Placement of conjugated oligomers in an alkanethiol matrix by scanned probe microscope lithography

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We report the in situ replacement of conjugated molecules in an insulating matrix by scanned probe microscope lithography. High yield, programmable patterning of a self-assembled monolayer of dodecanethiol was performed by applying voltage pulses from a scanning tunneling microscope. Conjugated oligomers were observed to be subsequently chemisorbed onto the patterned sites.

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Scanned probe microscopy offers a method for manipulating and modifying materials at the atomic level. Preliminary work has been done utilizing self-assembled monolayers (SAMs) as a substrate. In this letter, we demonstrate reproducible in situ scanning tunneling microscope (STM) lithographic SAM patterning and subsequent replacement of conjugated molecules in an insulating matrix at selectively patterned locations. A schematic of the process is shown in Fig. 1.

The initial substrate is a gold surface with an adsorbed SAM. An annealed Au/Cr (250 nm/50 nm) film on glass exhibited a Au(111) surface with terraces of up to several hundred nanometers with herringbone reconstruction. A dodecanethiol SAM was deposited from solution at room temperature. Subsequent mounting in a Teflon liquid cell of an ambient STM chamber allowed for immersion in 1,4-dioxane (Aldrich) solvent for the replacement experiments.

A Park Scientific [Universal SA1] tube-scanning ambient STM was used for both imaging and lithographic patterning. Tips were prepared from Pt/Ir (70%/30%) wires and then coated with epoxy leaving the end exposed. Images were taken in constant current mode with 1.5 V tip bias ($V_t$) and approximately 15 pA set-point tunneling current ($I_t$).

To perform lithography, we applied square voltage pulses to the SAM substrate with pulse amplitude ($V_p$) varying from 1.8 to 3.6 V and pulse duration ($T_p$) from 0.5 μs to 0.5 s. A constant dc bias voltage ($V_b$) of 0.1 V was kept on the STM tip during the voltage pulses. A larger time constant (relative to the pulse width) in the feedback electronics is used to both maintain the tip at a constant height and avoid tip crashing. Figure 2 is an image of a sample after three consecutive voltage pulses with $V_p$ of 3.0 V and $T_p$ of 0.5 s. The tip was biased at $V_t$ of 0.1 V with $I_t$ of 0.5 nA. To have a better view of the profile of the patterned structures (pits), we invert the image by plotting the depth of the pits as positive $z$. The diameters and depths of the patterned structures are approximately 10 and 1.4 nm, respectively. The pit depth is the length of dodecanethiol SAM to within the measurement resolution of the STM system. A lithographic positioning program was used to move the tip to obtain these distinct, spatially separated patterns.

We have studied the patterning yield of this STM lithography as a function of pulse voltage and pulse width (Fig. 3). The yield was defined as the percent of successful patterned structures (out of 50 pulses). Under appropriate pulse conditions patterns were produced with over 90% yield when positive pulses (sample positive with respect to the tip) were applied. There is an apparent threshold voltage for pattern formation [Fig. 3(a)]. Greater tip sample distance required higher threshold voltage. At pulse voltages above 3.4 V pits deeper than 1.4 nm were produced, suggesting that some of the underlying gold was removed. Pulse duration [Fig. 3(b)] was varied at various pulse voltages for fixed tip-sample separation. Reproducible patterns were created at pulse widths greater than about 0.1 s. We used positive pulses to remove the SAM in this work since we observed that negative pulses created mounds which may be the result of material deposition from the tip.

![Fig. 1. Schematic of the lithographic patterning and replacement of conjugated molecules in an alkanethiol matrix: (a) normal STM imaging of the SAM surface with tip bias $V_t$; (b) SAM removal by applying a pulse $V_p$ to the substrate; (c) the same as (b) in solution; (d) insertion of conjugated molecules on the patterned site.](image-url)
Subsequent to dodecanethiol removal, we introduced conjugated molecules into the liquid cell for "decoration" of the patterned sites. The conjugated molecule used is 2'-ethyl-4:1'-ethynylphenyl-4':1''-ethynylphenyl-1,4''-thioacetylbenezene (abbreviated as A). Upon addition of NH$_4$OH, the protective acetyl endgroups in A are removed and cause conversion into the thiolate form (abbreviated as B), a convenient method for utilizing oxidatively unstable dithiols and which have been shown to self-assemble onto Au. The mechanism for replacement adsorption is that the thiol group of the rigid rod aromatic, $\alpha$-w-dithiol binds to the exposed surface; at the concentration used (10 $\mu$M), one can estimate that there are approximately 40 collisions per second of molecule B with the exposed Au surface.

To demonstrate replacement, we added 10 $\mu$M of A in 1,4-dioxane along with several microliters of NH$_4$OH to the liquid cell under slightly positive Argon pressure. Lithography was performed on the dodecanethiol surface while in solution. The molecule B is similar to one previously investigated in which conductivity higher than alkanethiol was observed, resulting in bright spots in the STM image. Shown in Fig. 4(a) is an STM image after three spatially separated pulses on the SAM surface. The left of the image shows two bright peaks (indicating adsorbed molecules); the third pulsed site is a patterned pit that remained unadsorbed. Figure 4(b) is an STM image scanned minutes later showing adsorption at the third peak. The widths of the peaks are approximately 10 nm indicating the insertion of a bundle of approximately 400 B molecules. The height is about 0.7 nm above the background, which is consistent with the known length of B and dodecanthiol molecules of 2.2 and 1.4 nm, respectively. Shown in Fig. 4(c) is a defined grid pattern and Fig. 4(d) an image after chemisorption of the B molecules onto the pattern (as well as random adsorption of the molecules at defect sites). Several scans over one hour showed no pattern degradation.

We have demonstrated the controlled insertion of conjugated molecules into a self-assembled monolayer, providing a general method for generating intermixed SAMs of arbitrary shapes and compositions. The sharpness and regularity of the STM tip should play an important role; we suggest that future work utilizing a carbon nanotube STM tip might implement single molecule replacement.

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