Formation of Cocrystal Nanorods by Solid-State Reaction of Tetracyanobenzene in 9-Methylanthracene Molecular Crystal Nanorods

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Introduction

There is currently great interest in both the synthesis and characterization of one-dimensional nanostructures, commonly referred to as nanorods or nanowires. Nanowires composed of inorganic compounds have been grown using a wide variety of methods, and there has also been considerable effort devoted to growing multicomponent or doped inorganic nanostructures.\(^1\) Parallel to efforts in inorganic materials, there has also been progress in making one-dimensional organic nanostructures through both self-assembly methods\(^2\)\(^–\)\(^5\) and templated growth.\(^6\)\(^–\)\(^9\) For organic nanostructures, the ability to incorporate two different molecular species into a single nanorod would be especially beneficial, since molecular cocrystals can have unique properties relative to single component molecular crystals.\(^10\)

Usually such cocrystals are formed by slow precipitation from a mixed solution or slow solvent evaporation, resulting in large crystals. The first examples of nanococrystals were achieved using reprecipitation in conjunction with sonochemistry.\(^11\)

Milling (wet or dry)\(^12\)\(^–\)\(^14\) can also be used to grow small-scale cocrystals but at the expense of size and shape control. An alternative approach is to first grow a neat crystal of one molecular component with a specified size and shape and then react it with the second component to make a cocrystal that retains the morphology of the original crystal. This approach is not widely used for growing bulk cocrystals due to the scarcity of solid—solid diffusion reactions and problems with strain and crystal disintegration. While surface modification is quite common, chemical transformations occurring within nanostructures are usually limited to rearrangements of the species that are already present, for example, thermal elimination to form new crystal types.\(^15\)

Recently, it has been observed that crystalline organic nanostructures undergo various types of photochemical transformations and remain intact, while under identical conditions micron scale or larger crystals will fragment.\(^16\)\(^–\)\(^19\) This ability to withstand the strain of phase separation and crystal reconstruction is most likely related to the large surface-to-volume ratio of these small structures, which permits strain relief to occur at the surface in a facile manner.\(^20\)

An important question is whether the robustness of organic nanostructures with respect to photochemical transformations, which require only the penetration of photons, extends to chemical processes that require the penetration of molecules into the crystal lattice. If this proves to be the case, then a new class of structure-preserving chemical transformations for organic nanostructures will have been identified.

In this paper, we demonstrate that it is indeed possible to grow molecular crystal nanostructures by taking advantage of the crystalline nanorods’ ability to retain their morphology during chemical reactions. We examine the reaction of 9-methylnanthracene (9-MA) crystalline nanorods with 1,2,4,5-tetracyanobenzene (TCNB) to form a cocrystal with charge-transfer (CT) properties. 9-MA is known to be reactive in its crystalline state,\(^21\)\(^22\) and the current reaction, shown in Figure 1, involves...
a crystal-to-crystal transformation that occurs within the channels of a porous anodic aluminum oxide (AAO) template. In the case of 9-MA/TCNB cocrystal the driving force behind the solid–solid diffusion reaction is the formation of the thermodynamically stable ground-state charge-transfer complex. 9-MA is a good electron donor to the strong electron acceptor TCNB. The formation of ground-state charge-transfer complexes is typical for electron-rich planar polycyclic aromatic hydrocarbons (PAH) and planar electron acceptors such as TCNB.23-24 Organic charge-transfer nanorods have been previously synthesized using single component templated growth,25 electric-field methods,26-28 and photocrystallization.29 Our new approach demonstrates that it is possible to completely react a homogeneous nanorod with a second molecular species without disrupting its morphology. Furthermore, we show that this approach only works with nanostructures: attempting to apply the same method to macroscopic crystals leads to morphology changes and eventually fragmentation. The present work demonstrates how organic nanostructures provide new opportunities for doing solid-state chemistry in a controlled way, for example, the growth nanoscale molecular cocrystals with well-defined shapes and sizes.

Experimental Procedures

Synthesis of 9-MA/TCNB Charge-Transfer Nanorods. Twelve milligrams of 9-methylanthracene (9-MA) was dissolved in 0.1 mL of dry tetrahydrofuran (THF). The solution was deposited on the surface of an AAO template (Whatman Anodisc-13, 200 nm pore diameter, average membrane thickness of 60 µm, membrane diameter of 25 mm, with polypropylene supporting ring). The loaded AAO template was placed on a homemade Teflon holder that suspends it from the polymer support ring. The holder was then placed in a 16 oz glass jar containing 2 mL of THF and a 1 inch ring of filter paper to disperse the solvent vapors. The jar was tightly covered and a long needle was inserted through the cover until it touched the bottom of the jar. Another needle was inserted in the cover and was coupled with a gas bubbler. Argon gas was gently introduced into the jar at a rate of 1 bubble every 2 s. The slow room temperature vapor solvent annealing causes the 9-MA in the AAO template pores to form single crystalline nanorods. After all the solvent had evaporated, the solvent annealed AAO template was polished using 2000 grit sandpaper to remove excess 9-MA from the surface. Twenty milligrams of 1,2,4,5-tetracyanobenzene (TCNB) was sonicated in 20 mL of distilled water to form a suspension (TCNB is slightly water soluble). The 9-MA loaded AAO template is immersed in the AAO template pores to form single crystalline nanorods. After all the solvent had evaporated, the solvent annealed AAO template was washed away with distilled water and air-dried. The AFM tip was lowered back to the same approximate region and scanning commenced.

Absorption and Fluorescence Spectra. We could not directly measure the absorption spectra of the 9-MA and 9-MA/TCNB nanorods because of small sample volumes and scattering. In most cases, the fluorescence excitation spectrum is a good representation of the absorption. Both the emission and excitation spectra were collected using front-face geometry in a Spex Fluorolog fluorimeter. The nanorods were left in the alumina template to provide a flat, low-scattering surface normal to the exciting beam. The fluorescence spectra were collected using 365 and 500 nm excitation wavelengths for the 9-MA and 9-MA/TCNB nanorods, respectively. The excitation spectra were measured using detection wavelengths of 550 and 650 nm for the 9-MA and 9-MA/TCNB nanorods, respectively. Note that care was taken to minimize the exposure of the 9-MA samples to UV light, since 9-MA is known to undergo a solid-state [4 + 4] photodimerization.30 The 9-MA solid fluorescence was observed to slowly disappear under UV irradiation, in contrast to the 9-MA/TCNB cocrystal, which was completely photostable.

SEM and TEM. For the SEM measurements a drop of the aqueous CT nanorod suspension was placed on a copper tape and fixed on an SEM stub. The water was gently evaporated leaving behind dispersed bundles of nanorods. The SEM stub was sputter coated with Pt/Au for a cycle of 60 s. The SEM stub was placed inside a scanning electron microscope (XL30 FEG) and the data were collected using a 10 kV electron beam. Transmission electron microscopy of the nanorods was performed using a FEI-PHILIPS CM300 model. The specimen was prepared by depositing a drop of the aqueous 9-MA/TCNB nanorod suspension on the graphite grid sample holder and gently dried. Note that we were not able to perform electron microscopy measurements on pure 9-MA nanorods because the 9-MA rapidly sublimes under high vacuum and electron beam heating.

Powder X-ray Diffraction. All measurements were performed using a Bruker D8 Advance diffractometer with Cu Kα radiation (40 kV, 40 mA, λ = 1.5418 Å). In all cases a background scan of the polycarbonate sample holder was first performed. Then hexane extracted CT nanorods were scattered on the sample holder (without moving it) and scanned over a wide angle, typically between 5 and 70 deg at a scan rate of 0.1 deg/s. The calculated powder XRD of the CT nanorods was generated using the Mercury 1.4.2 program (Supporting Information).

Results and Discussion

The molecular cocrystal nanorods were formed using a variant of the solvent-annealing process developed by us previously to grow nanorods out of a variety of molecules.31 The procedure is outlined in Figure 2. After solvent annealing in the pores of an AAO template, 9-MA molecular crystal nanorods are formed. Excess 9-MA on the template surface is removed by polishing. Then an aqueous suspension of TCNB is deposited on the template and the sample is left in a sealed container for a period of several weeks. During this time, the TCNB slowly diffuses along the length of the channel, converting the 9-MA crystal into a 9-MA/TCNB cocrystal. The conversion is rarely 100% along the whole length of the rod, as judged from the fluorescence images. Even after a period of three weeks some unreacted 9-MA is left at the tip or along the length of the
If the reaction is interrupted, one can observe a sharp
transition in the XRD pattern. In cases where partially converted 9-MA and CT nanorods were investigated by TEM, electron beam heating would sublime out the unreacted 9-MA part of the nanorod, leaving behind a submicron boundary of the cocrystal regions. Thus, the reaction seems to occur via a well-defined diffusion front that slowly advances along the rod. The larger volume of the cocrystal results in extra CT material that is extruded from the pores and deposited on the template surface. In general, this red solid on the surfaces is mixed with unreacted TCNB and attempts to quantify the amount of extruded cocrystal were unsuccessful. It should be emphasized that this material must contain both 9-MA and TCNB, since its optical properties are consistent with the CT complex described below. Thus, 9-MA is displaced by the TCNB within the template, as expected.

The AAO template can be dissolved in a strong acid solution and the properties of the isolated rods examined in detail. The structural properties of the nanorods have been characterized using a variety of methods. The cocrystal rods are more fragile than the 9-MA rods (just as the bulk cocrystals are more brittle than bulk 9-MA crystals) and tend to break under prolonged electron beam illumination, but their rod morphology is clearly preserved, although the surface appears slightly roughened, perhaps due to residual strain induced by the reaction (see discussion below).

Further evidence for the crystalline nature of the rods is provided by powder XRD experiments. In Figure 4a,b, we show the powder XRD patterns of the 9-MA bulk crystals and 9-MA nanorods, respectively. The excellent correlation between the peaks is evidence of the crystalline nature of the 9-MA nanorods. The methyl units in 9-MA are arranged in a head to tail fashion with the anthracene units assuming a sandwich herringbone motif. After the nanorods are reacted with TCNB, the new motif is obtained. These new peaks line up with those from a bulk cocrystal powder obtained by either solution growth (Figure 4d) or by wet milling (Figure 4e). The relative peak intensities are different, possibly due to residual crystal alignment in the nanorod sample. Another possible complication is that in the solution-grown cocrystal the head-to-tail orientation of the 9-MA methyl groups is random, most likely because the extra separation introduced by the presence of the TCNB lessens the steric interactions between these methyl groups. This random orientation is not expected in the cocrystals grown directly from the 9-MA crystal, where the methyl orientations start out fixed in an alternating pattern. But comparison of the wet-milled and solution-grown bulk cocrystals in Figure 4d,e show no discernible difference in the powder XRD patterns between the two, suggesting that any ordering of the 9-MA methyls cannot be
discerned at this level of analysis. The important point is that the data in Figure 4, especially the narrow XRD peakwidths and good correspondence of the nanorod peaks to the bulk crystal peaks, provide strong evidence for the crystal-to-crystal nature of the reaction in the nanorods.

The change in crystal structure is accompanied by a change in optical properties. We were unable to directly measure the absorption of the nanorod samples due to strong light scattering in both transmission and reflection geometries. Fluorescence excitation spectra can be obtained more easily, but in this case self-absorption effects can complicate interpretation of the results. These effects are minimized by using front-face detection and, for excitation measurements, detecting at an emission wavelength far from the absorption edge. While we cannot rule out residual distortion of the spectra due to wavelength-dependent absorption and scattering in our solid samples, the close agreement of our excitation spectrum of 9-MA rods with absorption spectra of ultrathin spin cast 9-MA films suggests that such effects are minor. Figure 5a shows the excitation and fluorescence spectra of 9-MA nanorods, along with a fluorescence microscope image of a single rod. The structured excitation spectrum beginning at around 420 nm and the featureless excimer emission at 500 nm are both consistent with previous measurements on bulk 9-MA crystals. For the cocystals, a new excitation peak appears at around 520 nm, while the fluorescence peak is shifted to 610 nm. The red shift and broadening are typical of CT absorption lineshapes and reflects the increased coupling of this transition to environmental vibrational degrees of freedom. The spectroscopic data are consistent with a change from a neutral excited-state in the 9-MA crystal to a CT excited-state in the 9-MA/TCNB cocystal. This is expected since TCNB forms a CT complex with a variety of anthracene derivatives. The fluorescence we observe is consistent with that previously observed for both anthracene and 9-MA complexes with TCNB. The change in optical properties is immediately apparent by observing the nanorods with a fluorescence microscope, where the blue-green 9-MA rods change to the red-emitting CT rods.

We have shown how the 9-MA/TCNB cocystal nanorods can be grown from pristine 9-MA crystalline nanorods. Normally, macroscopic cocystals would be grown by precipitation from a mixed solution of 9-MA and TCNB. We attempted to use this mixed solution method for templated growth of cocystal nanorods in a single step. After solvent annealing using an equimolar mixed solution of 9-MA and TCNB, we find that no rods are formed, although bulk cocystals form on the surface of the AAO membrane. The low solubility of the CT complex in organic solvents such as THF prevents its dissolution in the solvent vapor which is necessary for crystal restructuring during the solvent annealing process. Instead of growing nanorods, we can try to use the same reaction conditions for our nanorods to transform bulk 9-MA crystals into bulk cocystals. In this case, the reaction is successful, but the structural integrity of the macroscopic crystal is lost. The TCNB diffuses into the bulk crystal, but the crystal quickly disintegrates. The effect of TCNB exposure on a pristine 9-MA crystal can be seen in Figure 6, which shows an AFM image of the crystal surface before and after exposure to a saturated aqueous solution of TCNB. The 9-MA crystal exhibits some 8–10 nm periodic ridges before exposure, but is otherwise featureless. After exposure, the surface undergoes dramatic changes, with eruptions that extend hundreds of nanometers above the surface. This type of surface roughening has been previously observed for 9-MA and other molecular crystals reacting with gases and undergoing photochemical reactions and has been ascribed to strain buildup between the heterogeneous phases within the crystal during the reaction.

It has been shown that the overall process of guest transport in a van der Waals molecule crystal to form a new intact crystals requires extensive cooperativity among the host molecules. It is possible that the presence of water aids the penetration of TCNB into the 9-MA crystal, but the fact that we could successfully make cocystals by dry milling the two solids together shows that the presence of water is not required. The surface roughening observed in our AFM experiments suggests that the formation of the 9-MA/TCNB cocystal does lead to significant strain within the crystal. Obviously some morphologies (e.g., macroscopic crystals) cannot accommodate this strain. One explanation for why the nanocrystals remain intact is that their high surface-to-volume ratios permit the strain to be alleviated at the crystal surface. While all previous examples of this size effect have centered on photochemical reactions, the present work demonstrates that bimolecular reactions can benefit from it as well. There are other factors that may also be important. Our reactions were carried out with the 9-MA rods still confined within the AAO template. It is possible that the template itself forces the nanorod to remain intact during the reaction. To examine the role of the template, we have reacted aqueous 9-MA nanorod suspensions with aqueous TCNB. 9-MA/TCNB nanorods form and remain intact even without the template, but tend to be much more heterogeneous, with 9-MA and 9-MA/TCNB regions highly intermixed. This
uncontrolled diffusion approach demonstrates that confinement is not necessary for the structure-preserving reaction in the nanorods, but it also resulted in more disordered samples and therefore was not pursued. A third possibility is that anisotropic diffusion within the nanorods helps them remain intact. The rate of molecular diffusion within a crystal is known to depend on the crystal axis,\cite{45,46} and it is possible that the crystal orientation of 9-MA in the rod, with only the ends exposed to TCNB, favors diffusion along a slow axis, which would lead to more gradual conversion and less fracture. Again, our results on the free rods suggest that even without constraints on the TCNB access to the rods, the rods can incorporate the TCNB and remain intact, albeit with less uniform reaction.

**Conclusion**

In this paper, we demonstrate a novel way to grow nanostructured molecular cocrystals. By taking templated nanorods of 9-MA and exposing them to aqueous suspensions of TCNB, we can form the 9-MA/TCNB cocrystal while retaining the structural integrity of the rods. The morphology of the cocrystal rods is characterized by electron microscopy and X-ray diffraction, while their charge-transfer nature is confirmed by optical spectroscopy. It is important to stress that this crystal-to-crystal reaction is the only route to the CT crystal nanorods, since nanorod growth from a mixed solution leads to phase separation and formation of random crystals on the surface of the template, while attempting to reproduce the reaction in bulk crystals leads to fragmentation. Use of nanostructured molecular crystals may provide opportunities for performing morphology-preserving chemical transformations that cannot be achieved in bulk crystals. Preliminary results in our laboratory suggest that TCNB cocrystal nanorods can be grown using the same method with different anthracene derivatives, including 9-anthracene carboxylic acid and 9-tert-butyl anthracene ester, as well as using different diameter templates. Our previous work has shown how single-component molecular crystal nanorods can exhibit interesting photomechanical properties, and the present work may lead to nanowires with interesting optoelectronic and electronic properties. Taken together, these results suggest that molecular crystal nanostructures provide a fruitful area for the exploration of solid state chemical reactivity.

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**Supporting Information Available:** Calculated powder XRD of the CT complex. TEM and SEM of CT nanorods. Crystal structure table of the CT complex. This information is available free of charge via the Internet at http://pubs.acs.org.

**References**


