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In situ Formation of Plasmonic Gold Nanoparticles in Ordered Block Copolymer Films

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Context

Metamaterials are of current interest for their unique optical properties. Although they are traditionally obtained by lithography, bottom-up ways of fabrication are now investigated for a precise nanometric control of the structure, through self-assembly and nanochemistry. To this end, organized films containing gold nanoparticles are widely studied for they combine the controlled structure of a film to the plasmonic properties of these nanoparticles.

We report here the *in situ* formation of gold nanoparticles (AuNPs) in a polystyrene-*block*-poly(vinylpyridine) (PS-*b*-PVP) copolymer solution, then cast as AuNP containing ordered films.

By use of an appropriate solvent, casting a block copolymer solution can form films of oriented cylinders (perpendicular or parallel to the substrate). The addition of gold salt *Au(III)chloride* ($AuCl_3$) followed by ultra-sound treatment before casting the film gives birth to spherical AuNPs (2-3 nm in size) located, after deposition, inside the cylinders. For parallel cylinders, AuNPs are found to re-direct them to a quasi-perpendicular orientation, while no major perturbation is found for the perpendicular cylinders. Larger spherical AuNPs (3-4 nm) exhibiting a plasmon resonance are also obtained by successive additions of gold salt to the AuNPs and locate equally inside the cylinders after casting the film.

This method of *in situ* insertion of AuNPs in patterned films provides nanocomposite materials with plasmonic properties and opens new ways for the preparation of metamaterials.

AuNPs in perpendicular cylinders

- Solution of PS₂₅-*b*-P4VP₇ in non-selective mixed solvent toluene/THF (80:20) spin-coated onto a silicon substrate → cylinders oriented normal to the substrate

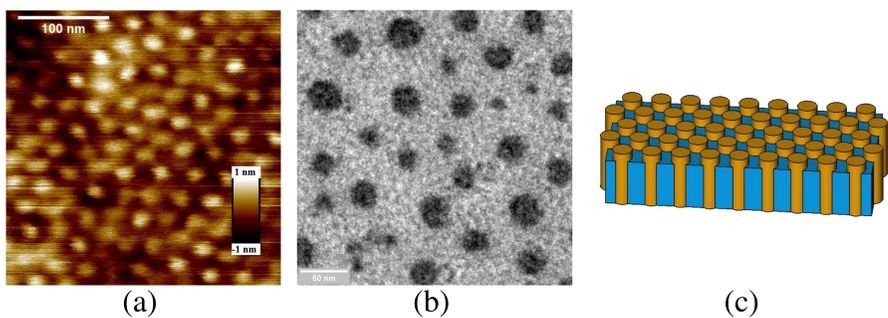


Figure 1: a/ AFM height picture and b/ TEM picture of PS-*b*-P4VP cylinders oriented normal to the silicon substrate after spin-coating. The cylinders have a 2D-hexagonal organization and their average diameter is 40 nm; c/ 3D-illustration of the cylinders of PVP (orange) in the PS matrix (blue) with an orientation normal to the substrate and a hexagonal organization.

►► Perpendicular order without gold addition

- Gold salt ($AuCl_3$) addition to the solution, previous to casting, followed by sonication, then spin-coating → reduction of the gold salt into AuNPs

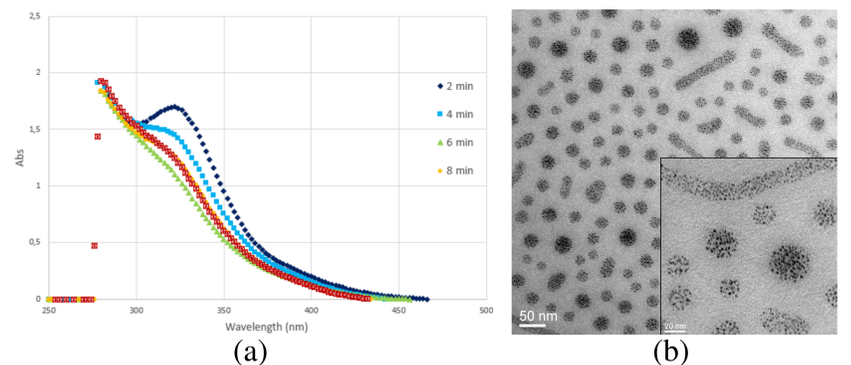


Figure 2: a/ UV-visible spectra showing the decrease of the gold salt peak over the sonication; b/ TEM picture of AuNPs formed by sonication and confined in PS-*b*-P4VP cylinders oriented normal to the substrate. The AuNP size is around 2 nm (see inset). The amount of gold per pyridine is 5 equivalents.

►► Formation of AuNPs (diameter 2-3 nm) located inside the cylinders

Re-orientation of parallel cylinders

- Solution of PS₃₄-*b*-P2VP₁₈ in chloroform ($CHCl_3$), spin-coated onto a silicon substrate + annealing in $CHCl_3$ vapors → cylinders parallel to the substrate
- $AuCl_3$ addition before spin-coating and annealing in $CHCl_3$ vapors → reduction of the gold salt into AuNPs + re-orientation of the cylinders

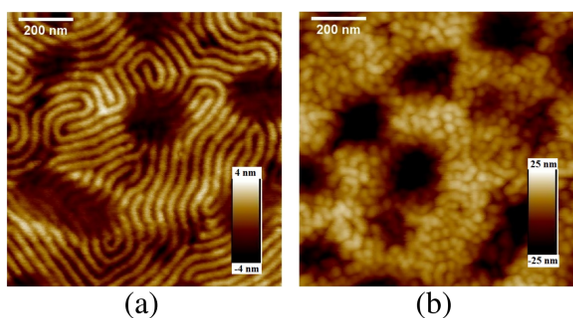


Figure 3: AFM height pictures of deposited and annealed PS-*b*-P2VP film a/ without and b/ with AuNPs formed by sonication. The presence of the AuNPs inside the cylinders triggers their re-orientation almost normal to the substrate.

►► Loss of the cylinder orientation while adding the AuNPs. Not seen with perpendicular cylinders

Formation of plasmonic AuNPs

- Further addition of $AuCl_3$ and sonication over a AuNPs/PS-*b*-P4VP solution in successive steps → growth of the AuNPs

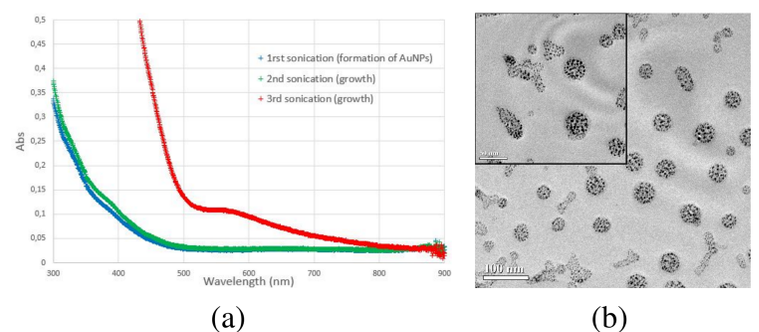


Figure 4: a/ UV-visible spectra: after the formation of AuNPs (blue) and the first addition/sonication (green), no plasmon is noticed. The plasmon peak (around 560 nm) appears after the second addition/sonication; b/ TEM picture of the film for a AuNPs/PS-*b*-P4VP solution after formation and growth of the AuNPs. The AuNPs are larger (3-4 nm, see inset), which explains their plasmon resonance and still located inside the cylinders. The amount of gold by pyridine is 10 equivalents and each growth step adds 2 more equivalents (for a total of 2 steps).

►► Formation of larger AuNPs (diameter 3-4 nm) located inside the cylinders and exhibiting a plasmon resonance

Conclusions and Perspectives

- Formation of AuNPs in copolymer solution by ultra-sound treatment and insertion in ordered and oriented cylinders in copolymer films
- Re-orientation of the parallel cylinder to quasi-perpendicular ones by addition of the AuNPs
- Formation of plasmonic, spherical AuNPs (3-4 nm) by a succession of gold salt additions
- Simple methodology for the fabrication of nanocomposite materials exhibiting plasmonic properties
- Quantification of the optical properties for these materials

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