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Valence-differential spectroscopy of Co–Fe cyanide films

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Electrochromism is extensively investigated for practical application of display and memory devices. To develop the material, reliable information on the optical and electronic properties of the solid film is indispensable. Here, we propose valence-differential spectroscopy that can selectively extract the spectral components related to the oxidized/reduced metal site. We applied the spectroscopy to Co2+–Fe2+ and Co2+/Fe3+ cyanide films with finely control of averaged valence (δ) of the transition metal by external electric pulses. The spectroscopy revealed transition energy E, width Γ, and oscillator strength f of the spectral components related to the transition metal.

Electrochromism, especially in films of transition metal complex, is attracting current interest of material scientists because the electronic structure (color of the film) can be widely tuned by substitution of the transition metal. For example, chemical substitution of Prussian blue analog, Na0.84M[Fe(CN)6]0.713.8H2O (M is a transition metal) changes the color from red (M=Co), blue (M=Fe), and yellow (M=Ni). Recently, Gotoh et al.1 reported a simple synthesis procedure of Prussian blue nanoparticle inks, which is a breakthrough toward a practical display device.2 The Prussian blue analog consists of a cyanobridged rock-salt-type network, −NC− Fe− CN− M− NC− Fe− CN− and guest species, Na+ ions and H2O molecules, accommodated in the nanopores of the network.3 The electrochromism in this system is ascribed to oxidation/reduction of the transition metal site and subsequent outgoing/incoming of the Na+ ions.4,5 In spite of the promising functionality, there exits few quantitative reports on the optical properties of the Prussian blue analog and assignment of the optical transitions is still controversial, especially in the higher energy region. So far, Nakada et al.7,8 synthesized a series of valence-controlled Co–Fe films (Na0.84–δCo[Fe(CN)6]δ0.713.8H2O and Na1.60–xCo[Fe(CN)6]0.90x2.9H2O) and investigated δ-dependence of absorption spectra. Based on the δ-dependence, they decomposed the absorption spectra and assigned the respective spectral components. However, the decomposition of rather broad spectra has ambiguities and is model dependent. Here, we propose a more accurate and unambiguous spectroscopy, i.e., valence-differential spectroscopy, that can extract the spectral components related to the oxidized/reduced metal site.

We synthesized valence-controlled Co–Fe films, Na0.84–δCo[Fe(CN)6]δ0.713.8H2O (δ=0.14:960 nm in thickness) and Na1.60–xCo[Fe(CN)6]0.90x2.9H2O (δ=0.16:650 nm in thickness) on an indium tin oxide (ITO) transparent electrode by an electrochemical method. Chemical compositions of the prepared films were determined by inductively coupled plasma and standard microanalytical methods. These compounds belong to the face-centered cubic (Fm3m: Z=4) with lattice constant of 1.03 nm. Details of the synthesis and characterization were described elsewhere.7,8 In these films, magnitude of δ can be controlled by oxidation process of nondoped film (δ=0). Electronic configuration of the nondoped film is high-spin Co2+ (t2g5eg1) and low-spin Fe2+ (t2g6). We confirmed a linear relation between δ and total current q during the oxidization process. Infrared absorption spectra revealed that the oxidization site is Fe (Co) for the former (latter) systems (vide infra); extended chemical formulas are Na0.84–δCo[Fe(CN)6]δ0.713.8H2O and Na1.60–xCo[Fe(CN)6]0.90x2.9H2O. For convenience of explanation, we will call the films as Co2+/Fe3+ and Co2+/Fe3+.

We schematically show in Fig. 1(a) the valence-differential spectroscopy. We sandwiched an aqueous solution containing 1 mol/l NaCl between Prussian blue analog films grown on ITO transparent electrode. Central part of the right-side film, which works as a counter electrode, was removed. Magnitude of δ is finely controlled by electric pulses, and was estimated from the induced total current q. Figure 1(b) shows relative change (Δδ) in the averaged valence in Na0.84–δCo[Fe(CN)6]δ0.713.8H2O (δ=0.14) film against the number of the electric pulse, 0.25 V in height and 1 s in duration. Magnitude of δ increases in a rate of 0.012 electrons/Co pulse. Reversely, δ can be reduced by negative electronic pulses (see inset of Fig. 2). Thus, we can finely and reversibly control the δ value by an external electric pulse. Absorption spectra a(hv) in the visible-violet region (1–4 eV) were measured with a conventional monochromator system with a xenon lamp. The transmitted light was

FIG. 1. (Color online) (a) Schematic illustration of valence-differential spectroscopy, (b) Relative change (Δδ) in the averaged valence against the number of the electric pulses, 0.25 V in height and 1 s in duration, in Na0.84–δCo[Fe(CN)6]δ0.713.8H2O (δ=0.14) film.

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focused on a Si-photodiode, and a lock-in detection was adopted to enhance the S/N ratio. $\alpha(hv)$ spectra in the infrared region were measured with a Fourier-transform-type spectrometer. We emphasize that the small sample size (2 mm in thickness and 2 cm$^2$ in area) enables our method to combine other microspectroscopic technique. This is an advantage point compared to an in situ electrochemical optical measurement.

Figure 2 shows valence-differential absorption spectra $\Delta\alpha(hv)$ against $\Delta\delta$ in $\text{Co}_{2+}^{2+}$–$\text{Fe}^{2+}$ at 300 K. Solid curves were measured in the $\Delta\delta$-increasing (oxidation) run, while broken curves were measured in the $\Delta\delta$-decreasing run (reduction process). The oxidation process of the Fe site increases (decreases) the density of the $\text{Fe}^{2+}$ ($\text{Fe}^{2+}$) state. Then, the positive (negative) spectral component at $\Delta\delta=0.015$ and 0.060 can be assigned to the optical transitions related to the $\text{Fe}^{2+}$ ($\text{Fe}^{2+}$) state. Reversely, the reduction process of the Fe site decreases (increases) the density of the $\text{Fe}^{3+}$ ($\text{Fe}^{3+}$) state, and suppresses $\Delta\alpha(hv)$ spectra magnitude of $\Delta\alpha$ at 2.4 eV and increases linearly with $\Delta\delta$, as shown in the inset.

Figure 3(a) shows $\alpha(hv)$ and $\Delta\alpha(hv)$ spectra around the CN stretching mode of $\text{Co}^{2+} \rightarrow \text{Fe}^{2+}$.$^6$ The $\Delta\alpha(hv)$ spectrum shows positive (negative) change in the CN stretching mode within $[\text{Fe}^{2+}(\text{CN})_6]^3^−$ ($[\text{Fe}^{2+}(\text{CN})_6]^3^−$),$^7$ indicating the hole doping on the Fe site. Figure 3(b) shows $\alpha(hv)$ and $\Delta\alpha(hv)$ spectra of $\text{Co}^{2+} \rightarrow \text{Fe}^{2+}$. The $\Delta\alpha(hv)$ spectrum shows a derivative profile, indicating a rigid blueshift$^8$ of the CN stretching mode within $[\text{Fe}^{2+}(\text{CN})_6]\text{Fe}^{3+}$. This indicates that the holes are introduced on the Co site. The rigid shift may be ascribed to the doping-induced volume change.$^8$

Figure 3(c) shows $\alpha(hv)$ and $\Delta\alpha(hv)$ spectra of $\text{Co}^{2+} \rightarrow \text{Fe}^{2+}$ at 300 K, respectively. In the $\alpha(hv)$ spectrum, three broad structures are observed at 2.4, 3.2, and 3.8 eV.

The $\Delta\alpha(hv)$ spectrum clearly indicates positive, negative, and positive signals around the respective structures. We further found a shoulder structure at 2.7 eV in the $\Delta\alpha(hv)$ spectrum, suggesting an extra component. We decompose the spectrum with four Gaussian functions, i.e., $P_1$, $P_2$, $N_1$, and $P_3$. Solid curves in the lower panel in Fig. 3(c) are results of the least-square fitting. The shoulder structure at 2.7 eV is satisfactory reproduced by the four-component fitting. Figure 3(d) shows $\alpha(hv)$ and $\Delta\alpha(hv)$ spectra of $\text{Co}^{2+} \rightarrow \text{Fe}^{2+}$ at 300 K, respectively. The $\Delta\alpha(hv)$ spectrum shows positive, negative, and positive signals at 2.2, 3.2, and 3.8 eV. So, we decompose the spectrum with four Gaussian functions, i.e., $P_1$, $N_1$, and $P_2$ [see solid curves in the lower panel of Fig. 3(d)]. Thus obtained optical constants, transition energy $E$, width $\Gamma$, and oscillator strength $f$, of the respective components are listed in Table I together with the assignments.
(vide infra). Magnitude of $f$ was estimated from the spectral weight and $\Delta \delta$.

Now, let us proceed to the assignment of the respective components observed in Figs. 3(c) and 3(d). The valence-differential spectroscopy indicates that the positive (negative) components in $\text{Co}^{2+} - \text{Fe}^{2+}$ [Fig. 3(c)] can be assigned to the optical transitions related to the $\text{Fe}^{3+}$ ($\text{Fe}^{2+}$) state, while those in $\text{Co}^{2+} - \text{Fe}^{2+}$ [Fig. 3(d)] can be assigned to the transitions related to the $\text{Co}^{3+}$ ($\text{Co}^{2+}$) state. The absorption edge of $\text{Co}^{2+} - \text{Fe}^{2+}$ ($\text{Co}^{2+} - \text{Fe}^{2+}$) is ascribed to charge-transfer transition of an electron from the $\text{Co}^{2+}$ to $\text{Fe}^{3+}$ state (from the $\text{Fe}^{2+}$ to $\text{Co}^{3+}$ state). The corresponding positive signals [P1 in Figs. 3(c) and 3(d)] are observed in both the films around the absorption edge. In $\text{Co}^{2+} - \text{Fe}^{2+}$ [Fig. 3(c)], the higher-lying positive components (P2 and P3) can be ascribed to charge-transfer of an electron from the ligand (CN group) to low-spin $\text{Fe}^{3+}$ ($t_{2g}^6$) site. The doublet structure is due to the ligand field splitting (10Dq) of the final states, since the observed value ($\approx 1.17$ eV) is comparable with the 10Dq for the $\text{Fe}^{3+}$ ion. The negative component (N1) is related to the low-spin $\text{Fe}^{2+}$ ($t_{2g}^6$) site, and is reasonably ascribed to intramolecular transition within [Fe$(\text{CN})_6$]$.^4$ In $\text{Co}^{2+} - \text{Fe}^{2+}$ [Fig. 3(d)], on the other hand, the higher-lying positive components (P2) is due to charge transfer of an electron from the ligand to low-spin $\text{Co}^{3+}$ ($t_{2g}^6$) site. The negative component (N1) is related to the high-spin $\text{Co}^{2+}$ ($t_{2g}^5e_g^2$) site. Looking at Table I, one may notice that the transition energy of the P3 component in $\text{Co}^{2+} - \text{Fe}^{2+}$ is very close to that of the P2 component in $\text{Co}^{2+} - \text{Fe}^{2+}$, suggesting a strong hybridization between the $\text{Co}^{2+}$ and $\text{Fe}^{2+}$ sites. Such a hybridization perhaps plays a significant role in the thermally and/or photoinduced charge-transfer transition observed in the $\text{Co} - \text{Fe}$ cyanides.

The valence-differential spectroscopy has several advantages over conventional spectroscopy. First, the spectroscopy selectively extracts the spectral components related to the oxidized/reduced metal site as positive/negative signals. The resultant sign modulation within the spectra enables accurate and unambiguous spectral decomposition as well as well-founded assignment. Actually, assignments listed in Table I are consistent with the assignments based on the conventional spectroscopy on a series of valence-controlled $\text{Co} - \text{Fe}$ films. In addition, magnitudes of the optical constants are more reliable in the present investigation, because all the parameters, i.e., $E$, $\Gamma$, and $f$, can be determined by least-square fitting of the spectra without any assumptions and/or constraints. Second, the spectroscopy significantly reduces light scattering effect as well as sample/position dependence of the spectra, because the spectra were measured at exactly the same position before and after the electronic pulses. Actually, small signal of the order of $10^{-3}$ can be discernible in the present experiment. Third, the spectroscopy is widely applicable to the other solid films. In the case of the film whose oxidized/reduced state is unstable, a transient spectroscopy under an external electric pulse is possible. Actually, we applied a transient spectroscopy to $\text{Fe}(\text{CN})_6$[$\delta$]$_3$/$\text{H}_2\text{O}$ film, and obtained reliable differential spectra. In addition to an intense charge-transfer transition at 1.8 eV, we observed a doublet structure at 2.9 and 3.8 eV. The doublet structure can be assigned to charge-transfer of an electron from the ligand to low-spin $\text{Fe}^{3+}$ ($t_{2g}^6$) site. Thus, the valence-differential spectroscopy is a promising tool to clarify complicated electronic structure of solid films, such as Prussian blue analogs and other electrochromic materials.

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