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STUDIES ON PHOTOPHYSICAL PROCESSES IN

POLY (N-VINYLCARBAZOLE) AND RELATED COMPOUNDS

(ボリ(N-ビニルカルバゾール)及びその関連化合物の) (光物理過程に関する研究

AKIRA ITAYA

List of Publications

The papers published by the author concerning this thesis are as follows.

- (1) Heterogeneous Triplet-Triplet Annihilation in Poly-Nvinylcarbazole Films Doped with Naphthalene and Fluorene.
 A. Itaya, K. Okamoto, and S. Kusabayashi
 Chem. Lett., 1975, 833.
- (2) Magnetic Field Effect on the Photoconductivity of Poly-Nvinylcarbazole.

K. Okamoto, N. Oda, A. Itaya, and S. Kusabayashi Chem. Phys. Lett., 35, 483 (1975).

(3) Magnetic Field Effect on the Exciplex Fluorescence of Poly-N-vinylcarbazole.

K. Okamoto, A. Itaya, and S. Kusabayashi

Chem. Lett., 1976, 99.

(4) The Photoconductivity of Poly(N-vinylcarbazole). VII. The Effect of the Singlet and Triplet Quenchers on the Photoconductivity.

K. Okamoto, N. Oda, A. Itaya, and S. Kusabayashi

Bull. Chem. Soc. Jpn., 49, 1415 (1976).

(5) Triplet Excitation Energy Transfer in the Vinyl Polymers with Pendant Carbazolyl Groups.

A. Itaya, K. Okamoto, and S. Kusabayashi

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(6) Emission Spectra of the Vinyl Polymers with Pendant Carbazolyl Groups.

> A. Itaya, K. Okamoto, and S. Kusabayashi Bull. Chem. Soc. Jpn., 49, 2082 (1976).

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(7) Polymerization of N-Vinylcarbazole, 5-Vinyl-benzo[b]carbazole, and 7-Vinyl-benzo[c]carbazole.

K. Okamoto, M. Yamada, A. Itaya, T. Kimura, and

S. Kusabayashi

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(8) Singlet Excitation Energy Transfer in the Vinyl Polymers with Pendant Carbazolyl Groups.

A. Itaya, K. Okamoto, and S. Kusabayashi

Bull. Chem. Soc. Jpn., 50, 22 (1977).

(9) Emission Spectra of the Vinyl Polymers with Pendant Phenanthryl Groups.

A. Itaya, K. Okamoto, and S. Kusabayashi

Bull. Chem Soc. Jpn., 50, 52 (1977).

(10) Emission Spectra of the Vinyl Polymers with Pendant Benzocarbazolyl Groups.

A. Itaya, K. Okamoto, and S. Kusabayashi

Bull. Chem. Soc. Jpn., 51, 79 (1978).

(11) Electronic Spectra of 1,3-Di(N-carbazolyl)propane in the Amorphous State

A. Itaya, K. Okamoto, and S. Kusabayashi

Chem. Lett., 1978, 131.

(12) Photo-oxidation Effect of Poly(N-vinylcarbazole) film on Photoconductivity and Electronic Spectra.

A. Itaya, K. Okamoto, and S. Kusabayashi

Bull. Chem. Soc. Jpn., 52, 2218 (1979).

(13) The Singlet-Excitation-Energy Migration in an Amorphous State Film of 1,3-Di(N-carbazolyl)propane.

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i-PrCz



DCzP



VCz Olig. I ; n~4 VCz Olig. II; n~10



PVCz



-(CH-CH₂)_n-

PVPh

COOCH₃

COOCH₃



PACz



PPhMA

Perylene

CC1₃

CC1₃

TCB





-(CH-CH₂)_n-

PCzEVE

-(CH-CH₂)_n-1 0 CH₂



PPhEVE

N(CH₃)₂

N(CH₃)₂

TMPD



P7VBCz

5EBCz



 $\bigcirc \bigcirc$

DMTP

Naphthalene

List of chemical structures and abbreviations of compounds used in this thesis.

INTRODUCTION

Recently, many kinds of organic compounds with special function have been developed. Among them, vinyl polymers with large pendant aromatic groups have received much attention because of high photoconductivity and interesting emission properties. Photoconductivity of this kind of polymers has been extensively studied with regard to poly(N-vinylcarbazole)(PVCz).¹⁾ However, electrical properties of a PVCz film still remain to be clarified. For the basic understanding of photoconductive properties, it is essential to have the information on the processes from lightabsorption to photo-carrier-generation (photophysical process).

Photophysical processes of aromatic hydrocarbons have been studied in detail and many phenomena have been found. They include excimer and exciplex formation, electron transfer, energy migration and transfer, and so on.²⁾ In the case of aromatic vinyl polymers, intramolecular excimer formation and energy migration along the polymer chain are particularly important and interesting.

Yanari et al.³⁾ were the first to suggest that the fluorescence emission of polystyrene(PSt) was excimeric in nature. The configurational basis of the intramolecular excimer formation was first described by Hirayama in his study on diphenyl and triphenyl alkanes.⁴⁾ His "n=3 rule" states that the intramolecular excimer formation occurs only when the aromatic chromophores are separated by a main chain segment of three carbon atoms. Vala et al. then studied on excimer formation in the diphenyl alkanes and paracyclophane series, atactic and isotactic PSt and atactic poly(1-vinylnaphthalene).⁵⁾ At the present time, the mechanism

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for the excimer formation in aromatic vinyl polymers is considered as follows. In rigid or high viscous solution and solid film, a singlet exciton generated in the polymer by exciting-light migrates over the pendant chromophores along the polymer chain and is trapped at an excimer-forming site, whose the chain conformation is geometrically suitable for excimer formation. On the other hand, in fluid solutions with low viscosity, the excimer formation is controlled by a dynamic process of chromophores. A chain conformation changes to another one such as an excimer-forming chain conformation by collision with solvent molecules. Such a conformational change occurs within the residence time of the migrating exciton on a particular chromophore with some probability, resulting in a rotational formation of excimer. Therefore, the excimer formation is closely related with the dynamic processes of polymers in solution. Studies on the excimer fluorescence provide many information on the segmental motion of polymer chains and on molecular arrangement in solution.⁶⁾

Singlet and triplet excitation energies migrate along the polymer chain because of a one-dimensional arrangement of pendant chromophores along the main chain in high density.⁷⁾ Singlet excitation energy migration in a polymer film was first investigated by Klöpffer.⁸⁾ He reported that the results of fluorescence quenching experiments in a PVCz film are consistent with a hopping model of monomer-exciton migration and that the singlet exciton migration is limited by excimer-forming sites.

Singlet exciton migration along the polymer chain in dilute and rigid solutions was ascertained by the experiment of energy transfer in homopolymer donor - low-molecular-weight acceptor

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systems. In this case, a singlet exciton migrates by a series of random walk due to Förster interactions between nearest neighbors.⁹⁾

From an observation of delayed fluorescence caused by triplettriplet annihilation and quenching experiments, Fox et al. showed that a triplet exciton migrates along the polymer chain in poly(1vinylnaphthalene) both in rigid solution and in the solid film at 77 K.¹⁰⁾ Thereafter, the triplet exciton migration has been observed for a number of aromatic vinyl polymers and values of the migration coefficient have been obtained by the quenching experiment.

Considering that photo-carrier-generation in the lowest $\pi-\pi^*$ absorption region is attributed to the interaction between a migrating excited state and any electron-accepting impurity,¹¹⁾ it is suggested that phenomena of the excimer formation and the above-mentioned energy migration are closely related to photo-carrier-generation and charge-carrier migration, because excimer-forming sites act as traps for the electronic energy migration and might act as traps for charge-carrier migration. Since both the triplet energy migration and the charge-carrier migration are due to the short-range interaction (electron-exchange interaction), the study on the triplet energy migration might provide some information on the charge-carrier migration.

In recent years, studies on ionic photodissociation in dilute homogeneous solution have been carried out in detail.¹²⁾ The ionic photodissociation is the basis of the photo-oxidationreduction reactions and closely related to the fields of solar cells and photosynthesis. In such fields, distribution of photoexcited species is non-uniform and the local concentration is high.

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That is, their systems are considered to be a kind of molecular aggregate ones. Therefore, it is interesting to see whether the results obtained in dilute homogeneous solution hold also for molecular aggregate systems or not.¹³⁾ The basic information on the above-mentioned problem is also given by the study of the photophysical process in aromatic vinyl polymers, which are a kind of molecular aggregate systems with respect to pendant aromatic groups.

Furthermore, the photophysical process of polymers is also closely related to phenomena of photostabilization and photo-degradation of polymers.¹⁴⁾

In this thesis, the author describes the photophysical process of aromatic vinyl polymers with different chemical and fine structures in solution and in film, and a few effects on the photo-carrier-generation process of a PVCz film.

Part I deals with fluorescence spectra of vinyl polymers with large pendant aromatic groups in solution. In chapter 1, 5-vinyl-benzo[b]carbazole and 7-vinyl-benzo[c]carbazole, which have more bulky pendant groups than N-vinylcarbazole(VCz), were synthesized. These two monomers and VCz were polymerized under various conditions. Solubilities, glass transition temperatures, and NMR spectra of these polymers indicated that the stereoregularity of the polymers varies depending on the polymerization method. In chapter 2, a relation between the fluorescence spectra and the stereoregularity of polymers has been obtained. From this relation, the conformational structures of the second and sandwich-like excimers were discussed. In chapter 3, the emission spectrum of poly[2-(N-carbazoly1)ethyl vinyl ether] with carbazolyl chromophores widely spaced on the

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main chain by -O-CH₂-CH₂- bonds has been investigated in comparing with those of PVCz and its related compounds. In chapter 4, emission spectra of the vinyl polymers with phenanthryl groups have been investigated in solution. The fluorescence spectra are of interest, because phenanthryl chromophore shows no excimer fluorescence in concentrated solutions as in the case of carbazolyl chromophore.

Part II deals with the photophysical process in polymer films. In chapter 1, singlet-excitation-energy migration in films of the vinyl polymer with pendant carbazolyl groups has been investigated in connection with chemical structure and tacticity of the polymers. The spectroscopic properties of an amorphous film of 1,3-di(N-carbazolyl)propane, which is a dimeric model compound of PVCz, has also been investigated and compared with those of the amorphous polymer films. In chapter 2, triplet-excitationenergy migration in films of the vinyl polymer with pendant carbazolyl groups has been investigated by measuring delayed emission spectra in films and in rigid solutions at 77 K.

In the last part, part III, in order to clarify the photocarrier-generation mechanism considered most appropriate at the present stage, the author has investigated the following effects on the photoconductivity of a PVCz film. In chapter 1, it is concluded, from the effects of triplet and singlet quenchers on the photoconductivity of PVCz and brominated PVCz films, that charge carriers are generated via the singlet exciton rather than the triplet one. In chapter 2, it is concluded, from magnetic field effects on the photoconductivity of PVCz films undoped and doped with dimethyl terephthalate, that the carriergeneration process with participation of a non-relaxed exciplex

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state plays an important role in the photo-carrier-generation in the lowest $\pi-\pi^*$ absorption region. In chapter 3, in order to elucidate what kind compound acts as an electron-accepting impurity, the photo-oxidation effects on the photoconductivity and electronic spectra of PVCz films have been investigated. From the results, it is concluded that the photo-oxidation products formed near the surface of PVCz films act as an electronaccepting impurity in the photo-carrier-generation mechanism.

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PART I

EMISSION SPECTRA OF AROMATIC VINYL POLYMERS IN SOLUTION

INTRODUCTION

It is known that aromatic vinyl polymers show excimer fluorescence in dilute fluid solution due to the intramolecular interaction between chromophores in a polymer chain. The excimer fluorescence provides some information on molecular motion or conformation of the polymer chain in solution. The excimer fluorescence is closely related also to the photoconduction of the polymers, because the excimer forming site acts as an energy trap for the migration of a singlet exciton.

Considerable data on the emission properties of PVCz have been accumulated because of an interest in high photoconductive polymers.¹⁻³³⁾ However, no relation between the tacticity and the fluorescence spectra of PVCz has been reported. Concerning the vinyl polymers with small aromatic chromophores such as a phenyl group, the relation has been reported in several investigations.³⁴⁻⁴¹⁾ However, such a relation has never hitherto been studied for the vinyl polymers with large and bulky aromatic chromophores such as carbazole and benzocarbazole.

In chapter 1, vinylcarbazole and vinyl monomer with benzocarbazolyl groups newly synthesized were polymerized under

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various conditions. A relation between the stereoregularity of the vinyl polymers and polymerization methods was found at the first time in this kind of polymers with large pendant groups.

In chapter 2, the fluorescence spectra of the polymers obtained in chapter 1 have been investigated in solution and a relation between the fluorescence spectra and the stereoregularity of the polymers is discussed.

In chapter 3, the fluorescence spectrum of poly[2-(Ncarbazolyl)ethyl vinyl ether] has been investigated in comparing with those of PVCz and its related compounds.

In chapter 4, fluorescence spectra of the vinyl polymers with phenanthryl groups, which show no excimer fluorescence in concentrated solutions as in the case of carbazolyl chromophore, have been investigated in order to elucidate characteristics of the excimer formation of aromatic vinyl polymers, model compounds of which show no excimer fluorescence.

Chapter 1

Polymerization of N-Vinylcarbazole, 5-Vinyl-benzo[b]carbazole, and 7-Vinyl-benzo[c]carbazole

Introduction

The polymerization of N-vinylcarbazole(VCz) has widely been investigated.⁴²⁾ This monomer can be easily polymerized by many kinds of initiators such as free-radical, cationic, Ziegler-Natta, and electron acceptor ones. Heller et al. have pointed out that VCz did not undergo a coordination polymerization with Ziegler-Natta catalyst systems, but merely underwent a cationic one. 43) The stereoregularity and the crystallinity of PVCz have been reported to be independent of the polymerization method from the NMR spectra and X-ray diffraction patterns. 44-47) Addition of a monomer onto a growing chain end has, therefore, been considered to be controlled sterically due to the bulkiness of the carbazolyl group, irrespective of the nature of the growing chain end. 44) This has been supported by the fact that poly(trityl methacrylate) prepared free radically has a pronounced isotacticity due to the steric hindrance of the pendant group. 48) On the other hand, this is clearly different from the polymerization behavior of N-acryloylcarbazole with a carbazolyl group widely spaced on a vinyl bond; the stereoregularity of this polymer has been reported to be dependent on the polymerization method. 49) It is, therefore, of interest to re-investigate in detail the relation between the stereoregularity of the vinyl

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polymers containing large pendant π -electron groups and the polymerization method.

In this chapter, 5-vinyl-benzo[b]carbazole(5VBCz) and 7vinyl-benzo[c]carbazole(7VBCz) were synthesized and polymerized by free-radical, cationic, and Ziegler-Natta initiators. The solubilities, glass transition temperatures, and NMR spectra of PVCz, poly(5-vinyl-benzo[b]carbazole)(P5VBCz), and poly(7vinyl-benzo[c]carbazole)(P7VBCz) were examined.

Experimental

 $5-\beta$ -Chloroethyl-benzo[b]carbazole(5CEBCz) was synthesized from 5H-benzo[b]carbazole, which was prepared from sodium 2naphthol-l-sulfonate.⁵⁰⁾ To a suspension of 5H-benzo[b]carbazole in 400 ml of acetone were added 20 g of β -chloroethyl p-toluenesulfonate and a solution of 10 g of NaOH in 7.5 ml of water with stirring. The mixture was stirred at 60 °C for 10 h; then another 10 g of β -chloroethyl p-toluenesulfonate and a solution of 4 g NaOH in 1.5 ml of water were added and stirring was continued for an additional 10 h. The resulting precipitate was filtered off. After the unreacted β -chloroethyl p-toluenesulfonate was separated off, the filtrate was evaporated and the residue chromatographed on silica gel using benzene as an eluent to give 7 g (54%) of the product. Recrystallization from methanol gave colorless needles of 5CEBCz, mp 121-122 °C.

To a suspension of 5CEBCz(7.2 g; 0.026 mol) in 360 ml of ethanol was added 75 ml of a 25% KOH-methanol solution. The mixture was refluxed for 15 h and poured into water. After standing over a night, a precipitate obtained was filtered and

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washed with water. Repeated recrystallizations from methanol gave pale yellow needles of 5VBCz(4.0 g, 62%), mp 89.5-90.5 °C.

Colorless plates of 7VBCz was synthesized from 7H-benzo[c]carbazole by the procedure described in the synthesis of 5VBCz. The product was recrystallized twice from methanol, 56% yield; mp 88-89 °C.

These new materials (5CEBCz, 5VBCz, 7CEBCz, and 7VBCz) were identified by elemental analysis, NMR and IR spectra.

All polymerization reactions were carried out using standard procedures under nitrogen or under vacuum.^{51,52)} Reagents were carefully purified and dried. Polymers were isolated by precipitation with methanol and purified by repeated reprecipitations followed by drying under vacuum.

The polymer molecular weights (M_n) were determined in ethylene chloride solution at 40 °C using a vapor pressure osmometer. Glass transition temperatures (T_g) were determined with a Shimadzu DSC-20 differential scanning calorimeter at the heating rates of 10-15 °C/min. The T_g for each sample was taken as the average of four or five determinations. NMR spectra were measured in deuteriochloroform at 60 °C and in tetradeuterioethylene bromide at 60-140 °C with a JEOL MH-100 spectrometer.

Results and Discussion

(1) Polymerization.

Representative polymerization results of VCz are shown in Table I-1. All of the PVCz samples are soluble in ordinary solvents irrespective of the polymerization method. Clear differences in T_g are, however, observed among the samples prepared free radically, cationically in a polar solvent, and

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cationically in a nonpolar one; the T_g increases in this order. The T_g is supposed to be independent of the M_n in the range of $M_n > 100000.^{54}$ It is considered that there is no serious difference in molecular weight distribution among these samples.^{55,56} Therefore, the differences in T_g are not attributable to the differences in M_n , but to some differences in the stereoregularity which will be discussed below.

5VBCz is easily polymerized by cationic initiators, as is shown in Table I-2. The yields, softening ranges(T_s), and M_n of the polymers increase with a decrease in polymerization

Table I-1,

Characterization of PVCz Prepared by Various Methods

			Polymerization	method				
		Concn	Initiator	Temp	 T	Ma	<i>I</i> _{H/}	′L ^b
Sample	Solvent	M	(mol %)	°C	°C	$\times 10^4$	-CHR-	-CH2-
Α	Benzene	0.5	AIBN (1)	70	228	20	0.35	0.38
в	Bulk		Thermally	80	226		0.46	
С	Toluene	0.26	$BF_{3}OEt_{2}(1)$	20	248	10	0.97	0.83
D	$(CH_2Cl)_2$	0.09	$BF_{3}OEt_{2}(1)$	20	238	11	0.58	0.60
\mathbf{E}	Toluene	0.09	$BF_{3}OEt_{2}(1)$	-78	244	77	1.00	
F	Toluene	0.2	$AlEt_3/TiCl_4$ (2/2)	20	245	33	0.73	

^a Determined from the specific viscosity.^{53 b} See text. The $I_{H/L}$ value for methylene signal could not be obtained from the spectra measured in CDCl₃ at 60 °C because of the broadness.

Table I-2.

Polymerization Results of 5VBCz

No.	Initiator (mol %)	Concn, M ^a	Time, h	Temp, °C	Yield, %	<i>T</i> ₅, °C	M_{n}
1	BF ₃ OE t ₂ (2)	0.14	4	40	14	212-219	
2	BF_3OEt_2 (2)	0.14	4	-20	83	230-240	1750
3	$BF_3OEt_2(2)$	0.42	3	-33	85	248 - 258	2000
4	BF_3OEt_2 (2)	0.42	3	-40	85	258-268	2030
5	$BF_{3}OEt_{2}(1)$	0.14	4	-78	93	264 - 274	2120
6	$BF_3OEt_2(2)$	0.10	6	-90	93	260 - 272	2030
7	$AlEtCl_2(2)$	0.14	4	-78	86	243 - 253	1770
8	$AlEt_3/TiCl_4$ (2/2)	0.20	10	20	71	245 - 250	
9	BPO (2)	0.35	15	80	0		
10	BPO (2)	Bulk	24	100	37	200-208	
11	AIBN (2)	0.34	48	75	30	230 - 240	3060
12	AIBN (2)	0.40	35	80	47	240 - 248	
13	AIBN (2)	Bulk	48	100	56	290-300	12000
14	None	Bulk	48	100	Trace		

^a Solvent: ethylene chloride for No. 3 and toluene for the others.

temperature. At the temperatures below -40 °C, the polymers are obtained almost quantitatively and the M_n values are constant at 2000-2100(DP=8-9). These polymers are readily soluble in ordinary solvents such as benzene and THF, and their T_s and T_g are 260-270 and 235 °C, respectively. Several attempts to obtain high molecular weight polymers by the use of cationic initiators have been unsuccessful. Ziegler-Natta initiators also give the low molecular weight polymers in high yields.

5VBCz is also polymerized by free-radical initiators, although the yields are very low as compared with those of cationic polymerizations. Benzoyl peroxide(BPO) gives the polymer having a low value of T_S only in bulk polymerization. Azobisisobutyronitrile(AIBN) gives the polymers even in solution. The yields increase with an increase in the monomer concentration up to 50% in the saturated solution(0.4 M), but the T_s and probably the Mn also hardly increase. The AIBN-initiated bulk polymerization at 95-100 °C gives the fairly high molecular weight polymer (DP=50) in an about 50% yield. The polymer thus obtained is pale yellow, and soluble in THF, but not in benzene, and the ${\tt T}_{\rm cr}$ is 271 °C. It is noted that the T_S values of the cationically polymerized samples(cationic polymers) are much larger than those of the free-radically polymerized samples (radical polymers) with the corresponding M_n values.

5VBCz is not polymerized by anionic initiators. This is due to the same reason as the case with VCz. $^{57)}$

7VBCz is easily polymerized by free-radical initiators. Some polymerization results are shown in Table I-3. The yields and the M_n depend a little both on monomer concentration and on temperature. The M_n values are much larger than those of the PV5BCz samples prepared under the similar conditions, but much

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smaller than those of the PVCz ones.

7VBCz is polymerized by cationic initiators in high yields with an exception of No. 7 in Table I-3. The T_g , M_n , and solubility of the cationic polymers vary remarkably with the polymerization condition. The M_n increases with an increase in the concentration of monomer or initiator, and there is an optimal polymerization temperature of the M_n at about -40 °C. The T_g increases with an increase in polymerization temperature and decreases with an increase in the solvent polarity. The T_g of the cationic polymers are much higher than those of the freeradical ones. These results are not explained by the differences in the M_n , although the T_g depends a little on the M_n even in the range of fairly high M_n . The polymers prepared at temperatures below -40 °C are readily soluble in ordinary solvents. However, the solubility becomes poor with an increase in the temperature, in spite of a decrease in the M_n . The polymers insoluble in all

Table 1-3.

	-	Dolumoninat	ion motho	a		•	Charac	cteriza	ation of p	olymers	
		r olymerizat.			X 72 X X			Solu	ıbility ^b	$I_{ m H/}$	L ^c
No.	(mol %)	Concn, M ^a	Time, h	°C	¥ ield, %	T _g , ℃	M _n	Bz	THF	-CHR-	CH2
1	AIBN (2)	0.10	5	75	39		11 000	0	0		
2	AIBN (2)	0.28	5	75	65	271	14 000	0	0	0.32	
3	AIBN (2)	0.56	5.5	75	92	271	$15\ 000$	0	0	0.33	0.39
4	$BF_{3}OEt_{2}(2)$	0.14	1.5	-78	99	283	>25 000	0	0	0.54	0.52
5	$BF_3OEt_2(2)$	0.14	1	-55	98	288	>25 000	0	0		
6	$BF_3OEt_2(2)$	0.14	1.5	-40	99	292		х	Δ		
7	$BF_3OEt_2(2)$	0.05	1	-78	19		10 000	0	0		
8	$BF_3OEt_2(2)$	0.05	3	-50	92	280	16 000	0	0	0.63	
9	$BF_3OEt_2(2)$	0.05	1	-40	99	287	20 000	0	0	0.68	
10	$BF_3OEt_2(2)$	0.05	1	0	93	287	14 000	0	0		
11	$BF_3OEt_2(2)$	0.05	1	10	94		8 500	0	Ō	0.99	
12	$BF_3OEt_2(2)$	0.05	1	25	93	291		х	Δ		
13	$BF_{3}OEt_{2}(1)$	0.15	3	10	99	289		Δ	0	0.92	0.78
14	$BF_{3}OEt_{2}(1)$	0.10	1	10	93	278		0	0	0.54	0.50
15	$AlEt_3/TiCl_4$ (1/1)	0.20	10	20	91	292		X	Δ		
16	AlEt ₃ /TiCl ₄ (3/2)	0.10	7	35	99			0	0	0.56	
17	AlEt ₃ /TiCl ₃ (1/1)	0.20	10	20	90	282		0	0	0.53	

Polymerization Results of 7VBCz

^a Solvent: ethylene chloride for No. 14 and toluene for the others. ^b (O) soluble; (Φ) soluble but opaque in cold benzene; (Δ) swollen; (\times) insoluble. ^c See Table I.

of the solvents examined are obtained at temperatures above a critical one, which is raised with an increase in the concentration of initiator or monomer. The solubility is also dependent on the solvent polarity; ethylene chloride gives the polymer having better solubility than toluene. Thus, there is a roughly parallel relationship between the $T_{\rm g}$ and the solubility.

7VBCz is easily polymerized by Ziegler-Natta initiators, but not polymerized anionically.

These three monomers are rich both in the electron-resonance stabilization and in the electron-donating character of the pendant groups. Their magnitudes are considered to be in the order of 5VBCz > 7VBCz > VCz. They were, therefore, predicted to be easily polymerized free radically and cationically to the high molecular weight polymers. In the case of free-radical polymerization, both the yields and the M_n are in the order of VCz > 7VBCz > 5VBCz. This is opposite to the order predicted from the electron resonance stabilization. The radical reactivity of these monomers seems to be controlled by the steric hindrance effect rather than by the resonance effect.

In the case of cationic polymerization, the yields are almost quantitative and the M_n is in the order of VCz > 7VBCz >> 5VBCz. The cationic reactivity of the growing chain end is predicted to be in the inverse order of that of the monomer, as is the case with vinyl ethers.⁵⁸⁾ This may be a reason for the latter fact (the order of M_n), if the approach of an incoming monomer to the growing chain end followed by the separation of the counterion is not the rate-determining step of the propagation reaction. However, this assumption seems very unlikely for these monomers. Therefore, the order of M_n suggests that the cationic reactivity

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of the growing chain end may be controlled by the steric hindrance effect rather than by the electron-donating effect. The cationic polymerization of 5VBCz, the most bulky monomer among these used, gives only the low molecular weight polymer in a high yield. This suggests that the cationic reactivity of the growing chain end decreases drastically with increasing sequence of the growing chain, resulting in the transfer reaction to a counterion and/or a monomer.

(2) NMR Spectra.

The T_g and solubility data mentioned above suggest that the stereoregularity and/or the crystallinity of the polymers vary depending on the polymerization method. The IR, NMR, and X-ray diffraction spectra of these polymers also were examined. The IR and X-ray diffraction spectra are essentially the same, irrespective of the polymerization method. All of the samples which were precipitated from solution and have never experienced heat treatment are of low crystallinity. The 100-MHz NMR spectra are shown in Figs. I-1-I-3. The spectra of P5VBCz and P7VBCz were assigned according to the case of PVCz.⁵⁹⁾ The assignment





Fig. I-2. NMR spectra(100 MHz) of P5VBCz in deoteriochloroform at 60 °C. 1 and 2 are for the samples of No. 13 and 5 in Table I-2, respectively.

NMR spectra (100 MHz) of PVCz in tetradeuterioethylene of No. 13 and 5 in Table I-2, bromide at 125 °C. The upper and lower curves are for the samples respectively. of No. C and A in Table I-1, respectively.

is listed in Table I-4. Both positions and relative intensities of all the signals are independent of temperature in the range of 60 to 140 °C, although the resolution of the signals of methylene protons are appreciably improved with an increase in temperature.

In the case of PVCz and P7VBCz, the aromatic regions of the spectra ($\tau < 6.0$) vary only a little with the polymerization method. The intensity of a peak appearing at about 5 τ always corresponds to about one proton. On the other hand, the methylene and methine regions of the spectra, which are composed of two peaks respectively, vary appreciably with the polymerization method; the intensity ratio of the higher field peak to the lower field one($I_{H/L}$) is dependent on the polymerization method, as is shown in the last columns in Tables I-1 and I-3, although the positions of the signals are hardly dependent on it. The $I_{H/L}$ value for methylene protons is in agreement with that for a

methine proton, taking into consideration the error (\pm 5%) in resolving a methylene signal into two components. In both PVCz and P7VBCz, the samples prepared cationically in a nonpolar solvent at room temperature have the largest $I_{H/L}$ value(unity) and the freeradical samples have the smallest value(one third). In the cationic polymerization, the $I_{H/L}$ value decreases appreciably in



NMR spectra (100 MHz) of P7VBCz in tetradeuterioethylene bromide at 125 °C. The upper, middle, and lower curves are for the samples of No. 13, 4, and 3 in Table I-3, respectively.

Table I-4. Chemical Shifts (τ) of Protons^a

Sample	Aromatic	Methine	Methylene
PVCz (cationic)	2.3–3.7, 4.95	6.7, 7.25	8.35, 8.83
PVCz (radical)	2.3–3.6, 5.0	6.73, 7.27	8.3, 8.85
P7VBCz (cationic)	2.0-3.8, 5.15	6.8, 7.4	8.4, 9.05
P7VBCz (radical)	1.9-3.7, 5.2	6.8, 7.4	8.36, 9.0
P5VBCz (radical)	2.3-3.9, 5.1	6.75	8.4

 a The spectra were measured in tetradeuterioethylene bromide at 125 °C.

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going from toluene to ethylene chloride. As the temperature decreases, it decreases down to about 0.50 at -78 °C in the case of P7VBCz, but does not vary in the case of PVCz. Thus, it varies with the polymerization method in a manner similar to the case of both the T_g and the solubility; the sample having a higher T_g and poorer solubility has a larger $I_{H/L}$ value. However, this relation is not held for the samples prepared by Ziegler-Natta initiators; the $I_{H/L}$ values are relatively small, in spite of the high T_g and the poor solubility.

In the case of P5VBCz, there seems to be a similar difference in the $I_{H/L}$ value between the cationic and the free-radical samples, but no quantitative discussion can be done because of the highly unresolved spectra.

Both the doublet signals of methylene and methine protons in the 100-MHz spectra are independent of measuring temperature but dependent on the polymerization method. On the other hand, Williams has reported that the 220-MHz NMR spectrum of PVCz shows clearly the multiplicity of the signals of methylene and methine protons and that the methylene signal in the 220-MHz spectrum varies with temperature owing to temperature-dependent conformational effects. 46) We have also measured the 220-MHz spectra of PVCz in several solvents and at a series of temperatures. The resolution of the spectrum of methylene protons varies with both solvent and temperature. In some cases, the 220-MHz spectra are unresolved rather than the 100-MHz spectra. However, the highly resolved spectra of methylene protons are composed of four peaks in both cases of the cationic and freeradical samples; the two higher field peaks and the two lower field ones correspond to the higher field peak and the lower

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field one in the 100-MHz spectra, respectively. Moreover, the intensity ratios of the two higher field peaks to the two lower field ones (for example, 0.35 and 0.89 for samples A and C, respectively) are in good agreement with the $I_{H/L}$ values of the corresponding samples listed in Table I-1. Therefore, the differences in $I_{H/L}$ values are attributable to some difference in the stereochemical configuration of the polymer chain.

The splittings of both doublet signals of methylene and methine protons are too large to be attributable to some triad tacticity in a usual manner. Therefore, it may be necessary to assume the presence of two kinds of both methylene and methine protons having the magnetically different environments, that is, the stereoblock arrangements of isotactic 3/1 and syndiotactic 2/1 helixes, as have been proposed by Mikawa et al..⁴⁴⁾

Heller et al. investigated the NMR spectra of polyvinylaromatics such as polystyrene and poly(1-vinylnaphthalene)(PlVN) in connection with the stereoregularity, and found that the aliphatic protons of PlVN have the following unusual behaviors probably owing to the bulkiness of the pendant group.⁶⁰⁾ (1) Both the methylene and methine signals of the isotactic sample (9.02 and 7.52 τ , respectively) are remarkably shifted to the field higher than those of the atactic one (8.57 and 7.24 τ). (2) The methine signals of both the samples are shifted appreciably to the lower field as compared with those of polystyrene etc..

In the present case, the aliphatic protons have the behaviors similar to those of PlVN. Therefore, the higher field peaks of both methylene and methine protons may tentatively be assigned to the isotactic sequence, and the lower field peaks to the syndiotactic one. According to this assignment, the $I_{\rm H/L}$ value

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is a measure of the relative amount of isotactic sequence to syndiotactic one of the polymer. In both PVCz and P7VBCz, the samples prepared cationically in a nonpolar solvent at room temperature have the largest amount of isotactic sequence (50%) and the free-radical samples have the smallest amount (25%). The difference in the amount of isotactic sequence explains reasonably the high T_g and poor solubility of the former samples as compared with the case of the latter ones, because the isotactic 3/1 helix is considered to be less flexible rather than the syndiotactic 2/1 helix. This is another evidence for the present assignment.

Crystal⁴⁷⁾ and Griffiths⁶¹⁾ have reported that in the lamellar type or rod type crystals of PVCz prepared by appropriate heat treatments a polymer chain is composed of an isotactic 3/1 helix but not of a syndiotactic 2/1 helix, although the PVCz sample used by Griffiths is considered to be a free-radical sample judging from the T_g (227 °C). This might suggest that the isotactic 3/1 helix segment is crystallized much more easily than the syndiotactic 2/1 helix segment, and the latter segment is primarily present in the poor crystalline and amorphous phases. The sample with the crystallinity of 60% has been obtained from the PVCz sample used by Griffiths.⁶¹⁾ This might support assigning the lower field peaks of both methylene and methine protons to the syndiotactic sequence rather than the atactic one.

Summary

5VBCz and 7VBCz having more bulky pendant groups than VCz were synthesized and polymerized. By cationic polymerization

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at low temperatures, 5VBCz gave only the low molecular weight polymer (DP \leq 9) in a high yield. The high molecular weight polymer (DP = 50) was prepared only by bulk polymerization initiated with azobisisobutyronitrile. 7VBCz was easily polymerized free-radically and cationically, as is the case with VCz. From the NMR spectra, the glass transition temperatures, and the solubility of the polymers, it was clarified that the stereoregularity varies depending on the polymerization method. Both methylene and methine protons of these three polymers show doublet signals, of which the higher field peaks were assigned to the isotactic sequence and the lower field peaks to the syndiotactic one. According to this assignment, it was estimated that the cationic polymers have the largest amount of the isotactic sequence (50%), and that the free-radical polymers have the smallest amount (25%).

Chapter 2

Fluorescence Spectra of the Vinyl Polymers with Pendant Carbazolyl and Benzocarbazolyl Groups

Introduction

Poly(N-vinylcarbazole)(PVCz), both in solution and in film, shows one broad fluorescence, which is not composed of only one component, but of two components. The broad emission at the longer wavelength(λ_{max} ; ca. 23900 cm⁻¹) is assigned to the sandwich-like excimer fluorescence. On the other hand, the assignment of the broad emission in the shorter-wavelength region $(\lambda_{max}; ca. 27000 \text{ cm}^{-1})$ is controversial because of the absence of the structure and of the large Stokes shift. David et al. and Yokoyama et al. have assigned it to the monomer fluorescence for PVCz in solution.^{5,17)} P. C. Johnson and Offen,⁴⁾ and Powell and Venikouas¹⁴⁾ have assigned it to the emission from a dimer trap for PVCz films. On the other hand, G. E. Johnson has recently assigned it to the second excimer fluorescence for PVCz in solution, and reported that the polymer conformation appropriate for the formation of this state exists prior to the initial excitation step.¹⁸⁾ Thus, the structure of the species emitting at ca. 27000 cm^{-1} has been ambiguous so far.

It is of interest to investigate how the difference in the tacticity is reflected in the fluorescence spectra.

Concerning the vinyl polymers with small aromatic chromophores, the relation between the tacticity of the polymer chain and

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the fluorescence spectrum has been reported in several For polystyrene(PSt), 34,35,40,41 poly(α -methylinvestigations. styrene),³⁶⁾ and poly(p-methylstyrene),⁴¹⁾ it has been reported that the ratio of the sandwich-like excimer to the monomer fluorescence intensity for the isotactic polymer is larger than that for the atactic polymer, although an opposite experimental result for PSt has been reported. 34) Y.-C. Wang and H. Morawetz have found, from the NMR spectra of polyacenaphthylene, that the polymers prepared by cationic and radical polymerizations have different configurations, and that this is reflected in the ease with which they form excimers.³⁹⁾ However, such a relation has never hitherto been studied for the vinyl polymers with large and bulky aromatic chromophores such as carbazole and benzocarbazole.

Experimental

The polymers used are listed in Table I-5. The low-molecularweight compounds were the same as used before⁶² or were of commercial origin and they were purified by the appropriate methods.

The emission spectra were measured

Table I-5. Characterization of Polymers Used ^{a)}						
Sample	Polymeri- zation temp/°C	Degree of polymeri- zation	I _{H/L}			
VCz Olig. I ^{b)} VCz Olig. II ^{b)} PVCz(r) PVCz(c) PCzEVE ^{c)}	20 20 70 20 -25	4 10 1040 520 33	0.35 0.97			
P7VBCz(r) P7VBCz(c)-I P7VBCz(c)-II P7VBCz(c)-III P5VBCz(r)-I	75 -78 -40 10	58 103 82 36 49	0.32 0.51 0.68 0.92			
P5VBCz(r)-II P5VBCz(c)	80 -78	13 8				

 a) Polymers prepared by the radical and cationic polymerizations are abbreviated as (r) and (c), respectively.

- b) VCz oligomers were prepared by the methods described in the literature.⁶²
- c) PCzEVE was the same as used before.⁶²⁾

with a spectrophotofluorometer constructed using a stabilized 500 W xenon lamp, monochromators, and a PM-55 (S-20) photomultiplier. The spectra were calibrated for the detector response by using a standard tungsten lamp. The fluorescence-decay times were measured with a pulsed N_2 gas laser as the exciting light source. Responses from the photomultiplier were led to a sampling oscilloscope (Tektronix 661) and recorded on a X-Y recorder, or led to a Tektronix 475 oscilloscope and the decay curve was photographed.

Results and Discussion

(1) Fluorescence Spectra of PVCz.

The fluorescence spectra of the various PVCz samples in aerated THF solution are shown in Fig. 1-4. The ratio of the fluorescence intensity in the shorter-wavelength region to that in the longer-wavelength region (sandwich-like excimer fluorescence, λ_{max} ; 23900 cm⁻¹) in PVCz(r) samples is remarkably larger than that in PVCz(c) samples. Figure I-5 shows the fluorescence spectra of PVCz(r) and PVCz(c), which have almost the same concentrations in degassed THF-MTHF solution, and their resolution spectra. The resolution was carried out on the assumption that the location of the sandwich-like excimer fluorescence band in solution was the same as that in film(see p.64). The fluorescence at the shorter wavelength in the resolution spectra (λ_{max} ; 26900 cm⁻¹) is not assigned to the monomer one because of the absence of the structure and of the large Stokes shift (2200 cm^{-1}). The fluorescence is not assigned also to the stable ground state dimer because of the absence of an

absorption due to such a dimer. The broad fluorescence seems to be assigned to a second excimer on the basis of a conformational consideration in section 4. It may safely be considered that the radiative rate constants of each fluorescence (second

and sandwich-like excimer bands) of PVCz(r) are equal to those of PVCz(c). Therefore, Fig. I-5 shows that the concentration of the sandwich-like excimer in PVCz(r) is nearly equal to that in PVCz(c), and that the concentration of the second excimer in PVCz(r) is higher than that in PVCz(c) by a factor of about 2.4.

In rigid glass at 77 K, a



Fig. 1-5.

Fluorescence spectra and the resolution spectra of PVCz(r) and PVCz(c) in MTHF-THF degassed solutions at room temperature. Optical densities at excitation wavelength (310 nm) are 0.132 and 0.131 for PVCz(r) and PVCz(c), respectively.

(1) ---; PVCz(r), (2) ---; PVCz(c), (3) \cdots ; the component of the sandwich-like excimer fluorescence of PVCz(r) and PVCz(c), (4) ----; the component of the second excimer fluorescence of PVCz(r), (5); the component of the second excimer fluorescence of PVCz(c).



Wave number/ 10^3 cm⁻¹ Fig. 1-4.

Fluorescence spectra of PVCz and VCzvinylnaphthalene(VN) copolymers polymerized by various methods in THF aerated solutions at room temperature. Excitation wavelength; 330 nm. (1) VCz-VN(0.5 mol% in feed) copolymer polymerized by AIBN (1 mol%) in benzene (1 M) at 70 °C for 2 h, (2) PVCz(r) polymerized by AIBN (1 mol%) in benzene (1.3 M) at 70 °C for 4.5 h, (3) PVCz(r) polymerized by AIBN (1 mol%) in benzene (0.5 M) at 70 °C for 7 h, (4) PVCz(c) polymerized by BF_3OEt_2 (1 mol%) in toluene (0.26 M) at 20 °C for 1 h, (5) VCz-VN (1 mol% in feed) copolymer polymerized by BF_3OEt_2 (1 mol%) in toluene (0.86 M) at 15 °C for 3 h, (6) PVCz(c) polymerized by BF₃OEt₂ (1 mol%) in toluene (0.09 M) at $-78 \degree \text{C}$ for 1 h, (7) PVCz(c) polymerized by BF_3OEt_2 (0.5 mol%) at 20 °C for 18,5 h.

significant difference in the fluorescence spectra between PVCz(r) and PVCz(c) was found, as is shown in Fig. I-6. The PVCz(c) shows mainly a monomer structured fluorescence, while the PVCz(r) shows mainly the second excimer fluorescence, although weak monomer fluorescence is observed. The fluorescence spectra both in fluid and in rigid solutions are considered to reflect the difference in the tacticity of PVCz. That is, the concentration of the second excimer in syndiotactic-rich polymer(PVCz(r)) is higher than that in isotactic-rich polymer(PVCz(c)).

(2) Fluorescence Spectra of P7VBCz.

The fluorescence spectra of 7EBCz and P7VBCz are shown in Fig. I-7. The 7EBCz shows no excimer fluorescence even in concentrated solutions up to 0.1 M, as is the case of ECz. The polymers show one broad fluorescence composed of two components $(\lambda_{max}; 21950 \text{ and } 24900 \text{ cm}^{-1})$, as is the case of PVCz. The



Emission spectra of (1) ----; VCz Olig, II, (2)----; PVCz(r), (3)-----; PVCz(c), and (4)-----; PCzEVE in MTHF-THF rigid glasses at 77 K.



Fig. I-7. Wave number/10³ cm⁻¹

Fluorescence spectra and their resolution spectra of 7EBCz, P7VBCz(r), and P7VBCz(c)-I in MTHF-THF degassed solutions at room temperature. Optical densities of both the polymers at excitation wavelength (350 nm) are 0.249. Fluorescence spectra: (1)——: 7EBCz, (2)——, P7VBCz(r), and (3)——: P7VBCz-(c)-I. Resolution spectra: (4)……: the component of the sandwich-like excimer fluorescence of P7VBCz(r) and P7VBCz(c)-I. (5)----, and (6)——: the component of the second excimer fluorescence of P7VBCz-(r) and P7VBCz(c)-I, respectively. energy differences between the peaks of these two bands and the 0-0 fluorescence band(26560 cm⁻¹) of 7EBCz are 4610 and 1660 cm⁻¹. These are almost the same as the corresponding energy differences between PVCz and ECz (4630 and 1630 cm⁻¹). The fluorescence excitation spectra of the polymers, obtained by monitoring at several wavelengths (390-500 nm), duplicate the absorption spectra of these solutions exactly. By analogy with the fluorescence spectrum of PVCz, the fluorescence bands can be assigned to the sandwich-like excimer fluorescence and the second excimer one. Figure I-7 shows that the difference in the fluorescence spectra between the radical and cationic polymers is attributable to the difference in the concentration of the second excimer.

The $I_{H/L}$ and \overline{DP} of the cationic polymers vary remarkably with the polymerization temperature (Table I-5). Thus, the fluorescence spectra of P7VBCz samples in aerated solution (5.9 x 10⁻⁵ M) were compared with one another in Fig. I-8. The second excimer fluorescence intensities are in the order of

(c)-III. This order is not correlated to the molecular weight of the polymers, but to the decrease in the I_{H/L} value. That is, the concentration of the second excimer increases with an increase in the syndiotactic sequence.

(r) > (c) - I > (c) - II >



Fluorescence spectra of (1) P7VBCz(r), (2) P7V-BCz(c)-I, (3) P7VBCz(c)-II, and (4) P7VBCz(c)-III in MTHF-THF aerated solutions at room temperature. Excitation wavelength; 350 nm.

The sandwich-like excimer fluorescence of each P7VBCz sample has a constant intensity at temperatures from ca. -110 to 20 °C, as is shown in Fig. I-9. At temperatures above ca. 20 °C, the intensity of the sandwich-like excimer fluorescence decreases and the intensity of the second excimer fluorescence increases with an increase in the temperature. This phenomenon is observed also for PVCz.¹⁸⁾ This behavior and the presence of the longlived component decay at the shorter wavelength (vide infra) indicate the dissociation of sandwich-like excimer to give back the second excimer.

The fluorescence-decay curves of the polymers at room temperature are multi-component in the shorter-wavelength region (36±2 ns and initial fast decay), while they are one-component (36±2 ns) in the longer-wavelength region. The long-lived component corresponds to the sandwich-like excimer fluorescence.

The spectral band of the polymers in rigid glass solutions at 77 K is shifted to a lower frequency and broadened as compared





Fluorescence spectra of (1) 7EBCz, (2) P7VBCz-(r), and (3) P7VBCz(c)-I in MTHF-THF rigid glasses at 77 K. Excitation wavelength; 350 nm.
with that of 7EBCz, as is shown in Fig. I-10. This deviation from the monomer-like fluorescence band shape seems to be attributable to a large contribution of the second excimer fluorescence. In the case of the PVCz samples, the cationic polymer shows a monomer-like structured fluorescence, while the radical polymers show mainly the second excimer fluorescence. The fluorescence decay times of both the PVCz samples (λ_{obsd} = 355-390 nm) are approximately the same as that measured for i-PrCz (17+2 ns), although a slight departure from a strict first-order decay is observed (Table I-6). This fact indicates that the lifetime of the second excimer fluorescence seems to be nearly equal to that of the monomer-like fluorescence. In the case of the P7VBCz samples, the fluorescence decay times of both the polymers at 380 nm are ca. 18 ns, while those at the longer wavelength(λ > 400 nm) are 25+2 ns. The decay curves are roughly single-exponential. This lifetime(25 ns) is not in agreement with those of the monomeric fluorescence of the model compound (13 ns) and of the sandwich-like excimer fluorescence(36 ns), and is attributed to the second excimer. The decay time(18 ns) at 380 nm seems to be an apparent value due to an overlap of the monomerlike fluorescence and the second excimer one.

(3) Fluorescence Spectra

of P5VBCz.

The fluorescence spectra of 5EBCz and P5VBCz are shown in Fig. I-11. The 5EBCz shows no excimer fluorescence even in concentrated solutions up

Table I-6. Fluorescence LIFETIME^{a)}

Sample	Monomer ^{b)}	Monomer- like ^{c)}	Second excimer ^{c)}	Sandwich- like excimer ^{d)}
<i>i</i> -PrCz	17 ± 2			
PVCz(c)		19 ± 2		45 ± 1
PVCz(r)		17 ± 2	17 ± 2	45 ± 1
7EBCz	13 ± 1	⁻		
P7VBCz(c)		(13) ^{e)}	25 ± 3	36 ± 2
P7VBCz(r)		(13) ^{e)}	25 ± 1	36 ± 2
5EBCz	24 ± 1			
P5VBCz(c)		23 ± 2		57 ± 3
P5VBCz(r)	·	26	65—75	57±3

a) Unit; ns. Solvent: MTHF-THF degassed solution.
b) At room temperature and 77 K. c) At 77 K. d) At room temperature. e) See text. to 0.1 M, as is the case with ECz and 7EBCz. The radical polymers show one broad fluorescence, the spectra of which were successfully resolved into two broad bands, with peaks of 20350 and 22550 cm⁻¹, in a manner similar to that in P7VBCz. The energy differences between the peaks of these two bands and the 0-0 fluorescence band(24330 cm⁻¹) of 5EBCz are 3980 and 1780 cm⁻¹ respectively. Therefore, the fluorescence bands can be assigned to the sandwich-like excimer fluorescence and the second excimer one .

The temperature dependence of the fluorescence spectrum of the P5VBCz(r)-II sample is shown in Fig. I-12. As may be seen, there is no evidence for the formation of an isoemissive point. The sandwich-like excimer fluorescence has a constant intensity over the temperature range from ca. -70 to 40 °C, as is shown in Fig. I-9. These phenomena observed for the radical polymers are



Normalized fluorescence spectra and their resolution spectra of 5EBCz, P5VBCz(r)-I, P5VBCz(r)-II, and P5VBCz(c) in MTHF-THF degassed solutions at room temperature. Excitation wavelength; 350 nm. Fluorescence spectra: (1)—: 5EBCz, (2)----; P5VBCz(r)-I, (3)—; P5VBCz(r)-II, and (4)—; P5VBCz(c). Resolution spectra: (5)……, and (6) ——: the component of the sandwich-like excimer fluorescence of P5VBCz(r)-II and P5VBCz(c), respectively. (7)……: the component of the second excimer fluorescence of P5VBCz(r)-II, and (8)—---; the component of the monomeric fluorescence of P5VBCz-(c).





Temperature dependence of the fluorescence spectra of P5VBCz(r)-II in MTHF-THF degassed solution. Excitation wavelength; 350 nm. (1) 143.5 K, (2) 163 K, (3) 203.5 K, (4) 222 K, (5) 244 K, (6) 263 K, (7) 281 K, (8) 294.5 K, and (9) 308 K.

quite similar to those observed for the PVCz and P7VBCz.

On the other hand, the fluorescence spectrum of the cationic polymer differs markedly from those of the above-mentioned polymers: (1) The monomer-like structured fluorescence (0-0; 23800 cm^{-1}) in the shorter-wavelength region is observed clearly even at room temperature, while the second excimer fluorescence is not observed, as is shown in Fig. I-11. (2) An isoemissive point is clearly observed, as is shown in Fig. I-13. (3) The intensity of the sandwich-like excimer fluorescence in the longerwavelength region increases with an increase in the temperature up to ca. 20 °C, as is also shown in Fig. I-9.

A dilute fluid solution of a VCz Olig. II at room temperature does not show any structured fluorescence at the shorter wavelength, but a broad fluorescence which is composed of two excimer fluorescences. Therefore, these three significant differences in the fluorescence spectra between the radical and cationic polymers are not attributable to any difference in the

molecular weight, but to the difference in their tacticities. This explanation seems to be supported also by the polymerization results: the cationic polymer is considered to be composed of only an isotactic sequence. The radical polymers are composed of stereoblock arrangements of the isotactic and syndiotactic helixes. Therefore, the



Fig. 1-13. Wave number/ 10^3 cm⁻¹

Temperature dependence of the fluorescence spectra of P5VBCz(c) in MTHF-THF degassed solution. Excitation wavelength; 350 nm. (1) 183.5 K, (2) 202.5 K, (3) 221 K, (4) 262.5 K, (5) 282 K, and (6) 304 K.

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above-mentioned discussion supports the idea that the second excimer site is not formed within the isotactic sequence, but only within the syndiotactic sequence.

The fluorescence-decay curves of both the radical and cationic polymers at room temperature are multi-component in the shorter-wavelength region $(57\pm3 \text{ ns} \text{ and an initial fast decay})$, while there is only one component with a long lifetime $(57\pm3 \text{ ns})$ in the longer-wavelength region. The long-lived component corresponds to the sandwich-like excimer fluorescence.

Figure I-14 shows the fluorescence spectra of P5VBCz and 5EBCz in rigid glass solutions at 77 K. The fluorescence spectrum of the cationic polymer is mainly composed of a monomer-like fluorescence, the lifetime of which at 420 nm was 23±2 ns. This is the same value as the decay time of 5EBCz. On the other hand, the fluorescence spectra of the radical polymers are markedly broadened as compared with that of the cationic polymer. This fluorescence is mainly emitted from the second excimer site,

although the fluorescences from the excited monomer state and the sandwich-like excimer site are slightly involved.

(4) <u>Structure of the Second</u> <u>Excimer</u>.

In the conformations for a three-skeletal carbon rotational dyad of syndiotactic and isotactic polymer





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chain sequences based on a threefold rotational potential,³⁷⁾ the conformational states with the lowest energy and those corresponding to the sandwich-like excimer site are shown in Fig. I-15 (a) and (b). In the syndiotactic case there are a doubly degenerate ground state (g^+g^+, tt) and a doubly degenerate sandwich-like excimer state (g^-t, tg^-) . On the other hand, in the isotactic case there are a doubly degenerate ground state (g^+t, tg^-) and a single sandwich-like excimer state(tt). The formation of the sandwich-like excimer conformation from the ground-state conformation requires rotational motion about the carbon-carbon bond. In the ground-state conformations, the tt conformation of the syndiotactic chain (Fig. I-15 (a)) has the

shortest distance between neighboring carbazolyl chromophores. Since the carbazolyl chromophore is bulky, it seems to be easy to approach and mutually overlap the one benzene ring of the neighboring carbazolyl chromophores in the excited state, as is shown in Fig. I-16. The ground-state molecules corresponding to this reduced separation have a repulsive interaction.

It has been reported that 1,2-di(1-anthry1)ethane









...ws a broad excimer fluorescence in solution at room temperature at wavelengths shorter than the fluorescence of a sandwichlike excimer formed by the decomposition of its photodimer in rigid glass at 77 K, and that the structure of the excimer seems to be the overlap of only one benzene ring in the mutual anthracene chromophores.⁶³⁾ Therefore, it seems reasonable to consider that the structure of the second excimer state may be the overlap of only one benzene ring in the neighboring chromophores. This structure seems to be supported by the fact that the formation of the sandwich-like excimer from the second excimer requires a rotational motion about the carbon-carbon bond.¹⁸⁾

(5) Formation of an Exciplex.

The fluorescence spectra of PVCz(r) and PCzEVE in solution in the absence and in the presence of DMTP (6 x 10^{-2} M) are

shown in Fig. I-17. In the cases of PVCz(r and c) and VCz Olig. II, the fluorescence is nonuniformly quenched by DMTP. That is, the sandwichlike excimer fluorescence is quenched more effectively than the second excimer fluorescence. On the other hand, the fluorescence of PCzEVE is uniformly quenched by DMTP. In all cases, the exciplex fluorescence





Fluorescence spectra of PVCz(r) $(3.9 \times 10^{-4} \text{ M})$ and PCzEVE $(1.9 \times 10^{-4} \text{ M})$ in MTHF-THF degassed solutions in the absence and the presence of DMTP $(1.9 \times 10^{-2} \text{ M})$ at room temperature. The intensities of the spectra are not comparative for the different polymers.

(1) ——; PVCz(r) in the absence of DMTP, (2) ----; PVCz(r) in the presence of DMTP, (3) ——; PCzEVE in the absence of DMTP, and (4) ……; PCzEVE in the presence of DMTP. appears in the longer-wavelength region. The quenching factor, Q, is defined by Q = $(I^{0}-I)/I$, where I^{0} and I are the fluorescence intensity in the absence and in the presence of the quencher molecule(DMTP) in a concentration of [Q] (mol/l), respectively. As is shown in Fig. I-18, the second excimer band is linearly quenched and the sandwich-like excimer band is superlinearly quenched. This phenomenon is the same as the case of fluorescence quenching by acid.¹³

From the results mentioned in the previous sections, the following reaction scheme is the most conceivable.



, where k_{f} , k_{r} , and k_{q} are the radiative, nonradiative, and quenching rate constants. respectively. The letters with

nonprime and prime show the second and sandwich-like excimers, respectively. The k_E and k_s are the rate constants of the conversion between the second excimer and the sandwich-like one. The τ_s and τ_E are the lifetimes of the second and sandwich-like excimer fluorescences in the absence of DMTP. Under steady state conditions, the intensities



Fig. I-18. DMTP concentration/M⁻¹

Quenching of PVCz(r) by DMTP in THF aerated solutions at room temperature. Excitation wavelength; 345 nm.

(1) The sandwich-like excimer band, and (2) the second excimer band.

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of the second and the sandwich-like fluorescence bands (I_s and $I_{\rm F}$, respectively) are given by the following equations;

$$I_{s} = \frac{k_{f}\tau_{s}(1 + k_{q}\tau_{E}[Q])}{(1 + k_{q}\tau_{s}[Q])(1 + k_{q}\tau_{E}[Q]) - k_{s}k_{E}\tau_{s}\tau_{E}} \cdot I_{a}$$
(1)

$$I_{E} = \frac{k_{f} k_{E} \tau_{s} \tau_{E}}{(1 + k_{q} \tau_{s}[Q])(1 + k_{q} \tau_{E}[Q]) - k_{s} k_{E} \tau_{s} \tau_{E}} I_{a}$$
(2)

From these equations, the following equations are obtained.

 \sim

$$Q_{s} = \frac{I_{s}^{0} - I_{s}}{I_{s}} = \frac{\tau_{s} \{k_{q}(1 + k_{q}^{\dagger}\tau_{E}[Q]) + k_{q}^{\dagger}k_{s}k_{E}\tau_{E}^{2}\}}{(1 - k_{s}k_{E}\tau_{s}\tau_{E})(1 + k_{q}^{\dagger}\tau_{E}[Q])} \cdot [Q]$$
(3)

$$\frac{Q_E}{[Q]} = \frac{I_E^0 - I_E}{I_E[Q]} = \tau_s (k_q + k_q^T \tau_E^T \tau_s^{-1} + k_q k_q^T \tau_E^{-1}]$$
(4)

$$\frac{\mathbf{I}_{s}}{\mathbf{I}_{E}} = \frac{\mathbf{k}_{f}}{\mathbf{k}_{E}\boldsymbol{\tau}_{E}} + \frac{\mathbf{k}_{q}\mathbf{k}_{f}}{\mathbf{k}_{f}\mathbf{k}_{E}} [Q]$$
(5)

The quantum yields of the second excimer(η_s) and the sandwichlike excimer(η_E) are given by equation (6);

$$\frac{\eta_{s}}{\eta_{E}} = \frac{k_{f}\tau_{s}}{k_{f}^{\dagger}\tau_{E}}$$
(6)

The plots of equations (4) and (5) are given in Figs. I-19 and I-20, respectively. The following values were measured for PVCz(r), 64 , $\tau_s \sim 4 \times 10^{-9}$ s, $\tau_E \sim 17 \times 10^{-9}$ s, and $\eta_s / \eta_E = 0.88$. Then, by using the equations (4), (5), and (6), the rate constants listed in Table I-7 were evaluated. Judging from the linear relation between Q_s and [Q] in Fig. I-18, the following relation should be held: $k'_q \tau_E[Q] << 1$. The relation was approximately satisfied under this experimental condition.

The quenching reaction is more efficient for the second



Fig. I-19. The ratio of the quenching factor of the sandwich-like excimer to the concentration of DMTP vs. the concentration of DMTP.



Fig. I-20. The fluorescence intensity ratio of the second excimer to the sandwich-like one vs. the concentration of DMTP.

excimer than for the sandwich-like one by a factor of ca. 4. The value of k_E is larger than that of k_S by a foctor of ca. 5, indicating that the sandwich-like excimer is formed easily from the second excimer. The value of k_E is the same order as that of the formation of the sandwich-like excimer from the monomer for DCzP in THF solution (1 x 10⁸ s⁻¹).⁶⁵⁾

Table I-7. Rate Constants for PVCz(r) in Aerated THF Solution.

$k_{q}/M^{-1}s^{-1}$	k ['] _q /M ⁻¹ s ⁻¹	k _E /s ⁻¹	k _s /s ⁻¹	$(k_{f}+k_{r})/s^{-1}$	$(k_{f}'+k_{r}')/s^{-1}$
4×10^9	1×10^9	2×10^8	4×10^{7}	5×10^{7}	2×10^{7}

Summary

The emission properties of PVCz, P7VBCz, and P5VBCz were investigated in solution in connection with the tacticity of the samples. PVCz(r and c), P7VBCz(r and c), and P5VBCz(r), which are composed of both the syndiotactic sequence and the isotactic one, showed one broad emission band composed of the second and the sandwich-like excimer fluorescences. The intensity of the second excimer fluorescence increased with an increase in the content of the syndiotactic sequence. P5VBCz(c), which is composed of only the isotactic sequence, showed structured emission composed of the monomer and the sandwich-like excimer These results indicate that the second excimer fluorescences. fluorescence is emitted from only the syndiotactic sequence. The conformational consideration suggests the structure of the second excimer that only one benzene ring in each neighboring carbazolyl chromophores overlaps with the other one (Fig. I-16).

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The parameters of the photophysical process for PVCz(r) were evaluated by the quenching experiment. The sandwich-like excimer fluorescence was quenched more effectively than the second excimer fluorescence. Fluorescence Spectra of Poly[2-(N-carbazolyl)ethyl vinyl ether]

Introduction

It is known that 1-ethylpyrene and 1-ethylnaphthalene show the excimer fluorescence in concentrated solutions. Poly(1-vinylpyrene)^{66,67)} and poly(1-vinylnaphthalene)^{34,68)} show, almost entirely, only the excimer fluorescence in dilute solutions. Poly(1-pyrenylmethyl vinyl ether)(PPyMVE),^{66,69)} poly(1-naphthylmethyl vinyl ether)(PNMVE),⁶⁹⁾ and poly(1-naphthyl methacrylate) (PNMA),⁷⁰⁾ in which the naphthyl and pyrenyl groups are widely spaced on the skeletal chains by -O-CH₂- and -CO-O- bonds, show both the structured monomeric fluorescence and the excimer one in dilute solutions. On the other hand, ECz shows no excimer fluorescence in concentrated solution.¹⁷⁾ However, PVCz shows excimer fluorescence in dilute solution. Therefore, spectroscopic studies on a vinyl polymer with carbazolyl chromophores widely spaced on the skeletal chains are of interest in comparing with those of the spectroscopic properties of PVCz.

In this chapter, the author investigated the spectroscopic behavior of PCzEVE with carbazolyl chromophores widely spaced on the main chain by $-0-CH_2-CH_2$ bonds. It has already been found, from the absorption and NMR spectra,⁷¹⁾ that PCzEVE has no appreciable ground-state interaction among the carbazolyl chromophores.

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The materials and the apparatus for measurments were the same as those in the previous chapter.

For the fluorescence polarization, two polarizers were used, and the data were corrected by the method described by Azumi and McGlynn.⁷²⁾

Results and Discussion

The fluorescence spectra of the compounds with the carbazolyl chromophores in MTHF-THF dilute solutions $(2.0 \times 10^{-4} \text{ M})$ are shown in Fig. I-21. The emission spectrum of DCzP consists of a monomeric structured fluorescence and a sandwich-like excimer one with a peak at ca. 23900 cm⁻¹, as is well-known.^{65,73-75)} VCz Olig. I and II show both a structureless fluorescence with a peak at 27000-27500 cm⁻¹ and a sandwich-like excimer fluorescence, as is the case of PVCz, although VCz

Olig. I also shows a monomer fluorescence as a shoulder. On the other hand, PCzEVE shows only a broad, structureless fluorescence with a peak at 27100 cm⁻¹. The Stokes shift (1990 cm⁻¹) is much larger than those of i-PrCz and DCzP (380 cm⁻¹). The fluorescence excitation spectra of PCzEVE, obtained





Normalized fluorescence spectra of the model compound and vinylpolymers in MTHF-THF degassed solution at room temperature. Excitation wavelength; 310 nm. (1) -----; DCzP, (2) ----; PCzEVE, (3) ----; VCz

(1) ----; DCzP, (2) ----; PCzEVE, (3) ----; VCz Olig. II, (5) ----; PVCz(r).

by monitoring at several wavelengths(350-450 nm), duplicated exactly the absorption spectrum of this solution.

The fluorescence decay curve of PCZEVE in MTHF-THF solution clearly consists of two-component in all wavelength regions (370-430 nm), as is shown in Fig. I-22. The decay constant of the long-lived component observed at 370 nm $(33\pm1 \text{ ns})$ is longer than that of i-PrCz $(17\pm2 \text{ ns})$ and shorter than that of the sandwich-like excimer fluorescence of PVCz $(45\pm1 \text{ ns})$. The initial fast decay constant is $13\pm2 \text{ ns}$. This two-component decay of PCZEVE is observed also in rigid glass at 77 K $(35\pm4 \text{ and } 12\pm2 \text{ ns})$. The intensity ratio of the short-decay component to the long one in rigid glass at 77 K is larger than that in fluid solution at room temperature.

Table I-7 shows the relative fluorescence quantum yields, taking the value of i-PrCz, which forms no excimer, as unity. The fluorescence quantum yield of PCzEVE is higher than that of DCzP, in which the sandwich-like excimer formation is minimum

among the materials used, as is shown in Fig. I-21.

The fluorescence spectra of PCzEVE at various temperatures are shown in Fig. I-23. At lower temperatures, two peaks appear, and they approach

Table I-7. RELATIVE FLUORESCENCE YIELDS

Substance	η
<i>i</i> -PrCz	1.00
PCzEVE	0.79
DCzP	0.57
VCz Olig. I	0.40
VCz Olig. II	0.37
PVCz(r)	0.33
PVCz(c)	0.27





Fluorescence decay curves of the model compound and vinylpolymers in MTHF-THF degassed solutions at room temperature.

- (1)]-; *i*-PrCz (λ_{obsd} = 370 nm); τ_f = 17 ns.
- (2) • •; PVCz(r) (λ_{obsd} =380 nm), -•-•; the long-lived component; τ_f =46 ns.
- (3) $\triangle \triangle \triangle$; PCzEVE ($\lambda_{obsd} = 370 \text{ nm}$), $-\triangle --\triangle -$; the long-lived component; $\tau_f = 31 \text{ ns}$, $--\triangle --\triangle --$; the short-lived component; $\tau_f = 11 \text{ ns}$; the difference between the observed decay curve and the long-lived component.

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each other with an increase in the temperature. At temperatures higher than ca. 273 K, only a structureless fluorescence spectrum is observed. The fluorescence intensity decreases with an increase in the temperature. From the measurement of the temperature dependence of the fluorescence spectra of PVCz, VCz Olig. I and II, and DCzP, it was found that the sandwich-like excimer fluorescence shows the maximum intensity at ca. 293 K. On the other hand, in the case of PCzEVE, no appreciable change in the fluorescence spectra near 23900 cm⁻¹ is observed over a wide temperature range, as is shown in Fig. I-23. These facts indicate that PCzEVE does not form the sandwich-like excimer.

The emission spectrum of PCzEVE in rigid glass at 77 K also is shown in Fig. I-23. The fluorescence spectrum of VCz Olig. II has, roughly, a so-called "mirror-symmetry" relation between the absorption and the fluorescence spectra (Fig. I-6 in chapter 2). However, PVCz and PCzEVE do not have the mirror-symmetry relation. In particular, the fluorescence spectrum of PCZEVE is Wavelength/nm observed at a longer wavelength 350 400 than that of PVCz and the shape is different from that of PVCz. Relative intensity The absorption spectra of PVCz 543 and PCzEVE in rigid glass at 77 K are sharper than those in fluid solution at room temperature, but no appreciable new 27 25 23 29 Fig. I-23. Wave number/10³ cm⁻¹ band is observed. The fluores-Temperature dependence of the fluorescence spectra of PCzEVE in MTHF-THF degassed solution. cence excitation spectra of Concentration of PCzEVE; 2.0×10^{-4} M. Excitation wavelength; 310 nm. these polymers in rigid glass (1) 168 K, (2) 208 K, (3) 249 K, (4) 298 K, and (5) 318 K. The fluorescence (6) at 77 K was also given

at 77 K are also in agreement

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for comparison.

with their absorption spectra. The resolution of the phosphorescence spectrum of PCzEVE is lower than those of i-PrCz and VCz Olig. I, and the spectrum is very similar to that of PVCz, although its lifetime $(6.6\pm0.1 \text{ s})$ is shorter than that of PVCz $(8.1\pm0.2 \text{ s})$. Therefore, the excited-state interactions are present among the carbazolyl chromophores in PCzEVE even at low temperatures, at which the solvent becomes rigid (ca. 102 K).

The degrees of polarization of the fluorescence are shown in Fig. I-24. The degrees are almost the same for PVCz and for PCZEVE. This fact indicates that the singlet energy migration in PCZECE occurs efficiently along the polymer chain, as is the case in PVCz.¹⁷⁾

A concentrated THF solution of PCzEVE (0.5 M) shows the same spectrum as in dilute solutions near 223 K and above.

As has been mentioned above, PPyMVE, PNMVE, and PNMA show both the normal structured monomeric fluorescence and the sandwich-like excimer one in dilute solutions. On the other hand, poly[2-(9-phenanthryl)ethyl vinyl ether](PPhEVE), with phenanthryl

chromophores widely spaced on the main chain by -O-CH₂-CH₂bonds, shows only the monomeric structured fluorescence in dilute solution at room temperature, as will be described in the following chapter. The fluorescence structure and lifetime in a dilute solutions are almost the same as those of 9-ethyl-



Fig. 1-24. Wavelength/nm

Fluorescence polarizations of MTHF-THF rigid glasses of (1) *i*-PrCz $(6.7 \times 10^{-5} \text{ M})$, (2) DCzP (7.3 $\times 10^{-5} \text{ M})$, (3) PCzEVE $(9.6 \times 10^{-5} \text{ M})$, and (4) PVC(r) $(1.0 \times 10^{-4} \text{ M})$ at 77 K. Excitation wavelength; 334 nm.

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phenanthrene. It is well-known that the excimer fluorescence of phenanthrene is not observed at room temperature.⁷⁶⁾ It is surprising that the normal structured monomeric fluorescence is not observed at all in the case of PCZEVE, in which aromatic chromophores are more widely spaced on the skeletal chain than PPyMVE, PNMVE, and PNMA. The fact that the sandwich-like excimer fluorescence of PCZEVE is not observed in either dilute or concentrated solution at room temperature also supports the idea that the sandwich-like excimer formation in PVCz is possible only between adjacent chromophores.¹⁷)

The above-described results indicate that the excited-state interaction among the carbazolyl chromophores in PCZEVE is quite different from that in DCzP, VCz Olig. I and II, and PVCz , and that only PCZEVE is unique among the vinyl polymers with aromatic chromophores widely spaced on the main chain. This behavior of PCZEVE may reflect a characteristic of the bulky carbazolyl chromophore.

The position of the PCZEVE fluorescence band in solution at room temperature is almost the same as that of the fluorescence in the shorter wavelength regions of PVCz and VCz Olig. I and II, although the appearance of the structure below ca. 273 K is different. Therefore, one of the emitting species of PCZEVE in solution might be the second excimer. At the present time, there is no idea about the other emitting species of PCZEVE.

Summary

PCzEVE showed an interesting, new, structureless emission band with a peak at 27100 cm^{-1} . This is different from both

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the monomeric structured fluorescence and the sandwich-like excimer one. The presence of two kinds of emitting species in this fluorescence was clarified by measuring the fluorescence decay curve. One of these emitting species is considered to be a second excimer. The emission properties of PCzEVE are unique among the vinyl polymers with aromatic chromophores widely spaced on the skeletal chain.

Chapter 4

Fluorescence Spectra of the Vinyl Polymers with Pendant Phenanthryl Groups

Introduction

The monomeric model compounds of the vinyl polymers investigated in the previous chapters show no excimer fluorescence. It is of interest to investigate the excimer formation of another aromatic vinyl polymer, whose the model compound shows no excimer fluorescence.

Phenanthrene crystal has only two molecules per unit cell arranged with their molecular planes almost perpendicular. It does not normally exhibit an excimer emission. However, this crystal under a high pressure exhibits a broad emission with a peak at ca. 440 nm. This emission is assigned to the excimer fluorescence.⁷⁷⁾ Despite extensive studies of the prompt and delayed fluorescences of phenanthrene, no excimer fluorescence has been observed in this compound at room temperature. 75) Studies on the emission of a phenanthrene sandwich dimer prepared by photolytic dissociation have also shown that phenanthrene does not form an excimer even under favorable condition. 78) Therefore, it is of interest to investigate the emission spectra of the vinyl polymers, where the phenanthryl(Ph) chromophores are directly connected to or widely spaced on the skeletal chains.

In this chapter, the author describes the spectroscopic behaviors of poly(9-vinylphenanthrene)(PVPh), poly(9-phenanthryl

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methacrylate) (PPhMA), and poly[2-(9-phenanthryl)ethyl vinyl ether](PPhEVE); in the latter two polymers, the Ph chromophores are widely spaced on the skeletal chains by -CO-O- and -O-CH₂-CH₂- bonds, respectively.

Experimental

9-Ethylphenanthrene (EPh) and 9-vinylphenanthrene were prepared according to the literature.^{79,80)} The PVPh was prepared by a radical polymerization (AIBN). The degree of polymerization ($\overline{\text{DP}}$) was about 18. This sample has sufficiently high molecular weight to examine the possibility of excimer formation, judging from the cases of DC2P and VCz Olig. I. 2-(9-Phenanthryl)ethyl vinyl ether was synthesized from 2-(9-Phenanthryl)ethanol by the route similar to 2-(N-carbazolyl)ethyl vinyl ether.^{51,81)} 9-Phenanthryl methacrylate was synthesized by the route similar to naphthyl methacrylate.^{82,83)} These monomers were identified by elemental analysis, NMR and IR spectra. The PPhEVE and the PPhMA used for spectral measurements were prepared by the cationic (BF_3OEt_2) and thermal polymerizations, respectively. The $\overline{\text{DP}}$ value of PPhEVE was about 87.

Results and Discussion

Fluorescence spectra of PVPh, PPhMA, PPhEVE, and EPh in solution are shown in Fig. I-25. The fluorescence spectrum of a concentrated EPh solution indicated that EPh forms no excimer, as is the case of phenanthrene. The fluorescence spectrum of PPhEVE is similar to that of EPh. Although the fluorescence

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spectra of PVPh and PPhMA are broad as compared with the spectrum of EPh, no clear excimer fluorescence in the longer-wavelength region is observed for PVPh and PPhMA. Therefore, it is suggested that the interaction between neighboring chromophores in these polymers is much weaker than that observed for other aromatic vinyl polymers.

Fluorescence spectra of PVPh, PPhEVE, and EPh in rigid glasses at 77 K are shown in Fig. I-26. The fluorescence band is shifted to lower frequencies in the order of EPh, PPhEVE, and PVPh. The clear vibrational structure bands observed for PVPh and PPhEVE are similar to that for EPh and the fluorescence lifetimes of PVPh and PPhEVE are much the same as that of EPh at 77 K (Table I-8). These facts suggest that there is little interaction between neighboring Ph chromophores in PPhEVE and PVPh in rigid glass at 77 K.

Table I-8 shows the relative fluorescence quantum yields, taking the value of EPh as unity, and the fluorescence lifetimes. The fluorescence quantum yields and lifetimes of PVPh and PPhMA



Normalized fluorescence spectra of (1); EPh, (2) -----; PPhEVE, (3) .---; PPhMA, and (4) -----; PVPh in solutions at room temperature. Solvents; (1), (2), (3); MTHF-THF (3:2) mixture, (3); chloroform containing 1.0% ethanol. Excitation wavelength; 345 nm. Normalized fluorescence spectra of (1); PPhEVE, (3) .----; PPhMA, and (4) ------; PVPh in solutions at room temperature. Solvents; fig. I-26. Normalized fluorescence spectra of (1); PPhEVE, (3) .----; PPhMA, and (4) ------; PVPh in solutions at room temperature. Solvents; fig. I-26. Normalized fluorescence spectra of (1); PPhEVE, (3) .----; PPhMA, and (4) ------; PVPh in solutions at room temperature. Solvents; fig. I-26. Normalized fluorescence spectra of (1); PPhEVE, (3); PP



Normalized fluorescence spectra of (1) EPh, (2) PPhEVE, and (3) PVPh in MTHF-THF rigid glasses at 77 K. Excitation wavelength; 330 nm.

at room temperature are much smaller than those of EPh and PPhEVE. The decay curve of PVPh has an initial fast component. The fluorescence spectra of PVPh in MTHF-THF solution at various temperatures are shown in Fig. I-27. The fluorescence intensity decreases and the vibrational bands broaden with raising temperature. Temperature dependence of the fluorescence yields are shown in Fig. I-28. The changes observed for EPh, PPhEVE, and PPhMA are small and their magnitudes are nearly equal. On the other hand, only PVPh shows a large decrease with raising temperature. The same tendency is observed for the temperature





Temperature dependence of fluorescence spectra of PVPh $(5.1 \times 10^{-4} \text{ M})$ in MTHF-THF solution. Excitation wavelength; 310 nm. (1) -130, (2) -86.5, (3) -37.7, (4) 6, and (5) 63.5 °C. The fluorescence (6) is 1.95 times the fluorescence (5).



Temperature dependence of the fluorescence yields. The value of η/η_{rt} exhibits the ratio of the fluorescence yield at some temperature to that at room temperature. (1) \bigcirc ; EPh, (2) \triangle ; PPhEVE, (3) \square ; PPhMA, and (4) \bigcirc ; PVPh. Solvent: (1), (2), (4); 1,4-dioxane, (3); chloroform containing 1.0% ethanol. These values were uncorrected for decrease in density of solvents with an increase in temperature.

 Table I-8.
 Spectroscopic data of vinyl polymers with pendant phenanthryl chromophores in MTHF-THF

Polymer	Relative yield at 20°C	Fluorescence lifetime ^b /ns				Phosphorescence
		77 K	-40°C	24 °C	50 °C	lifetime/s
EPh	1.00	59	50	50	48	4.4
PPhEVE	0.85	59	49	47	46	3.4
PPhMA ^a)	0.15			16 ^{c)}	15°)	
\mathbf{PVPh}	0.17	58	40	21°)	20°)	2.8

a) Solvent: chloroform containing 1.0% ethanol. b) The fluorescence was observed through Toshiba UV-DIC and UV-35 filters. Error is ± 2 ns. c) The initial fast decay component was observed.

dependence of the fluorescence lifetimes (Table I-8). Namely, the fluorescence lifetimes of EPh, PPhEVE, and PVPh in rigid glass at 77 K are all equal. Only the lifetime of PVPh, however, decreases unusually with an increase in temperature.

As mentioned above, the vibrational structure of fluorescence and its lifetimes of PVPh and PPhEVE are similar to those of EPh in rigid glass at 77 K. However, the phosphorescence of PVPh and PPhEVE is markedly different from that of EPh, as is shown in Fiq. I-29. The phosphorescence band shifts to lower frequency and the vibrational structure broadens in the order of EPh. PPhEVE, and PVPh. The phosphorescence lifetime shortens in the same order (Table I-8). The red shift of the phosphorescence band is much larger than that of the fluorescence. PVPh and PPhEVE showed the delayed fluorescence resulting from a T-T annihilation between two migrating triplet excitons. These facts indicate that an interaction between neighboring Ph chromophores in the triplet state is much larger than that in the singlet state in rigid glass at 77 K.

PPhEVE resembles EPh very closely in fluorescence properties, forms no excimer, and does not show the anomalous fluorescence observed for PCzEVE. The fluorescence spectra of PPhMA is markedly different from that of EPh and its lifetime is very short. The temperature dependence of the fluorescence yield, however, is similar to those of EPh and





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PPhEVE. This indicates that PPhMA forms no excimer. The fluorescence properties observed for PPhMA are more likely to be affected by the substituent groups (-O-CO-). On the other hand, PPyMVE, PNMVE, and PNMA show the excimer fluorescence in dilute solution at room temperature. Therefore, the failure of PPhEVE and PPhMA to form an excimer is considered to be attributable to the nature of Ph, that is, Ph forms an excimer with difficulty.

Emission properties of PVPh clearly indicate that the interaction between neighboring Ph chromophores in PVPh is the strongest in the polymers studied. It has been reported, from fluorescence yield and decay curve measurements, that phenanthrene does not show concentration quenching of fluorescence at room temperature.^{84,85)} Therefore, if PVPh forms no excimer, the fluorescence yield of PVPh is expected to be as large as that observed for EPh. The present result is not so. Namely, the fluorescence yield of PVPh is much lower than that of EPh (Table I-8) and decreases largely with an increase in temperature (Fig. I-28). These facts suggest that PVPh forms an excimer which has a very low fluorescence quantum yield. This is compatible with the fact that the phenanthrene excimer in the crystalline state under a high pressure has a low fluorescence quantum yield. 77)

ECz, 5EBCz, 7EBCz, and EPh show no excimer fluorescence in concentrated solution at room temperature. PVCz, P5VBCz, and P7VBCz, however, show the excimer fluorescence in dilute solution at room temperature. It is suggested from the above-mentioned results that PVPh forms an excimer, although it shows no clear excimer fluorescence. Thus, the following conclusions were drawn: in the case of vinyl polymers with the aromatic chromophores

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which form no excimer, an excimer is formed in the vinyl polymers in which the aromatic chromophores are directly connected to the skeletal chains, but it is not formed in the vinyl polymers in which the aromatic chromophores are widely spaced on the skeletal chains. PCZEVE is considered to be rather a special case.

Summary

The emission properties of EPh, PVPh, PPhMA, and PPhEVE were investigated in solution. These vinyl polymers showed no clear excimer fluorescence in the longer-wavelength region. Both the fluorescence quantum yield and lifetime of PVPh decreased largely with raising temperature in the higher temperature region, suggesting that PVPh forms an excimer which has a very low fluorescence quantum yield. The fluorescence properties of PPhMA and PPhEVE indicated little interaction between neighboring Ph chromophores in the singlet state. This originates from the nature of Ph, that is, Ph forms an excimer with difficulty. The phosphorescence spectrum and its lifetime of PPhEVE indicated a presence of a weak interaction between Ph chromophores in the triplet state.

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PART II

EMISSION SPECTRA OF AROMATIC VINYL POLYMER FILMS AND EXCITATION ENERGY MIGRATION IN THE FILMS

INTRODUCTION

It is known that PVCz shows the largest photoconduction among the vinyl polymers investigated so far.¹⁾ Okamoto et al. have suggested that the large photoconductivity of PVCz can be attributed both to the high efficiency of carrier generation due to the field-assisted thermal dissociation of an exciplex formed between the singlet exciton of PVCz and some electronaccepting impurity and to the ease of carrier migration due to the overlap of π -orbitals of neighboring Cz chromophores in one polymer chain.²⁻⁶⁾ Therefore, studies on singlet exciton migration in the aromatic vinyl polymer films are very important in understanding their photoconductivity, and also of interest in connection with the tacticity of the polymer chain.

It is known that triplet excitation energy also migrates very efficiently in aromatic vinyl polymers. The triplet excitation energy of PVCz is fairly high as compared with that of other aromatic vinyl polymers. Therefore, the photocarrier generation from the triplet excited state might be possible in PVCz.⁷⁾ Studies on the triplet exciton migration also are important, and of interest in connection with the singlet

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exciton migration.

In order to reveal an origin of the high photoconductive properties of PVCz films, it is of interest and important to compare the spectroscopic properties of an amorphous film of the model compound with those of an amorphous PVCz film.

In this part, emission spectra of the vinyl polymer films with pendant carbazolyl groups and migrations of the singlet exciton (chapter 1) and the triplet one (chapter 2) in their films have been investigated. In chapter 1, fluorescence spectrum of an amorphous DCzP film and the singlet exciton migration in the films also have been investigated and compared with those of the amorphous vinyl polymer films. Chapter 1

Singlet Excitation Energy Migration

Introduction

Klöpffer has reported that the results of fluorescencequenching experiments in an amorphous PVCz film are consistent with a hopping model of monomer exciton migration, in which excitons can migrate in a polymer film and both excimer-forming sites and guest molecules act competitively as exciton traps.⁸⁾ From a lifetime quenching observed in a PVCz film doped with perylene, Powell et al. have recently proposed a model containing dimer sites besides guest molecules and the excimer-forming sites.⁹⁾ Concerning an undoped PVCz film, Johnson and Offen have suggested the presence of a dimer site as a result of measuring the fluorescence decay time of a PVCz film at 77 K.¹⁰⁾ Klöpffer and and Fischer have showed that the two different types of the spectra were observed in the prompt fluorescence and phosphorescence at 77 K, independent of the polymerization methods.¹¹⁾

It is of interest to investigate how the difference in the tacticity of the PVCz samples affects the singlet-excitation-energy migration. The study on singlet-excitation-energy migration in vinyl polymer films with carbazolyl(Cz) chromophores widely spaced on the skeletal chains is of interest in connection with that of a PVCz film.

The author has found that DCzP, a dimeric model compound of PVCz, forms an amorphous state by a fast evaporation of

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solvent. It is of interest to investigate the spectroscopic properties of the amorphous DCzP(DCzP(a)) film and to compare them with those of an amorphous PVCz film. A few investigations have been carried out to ascertain how the efficiency of the singlet-excitation-energy migration changes when going from a crystalline to an amorphous state, 12-15) However, no comparison of the optical properties of an amorphous film of a model compound with those of the corresponding vinyl polymer has hitherto been performed.

In this chapter, the author describes the migration of the electronic-excitation energy in films of PVCz(r), PVCz(c), brominated PVCz(BPVCz), PCzEVE, poly(N-acryloylcarbazole)(PACz) and DCzP(a). DMTP, perylene, and p-bis(trichloromethyl)benzene (TCB) were used as the guest molecules. In the case of perylene, the long-range resonant energy transfer is somewhat possible.¹⁶⁾ However, the long-range resonant energy transfer from a Cz chromophore to a DMTP or TCB molecule is quite impossible, because DMTP and TCB have no absorption in the wavelength region where the fluorescence of the PVCz film is observed.¹⁶⁾ Therefore, DMTP and TCB are very useful in studying the phenomenon of the singlet exciton migration.

Experimental

PACz was the same as used in Ref. 17. BPVCz was prepared by the bromination of PVCz with N-bromosuccinimide and had one bromine atom per 1.1 Cz groups.¹⁸⁾

Films of the model compounds were cast on quartz glass plates from a benzene solution followed by a fast evaporation of the

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solvent at room temperature. Polymer films were cast on a quartz or pyrex glass plate from a benzene or 1,2-dichloroethane solution followed by slow evaporation of the solvent in a vessel nearly saturated with solvent vapor and dried in vacuo. The thickness of the film was about 0.1-0.06 μ m for DCzP(a), about 4 μ m for PCzEVE, and about 7 μ m for other polymers. The apparatus for measurements were the same as those in the previous part.

Results and Discussion

(1) Fluorescence Spectra.

The fluorescence spectra of PVCz(r) and PVCz(c) films are shown in Figs. II-1 and II-2. The difference in the spectra between PVCz(r) and PVCz(c) films is similar to that observed in fluid solution. That is, the fluorescence intensity in the shorter-wavelength region of PVCz(r) films is larger than that of PVCz(c) films. Therefore, the emission band in the shorterwavelength region was assigned to the second excimer fluorescence,





(1) 293 K, (2) 77 K, (3) the component of the sandwichlike excimer fluorescence at 293 K, (4) the component of the second excimer fluorescence at 293 K, (5) the component of the sandwich-like excimer fluorescence at 77 K, (6) the component of the second excimer fluorescence at 77 K.





(1) 293 K, (2) 77 K, (3) the component of the sandwichlike excimer fluorescence at 293 K, (4) the component of the second excimer fluorescence at 293 K, (5) the component of the sandwich-like excimer fluorescence at 77 K, (6) the component of the second excimer fluorescence at 77 K. and that in the longer-wavelength region, to the sandwich-like excimer one. The fluorescence spectra of both polymer films were resolved into two components. The resolution spectra thus obtained also are shown in Figs. II-1 and II-2. The intensity of the second excimer fluorescence relative to the sandwich-like excimer one and the fluorescence yields are listed in Table II-1.

The presence of two kinds Table II-1. of emitting species in the fluorescence of PVCz(r) and PVCz(c) was clarified also by measuring the fluorescence decay curves of the polymers. The fluorescence decay curves of

THE INTENSITY OF THE SECOND EXCIMER						
FLUORESCENCE RELATIVE TO THE SANDWICH-LIKE						
ONE $(I_{ m S}/I_{ m E})$ and the quantum efficiencies						
of the host fluorescence $(\eta_{\rm H})$						
Host	$I_{ m S}/I_{ m E}$	$\eta_{\rm H}$				
PVCz(r)	0.083	0.045				
PVCz(c)	0.013	0.041				
PVCz(r) ^a)	0.93					
PVCz(c) ^a)	0.55					
PCzEVE		0.10				
PACz		0.0002				
BPVCz		0.001				

both polymer films in air at 293 K clearly involve two-component (20+3 and 6+1 ns) in the shorter wavelength region, but onecomponent (22+2 ns) in the longer wavelength region. The intensity of the short-lived component relative to the long-lived one for PVCz(r) films in the shorter wavelength region is larger than that for PVCz(c) films. Therefore, the long- and shortlived components correspond to the sandwich-like excimer fluorescence and the second excimer one, respectively. Similar results were obtained also at 77 K; the long- and short-lived components are 17+2 and 6+2 ns, respectively. These decay constants are roughly in agreement with the values reported by Powell et al. (20 and 10 ns).⁹⁾ The lifetime of the sandwichlike excimer fluorescence of PVCz films is shorter than that of PVCz in fluid solution (ca. 40-42 ns)^{19,20)} and that of PVCz films obtained by Johnson and Offen (43 ns).¹⁰⁾

a) At 77 K.

The fluorescence spectra of PCzEVE films undoped and doped with DMTP are shown in Fig. II-3. PCzEVE films show only a

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broad, structureless fluorescence spectrum with a peak at ca. 26050 cm⁻¹. The fluorescence decay curve in air consists of one-component in both the shorter and longer wavelength regions (4-5 and 10 -11 ns, respectively). Two component decay curve is observed in the medium wavelength region. This fact



Fluorescence spectra of PCzEVE films doped with DMTP at 293 K. DMTP concentration; (1) 0, (2) 4.44×10^{-4} , (3) 8.88×10^{-4} , and (4) 1.18×10^{-2} (mol/ CzEVE unit mol). Fluorescence spectrum of the undoped film at 77 K is also given by a dotted line (5).

indicates that the two kinds of emitting species are present in PCZEVE films at 293 K. Although the fluorescence spectrum of PCZEVE films at 77 K has the structure, it does not seem to be the monomeric fluorescence of a Cz chromophore because of a large Stokes shift (1540 cm⁻¹).

The fluorescence spectra of PACz and BPVCz films undoped and doped with DMTP or perylene are shown in Fig. II-4. The fluorescence efficiency of PACz films is extremely low (Table

II-1) because of the presence of intersystem crossing from the lowest ${}^{1}\pi$, π^{*} state to the lowest ${}^{3}n$, π^{*} state due to a carbonyl group. PACz films show a fluorescence spectrum consisting of two components. The spectra of PACz and BPVCz films change rapidly under



Fig. II-4. Wave number/10³ cm⁻¹

Fluorescence spectra of PACz and BPVCz films at 293 K. PACz films; (1) undoped, (2) doped with DMTP (2.32×10^{-2} ACz unit mol). BPVCz films; (3) undoped, (4) doped with perylene (5.9×10^{-3} mol/mol basic unit).

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irradiation by an exciting light.

The fluorescence spectra of DCzP(a), polycrystalline DCzP (DCzP(c)), and polycrystalline i-PrCz films in air are shown in Fig. II-5. The spectrum of DCzP molecule in aerated benzene solution also is shown for comparison. The fluorescence spectrum of DCzP(a) films is remarkably different from those of amorphous PVCz and PCzEVE films. That is, DCzP(a) films show a structured fluorescence and a tailing in the long wavelength side of the band. The intensity of the second peak(26750 cm⁻¹) in the structured



Fig. II-5. Fluorescence spectra of (1) DCzP(a), (2) DCzP(c), (3) i-PrCz films and (4) DCzP in aerated benzene solutions at 293 K.

band is anomalously large, compared with those observed for polycrystalline i-PrCz films and the DCzP molecule in solution. The position of the second peak is nearly equal to that of the second excimer fluorescence in PVCz films. The fluorescencedecay curve of DCzP(a) films involves multi-components $(23\pm3 \text{ ns}$ and an initial fast decay) in the longer-wavelength region (λ_{obs} >400 nm) and one component (ca. 7 ns) in the shorter-wavelength region. The long-lived component in the longer-wavelength region is nearly equal in the lifetime to the sandwich-like excimer fluorescence in PVCz films (22±2 ns). Excitation spectra of DCzP(a) films, obtained by monitoring at several wavelengths(355-500 nm), duplicated the absorption spectrum of the films exactly. Therefore, the fluorescence spectrum of DCzP(a) films seems to be composed of three components; monomer, second excimer, and sandwich-like excimer fluorescences.

As is shown in Fig. II-5, thin DC2P(c) films show no sandwich-like excimer fluorescence. This is supported also by the fact that the long-lived component (ca. 23 ns) in the longerwavelength region was not observed. This is the same as the case of crystalline i-PrCz. However, the spectrum of thin DC2P(c) films is different from that of thin polycrystalline i-PrCz films. That is, the intensity of the second peak (ca. 26900 cm⁻¹) in the structured band is larger than that of the first peak, compared with the spectrum of i-PrCz films. The position of the second peak was the almost same as that observed for DC2P(a) films and that of the second excimer fluorescence in PVCz films. Therefore, thin DC2P(c) films may show both monomer and second excimer fluorescences. The presence of the second excimer site in DC2P(c) films seems to be also suggested by the fact that

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formation of a microcrystal of DCzP in benzene matrix at low temperature leads to formation of an excited state like that observed for PVCz (second excimer).¹⁹⁾

(2) Singlet Excitation Energy Migration.

The fluorescence spectra of PVCz(r), PVCz(c), and PCzEVE films doped with DMTP or perylene at 77 K are shown in Fig. II-6.

In the case of all polymer films studied at 293 and 77 K, by doping of DMTP, TCB, or perylene, the host fluorescence decreases and is replaced by the exciplex or guest fluorescence. The exciplex is formed between DMTP and a Cz chromophore. As DMTP and TCB have no

absorption in the wavelength



Fluorescence spectra of PVCz(r), PVCz(c) and PCzEVE films doped with DMTP or perylene at 77 K. (1) PVCz(r) doped with DMTP (3.26×10⁻³ mol/VCz unit mol), (2) PVCz(c) doped with DMTP (3.02×10⁻³ mol/VCz unit mol), and (3) PCzEVE doped with perylene (1.15×10⁻⁴ mol/CzEVE unit mol).

region where the host fluorescence of these films is observed, the dipole-dipole resonance cannot be responsible for the energy transfer to DMTP and TCB molecules.^{8,16)} In fact, no decrease in the fluorescence lifetimes of PVCz(r) films by doping of DMTP is observed. That is, the fluorescence lifetimes of the PVCz(r) film doped with a small amount of DMTP (ca. 3×10^{-3} mol/mol basic unit) in the shorter-wavelength region are 17 ± 3 and 5 ± 2 ns at 293 K and 19+2 and 6+1 ns at 77 K.

In the present case, therefore, the hopping model of monomerexciton migration is applicable at 293 and 77 K. We should consider the case of the presence of two kinds of traps (for

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example; the sandwich-like excimer-forming site, c_E [mol/mol basic unit], and the second excimer site, c_S [mol/mol basic unit], in the case of PVCz films) in a way similar to the treatment in Ref. 8. Then, the quenching factor of the host fluorescence, Q_{μ} , is expressed by the following equations;

$$Q_{\rm H} = (\eta_{\rm H,0} - \eta_{\rm H})/\eta_{\rm H} = 0.66 \, \text{nc} / [1 + 0.66 \, \text{n} (c_{\rm E} + c_{\rm S})] = n'c \qquad (1)$$

n' = 0.66 n/[1 + 0.66 n(c_{\rm E} + c_{\rm S})] (2)

$$n = n_{s} / (n_{e} + n_{1})$$
 (3)

where n_s , n_e , and n_1 are the relative probabilities (per unit time) of jumping, radiative, and nonradiative decay, $\eta_{H,0}$ and η_H are quantum efficiencies of the host fluorescence in the absence and in the presence of the guest molecule in a concentration of c [mol/mol basic unit], and n and n' are the number of jumps during the lifetime in the absence and in the presence of trap sites, respectively. The quenching factor of the monomer fluorescence, Q_M , may be expressed by;

 $Q_{\rm M} = 0.66n(c_{\rm E} + c_{\rm S}) = 0.66nc_{\rm T}$ (4) In the case of polymer films, where no monomer fluorescence can be observed, $Q_{\rm M} >> 1$. Therefore,

$$n' = 1/(c_{E} + c_{S}) = 1/c_{T}$$
 (5)

$$Q_{\rm H} = c/(c_{\rm E} + c_{\rm S}) = n'c$$
 (6)

The ratio of the fluorescence intensities guest/host(I_G/I_H) is;

$$I_{G}/I_{H} = c\eta_{G}/(c_{E}\eta_{E} + c_{S}\eta_{S}), \qquad (7)$$

where $n_{\rm G}$ is the quantum efficiency of the guest fluorescence in the host film under direct excitation. The relative intensity of the second excimer fluorescence to the sandwich-like excimer one is expressed by

$$I_s/I_E = c_s \eta_s/c_E \eta_E$$
. (8)
On the other hand, in the case of DCzP(a) films, the value of

 $1/c_{\rm T}$ seems to be smaller than n', since the value of $Q_{\rm M}$ could not be determined. Although the experimental values of $Q_{\rm H}$ scatter a little, the measurement of $Q_{\rm H}$ at various concentrations of a guest gives the value of n' and $c_{\rm T}$ according to Eqs. 5 and 6, respectively. In the case of PVCz films, the measurement of $I_{\rm G}/I_{\rm H}$ at various concentrations of the guest gives the values of $c_{\rm E}n_{\rm E}$ and $c_{\rm S}n_{\rm S}$ according to Eqs. 7 and 8, respectively, using the values of $\eta_{\rm G}$ and $I_{\rm S}/I_{\rm E}$ in Tables II-1 and II-2. Figure II-7 shows the bilogarithmic plots of $Q_{\rm H}$ or $I_{\rm G}/I_{\rm H}$ and c, as example, in the case of the PVCz(c)-DMTP system. The values thus obtained are listed in Table II-2.

In the case of the PVCz(r) and PVCz(c) films, the values of n', $c_{T} (=c_{E}+c_{S})$, $c_{E}\eta_{E}$, and $c_{S}\eta_{S}$ are obtained by the method mentioned above. The value of $c_{E}+c_{S}$ at 293 K is nearly equal to that of c_{F} obtained by Okamoto et al. in the PVCz(r)-DMTP system, on the assumption that the host fluorescence is emitted only from the sandwich-like excimer.¹⁶⁾ The values of n' and $c_{E}^{+}c_{s}^{-}$ are almost the same for PVCz(r) and PVCz(c) films. The value of $c_E^{}\eta_E^{}$ is almost the 10same for PVCz(r) and PVCz(c) 5 films, while the value of c_cn_c for PVCz(r) films is $Q_{
m H}$ or $I_{
m G}/I_{
m H}$ higher than that for PVCz(c) 0.5 films. It can safely be considered that the efficiencies of each 0.1 10^{-3} 10^{-2} Fig. II-7. DMTP concentration [mol/mol basic unit] Ratio of fluorescence intensities guest/host I_G/I_H fluorescence ($\eta^{}_{\rm E}$ and $\eta^{}_{\rm S})$ of PVCz(r) films are equal to (1) and quenching factor of host fluorescence $Q_{\rm H}$ (2) as a function of guest concentration for PVCz(c)-DMTP system at 293 K. those of PVCz(c) films.

	T TTUO	at 275	and // Kt				
Host	Guest	η _G b	c _T x 10 ³ (mol/mol asic unit)	n '	c _E n _E x 10 ⁵	c _s n _s x 10 ⁵	l/Å ^{c)}
PVCz(r)	perylene		0.83	1200			120
PCzEVE	**		0.27	3700			210
BPVCz		0.27	(19) ^{d)}	(52) ^{d)}			
DCzP(a)	11		<0.17	5900	· · ·		<u></u>
PVCz(r)	DMTP	0.043	2.7	370	10	0.83	67
PVCz(c)		0.039	2.2	450	(11) ^{b)}	(0.11) ^{b)}	74
PCzEVE	11	0.034	0.45	2200		 	
PACz	11	0.004	24 ^{e)}	42 ^{e)}			·
			(25) ^{f)}	(40) ^{f)}			
DCzP(a)	11	· · ·	<0.77	1300			
PVCz(r) ^a) ₁₁	0.17	3.5	290	40	37	60
PVCz(c) ^a) "	0.15	3.2	310	43	23	63
PVCz(r)	тсв	· <u>····</u>	1.4	690	······································	· · · · · · · · · · · · · · · · · · ·	92
PCzEVE	U		0.5	2000			
DCzP(a)	11		<0.28	3600			

Table II-2. Energy Transfer by the Exciton Diffusion Process in Amorphous Films at 293 and 77 K.

a) At 77 K. b) The values are less reliable because of the large error in I_s obtained by a resolution of the host fluorescence spectrum. c) $L = an'^{1/2}$, where L is the mean exciton migration length and a is the interchromophore sparation (3.5 Å). d) The values were obtained in a way similar to that used in the treatment in Ref. 8, on the assumption that the host fluorescence is emitted by one trap site. e) The values were obtained from Q_H , on the assumption that the fluorescences in the shorter and longer wavelength regions are emitted by some intrinsic traps and an excimer, respectively. f) The values were obtained from the ratio of (I_G/I_Hc) , on the assumption that the fluorescences in the shorter and longer wavelength regions are emitted by a monomer and an excimer, respectively.

Therefore, the concentration of the sandwich-like excimer site (c_E) in PVCz(r) films is nearly equal to that in PVCz(c) films. On the other hand, the concentration of the second excimer (c_S) in PVCz(r) films is higher than that in PVCz(c) films by a factor

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of about 1.6. This difference in the value of c_s does not clearly appear in the value of $c_E^+c_s^-$. This fact suggests that the value of c_E^- is larger than that of c_s^- in both films.

The values of $c_{_{\rm F}}\eta_{_{\rm F}}$ for PVCz(r) and PVCz(c) films obtained at 77 K are larger than those obtained at 293 K by a factor of 4. It may safely be considered that the values of $\eta_{\rm E}$ and $\eta_{\rm S}$ for PVCz(r) films change with the temperature in a manner similar to that for PVCz(c) films. The values of c_{E} and c_{S} are determined by the casting temperature of the film because of the high glass-transition temperature of PVCz. Therefore, this increase in the value of $c_E^{}\eta_E^{}$ seems to be caused mainly by a change in the value of η_{E} . On the other hand, the value of $c_{s}\eta_{s}$ obtained at 77 K is much larger than that obtained at 293 K. This increase in c_n with a decrease in the temperature is too large to be caused only by an increase in $\eta_{\rm s}$. The value of $\rm c_{s}$ should be considered to vary apparently with temperatures. This suggests that the second excimer site is a shallow trap and acts more effectively as a trap at low temperatures.

The value of $c_{\rm T}$ for PCZEVE films is much smaller than those for the PVCz(r) and PVCz(c) films. This suggests that it is difficult for Cz chromophores to form intrinsic trap sites because of the large distance between neighboring Cz chromophores. The value of n for PCZEVE films might be smaller than the values of n for PVCz(r) and PVCz(c) films, judging from the large distance between neighboring Cz chromophores. However, even in the case of PCZEVE films, $Q_{\rm M} >> 1$, that is, the singlet exciton migration is limited by the intrinsic trap sites. Thus, the value of n' for PCZEVE films is larger than that for PVCz films by a factor of ca. 5.

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In the case of the PACz and BPVCz films, the emitting species of the host fluorescence are ambiguous. Judging from the profile of the fluorescence spectrum of PACz films, the broad fluorescence band with a peak at ca. 25000 cm⁻¹ might be assigned to the excimer fluorescence. The fluorescence band in the shorter wavelength region might be attributable to a monomer or to some intrinsic trap sites. The host fluorescence of BPVCz films is considered to be emitted from some trap sites. Irrespective of the treatments used to obtain the value of n' (Table II-2), the values of n' for both polymer films are very This may be attributable to the short exciton lifetime small. because of the enhanced intersystem crossing and the presence of extrinsic trap sites (photoproducts and/or impurities introduced during a bromination reaction).

The concentration of the trap sites for DCzP(a) films was smaller than those for the polymer films, irrespective of the guest molecule, except for PCzEVE films doped with DMTP. This suggests that the excimer sites are formed with greater ease in a polymer film than in an amorphous film of the model compound, even if carbazolyl chromophores are widely spaced on the skeletal chain.

It has been reported that the singlet exciton migration length in crystalline films is more than one order of magnitude larger than that in amorphous films.¹²⁻¹⁵ In the present case, the value of n' in the crystalline film(DCzP(c)) was smaller than that in the amorphous film by a factor of about 3. The reasons for the poor energy transfer to a guest molecule in DCzP(c) films are probably as follows: (1) DCzP(c) has a trap site (second excimer site), and (2) the guest molecule may

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segregate in a DCzP(c) film and so may not be dissolved in the film uniformly.

The number of Cz chromophores covered by a singlet exciton during the lifetime decreases in the following order; DCzP(a) > PCzEVE > PVCz(r) ~ PVCz(c) > PACz ~ BPVCz.

Summary

The migration of electronic excitation energy in films of PVCz(r), PVCz(c), PCzEVE, BPVCz, PACz, and DCzP(a) has been studied by means of fluorescence-quenching experiments, using DMTP, perylene, or TCB as a guest molecule.

PVCz(r and c) films showed a broad structureless fluorescence composed of the second and sandwich-like excimer fluorescences. The second excimer fluorescence intensity was larger for PVCz(r) films than for PVCz(c) films. The lifetime of the second and sandwich-like excimer fluorescences were ca. 6 and ca. 21 ns, respectively. PCzEVE films showed only a broad, structureless fluorescence composed of two unknown components. DCzP(a) films showed fluorescence composed of three components; monomer, second excimer, and sandwich-like excimer fluorescences.

In the case of all films studied, the hopping model of monomer-exciton migration was applicable at 293 and 77 K. The concentration of the sandwich-like excimer site(c_E) in PVCz(r) films was nearly equal to that in PVCz(c) films, while the concentration of the second excimer site(c_S) in PVCz(r) films was higher than that in PVCz(c) films. The value of c_E was much larger than that of c_S for both films. The value of c_S increased apparently with a decrease in the temperature, suggesting that

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the second excimer site is a shallow trap. Even in PCZEVE films, the singlet exciton migration was limited by the intrinsic trap sites. DCzP(a) films had the lowest concentration of the trap sites (~3 x 10^{-4} mol/mol basic unit), suggesting that the excimer sites are formed with greater ease in a polymer film than in an amorphous film of the model compound, even if Cz chromophores are widely spaced on the skeletal chain.

The number of carbazolyl chromophores covered by a singlet exciton during the lifetime decreases in the following order: $DCzP(a) > PCzEVE > PVCz(r) \sim PVCz(c) > PACz \sim BPVCz$. This order was explained by the concentration of the intrinsic trap sites depending on the distance between neighboring Cz chromophores, and by the substituent effect determining the lifetime of the singlet exciton.

Chapter 2

Triplet Excitation Energy Migration

Introduction

The triplet energy migration and transfer in vinyl polymers with large aromatic rings, such as naphthyl and carbazolyl groups, in rigid solution at 77 K have been studied in detail. 21-23) The result indicates that the triplet excitons can migrate along the polymer chain, and that delayed fluorescence is caused by a homogeneous triplet-triplet (T-T) annihilation between two intramolecular migrating triplet excitons. David et al. reported the triplet energy transfer in vinyl polymers with small aromatic rings in solid film, which emit only phosphorescence and no delayed fluorescence. 24,25) They showed that the triplet exciton migration frequencies in such polymer films were smaller than those in the corresponding crystals. Concerning the vinyl polymer films with large aromatic rings, Fox at al., and Klöpffer and Fischer have qualitatively studied poly(1vinylnaphthalene) and PVCz, respectively. 11,26) Klöpffer and Fischer showed that the phosphorescence of PVCz films is governed by two kinds of traps, whose concentration was estimated to be 5 x 10^{-3} (mol of trap/mol of basic unit) from the saturated concentration of the sensitized dopant phosphorescence, and that PVCz films emitted no delayed fluorescence. However, the triplet energy migration in solid film has never hitherto been studied quantitatively on such vinyl polymers.

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Therefore, it would be interesting to investigate quantitatively the spectroscopic behavior of triplet excitons in the vinyl polymers with large aromatic rings and to compare the difference between in rigid solution and in solid film.

Experimental

Commercial naphthalene(NAPH) and fluorene of a scintillation grade were purified by zone-refining. Commercial 1,3-pentadiene (1,3-PD), a well-known triplet quencher, was used after distillation.

The apparatus and technique for measuring the emission spectra were described in the previous part. All the samples were excited by 345 nm light, which is not absorbed by the triplet quenchers. It was confirmed that the application of a phosphoroscope factor²⁷⁾ was not necessary in the case of film. In the case of rigid solution, the delayed fluorescence spectra were, however, corrected by using a phosphoroscope factor. Phosphorescence decay curves were recorded on a X-t recorder and delayed fluorescence lifetimes were determined from oscilloscope traces.

Results and Discussion

(1) Delayed Emission Spectra in Rigid Solution and in Film.

The delayed emission spectra of PVCz(r), PVCz(c), and PCzEVE in rigid solution and in film at 77 K are shown in Figs. II-8 and II-9, respectively. Concerning PVCz(r and c) in rigid solution, Yokoyama et al.⁷⁾ and Klöpffer¹¹⁾ have reported as follows : (1)

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the delayed emission spectrum consists of a delayed fluorescence and a phosphorescence; (2) the intensity and the lifetime of the delayed fluorescence are dependent on the molecular weight, and the delayed fluorescence is not observed for PVCz with a low molecular weight $(\overline{DP} \leq 50)$; (3) the phosphorescence is emitted from a trap site in the polymer chain because the phosphorescence (0-0 band; 420 nm) is located at a lower energy than that of i-PrCz (0-0 band; 412 nm). However, they have not reported on the relation between the spectra and the tacticity of the samples. In the present



Delayed emission spectra of (1) PVCz(r), (2) PVCz(c), and (3) PCzEVE in MTHF-THF rigid solutions at 77 K (uncorrected). The prompt fluorescence (4) of PCzEVE in MTHF-THF rigid solution at 77 K was also given for comparison.





results, the delayed fluorescence spectra are a little different between PVCz(r) and PVCz(c), as is shown in Fig. II-8, although the lifetimes (τ_{df}^{0}) are almost the same (ll ms for PVCz(r) and l2 ms for PVCz(c)). The phosphorescence spectra are almost the same between PVCz(r) and PVCz(c). The lifetime (τ_{ph}^{0}) is 8.1 \pm 0.2 s for PVCz(r and c).

In the case of film, Klöpffer¹¹⁾ showed that PVCz(r and c)

films emitted only a phosphorescence governed by two traps (two component decay; 7.9+1 and 1.4 s), and no delayed fluorescence, and that two different types of spectra were observed in the phosphorescence and in the prompt fluorescence at 77 K, independent of the polymerization methods. In the present results, the delayed emission spectra of PVCz(r and c) consist of two emission bands, as is shown in Fig. II-9. The prompt fluorescence spectra of the films and the example of the resolution spectrum are also shown in Fig. II-9. The further purification of the polymer (for example; reprecipitation up to ten times and/or extraction with hot methanol) gave no change in the spectra. The broad delayed emission in the longer wavelength region is invariably monophotonic in origin, and this lifetime consists of two components (7.9+0.7 and 1.6+0.5 s). Therefore, this emission is considered to be the phosphorescence emitted both from Trap I (the sandwich-like excimer forming site) and from Trap II, as has been reported by Klopffer.⁵⁾ Both the position and profile of the delayed emission band in the shorter wavelength region are the same as those observed for the normal fluorescence at 77 K, as is shown in Fig. II-8, and the emission intensity depends quadratically on the excitation light intensity. The lifetime of this emission is about 50 ms for PVCz(r and c). These facts indicate that this emission band can be assigned to the delayed fluorescence resulting from a homogeneous T-T annihilation between two migrating triplet excitons in film, as is the case with rigid solution. The delayed fluorescence of PVCz(r) differed from that of PVCz(c) in the similar manner observed for the prompt fluorescence.

The delayed emission spectra of PCzEVE in rigid solution

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and in film at 77 K also consist of two bands, as is shown in Figs. II-8 and II-9. Both the positions(λ_{max} ; 360 and 376 nm in rigid solution and 371, 388, and 409(shoulder) nm in film) and the profile of the emission band in the shorter-wavelength region are nearly the same as is observed for the normal fluorescence, and the emission intensity depends quadratically on the excitinglight intensity. Therefore, the emission bands of PCZEVE are identified as the delayed fluorescence. The τ^0_{df} of PCzEVE films is 45 ms, while that in rigid solution has not been determined because of low intensity. On the other hand, the delayed emission band at a longer wavelength (λ_{max} ; 418, 446 nm in rigid solution and about 495 nm in film) is monophotonic in origin. Therefore, this emission band is identified as the phosphorescence emitted from the trap site, by analogy with PVCz(r and c). The τ_{ph}^{0} in rigid solution is 6.6±0.1 s, while that in film consists of two components (3.5+0.2 and 0.7+0.05 s), which is smaller than that of PVCz(r and c) by a factor of ca. 2.

(2) Triplet Energy Migration in Film and in Rigid Solution.

The delayed emission spectra of PVCz(r) and PCZEVE films doped with NAPH of various concentrations are shown in Figs. II-10 and II-11. The prompt fluorescence spectra of the polymer films also are shown for the sake of comparison. In all the polymer films studied, by the doping of NAPH, the Cz chromophorephosphorescence(I_{ph}) decreases significantly and is replaced by a sensitized NAPH phosphorescence(I_{sph}). Although the Cz chromophore-phosphorescence lifetime(τ_{ph}) in PVCz(r and c) films has not been determined because of the overlapping of I_{ph} and I_{sph} , τ_{ph} in PCZEVE films was found, by observing the decay curve at

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444 nm, not to decrease by the doping of NAPH, Therefore, in these films, the triplet energy seems to transfer from the freely migrating triplet exciton to the dopant, and I_{ph} seems to be emitted from the trapped triplet state.



Delayed emission spectra of PVCz(r) films doped with NAPH at 77 K. NAPH concentration; (1) 0, (2) 1.41×10^{-4} , (3) 7.04×10^{-4} , (4) 1.41×10^{-3} , and (5) 7.04×10^{-3} mol/VCz unit mol. The prompt fluorescence (6) of a PVCz(r) film was also given for comparison.

The Cz chromophoredelayed fluorescence (I_{df}) increases with an increase in the concentration of NAPH. The changes in I_{df} anf I_{sph} as a fraction of the NAPH concentration in films are given in Fig. II-12. In the case of the highest concentration of NAPH, the I_{ph} is quenched completely and I_{sph} reaches saturation,



Fig. II-11. Wave number/10³ cm⁻¹ Delayed emission spectra of PCzEVE films doped with NAPH at 77 K. NAPH concentration;
(1) 0, (2) 2.62×10⁻⁴, (3) 1.64×10⁻³, and (4) 8.18 ×10⁻³ mol/CzEVE unit mol. The prompt fluorescence
(5) of a PCzEVE film was also given for comparison.



Fig. II-12 Naphthalene mol/VCz unit mol

Dependence of I_{df} and I_{sph} on concentration of NAPH in solid films at 77 K. (1), (2) —; PVCz(r), (3), (4) ----; PVCz(c), and (5), (6); PCzEVE. whereas I_{df} increases steeply and there was seen a second-order dependence of I_{df} on the exciting light intensity. The lifetime of the delayed fluorescence(τ_{df}) is



Naphthalene mol/VCz unit mol Fig. II-13. Dependence of $\tau_{\rm df}$ on concentration of NAPH in solid films at 77 K. (1) -O-; PVCz(r), (2) -- \Box --; PVCz(c), and (3) $\cdots \bigtriangleup \cdots$; PCzEVE.

gradually lengthened with an increase in the concentration of NAPH, as is shown in Fig. II-13. Similar phenomena are observed for the PVCz(c and r) and PCzEVE films doped with fluorene.

It is clear from the above-mentioned results that the dopants act as an effective triplet quencher for Cz chromophores in the The increase in I_{df} by doping of these triplet films. quenchers, therefore, strongly suggests that a heterogeneous T-T annihilation process occurs with a fairly good efficiency between freely migrating triplet excitons and the triplet state of the dopant.

In the case of PVCz(r and c) doped with NAPH in rigid solution (Fig. II-14), I_{ph} decreases and is replaced by I sph. On the other hand, I_{df} shows an anomalous phenomenon, a little different from that in solid film: it increases at low NAPH concentrations (up to 5 x 10^{-3} M) and decreases at



Fig. II-14. Quencher concentration (I_{ph}°/I_{ph}) and I_{dr}°/I_{dr} for the Stern-Volmer plots (I_{ph}°/I_{ph}) and I_{dr}°/I_{dr} for the PVCz(r) and quenching of the delayed emission of PVCz(r) and PVCz(c) by NAPH and 1,3-PD in MTHF-THF rigid solution at 77 K. (1) $-\triangle$ -; Phosphorescence in the PVCz(r)-1,3-PD system. (2) $- \bigstar$ ---; Delayed fluorescence in the PVCz(r)-1,3-PD system. (3)...O...; Phosphorescence in the PVCz(r)-NAPH system. (4) --@--; Delayed fluorescence in the PVCz(r)-NAPH system. (5) -----; Phosphoresceence in PVCz(c)-1,3-PD system. The point with arrow was corrected by the phosphoroscope factor which was estimated by assuming the lifetime to be 2 ms.

higher concentrations. The change in the $\tau_{\rm df}$ corresponds approximately to the change in the intensity; for example, the $\tau_{\rm df}$ values of PVCz(r) are about 11,12,10, and 3 ms for 0, 8.40 x 10⁻⁴, 2.10 x 10⁻³, and 1.45 x 10⁻³ M, respectively. In the case of PVCz(r and c) doped with NAPH, the heterogeneous T-T annihilation, therefore, seems to be present even in rigid solution at 77 K.

In the case of PVCz(r and c) doped with 1,3-PD, a wellknown triplet quencher in rigid solution, both I_{df} and I_{ph} decrease significantly with an increase in the concentration of 1,3-PD, as is shown in Fig. II-14, where I^0 and I are the Cz chromophore emission intensity in the absence and in the presence of the dopant, respectively. The τ_{df} is decreased steeply $(\tau_{df}^0 / \tau_{df})$ = 4-5 for 0.25 x 10^{-2} M). However, the $\tau_{\rm ph}$ is hardly decreased by the doping of 1,3-PD $(\tau_{ph}^{0}/\tau_{ph} = 1.0-1.1 \text{ for } 2.0 \times 10^{-2} \text{ M}).$ Therefore, it is considered that the triplet energy transfers mainly from freely migrating triplet excitons to the dopant, as is the case of solid films. The degree of quenching is much larger for I_{df} than for I_{ph} . This is the behavior usually observed for poly(1-vinylnaphthalene) and poly(1-naphthyl methacrylate).^{18,19)} When 1,3-PD is used as a triplet quencher, the heterogeneous T-T annihilation is not present

The I_{ph}^0/I_{ph} ratios are given in Fig. II-15 as a function of the NAPH mole ratio in polymer films. Since the density of the vinyl polymers was measured to be 1.19 g/cm³, a conversion of the unit on the abscissa in Fig. II-15 from mole ratio to molarity results in the Stern-Volmer plot:

 $I_{\rm ph}^0/I_{\rm ph} = 1 + k_{\rm t}\tau[Q]$

Here, k_{+} is the bimolecular quenching rate constant, τ is the

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excited state lifetime in the absence of a quencher, and [Q] is the quencher concentration in the molarity unit. In the case of PVCz(r and c) doped with 1,3-PD in rigid solution, Stern-Volmer polts also are obtained, as is shown in Fig. II-14. In the case of





most aromatic vinyl polymers containing the present polymers, the phosphorescence is emitted not from the freely migrating triplet but from the trapped triplet state, so that the lifetime (τ_{ph}) is considered to be that of the trapped triplet state. Two times the τ_{df} value should, therefore, be used as the value of τ_{ph} . The values of k_t were thus obtained from the slopes of the Stern-Volmer polts. Here, k_t is related to the migration coefficient, λ , in the following equation;

 $k_{t} = 4\pi N_{0}\lambda R_{0} \times 10^{-3} M^{-1}s^{-1}$

where N_0 is Avogadro's number and R_0 is the critical transfer distance between donor and acceptor. The value of 15 Å is used for R_0 .²⁸⁾ The migration frequency, k_{mig} , was estimated from the following equations;

 $k_{mig} = 2\lambda/a^2$ (for one-dimensional random walk)

 $k_{mig} = 6\lambda/a^2$ (for three-dimensional random walk)

where a is the interchromophore separation (3.5 Å). From the values of k'_{mig} , the number of Cz chromophores covered by the triplet exciton during its lifetime, n', and the mean exciton migration length, L, were obtained. These values are listed in

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Substance	k _t τ/M ⁻¹	$\tau_{\rm ph}^0/s$	τ_{df}^{0}/ms	$k_t/M^{-1}s^{-1}$	$\lambda/cm^2 s^{-1}$	kij/s ^{b)}	n'	l/Å ^{c)}
PVCz(r)	134	7.9 <u>+</u> 0.7 1.6 <u>+</u> 0.5	50 <u>+</u> 3	1300	1.2×10^{-12}	5.9 x 10^3	590	85
PVCz(c)	74	7.9 <u>+</u> 0.7 1.6 <u>+</u> 0.5	50 <u>+</u> 5	740	6.5×10^{-13}	3.2×10^3	320	63
PCzEVE	57	3.5 <u>+</u> 0.2 0.7 <u>+</u> 0.05	45 <u>+</u> 5	630	5.6 x 10^{-13}			
PVCz(r) ^{a)}	245	8.1 <u>+</u> 0.2	10 <u>+</u> 3	12000	1.1×10^{-11}	1.8×10^4	360	66
PVCz(c) ^{a)}	128	8.1 <u>+</u> 0.2	11 <u>+</u> 2	5800	5.1 x 10^{-12}	8.3 x 10^3	180	47

Table II-3. Triplet Energy Transfer in Film and in Rigid Solution.

a) In rigid solution. b) Three-dimensional random walk in film. c) L = an' $^{1/2}$ (a = 3.5 Å for PVCz). Table II-3.

The values of n' and L for PVCz(r) are higher than those for PVCz(c) both in film and in rigid solution. As described in the previous chapter, the singlet exciton migration is limited by the intrinsic trap sites (excimer-forming sites).^{8,12)} Similarly, the triplet exciton migration seems to be limited by the intrinsic trap sites, such as excimer-forming sites in the polymers, because the observed phosphorescence was emitted only from the trap sites. Therefore, the differences in n' and L between PVCz(r) and PVCz(c) seems to be attributable to the difference in the concentration of the intrinsic trap sites for the triplet exciton. That is, the concentration of the trap sites for a triplet exciton in PVCz(c) films is somewhat larger than that in PVCz(r) films. On the other hand, as has been described in the previous chapter, the concentration of the trap sites for a singlet exciton in PVCz(r) films is nearly equal to that in PVCz(c) films. This suggests that the conformation of the intrinsic trap sites is not necessarily the same between the singlet and triplet excitons. This explanation might be supported by the presence of a conformational difference between the singlet and triplet excimers of 1,3-di(l-naphthyl)propane.³⁰⁾

Although the conformation of the intrinsic trap sites seems to be difference between the singlet and triplet excitons, the number of the Cz chromophores covered by both excitons in the PVCz(r and c) films during the lifetimes ranges from 300 to 600 and the mean exciton migration length ranges from 60 to 80 Å.

As mentioned above, the phosphorescence of PCzEVE films is emitted from the trap sites. Judging from the presence of a shoulder at about 440 nm in the phosphorescence of PCzEVE films

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and the small degree of polymerization ($\overline{DP} = 33$), one of the emitting species seems to be assigned to a monomeric trap such as the Cz chromophore at the chain end. Therefore, the small value of λ in PCzEVE films may be explained by the assumption that a part of Cz chromophores at chain ends acts as an energy trap for a triplet exciton.

Summary

The triplet excitation energy migration in PVCz(r), PVCz(c), and PCzEVE has been investigated by measuring the delayed emission spectra composed of the delayed fluorescence and the phosphorescence in film and in rigid solution at 77 K.

The delayed fluorescence was attributed to a homogeneous T-T annihilation between two migrating triplet excitons. The phosphorescence was emitted from the trap sites. The delayed fluorescence spectra of PVCz(r) and PVCz(c) are different from each other as observed for the prompt fluorescence, because of the different tacticity.

In films of these polymers, the delayed fluorescence increased with an increase in the concentration of a triplet quencher, suggesting the presence of a heterogeneous T-T annihilation. Both the quenching of the Cz chromophore phosphorescence by doping with naphthalene in film and that by doping with 1,3-pentadiene in rigid solution obeyed the Stern-Volmer equation. The migration coefficient was in the following order; PVCz(r) > PVCz(c) > PCzEVE in film and PVCz(r) > PVCz(c) in rigid solution.

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PART III

PHOTO-OXIDATION EFFECT ON THE PHOTOCONDUCTIVITY IN POLY(N-VINYLCARBAZOLE) FILMS

INTRODUCTION

The photo-carrier-generation process in a PVCz film has widely been investigated in the last few years.¹⁻⁴⁾ In the case of the higher π - π * excitation, it is considered that photocarriers are generated from the singlet excited state via the intrinsic charge transfer exciton state.³⁾ In the case of the lower π - π * excitation, the following mechanism is considered to be most appropriate.

$D^* + A \longrightarrow (D^* \cdots A)^* \longrightarrow$	$^{1}\left(D^{+}\cdotsA^{-}\right)^{\ddagger}\overset{\varphi}{\longrightarrow}$	¹ (D ⁺ ···A ⁻) -	$\xrightarrow{\mathrm{E}}$ D ⁺	+ A
encounter complex	non-relaxed exciplex state	geminate ion pair		
	1-\$	·		
D + A + hv < exciplex fluoresce	l (D ⁺ -A ⁻) * exciplex state ence			

The singlet excited state (D^*) of PVCz migrates effectively through the Cz chromophores and encounters some electronaccepting impurity (A) during its lifetime to form an encounter complex $(D^* \cdots A)^*$. The complex goes through a rapid electron transfer and changes to a non-relaxed exciplex state ${}^1(D^+ \cdots A^-)^*$.

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The non-relaxed exciplex state has an excess kinetic energy and undergoes a thermalization of the excess energy to give either an electron-hole ion pair ${}^{1}(D^{+}\cdots A^{-})$ at a separation r_o or a relaxed fluorescent exciplex state ${}^{1}(D^{+}-A^{-})^{*}$. The electron-hole pair will either recombine geminately to give the relaxed fluorescent exciplex state or dissociate into free carriers (D^{+}, A^{-}) under an external electric field E. In this part, in order to confirm the mechanism described above, the following investigations have been done.

As described in the previous part, the triplet exciton has fairly high energy and migrates very efficiently in a polymer film as well as the singlet exciton. The effect of singlet and triplet quenchers on the photoconductivity in PVCz films has been described in chapter 1.

The use of magnetic fields is expected to be an invaluable tool for elucidating the mechanisms of photoconductivity.⁵⁾ Such investigations have been done successfully for molecular crystals such as anthracene and tetracene.⁶⁻⁹⁾ Therefore, it is interesting and practically important to investigate how the magnetic field affects the photoconductivity in a PVCz film. In chapter 2, the magnetic field effects on the photoconductivity in a PVCz film and on the prompt exciplex fluorescence of PVCz films doped with DMTP and N,N,N',N'-tetramethyl-p-phenylenediamine (TMPD) have been described.

In the last chapter, in order to elucidate what kind of compound acts as an electron-accepting impurity, the photooxidation effects on the photoconductivity and the electronic spectra of PVCz films have been investigated.

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Chapter 1

Effect of the Singlet and Triplet Quenchers on the Photoconductivity of Poly(N-vinylcarbazole)

Introduction

The triplet exciton of PVCz has a fairly high energy ($E_T = 3.02 \text{ eV}$; cf. the energy of the singlet exciton, $E_s = 3.55 \text{ eV}$), as compared with other aromatic vinyl polymers, such as poly(l-vinylnaphthalene) ($E_T = 2.5 \text{ eV}$, $E_s = 3.8 \text{ eV}$).¹⁰ Therefore, the photo-carrier-generation via the triplet exciton may be possible in PVCz films. In order to examine this possibility, the effect of the triplet and singlet quenchers on the photoconductivity in PVCz and brominated PVCz (BPVCz) films was examined.

Experimental

The copolymer of VCz with 1-vinylnaphthalene (VN) was prepared by free-radical polymerization. Naphthalene and fluorene were used as triplet quenchers of the Cz chromophore, while anthracene and perylene were used as both singlet and triplet quenchers. Gold-polymer-ness sandwich-type cells were used and the electrical measurements were made by the DC method in a vacuum of 10^{-5} Torr.²)

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Results and Discussion

BPVCz has photoconductive properties similar to those of PVCz.¹¹⁾ The magnitude of the photocurrent of BPVCz is twice as large as that of PVCz in the case of the hole current and is almost the same as that of PVCz in the case of the electron current.

In both PVCz and BPVCz, the triplet quencher does not affect the photoconductive properties at all. Anthracene and perylene acting as both singlet and triplet quenchers significantly reduce the photocurrent in magnitude, but do not affect other photoconductive properties, such as the voltage and temperature dependences of the photocurrent. Table III-1 shows a comparison of the magnitude of the photocurrent among the films undoped and doped with triplet or singlet quenchers. From this table, it is clear that, in the quencher concentrations of 1.5-5 mol%, the singlet quenchers reduce the photocurrent by a factor of ca. 5, while the triplet quenchers do not.

From the results in the previous part, it is clear that the concentration of the freely migrating triplet exciton in the films doped heavily (more than 1 mol%) with the triplet quencher is remarkably reduced as compared with that in the undoped films. The situation is the same as for the case of the singlet quencher. Therefore, the present effect of the singlet and triplet quenchers on the photoconductivity excludes the possibility of the photocarrier generation via the triplet state in these polymers under ordinary experimental conditions.

It is noted that the photocarrier generation via the singlet

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			Number	Magnitudes of the photocurrent ^a)				
		Hole current/10-10 A cm-2		Electron current/10 ⁻¹¹ A cm ⁻²				
Polymer	Dopant	mol %	samples	350 nm	330 nm	350 nm	330 nm	
PVCz	None	0	7	(4.0 ± 0.2)	(3.7 ± 0.3)	(4.3 ± 0.4)	(3.0±0.2)	
	Fluorene	1.5	3	1.1 ± 0.1	1.1 ± 0.1			
	Naphthalene	1.5	3	1.1 ± 0.1	1.2 ± 0.3	$1.1 {\pm} 0.3$	$0.93 {\pm} 0.23$	
	Naphthalene	5.0	3	$0.81 {\pm} 0.08$	$0.99 {\pm} 0.19$			
	Copolymerized	VN ^{b)}	2	1.4 ± 0.1	1.5 ± 0.1	1.5 ± 0.3	1.4 ± 0.2	
	Anthracene	3.0	2	$0.18 {\pm} 0.07$	$0.23 {\pm} 0.09$	$0.09 {\pm} 0.01$	$0.14 {\pm} 0.01$	
	Perylene	2.0	3	$0.20 {\pm} 0.06$	$0.22 {\pm} 0.09$	$0.13 {\pm} 0.05$	0.25 ± 0.08	
VCz–VN (5 mol %) copolymer		0	2	(5.5 ± 0.2)	(5.6 ± 0.1)	(6.3 ± 0.9)	(4.1 ± 0.4)	
	Anthracene	3.0	2	$0.29 {\pm} 0.01$	$0.36 {\pm} 0.01$	$0.29 {\pm} 0.06$	$0.40 {\pm} 0.07$	
BPVCz	None	0	3	(7.2 ± 0.2)	(7.0 ± 0.2)	(3.2 ± 0.2)	(2.6 ± 0.1)	
	Naphthalene	5.0	3	$0.95 {\pm} 0.06$	1.1 ± 0.2	$0.77 {\pm} 0.21$	0.81 ± 0.06	
	Anthracene	3.0	2	0.16 ± 0.03	0.19 ± 0.02	0.37 ± 0.03	0.54 ± 0.06	

Table III-1. COMPARISON OF MAGNITUDES OF THE PHOTOCURRENT¹¹

a) The relative photocurrent values in the doped films taking the photocurrent values in the corresponding undoped films as unity. The values in parentheses are the photocurrent values in the undoped films measured under the condition of 35000 V/cm, 1×10^{13} photons/cm² s, 20 °C, and 1×10^{-5} Torr. b) VCz-VN (5 mol % in feed) copolymer.

excited state is predominant even in BPVCz, which has a low efficiency of the singlet exciton migration. A similar example has been reported of the photoconductivity in tetracene crystals, where the low efficiency of the singlet exciton migration is attributed to the enhanced singlet fission into two triplet excitons. ⁸⁾ In tetracene crystals, the photocarrier generation via the triplet excited state is possible only in the presence of a strong electron acceptor like p-chloranil.

Under the present experimental conditions, the two photon processes are negligible because of the weak light intensity. In the case of the lower singlet $\pi - \pi^*$ excitation, the extrinsic photocarrier generation process with the participation of some electron-accepting impurities must be considered. From the simple energetic calculation, this process is considered to be endothermic in the case of a weak electron acceptor such as some photooxidation products of the Cz chromophore, the most conceivable acceptor. In such an endothermic process, the energy

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difference between the singlet and triplet excitons (0.5 eV) seems to determine the choice between the singlet and triplet channels, although it is much smaller than those of other aromatic vinyl polymers. This is the reason for the lack of the photocarrier generation process via the triplet exciton in these polymers.

Summary

The singlet quenchers significantly reduced the photocurrents of PVCz and BPVCz, but the triplet quenchers did not, regardless of the high efficiency of the polymer-to-quencher triplet energy transfer. This result excluded the possibility of the photocarrier generation via the triplet exciton in these polymers. Since the extrinsic photo-carrier-generation process is considered to be endothermic, the energy difference between the singlet and triplet excitons (0.5 eV) seems to determine the choice between the singlet and triplet channels.

Chapter 2

Magnetic Field Effect on the Photoconductivity of Poly(N-vinylcarbazole)

Introduction

Recently, Groff et al. and Frankevich et al. have, independently, found a new type of magnetically-sensitive process involving an intermediate charge-transfer (exciplex) state formed on the crystal surface, from observations of the magnetic quenching of the dye-sensitized delayed fluorescence in anthracene crystals¹²) and of the prompt exciplex fluorescence in finely-divided ruburene crystals¹³⁾ and in tetracene-adsorbed anthracene crystals, 14) respectively. This new type of magnetically-sensitive process has been considered to be due to a change in the rate of transformation between singlet and triplet states of the exciplex by a weak magnetic field. However, the sign of the change in the transformation rate is in controversy; the sign has been shown to be negative by Groff et al.¹²⁾ and conversely to be positive by Frankevich et al. 13,14) Groff et al. have also suggested that a similar mechanism is responsible for the positive magnetic field effect on the dye-sensitized photoconductivity in anthracene crystals.^{12,15)} However, details concerning this positive magnetic field effect on the photoconductivity has never been reported. Frankevich et al. have also suggested that a similar mechanism might be appropriate to the positive magnetic field effect which had been reported

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previously by these authors,¹⁶⁾ on the surface photoconductivity in anthracene crystals and on the low-temperature photoconductivity in a sandwitch-type cell of a sublimed tetracene film.¹⁴⁾ In these cases, the situation is, however, considered to be complicated, because of the possibility that the singlet fission may partly be responsible for this positive magnetic field effect. It has been confirmed by the observation of a small quenching of the prompt fluorescence by a magnetic field that the singlet fission is present both in anthracene crystals at 300 K and in tetracene crystals at 77 K.

In this chapter, the author examined magnetic field effects on the photoconductivity and on exciplex fluorescence in PVCz films, where the singlet fission is completely impossible energetically ($E_S = 3.55 \text{ eV}$, $E_{\pi} = 3.02 \text{ eV}$).

Experimental

An gold-PVCz-ness sandwich-type cell was set in a cryostat, mounted between the poles of an electromagnet, and excited by monochromatized light from a stabilized 500 W xenon lamp. The photocurrent was measured with a vibrating reed electrometer. The emission passed to a 1P28 photomultiplier through a light guide and appropriate filters. The photomultiplier current was measured with a differential electrometer. The current at zero magnetic field was compensated and only the variation of the current caused by magnetic field was measured. Small current changes up to 0.2 % could be detected by means of this measuring system. The results to be presented below were obtained for the arrangement of the magnetic field vector perpendicular to

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the electric field vector.

Results

(1) Photocurrent.

When a magnetic field H of up to 4000 G is applied to a PVCz film, the photocurrent, i, in the wavelength region shorter than 400 nm is increased by a quantity, Δi , although there is no change both in the dark current and in the photocurrent in the wavelength region longer than 400 nm. In all cases, $\Delta i/i$ increases monotonically with H and reaches saturation at about 500-1000 G, as is shown in Fig. III-1. The $\Delta i/i$ reaches a half value of the saturated one at 200-300 G. As is shown in Fig. III-2, $\Delta i/i$ increases with applied voltage, V, and reaches saturation. Judging from the i-V relationship² this means that the relation $i \propto V^n$ (n = 1.5 - 2.0) is valid over the whole range up to 100 V (10⁵ V/cm) for the hole current and in the low applied voltage range up to 40 V (3.3 x 10⁴ V/cm) for the electron current.



Fig. III-1.

Magnetic field dependence of the relative increase in the photocurrent under 35000 V/cm at 20°C. Undoped PVCz film: (1) 350 nm, hole current; (2) 260 nm, hole current; (3) 350 nm, electron current. PVCz film doped with DMTP (4 mol%): (4) 350 nm, hole current; (5) 350 nm, electron current.



Fig. III-2.

Applied voltage dependence of the relative increase in the photocurrent at 1500 G and 20°C. Undoped PVCz film: (1) 350 nm, hole current; (2) 340 nm, electron current. PVCz film doped with DMTP (2 mol%): (3) 350 nm, hole current; (4) 340 nm, electron current. The $\Delta i/i$ hardly depends on light intensity, L, in the range below about 2 x 10^{13} photons/cm²s ; the relation $i \ge L^1$ is valid. On the other hand, $\Delta i/i$ decreases with increasing L in the range above 2 x 10^{13} photons/cm²s with an exception of the case of illumination of the weakly-absorbed light (370-400 nm). In the case of illumination of the 370-400 nm light, $\Delta i/i$ does not depend on L over the whole range up to 5 x 10^{14} photons/cm²s.

Spectral dependence of $\Delta i/i$ is shown in Fig. III-3. In both cases of the hole and electron currents, $\Delta i/i$ is comparatively large in the ${}^{1}L_{a}$ and ${}^{1}L_{b}$ absorption regions (or the higher energy region above 4.6 eV). The $\Delta i/i$ is larger for the hole current (5--6 %) than for the electron current (3.5-4 %).





Spectral dependence of the relative increase in the photocurrent at 1500 G, 33000 V/cm, and 20°C. Undoped PVCz film: (1) hole current; (2) electron current. PVCz film doped with DMTP (2 mol%): (3) hole current; (4) electron current. The UV absorption spectrum of a PVCz film about 1 μ thick is shown by a dotted line (5).

The $\Delta i/i$ depends considerably on temperature. The positive magnetic field effect on the photoconductivity disappears below -30 °C. With increasing temperature above -30 °C, $\Delta i/i$ first increases sharply and then decreases gradually.

The DMTP-doped films show similar results to the undoped films mentioned above, with an exception that the magnitude of $\Delta i/i$ is enhanced by a factor of two with doped DMTP, as is shown in Figs. III-1 - -3.

(2) Exciplex Fluorescence.

The prompt excimer fluorescence of undoped PVCz films was not affected by magnetic field up to 4000 G at all. However, the prompt exciplex fluorescences of PVCz films doped with DMTP and TMPD were enhanced in intensity by magnetic fields. The ratio of this increment to the total prompt fluorescence intensity ($\Delta F/F$) increased monotonically with an increase in magnetic fields, H, and reached a saturation at about 400 G, as is shown in Fig. III-4. The $\Delta F/F$ reached a half value of the saturation at about 100 G. The data presented below were obtained for the saturated values.

The $\Delta F/F$ was independent both of the wavelength (280-360 nm) and of the intensity $(10^{13-14} \text{ photons/cm}^2 \text{s})$ of the exciting light. The $\Delta F/F$ apparently depended on the emission-wavelength, as is shown in Fig. III-5. In both doped systems, the $\Delta F/F$ for 400 nm was about 0.6 times as large as for 450-600 nm. This is because the excimer fluorescence is superimposed considerably on the exciplex fluorescence in the wavelength region shorter

in the PVCz film doped with 3.3 mol% DMTP, the ratio of the exciplex fluorescence to the total one was about 0.55 at 400 nm, and then the ratio of ΔF to the exciplex fluorescence intensity was evaluated to be about 2.1 % at 400 nm, being equal to the $\Delta F/F$ value for

than 400 nm. For instance,






450-600 nm (1.9 %) within the experimental error. Therefore, the magnitude of the magnetic field effect on the exciplex fluorescence is safely considered to be independent of the emission-wavelength; it is about 2.0 % and 1.1 % at 25 °C for the DMTP and TMPD doped systems, respectively.

The $\Delta F/F$ depended slightly on temperature, as is shown in Fig. III-6. With an increase in temperature, the F value decreased slightly and $\Delta F/F$ increased, resulting in an activation energy of about 0.04 eV.

The electric field quenching of the exciplex fluorescence of PVCz films doped with DMTP, which has been reported by Yokoyama et al.,⁴⁾ also was observed in the present study. However, the ratio of Δ F/F due to the magnetic field was independent of applied electric fields up to \pm 35000 V/cm, suggesting that magnetic and electric fields affect independently the exciplex fluorescence.

The exciplex fluorescence in solution has not been affected by magnetic fields up to 4000 G at all.

Discussion

The positive magnetic field effect on the photoconductivity in the undoped and DMTP-doped PVCz films is observed only for the wavelength region shorter than 400 nm and is not observed for the dark current and for the photocurrent in the wavelength region longer than 400 nm. This effect is, therefore, attributable to a carrier generation process rather than to a carrier migration process.

Doping of DMTP enhances the magnitude of the photocurrent

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by a factor of 1.5-3 in the $\pi-\pi^*$ absorption region of PVCz, because of additional carrier generation process via the nonrelaxed exciplex state.^{2,17)} The enhancement of $\Delta i/i$ with doped DMTP, therefore, clearly means that the efficiency of this carrier generation process is increased by a relatively weak magnetic field. This is also reasonable, judging from the fact that $\Delta i/i$ is very sensitive both to applied electric fields and to temperature as mentioned above.

The increase in both efficiency of the carrier generation and of the exciplex fluorescence is explained by means of a reduction of the rate of transformation between singlet and triplet states of the geminate ion pair $^{1}(D^{+}\cdots A^{-})$ by a weak magnetic field, which has been named magnetic hyperfine modulation, 12 provided that carriers are generated more efficiently via the singlet state than via the triplet state.

The ϕ has roughly been estimated to be 0.2 from the fact that the magnetic field effect on the prompt exciplex fluorescence (2%) is smaller than that on the photoconductivity (10%).

It is very interesting that a similar magnetic field effect on the photoconductivity is observed also for undoped PVCz films. This clearly suggests that a similar carrier generation process via an non-relaxed exciplex state plays an important role in photocarrier generation even in the undoped films.

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Summary

Magnetic fields were found to increase the photocurrent in PVCz films by 5-6 % at 1000 G. This positive magnetic field effect was sensitive both to applied voltage and to temperature, and was enhanced with DMTP doping by a factor of two. Magnetic fields were also found to increase the prompt exciplex fluorescence of PVCz films doped with DMTP and TMPD by 2 % at 500 G. This positive magnetic field effect was explained by means of a reduction of the rate of transformation between the singlet and triplet states of the geminate ion pair by a weak magnetic field, provided that carriers are generated more effectively via the singlet state than the triplet state. These results confirm that a carrier generation process with the participation of an non-relaxed exciplex state has an important role in photo-carrier-generation even in undoped PVCz films.

Chapter 3

Photo-oxidation Effect on the Electronic Spectra and the Photoconductivity of Poly(N-vinylcarbazole)

Introduction

In the carrier generation mechanism described in the previous chapter, the energy required to prepare an electron-hole ion pair at a separation r_0 is expressed by $(I_c - E_{ac} - e^2/\epsilon r_0)$, where I_c , E_{ac} , and ϵ are the ionization potential (5.9-6.1 eV), the electron affinity (1.5 eV), and the dielectric constant (3.0) of PVCz in film, respectively. The value for $r_0 = 25$ Å is estimated to be 4.2-4.4 eV, which is higher than the energy of the lowest excited singlet state (3.6 eV). Therefore, a compound having the electron affinity of 0.6-0.8 eV or above can act as an electron acceptor (A).

Some unknown materials contaminated in course of synthesis of monomers and/or polymers, dissolved oxygen, or photo-oxidation products of the polymer can be considered as the electronaccepting impurities in PVCz films. Judging from the migration efficiency of the singlet exciton, the concentration of electronaccepting impurities of about 10^{-3} mol/mol monomer unit seems to be necessary to interpret a high yield of photo-carrier-generation (ca. 0.1). Contamination by such a concentration of impurities is improbable in the course of the synthesis. In the case of contamination of triplet oxygen in the ground state, whose electron affinity is ca. 0.42 eV,¹⁸⁾ considerable thermal activation should be necessary to give an electron-hole ion pair.

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Therefore, it is impossible to explain the high yield of the photo-carrier-generation in PVCz films by considering only unknown contaminating impurities and oxygen. Any photo-oxidation products of PVCz should be considered as electron-accepting impurities which may be effective for the photo-carrier-generation. Therefore, detailed information on the photo-oxidation of PVCz films is very important in understanding the origin of the high photoconductive properties of polymers. It has briefly been reported that photo-oxidation products were obtained by irradiation of light on PVCz suspended in a sulfuric acid - ethanol mixture under a stream of air¹⁹ and of PVCz films in air.³ However, no detailed study has been reported on the photo-oxidation of PVCz films.

In this chapter, the author described the photo-oxidation of PVCz films and DCzP(a) films, by measuring the change in the absorption and fluorescence spectra caused by UV-irradiation. It has also been investigated how the photo-oxidation affects the photoconductivity in PVCz films.

Experimental

For photo-oxidation, films were irradiated in air under atmospheric pressure through a water filter (21 cm) with the full spectrum of a 500 W high pressure mercury lamp. The radiation density ($\lambda < 350$ nm) was about 1 x 10¹⁷ photons/cm²s.

Results and Discussion

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(1) Electronic Spectra.

Figure III-7 shows the change in the absorption spectra of PVCz films caused by UV-irradiation. With irradiation time, the absorption intensity of Cz chromophores decreases and new absorption (300-340 nm and 350-420 nm) appears and increases. Three isosbestic points are observed at 248, 341, and 303 nm. These new bands are attributed to photo-oxidation products, because no change in the absorption was observed for a PVCz film irradiated under a vacuum of 10^{-1} Torr. Further irradiation causes a deviation from these isosbestic points and also a decrease of the absorption intensities of the photo-oxidation product, as is shown in Fig. III-8. Similar phenomena were observed also for DCzP(a) films. The dependence of the absorption intensity (370 nm) due to the photo-oxidation product on irradiation time is shown in Fig. III-8. Assuming that there is no





Change in the absorption spectra of a PVCz film caused by photoirradiation. Irradiation time; (1) 0, (2) 6, (3) 12, (4) 18, and (5) 25 min.



Fig. III-8. Irradiation time/min



remarkable difference in the molar extinction coefficient of the photo-oxidation product between PVCz and DCzP(a), Fig. III-8 shows that the formation rate of the photo-oxidation product observed for PVCz films is larger than that for DCzP(a) films by a factor of ca.4. This may be attributed to the presence of the sites which are susceptible to photo-oxidation in PVCz films. Judging from the fact that the concentration of the sandwich-like excimer-forming site in PVCz films is higher than that in DCzP(a) films, the sandwich-like excimer-forming site seems to be susceptible. This is supported by the fact that the decrease of the sandwich-like excimer fluorescence (ca, 23900 cm⁻¹) caused by photo-oxidation is larger than that of the second excimer fluorescence (ca. 26500 cm⁻¹), as is clearly seen in Fig. III-10.

IR spectra of photo-oxidized PVCz films are shown in Fig. III-9. The increasing absorption observed at around 1700 cm⁻¹ is attributable to carbonyl groups. A similar phenomenon was observed also for DCzP(a) films. These results show that the photo-oxidation product formed in both PVCz and DCzP(a) films has carbonyl group, which is an electron-accepting substituent. The formation of the carbonyl group has been reported also in photo-oxidized polystyrene films.²⁰⁾

The change in the fluorescence spectra of PVCz films caused by photoirradiation in air is shown in Fig. III-10. Little change was observed in the fluorescence spectra of PVCz films by



Fig. III-9. Wave number/cm⁻¹

Change in the infrared absorption spectra of a PVCz film caused by photoirradiation. Irradiation time; (1) 0, (2) 20, and (3) 100 min.

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photoirradiation under a vacuum of 10^{-1} Torr. Remarkable change in the fluorescence spectra is observed for very slightly photo-oxidized PVCz films, compared with the case of the absorption spectra. With irradiation time, the excimer fluorescence of PVCz films decreases and a new fluorescence band in the longer-wavelength region ($v < 20000 \text{ cm}^{-1}$) appears. Further irradiation causes a decrease of both the excimer and the new fluorescence intensities. A quite similar phenomenon was observed for DCzP(a) films, as is shown in Fig. III-11. The broad and structureless emission band with a peak (or shoulder) at $19000-20000 \text{ cm}^{-1}$ of the slightly photo-oxidized PVCz and DCzP(a) films may be assigned to the exciplex fluorescence, which is formed between an excited Cz chromophore and the photooxidation product with carbonyl group. This assignment is supported by the following observation. The fluorescence observed at 515 nm for a slightly photo-oxidized PVCz film





Fig. III-11. wave number 10° cm⁻¹ Change in the fluorescence spectra of DCzP(a) films caused by photoirradiation. Irradiation time; (1) 0, (2) 0.17, (3) 0.5, (4) 1, (5) 1.5, and (6) 10 min.

Fig. III-10. Wave number/10³ cm⁻¹

Change in the fluorescence spectra of PVCz films caused by photoirradiation. Irradiation time; (1) 0, (2) 0.5, (3) 1, (4) 1.5, (5) 2.5, (6) 4.5, (7) 9, and (8) 15 min.

is partly quenched by applying an electric field and the decrease in the fluorescence shows a quadratic dependence on the field strength, as is shown in Fig. III-12. A similar phenomenon has been observed for the exciplex fluorescence of PVCz films doped with DMTP.⁴

A quenching factor, Q, is defined for convenience by the following equation;

 $Q_t = (I_0 - I_t) / I_t$

where I_o and I_t are fluorescence peak intensities before and after photo-irradiation, respectively. The dependence of the quenching factor of photo-oxidized PVCz and DCzP(a) films on the irradiation time is shown in Fig. III-13. The value of Q obtained for the thin films is larger than that obtained for the thick films. This result suggests that the process of the photooxidation proceeds near the surface of films. Therefore, the photo-oxidation product is not formed in the bulk of film uniformly. However, an approximately linear relationship exists





Electric field-strength dependence of the fluorescence decrease observed at 510 nm for the weakly Quence photo-oxidized PVCz film ($\approx 0.5 \,\mu$ m thick). Irradiation time; 25 s. Excitation wavelength; 335 nm. thick), (3)





Quenching factor vs. irradiation time. (1) PVCz film ($\approx 0.7 \,\mu$ m thick), (2) PVCz film ($\approx 0.2 \,\mu$ m thick), (3) DCzP(a) film ($\approx 0.3 \,\mu$ m thick), and (4) DCzP(a) film ($\approx 0.2 \,\mu$ m thick). Excitation wavelength; 330 nm. between Q and the irradiation time, t, in an early stage of photo-irradiation, suggesting that the photo-oxidation product acts as an exciton trap in competition with the excimer-forming sites. The value of Q obtained for DCzP(a) films is larger than that obtained for PVCz films. Considering that the concentration of the photo-oxidation product formed in PVCz films is higher than that formed in DCzP(a) films, this fact suggests that the number of Cz chromophores covered by a singlet exciton during its lifetime in a DCzP(a) film is larger than that in a PVCz film. This fact is consistent with the result obtained by the investigation of the excitation energy migration in chapter l of Part II.

The host fluorescence was depressed to half its original intensity at a guest concentration equal to the concentration of the excimer-forming sites, in the case of PVCz films. This concentration in a PVCz(r) film was ca. 3×10^{-3} mol/mol basic unit. This concentration level of the photo-oxidation product is reached by irradiating a thin PVCz film for one minute, as is seen in Fig. III-13. Considering the number of absorbed photons, the yield of the formation of the photo-oxidation oxidation product is estimated to be ca. 10^{-4} .

(2) Photoconductivity.

Dependence of the photocurrent in the photo-oxidized PVCz films on irradiation time is shown in Fig. III-14. The photocurrent observed in the π - π ^{*} absorption region increases steeply, by a factor of about 15, with photo-oxidation. It decreases when the film was exposed to radiation until a change in the absorption spectra by photo-oxidation could be observed.

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Voltage, spectral, light-intensity, and temperature dependences of the photocurrent in the photo-oxidized PVCz film were almost the same as those in the unirradiated PVCz film.¹⁷⁾

Figure III-15 shows the dependence of the positive magnetic field effect in the photo-oxidized PVCz films on irradiation time. The effect is enhanced by a factor of about 3 by slight photo-oxidation, as observed for the DMTP-doped system.

It is clear from these results that the yield of photocarrier-generation in the slightly photo-oxidized PVCz films is much higher than the yield in the unirradiated films. That is, even if the concentration of the photo-oxidation product is too low to be detected in the absorption spectra, the product acts effectively as an electron-accepting impurity (A) in the photocarrier-generation mechanism described in the previous chapter, to enhance effectively the yield of the photo-carrier-generation process in a PVCz film.





(1) 400, (2) 360, and (3) 330 nm.





Dependence of the positive magnetic field effect in a photo-oxidized PVCz film on irradiation time. Magnetic field; 1000 G. Applied voltage; 35000 V/cm. (1) 400, (2) 360, and (3) 330 nm.

Summary

When PVCz and DCzP(a) films were irradiated by UV light in air, a photo-oxidation product with carbonyl group was formed near the surface of the films with a yield of 10^{-4} . This photooxidation product acted as a singlet exciton trap and formed an exciplex with an excited Cz chromophore. A very slight amount of the photo-oxidation product, whose concentration is too low to be detected in the absorption spectra, enhanced the photoconductivity in PVCz films by a factor of ca. 15 and the positive magnetic field effect by a factor of ca. 3. These results suggest that the product acts as an electron-accepting impurity in the photo-carrier-generation mechanism in the lowest $\pi-\pi^*$ absorption region of PVCz.

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CONCLUSION

The purpose of this thesis is to present information with regard to the photophysical process of aromatic vinyl polymers and to clarify the mechanism of photoconductivity of the polymers on the basis of this information. The following conclusions were obtained in this thesis.

1) It was revealed by the NMR spectroscopy and others that the tacticity of PVCz, P5VBCz, and P7VBCz varies depending on the polymerization method. The cationic polymers were rich in isotacticity and the radical polymers were rich in syndiotacticity.

2) The polymers composed of both the isotactic and the syndiotactic sequences (PVCz, P7VBCz, and P5VBCz(r)) showed one broad emission band composed of the second and the sandwich-like excimer fluorescences. The intensity of the second excimer fluorescence increased with an increase in the content of the syndiotactic sequence. The intensity of the sandwich-like excimer fluorescence was independent of the tacticity. The polymer composed of only the isotactic sequence (P5VBCz(c)) showed the monomeric and the sandwich-like excimer fluorescences. The second excimer fluorescence was emitted from only the syndiotactic sequence. The structure of the second excimer was suggested that only one benzene ring in each neighboring Cz chromophores overlaps with the other one.

3) The parameters of the photophysical process for PVCz(r) were evaluated by the quenching experiments. The parameters indicate that the sandwich-like excimer is formed easily from

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the second excimer.

4) PVPh, PPhEVE, and PPhMA showed the structured monomeric fluorescence and no excimer one. The fluorescence properties of these polymers indicate the formation of a non-fluorescent excimer in PVPh. The excimer was easily formed in the vinyl polymers having aromatic chromophores connected directly to the skeletal chain, even if the aromatic chromophores did not form any excimer between those.

5) PCzEVE showed an anomalous and structureless fluorescence composed of two components. One of the emitting species may be the second excimer.

6) The efficiency of the singlet exciton migration in polymer films was determined by the concentration of the intrinsic trap sites such as the excimer-forming sites. It decreases in the following order: DCzP(a) > PCzEVE > PVCz(r) ~ PVCz(c) > PACz ~ BPVCz.

7) The concentration of the second excimer-forming site, c_s , was higher in the syndiotactic rich polymer than in the isotactic rich polymer. As the concentration of the sandwich-like excimer-forming site, c_E , was much higher than c_s , there was little difference in $c_F + c_s$ between PVCz(r) and PVCz(c).

8) The efficiency of the triplet exciton migration also was determined by the concentration of the intrinsic trap sites. The decreasing order is; PVCz(r) > PVCz(c) > PCzEVE in film, and PVCz(r) > PVCz(c) in rigid solution.

9) The delayed fluorescence increased with an increase in the concentration of triplet quenchers such as naphthalene and fluorene, suggesting the presence of a heterogeneous T-T annihilation.

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10) The singlet quenchers significantly reduced the photocurrent of PVCz and BPVCz, but the triplet ones did not, indicating that charge carriers are generated via the singlet exciton rather than the triplet one.

11) The study of magnetic field effect on the photoconductivity and the exciplex fluorescence of PVCz revealed that the photo-carriers are generated by field assisted dissociation of the singlet non-relaxed exciplex.

12) UV-irradiation on PVCz films yielded the photo-oxidation product with carbonyl group near the surface of the film with a yield of 10^{-4} . This product acts as an effective electron-acceptor and enhances the photo-carrier-generation.

As mentioned above, many information were obtained on the photophysical process of PVCz and its related compounds. The author believes that this thesis would contribute to advanced understanding of the optical and electrical properties of organic materials.

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