Self-assembly of highly fluorescent semiconductor nanorods into large scale smectic liquid crystal structures by coffee stain evaporation dynamics

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Received 11 November 2008, in final form 19 January 2009
Published 11 June 2009
Online at stacks.iop.org/JPhysCM/21/264013

Abstract
We deposit droplets of nanorods dispersed in solvents on substrate surfaces and let the solvent evaporate. We find that strong contact line pinning leads to dense nanorod deposition inside coffee stain fringes, where we observe large scale lateral ordering of the nanorods with the long axis of the rods oriented parallel to the contact line. We observe birefringence of these coffee stain fringes by polarized microscopy and we find the direction of the extraordinary refractive index parallel to the long axis of the nanorods.

Supplementary data are available from stacks.iop.org/JPhysCM/21/264013

(Some figures in this article are in colour only in the electronic version)
the pinning of the contact line and its retraction motion, which can be exploited in order to achieve ordered assemblies of nanorods. We use polarized microscopy to characterize the refractive properties of the obtained nanorod assemblies.

In the coffee stain effect fringes consisting of deposited material form due to pinning and subsequent retraction of the contact line of a drying droplet. The pinning of the contact line induces an outward flow of the liquid from the inner regions towards the edge of the droplet in order to compensate for the evaporated solution from the droplets surface. The radially outward flow transports particles that are dissolved in the solution towards the contact line of the droplet. When the contact line eventually retracts, it leaves a ring-like structure of dense material on the surface which is called the coffee stain.

In this work we investigate the coffee stain effect and the resulting patterns for solutions of nanorods dispersed in toluene. We find that the assembly pattern of the nanorods depends significantly on the nanorod density in the solution and on the strength of the contact line pinning. We start with discussing our experiments for solutions with nanorod concentrations below $10^{-6}$ M. We deposited small droplets of toluene (10–30 µl) with different nanorod concentrations ($6 \times 10^{-7}$, $6 \times 10^{-8}$, $1 \times 10^{-8}$ M) on a SiO$_2$ substrate and let the solvent evaporate under ambient conditions. We find that the contact line retracts, undergoing pinning and depinning, in an overall concentric motion, although locally some ‘arch-shaped’ irregularities and fingering occur, as observed in [22]. After drying we observed fringes on the substrate consisting of densely deposited nanorods, as shown in figure 1(b). We find ribbon-like nanorod assemblies within the dense fringes in which a bunch of nanorods is aligned side by side, as displayed in figure 1(d), whereas in the area in between the fringes the rods are deposited in a layer of random order (figure 1(c)). The nanorod ribbons displayed in figure 1(d) form ‘spaghetti-like’ entangled structures, and we find that with decreasing nanorod concentration in the solution the length and the density of the ribs in the fringes decreases (see figure S2 of the supporting information (available at stacks.iop.org/JPhysCM/21/264013)). Similar ribbon structures formed by nanorods have also been observed for hard, rod-like β-FeOOH suspensions [25–27], Se and CdSe nanorods [17, 28], and also for gold nanorods [29].

Figure 2 shows what happens if we deposit droplets of toluene solutions with nanorod concentrations of $10^{-6}$ M on SiO$_2$ or glass. Here SEM inspection of the coffee stain fringes (figure 2(a)) shows that the nanorods assemble in much higher order. Higher magnification images show (see figure 4) that the nanorods are aligned in smectic structures with their long axis parallel to the circular fringe. Within the fringe we find stripes with differing density. We observe that the inner border of the stripe with the highest density (see dotted lines in figures 2(a) and (b)) is well defined and that this part of the coffee stain shows the highest order of the nanorod assembly. We find that the increased nanorod density in the solution leads to stronger pinning of the contact line and a thicker fringe of deposited nanorod material within the fringe. The thick fringe of nanorod material prolongs the pinning of the contact line and, subsequently, leads to the rupture of the liquid film behind the fringe (see figure 4(c)), similar to the mechanism reported in [30]. Following this model, the ring of deposited material
Figure 2. SEM images of the coffee stain fringe for nanorod deposits with higher (a) and lower (b) density. We observe high lateral order with the individual nanorods aligned parallel to the contact line. We note the well-defined inner border of the dense stripe of the fringe and the large degree of lateral rod alignment near this border.

behind the contact line prolongs the pinning until the minimum of the droplet height behind the deposit leads to a rupture of the fluid.

The nanorods in the coffee stain fringes shown in figure 2 align parallel to the contact line. This arrangement results from the balance of the torque exerted on the ends of the nanorods by the solvent flow towards the pinned contact line. In this picture the first rods that encounter the pinned contact line orient along the contact line and act as nucleation centers for the following parallel arrangement. We know from previous studies [10] that the CdSe/CdS core–shell rods arrange easily into ordered lateral arrays. Figure 3 shows SEM images where the nanorods align parallel to the border line of voids. Here the nanorods were dissolved in toluene and GaAs substrates were immersed at a tilted angle of 45° in the nanorod solution and the solvent was allowed to evaporate. We find that the tendency of the rods to align parallel to the border line and the smectic order of the deposited nanorods increase with increasing rod density. Theoretical modeling of the evaporation dynamics of solvents containing colloidal nanocrystals shows that the voids start to form well before the complete evaporation of the solvent [31]. Consequently, also in this case the torque argument for the nanorod that encounters a contact line applies. We note that the case of the pinned contact line in our coffee stain experiments differs significantly from a continuously retracting contact line, where the rods align along the flow direction due to capillary forces.

We find that the nanorod alignment in the coffee stain region depends mainly on two factors. The first is the nanorod density in the solution. Higher nanorod concentrations lead to

Figure 3. SEM images of CdSe/CdS core/shell nanorods deposited on tilted GaAs substrates by slow solvent evaporation (of the order of 5 nm s⁻¹) for increasing nanorod density from (a) to (c). The scale bar corresponds to 500 nm for all images.
Figure 5. (a)–(c) SEM images of areas within the coffee stain fringes showing large scale assembly of nanorods in liquid crystal structures forming multilayered stripe patterns. (b) Higher SEM magnification of stripes with different thicknesses, i.e. different numbers of layers. (c) Magnification which resolves individual rods within the nanorod ribbons. The scale bar for all three images corresponds to 500 nm. Some nanorods are highlighted in red as a guide to the eye.

We can stimulate such a circular fluid retraction by applying an electrical dipole field at the center of the drop, as illustrated in figure 4(a). If we position two electrical micromanipulators on the substrate surface at the center of the drop we find that the contact line pinning related to the outermost coffee stain is enhanced. After the rupture of the fluid film we observe a circular retraction of the fluid, as illustrated in figure 4(b) (see figure S3 of the supporting information (available at stacks.iop.org/JPhysCM/21/264013)). Two films showing the drop drying dynamics with, and without, voltage bias of 150 V are available online at stacks.iop.org/JPhysCM/21/264013.

Figure 5 shows the nanorod alignment that we obtained by positioning two electrical micromanipulators (tungsten needles) with 150 V voltage bias at the center of the drop as illustrated in figure 4(a). We find multilayers of nanorods that are aligned in a smectic structure, but in addition the smectic domains arrange in stripes with almost constant width over several micrometers in length. The width of these stripes ranges from some hundreds of nanometers to some microns. Figures 5(b) and (c) show larger magnification SEM images that resolve the ribbons and the individual rods within the ribbons. We note that these stripe-like superstructures have relatively sharp, continuous boundaries (less than 50 nm across) that extend over several microns. Patterns like the ones displayed in figure 5 can extend up to the millimeter scale as shown in figure 6. We think that these stripe-like patterns can be attributed to pinning and depinning of the contact line on a comparatively short length scale that resemble a stick–slip movement [30]. We would like to mention that the pinning of the contact line can also be influenced by patterning the surface of the substrate. For example, we evaporated a ring-shaped Au layer (thickness 30 nm) with a diameter of 5 mm on a glass substrate and found that the contact line pinning was enhanced on the Au a short distance from the Au/glass border. Also in this case we observed the highly ordered lateral assemblies of nanorods.

Figure 6. Polarized microscopy images taken in crossed polarization with a 530 nm retardation plate inserted in the optical path at 45°. The sample was prepared by drop evaporation with 150 V bias voltage as described in the text. The sample has been rotated by 90° from (a) to (b). The dashed black arrows indicate the orientations of the polarizer and analyzer, and the green arrow shows the orientation of the 530 nm retardation plate. ‘Inside’ and ‘outside’ refer to which side of the fringe faces the inside/outside of the coffee stain.
For samples with a uniaxial anisotropy two refractive indices can be assigned, the ordinary refractive index $n_o$ with orientation perpendicular and the extraordinary index $n_e$ with orientation parallel to the axis of anisotropy, respectively. The birefringence is then $B = |n_e - n_o|$, and the sign of birefringence can be evaluated using polarized microscopy by inserting a first order retardation plate (530 nm) in the optical path in crossed polarization, with the axis of anisotropy of the sample aligned parallel to the slow axis of the retardation plate. In this configuration positive birefringence leads to blue color, and negative birefringence to red color in the polarized microscopy image.

Figure 6 shows part of the outer coffee stain fringe where a bias voltage of 150 V was applied to the manipulator tip during solvent evaporation (as described above). In this case we used a microscope glass slide as substrate. We clearly observe the dense part of the fringe in blue color revealing macroscopic ordering of nanorods arranged in liquid crystal phase. Rotation of the sample by 90° leads to red/orange color of the fringe as it should (figure 6(b)). We conclude that the direction of $n_e$ is oriented parallel to the contact line, and from comparison with SEM images we find that $n_e$ is directed along the long axis of the nanorods.

In conclusion, we formed highly ordered multi layers of laterally organized nanorod assemblies inside the coffee stain fringes of an evaporating droplet of dense nanorod solution. In detail, we observed that the individual nanorods form ribbons and that these ribbons arrange in smectic order in the dense central region of the coffee stain fringe. We found that the individual nanorods in this arrangement are oriented parallel to the contact line and we could also assign the slow axis of refraction of the nanorod assemblies along this direction.

Acknowledgments

The authors thank Benedetta Antonazzo for technical assistance in the nanochemistry laboratory. The work was supported by the SA-NANO European project (contract No STRP013698).

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