Photochemical Control of Dark Conductivity - A New Approach to Devices Based on Molecular Crystals

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Thermal analysis of Ag(DM)\textsubscript{2}, where DM = 2,5-dimethyl-N,N'-dicyanoquinonediimine, clarified that the salt had an insulating amorphous phase (\textgeq 155\textdegree C). Characterization of this and related solid states of Ag(DM)\textsubscript{2} indicated that a photo-induced process should be essential in controlling the number of carriers and thus conduction behavior of the salt by illumination. In fact, while heating could do nothing but make the salt insulating when the sample temperature exceeded 155\textdegree C, ultraviolet-visible light illumination (\textlt 155\textdegree C) could gradually change the properties to be semiconducting with retaining the crystal lattice (average structure).

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1. INTRODUCTION

Recently, we developed a novel photochemical way of fabrication of molecular devices, which may enable simultaneous (persistent) carrier doping and photolithography\textsuperscript{1}. At present, we consider that this method is principally based on the same chemistry of silver halides photography. Instead of AgBr, a silver salt of an organic \pi-acceptor Ag(DM)\textsubscript{2}, where DM = 2,5-dimethyl-N,N'-dicyanoquinonediimine, is illuminated with ultraviolet-visible (UV-Vis) light. Due to the photosensitivity, the illuminated Ag\textsuperscript{+} ions deprive the DM species of some \pi-conduction electrons. Let us suppose here that only a part of the sample be illuminated and thus the rest (non-illuminated part) of the solid remain unchanged. One could make different parts with different conducting properties in a single crystal, \textit{i.e.} a junction structure; a fundamental form of current devices. Consequently, a single crystal of Ag(DM)\textsubscript{2} can be transformed to have a junction structure after appropriately partial illumination. Irreversibility and high spatial resolution
of photochemistry are found to be advantageous in this kind of fabrications. In order to establish and generalize this method, we should understand more detailed mechanism of the “optical doping” method. In this paper we will briefly discuss whether heat should involve and whether the crystal structure should retain during the UV-Vis illumination on Ag(DM)$_2$ based on the results of chemical, spectral and thermal analyses$^2$.

2. EXPERIMENTAL

The crystalline sample of Ag(DM)$_2$ were obtained by a reported method$^3$. Illumination was carried out in an open atmosphere at room temperature using a Hg/Xe lamp (200 W) equipped with a multimode optical fiber (8 mmΦ), a heat ray cutoff filter and a cold mirror. Also examined were dependencies on wavelengths and intensities of the incident light as well as duration of illumination. Thermal analysis (DSC and TGA) were conducted on the samples sealed in aluminum pans (7×7×0.25 mm$^3$).

3. RESULTS AND DISCUSSION

We obtained four different kinds of resultant states depending on the illumination conditions. Among them including the pristine material, let us concentrate our discussion on the following three, and tentatively call these series of states $\alpha$, $\beta$ and $\gamma$, respectively (Fig. 1).

On UV-Vis illumination around room temperature (RT), the original metallic properties of the pristine material ($\alpha$) gradually turned to be semiconducting ($\beta$). Compared with the pristine ($\alpha$) state, the $\beta$-state exhibited differed electrical behavior, Raman shifts (C=C and C=N stretching modes) and X-ray photoelectron spectra (Ag, N) with unchanged XRD pattern, elemental analyses (C, H, N) and infrared (IR) absorption spectra. When the sample temperature exceeded 155ºC during the illumination, an irreversible structural transition occurred; the dark blue crystalline sample ($\alpha$) suddenly turned to be light brown amorphous solid retaining the original clear-cut shape of each single crystal ($\gamma$). The $\gamma$-state turned out to be insulating and diamagnetic from the electrical resistivity and the magnetic susceptibility measurements. From the experimental results below, it is concluded that at 155ºC a solid state redox reaction, i.e. electron transfer, occurs between the Ag$^+$ ions and the DM radical anions, leading both chemical species to neutral states. Because the heat of reaction is considered to overwhelm that of amorphization in this case, a clear exothermic heat ($\Delta H = -126.8$ kJmol$^{-1}$) was observed in the measurements.
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Fig. 1. Scanning Electron Microprobe (SEM) photographs of different states after UV-Vis illumination on single crystals of Ag(DM)$_2$; (a) $\beta$−, and (b) $\gamma$−states, respectively. The SEM photographs of the pristine (α−) samples (not shown here) much resemble those of the $\beta$−state.

of differential scanning calorimetry (DSC) (Fig. 2). Elemental analyses and thermogravimetric and differential thermal analysis (TG-DTA) indicated that the chemical stoichiometry of Ag(DM)$_2$ was substantially retained through the structural transition. Solid state NMR ($^{13}$C) and IR absorption spectra indicated that the molecular structure of DM remained without decomposition in the $\gamma$-state. Furthermore, X-ray absorption fine structure (XAFS) indicated that the $\gamma$-state included neutral silver instead of Ag$^+$ ions. Yet the material exhibited different spectroscopic and chemical properties from a mixture of bulk silver and neutral DM species. The $\gamma$-state could not be separated into the two components by standard chemical procedures of purification. All these data suggest that the $\gamma$-state should correspond to a single phase comprised of neutral Ag and DM species.

In Fig. 2, except for the exothermic peak, only clear plateau was observed at all the temperature, which means that nothing should occur below 155°C. Accordingly, heat can only produce the totally insulating $\gamma$−state at 155°C or can not make any difference in the physical properties of Ag(DM)$_2$ below 155°C. On illumination, the pristine solid, whether it is a single crystal or powder, turns into the amorphous state all at once. The findings about the $\gamma$-state indicate that the $\beta$−state could NOT be produced by heat, because the single crystals of Ag(DM)$_2$ in the $\beta$−state are known to progressively become semiconducting in accordance with illumination$^1$. Detailed structural information of the $\beta$−state, particularly the difference from the $\alpha$−state, is evidently required, which is now under investigation.
Fig. 2. Differential scanning calorimetry of Ag(DM)$_2$ (6.600 mg). Heating rate was 3 °Cmin$^{-1}$. Identical results were obtained when the measurements were carried out in air and in dry nitrogen.

4. CONCLUSIONS

At 155°C, an irreversible crystal-amorphous transition of Ag(DM)$_2$ is found and the amorphous state is characterized to be diamagnetic and insulating, comprised of neutral silver and DM species. We did not observe an intermediate or mixed situation between the pristine and the $\gamma$-states that could be produced by heating, regardless of the heating conditions. Thus it can be concluded that a photo-induced process, instead of thermal effects, has turned out to be essential in the spectral and electrical property changes observed in the illuminated Ag(DM)$_2$. In other words, the photochemical electron transfer (redox reaction) between the Ag$^+$ and the DM species is important in the phenomena in question, and no heating effects such as lattice defects or thermal decomposition could explain the phenomena as dominant origins.

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REFERENCES

2. More detailed manuscript is now in preparation.