Poly (N-Isopropylacrylamide) Microgel-Based Etalons for Optical Sensing

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Introduction

Poly (N-isopropylacrylamide) (pNIPAm) is one of the most completely studied "smart" polymers due to its unique reversible thermo-responsivity. That is, when pNIPAm in water is heated > ~31°C, it transits from a random coil to a globule conformation; this transition is reversed when T < ~31°C. This conformational change is accompanied by water exchange process. When pNIPAm undergoes the coil to globule transition, water is expelled, while water is "sorbed" when the polymer undergoes the opposite process. Researchers have synthesized various structures from pNIPAm, e.g., block [1], random [2], and star [3] polymers, as well as crosslinked hydrogels, and ultimately colloidal-stable particles (microgels [4], and nanogels [5]). Of these, microgels are generally defined as cross-linked networks of water-soluble polymers. The special network structure leads microgels to swell with water. Various optical sensors have been designed with stimuli-responsive hydrogels/ microgels. For example, Asher et al. [20,21] developed three dimensional (3D) polymerized colloidal crystal array hydrogel sensing materials. Composed of stimuli-responsive hydrogels, these photonic materials (PMs) change their volume in response to external stimuli. Additionally, Asher et al. [22] prepared close-packed two dimensional (2D) poly styrene particle arrays by self-assembly of spreading particle monolayers on mercury surfaces. Color alterations of high diffraction efficiency 2D photonic crystals indicate environmental changes, and hence allowing them to be developed for sensing. PMs fabricated by a number of groups [23-27] have refractive index order periodicity in 3D and 2D, however PMs can also have structural periodicity in one dimension (1D), yielding their own interesting optical properties and colors as well. In this submission, we review our recent work on pNIPAm microgels based the 1D PM structure [28-33].

1D PMs are generally composed of periodically arranged layers of alternating refractive index in only one dimension (e.g. x, y, or z axis only) such as Bragg mirrors, interferometers, waveguides, and Fabry–Pérot etalons; the reflected/refracted light at the interface of each layer leads to constructive/destructive interference, resulting in color. Recently, Fabry–Pérot etalons (simply etalons) were investigated in our group [28-34] and others. [35-37] Etalons are composed of a dielectric cavity confined between two reflective surfaces (Scheme 1). After entering the etalon, light resonates in the dielectric cavity, and therefore produces light interference. This interference gives rise to specific wavelengths of light that are reflected. Interestingly, the material can exhibit visible color tunability if the dielectric layer is of the appropriate thickness, and can be made to change thickness. Furthermore, the refractive index of the dielectric layer can also affect the visible color. This is accounted for by equation (1):

\[ \lambda m = 2nd \cos \theta \] (1)

where the specific wavelength maximum of the peak (\( \lambda \)) depends on the peak order (m), refractive index of the dielectric(n) and the spacing between the mirrors (d), as well as the angle of incidence(\( \theta \)). [38] In our etalons, Au and pNIPAm-based microgels serve as the mirrors and the dielectric layer, respectively.

Fabrication of pNIPAm Microgel Based Etalons

To fabricate an etalon, a Au coated glass cover slip (2 nm Cr was used as an adhesion layer followed by 15 nm Au) was made using a thermal evaporation system (New Windsor, NY) at a rate of ~1 Å s\(^{-1}\) (Cr), and ~0.2 Å s\(^{-1}\) (Au), respectively. Then, the Cr/Au coated substrates (simply Au substrates) were annealed at 250°C for 3 h, followed by rinsing with ethanol, dried with \( \text{N}_2 \) prior to use. After pNIPAm based microgels were deposited on Au substrate, followed by deposition of...
an additional 2 nm Cr/15 nm Au layer using the aforementioned instrument. Herein, we desire “perfect” microgel monolayers on the Au substrate. To accomplish this, a so-called “paint-on” protocol was developed by our group. As seen in Scheme 2, the paint-on protocol requires centrifuging a solution of microgels (microgels synthesized according to [30]) until they were concentrated at the bottom of the centrifuge tube. This takes 30–60 min, depending on the diameter of the microgels.
Following centrifugation, the supernatant solution was removed, and a 40 µL aliquot of the concentrated microgels was deposited onto the Au substrate at 30°C. This aliquot was then spread toward each edge using the side of a micropipette tip until the microgels covered the entire Au substrate. The spreading continued until the microgel solution was too viscous to spread over the surface. At that point, the microgels were allowed to dry completely on the substrate for 2 h at 35°C. After 2 h, the dry film was rinsed with copious amounts of DI water and soaked in DI water overnight to remove microgels not bound directly to the Au. This method yields an extremely uniform etalon both spectrally and visually. [29] Moreover, homogeneity of the response of the etalons was also significantly enhanced from spot to spot using our painting protocol. In addition, this painting method can be applied to coat a variety of different microgels on a variety of surfaces [31].

Application

To investigate the thermo- and pH-responsivity of the etalon, we secure it in a special chamber, Figure 1. The chamber allows us to directly measure the reflectance spectrum from the etalon, and allows us to very precisely control the temperature of the solution the etalon is in. Using this setup, we can easily monitor the wavelength maximum (λmax) in the reflectance spectrum for a given peak order as a function of temperature and pH. From Figure 2, we see that when the etalon is immersed in pH 3.0 solution, the λmax shows a blue-shift of approximately 300 nm over the given temperature range. This shift is attributed to the collapse of the microgels at high temperature bringing the Au mirrors closer to one another shifting λmax accordingly based on equation 1. Moreover, the most dramatic shift occurs between 29 and 35°C, which correlates with the behavior of the microgels in solution. At pH 6.5, this behavior is suppressed due to Coulombic repulsion deprotonated AAc in the microgel structure. This behavior shows the sensitivity of etalons toward temperature and pH changes.

To assess whether etalons can potentially be developed for point-of-care (POC) diagnostic applications, we used glucose sensitive etalons for the proof-of-concept. [32] Glucose sensitive microgels were synthesized by modification of the microgels with aminophenylboronic acid (APBA) APBA-functionalized pNIPAm-based microgels have been fabricated, and we studied their spectral responsive properties with glucose. As seen in Figure 4, there is a significant spectral red-shift (~134 nm) in the presence of 3 mg/mL solution of glucose (pH 9 carbonate buffer) and the majority of the spectral shift occurs within 30 minutes of glucose introduction, therefore yielding a visible color changes (Figure 5). Given that APBA can bind to other diols as well, APBA modified microgel etalons can exhibit a colorimetric and spectral response to other biologically relevant molecules as well, thus, further specificity will need to be built in for future biosensing efforts.

Thus far, our group and others have focused on investigating the color tunability of a complete PM [29-33]. To move the field forward, one challenge is modulating the color of the material in specific regions, while not affecting the optical properties of another, spatially isolated, region. Recently our group investigated the ability of the etalon to be solvated, and the color modulated, in spatially isolated regions. Specifically, pH 3.0, 4.0, 7.0 solutions were deposited between three separate reflectance probes and an etalon, and the reflectance spectra for the different regions were collected as a function of temperature. As seen in Figure 6, the peak for the spot at pH 3.0 significantly blue-shifts with increasing temperature, while the spots at pH 4.0 and 7.0 shift minimally over the same temperature range. Additionally, the colors of the individual spots are visibly distinct, and change independently as a function of temperature. For example, the pH 3.0 spot visibly changes color from green to red as the temperature increases, while the others do not significantly change. Although three spots are close to each other, independent responsive behavior is observed as expected. This independent responsive behavior on a single etalon enhances the utility of the materials for sensing applications.

So far, our pNIPAm microgel based etalons also show potential applications in humidity sensing, pressure sensing, and organic solvent titration (data not shown).

Conclusions

With the fascinating properties of pNIPAm microgels and the invention of the “paint-on” protocol, it is quite simple to fabricate a variety of pNIPAm microgels based etalons. Given that tunable “lattice” spacing dominates colors in ordered materials, pNIPAm microgels based
etalons can be made to respond to pH and temperature, changing color visually and spectrally. APBA functionalized pNIPAm microgels rendered etalons responsive to biologically molecules (i.e., glucose), thus the color of the etalon changes in the presence of biological analytes. Also, the independent tunable optical properties at different regions on a single etalon are realized. Even though pNIPAM microgels have been widely studied, pNIPAm microgel based etalons require further research. Combining promising physical and chemical properties of pNIPAm microgels with optical properties of etalons, pNIPAm microgel based etalons are likely to become quite appealing biosensors in the near future.

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