Growth modes of nanoparticle superlattice thin films

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Abstract: We report about the fabrication and characterization of nanoparticle thin film superlattices, which show two of the three thin film growth modes, i.e. Volmer-Weber and Frank van der Merwe growth. In addition three-dimensional mesocrystallite growth is evidenced. The formation into different systems is accomplished by tuning the particle-to-substrate interaction. The understanding of the mechanisms ruling nanoparticle self-assembly represents an important step toward the fabrication of novel materials with tailored optical, magnetic or electrical transport properties.

The advent of controlled thin film growth about seven decades ago revolutionized many areas of science and technology1-6. Examples are optical coatings1,2, magnetic layers and multilayers3,4 or semiconductor thin films5,6. Thin films enabled the development of important applications and lead to the discovery of novel effects which arise from the structure of materials and the presence of surfaces7,8 or interfaces between different layers9,10. In the early stage of research on thin films it soon became clear that it was imperative to understand the mechanisms which control and define the growth of thin films to achieve good control over these novel materials. Hence the huge effort of the scientific community to characterize, optimize and understand film growth. Evidently thin films are composed of atoms, which are their zero-dimensional building blocks. Extending this concept, nanoparticles (also termed 'nanocrystals') can also serve as zero-dimensional building blocks
which by self-assembly may form two-dimensional thin films or three-dimensional crystals (so-called 'nanoparticle superlattices'), analogous to atomic films and crystal lattices\textsuperscript{11,12,13,14,15,16,17,18,19,20,21}. Atomic thin film growth is understood to occur in form of three basic growth modes, which result from competing energy terms during the film deposition, i.e. Frank van der Merwe, Stranski-Krastanov, and Volmer-Weber growth\textsuperscript{8,22,23}. Various processes occur when atoms arrive at a substrate during thin film growth. That is adsorption, desorption, diffusion, finding or leaving of equilibrium positions. These processes occur simultaneously averaged over the ensemble of arriving atoms. Consequently, textbooks describe these processes in the framework of macroscopic thermodynamics\textsuperscript{8,22,23}.

An important quantity, e.g., is the equilibrium vapor pressure $p_e$ of atoms, being defined as the pressure, where condensation and evaporation of atoms at the substrate surface occurs at the same rate. Thin film growth is possible, when the pressure of the vapor near the substrate, $p$, is larger than the equilibrium vapor pressure $p_e$. In the case of a supersaturated vapor, a free energy difference exists between the vapor and the condensed atoms in the film. Consequently a chemical potential exists, which is responsible for further condensation of atoms into the film\textsuperscript{23}. This concept can also be transferred to nanoparticle (NP) films and superlattices. While NPs are in the solvent, the solvent constitutes the confining volume. Upon evaporation of the solvent the volume shrinks and leads to an increase of the NP 'vapor' pressure. Eventually NP film growth is possible.

As in the case of atoms, the interplay of various free energy terms determines the way on how the NP films will grow. These are in detail\textsuperscript{23}: an entropic contribution $E_T S$, an inter-particle energy term $E_p$, summing up all relevant types of interactions between NPs\textsuperscript{24}, and a NP-to-substrate interaction energy, $E_s$. A further important factor is the diffusion energy barrier, $E_d$, which can be overcome by 'thermal' energy $k_B T_s$. Hereby $T_s$ is a quantity comparable to a substrate 'temperature', which precise physical meaning still needs to be understood for NP
systems. Depending on the relative magnitudes of $E_d$ and $k_B T_s$ the NPs will either stay fixed at one place once they are attached to the substrate or move freely to seek energetically more favorable locations considering the two extreme cases.

From a comparison of these free energy terms one finds$^{22,23}$ that three different growth modes follow: in the case where NP-to-substrate energy, $E_s$, dominates a layer-by-layer growth is found, viz. the so-called *Frank-van-der-Merwe growth mode* (FvdM). This case is depicted in Fig. 1. Once a stable cluster (or 'nucleus') of NPs is formed, the following NPs prefer to attach at the periphery of the nucleus in contact with the substrate. Accordingly this leads to the advancement of planar film growth$^{23}$. Depending on the ratio of $E_d$ and $k_B T_s$ (i.e. the mobility of NPs) one will either obtain polycrystalline or single-crystalline superlattices. In the first case, the immobility of NPs leads to the nucleation of many independent superlattice crystallites, while, in the other case, the large mobility enables the NPs to seek equilibrium positions and hence promotes single-crystal growth.

![Figure 1: Schematic representation of the three known film growth modes$^{23}$. The spheres can be either atoms or NPs. When NPs approach a substrate, either adsorption, desorption or diffusion occurs. Depending on the free energy contributions one finds Frank-van-der-Merwe (FvdM), Volmer-Weber (VW), or Stranski-Krastanov (SK) growth.](image)

If, however, the NP-NP energy dominates, the NPs tend to attach on top of existing NPs and hence favor three-dimensional or island growth (Fig. 1). This mode is termed *Volmer-Weber*
growth (VW). Whether polycrystalline or single-crystalline islands arise also depends on the mobility as in the FvdM case.

The third growth mode constitutes in some sense a mixed situation, where layered and island growth coexist. It can be understood by a change of growth mode during self-assembly. As the first layer experiences a large interaction to the substrate the growth starts in a planar fashion. Once the substrate is covered by a monolayer of particles the ratio of interaction energies changes so that island growth is preferred. This mode is termed Stranski-Krastanov mode (SK) and is shown schematically in Fig. 1.

An alternative approach toward the distinction of growth modes is based on the analysis of 'surface tension' energies. By comparing various surface tension contribution one basically arrives at the same distinction of three basic growth modes 23.

The aim of the present article is to explore thin films and superlattices composed of iron oxide NPs. By varying the particle – substrate interaction energy we obtain two of the three growth modes in analogy to atomic thin films, viz. FvdM and VW growth. In addition we obtain three-dimensional growth comparable to seed-mediated bulk crystal growth. These growth modes are verified through a series of case studies reported in this article. The third film growth mode, SK growth, still evades experimental realization. Moreover, as increasingly revealed by other studies, 25 it turns out that the influence of the drying solvent plays a considerable additional role in the formation of NP superlattice films.

The first case to be considered is a monolayer thin film of NPs, spin-coated onto a Si substrate (sample 'Si'), the scanning electron microscopy (SEM) image of which is shown in Figure 2a. A complete monolayer of NPs is clearly visible, covering the entire substrate surface. The NPs self-assemble into a 2D hexagonal lattice, showing domain boundaries, vacancies and other structural defects analogous to those found in polycrystalline solids. Moreover, in some places it is possible to observe an incomplete second layer forming a
system of effectively 1.2 to 1.5 monolayers. The growth mode of this film clearly resembles that of layer-by-layer or FvdM growth.

Figure 2: SEM images of NPs spin-coated on (a) silicon substrate with native oxide (sample ‘Si’) (b) PMMA coated silicon (sample ‘PMMA_4P’) showing a completely different NP arrangement on the substrates. The corresponding GISAXS patterns are shown in (c) and (d) respectively.

Although SEM imaging provides us with a useful visual inspection of the sample, it alone is insufficient to address a comprehensive study of long-range correlations of NPs across the substrate. A more powerful characterization technique is needed, providing information on both the in-plane as well as the in-depth structural ordering. These requirements are met by the use of Grazing Incidence Small Angle X-ray Scattering (GISAXS), a surface sensitive x-ray scattering technique which provides electron density profiles statistically averaged over a
large lateral area. By the glancing incident and exit angles of the x-ray beams the surface sensitivity is enhanced.

The corresponding GISAXS pattern of NPs on Si, measured at an incident angle ($\alpha_i$) of 0.5°, is shown in Figure 2c, with the intensity plotted on a logarithmic scale coded in color scale shown on the right hand side. Two distinct features can be observed in the scattering pattern. In first place, the intensity is distributed in ring like patterns (although not continuous). At least three such rings can be seen in this pattern. Secondly, on top of the rings high intensity peaks, which modulate the ring intensity, can be observed. The peaks are extended along $q_y$ and $q_z$ directions and appear symmetrically on both sides of the $q_z$ axis along the $\pm q_y$ axis. These features represent the Fourier transform of the in- and out-of-plane electron density variations, which basically depend on two factors: the morphology (shape and size) of the NPs and the particle-particle correlations (NP ordering). The ring like pattern is a manifestation of the Fourier transform of the morphology, i.e. the form factor of the spherical NPs. Simultaneously, the intense Bragg peaks seen as streaks extending in the $q_z$ direction are manifestations of the scattering function, the Fourier transform of the particle-particle correlation function and hence indicate the presence of long-range ordering of the NPs. In other words, the high intensity Bragg peaks arise from long-range NP ordering in the GISAXS geometry, where the momentum transfer vector or the scattering vector (Q) is of the order of the reciprocal lattice vector ($\sim G$) of the NP in-plane lattice. The Bragg peaks are an indication of the crystal structure and can be used to determine the NP unit cell structure since in small angle geometry the $|Q|^{-1}$ correspond to few tens of nm, which is similar to the inter-particle distances for the NPs used in this study. The pattern shown in Figure 2c can therefore be assigned to a hexagonal close packed (HCP) lattice of lattice constant 20.38 nm, which is larger than the NPs average diameter of 18 nm as found from SEM images. The discrepancy can be explained by the fact that the SEM image is less sensitive to the organic oleic acid shell, because the element contrast in SEM scales with the atomic number of the elements.
Consequently, the SEM will rather reveal the diameter of the individual iron oxide NPs, while GISAXS probes correlation lengths, i.e. periodicities and hence the distances between NPs including the organic shells.

The second system proposed in this study is shown in Figure 2b and it was prepared by spin-coating nanoparticles on top of a Si substrate pre-coated with a few nm of polymethyl methacrylate (PMMA) with 4% solid contents (sample 'PMMA_4P'). In this case the NPs present a completely different ordering compared to the previous substrate. The NPs form islands (mostly disc like) of approximately 1μm in size. The inset shows one of the islands, where the NPs are arranged in a close packed structure. The corresponding GISAXS pattern is shown in Figure 2d. Unlike for NPs spin-coated onto Si, the GISAXS pattern of NPs on PMMA does not show any in-plane Bragg peaks indicating that the NPs within the islands are arranged in an amorphous fashion. The ring like structure only arises from the short-range ordering of the NPs (form factor) and does not show any preferred crystallite formation. This growth mode can clearly be assigned to the VW growth mode, where the NPs prefer to aggregate into individually isolated islands. A feature of particular interest is the agglomeration of densely packed NPs without any crystalline structure. It is clear from these observations that the surface chemical potential indeed influences the self-assembly process. This will become even more obvious from the observation of systems presented in the following paragraphs, where it will be shown that the surface interaction of the solvent and the substrate is the driving force behind these different growth modes.
In the following we consider two systems composed of NPs spin-coated on Si pre-coated with copolymer polymethyl methacrylate/ methacrylic acid (PMMA/MA) with 33 % solid contents (sample 'PMMA_33P') and on Si substrates with 300 nm of thermal SiO₂, shown in Figure 3a and 3b, respectively.

The use of a PMMA/MA that differs in composition from the one considered in the previous section brings forward important changes in the self-assembly process. The NPs on PMMA/MA 33% substrates form islands partially interconnected with each other to create a meander like pattern.

The inset in Figure 3a shows a magnified view of one of the islands, composed of several layers of NPs. Remarkably, the NP layers form terraces, which have preferred orientations of crystalline order, although the orientations of the crystallites inside the terraces are different
in between any two islands and are probably randomly distributed over the whole substrate.

The composition of the PMMA/MA layer therefore drastically modifies the surface interaction forces leading to different NP ordering mechanisms. The GISAXS pattern shown in Fig. 3c is once again representative of an HCP lattice. Note, while the average electron density variation resembles that of a hexagonally arranged monolayer, the formation of randomly oriented and shaped islands add a considerable amount of diffuse scattering around the Bragg peaks, which nevertheless still remain clearly visible and resolvable. For instance, the (30) peak can also be easily identified, confirming the high hexagonal ordering of the NPs inside the islands. Overall, this growth process can also be associated with a VW growth mode. Table 1 shows the comparison of intra-planar distance $d_{hk}$ measured from GISAXS patterns and the value calculated assuming a hexagonal lattice of lattice constant 20.38 nm for these NPs. One finds a relatively good correspondence between the experimental and calculated values from the various peak positions.

Table 1: Comparison of inter-planar distances measured from GISAXS patterns (2nd column) and calculated assuming a hexagonal lattice (3rd column).

<table>
<thead>
<tr>
<th>Miller indices (hk)</th>
<th>$d_{hk} = \frac{2\pi}{q_0 b}$ (nm)</th>
<th>$d_{hk} = \frac{a}{\sqrt{4(h^2+k^2)}}$ (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(10)</td>
<td>17.79 (± 0.023)</td>
<td>17.65</td>
</tr>
<tr>
<td>(11)</td>
<td>10.3 (± 0.01)</td>
<td>10.19</td>
</tr>
<tr>
<td>(20)</td>
<td>8.85 (± 0.008)</td>
<td>8.825</td>
</tr>
<tr>
<td>(21)</td>
<td>6.61 (± 0.004)</td>
<td>6.67</td>
</tr>
<tr>
<td>(30)</td>
<td>5.87 (± 0.005)</td>
<td>5.88</td>
</tr>
<tr>
<td>(22)</td>
<td>4.9 (± 0.006)</td>
<td>5.095</td>
</tr>
</tbody>
</table>

As mentioned in the early paragraphs, the formation of NP superstructures can be described by the competition of energy terms which account for the complete set of interactions.
between particle-particle, particle-solvent and particle-substrate\textsuperscript{23,24}. When preparing NP superstructures by spin-coating methods, as in the case of the system presented in this study, further considerations on the wetting ability of the solvent on the substrate should be addressed. As a matter of fact, the colloidal particles are dispensed in a solvent, in this case toluene, and the self-assembly process will also be affected by the solvent evaporation, solvent wetting and its viscosity. The last stage of spin-coating is characterized by evaporation and subsequent thinning of the solvent film\textsuperscript{27}.

The self-assembly achieved by spin-coating is a dynamic process, where the close packing is mostly determined by the spin speed. By changing the wetting properties of the substrate, it is possible to induce dewetting, occurring simultaneously to evaporation. Dewetting is a process that causes the formation of voids within the uniform solvent film, and to force the surrounding liquid and NPs to move away from them. In this case, the formation of NP superstructures is strongly affected by the solvent evaporation rate\textsuperscript{28}.

The image of NP monolayers shown in Figure 3b provides an excellent illustrative example of the effect of dewetting during the formation of NPs superstructures. In this system, the NPs in solution are spin-coated in the same fashion as the system reported in Figure 2a. However, the substrate was changed from Si(100) (with few nm of native oxide layer) to a hydrophobic Si substrate coated with 300 nm of SiO\textsubscript{2}. The NPs form a monolayer presenting defects and holes, often few hundreds of nm in diameter, as a consequence of dewetting. Still the NPs are packed in a HCP like lattice as confirmed by the GISAXS pattern in Figure 3d. The imperfections in the layer, having a large shape and size distribution, do not affect the Bragg peak positions, however, the absolute intensities of the Bragg peaks are lowered by one order of magnitude with respect to those found in the other GISAXS patterns. This reduction can be accounted for by the penetration of the x-rays into the substrates. At $\alpha_i$ of 0.5° the incident angle is, in fact, above the critical angle of SiO\textsubscript{2}, so that the x-rays penetrate into the substrate.
and hence less x-ray photons contribute to the diffuse pattern. This growth mode can therefore still be classified as FvdM growth with incomplete coverage of the substrate due to dewetting.

So far we have presented the self-assembly of NPs which is characterized by a very fast evaporation rate of the solvent. The opposite scenario is represented by the extreme case when complete wetting is ensured and solvent evaporation is so slow that it seems virtually non-existent. In this case, the NPs which are free to move within the toluene, given sufficient time, will assemble to an equilibrium position. Such case is readily obtained by means of sedimentation of NPs, a process by which the solvent is deposited on PMMA coated silicon substrates (either type of PMMA layer) and let to evaporate over an extended time. Complete evaporation takes from few minutes to few hours depending on the volume of toluene, the temperature of the system and the atmospheric pressure conditions.
Figure 4: The SEM images of NPs obtained by sedimentation of NPs on PMMA coated silicon substrates showing (a) islands (supercrystals) with random orientations. The inset shows a single supercrystal with HCP type NP ordering. The corresponding GISAXS pattern is shown in (b).

Figure 4a and b show the SEM images and the GISAXS pattern of NP self-assembly after sedimentation. Due to the formation of distinct islands the growth process resembles - at the first glance - a VW growth mode, but should rather be considered as seed-mediated growth of three-dimensional "supercrystals" or "mesocrystals". The dark contrast, which appears as bridges between the islands, is due to impurities of the toluene, which concentrate in the remaining solvent before complete evaporation. The image of networks of residual toluene also gives an impression of the complex fluid flow during the evaporation process, driven by
the local air-flow, toluene over-pressure and temperature fluctuations over the substrate. As found from the SEM image, the formation of mesocrystals results in this case from another manifestation of a dewetting process during evaporation at very small NP concentration. By examination of the black contrast of the remaining solvent one can easily comprehend the formation of dry cells growing ever larger until coalescing into each other. Hence, the formation of mesocrystals is restricted mostly to areas of dry cells contacting each other, as not until then the particle concentration and/or solvent pressure have overcome a critical threshold.

Each supercrystal is like a single crystal formed by NPs, with different facets. The nearly triangular shape of the supercrystal shows that the out-of-plane growth direction is primarily along the (0001) axis and further corroborates the assumption that 3D supercrystal growth occurs. From Figure 4a it is clear that the planar orientation of the islands is random with respect to each other. Therefore, in the GISAXS pattern the scattered intensity is averaged over all planar orientations. Statistically one can observe all possible lattice planes similar to a powder sample and hence represents a 2d-pattern.

The GISAXS patterns measured at a glancing angle of 0.1° is shown in Figure 4b. It was not possible to observe any in-plane Bragg peaks at higher angles of incidence. At this very shallow angle of incidence, i.e. below the critical angles of Si and PMMA, the scattering is dominated by (or originates from) the surface features, viz. the NP assembly, while the x-rays do not penetrate the substrate material. Consequently, the shallow incident angle increases considerably the footprint and therefore the contributing scattering volume of the NPs. Hence the best information about the NP ordering inside the islands is obtained at low angles of incidence.

The Bragg peaks are much sharper compared to other GISAXS patterns. Except the (10) peak other peaks have less extension along the qz axis, indicating a 3-dimensional nature of the islands. As the number of layers perpendicular to the substrate grows, the Bragg peaks
become more point-like. Instead of Bragg rods, Bragg spots are seen. The HCP lattice of the NPs is confirmed from the observation of (11), (20), (21) and (30) peaks. The (22) peak is not visible in this case.

Figure 5: The SEM images of NPs spin-coated on (a) electropolished aluminum ('Al') (b) a-plane sapphire substrates ('Al₂O₃'). The corresponding GISAXS patterns are shown in (c) and (d) respectively.

The analogy between atomic growth modes and NPs growth modes would not be complete without mentioning the role of the substrate surface roughness on the NPs self-assembly. Obviously with regards to the substrates presented up to now, lattice mismatch or the atomic roughness can be neglected as the NP size is much bigger than the atomic size roughness. The surface interaction between substrate and solvent represents the driving force for the growth modes earlier considered, while the NP and substrate interaction plays only a minor role. However, with the appropriate choice of substrate, surface roughness becomes a major player.
during the ordering mechanism process of NPs. The effect of surface roughness is seen when NPs are \textit{spin-coated} onto electropolished Al and Al$_2$O$_3$ substrates, as shown in Figure 5. From the SEM image a close packed monolayer of particles can be observed. The Al substrate, electropolished prior to the spin-coating process, has a roughness in the order of a few nm and gives rise to a wavy appearance to the substrate. (Al$_2$O$_3$ substrate is an insulator and hence one loses resolution at high magnifications in SEM).

Once again, by the combination of both GISAXS and SEM imaging one is able to obtain the complete picture of growth modes and self-assembly. The GISAXS pattern in Figure 5c only shows a few broad Bragg peaks. The resolution of this system is complicated by the extreme ductility of the substrate which causes it to slightly bent and deform even during careful manipulation, since the thickness of Al foil is less that 0.5 mm and the discs are very malleable. On the other hand, the GISAXS pattern of the NPs on Al$_2$O$_3$ substrate (Figure 5d) evidences clearly a HCP lattice of the monolayer. The (11) and (20) peaks are well resolved. Even at high $q_z$ values it is possible to resolve the (21) and the (30) peaks as well, which was not possible in previous measurements. The narrower peak width along $q_y$ axis indicates larger coherence length or larger crystallite sizes. The growth modes in these cases also resemble to that of a layer-by-layer or FvdM growth process, although the SEM images show that the shape and size of imperfections in the monolayer differ considerably from the case of NPs spin-coated on Si.

Having presented a number of case studies of different NP growth conditions, it is clear that the hexagonal close packing is the geometry naturally preferred by NPs for self-assembly, as it obviously provides the highest packing density. One might expect this behavior for systems with negligible entropic effects that cause other geometries to be energetically favorable. In order to gain further understanding on how the substrate composition and surface roughness affect the degree of packing, we need to look into one-dimensional line cuts at a constant $q_z$ values of the GISAXS patterns presented in this work. The line cuts, shown in
Figure 6 for all samples, were taken at $q_z = 0.9$ nm$^{-1}$, with the exception of the NPs prepared by sedimentation (see Figure 6d) taken at $q_z = 0.166$ nm$^{-1}$. The Bragg peaks have been indexed assuming a HCP lattice. For simplicity, the negative $q_y$ axis alone is shown, together with Lorentzian fits to the Bragg peaks. The inverse of the width of the Lorentzian yields the coherence length of the NP crystallites.

Figure 6: Line cuts at constant $q_z$ for NPs self-assembled on: (a) Si, (b) SiO$_2$, (c) Si$^+$ PMMA/MA (33 %), (d) Sedimentation, (e) Al and (f) Al$_2$O$_3$ substrates. Indexing is given according to a 2-dimensional HCP lattice. The open circles are measured data points and the solid lines are Lorentzian profile fits to the peaks.
The position of the first Bragg peak is similar for all samples. That means that the hexagonal unit cell has the same lattice constant for different growth modes. The main difference between the samples is found in the values of long range ordering of the NPs.

Table 2. Comparison of intra-planar distance \( d_{10} \) (calculated from GISAXS pattern) and the coherence length \( \xi_{10} \) of (10) planes (calculated from the Lorentzian fit to the peak of width \( \omega_{10} \)) for NPs self-assembled on different substrates. The corresponding values for the substrate PMMA (4%) is not included in the table because it has amorphous like structure.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>( d_{10} = \frac{2\pi}{q_{10}} ) (nm)</th>
<th>( \xi_{10} = \frac{2\pi}{\omega_{10}} ) (nm)</th>
</tr>
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<tr>
<td>Si</td>
<td>17.55 (± 0.02)</td>
<td>70 (± 0.53)</td>
</tr>
<tr>
<td>Al(_2)O(_3)</td>
<td>17.95 (± 0.02)</td>
<td>90 (± 0.71)</td>
</tr>
<tr>
<td>Al</td>
<td>17.9 (± 0.04)</td>
<td>87.26 (± 5)</td>
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<tr>
<td>SiO(_2)</td>
<td>17.79 (± 0.01)</td>
<td>73 (± 0.93)</td>
</tr>
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<td>PMMA.4P</td>
<td>17.95 (± 0.06)</td>
<td>16.5 (± 1.2)</td>
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<tr>
<td>PMMA.33P</td>
<td>17.65 (± 0.023)</td>
<td>100 (± 4)</td>
</tr>
<tr>
<td>Sedimentation</td>
<td>17.9 (± 0.008)</td>
<td>282 (± 10)</td>
</tr>
</tbody>
</table>

Table 2 lists the intra-planar distance \( d_{10} \) calculated for (10) peak from the Lorentzian fitting and the coherence length of the films from the width of the Lorentzian \( (\omega_{10}) \) on different substrates. As expected the sedimentation sample has the maximum coherence length of the order of 300 nm. In this case the crystal was formed under an equilibrium condition and shows the maximum ordering.

In all other cases the crystals were formed at a much faster rate and did not have enough time to form a single crystal, rather forming a continuous film with polycrystalline grain boundaries. Curiously, the rougher Al and Al\(_2\)O\(_3\) substrate, although promoting monolayer
growth with higher density of defects, present a slightly longer correlation length than their Si sample counterpart.

We can therefore conclude that NP ordering is different on different substrates, revealing a striking resemblance to atomic growth modes.

The interaction between the solvent and the substrate is the most dominant of all the interactions. The surface interaction defines the wetting ability of the solvent on different substrates. Since in the present case toluene is used as a solvent, the "Toluenophobicity" of the substrate determines, whether the solution will completely wet the surface or form islands on it. Only in a second step, the ordering or crystallization of NPs takes place. The solvent (toluene) starts evaporating and NPs form a close packed structure. The crystallization process is a slow process. Hence, with spin-coating there is not sufficient time for the formation of large crystals leading to the formation of domains or crystallites. If sufficient time is allowed, as in the case of a sedimentation process, it is possible to generate three-dimensional "super-crystals".

Methods:

The iron oxide NPs are chemically synthesized and are purchased from Ocean NanoTech LLC Company. The mean diameter is 18 nm with a size distribution of 6.5 %. The iron oxide core is surrounded by oleic acid shell with 2 nm thickness. This surfactant prevents the NPs from agglomeration. The NPs are dispersed in toluene solvent and are stored in a sealed bottle.

Samples: The samples were prepared by spin-coating as described in the literature\textsuperscript{30,31}. The concentration of NP in toluene determined the number of layers formed during spin-coating. In order to obtain a monolayer the NP dispersion was diluted by pure toluene solvent in 1:1 ratio. Then 0.01 ml of this diluted solution with NPs was taken from the bottle in an oil and
silicon free plastic syringe purchased from NORM-JECT® with EROSA disposable hypodermic needles. Different substrates with typical dimensions of 10 mm² were used. The various substrates used for the investigation are silicon (100) substrate with natural oxide ('Si'), silicon (100) with 300 nm silicon dioxide ('SiO₂'), silicon(100) spin-coated with copolymer polymethyl methacrylate/ methacrylic acid (PMMA/MA) with 33 % solid contents ('PMMA_33P'), polymethyl methacrylate (PMMA) with 4% solid contents ('PMMA_4P'), polished Aluminum ('Al') and a-plane sapphire ('Al₂O₃'). The Al substrates were prepared starting from high purity Al foils (Goodfellow, 99.999%), cleaned by sonication in isopropanol and ethanol for 10 min. The electropolishing of the surface was performed in a 1:3 vol. perchloric acid and ethanol mixture at 10 °C during 10 min, with a constant dc potential of 20 V applied between the sample and a Pt mesh. The Si and SiO₂ substrates were purchased from CrysTech. The PMMA_33P and PMMA_4P substrates were prepared by spin-coating PMMA/MA with 33 % solid contents and PMMA with 4% solid contents on Si substrates at 4000 revolution per minute (rpm) for 30 seconds, respectively. After spin-coating it was heat-treated at 80° C in air for 20 minutes on a hotplate (Przitherm PR35 with microprocessor control). The PMMA/MA and PMMA were purchased from Allresist. All spin-coating processes were performed on a commercial spin-coater from SPIN. The NPs were spin-coated in two steps. First, the substrate was spun at 300 rpm for 3 seconds and within 3 seconds the NP solution was drop-cast onto the substrate. This step helps in spreading the solution uniformly throughout the substrate. In the second step, the rotation was increased to 4000 rpm with an acceleration of 1000 rpm per second. This step determines the thickness of the film. Our effort was to keep the volume of NP solution used and the rotational speed for all the substrates constant. This yields a uniform nominal thickness, irrespective of the arrangement of the particles. After spin-coating, the samples were also heat treated at 80° C in air for 20 minutes for the evaporation of the toluene solvent. The sedimentation sample was prepared in a slightly different way. A PMMA_4P substrate was
immersed in a highly diluted solution of NPs. The volume ratio of NP solution to toluene was 1:40000. The beaker containing the substrate and the solution was tightened completely and the solvent was allowed to evaporate slowly overnight through a small opening at the top.

**Characterization:** A FEI Quanta 200 FEG scanning electron microscope was used for imaging the NPs, which was equipped with an Everhart-Thornley secondary electron detector. The energy was 20 keV with a 30 µm aperture for better resolution. The GISAXS experiments were performed at Hasylab (Hamburg, Germany) beam line BW4 at photon energy of 8.798 keV (λ = 0.138 nm)\(^{32}\). A MAR CCD camera with pixel size 79.1 µm and resolution of 2048 × 2048 pixels was used to capture the two dimensional images. An aperture (0.4 mm × 0.4 mm) and focusing lens system reduced the beam size to 36 µm × 23 µm (horizontal × vertical). The sample to detector distance was found to be 210.44 cm using silver behenate as a calibrant. This gives a q-space resolution of 1.7 × 10\(^{-3}\) nm\(^{-1}\). The intensities were normalized to the monitor intensity. The geometry of the GISAXS experiment is also shown in Fig 8. The angle of incidence for spin-coated samples was 0.5° and for the sedimentation sample was 0.1°.

![Fig. 8. Geometry of the GISAXS measurement. \(\alpha_i\) is the angle of incidence, \(\alpha_f\) is the angle of reflection, \(2\theta_f\) is the in-plane angle. The specular condition is satisfied for \(\alpha_i = \alpha_f\).](image)

Indexing the lattice: The indexing was performed assuming a two dimensional hexagonal lattice. The first Bragg peak is the (10) peak and the lattice constant were calculated from the position of this peak and using the following equation.

\[ d_{hh} = \frac{a}{\sqrt{\frac{4}{3}(h^2 + hk + k^2)}} = \frac{2\pi}{q_y^{hk}} \]

Here \((hk)\) corresponds to Miller indices, \(d_{hh}\) corresponds to the inter-planar distance, \(a\) is the lattice constant and \(q_y^{hk}\) is the position of Bragg peak as observed in the experiment.

References:


