Elaboration of Nanomagnet Arrays: Organization and Magnetic Properties of Mass-Selected FePt Nanoparticles Deposited on Epitaxially Grown Graphene on Ir(111)

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The moiré pattern created by the epitaxy of a graphene sheet on an iridium substrate can be used as a template for the growth of 2D atomic or cluster arrays. We observed for the first time a coherent organization of hard magnetic preformed FePt nanoparticles on the 2D lattice of graphene on Ir(111). Nanoparticles of 2 nm diameter have been mass selected in a gas phase and deposited with low energy on the hexagonal moiré pattern. Their morphology and organization have been investigated using grazing incidence small angle x-ray scattering, while their magnetic properties have been studied by x-ray magnetic circular dichroism, both pointing to a FePt cluster-graphene surface specific interaction. The spatial coherence of the nanoparticles is preserved upon annealing up to 700 °C where the hard magnetic phase of FePt is obtained.

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Nanoparticles are intensively studied for optical [1-4], catalytic [5–7], magnetic [8–12], and storage applications [13–18]. FePt nanoparticles are interesting candidates for ultrahigh density storage applications due to their extremely high magnetocrystalline anisotropy when chemically ordered in the $L1_0$ phase [19,20]. Another requirement for such applications, as well as for fundamental studies, is to organize magnetic nanoparticles in a 2D array. A great effort is devoted to the bottom-up elaboration of periodic patterned arrays of nanoparticles [21-25]. In this domain, the moiré pattern appearing from the epitaxy of graphene (q) on a transition metal (q/TM) has been used successfully to organize by physical vapor deposition (PVD) on $g/\mathrm{Ru}(0001)$, $g/\mathrm{Cu}(111)$, or $g/\mathrm{Ir}(111)$ a wide variety of pure metallic nanoparticles [21,26-28]. However, organized islands of Fe cannot be grown by PVD on g/Ir(111) [21], so that it is not possible to obtain a superlattice of FePt alloy nanoparticles, even using Pt seeding [26,29].

The MSLECBD (mass-selected low energy cluster beam deposition) technique [30–32] may be used to deposit preformed alloy nanoparticles having a chosen chemical composition. MSLECBD offers opportunities not accessible by PVD: the cluster size can be controlled and is independent of the surface coverage. Recent studies point out the possibility to organize pure nanoclusters on a graphene moiré using soft-landed clusters of Pt on g/Ir(111) or Pd on g/Ru(0001) [33,34], where one of the three high-symmetry sites of the surface is more favorable for adsorption. The same approach could enable to order metallic alloys on graphene, and especially hard-magnetic ones ($L1_0$ FePt)

with a strong uniaxial magnetization. Such a possibility would rely on a preferential adsorption of deposited particles on specific sites of the moiré superlattice. In order to study the effect of the graphene substrate on FePt particle organization and magnetic properties, we consider here diluted samples, thus avoiding interparticle interactions. In this Letter, we report on the low-energy deposition of mass-selected FePt clusters (around 2 nm diameter) on the q/Ir(111) moiré superlattice, and their characterization by GISAXS (grazing incidence small angle x-ray scattering) and XMCD (x-ray magnetic circular dichroism). These two complementary experimental techniques allow us to probe, respectively, the organization and magnetic properties (and hence to detect the chemical ordering phase transition upon annealing) of the FePt nanoparticles. A clear tendency to preferential pining and coherent organization is found, as well as specific properties due to the cluster-surface interaction.

The substrate of graphene on a Ir(111) monocrystal has been prepared using chemical vapor deposition (CVD) following a procedure well known in the literature [35]. Chemically disordered FePt clusters of 2 nm diameter, synthesized by the MSLECBD technique, are deposited on the g/Ir(111) surface in a soft-landing regime at various temperatures (see Supplemental Material [36]) with a density of 3×10^4 cluster/ μ m². With this surface coverage and nanoparticle size, a compromise is obtained between a low proportion of multimers [37] and a sufficient signal in x-ray experiments. An amorphous carbon capping layer (a few nanometers thick) is grown after cluster deposition,



FIG. 1. (a) GISAXS intensities (at room temperature, after deposition at 150 °C) along the $\langle 100 \rangle$, $\langle 010 \rangle$, and $\langle 1-10 \rangle$ directions of the hexagonal moiré lattice. The presence of a correlation peak (highlighted in red) in those directions reflects the hexagonal organization of FePt nanoparticles on the moiré lattice. Out of azimuth, no correlation is visible. (b) GISAXS intensity in the $\langle 100 \rangle$ direction at 700 °C. (c) GISAXS intensity profile (fixed α_f) of the correlation peak for three different temperatures with their respective fits represented with the black line. The curves have been shifted for clarity. The fraction f_{order} of particles coherently located on the moiré lattice remains constant around 35% ±3%.

using carbon wire sublimation under vacuum, to avoid any contamination and sintering.

First, the organization and the nanoparticles morphology have been investigated by GISAXS on the BM32 beam line at the European Synchrotron Radiation Facility (ESRF), with an incident x-ray energy of 11 keV and a critical angle of 0.42°. Measurements have been performed on a capped sample, after FePt cluster deposition at 150°C. Once in the beam line UHV chamber, the sample is heated up from room temperature to 700°C.

GISAXS measurements provide information on the lateral correlation, size, and shape of the nanoparticles [38–40]. The central intensity reflects the size and shape of the nanoparticles while the correlation peak in Fig. 1(a) (highlighted in red) corresponds to the coherent organization of the nanoparticles across the sample. The correlation peak appears only in preferential directions (remarkably, in the $\langle 100 \rangle$, $\langle 010 \rangle$, and $\langle 1-10 \rangle$ directions with the *h*, *k*, *l* = 0 surface hexagonal lattice notation), and in the vicinity of $2\theta = 51$ mrad which corresponds to the moiré lattice parameter of 2.53 nm [34,35,41]. In other directions (out of azimuth), only the central intensity remains. This indicates that, at room temperature, a 2D hexagonal organization of

FePt clusters on the moiré lattice, similar to size-selected Pt clusters on q/Ir(111) [34] is obtained, and excludes a simple preferential nearest-neighbor distance. This demonstrates that clusters can diffuse on the surface (since they are initially randomly deposited [42,43]) before being pinned on specific sites reflecting the moiré periodicity. The cluster-surface interaction is thus favorable for self-organization and may be described as a chemisorption involving π -d hybridization [44] and local rehybridization from sp^2 carbon to diamondlike $s p^3$ [33]. The sample was then progressively annealed up to 700 °C, high enough to promote the FePt $L1_0$ chemical ordering [45,46]. As seen in Fig. 1(b), the correlation peak is less intense but a coherent organization still remains. Qualitatively, one can detect moderate changes in the form factor of the particles: the GISAXS signal is more concentrated and intense at low θ angle (near the specular beam).

A quantitative analysis of the GISAXS patterns (using inplane and out-of-plane line cuts) gives a precise estimation of the form factor (size and shape) and the lateral correlation of the cluster superlattice. Furthermore, it is possible to link the correlation peak intensity to the proportion f_{order} (supposed to be homogeneous over the entire sample) of nanoparticles coherently pinned on the moiré lattice [34]. The total GISAXS intensity can indeed be separated in two contributions: one coming from particles with random locations [47] on the surface (i.e., with no constructive interference and thus no correlation peak) and the other from organized particles on the moiré lattice.

The incident cluster size deduced from transmission electron microscopy (TEM) observations can be modeled with a Gaussian centered on the mean equivalent diameter $D_{\text{TEM}} = 1.9 \text{ nm}$ (see Supplemental Material [36]). First, we want to determine if the incident particle size is preserved on the surface, since diffusion-coalescence processes could produce larger particles. A first quick analysis of the out-of-azimuth GISAXS cuts, by fitting with a simple Gaussian size distribution, indicates that the diameter is globally conserved (mean diameter around 1.9 nm), however, with a presence of some larger particles (relative dispersion of 50%). In fact, it is not surprising to find a particle size distribution different from the incident monomers because even without diffusion, there is a probability (which depends on the cluster density) that a cluster lands on another one and forms a dimer (or multimers) [48]. If clusters diffuse on the surface, they can form additional multimers. Therefore, we can use a better description to analyze GISAXS measurements using two Gaussians: a first one for monomers (main contribution) and a second one corresponding to dimers and multimers. The respective proportion of monomers (X) and dimers (1-X) can be inferred from a best fit, together with the monomers mean size and the dimers relative diameter dispersion [49]. Moreover the particle shape is modeled by truncated spheres with an adjustable H/D ratio, where H is the height and D the in-plane diameter. A value H/D = 0.74, coherent with the wetting parameter found for Pt clusters on g/Ir(111) [34] is obtained. The monomers mean size (spherical equivalent diameter [50]) is found to be $D_{eq} =$ 1.9 nm which is in full agreement with the TEM size histogram while the monomers proportion amounts to 73%. Most of FePt particles on the surface have thus kept their incident size which shows that diffusion, during cluster deposition, is limited but still present. Note that diffusion is indeed required to explain the observed coherent organization and the monomers proportion, slightly lower than expected with a random pinning. Once the size and shape parameters have been determined from the optimal fit of out-of-azimuth GISAXS patterns (see Supplemental Material [36], S3), the proportion f_{order} of particles located on moiré sites is determined from a GISAXS fit along the $\langle 100 \rangle$ direction of the moiré lattice. Here, the form factors are fixed while the interference function is adjusted (it is directly related to the particle organization, i.e., to the moiré crystallographic parameter and the proportion f_{order} [39]).

The intensity of the correlation peak (which depends both on the form factor and on the interference function) and calculated fits are pictured on Fig. 1(c) for three different temperatures. At room temperature $f_{order} = 35\% \pm 3\%$, which is slightly lower but comparable to Pt particles deposited on q/Ir(111) where a value of around 50% is found [34]. This difference is probably due to the larger size of nanoparticles and/or to the alloy nature of FePt clusters. The evolution of the form factor between RT and annealing at 300 °C slightly influences the intensity of the correlation peak but f_{order} is in fact almost the same. This is coherent with the fact that at 300 °C the temperature is relatively low and does not induce any important changes in the nanoparticles, graphene, or amorphous carbon capping layer morphology. Remarkably, at 700 °C the organization is preserved and remains visible, with $f_{\text{order}} = 38\% \pm 3\%$. The nanoparticle shape is unchanged, however, with a decrease of the monomers proportion down to 65%. At such a high temperature, atomic displacements can occur allowing some particles to diffuse over a small distance.

The fact that $f_{\text{order}} < 100\%$ indicates that two types of pinning sites coexist: coherent location(s) within the moiré cell and other positions (incoherent contribution). f_{order} is then linked to the relative density of the different possible pinning sites. However, a full description of surface energy potential together with diffusion and coalescence processes is required to infer their density from the f_{order} value, which is out of the scope of this work. Nevertheless, the estimated density of defects in the moiré lattice (moiré domains have sizes in the micrometer range) and/or graphene sheet (wrinkles, step edges...) [35,51,52] is too low to account for the proportion of FePt nanoparticles randomly located. This means that clusters landing in some places inside a moiré cell have a non-negligible probability to end up at many various locations (this supposes some kind of rough potential energy landscape, more complex than with only three high-symmetry pining sites), while other landing areas enable the FePt particles to reach specific pinning sites. In order to reach ultradense (>1 Tbit/in²) magnetic bit arrays, it is important to understand the pinning mechanisms of preformed clusters as well as the surface energy potential, which governs the diffusion of the nanoparticles on the moiré surface. This may be achieved through theoretical studies or additional experiments (such as STM investigations).

From the GISAXS study, we can tell that the incident particles have been preserved but the surface has a clear influence on their behavior: the random deposition produces, in the end, a partially organized array of particles, even at 700 °C. This cluster-surface interaction may as well have an impact on the magnetic properties, which can be investigated by XMCD. Moreover, if chemical ordering has occurred upon annealing, it should be visible through an evolution of the magnetic properties.

XMCD measurements have been carried out at the DEIMOS beam line at the SOLEIL Synchrotron in Paris-Saclay [53]. By using circularly polarized x rays, the Fe $L_{2,3}$ edges have been probed using total electron yield (fluorescence mode for the hysteresis loops) for several incidence angles, from normal to the sample (0°) to 60° from normal.



FIG. 2. Top (a) and (b) are, respectively, before and after annealing XAS spectrum at the Fe $L_{3,2}$ edges taken at 4 *K* for light circularly polarized left (+) and right (-) with the isotropic XAS signal shown in dotted line. Both averaged XAS signals have been normalized for a direct comparison of the XMCD spectrum. The XMCD spectrum [bottom (a) and (b), respectively before and after annealing] is the difference between the two XAS spectra recorded with opposite orientation of the magnetic field and the x-ray polarization (respectively, blue and red for left and right helicity). The dotted lines are the integral of the XMCD spectrum to highlight the modification of the intensity as well as the evolution of the m_L/m_S ratio. (c) Hysteresis loops taken at 4 *K* for two incidence angles 0° (normal to the surface, in red) and 60° from the normal (in blue), before annealing (top) and after annealing (bottom). The fits are pictured in green for both phase of the nanoparticles.

Those measurements were investigated from room temperature to low temperature (4 K), before and after annealing of the sample up to 700 °C. The sample used for the XMCD characterization, (different from the GISAXS measurements) has been prepared in the same conditions with a higher deposition temperature of 300 °C (the organization is identical, see Supplemental Material [36]).

Figure 2(c) shows the hysteresis loops at low temperature (4 K, i.e., in the blocked regime) before (top) and after annealing at 700 °C (bottom), for two different x-ray incidence angles. The hysteresis loops are completely isotropic which shows that there is neither demagnetizing factor effect, nor any preferential orientation of the nanoparticles. Interactions among the 2D layer of nanoparticles (as for a thin magnetic film) or interface anisotropy with the graphene sheet would have introduced an orientation dependence (anisotropy) of the hysteresis loops. Since none has been observed, this is an other evidence that FePt nanoparticles have kept their individuality even after annealing. Note that the magnetic remanence at 4 K is around 50% of the saturation, which is consistent with the model of isolated nanoparticles having a random distribution of their easy axis.

The effect of annealing on the opening of the loops is clear: the coercive field H_C is increasing from 122 to 650 mT, reflecting an increase of the magnetic anisotropy constant. From the room temperature magnetization curves (in the superparamagnetic regime, see Supplemental Material [36]), we observe that, as expected, the magnetic size distribution is almost unchanged upon annealing and is in full agreement with the geometrical size distribution deduced from GISAXS. To go further, a theoretical model based on a combined Stoner-Wohlfarth and Néel relaxation description, has been used to fit the hysteresis loops, taking into account the magnetic size distribution [54–58]. A biaxial description has been adopted for the magnetic anisotropy [59], in order to reflect the non ideal morphology of the nanoparticles. Before annealing, the mean anisotropy constant is $K_1 = 300 \text{ kJ/m}^3$ with a relative dispersion of 40% (we use a Gaussian distribution, which reflects the variation of the anisotropy constant among the particles) and a biaxial ratio of $K_2/K_1 = 1.2$.

After annealing, the mean K_1 is 1.4 MJ/m³ with an increased relative dispersion of 70% and an unchanged biaxial ratio. This large anisotropy dispersion must be due to a chemical order distribution among the FePt particles, in addition to the existence of a variety of geometries (and possible defects) [45]. The two branches of the hysteresis loop only merge at very high field which implies that some particles in the assembly have an anisotropy field higher than 3 T. We estimate the highest anisotropy constant around 3 MJ/m³ which is close to the bulk value for $L1_0$ FePt [60,61]. The value found for the magnetic anisotropy constant is very large for 2 nm FePt nanoparticles, indicating a transition towards the chemically ordered $L1_0$ phase, already observed for particles embedded in a carbon matrix (with possible defects and multiply twinned chemically ordered particles) [45].

Before annealing, the x-ray absorption spectroscopy (XAS) and XMCD measurements depicted on Fig. 2(a), show well-defined Fe- $L_{2,3}$ absorption edges with no sign of oxidation [62,63] and a clear magnetic Fe signature. All the spectra have been acquired at the saturation regime, respectively 2 and 5 T before and after annealing. The maximum intensity of the XMCD spectrum decreases for annealed FePt nanoparticles [Fig. 2(b)] reflecting a reduced magnetic moment as theoretically expected for $L1_0$ FePt compared to the A1 phase [64,65].

The Fe magnetic moments (spin and orbital moments, respectively m_S and m_L) have been determined by applying the sum rules [66–68] with a number of holes $N_h = 3.705$ [64,69]. The orbital moment is 0.11 ± 0.01 before annealing and decreases to 0.06 ± 0.01 after annealing. The spin moment is 3.0 ± 0.2 before annealing and 2.5 ± 0.2 after annealing. This corresponds to a m_L/m_S ratio of 0.037 ± 0.005 before annealing and 0.024 ± 0.005 after annealing.

The Fe magnetic moment for the disordered A1 FePt nanoparticles is close to the bulk [70]. Such a high value has never been reported for disordered FePt nanoparticles. Nanoparticles are very sensitive to their environment because of the high surface-to-volume ratio, so that an influence of the graphene on the orbital and spin moments cannot be excluded [71]. After annealing, the spin moment is lower and is similar to the value found in the literature for FePt nanoparticles [64,72]. However, m_L is surprisingly low compared to previous studies [64,73], including our results on 3 nm FePt nanoparticles embedded in carbon matrix (0.18 $\mu_{\rm B}/{\rm at}$) [72]. m_L/m_S is very close to the value of $L1_0$ bulk (around 2.5% [65,69,74–76]) and smaller than reported for thin films [77–79], which is unexpected, since it is usually assumed that the orbital moment in nanoobjects is higher than in bulk due to the broken symmetry. Thus it seems that the presence of graphene has a strong effect on Fe orbital moment in the chemically ordered nanoparticles through modification of the electronic structure. The particle flattening and pining, as revealed by GISAXS, imply that a sizable interaction exist between the FePt clusters and the graphene surface. Charge transfers can occur between the nanoparticles and the graphene and it is very likely to observe a mixing of electronic states as it is observed for clusters and adatoms [80-83]. Magnetic anisotropy energy and orbital moment are closely related: the origin of the MAE comes from the anisotropy of the density of state resulting from the spin-orbit interactions for two distinct directions of the system. The equations derived by Bruno [84] state that for a sizable contribution to the magnetic anisotropy energy, strong spin-orbit coupling and large orbital moment are needed, which is especially true for the Pt atoms in a FePt alloy. Here, we find that the MAE is not directly proportional to the Fe orbital moment in FePt nanoparticles, in full agreement with theoretical studies [69,75,76,78,85,86], even if the question may still be debated [77].

In this study, we have reported the organization of sizeselected FePt magnetic nanoparticles on a moiré lattice from the epitaxy of graphene on an iridium monocrystal. The organization has been studied using GISAXS measurements and simulations. We found that after deposition, 38% of the nanoparticles are coherently pinned on the hexagonal moiré lattice. Further theoretical investigation is needed to understand the adsorption energy landscape and the resulting cluster-surface interaction responsible for the organization. A subsequent annealing leads to an impressive increase of the magnetic anisotropy (deduced from XMCD measurements), which can be ascribed to a $L1_0$ chemical ordering transition, while the FePt nanoparticles remain organized on the surface. For the first time, a system made of small FePt nanoparticles, chemically and spatially ordered, has been synthesized and characterized. Moreover, we have illustrated that the magnetic anisotropy and the Fe orbital magnetic moment can exhibit a complex relationship and a subtle dependence on interface effects. Future *ab initio* calculations may give more insight on the electronic and magnetic interactions between FePt nanoparticles and the *g*/Ir(111) substrate.

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