Fabrication of Patterned Fluorescence Images in Electrospun Microfibers

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Recently, generation of patterned fluorescence images based on the "precursor approach" has gained much attention. The concept of the 'precursor approach' is to use different electronic properties between the protected and unprotected forms. For example, a dye molecule is non-fluorescent when the key functional group of the dye molecule is protected with a protecting group. If the protecting group is removed under photoinduced chemical transformation, the fluorescence can be regenerated, allowing patterned fluorescent images in the polymer film by selective removal of the protecting group in the exposed areas. The "precursor approach" allows rapid and cost-effective generation of patterned images in one step, without the need for additional wet developing processes.

We have previously reported that the fluorescent quinizarin molecule can be readily converted to a nonfluorescent protected molecule \( t\)-BocQ by introduction of acid-labile tert-butoxycarbonyl (\( t\)-Boc) groups to the quinizarin hydroxyl moieties (Scheme 1). When the \( t\)-Boc groups were selectively removed by photogenerated acids, yellow fluorescent patterns were appeared in the UV exposed areas. Thus, the nonfluorescent \( t\)-BocQ precursor molecule was transformed to unprotected quinizarin by the photoinduced chemical transformation.

The vast majorities of the fluorescence patterns generated based on the "precursor approach" have been fabricated in polymer film. During the course toward the development of fiber-based chemosensor systems, we have discovered a new strategy for the fabrication of patterned fluorescence images in microfibers. The key strategy and procedure employed for the fluorescence patterning is schematically presented in Figure 1. A viscous chloroform solution containing the precursor molecule \( t\)-BocQ (1.3 wt%), poly(methyl methacrylate) (PMMA) (19.6 wt%), and a photacid generator (PAG), triphenylsulfonium triflate (0.6 wt%) is placed in a syringe. A high voltage (18 kV) is then applied to the metal syringe needle causing ejection of a charged polymer jet from the polymer solution. The microfibers are collected on the surface of a grounded aluminum plate (15 cm from the needle). A scanning electron microscope (SEM) image confirms the formation of microfibers (Figure 1). Photomasked UV irradiation of the electrospun fiber should generate strong acids from the PAG in the exposed areas. The photochemically produced strong acids should promote catalytic deprotection of the \( t\)-Boc groups from the precursor molecules, allowing patterned fluorescence images in the polymer fibers.

In order to investigate the feasibility of fluorescence patterning, the polymer fiber mats obtained as described above were irradiated with 254 nm UV light through a photomask for 2 min. The UV-treated fiber mats were then placed on a hot plate (100 °C) for 1 min for a post-exposure...
We also explored fluorescence patterning in a single electrospun fiber. If patterned images can be created photo-lithographically in a single fiber, the strategy can be potentially employed to introduce other functional groups into the polymer fiber. In order to investigate the feasibility of fluorescence patterning in a single polymer fiber, the t-BocQ containing single electrospun fiber was placed under a photomask and was subjected to the UV irradiation-PEB process. As displayed in Figure 3, yellow fluorescence patterns can be clearly seen under a fluorescence microscope.

In order to measure the yield of the t-Boc protected quinizarin to its unprotected quinizarin in the electrospun fibers, the precursor-embedded fiber mat was irradiated with UV light for 2 min without using the photomask. The resultant fiber mat was dissolved in chloroform and the UV-visible spectrum of the chloroform solution was recorded (Figure 4a). To this chloroform solution was added trifluoroacetic acid (20 %, v/v) and the solution was stirred for 60 min to remove the residual t-Boc groups. After concentration in vacuo, the residue was dissolved in chloroform and UV-visible spectrum was monitored (Figure 4b). The absorption difference at 480 nm allows calculation of the yield of quinizarin and is found to be 61%. Thus, fairly significant amounts of the precursor molecules were converted to quinizarin in the electrospun fibers.

In summary, we have developed a new and straightforward strategy for the generation of patterned fluorescence images in microfibers. By employing the photolithographic technique, we were able to create patterned fluorescence images in polymer fibers based on a t-Boc protected quinizarin as a precursor molecule. Considering the significance of the fluorescence patterning technology in the field of sensor, molecular switch as well as memory device and display areas, the results described above should find great utility in those areas. In addition, photolithographic method for patterned images in a single microfiber should be useful for selective introduction of functional groups to the polymer microfibers.

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References
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