THE FILE MAPY

OFFICE OF NAVAL RESEARCH

Contract N00014-87-J-1118

R & T Code 4133016

Technical Report No. 8

A Raman Spectroscopic and Electrochemical Study on Photoinduced Crystal Growth of DABCO Polyiodide Upon a Silver Electrode

by

T. Ozeki and D.E. Irish

Prepared

for

1989 International Congress of Pacific Basin Societies

Guelph-Waterloo Center for Graduate Work in Chemistry Waterloo, Campus Department of Chemistry University of Waterloo Waterloo, Ontarlo Canada, N2L 3G1

June 13, 1989



05 066

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.

30

SECURITY CLA	SSIFICATION O	THIS PAGE					
		• •	REPORT DOCUM	AENTATION A	PAGE		
TA REPORT S	IN REPORT SECURITY CLASSIFICATION				16. RESTRICTIVE MARKINGS		
Unclas	Safied	N AUTHORITY			AVAILABILITY OF	AFPORT	
Unclas	sified	· · · · · · · · · · · · · · · · · · ·					
20 DECLASSIF	KATION / DOW	INGRADING SCHEDU	LE	Public Release/Unlimited			
4 PERFORMIN	G ORGANIZAT	ION REPORT NUMBE	R(S)	S. MONITORING ORGANIZATION REPORT NUMBER(S)			
ONR Technical Report #8							
6. NAME OF	PERFORMING	ORGANIZATION	6b. OFFICE SYMBOL (If applicable)	78 NAME OF MO	NITORING ORGAN	IZATION	
Univer	sity of Wa	aterloo		Office of	f Naval Rese	earch	
6c. ADDRESS (City, State, and	d ZIP Code)		7b. ADDRESS (City, State, and ZIP Code)			
Depart	ment of Cl	hemistry		The Ohio State University, Research Center			
Water1	oo. Ontar:	io. Canada. Ni	2L 3G1	Columbus, Ohio, U.S.A., 43212–1194			
Ba. NAME OF FUNDING / SPONSORING			85. OFFICE SYMBOL	9. PROCUREMENT INSTRUMENT IDENTIFICATION NUMBER			MBER
ORGANIZATION Office of Naval Research			(m appircable)	N00014-8	7-J-1118		
BC. ADDRESS (City, State, and	I ZIP Code)	1	10 SOURCE OF FUNDING NUMBERS			
Chemis	try Divis	ion		PROGRAM	PROJECT NO.	TASK NO	WORK UNIT
Arling	ton, VA, I	U.S.A., 22217-	-5000				
11 TITLE (Incl A Rama DABCO	ude Securny C n Spectros Polyiodide	Temfication) scopic and Ele e upon a Silve	ectrochemical S er Electrode	tudy on Photo	oinduced Cry	vstal Growth	ı of
12 PERSONAL T. Oze	AUTHOR(S) ki, and D.	.E. Irish	· •				
13a. TYPE OF Techni	REPORT cal	136 TIME CO	DVERED 3/38 to 06/89	14. DATE OF REPO 1989-06-13	RT (Year, Month, I	Day) 15. PAGE 11	COUNT
16 SUPPLEME To be Decemb	ntary notal presented er 17-22,	non at the 1989 1 1989.	International C	ongress of P	acific Basir	n Societies,	Honolulu,
17	COSATI	CODES	18 SUBJECT TERMS (Continue on reverse	e if necessary and	l identify by bloc	k number)
FIELD	GROUP	SUB-GROUP	> Laser-induce Spectroscopy	d crystal gr	owthe DABCOR	$I_2(I_3)_2$; Ran $F_A \sigma / DABOOH_2$	$an_{2+/T}$
			opectroscopy				. / -
9 ABSTRACT	(Continue on	reverse if necessary	and identify by block i	number)	GAG	ik	
$\sum_{i=1}^{n}$						dia solution	containing
Surface Raman spectra of a silver electrode contacting with an acidic solution containing							
triethylenediamine (DABCO) and sodium lodice showed some new remning perception of dissolved DABCO species nor of iodide. These were ascribed to diprotonated DABCO polyiodide							
crystal, formed and attached onto the laser-irradiated site of the silver electrode. Several facts implied that							
laser illumination used for the Raman measurement promoted the growth of the crystal, the possibility of the last of the last of the state of the st							
of diprotonated DABCO polyiodide on the silver electrode was examined, as to its dependence upon the							
solution composition and applied potential.							

....

20. DISTRIBUTION / AVAILABILITY OF ABST	RACT	21. ABSTRACT SECURITY CLASSIFICATION
UNCLASSIFIED/UNLIMITED SAM	E AS RPT. DTIC USERS	Unclassified
228 NAME OF RESPONSIBLE INDIVIDUAL		22b. TELEPHONE (Include Area Code) 22c. OFFICE SYMBOL
Dr. Robert J. Nowak		(519) 885-1211, ext, 2500
	82 A 88 edition may be used up	til anhausted

DD FORM 1473, 84 MAR

83 APR edition may be used until exhausted. All other editions are obsolete.

SECURITY CLASSIFICATION OF THIS PAGE Unclassified

A RAMAN SPECTROSCOPIC AND ELECTROCHEMICAL STUDY OF THE

PHOTOINDUCED CRYSTAL GROWTH OF DABCO POLYIODIDE

UPON A SILVER ELECTRODE

Toru Ozeki and Donald E. Irish*

Hyogo Univ	versity of Tead	cher Education	
942-1 Shimokume,	Yashiro, Kato), Hyogo, Japan	673-14

*Department of Chemistry University of Waterloo Waterloo, Ontario Canada N2L 3G1

OT1C

COPY NSPECTE

Accession For
NTIS GFASI
DTIC TAP
Unannouries
Justification
By
Distribution
A
Avall Codes
Die A - Di/or
Dist Special
1

A large number of studies of Surface Enhanced Raman Scattering (SERS) from organic molecules adsorbed on silver electrodes have been reported. Recent studies of triethylenediamine (DABCO) carried out in our laboratory (1-3) have been extended and the results are presented below. When the Raman spectrum of an acidic solution of DABCO was measured in the presence of sodium iodide new Raman peaks were observed in addition to those due to the dissolved DABCO species and electrode-adsorbed iodide, as shown in Fig. 1-C. These new bands were ascribed to the diprotonated DABCO polyiodide crystal, formed and attached onto a laser-irradiated site of the silver electrode. Several facts suggest that the laser irradiation, applied for the Raman measurement, promoted the crystal growth; the possibility of so-called 'photoinduced electrochemical crystal growth' was suggested.

Experimental

A Dilor OMARS-89 spectrometer with microscope, interfaced to an IBM-AT computer, was used for the measurements. The 514.5 nm line of an Argon ion laser was used as an excitation light source. The working electrode was a silver rod of 5 mm diameter sealed inside an 8 mm diameter Teflon sheath, which was assembled in an electrochemical Raman cell specially designed for use under the microscope. The surface of the silver was polished with 0.3 μ m alumina slurries on metron cloths. A platinum wire was used as counter electrode; silver wire, immersed in 1 M NaI solution, was used as reference electrode; a salt bridge separated the sample solution from the inner solution of the reference electrode.

Results and Discussion

Raman spectra of an alkaline solution (pH 10.77, Fig. 1-A) and a slightly acidic solution (pH 6.23, Fig. 1-B), both containing 0.1 M DABCO and 0.5 M NaI, gave peaks ascribed to the bulk species of DABCO or to the electrode-adsorbed iodide ion. However, two additional peaks 220 and 160 cm⁻¹, were observed in the Raman spectra of an acidic solution such as pH 0.23 (Fig. 1-C); the peak at 110 cm⁻¹, which was observed for all pH values, was also strongly enhanced in the acid solution.

After keeping the potential \sim the silver electrode at +100 mV for several hours and simultaneously irradiating the surface with the 514.5 nm laser line at 300 mW, the growth of needle-like crystals was observed in the acidic solution, radiating out from the focal center of the laser beam on the electrode surface as shown in Fig. 2. This crystal did not disappear even at -100 mV, but completely disappeared at -150 mV. The cyclicvoltammogram of the acidic solution differs significantly in shape from those of the near-neutral and alkaline solutions as shown in Fig. 3. In the acidic solution, oxidation current starts to flow already at -0.08 V; and reduction current remains small up to -0.15 V. This potential was consistent with the

- 2 -

potential where the crystal disappeared. The Raman peaks at 200 and 160 cm^{-1} observed at anodic potential also disappeared at -150 mV as shown in Fig. 4.

In order to identify these Raman peaks, several Raman measurements were carried out: (1) for the silver iodide film covering silver metal in air, (2) for the solution containing only sodium iodide, (3) for the solution containing iodide ion and iodine (the formation of I_3^- is observed), (4) for pure iodine crystal, etc. (Fig. 5). Experimental results suggest that the 220 cm⁻¹ peak should be ascribed to iodine (I_2), formed by the photodecomposition of the silver iodide; the 120 and 160 cm⁻¹ peaks are ascribed to the symmetric and asymmetric stretching vibrations of i_3^- , formed by the reaction between I_2 and I^- .

A calculation of equilibria as to the dissolved DABCO species suggests that the major species in the solutions with pH 10.77, 6.23 and 0.23 are unprotonated, monoprotonated, and diprotonated DABCO, respectively.

These facts suggest the following reaction mechanism for the crystal growth, observed for the acidic solution under laser light irradiation:

$$2 Ag + 2 I^{-} = 2 AgI + 2 e^{-}$$
 (1)

$$2 \operatorname{Agl} + \operatorname{hv} = 2 \operatorname{Ag} + I_2 \tag{2}$$

$$I_2 + I^2 = I_3^2$$
 (3)

$$DABCO-H_2^{2+} + 2I_3 = DABCO-H_2(I_2)_2$$
(4)

Thus a net reaction can be represented as follows:

(Ag)

$$DABCO-H_2^{2+} + 6 I^- + 2 hv = DABCO-H_2(I_3)_2 + 4 e^-$$
(5)

Electrochemically-formed silver iodide (Eq. 1) is photodecomposed to atomic silver and iodine (Eq. 2). The iodine forms triiodide ion with another iodide ion which comes by diffusion from the bulk solution (Eq. 3), and is fixed upon DABCO ion (Eq. 4). These steps (Eq. 1 to 4) are repeated (Fig. 6), and the needle-like crystal growth results at the laser-irradiated site of the silver surface. Light illumination, acidic solution, and anodic potential are required to cause the crystal growth to proceed.

In the above explanation, the participation of the trilodide ion is assumed. However, there is the possibility that more highly aggregated iodide ion such as I_5^- is participating in the reaction. So far this point remains unclear.

Acknowledgement

This work was supported by grants from the Natural Sciences and Engineering Research Council of Canada and the Office of Naval Research.

References

- 1. D.E. Irish, D. Guzonas, and G.F. Atkinson, Surface Science, 1985, 158, 314.
- 2. D.A. Guzonas, D.E. Irish, and G.F. Atkinson, Langmuir, 1989, 5, 787.
- 3. D.A. Guzonas, Ph.D. Thesis, University of Waterloo, Waterloo, Ontario, Canada, 1987.



Fig.1 Surface Raman spectra of silver metal in 0.5 M NaI and 0.1M DABCO solutions with A:pH 10.77, B:pH6.23 C:pH0.23. B-2 is the spectrum after the illumination of laser light of 514.5 nm 100mW for 60min.



Fig.2 Typical examples of diprotonated triiodide crystal upon silver electrode found 3h after kept at +100mV under 514.5 nm 300mW laser illumination.



Fig.3 Cyclicvoltammograms of 0.5 M NaI and 0.1M DABCO solutions with A:pH10.72, B:pH5.40 and C: pH 0.93. Scan rate is 10 mV / s.



Fig.4 Dependence of surface Raman spectrum of silver electrode upon the electrode potential for 0.5 M NaI and 0.1M DABCO solution with pH0.85.



Fig.5 Comparison of several Raman spectra: A, surface Raman spectrum of silver metal in 0.5M NaI and 0.1M DABCO solution with pH0.23; B, bulk Raman spectrum of 1M NaI + 0.05M I₂ solution; C, bulk Raman spectrum of 1M NaI solution; and D, surface Raman spectrum of pure iodine crystal.



Figure 6 Schematic illustration of diprotonated DABCO triiodide crystal formation upon silver electrode.

QL/1113/87/2

•

TECHNICAL REPORT DISTRIBUTION LIST, GEN

-

. .

	No.		No.
	Copies		Copies
Office of Naval Research Attn: Code 1113 800 N. Quincy Street Arlington, Virginia 22217-5000	2	Dr. David Young Code 334 NORDA NSTL, Mississippi 39529	1
Dr. Bernard Douda Naval Weapons Support Center Code 50C Crane, Indiana 47522-5050	1	Naval Weapons Center Attn: Dr. Ron Atkins Chemistry Division China Lake, California 93555	1
Naval Civil Engineering Laboratory Attn: Dr. R. W. Drisko, Code L52 Port Hueneme, California 93401	1	Scientific Advisor Commandant of the Marine Corps Code RD-1 Washington, D.C. 20380	1
Defense Technical Information Center Building 5, Cameron Station Alexandria, Virginia 22314	12 high quality	U.S. Army Research Office Attn: CRD-AA-IP P.O. Box 12211 Research Triangle Park, NC 27709	1
DTNSRDC Attn: Dr. H. Singerman Applied Chemistry Division Annapolis, Maryland 21401	1	Mr. John Boyle Materials Branch Naval Ship Engineering Center Philadelphia, Pennsylvania 1911	1 2
Dr. William Tolles Superintendent Chemistry Division, Code 6100 Naval Research Laboratory Washington, D.C. 20375-5000	1	Maval Ocean Systems Center Attn: Dr. S. Yamamoto Marine Sciences Division San Diego, California 91232	1