

OFFICE OF NAVAL RESEARCH

Contract N00014-84-G-0201

Task No. 0051-865

Technical Report #39

Molecular Orientation of the Silver Tetraneopentoxyphthalocyanine and Stearic Acid Mixed Langmuir-Blodgett Film and Its Chemical and Electrochemical Behaviour

Ву

Yansong Fu and A.B.P. Lever*

in

Journal of Physical Chemistry



York University
Department of Chemistry, 4700 Keele St., North York
Ontario, Canada M3J 1P3

Reproduction in whole, or in part, is permitted for any purpose of the United States Government

*This document has been approved for public release and sale; its distribution is unlimited

*This statement should also appear in icem 10 of the Document Control Data-DD form 1473. Copies of the form available from cognizant contract administrator

91 4 17 094

1a. REPORT SECURITY CLASSIFICATION		MENTATION I	PAGE			
Za. SECURITY CLASSIFICATION AUTHORITY		TO RESTRICTIVE	MARKINGS			
		3. DISTRIBUTION / AVAILABILITY OF REPORT				
Unclassified 2b. DECLASS:FICATION / DOWNGRADING SCHEDUL	Ē	As it appears on the report 5. MONITORING ORGANIZATION REPORT NUMBER(5)				
4. PERFORMING ORGANIZATION REPORT NUMBER	(S)					
Report # 39						
NAME OF PERFORMING ORGANIZATION 66. OFFICE SYMBOL		7a. NAME OF MO	NITORING ORGA	NIZATION	 	
A.B.P. Lever, York University	(If applicable)	Office of Naval Research				
Chemistry Department & ADDRESS (Gty, State, and ZIP Code)		7b. ADDRESS (City	. 54344 344 710	Codel		
4700 Keele St., North York, Ont	ario M3J 1P3		Division	Code)		
Canada			incy Street			
			, VA 22217			
OE: TEATURE OF TOTAL AND A STREET	8b. OFFICE SYMBOL (If applicable)	9. PROCUREMENT		ENTIFICATION	NUMBER	
ORGANIZATION	/ abbama/	N00014-84	-G-0201			
8c. ADDRESS (City, State, and ZIP Code)		10 SOURCE OF F	UNDING NUMBE	२९	•	
		PROGRAM ELEMENT NO.	PROJECT	TASK	WORK UNIT	
		ELEMENT NO.	NO.	NO.	ACCESSION NO	
Molecular Orientation of the S Mixed Langmuir-Blodgett Film a	Silver Tetraneo and Its Chemica	pentoxyphthal 1 and Electro	locyanine at ochemical Be	nd Steario havior	e Acid	
12. PERSONAL AUTHOR(S)	* * * * * * * * * * * * * * * * * * *					
	A.B.P. Lever	14. DATE OF REPO	ST /You March	Om. 115 24	GE COUNT	
13a. TYPE OF REPORT 13b. TIME CO Technical FROM:Aug.	. '90 to Aug. '91	April 4,	.991	Day) IIS. FA	30	
16. SUPPLEMENTARY NOTATION						
17 COSATI CODES	18. SUBJECT TERMS (
55.5	Phthalocyan		-Blodgett.#	Flactrock		
FIELD GROUP SUB-GROUP 5	p-Fillenaiocyan	ine, Langmuii		LIECTION	iemistry,	
HELD GROUP SUR-GROUP G	Polarizatio	ine, Langmuli n,≁Thin FilmS	4	Diectioci	emistry,	
	Polarizatio	n,≯Thin Films	4	Liection	emistry,	
19. ABSTRACT (Continue on reverse if necessary	Polarization	n, Thin Films	4			
19. ABSTRACT (Continue on reverse if necessary a	Polarization and identify by block ms comprised of	n, Thin Films number) various rel	ative conce	ntrations	of	
Mixed Langmuir-blodgett fil	Polarization and identify by block ms comprised of locyanine (AgT)	n, Thin Films number) various rel NPC) and stea	ative conce	ntrations e reporte	of	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po	Polarization ms comprised of locyanine (AgT) larisation data gTNPc domains	ry Thin Films Townser Various rel NPC) and stea I are discuss In the film.	ative conce ric acid ar ed in terms Electroche	ntrations e reporte of the r mical dat	of d. elative a	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated Aconfirm the oxidation of Ag	Polarization ms comprised of locyanine (AgTM larisation data gTNPc domains in IITNPc(-2) succession.	ry Thin Films Thin Various related are discussion the film.	ative conce ric acid ar ed in terms Electroche [Ag IIITNPc(ntrations e reporte of the r mical dat -2)], and	of d. elative a	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ , in the fi	Polarization and identify by block ms comprised of locyanine (AgTN larisation data gTNPc domains if ITNPc(-2) success lm phase withou	ry Thin Films Townser)	ative conce ric acid ar ed in terms Electroche [AgIIITNPc(ientation.	ntrations e reporte of the r mical dat -2)] and The film	of d. elative a was	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the file also subject to chemical ox	Polarization ms comprised of locyanine (AgTM larisation data gTNPc domains if ITNPc(-2) succession by the succession of the phase without idation and recommend to the succession of the succes	ry Thin Films Townser)	ative conce ric acid ar ed in terms Electroche [AgIIITNPc(ientation. he degree o	ntrations e reporte of the r mical dat -2)] and The film f organis	of d. elative a was ation	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the fill also subject to chemical ox	Polarization ms comprised of locyanine (AgTM larisation data gTNPc domains if ITNPc(-2) succe lm phase without idation and rece discussed. As	ry Thin Films rumoer) various rel NPC) and stea are discuss in the film. cessively to it loss of or duction and t	ative conce ric acid ar ed in terms Electroche [AgIIITNPc(ientation. he degree o	ntrations e reporte of the r mical dat -2)] and The film f organise even when	of d. elative a was ation diluted	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the fi also subject to chemical ox following such chemistry is with stearic acid. do not d	Polarization and identify by block ms comprised of locyanine (AgTh larisation data gTNPc domains if ITNPc(-2) succe lm phase without idation and rec discussed. Ag isplay resolved	ry Thin Films rumoer) real various related are discussion the film. ressively to a loss of or duction and to sell the	ative conceric acid ared in terms Electroche [AgIIITNPc(ientation.he degree oggregates, but ternar	ntrations e reporte of the r mical dat -2)] and The film f organise even when	of d. elative a was ation diluted	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the fill also subject to chemical ox following such chemistry is	Polarization and identify by block ms comprised of locyanine (AgTh larisation data gTNPc domains if ITNPc(-2) succe lm phase without idation and rec discussed. Ag isplay resolved	ry Thin Films rumoer) real various related are discussion the film. ressively to a loss of or duction and to sell the	ative conceric acid ared in terms Electroche [AgIIITNPc(ientation.he degree oggregates, but ternar	ntrations e reporte of the r mical dat -2)] and The film f organise even when	of d. elative a was ation diluted	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the fi also subject to chemical ox following such chemistry is with stearic acid. do not d	Polarization and identify by block ms comprised of locyanine (AgTh larisation data gTNPc domains if ITNPc(-2) succe lm phase withou idation and rec discussed. Ag isplay resolved stearic acid do	rumoer) ? various rel ¡Pc) and stea a are discuss in the film. cessively to it loss of or duction and t ¡IITNPc(-2) a i ESR spectra o exhibit suc	ative conceric acid ared in terms Electroche [Ag III TNPc(ientation.he degree of gregates, but ternarh resolved]	ntrations e reporte of the r mical dat -2)] + and The film f organis even when y film mi spectra.	of d. elative a was ation diluted	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the fi also subject to chemical ox following such chemistry is with stearic acid, do not d of AgIITNPc(-2)/H ₂ TNPc(-2)/	Polarization and identify by block ms comprised of locyanine (AgTh larisation data gTNPc domains if ITNPc(-2) succe lm phase withou idation and rec discussed. Ag isplay resolved stearic acid do	rumber) Thin Films Thin Films Various rel NPC) and stea are discuss in the film. Sessively to it loss of or duction and t SITNPC(-2) a it ESR spectra be exhibit suc	ative conceric acid ared in terms Electroche [AgIIITNPc(ientation.he degree of gregates, but ternarh resolved]	ntrations e reporte of the r mical dat -2)]+ and The film f organis even when y film mi spectra.	of d. elative a was ation diluted xtures	
Mixed Langmuir-blodgett fill silver tetraneopentoxyphtha Electronic spectroscopic po orientation of aggregated A confirm the oxidation of Ag [AgIIITNPc(-1)] ²⁺ in the fi also subject to chemical ox following such chemistry is with stearic acid, do not d of AgIITNPc(-2)/H ₂ TNPc(-2)/	Polarization and identify by block ms comprised of locyanine (AgTh larisation data gTNPc domains if ITNPc(-2) succe lm phase withou idation and rec discussed. Ag isplay resolved stearic acid do	rumoer) ? various rel ¡Pc) and stea a are discuss in the film. cessively to it loss of or duction and t ¡IITNPc(-2) a i ESR spectra o exhibit suc	ative conceric acid ared in terms Electroche [AgIIITNPc(ientation.he degree of gregates, but ternarh resolved]	ntrations e reporte of the r mical dat -2)]+ and The film f organis even when y film mi spectra.	of d. elative a was ation diluted xtures	

TECHNICAL REPORT DISTRIBUTION LIST - GENERAL

Office of Naval Research (2) Chemistry Division, Code 1113 800 North Quincy Street Arlington, Virginia 22217-5000

Commanding Officer (1)
Naval Weapons Support Center
Dr. Bernard E. Douda
Crane, Indiana 47522-5050

Dr. Richard W. Drisko (1)
Naval Civil Engineering
Laboratory
Code L52
Port Hueneme, CA 93043

David Taylor Research Center (1)
Dr. Eugene C. Fischer
Annapolis, MD 21402-5067

Dr. James S. Murday (1) Chemistry Division, Code 6100 Naval Research Laboratory Washington, D.C. 20375-5000 Dr. Robert Green, Director (1) Chemistry Division, Code 385 Naval Weapons Center China Lake, CA 93555-6001

Chief of Naval Research (1)
Special Assistant for Marine
Corps Matters
Code 00MC
800 North Quincy Street
Arlington, VA 22217-5000

Dr. Bernadette Eichinger (1)
Naval Ship Systems Engineering
Station
Code 053
Philadelphia Naval Base
Philadelphia, PA 19112

Dr. Sachio Yamamoto (1) Naval Ocean Systems Center Code 52 San Diego, CA 92152-5000

Dr. Harold H. Singerman (1)
David Taylor Research Center
Code 283
Annapolis, MD 21402-5067

Defense Technical Information Center (2) Building 5, Cameron Station Alexandria, VA 22314

ONR Electrochemical Sciences Program Robert J. Nowak, Program Manager

Professor Hector Abruña Department of Chemistry Cornell University Ithaca, NY 14853 413d018

Professor Allen Bard Department of Chemistry The University of Texas at Austin Austin, TX 78712-1167 413a002

Professor James Brophy Department of Physics University of Utah Salt Lake City, UT 84112 413d015

Professor Bruce Dunn Departement of Materials Science and Engineering University of California, Los Angeles Pennsylvania State University Los Angeles, CA 90024 413d011

Professor Gregory Farrington Laboratory for Research on the Structure of Matter 3231 Walnut Street Philadelphia, PA 19104-6202 413d003

Professor Martha Greenblatt Department of Chemistry Rutgers University Piscataway, NJ 08854 413d008

Professor Adam Heller Department of Chemical Engineering University of Texas at Austin Austin, TX 78712-1062 413h007

Professor C. A. Angell Arizona State University Department of Chemistry Tempe, AZ 85287 413d007

Professor Lesser Blum Department of Physics University of Puerto Rico Rio Piedras, PUERTO RICO 00931 4133002

Professor Daniel Buttry Department of Chemistry University of Wyoming Laramie, WY 82071 4133019

Professor Andrew Ewing Department of Chemistry 152 Davey Laboratory University Park, PA 16802 4133030

Professor W. R. Fawcett Department of Chemistry University of California, Davi Davis, CA 95616 4133020

Professor Joel Harris Department of Chemistry University of Utah Salt Lake City, UT 84112 413a005

Professor Pat Hendra The University Southampton SO9 5NH ENGLAND 4134001

ONR Electrochemical Sciences Program Robert J. Nowak, Program Manager

Professor Joseph Hupp Department of Chemistry Northwestern University Evanston, IL 60208 4133025

Professor A. B. P. Lever Department of Chemistry York University 4700 Keele Street North York, Ontario M3J 1P3 4131025

Professor Rudolph Marcus Division of Chemistry and Chemical Engineering California Institute of Technology Pasadena, CA 91125 4133004

Professor Royce W. Murray Department of Chemistry University of Morth Carolina Chapel Hill, NC 27514 4133015

Professor Richard Pollard Department of Chemical Engineering Department of Chemistry University of Houston, University Park University of Utah 4800 Calhoun, Houston, TX 77004 413d016

Dr. Donald Sandstrom Boeing Aerospace Company P.O. Box 3999, M/S 87-08 Seattle, WA 98124-2499 4133007

Professor D. E. Irish Department of Chemistry University of Waterloo Waterloo, Ontario, CANADA N21 3G1 4133017

Professor Nathan S. Lewis Division of Chemistry and Chemical Engineering California Institute of Technology Pasadena, CA 91125 413d017

Professor Charles Martin Department of Chemistry Texas A&M University College Station, TX 77843 413d005

Dr. Michael R. Philpott IBM Research Division Almaden Research Center 650 Harry Road San Jose, CA 95120-6099 4133011

Professor B. S. Pons Salt Lake City, UT 84112 4133010

Professor Jack Simons Department of Chemistry University of Utah Salt Lake City, UT 84112 4131050

ONR Electrochemical Sciences Program Robert J. Nowak, Program Manager

Dr. H. Gilbert Smith
EG&G Mason Research Institute
57 Union Street
Worcester, MA 01608
413k003

Dr. Stanislaw Szpak
Code 634
Naval Ocean Systems Center
San Diego, CA 92152-5000
4131006

Professor Michael Weaver Department of Chemistry Purdue University West Lafayette, IN 49707 4133001

Professor Geroge Wilson Department of Chemistry University of Kansas Lawrence, KS 66045 413k002

Professor Ernest Yeager Case Center for Electrochemical Sciences Case Western Reserve University Cleveland, OH 44106 4133008 Professor Ulrich Stimming
Department of Chemical Engineering
and Applied Chemistry
Columbia University
New York, NY 10027
4133014

Professor Petr Vanýsek
Department of Chemistry
Northern Illinois University
Dekalb, IL 60115
413k001

Professor Henry White Department of Chemical Engineering and Materials Science 421 Washington Ave., SE Minneapolis, MN 55455 4000027yip

Professor Mark S. Wrighton
Department of Chemistry
Massachusetts Institute of Technol
Cambridge, MA 02139
4131027

Dept. of Chemistry, York University,
North York, Toronto, Ontario, Canada, M3J 1P3

Molecular Orientation of the Silver Tetraneopentoxyphthalocyanine and Stearic Acid Mixed Langmuir-Blodgett Film and Its Chemical and Electrochemical Behavior

by Yansong Fu and A.B.P. Lever

Abstract

Mixed Langmuir-blodgett films comprised of various relative concentrations of silver tetraneopentoxyphthalocyanine (AgTNPc) and stearic acid are reported. Electronic spectroscopic polarisation data are discussed in terms of the relative orientation of aggregated AgTNPc domains in the film. Electrochemical data confirm the oxidation of AgIITNPc(-2) successively to [AgIIITNPc(-2)]⁺ and [AgIIITNPc(-1)]²⁺ in the film phase without loss of orientation. The film was also subject to chemical oxidation and reduction and the degree of organisation following such chemistry is discussed. AgIITNPc(-2) aggregates, even when diluted with stearic acid, do not display resolved ESR spectra but ternary film mixtures of AgIITNPc(-2)/H₂TNPc(-2)/stearic acid do exhibit such resolved spectra.

Access	ion For				
NTIS	GRA&I				
DTIC T	IAB				
Unannounced 🔲					
Justification					
By					
Dist	Specia.				
A-1					

Introduction

The phthalocyanines, ^{1,2} known for some 60 years, and used primarily in the dye, paint and ink industry, now seem ready for exploitation in a wide range of other diverse applications. ³ The Langmuir-Blodgett (LB) technique for generating monolayer structures has considerable promise for commercial applications involving these phthalocyanine species. ⁴ Such applications include electrochromic devices, ⁵ molecular electronics ^{6,7}, switching devices ⁸ and, especially, chemical gas sensors. ⁹⁻¹² Success is largely dependent on the degree of the molecular organization within the film and the ability to control and reproduce such organization. Recently, Vandevyver and Barraud ¹³ have reviewed the techniques for the LB film characterization, among which linear dichroism measurement, ^{14,15} grazing incidence reflection infrared spectroscopy ¹⁶ and ESR spectroscopy ^{17,18} are the most convenient and commonly available methods to investigate molecular orientation in the film.

We have recently described ¹⁹ the preparation and characterization of a new soluble silver ^{II} phthalocyanine, Ag ^{II}TNPc (TNPc = tetraneopentoxyphthalocyanine). The solubility in a range of non-donor organic solvents provides a means of studying silver phthalocyanine in a fashion unavailable previously for the insoluble unsubstituted analogue. The neopentoxy groups are randomly distributed among the 3 and 4 positions of the outer benzene rings so that this species is isolated as a mixture of geometric isomers which are not separable by the usual chromatographic procedures. Nevertheless it has proven possible to generate organized films therewith.

The present study describes the formation of a mixed LB film of Ag IITNPc and stearic acid which is characterized using dichroism measurements to show the average molecular orientation of the film. Strong intermolecular aggregation is present as monitored by UV-Vis and ESR spectra.

While it is sometimes difficult to transfer a pure phthalocyanine monolayer onto a solid substrate, a mixture of the phthalocyanine with some long chain amphiphilic

species, such as octadecanol, ²⁰ stearic ²¹⁻²³ or arichidic acid, ²⁴ has proven effective. It would be interesting to know how the long chain species such as stearic acid functions in the film, and how this might affect the molecular orientation and the chemistry of the co-partner. The understanding of this issue is important to the utility of those long chain hydrocarbon species in making mixed LB films.

To further characterize the film, linear cyclic voltammetry (LCV) and differential pulse cyclic voltammetry (DPV) were carried out on the film deposited on a SnO2 film-coated glass slide. The stability of the molecular organization within the film was also investigated by monitoring the change in dichroic ratio of the film upon electrochemical or chemical modification.

Experimental Details

Materials: Metal-free tetraneopentoxyl phthalocyanine (H2TNPc) was prepared following the literature methods. 25 Silver II tetraneopentoxyphthalocyanine (Ag^{II}TNPc) was prepared according to the recently described method. ¹⁹ Toluene, as spreading solvent, was Aldileh HPLC grade and glass distilled. The subphase was water purified by double distillation over KMnO₄ followed by passage through a Barnstead organic removal cartridge and two Barnstead mixed resin ultrapure cartridges.

Spectroscopic and Electrochemical Methods: The film electronic spectra and dichroic ratio were measured with a Hitachi Perkin - Elmer Microprocessor 340 spectrometer. Polaroid film polarizers were mostly employed, but some late experiments used calcite polarizers and a Cary 2415 spectrometer. A Varian E4 EPR spectrometer was used to record ESR spectra. Cyclic and differential pulse voltammetry were performed with a Princeton Applied Research (PARC) model 173 Potentiostat/Galvanostat, controlled by a PARC 175 Universal Programmer, and a PARC 174A Polarographic Analyzer. Spectro-electrochemical data were collected with a Guided Wave Inc. model 100 Spectrum Analyzer and the PARC model 173 Potentiostat.

Preparation of the LB Film: The LB trough was built at York following a described design. 26 The measurements of surface pressure (π) versus area per molecule (A) isotherms and multilayer deposition were carried out with an Apple II+ microcomputer controlled LB system, using a local program. In general, a monolayer was compressed at a speed of 1.5 Å^2 /molecule/min. and then transferred onto a substrate at 5 mm/min. Before the actual deposition, about 10 minutes are allowed for the self-organization of the molecules after forming the monolayer. Surface pressure was measured based on Wilhelmy's principle, 27 calibrated with a stearic acid monolayer, whose collapse pressure was reported to be 40 mN/M. 27 The spreading solution was usually of $8 \times 10^{-4} \text{ M}$ and the deposition pressure was about 20 mN/M.

Glass and quartz slides were first treated with boiling CH₂Cl₂ or CHCl₃ and then sonicated in a 1 M NaOH aqueous solution. The resulting hydrophilic surface of the slide was made hydrophobic by immersing the slide in a 5% (Vol.) Me₂Cl₂Si/CCl₄ (Me-methyl) solution for a few minutes. For SnO₂ film-coated glass slides, a slightly basic Decon solution was used instead of the NaOH solution. A carefully stored, cleaned slide is usable for a few days before re-cleaning would be necessary.

Results and Discussions

Pure Ag^{II} TNPc Monolayer: Figure 1 (curve a) shows the surface pressure (π) versus area per molecule isotherm for a pure Ag^{II} TNPc monolayer. The monolayer has a fairly large compressibility, as implied by the fact that the surface pressure begins to rise at about 90 Å² and collapses at about 55 Å², giving an area of 35 Å² for the compression. The large compressibility is attributed to the interaction between the neopentoxyl groups of the molecule. It can also be seen that the monolayer collapses at a relatively low pressure, around 16 mN/M, suggesting that a fragile monolayer has been formed on the subphase. The fragility of the film led to a poor deposition ratio (the area loss on the subphase surface over the area coated on the solid substrate), being

only 0.2 or so, and thus, no useful film of pure Ag^{II} TNPc was obtained. The low collapse pressure of the Ag^{II} TNPc monolayer is due to the large compressibility causing the monolayer to fold easily. In addition, the area per molecule of the monolayer at zero surface pressure was found to be $76 \, \text{Å}^2$, indicating a tilted orientation of the phthalocyanine macrocyclic plane with respect to the subphase surface.

Mixed Films of Ag^{II} TNPc and Stearic Acid: When a single-component monolayer shows a poor deposition property, improved behaviour can often be observed by mixing the pure monolayer with some long chain hydrocarbon alcohol or acid. In our study, stearic acid $(C_{18}H_{36}O_2, SteA)$ was mixed with the Ag^{II} TNPc species to improve the deposition ratio. The pressure isotherm of the Ag^{II} TNPc(-2)/SteA mixed monolayer with a molar ratio of $1:2 = Ag^{II}$ TNPc:SteA is also shown in Figure 1 (curve d). Compared to curve (a) in the same figure, the mixed monolayer showed a much smaller compressibility and a higher collapse pressure (26 mN/M), indicating that a more compact monolayer had been obtained.

The mixed monolayer can be readily deposited onto hydrophobic surfaces of different kinds of solid substrates, such as a glass/quartz slide, SnO₂ film-coated glass slide, and an HOPG (highly-oriented pyrolytic graphite) plate, all with a deposition ratio of nearly unity. The total molecular area occupied by the mixed monolayer on the water surface equals the sum of the individual molecular areas of the two components within experimental error when assuming the molecular areas to be 76 Å² for Ag^{II}TNPc and 21 Å² for SteA. This observation suggests that the Ag^{II}TNPc molecules and SteA molecules all have direct contact with subphase surface, indicative of immiscibility of the Ag^{II}TNPc and stearic acid components.

The Ag IITNPc molecules are believed to be packed tightly by the SteA molecules; when a hydrophobic substrate is dipped into the water, its surface interacts with the

SteA hydrocarbon long chain strongly enough to bring the entire mixed monolayer onto the substrate. X-type transference (the monolayer can only be transferred upon the downward stroke of a substrate) was observed, which probably suggests that after the first layer was transferred, the deposition of the successive layers involved only the interaction between the neopentoxyl groups of the Ag^{II}TNPc and the hydrocarbon long chains of the SteA.

In Figure 2, the electronic spectrum (curve a) of the mixed film displays a normal Soret band at 350 nm and a Q-band at about 620 nm with a shoulder at about 680 nm. The 620 nm Q-band is blue shifted from the corresponding band in solution, ¹⁹ indicative of strong inter-molecular aggregation; ^{23,28,29} the 680 nm shoulder probably indicates a small amount of monomer species exists in the film, though it could arise from transitions within low symmetry aggregated species. ²⁹

One of the Ag^{II}TNPc(-2)/SteA mixed films was washed off the glass slide with toluene and the spectrum of the resultant solution (about 10⁻⁵M) was taken, as shown in Figure 2-b. The spectrum looks identical to that of a newly-prepared Ag^{II}TNPc(-2)/toluene solution, ¹⁹ suggesting that no chemical change has occurred to the Ag^{II}TNPc molecules.

The inset in Figure 3, an ESR spectrum taken from an Ag^{II}TNPc(-2)/SteA (molar ratio, 1:2) mixed film, shows only a single broad peak arising from the unpaired electron in the Ag^{II} ion. The strong inter-molecular interaction has obscured the hyperfine coupling due to the silver nucleus and the four nitrogen nuclei, which otherwise should yield eleven peaks. ^{19,30}

However, using a ternary film composed of $\operatorname{Ag}^{IL}\operatorname{TNPc}(-2)/H_2\operatorname{TNPc}(-2)/\operatorname{SteA}$ (1:11:24), hyperfine coupling was observed with anisotropic g values (Figure 3). Thus in this case, the $H_2\operatorname{TNPc}$ molecules are able to slip between adjacent $\operatorname{Ag}^{IL}\operatorname{TNPc}$ molecules and reduce the degree of inter- Ag^{II} relaxation.

Dichroic Ratio Measurements: When polarized light is used to record the electronic spectrum of a well-organized film, the absorption of the light is expected to vary with the angle between the direction of the electric vector of the light and the dipping direction when making the film. Shown in Figure 4 are the electronic absorption spectra of two Ag ITNPc(-2)/SteA mixed films with the electric vector perpendicular (E₁) and parallel (E₁) to the dipping direction. The fact that Abs is larger than Abs suggests that most of the Ag ITNPc molecules in the film tend to sit on the substrate surface with their macrocyclic plane facing the dipping direction. Moreover since this film is 12 layers thick, the films must lay down on top of one another in a well organised fashion.

A series of Ag^{II}TNPc(-2)/SteA mixtures have been investigated with the molar ratios being 2:1, 1:1, 1:2, 1:4, and 1:9, in order to determine how the SteA molecules affect the deposition of the Ag^{II}TNPc monolayer and their molecular orientation. The π-A isotherms of the five mixtures are shown in Figure 1. These illustrate that the collapse pressure of the mixed monolayer increases with the amount of SteA in the mixture; the more SteA a monolayer contains, the more difficult it will be to fold the monolayer. The experiment has also shown that when the Ag^{II}TNPc(-2)/SteA molar ratio was larger than one, the deposition ratio became less than unity, with the 2:1 mixture only exhibiting a value of 0.5. This could be understood by recalling that the pure Ag^{II}TNPc monolayer can only be transferred with difficulty.

Each film shows a dichroic effect but to a varying degree. The film with a higher AgIITNPc(-2)/SteA molar ratio gave a higher value of the dichroic ratio (Abs/Abs_{||}), indicative of a higher degree of molecular organization. Two examples are shown in Figure 4 with (a) from the 2:1 mixed film and (b) from the 1:4 mixed film, and in Figure 5 the dichroic ratios for five mixtures are plotted versus the molar fraction of AgIITNPc in the mixture. This clearly illustrates that the molecules in the mixed film become more organized with an increase in the relative concentration of

Ag^{II}TNPc. Thus the SteA molecules do not help to organize the Ag^{II}TNPc molecules but instead reduce the orderliness of the film.

Previous studies with stearic acid diluted phthalocyanine films, ²³ have demonstrated the existence of aggregated domains within the stearic acid arrays. Our data suggest a similar situation. Individual aggregates show high organisation and dichroism, and at low stearic acid concentrations, neighboring domains may serve to orient each other in a parallel fashion. As the relative concentration of phthalocyanine is reduced, the domains are further apart and their orientation more random; thus the dichroism is reduced.

To gain further insight into the molecular orientation in the film, two angles, θ and ϕ , are proposed to represent the average molecular orientation. This is shown schematically in Figure 6, in which Y corresponds to the dipping direction when making the film, Z is the normal of the substrate surface and X is perpendicular to the YZ plane. If a flat circle is used to represent the phthalocyanine transition dipole plane, the angle formed by the normal (z') to the circle with respect to the Z axis will be called θ , and the angle between the projection of the normal in the XY plane and the X axis called ϕ .

Following the procedure developed by Yoneyama et al., 31 orientation angles, θ and ϕ , for each of the five mixed films can be calculated and the results are listed in Table 1. There is no consistent change in the θ value, but the value of ϕ decreases with increasing relative concentration of SteA. Evidently the rotation of Ag TNPc molecules about their x' axis, which is parallel to the X axis, is not affected by the presence of SteA, but the rotation about the r' axis, which is perpendicular to the normal of the phthalocyanine circle and parallel to the YZ plane, will be easier if the amount of SteA is increased. Considering that a film with randomly ordered molecules should have an average ϕ value of 45° ($\int_{0}^{90} \dot{\phi} d\phi / \int_{0}^{90} \dot{d}\phi = 45^{\circ}$) and a perfectly ordered film should show 90° for ϕ , Table 1 also suggests that the molecular organization in the

film becomes worse in SteA mixed films.

In summary, the above observations reveal that although the presence of SteA is necessary to obtain uniform films, the molecular organization of the film is reduced by giving more freedom of orientation for the Ag^{II}TNPc molecules.

Cyclic Voltammetric Studies: An $Ag^{II}TNPc(-2)/SteA$ (1:2) mixed film (3 layers thick) was deposited onto a SnO_2 film-coated glass slide, and studied by cyclic voltammetry in a 0.2M Na ClO_4 aqueous solution. The film voltammogram exhibited a broad and irreversible redox wave with the half-potential, $E_{1/2} = 0.65V$ (vs. SCE), as shown in Figure 7a. The redox process is a one-electron oxidation, leading to the species $[Ag^{III}TNPc(-2)]^+$ based upon the following arguments.

- (1) In the solution voltammetry, Ag^{II}TNPc is successively oxidized in two one-electron steps to [Ag^{III}TNPc(-2)]⁺ and [Ag^{III}TNPc(-1)]²⁺ at 0.71 and 1.19 V (vs SCE) respectively.¹⁹
- (2) When the same kind of film, deposited on a glass slide, was oxidized chemically with chlorine gas, oxidation occurred to yield [Ag^{III}TNPc(-2)]⁺, as indicated by electronic spectroscopy (vide infra).
- (3) A spectroelectrochemical study of the film polarized at +0.75V showed a spectrum with a red shifted and slightly less intense Q-band and a new band around 440 nm (Figure 8). This is consistent with the solution spectroelectrochemical data wherein the formation of [Ag^{III}TNPc(-2)]⁺, from Ag^{II}TNPc in DCB, is associated with a red shifted and less intense Q band, and the appearance of a new absorption at 430 nm. ¹⁹ Upon re-reduction of the oxidized film by polarization at -0.05 V, the 440 nm band disappeared, and the Q band shifted back to its original position with increased intensity. This cycle can be repeated many times, but with some loss of overall Q band intensity each time, probably due to some film loss from the electrode. These observations are inconsistent with the postulate that polarization merely changes the aggregate-monomer

ratio, since such a change towards the monomer would result in an increase in intensity of the red shifted Q band.

In the solution electrochemistry, the first oxidation step of $Ag^{II}TNPc(-2)$ to $[Ag^{III}TNPc(-2)]^+$ is very broad and ill-resolved due to extensive aggregation and a kinetically slow electron transfer step. The $Ag^{II}TNPc(-2)$ LB film is also greatly aggregated, as discussed above. This molecular interaction also results in a broad CV peak for the LB film whose irreversibility is indicated by the large peak separation $(E_a^-E_c^-)$, of over 200mv. The differential pulse cyclic voltammogram, (in Figure 7b), shows that the peak is much smaller upon reversing the scanning. This may be associated with the movement of ClO_4^- anions through the film. Upon oxidation of the film, the ClO_4^- anions penetrate into the film to balance the positive charge. However, after their penetration, the anions are locked inside the film and have difficulty leaving on the time scale of the experiment. The scan rate dependence of the redox wave shows a nonlinear relationship between peak current and scan rate, consistent with this supposition.

Chemical Stability of the LB Film: An important property for an LB film is the stability of the molecular organization of the film against electrochemical or chemical change. We have therefore studied the change in dichroic ratio of the LB film upon electrochemical or chemical modification.

The polarized UV-Vis spectra of a Ag^{II}TNPc(-2)/SteA (1:2) mixed LB film (6 layers) deposited on a SnO₂-film coated glass slide is shown in Figure 9a. They resemble the spectra of the film on a regular glass slide, except for a small broad band around 500 nm, which is believed to be due to a specific interaction with the SnO₂ surface since the absorption remains if the film is oxidized or reduced electrochemically, but is not present in the solution spectrum of the film washed off the surface.

Electrochemical studies were carried out under the same conditions as those employed in the cyclic voltammetric studies. After repeating the cyclic potential scanning in the range of -0.4 to 1.0 V for 9 times, the polarized spectra were recorded (Figure 9b). The dichroic ratio when measured at the main Q-band measured close to 100 % of its original value, although the interaction absorption at 500 nm had increased. The invariance of the dichroic ratio suggests that the molecular organization within the film is stable in terms of general cyclic voltammetric measurement.

Chemical Cycling of LB Film: An Ag^{II}TNPc(-2)/SteA (1:2) film (10 layers) deposited on a glass slide was treated with molecular Cl₂ gas either in the gas or aqueous phase. In the gas phase experiment, the slide was placed, for a few minutes, inside a bottle containing Cl₂ gas, approximately 0.5% by volume in air, and then removed and washed with distilled water. The resulting spectrum is shown in Figure 10b. Compared to the original film spectrum (Figure 10a), the Cl₂-treated film showed a red-shifted Q-band at 640 nm but no new band around 500 nm, consistent with the one-electron oxidation of Ag^{II}TNPc(-2) to [Ag^{III}TNPc(-2)]⁺. The Cl₂-treated film maintained the same degree of molecular organization as before, indicated by the almost unchanged dichroic ratio (1.23 in Figure 10a, and 1.19 in Figure 10b).

When the gas phase Cl₂-oxidized film was immersed in a 10% (Vol) N₂H₄ aqueous solution for 15 to 30 min., the initial Q band envelope was restored entirely with a slight decrease in dichroic ratio (to 1.17), as shown in Figure 10c. Thus the LB film can be oxidized by chlorine gas and reduced back with aqueous hydrazine without change in film organization. The spectra of the [Ag^{III}TNPc(-2)]⁺ species in Figures 8 and 10 likely differ because the counter-ions and, possibly, coordination number are different in the two cases.

In a solution phase oxidation, the same film as above was exposed to a 0.1 M NaClO₄ aqueous solution bubbled with chlorine gas (10 - 20 s) for 2 minutes. In this

case, the electronic spectrum again demonstrated oxidation to $[Ag^{III}TNPc(-2)]^+$ but with a slightly larger red shift in the Q band perhaps because the reaction was more complete. However the dichroic ratio was greatly reduced (Figure 11a), implying substantial loss of film organization. When this film was reduced back to $Ag^{II}TNPc$ (Figure 11b) with hydrazine, the dichroic ratio declined almost to unity, indicating loss of order. A much thinner film (4 layers) was used to repeat the liquid phase experiment, with the same results being obtained.

Thin film reorganization upon redox sampling undoubtedly involves solvent transfer through the film. ^{32,33} Thus the loss of organization upon solvent phase oxidation, but retention during gas phase oxidation may reflect the disruptive effect of such solvent transport in the former case.

Langmuir-Blodgett films can be co-deposited with stearic acid onto suitable substrates, with a deposition ratio of unity. Increasing stearic acid facilitates the transfer of a compact film, but decreases the degree of organization of the Ag IITNPc(-2) component. The ESR spectra of the mixed Ag IITNPc(-2)/SteA films show that aggregation is still occurring, evidence that the stearic acid does not interleave between the Ag IITNPc(-2) molecules. Thus dilute films are likely composed of domains of aggregated Ag IITNPc(-2) which are internally ordered, separated by stearic acid, but with some loss of relative organization between the domains. The presence of the dichroism in the film spectra suggests that each domain consists of Ag IITNPc(-2) molecules with their macrocyclic plane tending to face the dipping direction. In the films of higher relative Ag IITNPc concentration, each of the domains has a similar orientation, resulting in the anisotropic optical property. The degree of freedom with which the domains orient is dependent upon the relative amount of SteA in the mixture. A larger relative amount of SteA results in a larger degree of freedom for domain

orientation and, therefore, worsens the film organization.

Film organization is largely unaffected by electrochemical cycling or by chemical cycling using gaseous chlorine and aqueous hydrazine, but the order is rapidly lost if the film is cycled using a solution chemical oxidant, perhaps due to solvent transport. Retention during electrochemical oxidation in solution shows that different solvent transport and likely anion compensation pathways occur depending upon whether the method of oxidation is chemical or electrochemical. Clearly the detailed mechanisms require further study.

Acknowledgements We are indebted to the Natural Sciences and Engineering Research Council (Ottawa), and the Office of Naval Research (Washington) for financial support.

Reference

- (1) Lever, A. B. P. Adv. Inorganic Chem. and Radiochem 1965, 7, 27.
- (2) Leznoff, C. C.; Lever, A. B. P. Eds. <u>Phthalocyanines</u>. <u>Properties and Applications</u>; VCH: New York, 1989.
- (3) Lever, A. B. P. <u>CHEMTECH</u> 1987, <u>17</u>, 506.
- (4) Fourth International Conference on Langmuir-Blodgett Films, Tsukuba, Japan, April 1989, Thin solid Films 1989, 179, 1-554.
- (5) Yamamoto, H.; Sugiyama, T.; Tanaka, M. Japn. J. Appl. Phys. 1985, 24, L305.
- (6) Sugi, M. J. Molecular Electronics 1985, 1, 3.
- (7) Hann, R. A.; Gupta, S. K.; Fryer, J. R.; Eyres, B. L. Thin Solid Films 1985, 134, 35.
- (8) Vandevyver, M.; Ruaudel-Teixier, A.; Barraud, A. <u>J. Chim. Phys.</u> 1985, 82(6), 707.
- (9) Tredgold, R. H.; Young, M. C. J.; Hodge, P.; Hoorfar, A. <u>IEEE Proc. I.</u> 1985, 132, 152.
- (10) Fujiki, M.; Tabei, H.; Imamura, S. Japn. J. Appl. Phys. 1987, 26, 1224.
- (11) Reswick, R. B.; Pitt, C. W. Chem. Phys. Letters 1988, 143, 589.
- (12) Palacin, S.; Ruaudel-Teixier, A.; Barraud, A. <u>J. Phys. Chem.</u> 1986, 90, 6237.
- (13) Vandervyver M.; Barraud, A. J. Molecular Electronics 1988, 4, 207.
- (14) Kalina, D. W.; Crane, S. W. Thin Solid Films 1985, 134, 109.
- (15) Ogawa, K.; Kinoshita, S.; Yonehara, H.; Nakahara, H.; Fukuda, K. <u>J.</u>
 Chem. Soc., Chem. Commun 1989, 477.

- (16) Mumby, S. J.; Swalen, J. D.; Rabolt, J. F. Macromolecules 1986, 19, 1054.
- (17) Cook, M. J.; Daniel, M. F.; Dunn, A. J.; Gold, A. A.; Thomson, A. J. J. Chem. Soc., Chem. Commun. 1986, 863.
- (18) Pace, M. D.; Barger, W. R.; Snow, A. W. J. Mag. Res. 1987, 75, 73.
- (19) a) Fu, G.; Fu, Y.; Lever, A. B. P. <u>Inorg.Chem.</u> 1990, 29, 4090. b)
 Ouyang, J.; Lever, A. B. P. <u>J. Phys. Chem.</u> 1991, 95, in press. c)
 Ouyang, J.; Lever, A. B. P. <u>J. Phys. Chem.</u> 1991, 95, accepted.
- (20) Barger, W. R.; Snow, A. W.; Wohltjen, H.; Jarvis, N. L. <u>Thin Solid</u> <u>Films</u> 1985, 133, 197.
- (21) Palacin, S.; Ruaudel-Teixier, A.; Barraud, A. <u>J. Phys. Chem.</u> 1989, 93, 7195.
- (22) Watanabe, I.; Hong, K.; Rubner, M. F. <u>J. Chem. Soc., Chem. Commun</u> 1989, 123.
- (23) Petty, M.; Lovett, D. R.; O'Connor, J. M.; Silver, J. Thin Solid Films 1989, 179, 387.
- (24) Aroca, R.; Battisti, D. Langmuir 1990, 6, 250.
- (25) Leznoff, C. C.; Greenberg, S.; Marcuccio, S. M.; Minor, P. C.; Seymour, P.; Lever, A. B. P.; Tomer, K. B. Inorg. Chim. Acta 1984, 89, L35.
- (26) Van Ryswyk, H.; Eisenhart, J. M.; Blom, N. Rev. Sci. Instrum. 1985, 56, 2339.
- (27) Gaines, Jr., G. L. <u>Insoluble Monolayers At Liquid-Gas Interface</u>; Interscience: New York, 1966.
- (28) Snow, A. W.; Jarvis, N. L. J. Am. Chem. Soc. 1984, 106, 4706.
- (29) Dodsworth, E. S.; Lever, A. B. P.; Seymour, P.; Leznoff, C. C. <u>J. Phys.</u>
 Chem. 1985, 89, 5698.
- (30) Kneubuhl, F. K.; Koski, W. S.; Caughey, W. S. J. Am. Chem. Soc. 1961,

<u>83</u>, 1607.

- (31) Yoneyama, M.; Sugi, M.; Saito, M. Japn, J. Appl. Phys. 1986, 25, 961.
- (32) Bruckenstein, S.; Hillman, A. R.; <u>J. Chem. Phys.</u> 1988, <u>92</u>, 4837.
- (33) Bruckenstein, S.; Wilde, C. P.; Hillman, A. R.; <u>J. Chem. Phys.</u> 1990, <u>94</u>, 6458.

Table 1 Molecular Orientation as a Function of Molar Fraction

Mol.Frac. Ag ^{II} TNPc	θ	φ	
0.10	52	46	
0.20	52	53	
0.33	42	63	
0.50	46	72	
0.67	48	79	

See text and Figure 6 for coordinate system.

Figure Legends

Figure 1. Surface pressure vs. area per molecule isotherms for a) pure $Ag^{II}TNPc$, and the five $Ag^{II}TNPc(-2)/SteA$ mixtures with molar ratios being $(Ag^{II}TNPc:SteA)$: b) 2:1, c) 1:1, d) 1:2, e) 1:4, f) 1:9.

Figure 2. Electronic absorption spectra of a) Ag^{II}TNPc(-2)/SteA (1:2) mixed LB film, with a thickness of 12 layers, and b) Ag^{II}TNPc(-2)/toluene solution, about 5x10⁻⁵M, path length 0.2 cm, made by washing off the LB film with toluene.

Figure 3. ESR spectra of Ag^{II}TNPc(-2)/H₂TNPc(-2)/SteA (1:11:24) mixed LB film of 40 layers on quartz slide. The inset in the figure:

Ag^{II}TNPc(-2)/SteA (1:2) mixed LB film of 50 layers on quartz slide.

Figure 4. Polarized electronic spectra of $Ag^{II}TNPc(-2)/SteA$ mixed LB film (12 layers thick) with molar ratios (AgTNPc:SteA) being a) 2:1 and b) 1:4, and the electric vector of the incident light being α) perpendicular (E_{II}) and β) parallel (E_{II}) to the dipping direction.

Figure 5. A relationship between dichroic ratios of Ag^{II}TNPc(-2)/SteA mixed LB films and the molar fraction of Ag^{II}TNPc.

Figure 6. Coordinate system for showing the molecular orientation of Ag^{II}TNPc in the film. Y is the dipping direction, Z is the normal of the substrate surface and X is perpendicular to the YZ plane.

Figure 7. a) Linear cyclic voltammogram (scan rate 100 mV/s) and b) differential pulse cyclic voltammogram (scan rate 2 mV/s) of Ag^{II}TNPc(-2)/SteA (1:2) mixed LB film on a SnO₂ film-coated glass slide (8 layers thick) in 0.2M NaClO₄ aqueous solution.

Figure 8. Electronic spectra of Ag ^{IL}TNPc(-2)/SteA (1:1) mixed LB film (12 layers thick) and the oxidized species generated by potentiostatic electrolysis, the potential was set at 0.75V vs. SCE.

Figure 9. Dichroic measurements of Ag ^{II}TNPc(-2)/SteA (1:2) mixed LB film (6 layers) on a SnO₂ film-coated glass slide with a) before and b) after cyclic voltammetric studies (9 cycles).

Figure 10. Polarized spectra of $Ag^{II}TNPc(-2)/SteA$ (1:2) mixed LB film (10 layers) on glass slide upon chemical reactions: a) original film; b) 5 minute exposure to 0.5%(Vol.) Cl_2 gas in air; c) 25 minute immersion of the Cl_2 -treated film in a 10%(Vol.) N_2H_4 aqueous solution.

Figure 11. Polarized spectra of $Ag^{II}TNPc(-2)/SteA$ (1:2) mixed LB film (10 layers) on glass slide. a) The film has been treated with a Cl_2 -bubbled (15 secs) 0.1M $NaClO_4$ aqueous solution for 3 minutes; b) the oxidized film was treated with a 10%(Vol.) N_2H_4 aqueous solution for 25 minutes.





















