学位論文(要約)

Study on Reversible Photo- and Electro- Isomerization Systems of Benzodimethyldihydropyrene Derivatives

(ベンゾジメチルジヒドロピレン類を用いた

可逆な光・電気化学異性化反応系の研究)

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Introduction

Photochromic molecules change their molecular structures and colors by light irradiation, thereby forming two metastable isomers. The photoisomerization accompanies the change of the molecule's electronic structure and physical properties, such as luminescence, magnetism, and electronic conductivity, and it is useful to combine the photo-functionality with the electro-functionality for the purpose to fabricate molecular devices. In this sense, to find unique electrochemical properties of photochromic molecules is important to establish dual electro- and photo responsive systems. In my Ph.D course study, I focused on benzodimethyldihydropyrne (BzDHP) derivatives. Benzodimethyldihydropyrne is one of the most efficient photochromic molecules; upon visible light irradiation, purple-colored, BzDHP, is transformed into a colorless benzocyclophanediene, BzCPD. The reverse isomerization is induced by UV light irradiation or heat treatment. It was known that BzDHP and BzCPD exhibited different redox properties, but the relationship between the photoisomerization and the redox properties have not been clarified. I discovered a new type of isomerization which is induced by oxidation, which allowed to fabricate both photo- and electro-responsive molecular modified electrode.

Electronic communication between the ferrocene moieties in bis (ferrocenyl)DHP and its electronic stateIn Chapter 2, I studied about the electronic state of DHP structure as a spacer unit with comparing three molecules (Figure 1). Previously it is

was

known that 2 shows the

communication between two ferrocene. But the

confirmed by only cyclic

measurement. In order to

reveal the effect of π

conjugation of the linker, I

measured the UV-vis-NIR

electronic

electronic

communication

voltammetry

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Figure 1 Molecule structures of 1-3.



Figure 2 (A) Energy levels of HOMOs of **2** and **3** estimated by DFT calculation. The blue orbitals have d-orbital characters, whereas the red ones are chiefly composed of the π orbitals of the linkers. (B) Selected molecular orbitals of **2** and **3**.(Kishida, M.; Muratsugu, S.; Sakamoto, R.; Kusamoto, T.; Nishihara, H.; *Chem. Lett.* **2013**, *42*, 361-362)

spectrometry and carried out DFT calculation to the three molecules. When an oxidant, $[(4-BrC_6H_4)_3N][SbCl_6]$, was added into the dichloromethane solution of **2**, UV-vis-NIR spectrum showed the broad band in the NIR region. This band is considered to compose of two different

bands; the first band is an intervalence charge transfer transition from the ferrocene moiety to the ferrocenium moiety, and the second band is an charge transfer transition from the DHP moiety to the ferrocene moiety. This spectrum confirms the electronic communication between the two ferrocene in **2**. DFT calculation revealed the energy level of MOs as shown in Figure 2A. The energy level of the π orbital of DHP is near that of d orbital of ferrocene, while the energy level of the π orbital of diethynelphenylene is far from that of d orbital of ferrocene. These different energy diagrams support that DHP structure enhances the electronic interaction between ferrocenes. These results suggest the π conjugation of the linker is important to communicate the two redox active units.

Fabrication of a modified electrode with reversible photo- and electro- responsive BzDHP molecules



Ι examined the photoisomerization of 4c and 5c on an ITO electrode (Figure 3). The modified electrode ITO-4c and ITO-5c was fabricated by immersing an ITO electrode into a 0.1 mM chloroform solution of 4c for over night, in that of 5c for 1h. The immobilization of 4c and 5c were confirmed by cyclic voltammetry and UV-vis spectrometry. The reversible isomerization of ITO-4c was not observed, but that of ITO-5c was observed. Figure

Figure 3 (A) Structure of 4c and 5c (B) Scheme of ITO-5c and ITO-5o.

4A shows the cyclic voltammogram of ITO-**5c** after the visible light irradiation ($\lambda = 550$ nm). After the several redox reactions, the oxidation peak was observed at 0.30 V vs. Ag⁺/Ag. The change of the volatmmogram by the oxidation was similar to that observed in the cyclic voltammogram of **5o** dissolved in a dichloromethane solution. This suggests that the same oxidation-induced cyclization of **5o** occurred for a molecule immobilized on ITO.

The UV-vis spectral change was observed by the 550 nm visible light irradiation as displayed in Figure 4B, where the peak absorbance at 400 nm was decreased. The following oxidation and the reduction of **50** recovered the absorbance at 400 nm partially. These results suggest that visible light photoirradiation change ITO-**5c** to ITO-**50** and the oxidation of **50**⁺/**50** change ITO-**50** to ITO-**50** as confirmed by the cyclic voltammograms.



Figure 4 (A) Cyclic voltammograms of ITO-**50** in the first scan (plain line) and in the second scan (dashed line) in 0.1 M Bu₄NClO₄ – dichloromethane. (B) Changes of Δ Absorbance after visible light irradiation (plain line) and after oxidation (dashed line). Δ Absorbance indicates difference of spectrum from a spectrum before the photoirradiation or oxidation.





reverse isomerization

In Chapter 4, I studied the photochemical and electrochemical properties of 6c, 60 and their analogs 7c, 70 (Figure 5). Figure 6 shows the cyclic voltammograms of 6c, 6o, 7c, and 7o. An irreversible redox wave was observed at $E_{p,a} = 0.39$ V (0.44)V) vs. ferrocenium/ferrocene in the cyclic voltammogram of 60 (70), indicating an occurrence of a chemical process followed by oxidation (EC reaction). In the reverse scan in the negative

Figure 5 Scheme of the catalytic oxidation cyclization of 6 and 7.

direction, a cathodic peak was

observed at the similar potential to that of **60** (**70**), which suggests that **60** (**70**) was converted into **6c** (**7c**) by oxidation.

This oxidative **60** to **6c** cyclization is further supported by UV-vis spectroscopy, where the unique catalytic nature of the **60** to **6c** isomerization was revealed by monitoring the change of the absorbance after adding 0.3 equiv. of $[(4-BrC_6H_4)_3N][SbCl_6]$ to a dichloromethane solution of **60** (Figure 7A). The spectrum obtained at 2 hours after the addition of the oxidant was similar to that expected for **6c** + **6c**⁺ with the **6c**⁺/**6c** ratio of ca. 3/7, indicating that most **60** were converted into **6c** (Figure 7B). The reaction rate of the electro-catalytic **60** to **6c** isomerization was remarkably faster



than that of (non-catalytic) thermal **60** to **6c** isomerization which takes 4 days to terminate the isomerization.

The mechanism of the catalytic to 6c isomerization is 60 explained as follows (Figure 5): 10 is oxidized by [(4-BrC₆H₄)₃N][SbCl₆] to form 60^+ (eq. 1). 60^+ is thermally unstable to isomerize immediately to $6c^+$ (eq. 2). Then, intermolecular electron transfer occurs from $6c^+$ to neutral 6o, yielding neutral 6c and $6o^+$ (eq. 3). The generated **60⁺** isomerizes to $6c^+$ according to eq. 2. Consequently, the chemical reactions shown in eq. 2 and eq. 3 successively occur so as to convert all 60 into 6c.

To confirm the reaction shown

Figure 6 Cyclic voltammograms of 6c and 6o (A), and 7c and 7o (B), in 0.1 M Bu₄NClO₄ – dichloromethane.

in eq. 3, 0.3 equiv. of $6c^+$ was added to the solution of 6o, and the reaction was monitored by UV-vis spectroscopy. Predictably the absorbance of the peak derived from 6c was increased, and eventually almost all 6o was converted into 6c. This result supports the reaction in eq. 3, in which $6c^+$ plays a key role in this chain reaction. The rate of the catalytic oxidative cyclization was higher in 6o than in 6o. $\Delta E^{0'}_{oxpeak}$ ($\Delta E^{0'}_{oxpeak}$ indicates the difference of $E^{0'}_{oxpeak}$ between 6o (7o) and 6c (7c)) is 0.10 V smaller in 7o than in 6o; thus, the intermolecular electron transfer shown in eq. 3 proceeds more efficiently in 7o than in 6o. The results were consistent with the expectation that the eq. 3 is a rate-determining step in the chain reaction.



Figure 7 (A) Changes of Δ Absorbance after addition of 0.3 equivalent of $[(4-BrC_6H_4)_3N][SbCl_6]$ to 0.1 M dichloromethane solution of **10**. Δ Absorbance indicates difference of spectrum from a spectrum just after the addition. (B) Time dependence of ratio of **1c** and **2c**.

The combination of the catalytic isomerization with the photochromic nature allowed me to establish а novel photo-sensing system. The introduction of photons (light signal) was detected by electrode rest potential (E_{rest} ; electric signal). In the presence of 0.1 equiv. of $6c^+$, E_{rest} of dichloromethane the solution of 6c was shifted from 0.20 V to 0.23 V vs. Ag^+/Ag by successive switching of the visible light irradiation (Figure 8A). This behavior was repeated over 10 cycles. $E_{\rm rest}$ is briefly determined by the 6c⁺/6c ratio according to Nernst equation. The visible



Figure 8 Change of electrode rest potential of (A) 6c and (B) 7c with 0.1 eq. of $[(4-BrC_6H_4)_3N][SbCl_6]$ in Bu_4NClO_4 -dichrolomethane.

light irradiation converted **6c** into **6o** and increased the **6c**⁺/**6c** ratio. Eventually E_{rest} value was positively shifted. After the irradiation was stopped, the catalytic **6o** to **6c** isomerization reaction was induced by the presence of **6c**⁺. Then the **6c**⁺/**6c** equilibrium reverted to the initial state to decrease E_{rest} to the initial value. As shown in Figure 8B, E_{rest} of **7c** was changed between 0.27 V and 0.31 V in the cycles. The range of E_{rest} was different from that of **6c**, while **6c** and **7c** were similar in ΔE_{rest} . The results demonstrate a new photo-sensing system on the basis of the photoisomerization and electro-catalytic isomerization.

Conclusion

In my Ph.D study, I studied that electronic and photonic properties of DHP derivatives. It is found that DHP enhances the electronic communication between the redox sites, and that BzDHP derivative shows electro-catalytic isomerization. I fabricated reversible photo- and electro-responsive electrode system in chapter 3 and constructed a new photo-sensing system using this catalytic reaction in chapter 4. In this study, I proposed a new photo- and electro- functional system, which provides the basis to the construction of the functional molecular devices.