Growth of (Ge,Mn) nanocolumns on GaAs(100): the role of morphology and co-doping on magnetotransport

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Changing the morphology of the growing surface and the nature of residual impurities in (Ge,Mn) layers - by using different substrates - dramatically changes the morphology of the ferromagnetic Mn-rich inclusions and the magnetotransport properties. We obtained p-type layers with nanocolumns, either parallel or entangled, and n-type layers with spherical clusters. Holes exhibit an anomalous Hall effect, and electrons exhibit a tunneling magnetoresistance, both with a clear dependence on the magnetization of the Mn-rich inclusions; holes exhibit orbital MR, and electrons show only the normal Hall effect, and an additional component of magnetoresistance due to weak localization, all three being independent of the magnetic state of the Mn rich inclusions. Identified mechanisms point to the position of the Fermi level of the Mn-rich material with respect to the valence band of germanium as a crucial parameter in such hybrid layers.

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Research on ferromagnetic semiconductors triggered enormous activity due to their potential use in spintronics [1, 2]. Up to now, efforts have mainly focused on diluted magnetic semiconductors (DMS) in which magnetic atoms randomly substitute the host matrix atoms [3]. Their magnetic properties can be manipulated by electric fields making them suitable materials for spintronic applications provided that they can be made ferromagnetic above room temperature. However DMS based on II-VI and III-V semiconductors still exhibit very low magnetic values above room temperature. However DMS based on DMS based on II-VI and III-V semiconductors still exhibit very low magnetic above room temperature. However DMS based on II-VI and III-V semiconductors still exhibit very low magnetic above room temperature. Therefore, research on DMS is continuing, and DMS remains an active research area.

Many groups have reported $T_C$ values well above room temperature, along with remarkable magneto-transport and magnetooptical properties, in semiconductors doped with magnetic transition metals (TM). It is now admitted that these properties may be attributed to TM-rich areas resulting from spinodal decomposition [4]. Such features have been theoretically predicted [5] and reported in (Ge,Mn) [6, 7, 8, 9], and in Cr and Fe-doped GaN [10, 11] or ZnTe [12]. In this field of intense materials research, goals are now: (i) controlling spinodal decomposition to reproducing stabilize high-$T_C$ TM-rich areas and tailor desirable magnetic properties, and (ii) enhancing the coupling with carriers to give rise to strong magneto-resistance (MR) or anomalous Hall effect (AHE).

In this paper, we demonstrate the fine control of spinodal decomposition in (Ge,Mn) films grown on GaAs(001) substrates. We focus on (Ge,Mn) because it is compatible with mainstream silicon technology, and spinodal decomposition leads to high $T_C$ values in layers grown on Ge substrate. Growing (Ge,Mn) films on GaAs(001) semi-insulating ($\rho > 10^7 \Omega\text{cm}$) substrates makes in-plane transport measurements easier, and constitutes a first step towards spin injection from (Ge,Mn) into a GaAs-based spin-LED [13]. Using different surface preparations, we clearly identify the role of surface morphology and the role of impurity diffusion from the substrate (either Ga or As atoms), on the nanocolumns growth, on one hand, and on the electrical properties, on the other hand. We thus address the major issue of the influence of co-doping (either n-type or p-type) on spinodal decomposition in group IV magnetic semiconductors, demonstrating a major influence on the shape of the Mn-rich precipitates. We also provide new hints to control and optimize magneto-transport properties of the (Ge,Mn) films. We show that AHE and MR are not optimized simultaneously, and we propose a general picture based on the electrical doping of the matrix and on the position of Fermi level in the precipitates with respect to the valence band of Ge.

(Ge,Mn) films were grown by low temperature Molecular Beam Epitaxy (MBE), using growth conditions as described in Ref. [6], with the substrate temperature $T_g = 100^\circ\text{C}$ and deposition rate $\sim 0.2 \text{ A.s}^{-1}$. We have used two different methods to prepare the GaAs surface. In the first one, the native oxide was thermally desorbed from an epiready substrate, by raising the substrate temperature up to almost 600$^\circ$C. The (Ge,Mn) layers was grown directly on the resulting Ga-rich GaAs surface, which was rough as observed by RHEED. Such samples will be labelled Ga-(Ge,Mn). In the second case, As-(Ge,Mn) samples, a thin undoped GaAs buffer layer was grown first in a separate III-V system, protected with an amorphous As capping, and transferred in air to the IV-IV MBE machine. Desorbing the As capping layer at 200$^\circ$C results in a very flat, (2x4) reconstructed, As-rich surface. Samples grown on Ge substrates, labelled Ge-(Ge,Mn), constitute our reference samples.

Magnetization was measured using a Superconducting QUantum Interference Device (SQUID). Magnetotrans-
port properties (magnetoresistance and Hall effect) were investigated using Hall bars defined by optical lithography, aligned along a <110> direction, of width 20 µm, with voltage probes separated by 140 µm.

By contrast, in Fig. 1d, the As-(Ge,Mn) layers feature randomly distributed Mn-rich precipitates. In addition, a few Ge$_2$Mn$_5$ clusters already start to form, as evidenced by their typical Moiré contrast (see inset of Fig. 2b). The same random distribution of nanoclusters is observed when increasing the nominal Mn content from 2% to 6% and 10% at $T_c=10^6$ C, but for an increase of the average precipitate density and length along the growth direction. This suggests that the decomposition is of 3D character and mostly driven by nucleation.

Secondary Ion Mass Spectroscopy (SIMS, not shown) performed on the As-(Ge,Mn) samples evidences an Mn-rich topmost layer, extending over 3±1 nm below the sample surface and containing up to 6×10\(^{19}\) As cm\(^{-3}\), i.e., an integrated amount of almost 1 ML As. This is a consequence of the segregation of As atoms, initially present at the GaAs surface, during the growth of Ge, with a well-known surfactant effect [14]. Accordingly, as described below, the As-(Ge,Mn) films appear as n-doped: As atoms are shallow donors in Ge, and in this topmost layer they compensate p-type doping by substitutional Mn.

The presence of As near the surface of the growing layer offers a possible explanation for this change of character of the spinodal decomposition, from 2D to 3D. According to Ref. [15], Mn atoms are incorporated into germanium in a subsurface interstitial position, and further diffuse within the growth plane: this offers a mechanism for 2D spinodal decomposition [5]. Codoping with As changes the charge state of Mn atoms, thus reducing Coulomb repulsion and enhancing the effect of attractive Mn-Mn pair interaction, making the nucleation of Mn-rich precipitates easier [4]. In addition to that mechanism, the presence of donors like As is expected to displace the equilibrium between interstitial Mn (another donor) and substitutional Mn (an acceptor), enhancing the amount of substitutional Mn (which form nucleation centers for further Mn aggregation [16]), and reducing the amount of interstitial Mn (thus decreasing the incorporation into already existing clusters). These different mechanisms induced by the presence of As conspire to favor a growth process dominated by nucleation, contributing to make the spinodal decomposition 3D.

A complete study of the magnetic properties will be published elsewhere [17]. All samples exhibit two [case of Ga-(Ge,Mn), Fig. 2a] or three [case of As-(Ge,Mn), Fig. 2b] magnetic phases, as evidenced from the temperature dependence of the saturation magnetization $M_s$ (inset of Fig. 2) and the remanent magnetization $M_r$, and the ZFC-FC curves. They exhibit: (i) a strong paramagnetic signal with a 1/T temperature dependence at low temperature, attributed to Mn atoms diluted in the Ge matrix, and well fitted using a 3/2-Brillouin function [18]; (ii) a contribution attributed to the superparamagnetic Mn-rich nanoclusters or precipitates, with finite $T_C$ and blocking temperature $T_B$; (iii) in As-(Ge,Mn) only,
a contribution from Ge$_3$Mn$_5$ clusters with a broad range of blocking temperatures. A detailed analysis [17] of the Ga-(Ge,Mn) sample in Fig. 2a shows that 40±6% of the magnetic moments are in nanocolumns, with $T_C = 150$ K and $T_B = 15\pm5$ K, 1.0±0.1 $\mu_B$/Mn, and an average magnetic moment of a nanocluster 520±50 $\mu_B$. For the As-(Ge,Mn) sample in Fig. 2b, we found that 52±3% of the magnetic moments are in the matrix, and 22±2% in the Mn-rich precipitates with $T_C = 50$ K, $T_B = 15\pm5$ K, 1.2±0.2 $\mu_B$/Mn, and ≈100±20 $\mu_B$ per precipitate.

Then the magnetotransport properties can be understood as follows: (i) as the nanocolumns configuration is well below the percolation threshold, holes have to propagate through the germanium matrix; that makes the basis of the conductivity; (ii) however, the electric field pattern drags the holes through the nanocolumns, where the conductivity is higher; applying a magnetic field suppresses this effect, creating the geometrically enhanced orbital MR, or Extraordinary MR (EMR)[21] which we observed to be strong in Ge-(Ge,Mn) [6]; (iii) finally, the absence of Schottky barrier enhances the interaction of holes with Mn atoms in the nanocolumns, thus allowing a spin polarization and a strong AHE to appear [6].

In the Ga-(Ge,Mn) samples, we still observe the EMR (Fig. 3b), with the same temperature dependence as in Ref. [6], but much weaker (although much higher than classical Lorentz MR). This is readily explained by considering the dependence of EMR on the carrier mobility (it scales as $\mu^2$): as seen in Fig. 3a, the zero-field resistivity $\rho_0$ in Fig. 3a is of the order of $10^{-2}$ $\Omega$ cm, so that the mobility is lower than in Ge-(Ge,Mn) by more than one order of magnitude, possibly due to the higher disorder and induced defects. Finally, at very low temperature, negative MR is observed, which may be due to spin disorder scattering [22] on Mn atoms diluted in the Ge matrix (as typical in metallic DMS), or to GMR on the Mn-rich nanocolumns. In both cases, this effect is expected to be very weak; in addition the spin diffusion length of holes is very short due to spin-orbit coupling, making GMR on Mn-rich nanocolumns unlikely.

AHE in Ga-(Ge,Mn) nicely matches the magnetization of the nanocolumns (Fig. 3c-d). Again, the effect is weaker than in Ge-(Ge,Mn), by almost two orders of magnitude. This is expected from the lower mobility: it was pointed out in Ref. [23] that scattering on impurities such as Ga atoms (SIMS measurements have indeed shown that Ga out-diffused from the GaAs substrate) partly suppresses the effect of skew scattering.

As-(Ge,Mn) films exhibit metallic n-type conductivity. This is clearly due to the presence of As donors in the topmost layer. Hence magnetotransport essentially measures the properties of this 3 nm thick layer. We observe no AHE, as expected since spin-orbit scattering is small for electrons in the conduction band of germanium, and also because the same assumptions as above (Fermi level of precipitates lying below the top of the valence band) creates a high Schottky barrier for electrons.

MR is highly anisotropic (Fig. 4a): we show now that this is due to 2D weak localization in the As-doped layer, which vanishes when the field is applied in-plane ($\theta = 0$). The isotropic part (Fig. 4c-d) will be analyzed later on.

The MR of non-interacting electrons in the 2D weak localization regime is $\Delta\rho/\rho \approx -\Delta\sigma/\sigma = -A f_2(4\pi\mu_0 H\sin(\theta)L_0/h)$ [24], where $A = e^2/2\pi^2hL_0$ is the phase relaxation length, and the function $f_2(x)$ is defined in [24]. Here we neglect
the effect of spin-orbit coupling and the anisotropy of $L_\phi$ [17]. Fits in Fig. 4a were obtained with only two adjustable parameters: $A = 4\%$, close to the value $A = (6 \pm 2)\%$ calculated using the experimental value of the 2D conductivity at $H = 0$, and $L_\phi = 11.5$ nm, large enough with respect to the thickness of the conducting layer to justify the use of the 2D regime of weak localization. Moreover, fits of the temperature dependence of the anisotropic MR (Fig. 4b) were obtained by simply writing $L_\phi = \sqrt{D\tau_\phi}$, using the temperature dependence of the diffusion coefficient $D$ for an n-doped degenerate semiconductor [25], and a temperature dependence of the phase relaxation time $\tau_\phi \propto T^{-\alpha}$ with $\alpha \approx 1.7$, similar to that obtained in [26] for Ge:Sb ($\alpha=1.5$) and in [27] for Si-MOSFETs ($\alpha=1.6$), and currently attributed to both electron-electron and electron-phonon collisions.

Turning back to the isotropic MR, Fig. 4c-d, it contains a negative contribution, which features two maxima at the coercive field of Mn-rich precipitates, and vanishes above 50 K as does their magnetization. Hence, we tentatively ascribe it to tunneling MR (TMR) through the precipitates and the Schottky barriers formed around them. By analogy with spin injection from a ferromagnetic metal to a semiconductor [28], efficient spin injection from the precipitate to the matrix requires an interface resistance provided by the Schottky barrier. This barrier must be high enough to prevent full spin relaxation inside the precipitate but reasonably transparent to allow tunnel MR to occur.

To summarize, we have shown that surface morphology and co-doping have major influence on spinodal decomposition in (Ge,Mn) films grown on GaAs(001) substrates. For films grown on Ga-rich rough surfaces, we recovered 2D spinodal decomposition with bent nanocolumns. Electrical properties are similar to what we obtained on Ge(001) substrates except that the presence of defects in the films leads to weaker positive MR and AHE. For films grown on As-rich flat surfaces, 3D spinodal decomposition is observed due to As co-doping and magnetotransport is dominated by TMR and weak localization, AHE is negligible. These results are consistent with the assumption that the Fermi level of Mn-rich precipitates lies in the valence band of the Ge matrix.

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