Mass production of polymer nano-wires filled with metal nanoparticles

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The ongoing progress in nanotechnology and its applications, the development of strategies for connecting nano-scale systems to micro- or macroscale elements is hampered by the lack of structural components that have both, nano- and macroscale dimensions. The production of nano-scale wires with macroscale length is one of the most interesting challenges here. There are a lot of strategies to fabricate long nanoscopic stripes made of metals, polymers or ceramics but none is suitable for mass production of ordered and dense arrangements of wires at large numbers. In this paper, we report on a technique for producing arrays of ordered, flexible and free-standing polymer nano-wires filled with different types of nano-particles. The process utilizes the strong response of photosensitive polymer brushes to irradiation with UV-interference patterns, resulting in a substantial mass redistribution of the polymer material along with local rupturing of polymer chains. The chains can wind up in wires of nano-scale thickness and a length of up to several centimeters. When dispersing nano-particles within the film, the final arrangement is similar to a core-shell geometry with mainly nano-particles found in the core region and the polymer forming a dielectric jacket.

The ongoing minimization of functional elements in nanoscience and nanotechnology raises considerable challenges of connecting them to macroscopic components. It would be highly desirable if one could dispose of wires with nano-scale cross section but that extend to macroscale dimensions without flaws until they ultimately connect to a macroscale contact or patch, which can be easily accessed¹. For the above problem one might immediately think of e-beam lithography as one (albeit expensive) solution for purely metallic wires, but what is in fact most interesting for electronic²⁻⁴, optoelectronic⁵⁻⁷, electrochemical⁸⁻¹⁰ and information processing industry¹¹⁻¹³ are hybrid materials such as inorganic materials/nanoparticles integrated into soft polymeric matrices. There are currently several methods to produce hybrid nano-wires ranging from simple procedures of molecular self-assembly to specialized synthetic routes. In the first case the specific interactions such as hydrogen bonding, electrostatic forces or π−π stacking can lead to self-organization of cyclic peptides, polypeptides, amino acids, rigid organic molecules, conjugated polymers and polyelectrolytes to nanofibers and nanowires¹⁴⁻²⁰. One can also synthesize multicomponent rod-like structures that contain metals, inorganic semiconductors, and conducting polymers via template assisted in situ electrochemical deposition²¹,²². Using a self-assembly approach of mesoscopic metal-polymer amphiphiles it was also possible to organize them into curved structures²³,²⁴. In all above mentioned approaches the alignment and simultaneous connection to macroscopic components seems to still be out of reach.

With our approach, the above mentioned problems could at least partly be bypassed. With our discovery, how nano-wires can be produced in a self-organized manner, we can provide a simple, fast and cheap method to fabricate a wide variety of different nano-fiber systems in mass production. At the same time, the problem of achieving a very high aspect ratio is solved, and the dielectric cover of the wires is provided automatically, and can possibly be influenced separately by a simple choice of process parameters.

At the heart of our fabrication process is the careful synthesis and design of special photosensitive polymer brushes. These are thin polymer films where polymer chains are constrained by covalent bonding to a solid

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As a molecular actuator that converts optical energy into mechanical work. Under irradiation with UV light the azobenzene undergoes reversible photo-isomerization from the trans- to the cis- conformation. The photo-isomerization is associated with significant alteration of the azobenzene properties: the length of the molecules changes by almost a factor of two, the dipole moment changes from 0 Debye for the trans- (the more stable configuration) to ~3 Debye for the cis-isomer. When azobenzene molecules are attached to a polymer, they act as active elements that transform optical energy through molecular mechanisms to the global deformation of the polymer matrix. Among the many unexpected physical phenomena related to azobenzene containing polymers is the formation of so-called surface relief gratings (SRG) upon irradiation with an interference pattern of UV light. During this process the polymer film deforms according to the distribution of intensity or a polarization pattern. Opto-mechanical forces generated during the absorption of UV light are strong enough in order not only to deform glassy polymer film, but also to rupture metal films and stretch graphene multilayers deposited on top of the polymer film. In polymer brushes, the influence of the photosensitive side chains led to the phenomenon of opto-mechanical scission of the covalently bound chains. The polymers are ruptured locally during the deformation process at the place where the topography minima are formed. We have used this property for instance, in order to pattern brushes on a nano-meter scale. Indeed, irradiating the brush with an interference pattern of UV light, most of the polymer material after treatment with good solvent is removed from the surface, leaving characteristic line or dot patterns behind.

Here we report on how one can use the process of the local rupturing of polymer chains within photosensitive polymer brushes in order to produce large amounts of ordered wires of nano-scale thickness and a length of up to several centimeters filled with nano-particles of different nature.

**Results and Discussion**

In our study we utilize a poly(methacrylic acid) (PMAA) brush (prepared as described in the Material Section) with the following molecular parameters: the initial thickness in the dry state was $h = 84$ nm, molecular weight $M_n = 3.1 \times 10^6$ g/mol, and grafting density $\sigma = 0.018$ nm$^{-2}$ resulting in a distance between neighboring chains of $D = 7.4$ nm (Fig. 1). The parameters of the brush were determined as described in ref. 29. The PMAA brush is negatively charged. To integrate negatively charged gold nano-particles (AuNPs) into the negatively charged polyelectrolyte brush and to simultaneously load the film with photosensitive azobenzene containing cationic surfactants (Fig. 1), we first prepare a mixture of water suspension of AuNPs and gold nano-particles, respectively (Fig. 1). In this way, the complex of the Azo-AuNPs is positively charged, allowing for ionic interaction with the negatively charged polymer brush. When the brush was exposed to the azobenzene modified AuNPs, these particles diffuse into the brush and adsorb to the polymer matrix resulting in an increase of the brush dry thickness. It was recently reported that monovalent as well as multivalent ions cause shrinkage in swollen polyelectrolyte brushes. In our case the increase in the brush dry thickness is explained by increase of the overall molecular weight of the polymer chains when surfactant and Azo-AuNPs complex are loaded in the film. The increase in dry thickness of the brush depends on the ratio $C_{\text{azo}}/C_{\text{Au}}$, where $C_{\text{azo}}$ and $C_{\text{Au}}$ are the molar concentrations of the azobenzene containing cationic surfactant and gold nano-particles, respectively (Fig. 2a). In our study we use four solutions with ratios of $C_{\text{azo}}/C_{\text{Au}}$ equal to 0.3, 0.5, 1 and 1.2. To vary $C_{\text{azo}}/C_{\text{Au}}$ we change the concentration of surfactant while keeping the concentration of gold particles constant at $C_{\text{Au}} = 1$ mM. In all four cases there is a formation of an azobenzene containing shell around the AuNPs rendering them positively charged as determined by measuring the surface potential of the particles. Larger values of $C_{\text{azo}}/C_{\text{Au}}$ indicate an excess of surfactant in the system. The reason for using different concentrations is to find the optimal conditions at which one can load the brush with the maximum number of...
AuNPs while keeping the brush photosensitive. As can be inferred from Fig. 2a (black curve), the brush height increases up to (300 ± 50) nm after absorption of azo-AuNPs for $C_{\text{azo}}/C_{\text{Au}}$ starting from 0.3. For higher surfactant concentrations, the brush thickness increases even more up to 420 nm, i.e. 5 times of its initial thickness.

Since the absorption peaks of azobenzene and Au nano-particles are well separated ($\lambda_{\text{azo}} = 338$ nm\(^4\), $\lambda_{\text{Au}} = 545$ nm, Fig. 2c), we determine the amount of gold nano-particles absorbed within the brush using absorption spectra of the azo-AuNPs solution before and after immobilization. Figure 2c shows corresponding spectrum of brush loaded with azo-AuNPs. The absorption peak of the surfactant is clearly observed in Fig. 2c and can be used to calculate the surfactant concentration within the brush. The surface plasmon absorption peak of the gold nano-particles within the brush is shifted to 545 nm due to the change in refractive index with respect to the water solution\(^5\). Figure 2b shows the dependence of the amount of gold nano-particles within the brush, $\alpha$, as a function of $C_{\text{azo}}/C_{\text{Au}}$. $\alpha$ is the fraction of the brush surface area eclipsed by projecting the cross-sectional area of the nano-particles onto it. For instance, $\alpha = 1$ means that if the gold nano-particles would form a monolayer within the brush, the coverage of this monolayer would be 100%.

With increasing the $C_{\text{azo}}/C_{\text{Au}}$ ratio the number of absorbed gold nano-particles decreases by almost a factor of three (Fig. 2b), while the amount of adsorbed surfactant increases. To compare the projected concentration of particles to the volume concentration, we calculated also the number of nano-particles contained within the volume of a cube of 100 nm side length. For instance, for brush with maximum particle concentration one can find 160 nano-particles within a volume of $10^6$ nm\(^3\). With a particle diameter of 10 nm, one can estimate that the particles occupy 8% of the brush volume. This seems to be quite a low value, however, the fact that the bulky particles are not homogeneously distributed within the brush and only slightly penetrate it results in the formation of a dense layer of nano-particles not far away from the brush/air interface (Fig. 2d). In general, we find that the particles are distributed within a slab of thickness equal to roughly a fourth of the brush height (and adjacent to the air-brush interface), as can be inferred from the TEM micrograph of the brush film cut by the FIB (Fig. 2d). These results will be further supported by the SEM experiments in the following.

After loading the brush with azo-AuNPs its topography is still flat (Fig. 3a). Here we show the results with a brush containing the maximum concentration of gold particles. One can still see a small number of gold nano-particles (white spots in Fig. 3a and b) of 2–3 nm in height that are partially embedded in the brush but still covered with polymer material as could be inferred from a measurement of the adhesion force by means of AFM. During irradiation ($E = 12.4$ J/cm\(^2\)) with an interference pattern (IP) generated of cross-propagating beams of UV light ($\lambda = 325$ nm) a surface relief grating (SRG) forms (Fig. 3b). The height and the periodicity of

**Figure 2.** (a) Dependence of the height of the PMAA brush after absorption of azo-AuNPs, as a function of $C_{\text{azo}}/C_{\text{Au}}$ ratio (black curve). The red curve shows the height of the nano-wires produced from the brushes in (b). (b) Dependence of the amount of gold nanoparticles within the brush as a function of $C_{\text{azo}}/C_{\text{Au}}$. (c) Absorption spectrum of the PMAA brush loaded with azo-AuNPs. The inset depicts a surface plasmon peak of the gold aggregates at $\lambda = 545$ nm. (d) TEM cross sectional images of the brush film obtained by cutting it using a FIB.
the stripes are \((25 \pm 5)\) nm and \(1.5\ \mu m\), respectively. We have previously shown\(^{45–47, \ 53}\) that during SRG formation the polymer chains rupture within areas where the polymer material is receding from, i.e. at the minima of the topography. To check whether chain scission also occurs in brushes loaded with gold nano-particles we exposed the irradiated brush to DMF which is a good solvent for both, the surfactant and the PMAA. Figure 3c shows the same area as in Fig. 3b after DMF treatment. An array of polymer stripes of \(w = 450\) nm in width and \(h = 300\) nm in height remains. In between the stripes one can find a thin layer of polymer material of only \(5\) nm in thickness. Apparently, during irradiation part of the polymer chains are ruptured and during the treatment with DMF, the ruptured chains are removed leaving behind the nano-bush like structure. The height of the stripes formed is smaller than the initial brush height. Exposing the non-irradiated photosensitive polymer brush to a good solvent results in a partial desorption of the surfactant molecules. As a consequence, the height of the brush decreases, in the present case from \(400\) nm to \(300\) nm.

Applying irradiation with larger intensity, we were able to produce nano-wire like structures, i.e. stripes which are not covalently connected (i.e. free standing) to a surface. Indeed, as can be seen in Fig. 4 nano-wires are displaced from their initial position just by capillary forces. The nano-wires are formed during irradiation with polarization interference pattern at irradiation intensity of \(20\ \text{J/cm}^2\). Starting from the periodicity of \(2.5\ \mu m\) the nano-bush like structure (see for schematic representation Fig. 5g) is formed even at high irradiation intensity \((20\ \text{J/cm}^2)\).

The width and height of the nano-wires remain unchanged after exposure to DMF solution. This indicates that the nano-wires consist of the cross-linked polymer chains filled with AuNPs. The possible mechanism of cross-linking could be a recombination of two radicals of ruptured chains. During irradiation, the opto-mechanical stress generated within the brush results in scission of covalent bonds that right after scission form shorter chains with terminal free radicals. The mutual recombination of nearby radicals may lead to cross-linking within the polymer shell of the nano-wires, explaining their stability.

The width and the periodicity of the nano-wires can be varied by adjusting irradiation conditions such as optical periodicity of the interference pattern. Figure 5(a–e) show four nano-wires of the same height but differing in the width: \(320\) nm, \(400\) nm, \(580\) nm, \(780\) nm and \(1.2\ \mu m\). The length of the nano-wires is controlled by the size of the laser spot and can be as large as several centimeters implying very high aspect ratio. The number of the produced nano-wires during one irradiation process is also limited only by the size of the laser spot and can be calculated as the size of the laser spot divided over the periodicity of the interference pattern, for instance, with laser spot diameter of \(1\) millimeter and optical periodicity of \(1.5\ \mu m\) one obtains in one irradiation step ca. \(666\) nano-wires.

The nano-wires are uniform in height and width along the whole length and no gold nano-particles are found at or close to the surface of the nano-wires (Figs 3 and 4). The AuNPs are immersed within the nano-wires as can be seen in Fig. 6. To visualize gold nanoparticles within the nano-wires, we gently etch the upper polymer layer with a focused electron beam, simultaneously recording the scanning electron microscopy images. The nano-particles are distributed only within the nano-wires as a single specimen (Fig. 6d) as well as aggregates. The side view shows that the nano-particles do not penetrate the whole film and are situated in the upper fourth of the brush (Fig. 2d). The nano-wires, however, are separate independent entities with the nano-particles well embedded in the polymer, as can be inferred from Fig. 6c where one nano-wire filled with gold particles is removed from its initial position and accidentally placed by capillary forces across the others.
In the very same way one can load the brush with any small molecule or particle. The requirement is that the loading substances interact with azobenzene containing surfactant and become both photosensitive and positively charged. For instance, Fig. 7 shows the polymer nano-wires loaded with gold nano-rods. The samples were prepared using the same procedure as described above.

In the next example, we have functionalized the nano-wires with quantum dots (QDs) (Fig. 8). In this case, it was possible to produce nano-wires emitting at 610 nm.

Conclusions
We report on a mass production of spatially ordered, flexible, free-standing nano-sized wires of macroscale length filled with different types of nano-particles. The process is based on the opto-mechanical scission of chains of photosensitive polymer brushes upon irradiation with UV-interference patterns. The polymer brushes were turned photosensitive by loading with surfactant containing azobenzene. Upon irradiating the photosensitive polymer brushes with the interference pattern, a substantial mass redistribution of the polymer material occurs resulting in a formation of a relief grating. Along with the polymer film deformation a rupturing of the polymer chains takes place. During treatment of the irradiated brush with a good solvent, the chains wind up in wires of nano-scale thickness and a length of up to several centimeters. We have shown that the process of nano-wire formation proceeds in the same way when nano-particles of different structure, e.g. gold nano-spheres, nano-rods, quantum dots, are dispersed within the film prior irradiation. The final arrangement is similar to a core-shell geometry, where nano-particles were mainly found in the core region and the polymer is forming a dielectric jacket.

Materials and Methods
Synthesis of poly(methacrylic acid) brushes. Polymethacrylic acid (PMAA) brushes were synthesized using a surface initiated free radical chain polymerization of methacrylic acid utilizing a “grafting from” technique on a silicon wafer. PMAA chains grafted on the surface are schematically shown in Fig. 1. The brush sample was divided in several pieces of equal size, which then were loaded with different amount of nano-particles.

To integrate complexes of gold nano-particles (AuNPs), the brush was placed in a mixture of water suspension of AuNPs (d = 10 nm) with surfactant solution of different ratios for 30 minutes followed by drying with nitrogen flow.

Synthesis and characterization of azobenzene containing surfactants. Azobenzene containing trimethylammonium bromide surfactant (Fig. 1) was synthesized as described elsewhere. The surfactant was
dissolved in water (MilliQ) to different initial concentrations and kept in the dark for several days to ensure complete relaxation to the trans-configuration. Then, the solutions were further diluted to an appropriate concentration for the complex preparation. All experiments were carried out under yellow light to avoid premature isomerization of the surfactant.

**Nano-particles.** Gold nanoparticles (AuNPs) of diameter (10 ± 3) nm were generated by laser ablation. At basic pH values the nano-particles are negatively charged as measured by a Zeta-sizer. The characteristic absorption peak of AuNPs in water originating from the excitation of surface plasmon resonance is λsp = 523 nm. Quantum dots eFluor® 650 nanocrystal (CdSe) were purchased from eBioscience. Gold nano-rods were synthesized as described in ref. 56.

**Optical set-up for generation of interference patterns.** The Lloyd’s mirror scheme with He-Cd laser (Kimmon) operating at λ = 325 nm (total power of ~3 mW) was used for UV interference lithography. The periodicity of the interference pattern is given by \( d = \frac{\lambda}{2\sin(\theta)} \), where \( \lambda \) is the wavelength of the incident light, and \( \theta \) is the angle between the incoming laser beam and the mirror surface. The irradiation time was 15 minutes.

**AFM measurements.** Atomic Force Microscopy (AFM) (Nanoscope V, Veeco) was used to measure the height of the brushes. For thickness measurements the brushes were scratched by a glass pipette to remove the material and the step height between the top of the brush and the carried substrate were measured from the AFM cross-section analysis. Scanning electron microscopy (SEM) measurements were performed using an Ultraplus 4061 microscope (Zeiss). Cross sections for investigation by transmission electron microscopy (TEM) were prepared by cutting and thinning thin lamellae using a Ga focussed ion beam (Zeiss Crossbeam 1540 EsB). Bright field images were obtained in a Zeiss Libra 200 FE operated at 200 kV accelerating voltage. The in-column Omega type energy

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**Figure 5.** AFM micrographs of the polymer nano-wires filled with gold nano-particles (a–d). Four different periodicities and diameter of the nano-wires were obtained by adjusting the periodicity of the optical interference pattern. (e) At larger periodicity, not all polymer chains are ruptured, but a small amount stays still bound to a surface forming “nano-bushes” filled with gold nano-particles. (f) Scheme of the nano-wires, which can be formed during irradiation with polarization interference pattern. (g) Scheme of the “nano-bushes”; i.e. nano-structured polymer brushes. In both scheme yellow spots represent AuNP.
Figure 6. SEM micrographs of the polymer nano-wires filled with gold nano-particles. The gold nano-particles can be clearly identified within the nano-wires by their strong white contrasts. One nano-wire removed from its initial position during treatment with DMF is shown in c (marked by white arrow).

Figure 7. SEM micrographs of the polymer wires filled with gold nano-rods. (a) The gold nano-rods are randomly situated within the nano-wires. (b and c) Larger overview of the ensemble of many nano-wires.
filter was used to exclude most of the inelastically scattered electrons to improve image contrasts (ZLF, zero-loss filtered imaging). UV absorption spectroscopy. UV-Vis spectra were obtained using a Cary 5000 UV-Vis-NIR spectrophotometer (Varian Inc.). The Zeta-potential was measured with a Zetasizer Nano ZS (Malvern Instruments Ltd.) at a scattering angle of 173°.

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Acknowledgements
We thank the V.W.- Stiftung for the financial support. We are grateful to Prof. A.V. Kabashin for providing us with gold nano-particles and to Dr. Qianling Cui for preparing the gold nano-rods.

Author Contributions
Conceptualization: S.S.; investigation: N.L., A.K., M.W.; resources: M.B.; writing—original draft: N.L., A.K., S.S.; writing—review and editing: S.S., M.W., M.B.; funding acquisition: S.S.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

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