The Concentration Effects for the Adsorption Behavior of Heptyl Viologen Cation Radicals on Indium-Tin-Oxide Electrode Surfaces

Yusuke Ayato, Takashi Itahashi, Akiko Takatsu, Kenji Kato, and Naoki Matsuda[†]

Abstract—In situ observation of absorption spectral change of heptil viologen cation radical (HV+) was performed by slab optical waveguide (SOWG) spectroscopy utilizing indium-tin-oxide (ITO) electrodes. Synchronizing with electrochemical techniques, we observed the adsorption process of HV+ on the ITO electrode. In this study, we carried out the ITO-SOWG observations using KBr aqueous solution containing different concentration of HV to investigate the concentration dependent spectral change. A few specific absorption bands, which indicated HV+ existed as both monomer and dimer on ITO electrode surface with a monolayer or a few layers deposition, were observed in UV-visible region. The change in the peak position of the absorption spectra from adsorption species of HV+ were correlated with the concentration of HV as well as the electrode potential.

Keywords—absorption phenomena, heptil viologen, indium-tin-oxide (ITO) electrode, in situ, slab optical waveguide (SOWG) spectroscopy,

I. INTRODUCTION

WE have been studying the adsorption process and adsorbed states of various molecules on solid/liquid interfaces by slab optical waveguide (SOWG) spectroscopy [1]-[7]. Our recent improvement of the SOWG instrument based on the light incidence method utilizing a glycerol drop has enabled us to observe time dependent adsorption behavior [8]-[11], and to obtain the standard Gibbs free energy (Δ G) value due to adsorption process of proteins on solid/liquid interfaces. The development of an indium-tin-oxide (ITO) coated SOWG (ITO-SOWG) has also allowed us to carry out the spectroelectrochemical observation of adsorbed molecules accompanied by redox reaction at electrochemical interfaces [12]-[14].

N,N'-Disubstituted-4,4'-bipyridinium salts (viologens) were first studied for electrochromic devices (ECDs) in 1970s

and have been attracted much attention for the application owing to their high transmittance changes and fast responses [15]-[17]. Recent studies have been performed for applications of viologens to herbicides, electron mediators and catalysts [18]-[20]. 1,1'-Dimethyl-4,4'-bipyridinium (methyl viologen; MV) modified electrodes exhibit measurable rates of heterogeneous electron transfer with several biological molecules [21], [22]. 1,1'-Diheptyl-4,4'-bipyridinium (heptyl viologen; HV) is usually studied for electrochromic applications because its one-electron reduction of the dication HV²⁺ to the mono-cation radical salt HV⁺ in aqueous solutions can form a precipitate with violet color on electrode surfaces [23]-[32].

 HV^{2+} has been known to receive two-step one-electron reactions resulting in the formation of HV^{+} and HV.

$$HV^{2+} + e^{-} \rightarrow HV^{+}. \tag{1}$$

It is reversible reaction and immediately follows chemical reaction with proper counter anions (X'), which produced a cation radical salt film on electrodes with strong violet color. Jasinski reported that the precipitates film of HV⁺ X' has amorphous structure in the primary stage of the film formation process, then, the phase gently and unsteadily changes in successive reactions, and the aging process follows [33].

The second reduction step of HV^{+} can be expressed by Eq. (2):

$$HV^{+} + e^{-} \rightarrow HV \tag{2}$$

It is less reversible reaction and forms a solid deposit with yellow color on the electrode surface. These film formation processes have been solved by a number of researchers using various analytic techniques [31], [34]-[38]. However, so little report included the time and the potential dependent adsorption states change on the electrode surfaces covered with the monolayer or a few layers deposition.

Although a number of researchers have demonstrated for the reduction process of the first reduction step of HV^{2+} , they mainly observed the adsorption states of HV^{+-} in multilayer deposition on the electrode surface. In addition, they usually discussed the behavior of the absorption bands at around 550 nm, so little reports included the results of the absorption bands at around 400 nm. The information of the absorption band only at around 550 nm is insufficient to comprehend the potential

Y. Ayato and N. Matsuda† are with On-site Sensing and Diagnosis Research Laboratory, AIST, 807-1, Shuku, Tosu, Saga, 841-0052, Japan (corresponding author† to provide phone: +81-942-81-3623; fax: +81-942-81-3690; e-mail: naoki.matsuda@aist.go.jp).

T. Itahashi is with Mechanical and Electrical Systems Engineering Advanced Course, Kurume National College of Technology, 1-1-1, Komorino, Kurume, Fukuoka, 830-8555, Japan.

A. Takatsu and K. Kato are with National Metrology Institute of Japan, AIST, Central 3, 1-1-1, Umezono, Tsukuba, 305-8563, Japan.

dependent adsorption process of HV⁺⁻ on the electrode surface, because the absorption bands of HV⁺⁻ monomer and dimer are observed in one broad band consisting of a few absorption bands.

The main aim of present study is to discuss the adsorption behavior of HV⁺ monomer and dimer, especially in a monolayer deposition, separately by employing the absorption bands at around 400 nm. And we will note and investigate the potential dependent adsorption behavior of HV⁺ molecules by the spectroelectrochemical SOWG using different concentration of HV.

II. EXPERIMENTALS

A Materials

Heptylviologen dibromide (HVBr₂) purchased from Tokyo Kasei Industry Co., Ltd. was used as received without further purification. Potassium bromide (KBr) used as the supporting electrolyte was guaranteed grade and purchased from Kanto Chemical Co., Inc. The sample solutions were prepared using Milli-Q water (resistivity > 18 M Ω cm). The concentrations of HVBr₂ were adjusted to 1×10⁻³, 5×10⁻⁵ and 1.25×10⁻⁵ mol dm⁻³ and that of KBr was 0.3 mol dm⁻³.

B. Spectroelectrochemical SOWG Systems

The experimental details of SOWG have been given elsewhere [11], [13], [39]. As shown in Fig. 1, a 150-W xenon lamp (System Instruments Co., Ltd., Japan) was used as the light source, and the light was guided by an optical fiber. The SOWG spectra were detected using a CCD detector with monochrometer (PMA-11, Hamamatsu Photonics, Japan). The minimum time resolution of the CCD detector is 20 ms. The ITO film was formed by vapor deposition on a glass plate (50 mm × 20 mm × 0.05 mm, Matsunami Glass Industry, Ltd., Japan) giving a thickness of approximately 20 nm. ITO-SOWG plates were cleaned by soaking in ethanol for 1 h and then rinsed extensively with Milli-Q water. spectroelectrochemical surface area of ITO-SOWG working electrode covered with sample solution was approximately 1.3 cm². A platinum wire and an Ag/AgCl electrode were used as the counter and reference electrodes, respectively. All the potentials in this paper are quoted against Ag/AgCl. The electrode potential was controlled with a potentiostat (EG&G Princeton Applied Research, Model 273) synchronizing with the spectrometer.

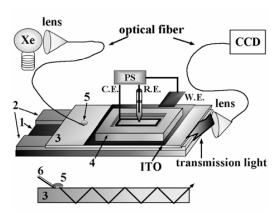


Fig. 1 Illustration of ITO-SOWG systems. 1: slide glass base, 2: silicon rubber strip acting SOWG supports, 3: SOWG (50×20×0.05 mm), 4: leak-proof silicone rubber solution cell, 5: glycerol drop, and 6: optical fiber from the light source. PS: Potentiostat, W.E., C.E., and R.E.: working (ITO), counter (Pt wire), and reference (Ag/AgCl) electrode, and ITO: indium-tin-oxide electrode.

C. Measurements

The resulting SOWG spectra are shown in terms of absorbance units defined as follows:

$$A = -\log(I_S/I_R) \tag{3}$$

Where I_S/I_R represent the relative change in the intensity and I_S and I_R are the sample and reference intensities, respectively. All reference spectra on the SOWG analysis were observed at -200 mV in this study.

In situ spectroelectrochemical SOWG observations were carried out with synchronized with cyclic voltammetric measurements. The electrode potential was swept to the cathodic direction from -200 to -600 mV, and back to -200 mV at a sweep rate of 10 mV s $^{-1}$. The time width of each spectrum was 600 ms and SOWG spectra were obtained continuously.

A potential step method was also applied to in situ SOWG measurements. Before the sample measurements, the electrode potential was held at -200 mV for 30 s and the reference spectrum was measured with the exposure time of 300 ms. The resulting spectra were measured at -600 mV for 60 s with the exposure time of 300 ms.

III. RESURUTS AND DISCUSSION

A. Cyclic voltammetry

The first one-electron reduction of HV^{2+} giving the radical salt is known to a simple, soluble and reversible reaction. Since the electron transfer reaction is fast, it is effectively diffusion controlled at enough large current densities [40]. Fig. 2 shows the cyclic voltammogram measured in 0.3 mol dm⁻³ KBr aqueous solution containing 1×10^{-3} mol dm⁻³ HV^{2+} . A pair of the peaks was observed at around -400 and -500 mV. One peak at around -500 mV corresponds to the one-electron reduction shown in Eq. (1), and another peak at around -400 mV indicates

the oxidation from HV^{+} to HV^{2+} , which is the reversal reaction of Eq. (1). A strong violet color was observed directly by eyes from the precipitates film was formed on the electrode surface below -500 mV.

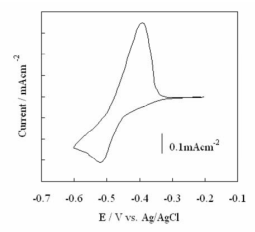


Fig. 2 Cyclic voltammograms at ITO electrodes for 1×10^{-3} mol dm⁻³ HVBr₂ in 0.3 mol dm⁻³ KBr aqueous solution. Sweep rate: 100 mV s⁻¹.

The surface coverage of the electrode can be estimated from the redox current of the cyclic voltammograms. The charge density in 1×10⁻³ mol dm⁻³ HV²⁺ containing solution was approximately 140 µC cm⁻². The charge density for the reduction of HV+ of monolayer coverage, assuming one molecule occupies 0.5 or 1 nm², is about 32 or 16 μC cm⁻². Thus, the electrode surface should be covered with multilayer precipitates film of HV+. To investigate the change in the adsorption states of HV+ molecules in a thin deposition layer, we carried out the cyclic voltammetric mesurments with following experimental condition. The electrode potential was held at -600 mV for 60 s in 0.3 mol dm⁻³ KBr aqueous solution containing 5×10⁻⁵ mol dm⁻³ HV²⁺, and then swept at 20 mV s⁻¹ from -600 to -200 mV. The charge density was obtained approximately 40 µC cm⁻² from the first cycle of the cyclic voltammograms. Thus, all spectroelectrochemical SOWG data obtained in this study should be based on the potential dependent change of adsorbed HV+ · molecules in the monolayer or a few deposition layers on the electrode surface.

B. In Situ Spectroelectrochemical Observations by SOWG Spectroscopy Synchronizing with Cyclic Voltammetric Measurements

Fig. 3 shows SOWG absorption spectra obtained by synchronizing with cyclic voltammetric measurements using 0.3 mol dm⁻³ KBr aqueous solution containing 5×10^{-5} mol dm⁻³ HV²⁺. It has been known that several absorption bands assigned to HV⁺ species are observed in UV-visible region [41]. As shown in the Fig. 3, several positive-going bands were observed at around 400 and 550 nm. The absorption band at around 380 nm, which was assigned to the dimer of HV⁺ [41],

appeared in UV-visible region below about -400 mV (20 s) and another absorption at around 400 nm, which was assigned to a monomer of HV⁺⁺ [41], seems to arise from more positive electrode potential. The absorption band at around 530 nm appeared as a broad-band below about -400 mV, which was assigned to dimer of HV⁺⁺ [42]. The absorption band at around 550 nm might be observed due to the different aggregation from monomer and dimer of HV⁺⁺ [43].

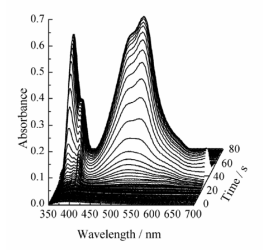


Fig. 3 Time- resolved SOWG spectra synchronized with cyclic voltammetric measurements in $0.3 \text{ mol dm}^{-3} \text{ KBr}$ aqueous solution containing $5 \times 10^{-5} \text{ mol dm}^{-3} \text{ HV}$. Sweep rate: 10 mV s^{-1} .

Fig. 4 shows the potential dependent change in the intensity and peak position of the absorption bands at around 380 and 400 nm obtained by SOWG data synchronized with cyclic voltammetric measurements. The intensities of adsorption bands at around 400 and 380 nm started to increase monotonically with decreasing the electrode potential at around 16.8 and 21 s passed (at -368 and -410 mV). Contrary to the results for the cathodic polarization, the intensities of them decreased with increasing the electrode potential and disappeared completely at around 63.6 and 60.6 s (- 364 and -394 mV) in the anodic polarization. Thus, it was confirmed that the dimer of HV⁺⁻ adsorbed at lower electrode potential region than the monomer of HV⁺⁻ on the electrode surface.

In the potential dependent peak position change, the band at around 400 nm showed no peak shift of absorption band under this experimental condition, as shown in Fig. 4(a). This result was very similar to that observed in our former study [41]. On the other hand, the peak position at around 380 nm has been known to show the peak shifts depended on the electrode potential [41]. The absorption band at around 380 nm appeared at 374 nm and exhibited a red shift to 380 nm with decreasing electrode potential. In the positive polarization, a blue shift of the maximum absorption wavelength was observed from 380 to 374 nm. These results were also similar to those observed in our former study [41]. The peak shift of the absorption band at

around 380 nm implied that the different aggregation of HV⁺ adsorbed on the electrode surface.

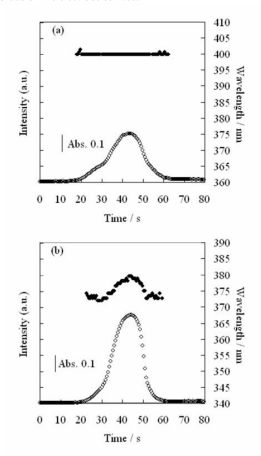


Fig. 4 Change in the intensity and peak position at around 400 nm (a) and 380 nm (b) obtained from time-resoluved SOWG analysis synchronized with cyclic voltammetric measurements in 0.3 mol dm⁻³ KBr aqueous solution containing 5×10⁻⁵ mol dm⁻³ HV. Sweep rate: 10 mV s⁻¹.

C. In Situ SOWG Measurements using Potential Step Methods

To investigate whether the absorption band at around 380 nm shows the peak shift depending on the concentration of HV or not, we carried out time-resolved SOWG observations by synchronizing with potential step measurements in 0.3 mol dm⁻³ KBr aqueous solutions containing 5 and 1.25×10⁻⁵ mol dm⁻³ HV²⁺. Fig. 5 shows the change in the peak position of absorption band at around 380 nm obtained by SOWG analysis synchronizing with potential step measurements from -0.2 to -0.6 V for 60 s. In potential step measurements, the peak position of the absorption band at around 380 nm exhibited a red shift from around 375 to 390 nm from 0.3 to 60 s in 5×10⁻⁵ mol dm⁻³ HV²⁺ containing solution. On the other hand, clear peak shift of the absorption band at 380 nm was not observed

for 60 s in the case of using $1.25 \times 10^{-5} \text{ mol dm}^{-3} \text{ HV}^{2+}$ containing solution. These results indicate that the change in the peak position of the absorption band at around 380 nm probably depended on the concentration of HV as well as the electrode potential.

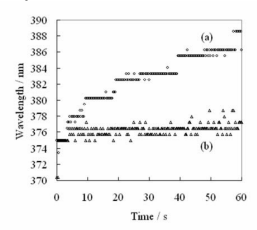


Fig. 5 Potential dependence of the peak position at around 380 nm obtained from time-resoluved SOWG analysis synchronized with potential step measurements from -200 to -600 mV for 60 s in 0.3 mol dm⁻³ KBr aqueous solutions containing 5 (a) and 1.25 (b) ×10⁻⁵ moldm⁻³ HV.

Consequently, we carried out a very successful observation for the heptilviologen cation radicals on the electrode surface by using spectroelectrochemical SOWG. The adsorption behavior of HV^{++} could be clearly characterized and distinguished both monomer and dimer by employing the absorption bands at around 400 nm.

This technique we have presented allowed the very successful observation of both the spectroscopic and electrochemical properties of molecules on electrode/electrolyte interfaces. Highly sensitive in situ SOWG spectra yield essential molecular information for coverage of monolayer or a few layers.

IV. CONCLUSION

The ITO-SOWG successfully observed the potential and concentration dependent structural changes. It was found from SOWG studies that several different species of HV⁺⁻ adsorbed on the electrode surface. The dimer of HV⁺⁻ was adsorbed more negative electrode potential than the monomer of HV⁺⁻ on the ITO electrode surfaces. These monomeric and dimeric HV⁺⁻ molecules co-adsorbed on the ITO electrode surface in all applied negative electrode potential region. The adsorption states of HV⁺⁻ molecules depended on the concentration of HV as well as the electrode potential.

We presented an application of the SOWG spectroscopy, which can easily and directly yield in situ molecular information on electrode/electrolyte interfaces. This powerful technique will disclose the configurations of many other

molecules adsorbed on the electrode surfaces.

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