\rho_{xx}$, was measured as a function of temperature between 4 and 350 K. As quantum effects, such as hopping conduction or electron localization, are more dominant at low temperatures, we identify the (classical) percolation threshold by focusing on the temperature dependence of room temperature (300 K) resistivity [7]. We find the room temperature $d \ln \rho / d \ln T$ to change sign at $x = x_c = 0.43$, indicative of the percolation threshold. This is corroborated by the fact that, at $x = 0.42$, $\rho_{xx}$ exhibits the $\ln \sigma \propto -1/\sqrt{T}$ hopping conduction behavior [8] and is 2 orders of magnitude larger than that for $x = 0.45$. While not shown, we have also measured the magnetoresistance. The negative MR = $[\rho(H) - \rho(0)]/\rho(0)$, observed at 5 K for various compositions, is indicative of the electron weak localization effect. In particular, the MR magnitude is noted to peak above the percolation threshold.

The Hall resistivity, $\rho_{xy}$, for all samples was measured in a magnetic field $-4.6 \leq H \leq 4.6$ T and at temperatures ranging from 2.5 to 50 K. A perfect linear dependence of $\rho_{xy}$ on the magnetic field was observed. In Fig. 1, the measured $\rho_{xy}$ at 5 K is plotted versus $x - x_c$. It is seen that $\rho_{xy}$ increases monotonically with decreasing $x$, reaching a maximum of $R_0 = 1.84 \times 10^{-8}$ m$^2$/C at $x \sim 0.513$, and plateaus after that. The ratio of the highest value to the lowest is 700, almost 3 orders of magnitude, and this enhancement is nearly independent of the temperature up to 50 K. In magnetic granular films, the ordinary Hall coefficient is noted to increase by about 640 times [4], similar to what we find here. The maximum
$R_0 = 1.84 \times 10^{-8}$ m$^3$/C in our sample is larger than that observed in Al-Ge [9], Au-SiO$_2$, W-SiO$_2$ [10], and Al granular films [11], where the enhancement is about 100 or less. However, we should note that there are large fluctuations in the measured maximum $R_0$. Thus, a careful search in other granular systems might yield enhancement factors similar to what we have observed.

The measured enhancement in Hall coefficient is much larger than that expected by the classical percolation theory, where the maximum predicted enhancement factor (occurring at the percolation threshold) is on the order of $(T/a)^{g/n}$, where $T$ is the thickness of the film, and $a$ is the size of granules. Here, $g$ is the critical exponent of 3D Hall resistivity, and $n$ is the exponent for the 3D correlation length. The ratio $g/n$ is approximately given by $0.4/0.9 \sim 0.45$ [1]. By taking $T = 1$ µm, the thickness of our films, and a minimum value of $a = 1$ nm, we get $(T/a)^{g/n} \sim 30$ [2].

Besides the magnitude of the enhancement factor, another intriguing feature is that the maximum Hall coefficient does not occur at the percolation threshold, $x_c = 0.43$, but rather at a higher value of $x = 0.51$, unexpected within the classical picture. Below, we show that MR data, when coupled with electron dephasing length measurements and TEM characterization of the microstructures, identify $x \sim 0.51$ to be close to the quantum percolation threshold.

Electron dephasing length of our samples was obtained by the standard approach of measuring the magnetoresistance in low magnetic fields ($<0.5$ T), over a temperature range between 5–35 K, and then least-squares fitting the data to the 3D weak-localization predictions [12–15] for extracting the electron dephasing length $L_\varphi(T)$. The results are shown in the inset of Fig. 1. It is seen that $L_\varphi$ increases with decreasing temperature and increasing metal volume fraction. It varies from ~10 nm up to slightly below 100 nm. We have also characterized the microstructure of our samples by means of TEM pictures. In Fig. 2(a), it is seen that, for the sample with $x = 0.513$ and a corresponding large Hall coefficient, there are many small substructures on the order of 1 nm. Since this is considerably smaller than the measured $L_\varphi = 8$ nm, it follows that, within a footprint on the order of $L_\varphi \times L_\varphi \times L_\varphi$ [seen as a dashed circle on Fig. 2(a)], the local quantum interference must necessarily be present. In Fig. 2(b), the high resolution TEM picture of the same sample, annealed at 450 °C for 1 h is seen to have a grain size ~10 nm, with a clear suppression of the small substructures. The Hall coefficient of the annealed sample was measured to be more than 2 orders of magnitude smaller than before; i.e., the GHE has disappeared. There is thus a clear-cut correlation between the presence of small substructures, dephasing length, and the GHE.

An important indicator of electron scattering/localization effect is the product $k\ell$, where $k$ is the electron wave vector (renormalized by multiple scattering effects) at the Fermi level, and $\ell$ is its scattering mean-free path. From the measured MR and the dephasing length, and by setting the scattering mean-free path to be the size of the small substructures (1 nm at $x = 0.513$), we can directly deduce the value of $k\ell$ [16]. This is shown by the open circles in Fig. 1. It is seen that the composition where the Hall coefficient peaks corresponds with $k\ell \sim 1$, i.e., the quantum percolation threshold (or the mobility edge) [16]. For high metal concentrations, the $k\ell$ increases considerably even if we still assume the

![Graph showing the Hall resistivity plotted as a function of $x - x_c$, on a log-linear scale. The values of $1/k\ell$ for several typical samples are also shown; open circles are for the lower bound of $\ell = 1$ nm, solid diamonds are for the upper bound of $\ell = 10$ nm. The inset shows the temperature dependence of the dephasing length for several samples.](image)
For concentrations close to the quantum percolation threshold, the distribution is very symmetrical and peaks approximately at zero. While not shown, distributions of local (longitudinal) conductivities can also be similarly obtained [6]. The knowledge of the local (Hall and longitudinal) conductivity distributions at every concentration enables us to calculate the macroscopic conductivity tensor, where we use the classical (Kirchhoff) conditions of $\nabla \times \vec{E} = 0$ and $\nabla \cdot \vec{j} = 0$, where $\vec{j} = \vec{\sigma} \vec{E}$. Here, $\vec{\sigma}$ is the local conductivity tensor (with components $\sigma_{xx} = -\sigma_{xy} = -\sigma_{yx}$ and $\sigma = \sigma_{xx} = \sigma_{yy} = \sigma_{zz}$). By combining the two conditions to yield $\nabla \cdot \phi \vec{\nabla} \phi = 0$, where $\phi$ denotes the electrical potential, and discretizing the model, we have numerically solved the problem by allocating on each discretized node a local $\vec{\sigma}$ in accordance with the prior-calculated distributions. The solution of the classical network problem yields the macroscopic Hall resistivity $\rho_H = 1/\rho^\text{macro}$ and the macroscopic resistivity $\rho = 1/\rho^\text{macro}$. To compare the predictions of the theory with the experiment, we eliminate the concentration $x$ by cross-plotting the Hall coefficient as a function of $\rho$ in Fig. 3. Excellent quantitative agreement is seen. The GHE can therefore be heuristically understood as follows. Since $R_0 \sim 1/\rho_H \sim 1/(n)ec$, a symmetric distribution in local Hall conductivities effectively implies an overall decrease in the averaged charge carrier density and, hence, an increase in the Hall coefficient, because $R_0 \sim 1/(n)ec$, where $(n) \sim 0$ denotes the average charge carrier density. This also gives a plausible explanation to the temperature independence of the observed GHE up to 50 K, because the temperature variation is associated with the magnitude of the dephasing length. While the distribution of the local Hall conductivity can vary with the size of coherence region as defined by the dephasing length, the mean of the distribution $(\sim(n))$, which depends on the degree of asymmetry in the distribution, may remain relatively robust up to a point. Moreover, the interference component increases in importance as $x$ approaches the quantum percolation threshold, which is the point at which the interference-induced wave function localization occurs (at the Fermi level). Below the quantum percolation threshold, interference has the effect of changing the localization length. As $L_x$ is only a few feature sizes in our case, the localization length variation is not expected to be significantly perceivable on that scale, and a plateau in Hall resistivity results. As the quantum percolation threshold is above the classical percolation threshold, it follows that the GHE does not “see” the classical $x$, but rather peaks at or close to the quantum percolation threshold. Local interference is also expected to affect the resistivity by modifying the two-component composite picture to one that has a distribution of local conductivities. Moreover, this distribution is expected to have an increased number of small conductivities due to the localizing nature of the interference effect. Nonuniversal critical behavior may therefore result.

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FIG. 3. The Hall coefficient cross-plotted as a function of $\rho_{xx}$. Experimental data are shown by solid circles and the theory points are indicated by crosses. The theory and experimental data are normalized at one point—the peak value of the Hall coefficient and its associated (longitudinal) resistivity. Excellent quantitative agreement is seen. Inset: Distributions of local Hall conductivities at two metal concentrations, calculated from the $10 \times 10 \times 10$ quantum percolation model through the Kubo formula. The distribution is noted to be nearly symmetrical for the concentration close to the percolation threshold. Here $x_c = 0.31$ is the theoretical model.

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[7] It is common in granular systems to see insulating temperature variation at low temperatures and metallic behavior at high temperatures, for metal concentrations that fall between the quantum and classical percolation thresholds. This is due to an increased local quantum interference effect at low temperatures, associated with the increased dephasing length. Thus, at $T = 0$ K, the conductivity threshold should be at the quantum percolation threshold, whereas, at $T = \infty$, the conductivity threshold should be at the classical, or geometric (connectivity) threshold.

[18] If the grain size is large (such as in the annealed sample), then the distribution would be bimodal in character (either metal or insulator), and classical behavior prevails.