

HOKKAIDO UNIVERSITY

| Title | Mechanisms of photo-induced degradation of polythiophene derivatives: re-examination of the role of singlet oxygen | |
|------------------|--|--|
| Author(s) | Ohta, Hiroki; Koizumi, Hitoshi | |
| Citation | Polymer bulletin, 74(6), 2319-2330 https://doi.org/10.1007/s00289-016-1837-6 | |
| Issue Date | 2017-06 | |
| Doc URL | http://hdl.handle.net/2115/70626 | |
| Rights | The final publication is available at link.springer.com. | |
| Туре | article (author version) | |
| File Information | 160630Photodegradation_P3AT.pdf | |



Hokkaido University Collection of Scholarly and Academic Papers : HUSCAP

Mechanisms of photo-induced degradation of polythiophene derivatives: Re-examination of the role of

singlet oxygen

Hiroki Ohta and Hitoshi Koizumi

Division of Applied Chemistry, Faculty of Engineering, Hokkaido University,

Kita-ku, Sapporo, Hokkaido, 060-8628, Japan

Corresponding author: Hitoshi Koizumi

Tel.:+81-11-706-6748

E-mail address: koizumih@eng.hokudai.ac.jp

Abstract

Roles of singlet oxygen (${}^{1}O_{2}$) in photo-induced degradation of poly(3-hexylthiophene) (P3HT) and poly(3-hexyloxythiophene) (P3HOT) were examined. The results indicate that ${}^{1}O_{2}$ has a considerable role in the degradation of P3HT and P3HOT. Photo-irradiation of P3HT in air resulted in reduction of the π -conjugation. The photo-induced reduction also occurred for P3HOT, and it was much faster than that of P3HT. Manceau et al. have reported that ${}^{1}O_{2}$ was generated by photo-irradiation of P3HT in the presence of oxygen and nevertheless ${}^{1}O_{2}$ is not the principal photo-oxidative degradation intermediate of P3HT [Manceau et al., Macromol Rapid Commun 29 (22):1823-1827 (2008)]. However, exposures of solid P3HT and P3HOT to chemically generated ${}^{1}O_{2}$ provided decreases in the $\pi\pi$ *absorption in our study. The decrease for P3HOT was much larger than that for P3HT. The exposure of P3HOT showed formation of polarons. The difference in P3HT and P3HOT for the effects of the photo-irradiation and of ${}^{1}O_{2}$ will be ascribed to the difference in their ionization potentials.

1. Introduction

Conducting polymers have potentials for various applications in electronics, such as organic field effect transistor (OFET) [1], organic solar cells (OSC) [2], organic light emitting diode[3], and actuators[4]. In particular, soluble conducting polymers are key materials to enable flexible, large area, and low-cost products by printing processes[5].

Poly(3-alkylthiophene)s (P3AT) are one of the most common polymers in researches on OFET[6] and OSC[7]. For applications exposed to light, such as OSC, it is important to understand mechanisms of their photo-induced degradations[8]. Photo-induced chemical reactions of P3AT have been studied from the early 1990's[9-13]. The reports are, however, relatively sparse, and the mechanisms of the photo-induced degradation is still ambiguous[14-16].

Two major mechanisms have been proposed: reactions of singlet oxygen $({}^{1}O_{2})$ generated by photosensitization with P3AT [9,10,17,11,12] and of photolytically or thermally- generated hydroxyl radicals [14-16]. The latter mechanism seems to be presently accepted as a principal photodegradation process of P3AT [18,19].

Holdcroft and Abdou investigated the mechanism with ${}^{1}O_{2}$ [9-12]. They examined degradations of poly(3-hexylthiophene) (P3HT) in solutions by photolysis and by reactions with chemically generated singlet oxygen(${}^{1}O_{2}$) [10-12], and photo-degradations of solid P3HT[9]. They have concluded that the photo-degradations in the presence of oxygen involve photosensitized generation and reactions of ${}^{1}O_{2}$.

Photolysis of poly(3-hexylthiophene) (P3HT) in O_2 saturated solutions[11,12] resulted in reduction of the π -conjugation and chain scissions, while those of solid P3HT provided the reduction and crosslinkings. They explained that 1,4-Diels-Alder addition of 1O_2 to a thienyl ring followed by rearrangements yielding a sulfine or a trans-diketone caused the reduction as shown in Fig. 1. The chain scissions and crosslinkings were ascribed to formation of P3HT radicals initiated by photolysis of impurity iron (III). Caronna et al. have also suggested that addition of 1O_2 to the conjugated double bonds for the mechanism of photo-degradation of poly(3-butlythiophene) in the solid state[17]. They analyzed low

molecular weight species produced by the photo-irradiation.

Manceau et al. have proposed the mechanism with hydroxyl radicals. They investigated the role of ${}^{1}O_{2}$ in photolysis of solid P3HT[15], and concluded that ${}^{1}O_{2}$ is not the principal photo-oxidative degradation intermediate of P3HT. They have confirmed that ${}^{1}O_{2}$ could be generated by the photolysis of P3HT in presence of oxygen. They have reported that P3HT, however, did not decompose by reactions with chemically generated ${}^{1}O_{2}$. They also investigated photodegradation of P3HT based on infrared spectra[14] and X-ray photoelectron spectra[16], and proposed a mechanism of the degradation through reactions of radicals as shown in Fig. 2: generation of OH radicals through photo or thermal C-H bond scissions at α -position of hexyl groups and reactions with oxygen (a), addition of the OH radicals to a sulfur of thienyl rings forming sulfone (b), and ring-opening of the sulfone (c).

Aoyama et al. examined photodegradation of polythiophene derivatives with no α -hydrogen at side

chains: poly(3-octyloxythiophene) (P3OOT), poly(3-phenylthiophene), and

poly[3-(1,1-dimethylpropyl)thiophene) [20]. P3OOT was converted to polarons by photo-irradiation in air, and this could be reverted to the pristine one by reduction with hydrazine. The other polythiophene derivatives degraded in the same way as P3HT.

Although the mechanism in Fig.2 seems to be presently accepted as a principal photodegradation process of P3AT [18,19], we have questions about it. P3HT and OH radial formation may proceed through reactions (a) in Fig.2. However, a theoretical study suggested that an attack of OH radial on sulfur in P3HT (b) was thermodynamically unfavorable[21]. Furthermore, the ring-opening of the sulfone (c) is questionable. To our knowledge, no experimental evidence for the ring-opening of thienyl-1,1-dioxide moiety has been reported. In addition, reactions in the mechanism is initiated by C-H bond rapture and addition of oxygen to α -position of the alkyl side chains. The photodegradation of polythiophene derivatives with no α -hydrogen, however, occurs in the same way as P3HT [20]. It is inconsistent with the mechanism.

This paper re-examines the role of ${}^{1}O_{2}$ in the photodegradation of P3AT. Reactions of ${}^{1}O_{2}$ with low-molecular unsaturated compounds are well established [22-25]. The Diels-Alder addition followed by ring-opening seems to be more plausible than that of thienyl-1,1-dioxide moiety.

2. Experimental

Regioregular P3HT was purchased from Aldrich. The weight-averaged molecular weight (M_w) and polydispersivity (M_w/M_n) determined by gel permeation chromatography against polystyrene standards were 55,800 and 1.94, respectively. P3HT was also obtained by oxidative polymerization of 3-hexylthiophene with FeCl₃[26]. The values of M_w and M_w/M_n were 8,300 and 1.60, respectively.

Poly(3-hexyloxythiophene) (P3HOT) was obtained by polymerization of 3-hexyloxythiophene with the Grignard metathesis(GRIM) method[27]. 3-Hexyloxythiophene was synthesized from 3-methoxythiophene and hexanol with sodium hydrogen sulfate (NaHSO₄) as a catalyst [28,29]. The monomer was dibrominated with *N*-bromosuccinimide (NBS) to afford

2,5-dibromo-3-hexyloxythiophene, which was subsequently transformed into the Grignard derivative with methyl magnesium bromide and polymerized with a Kumada coupling reaction, catalyzed by [1,3-bis(diphenylphosphino)propane]nickel(II) dichloride [Ni iCl₂(dppp)]. The values of M_w and M_w/M_n

for P3HOT were 7,900 and 1.15, respectively.

Films of P3HT and P3HOT were prepared by spin-coating from their solutions in chloroform on glass plates.

 $^{1}O_{2}$ was generated with the reaction between hydrogen peroxide and sodium hypochloride [30,31,10,15]. The polymer films were exposed to $^{1}O_{2}$ by two methods. One is the same as that by Manceau et al.[15]: 100 µL of 0.6 mol dm⁻³ aqueous solution of hydrogen peroxide was dropped on the films, and then 100 µL of 0.6 mol dm⁻³ aqueous solution of sodium hypochloride was added. Manceau et al. made this operation only once. However, we repeated this operation several times about the same film after rinsing with methanol and drying it. The other one is similar operation with the same solutions. The mixings were, however, made in a small petri dish. The polymer coated plates were put on the dish with the coated side down. The distance between the surface of the solution mixture and the polymer films were about 1 cm.

Light from a xenon lamp (Hamamatsu L2275) was irradiated through an optical filter (Toshiba Glass Co. UV-31) that transmits light longer than 310 nm. The illuminances at the surfaces of the polymer films were measured with an illuminometer (AS ONE LM-332).

Ultraviolet-visible-near-infrared (UV-Vis-NIR) spectra of the polymer films were measured with a Shimadzu UV-3600 spectrophotometer.

3-Hexylthiophene, 3-methoxythiophene, and methyl magnesium bromide were purchased from Aldrich, NBS and NiCl₂(dppp) from Tokyo Chemical Industry Co., Ltd., and 1-hexanol, NaHSO₄, hydrogen peroxide, sodium hypochloride, and methanol from, Junsei Chemical Co., Ltd.

3. Results

3.1 Photo-irradiation of Poly(3-heyxylthiophene) and poly(3-hexyloxythiophene)

We compared the rates of photo-degradation of P3HT and P3HOT. If photo-degradation of polythiophene derivatives occurs through the radical mechanism in Fig. 2, the rate of the

photo-degradation should depend on the chemical structures of the side chains. Since the initial step is C-H bond scission at a α -position of alkyl side chains, the rate for polythiophene derivatives having side chains without α -hydrogens should much decrease.

UV-Vis-NIR spectra of P3HT and P3HOT films before and after irradiation with light of the same intensity from a xenon lamp in air are shown in Fig.3 and 4, respectively. The absorption bands around 500 nm for P3HT and around 550 nm for P3HOT are due to the $\pi\pi*$ transition. These bands decreased by the irradiation, and the rate of the decrease for P3HOT was faster than that for P3HT. It is not explained by the mechanism in Fig. 2; the rate for P3HOT should be slower than that for P3HT according to this mechanism.

The absorbance of bands around 1000 and 1650 nm for P3HOT increased by the photo-irradiation. This result agrees with that by Aoyama et al. [20] for poly(3-octyloxythiohene) (P3OOT) . These bands can be assigned to polarons [29]. It was confirmed by doping of P3HOT with iodine vapor. Absorption spectra of a P3HOT film before and after the doping are shown in Fig.5. Absorbance of these bands increased by the doping. Polarons were generated by the photo-irradiation of P3HOT.

Only a small fraction of the decrement of neutral P3HOT was converted to polarons by the photo-irradiation. The increment at 1650 nm and the decrement at 600 nm in absorbance for a film after the doping in Fig. 4 are 0.068 and 0.094, respectively. The ratio of the increment to the decrement is 0.72. On the other hand, the increment, the decrement and the ratio for a film after 24 hours of the

photo-irradiation in Fig.4 are 0.018, 0.247 and 0.075, respectively. The conversion efficiency from neutral P3HOT to polarons by the photo-irradiation is hence about 1/10 of that by the doping. Most of neutral P3HOT decomposed through processes other than the conversion to polarons by the photo-irradiation.

3.2 Exposure of poly(3-heyxylthiophene) and poly(3-hexyloxythiophene) to singlet oxygen

Exposure of P3HT to ${}^{1}O_{2}$ resulted in considerable decrease in the π -conjugation. When only once ${}^{1}O_{2}$ was generated directly on P3HT film following the procedure of Manceau et al. [15], UV-Vis-NIR-spectra showed little change. However, they showed considerable decrease in the absorbance for the $\pi\pi$ *-transition after repeating this procedure several times. Manceau et al. estimated that the reaction between 100 µL of the solutions would produce 60 µmol of ${}^{1}O_{2}$, and this is 900-fold excess [15]. However, we observed that bubbles generated in the solution mixture rose to the solution surface, implying that large part of ${}^{1}O_{2}$ escaped to the atmosphere and only a small fraction of ${}^{1}O_{2}$ reached to the surface of the film.

The direct contact of the solutions to P3HT films may cause effects other than those of ${}^{1}O_{2}$. We then put a P3HT coated glass plate on a petri dish with the coated side down, and mixed the solutions in the dish without contact to the film. Fig. 6 shows the spectra before and after this operation. Absorbance of the $\pi\pi^*$ -transition around 500 nm decreased by repeating the exposure. Hence, ${}^{1}O_{2}$ can reduce the π -conjugation. We also examined reactivity of P3HOT with ${}^{1}O_{2}$. Fig.7 shows absorption spectra of P3HOT before and after exposure to ${}^{1}O_{2}$, which was performed by the second method.. It showed a significant decrease in the $\pi\pi$ * absorption and an increase in polarons. It indicates neutral P3HOT efficiently converted to the polarons by reactions with ${}^{1}O_{2}$. The spectrum after the exposure was stable at least one hour.

4. Discussion

The π -conjugation of P3HOT was reduced faster than that of P3HT by the photo-irradiation in air. This result cannot be explained by the radical mechanism in Fig.2. The reactions are initiated by C-H bond scission of a side chain in this mechanism, and this step will be a rate-determining one. Since the scission of C-H bonds at α -position of the hexyl chains is easier than that of C-H bonds of the hexyloxy chains, the degradation of P3HT should be faster than that of P3HOT.

Exposures of P3HT and P3HOT to ${}^{1}O_{2}$ also induced the reduction of the π -conjugation. The reduction for P3HOT was faster than that for P3HT. In addition, the reactions of ${}^{1}O_{2}$ caused increase in polarons only for P3HOT. These results indicate that ${}^{1}O_{2}$ have a role in the photo-induced degradation of P3HT and P3HOT in air.

The difference in the reactions of ${}^{1}O_{2}$ with P3HT and P3HOT will arise from the difference in their oxidation potentials. They are 0.94 V vs. SCE for P3HT and 0.27-0.73 V vs. SCE for P3HOT, respectively [32-34].

Polarons can be generated by reactions of P3HOT with ${}^{1}O_{2}$, whereas cannot by those of P3HT. This may be evaluated by the following.

The energy associated with excitation of a charge transfer complex of an electron donor and an acceptor is given by

$$E_{\rm e} = IP_{\rm D} - EA_{\rm A} - W, \tag{1}$$

where IP_D and EA_A are the ionization potential of the doner and the electron affinity of the acceptor, respectively [35]. *W* is the Coulombic attraction energy of the complex [35].

 $E_{\rm e}$ can be estimated by the following empirical relation [36]:

$$E_{\rm e}({\rm eV}) = E({\rm D}/{\rm D}^+) - E({\rm A}^-/{\rm A}) + 0.15 \pm 0.10, \tag{2}$$

where $E(D/D^+)$ and $E(A^-/A)$ are the oxidation potential of the donor and the reduction potential of the acceptor, respectively. The values of $E(D/D^+)$ for P3HT [35] and P3HOT[32-34] were reported to be 0.94 and 0.27-0.73 V vs. SCE, respectively. $E(A^-/A)$ was -0.82 V vs. SCE [35]. E_e for P3HT⁺-O₂⁻ has been estimated to be 1.9 eV above the ground state [35], while E_e for P3HOT⁺-O₂⁻ is estimated to be between 1.2 and 1.6 eV with the equation (2).

The electron affinity of oxygen molecule is 0.4480 eV [37], and the excitation energies for ${}^{1}O_{2}$ (${}^{1}\Delta_{g}$) is 0.979 eV[38]. The sum of these energies, which can be used for the formation of the complexes, is 1.43 eV. Hence the formation of P3HT⁺-O₂⁻ with E_{e} of 1.9 eV is difficult, whereas P3HOT⁺-O₂⁻ with E_{e} of 1.2 and 1.6 eV can be generated by the reaction between P3HOT and ${}^{1}O_{2}$.

The formation of polarons by photo-irradiation of P3HOT will be also due to ¹O₂ generated by

photosensitization. The difference in the conversion efficiencies to polarons by the photo-irradiation and exposure to ${}^{1}O_{2}$ will be caused by photoexcitation of polarons. Fig. 8 shows probable mechanisms for the photodegradation of P3HOT: a polaron (P3HT⁺-O₂⁻) is converted to a pair of neutral P3HOT and ${}^{1}O_{2}$ by the photoexcitation, and it returns to a polaron or cause the Diels-Alder addition followed by ring-opening. The polaron can be photoexcited again, and may convert to the neutral pair, whereas the addition and ring-opening is irreversible. The π -conjugation thereby reduces.

Photo-irradiated P3HT will also degrade through the Diels-Alder addition of ${}^{1}O_{2}$. The addition of ${}^{1}O_{2}$ to thienyl ring is probable for a reaction path to ring-opening[39,40].

The degradation through the complex of P3HT⁺ and O₂⁻ has been suggested[41,42,40]. Aoyama et al. have proposed that O₂⁻ abstracts a proton at a α -position of hexyl groups of P3HT⁺, a polyenyl radical of P3HT forms, and this decomposes [43,42]. However, we believe that ring-opening of this radical is improbable. As far as we know, thermal or photolytic reduction of π -conjugation of polyenyl radicals has not been reported. Furthermore, this mechanism cannot explain the degradation of poly(3-phenylthiophene) and poly[3-(1,1-dimethylpropyl)thiophene], which decomposed in the same way as P3HT [43].

Conclusions

 $^{1}O_{2}$ reacted with P3HT, and resulted in the decrease in the π -conjugation in contrary to the results by

Manceau et al. [15]. We believe that ${}^{1}O_{2}$ play a considerable role in the photodegradation of

polythiophene derivatives, and the addition of ${}^{1}O_{2}$ to thienyl ring followed by ring-opening is a most

probable mechanism for the degradation.

Figure Captions

Fig. 1 Ring-opening of thienyl rings through Diels-Alder addition of singlet oxygen[11]

Fig. 2 Ring-opening of thienyl rings through (a) generation of hydroxy radicals, (b) formation of suloxide (thienyl-1-oxide) and sulfone (thienyl-1,1-dioxide) by reaction with hydroxy radicals, and (c)

ring-opening of thienyl-1,1-dioxide [14]

Fig. 3 Photo-absorption spectra of a poly(3-hexylthiophene) film before (---) and after 5 h (---), 15 h (---)

and 24 h (---) irradiation with a xenon lamp

Fig. 4 Photo-absorption spectra of a poly(3-hexyloxythiophene) film before (—) and after 5 h (—), 15 h

(—) and 24 h (—) irradiation with a xenon lamp

Fig. 5 Photo-absorption spectra of a poly(3-hexyloxythiophene) film before (—) and after (—) doping

with iodine vapor

Fig. 6 Photo-absorption spectra of a poly(3-hexylthiophene) film before (---) and after the 2nd (---), 4th

(---), 6th (---), and 8th (---) eposures to singlet oxygen

Fig. 7 Photo-absorption spectra of a poly(3-hexyloxythiophene) film before (—) and after the 2nd (—),

4th (---), 6th (---), and 8th (---) eposures to singlet oxygen

Fig.8 Reactions between poly(3-alkoxythiophene) (P3AOT) and singlet oxygen under photo-irradiation

References

1. Klauk H (2010) Organic thin-film transistors. Chem Soc Rev 39 (7):2643-2666.

2. Chen L-M, Hong Z, Li G, Yang Y (2009) Recent Progress in Polymer Solar Cells: Manipulation of Polymer:Fullerene Morphology and the Formation of Efficient Inverted Polymer Solar Cells. Adv Mater 21 (14-15):1434-1449.

 Christian-Pandya H, Vaidyanathan S, Galvin M (2007) Polymers for Use in Polymeric Light-Emitting Diode: Structure-Property Relationships. In: Skotheim TA, Reynolds JR (eds) Conjugated polymers: processing and applications. 3rd, edn. CRC Press, Boca Raton,

4. Otero TF (2007) Artificial Muscles. In: Skotheim TA, Reynolds JR (eds) Conjugated polymers: processing and applications. CRC Press, Boca Raton,

 5. Arias AC, MacKenzie JD, McCulloch I, Rivnay J, Salleo A (2010) Materials and Applications for Large Area Electronics: Solution-Based Approaches. Chem Rev 110 (1):3-24
6. Nielsen CB, McCulloch I (2013) Recent advances in transistor performance of polythiophenes. Prog Polym Sci 38 (12):2053-2069.

7. Günes S, Neugebauer H, Sariciftci NS (2007) Conjugated Polymer-Based Organic Solar Cells. Chem Rev 107 (4):1324-1338.

8. Jørgensen M, Norrman K, Krebs FC (2008) Stability/degradation of polymer solar cells. Sol Energy Mater Sol Cells 92 (7):686-714.

9. Abdou MSA, Holdcroft S (1995) Solid-state photochemistry of π-conjugated poly(3-alkylthiophenes). Can J Chem 73 (11):1893-1901.

10. Abdou MSA, Holdcroft S (1993) Mechanisms of photodegradation of

poly(3-alkylthiophenes) in solution Macromolecules 26 (11):2954-2962

11. Holdcroft S (1991) A photochemical study of poly(3-hexylthiophene). Macromolecules 24 (17):4834-4838.

12. Holdcroft S (1991) Photochain scission of the soluble electronically conducting polymer: poly(3-hexylthiophene). Macromolecules 24 (8):2119-2121.

13. Abdou MSA, Holdcroft S (1994) Photochemistry of Electronically Conducting Poly(3-alkylthiophenes) Containing FeCl₄⁻ Counterions. Chem Mater 6:962-968

14. Manceau M, Rivaton A, Gardette J-L, Guillerez S, Lemaître N (2009) The mechanism of photo- and thermooxidation of poly(3-hexylthiophene) (P3HT) reconsidered. Polym Degrad Stab 94 (6):898-907.

 Manceau M, Rivaton A, Gardette J-L (2008) Involvement of Singlet Oxygen in the Solid-State Photochemistry of P3HT. Macromol Rapid Commun 29 (22):1823-1827.
Manceau M, Gaume J, Rivaton A, Gardette J-L, Monier G, Bideux L (2010) Further

insights into the photodegradation of poly(3-hexylthiophene) by means of X-ray

photoelectron spectroscopy. Thin Solid Films 518 (23):7113-7118.

17. Caronna T, Forte M, Catellani M, Meille SV (1997) Photodegradation and Photostabilization Studies of Poly(3-butylthiophene) in the Solid State. Chem Mater 9 (4):991-995.

18. Griffini G, Turri S, Levi M (2011) Degradation and stabilization of

poly(3-hexylthiophene) thin films for photovoltaic applications. Polym Bull 66 (2):211-222. 19. Manceau M, Rivaton A, Gardette J-L (2013) Photochemical Stability of Materials for OPV. In: Krebs FC (ed) Stability and Degradation of Organic and Polymer Solar Cells. John Wiley & Sons, Ltd., Chichester, pp 71-108

20. Aoyama Y, Yamanari T, Koumura N, Tachikawa H, Nagai M, Yoshida Y (2013) Photo-induced oxidation of polythiophene derivatives: Dependence on side chain structure. Polym Degrad Stab 98 (4):899-903.

21. Sai N, Leung K, Zador J, Henkelman G (2014) First principles study of photo-oxidation degradation mechanisms in P3HT for organic solar cells. PCCP 16 (17):8092-8099.

22. D'Auria M (2011) Photochemical and Photophysical Behavior of Thiophene. Adv Heterocycl Chem 104:127-390

23. Turro NJ (1978) Modern Molecular Photochemistry. The Benjamin/Cummings Publishing Company Inc., Menlo Park

24. Wasserman HH, Strehlow W (1970) Reactions of singlet oxygen with thiophenes. Tetrahedron Lett 11 (10):795-796.

25. Skold CN, Schlessinger RH (1970) The reaction of singlet oxygen with a simple thiophene. Tetrahedron Lett 11 (10):791-794.

26. Sugimoto R, Takeda S, Gu HB, Yoshino K (1986) Preparation of Solubule Polythiophene Derivatives Utilizing Transition Metal Halides as Catalysts and Their Property. Chem Express 11 (1):635-638

27. Loewe RS, Ewbank PC, Liu J, Zhai L, McCullough RD (2001) Regioregular, Head-to-Tail Coupled Poly(3-alkylthiophenes) Made Easy by the GRIM Methos: Investigation of the Reaction and the Orgin of Regioselectivity. Macromolecules 34 (13):4324-4333

28. Koeckelberghs G, Vangheluwe M, Samyn C, Persoons A, Verbiest T (2005) Regioregular Poly(3-alkoxythiophene)s: Toward Soluble, Chiral Conjugated Polymers with a Stable Oxidized State. Macromolecules 38:5554-5559

29. Hatakeyama K, Koizumi H, Ichikawa T (2009) Stability of a Conductive State of Poly(3-alkoxythiophene)s. Bull Chem Soc Jpn 82 (2):202-205

30. Seliger HH (1960) A photoelectric method for the measurement of spectra of light sources of rapidly varying intensities. Anal Biochem 1 (1):60-65.

31. Khan AU, Kasha M (1963) Red Chemiluminescence of Molecular Oxygen in Aqueous

Solution. J Chem Phys 39 (8):2105-2106

32. Köse M (2011) Theoretical prediction of ionization/oxidation potentials in conjugated polymers. Theoretical Chemistry Accounts: Theory, Computation, & Modeling 128 (2):157-164.

33. Shi C, Yao Y, Yang, Pei Q (2006) Regioregular Copolymers of 3-Alkoxythiophene and Their Photovoltaic Application. J Am Chem Soc 128 (27):8980-8986.

34. Scharber MC, Mühlbacher D, Koppe M, Denk P, Waldauf C, Heeger AJ, Brabec CJ (2006) Design Rules for Donors in Bulk-Heterojunction Solar Cells—Towards 10 % Energy-Conversion Efficiency. Adv Mater 18 (6):789-794.

35. Abdou MSA, Orfino FP, Son Y, Holdcroft S (1997) Interaction of Oxygen with Conjugated Polymers: Charge Transfer Complex Formation with Poly(3-alkylthiophenes). J Am Chem Soc 119 (19):4518-4524.

36. Rehn D, Weller A (1970) Bindungszusatnd und Fluoreszenzspektren von Heter-Eximeren. Z Phys Chem 69:183-200

37. Ervin KM, Anusiewicz I, Skurski P, Simons J, Lineberger WC (2003) The Only Stable State of O_2 . Is the X ${}^{2}\Pi_{g}$ Ground State and It (Still!) Has an Adiabatic Electron Detachment Energy of 0.45 eV. J Phys Chem A 107 (41):8521-8529.

38. Badger RM, Wright AC, Whitlock RF (1965) Absolute Intensities of the Discrete and Continuous Absorption Bands of Oxygen Gas at 1.26 and 1.065 μ and the Radiative Lifetime of the 1Δg State of Oxygen. The Journal of Chemical Physics 43 (12):4345-4350.

39. Song X, Fanelli MG, Cook JM, Bai F, Parish CA (2012) Mechanisms for the Reaction of Thiophene and Methylthiophene with Singlet and Triplet Molecular Oxygen. The Journal of Physical Chemistry A 116 (20):4934-4946.

40. Chen L, Yamane S, Mizukado J, Suzuki Y, Kutsuna S, Uchimaru T, Suda H (2015) ESR study of singlet oxygen generation and its behavior during the photo-oxidation of P3HT in solution. Chem Phys Lett 624:87-92.

41. Chen L, Mizukado J, Suzuki Y, Kutsuna S, Aoyama Y, Yoshida Y, Suda H (2014) An ESR study on superoxide radical anion generation and its involvement in the photooxidative degradation of poly-3-hexylthiophene in chlorobenzene solution. Chem Phys Lett 605–606 (0):98-102.

42. Aoyama Y, Yamanari T, Murakami TN, Nagamori T, Marumoto K, Tachikawa H, Mizukado J, Suda H, Yoshida Y (2015) Initial photooxidation mechanism leading to reactive radical formation of polythiophene derivatives. Polym J 47 (1):26-30.

43. Aoyama Y, Hatakeyama K, Mukai T, Koizumi H (2013) Electronic Properties of Poly[3-(1,1-dimethylpropyl)thiophene] and Poly(3-cyclohexylthiophene). Bull Chem Soc Jpn 86 (1):51-56

<u>Errata</u>

Mechanisms of photo-induced degradation of polythiophene derivatives: Re-examination of the role of singlet oxygen Hiroki Ohta and Hitoshi Koizumi

Manuscript accepted for publication in Polymer Bulletin.on 21 October 2016.

| | Original Text | Correction |
|-------------------------|---------------|--------------|
| page 4, line 5 | impurity | residual |
| page 6, lines 15 and 18 | hypochloride | hypochlorite |
| page 8, line 17 | Fig.4 | Fig.5 |

page 15, reference 3

Original Text

Christian-Pandya H, Vaidyanathan S, Galvin M (2007) Polymers for Use in Polymeric Light-Emitting Diode: Structure-Property Relationships. In: Skotheim TA, Reynolds JR (eds) Conjugated polymers: processing and applications. 3rd, edn. CRC Press, Boca Raton,

Correction

Christian-Pandya H, Vaidyanathan S, Galvin M (2007) Polymers for Use in Polymeric Light-Emitting Diode: Structure-Property Relationships. In: Skotheim TA, Reynolds JR (eds) Conjugated polymers: processing and applications. 3rd, edn. CRC Press, Boca Raton, pp.5.3-5.35

page 15, reference 4 Original Text

Otero TF (2007) Artificial Muscles. In: Skotheim TA, Reynolds JR (eds) Conjugated polymers: processing and applications. CRC Press, Boca Raton,

Correction

Otero TF (2007) Artificial Muscles. In: Skotheim TA, Reynolds JR (eds) Conjugated polymers: processing and applications. CRC Press, Boca Raton, pp.16.1-16.33

page 16, reference 21 <u>Original Text</u> Sai N, Leung K, Zador J, Henkelman G (2014) First principles study of photo-oxidation degradation mechanisms in P3HT for organic solar cells. PCCP 16 (17):8092-8099.

Correction

Sai N, Leung K, Zador J, Henkelman G (2014) First principles study of photo-oxidation degradation mechanisms in P3HT for organic solar cells. Phys Chem Chem Phys 16 (17):8092-8099.

page 17, reference 36

Original Text

Rehn D, Weller A (1970) Bindungszusatnd und Fluoreszenzspektren von Heter-Eximeren. Z Phys Chem 69:183-200

Correction

Rehn D, Weller A (1970) Bindungszustand und Fluoreszenzspektren von Heter-Eximeren. Z Phys Chem 69:183-200























