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Raman, Surface Enhanced Raman Scattering and
Infrared Studies of Phthalocyanines: Thin Films and
Langmuir-Blodgett Monolayers

BY

DORIANO BATTISTI

A Dissertation
Submitted to the Faculty of Graduate Studies through the
Department of Chemistry and Biochemistry in partial fulfillment
of the requirements for the Degree of Doctor of Philosophy at
The University of Windsor
1992
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ABSTRACT

Raman scattering of thin films, and surface enhanced \{resonance\} Raman scattering(SE\{R\}RS), mostly of Langmuir-Blodgett(LB) monolayers are interpreted for various phthalocyanine derivatives. SERS and SERRS are reported for phthalocyanines transferred to silver and gold island films and silver coated tin spheres. The Raman data were complemented with the infrared(IR) data and the vibrational assignment of the phthalocyanine was attempted. The Raman spectra of vanadyl phthalocyanine sublimed as an ultra thin film onto reflecting surfaces are reported. Energy transfer and enhanced absorption was observed in a two component system composed of one LB of a perylene derivative and one LB of zinc tetra-tert-butylphthalocyanine. The LB technique was applied to control the coverage on the surface in the investigation of SERRS of vanadyl tetra-tert-butylphthalocyanine and surface coverage dependence. SERRS, IR, ultraviolet-visible(uv-vis) spectroscopy were used to characterize stable green and blue forms of lanthanide bisphthalocyanine complexes. SERRS was applied to investigate the interaction of nitrogen dioxide and one LB of metal-free tetra-tert-butylphthalocyanine, copper tetra-tert-butylphthalocyanine and ytterbium(and lutetium) bisphthalocyanines. UV-VIS spectroscopy was also valuable in the investigation of nitrogen dioxide and the phthalocyanine derivatives.
DEDICATION

To my wife Carmela, our parents and our brothers
ACKNOWLEDGEMENTS

During the course of almost five years it becomes very easy to realize that there was good will from many people. The excellent supervision and encouragement from Dr. R. Aroca led to interesting and dedicated work. His weekly group meetings kept us sharp and "up with the times". I am also grateful to my labmates past and present (C. A. Jennings, K. Akers, F. Martin, U. Guhathakurta, Dr. E. Clavijo, E. Johnson, and J. Soto). They have always provided a good working relationship and challenging discussions. I would also like to express my thanks to our collaborators. They include Drs. G. J. Kovacs, Dr. A. M. Hor, Dr. R. O. Loutfy at the Xerox Research Centre of Canada, Drs. T. Tomilova and E. Luk’yanets at the Organic Intermediate and Dyes Institute, Moscow, and Dr. A. Tsivadze at the Institute of General and Inorganic Chemistry Academy of Science, Moscow. The technical staff at the University of Windsor is also acknowledged. They include Mr. M. Fuert and Mr. J. Olsen from the Department of Chemistry and Biochemistry, Mr. J. Robinson form the Department of Mechanical Engineering and Mr. W. Grewe from the Department of Physics.

Finally I would like to thank my "human catalysts", Carmela for her constant support and understanding, and our parents (Ferruccio, Maria, Angelo) and brothers (Roberto, Alessandro, Carlo).
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<tr>
<td>A</td>
<td>depolarization factor</td>
</tr>
<tr>
<td>Ag-Sn spheres</td>
<td>Silver coated tin spheres</td>
</tr>
<tr>
<td>bend</td>
<td>bending</td>
</tr>
<tr>
<td>CCD</td>
<td>charge coupled device</td>
</tr>
<tr>
<td>CuPc</td>
<td>copper phthalocyanine</td>
</tr>
<tr>
<td>CuPcTTb</td>
<td>copper terta-tert-butylphthalocyanine</td>
</tr>
<tr>
<td>d</td>
<td>metal molecule separation</td>
</tr>
<tr>
<td>def.</td>
<td>deformation</td>
</tr>
<tr>
<td>EPR</td>
<td>electron paramagnetic resonance</td>
</tr>
<tr>
<td>$E_{\text{op}}$</td>
<td>electric field of P-polarized light</td>
</tr>
<tr>
<td>$E_{\text{os}}$</td>
<td>electric field of S-polarized light</td>
</tr>
<tr>
<td>$\varepsilon$</td>
<td>dielectric constant of metal</td>
</tr>
<tr>
<td>$\varepsilon_0$</td>
<td>dielectric constant of medium</td>
</tr>
<tr>
<td>FT</td>
<td>Fourier Transform</td>
</tr>
<tr>
<td>FTIR</td>
<td>Fourier Transform Infrared</td>
</tr>
<tr>
<td>H$_2$Pc</td>
<td>metal free phthalocyanine</td>
</tr>
<tr>
<td>HoPc$_2^+$</td>
<td>holmium octa-tert-butylbisphthalocyanine</td>
</tr>
<tr>
<td>IR</td>
<td>infrared</td>
</tr>
<tr>
<td>LB</td>
<td>Langmuir-Blodgett</td>
</tr>
<tr>
<td>LL</td>
<td>laser line</td>
</tr>
<tr>
<td>Ln</td>
<td>lanthanide</td>
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</table>
LnPc₂  lanthanide bisphthalocyanine
LnPc'₂  octa-tert-butylbisphthalocyanine of lanthanide element
LuPc₂  lutetium bisphthalocyanine
LuPc'₂  lutetium octa-tert-butylbisphthalocyanine
m  medium intensity
ML  monolayer
N₀  number of molecules spread on subphase
NdPc₂  neodymium octa-tert-butylbisphthalocyanine
NOX  [2NO₂ (gas)→N₂O₄ (gas)]
NO₂  nitrogen dioxide gas
MPc  metallated phthalocyanine
PBDC  N’octyl,N-isobutyl-3,4:9,10-perylene-bisdicarboximide
Pc  phthalocyanine ligand
P-polarization  electric field parallel to the plane of incidence
PP  P-polarized incident field, and P-polarized scattered field
PS  P-polarized incident field, and S-polarized scattered field
PT  P-polarized incident field, and total scattered collection
r  radius of metal particle
rₛ, rₚ  Fresnel coefficient for S-polarized and P-polarized light
RS  Raman spectrum
RRS  resonance Raman spectrum
s  strong intensity

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<table>
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<tr>
<th>Abbreviation</th>
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<tr>
<td>SEM</td>
<td>scanning electron microscopy</td>
</tr>
<tr>
<td>SCE</td>
<td>standard calomel electrode</td>
</tr>
<tr>
<td>SERS</td>
<td>surface enhanced Raman scattering</td>
</tr>
<tr>
<td>SERRS</td>
<td>surface enhanced resonance Raman scattering</td>
</tr>
<tr>
<td>S-polarization</td>
<td>electric field perpendicular to the plane of incidence</td>
</tr>
<tr>
<td>SP</td>
<td>S-polarized incident field, and P-polarized scattered field</td>
</tr>
<tr>
<td>SS</td>
<td>S-polarized incident field, and S-polarized scattered field</td>
</tr>
<tr>
<td>ST</td>
<td>S-polarized incident field, and total scattered collection</td>
</tr>
<tr>
<td>sh</td>
<td>shoulder</td>
</tr>
<tr>
<td>SP</td>
<td>surface plasmon</td>
</tr>
<tr>
<td>t-butyl</td>
<td>tert-butyl</td>
</tr>
<tr>
<td>tb</td>
<td>tert-butyl</td>
</tr>
<tr>
<td>TEM</td>
<td>transmission electron microscopy</td>
</tr>
<tr>
<td>uv-vis</td>
<td>ultraviolet-visible</td>
</tr>
<tr>
<td>v</td>
<td>very</td>
</tr>
<tr>
<td>VOPc</td>
<td>vanadyl phthalocyanine</td>
</tr>
<tr>
<td>VOPcTTb</td>
<td>vanadyl tetra-tert-butylphthalocyanine</td>
</tr>
<tr>
<td>w</td>
<td>weak intensity</td>
</tr>
<tr>
<td>YbPc₂</td>
<td>ytterbium bisphthalocyanine</td>
</tr>
<tr>
<td>ZnPcTTb</td>
<td>zinc tetra-tert-butylphthalocyanine</td>
</tr>
<tr>
<td>λ</td>
<td>wavelength</td>
</tr>
<tr>
<td>Φ</td>
<td>angle of incidence</td>
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CHAPTER 1: Introduction

The intention of this chapter is to provide a general introduction regarding the molecular structure, the properties, and the potential applications of a class of organic molecules known as phthalocyanines (Pcs). They are unique molecules that find their place in many areas of science. Their origin begins in the synthetic laboratory and their science becomes enriched by the study of their structural and physical properties. Notably, the Pc structure resembles that of haemoglobin and chlorophyll.

Phthalocyanine Discovery

The name phthalocyanine, from the Greek naphtha (rock-oil) and cyanide (dark blue) and its structure, was first proposed by Linstead and his students in 1933-1934.\(^4\) They synthesized a new class of highly coloured and unusually stable organic compounds. Metallated phthalocyanines (MPc) were first produced in the reaction of heating o-cyanobenzamide or phthalonitrile with iron, copper, magnesium and other metals. The metal free phthalocyanine or di hydrogen phthalocyanine (\(\text{H}_2\text{Pc}\)) was however first identified in 1907 as a product in the preparation of o-cyanobenzamide from phthalic acid and acetic acid.\(^5\) The first X-Ray crystal studies by Roberston on nickel phthalocyanine, copper phthalocyanine (CuPc) and platinum phthalocyanine showed that the metal and the four isoindole groups were contained in one plane.\(^6,9\)

In air, most Pcs are extremely stable and begin to show signs of degradation at temperatures of 400-500 °C. In vacuum, temperatures in excess of 900 °C are required to decompose Pc\(^10\). The high stability of Pc towards light/heat, acid/alkalies is attributed to the coordination of the metal with the four pyrrole nitrogen atoms and
the conjugated π system.

Applications of Phthalocyanine

Pcs are insoluble in water and organic solvents. This makes them suitable for commercial applications in areas such as electrophotography, dyes, and xerography. In 1987 it was estimated that 16 million pounds of Pc blue toner and two million pounds of green toner were produced in the United States at a commercial value of about ten dollars per pound\textsuperscript{11}. Phthalocyanines have also been used as stains in histology\textsuperscript{12}. The lack of toxicity in Pcs has extended their use as dyes in food colouring, contact lenses and polypropylene sutures. Thin films of H\textsubscript{2}Pc are stable against moderate temperatures, oxidation, humidity and were reported as "write-once" media for optical data storage.\textsuperscript{13} Vanadyl phthalocyanine was also reported as a potential candidate for optical data recording.\textsuperscript{14}

Pc, like many organic pigments, has a small band gap. They have been fabricated into photovoltaic devices where a Pc thin film is commonly sandwiched between two dissimilar electrodes. In the photovoltaic effect light is absorbed by the organic layer, which leads to exciton creation and dissociation at the surface and eventually to charge separation and transport.\textsuperscript{15} The photovoltaic activity in Pc devices has led to potential Pc based solar cells.\textsuperscript{16-21} The photoactivity was influenced by the molecular organization within the film. For example, the crystalline phase of vanadyl \textit{tetr}-butylphthalocyanine, denoted as phase(II), was more effective in photogeneration than phase(I), which was classified as polycrystalline.\textsuperscript{20} Along the same context, sublimed films of titanyl phthalocyanine showed two distinct phases.
Those films with an absorption at 820 nm were better photoconductors than those with the absorption band centred at 720 nm.\textsuperscript{21}

Investigations in Pcs as catalysts for the reduction of molecular oxygen to water have also been reported.\textsuperscript{22} The reaction was highly exothermic and could apply to fuel cells. Cobalt tetrasulfophthalocyanine chemically bound to a titanium dioxide surface has been reported to show photocatalytic activity.\textsuperscript{23} In the latter work, excitation at 366 nm (in the bandgap of titanium) produced hydrogen peroxide in an oxygen saturated solution and hydrogen peroxide depletion in the absence of oxygen. Electroanalytical studies on chemically modified carbon electrodes of cobalt phthalocyanine showed sensitive limits to the detection of sugars(1-5 nmol), oxalic acid(0.3 pmol), and various $\alpha$-keto acids(< 1 nmol).\textsuperscript{24,25}

A recent interest in Pc is in their use as chemical sensors for gases. Chapter 7 is devoted to this area, and our results on the chemical interaction of nitrogen dioxide and various Pc will be presented in that section.

Three books have been written about Pcs which cite the journal and patent literature.\textsuperscript{11,26,27} Recently, another book was published which extensively accounts for the properties and applications of Pc.\textsuperscript{28} The monographs in the latest book are detailed, and suggest that the science of Pc has not been exhausted.

**Molecular Phthalocyanine Structure**

In Figure 1 the structures of $\text{H}_2\text{Pc}$ and $\text{MPc}$ are presented where the basic Pc unit consists of four isoindole groups joined by meso-bridged nitrogen atoms. The aromatic core or macrocycle is composed of 18 $\pi$ electrons which leads to intense
Figure 1.1 Molecular structure of metal free phthalocyanine($H_2Pc$) and metal(M) coordinated phthalocyanine(MPc). The pyrrole nitrogen(1), and meso-bridged nitrogen(2) are also depicted.
$H_2Pc$

$MPc$

Figure 1.1
absorption. A molar absorptivity in the order of $10^4 \text{ (Lcm}^{-1}\text{mol}^{-1})$ is typical for Pc in the red region of the visible absorption spectrum. Substitution on the benzene rings of Pc leads to substituted phthalocyanines and the organic synthesis of these complexes has been reviewed by Leznoff. The metal and the interaction of the $\pi$ system with the substituents on the benzene rings are significant in Pc redox chemistry. For example, in octacyanophthalocyanine complexes of Zn and Cu, the electron withdrawing cyano groups reduce the electron density on the ligand. At present, seventy-six different metals form complexes with Pc. They include Group 1-15, transition elements and all the members in the lanthanide and actinide blocks.

Crystallographic data for most metal phthalocyanines reveal that the macrocycle is planar and that the bond between the carbon and pyrrole nitrogen is larger by (about 0.038 Å in the 6 form of CuPc) than the bond formed between the carbon and meso-bridging nitrogen atom. For CuPc in the 6-phase the mean displacement of the carbon and nitrogen atoms is 0.03 Å above or below the macrocycle. The metal in CuPc is bonded to all four pyrrole nitrogen with a mean bond length of 1.935 Å. The average C-N bond length is 1.347 Å, which gives a bond order of ca. 1.5 and supports the extensive $\pi$ conjugation in the central macrocycle. The cavity formed by the four isoindole nitrogen atoms is approximately a square with a length of 2.7 Å. In $D_{4h}$ complexes of Pc[i.e., MPc(M=Cu, Zn, Fe)], the metal is symmetrically positioned inside the cavity. Cations of Pb and Sn are, however, too large to fit into the cavity. They form four coordinated complexes with C$_{4v}$ molecular symmetry. In passing, it is also
interesting to note that the electrical conductivity in crystals of triclinic PbPc and monoclinic PbPc differed by eight orders of magnitude.\(^{33}\) Five coordinate square pyramidal complexes have also been observed in such molecules as chlorogallium phthalocyanine(CIGaPc), and chloroaluminium phthalocyanine.\(^{34}\) In these complexes, the Group III element was coordinated to the four pyrrole nitrogen and to the chlorine atom. In the square pyramidal complex of vanadyl phthalocyanine(VOPc) the VO\(^{2+}\) cation sits 0.575 Å above the ring and vanadium is coordinated to the pyrrole nitrogens.\(^{35}\) Six coordinated tetragonal complexes have been synthesized. These include molecules like SnPcCl\(_2\) \(^{36}\) and FcCo(picoline)\(_2\) \(^{37}\). The planarity of the macrocycle is destroyed in molecules that show seven and eightfold coordination. In the superphthalocyanine(SPc) as in SPcUO\(_2\) there are five isoindole groups joined by meso bridged nitrogen atoms, and the entire macrocycle is distorted from planarity due to the large angle strain within the molecule.\(^{38}\) The coordination geometry for the uranium atom approximates a pentagonal bipyramid. Bisphthalocyanines have been documented. In these structures, the metal cation is eightfold coordinated to two Pc rings. X-ray crystal data for bis(phthalocyanato)tin(IV)\(^{39}\), bis(phthalocyanato)uranium(IV)\(^{40}\), and bis(phthalocyanato)neodymium(III)\(^{41}\) show that the rings are rotated approximately 45° with respect to one another. The separation between the two rings is about 2.8 Å. All of the lanthanide elements form dimeric phthalocyanine complexes and further discussion is presented in Chapter 6 where our optical spectroscopic data are presented.
Polymorphs of Phthalocyanine

Another well known structural property of Pc in the solid state is polymorphism. It results from the different stacking of the molecules along a certain direction. Polymorphs also have unique physical properties as observed in the different electrical conductivities of triclinic and monoclinic PbPc. It is known that Pcs form columnar stacks with the macrocycle inclined to the axis of the stack. For example, the normal on the Pc molecule forms an angle with the b axis of the crystal of 26.5° and 45.8° in the α and the β form of CuPc respectively.

The polymorphic form is dependent on the method of preparation. The α-form is normally produced by dissolving the Pc complex in concentrated sulfuric acid and precipitating it out with a large amount of cold water. A pressure dependent phase change was reported for thin films of CuPc, where the α-form was produced at sublimation pressures up to 13.3 Pa, while the β-phase required higher sublimation pressure. Polymorphs of Pc also give different X-ray diffraction patterns, infrared(IR) and visible absorption spectra. Their solution spectra are, however, identical.

In high vacuum(10⁻⁷ -10⁻⁸ Pa), the α-form is produced for thin films of MPc prepared at room temperature. Conversion from α→β form required heating to temperatures of 300 °C. Mindorff et al. reported that annealing the α form of H₂Pc to produce the β form was, however, irreversible. Vapours of alcohol can also produce the α→β transformation in ZnPc. Sidorov and Kotlyar observed new IR bands at 780, 877, 1100 and 1173 cm⁻¹ in the spectra of the β-form of MPc.
Spectral characterization of $\alpha$, $\beta$, $\gamma$ (produced by milling the $\alpha$-form) polymorphs of $\text{H}_2\text{Pc}$ showed distinct differences in the visible, near-infrared and infrared spectra. Changes in the infrared were observed in the 700-800 cm$^{-1}$ and in the N-H region. The red shift in the N-H stretching frequency observed only in the $\beta$ form was attributed to intermolecular hydrogen bonding which could weaken the N-H stretching bond.

Electronical Spectra of Phthalocyanine

A recent overview of the optical properties of Pc has been published by Stillman and Nyokong. The absorption spectra of Pc derivatives have received considerable attention. The vapour absorption or "unperturbed" spectra in a series of MPc contained 5 characteristic bands denoted as: Q(ca.660 nm), B(ca.320 nm), N(ca. 275 nm), L(ca. 245 nm) and C(ca. 210nm). The N, L, and C bands are not always resolved in the solution spectra. The Q and B(or Soret) bands are the main absorption bands in Pc and arise from as $\pi-\pi^*$ ($a_{1u} \rightarrow e_g$ in $D_{4h}$) and $\pi-\pi^*$ ($a_{2u} \rightarrow e_g$ in $D_{4h}$) transitions respectively. The energy diagram for CuPc($D_{4h}$ point symmetry) is shown in Figure 1.2. Their assignment is mostly based on the theoretical and experimental work by Gouterman and coworkers. In $D_{4h}$, the $a_{1u}$ is the highest occupied molecular orbital(HOMO) and the $e_g$ is the lowest unoccupied molecular orbital(LUMO).

They classify the absorption spectra of Pc derivatives to one of five groups. The first type of spectrum is denoted to the distinct absorption features seen in metal free phthalocyanine($D_{4h}$) or its derivatives. For these Pcs, the Q-band is seen as two intense bands($Q_x$ and $Q_y$). The two weaker absorptions which are also observed in the
Figure 1.2    Molecular orbitals of CuPc. Taken from Reference 51.
Energy Diagram of CuPc

\[ \uparrow \quad b_{1g} \quad (d_{z^2}) \]

\[ \downarrow \quad e_g \quad (\pi^*) \]

\[ \uparrow \downarrow \quad a_{1g} \quad (d_{z^2}) \]

\[ \uparrow \downarrow \quad a_{1u} \quad (\pi) \]

\[ \uparrow \downarrow \quad e_g \quad (d_{xz}, d_{yz}) \]

\[ \uparrow \downarrow \quad b_{2g} \quad (d_{xy}) \]

\[ \uparrow \downarrow \quad a_{2u} \quad (\pi) \]

Pc MO's

metal d orbitals

Figure 1.2
Q-band envelope were considered as vibrational overtones of the $Q_x$ and $Q_y$. The two component Q band is attributed to the lower symmetry in $D_{2h}$, i.e., to the splitting of the degenerate $e_g$ when the symmetry is lowered from $D_{4h} \rightarrow D_{2h}$. Distinct transition moments were observed by Lyons et al. in the polarized absorption spectrum of $H_2Pc$ crystals. Deprotonation in metal free phthalocyanine gives an absorption spectrum similar to that found in Pc complexes of $D_{4h}$ symmetry.

The second type of absorption spectra is given to Pc complexes with $D_{4h}$ symmetry in which only π-π*(B, and Q bands) are observed. The spectra may contain a weak vibronic band{Q(0,1)} at the high energy end of the Q band. The third type of spectra contains an additional band (assigned to a charge transfer transition) in between the Soret and Q-band. As an example, reduction of CoPc with borohydride gave a new band at 467 nm, which was assigned to a metal-ligand charge transfer band based on the magnetic circular dichroism spectrum. The fourth type of spectra is assigned to Pc complexes that show dimerization or higher order associations in solution.

This phenomenon is characterized by a new band appearing near the high energy side of the Q-band as the dye concentration is increased. Aggregation can be diminished in solvents that give better dye solubility since greater solvation of the monomer would reduce the dye-dye interactions. Alternatively, solvents with high dielectric constants were reported to reduce the dye-dye interaction. For example, the dimerization constant of 4,4',4''',4'''-tetraoctadecylsulfonamidophthalocyaninecopper(II)(CuPc-$X_4$) in benzene was smaller
than in carbon tetrachloride,\textsuperscript{57} and the dimerization constant of CuPc-X\textsubscript{4} was lower in tetrahydrofuran than in benzene.\textsuperscript{58} The degree of association was also shown to be dependent on the metal cation (M=Mg, Co, Zn, Bi, Cu, Ni, Pd, Pt) in a series of tetrakis(cumylphenoxy)-phthalocyanine complexes.\textsuperscript{59}

In dimeric silicon phthalocyanine, the Q and B bands are blue shifted with respect to the monomer.\textsuperscript{60} The shift in the electronic bands was attributed to the exchange interaction (or exciton coupling) of the transition moment of one ring with the transition moment on the other ring. In the dipole approximation, the coupling is proportional to the dot product of the transition moments of the two chromophores and inversely proportional to the cube of the separation distance.\textsuperscript{61} For \textit{translational equivalent} molecules, the exciton theory predicts a displacement of the absorption band relative to that of the free molecule. A blue-shifted and a red-shifted band are expected for parallel and in-line transition moments respectively.\textsuperscript{62} For \textit{translational inequivalent} molecules exciton coupling leads to a splitting of the optical transition, known as Davydov splitting. The magnitude of the splitting is a measure of the interaction energy. For molecules with molar extinction coefficients larger than \(10^4\text{Lcm}^{-1}\text{mol}^{-1}\) Davydov splitting in the order of \(10^3-10^6\ \text{cm}^{-1}\) may be expected and may account for band broadening, especially in the solid state.\textsuperscript{61} Lever et al. have reported exciton splitting of the order of 2300 cm\textsuperscript{-1} in metal free binuclear phthalocyanines.\textsuperscript{54}

The last type of absorption spectra pertains to Pc in the solid state, where the normally narrow absorption bands in the solution spectra are seen as broad bands.
Davidson noted that in MPc(M = H₂, Mg, Fe, Co, Cu, Zn), prepared as thin solid films, the splitting and broadening of both Q and Soret bands could be attributed to a vibronic effect; however, Davydov splitting was not ruled out. This report also showed that the N band was the most sensitive to metal substitution. Thin film preparation also has been reported to affect the spectral position and shape of the Q Band. For example, in thin films of ClGaPc, the prismatic block phase was grown under slow rates of evaporation and the platelet block phase was formed at higher evaporation rates.

Outline of Thesis

The work that remains is divided into chapters. Chapter 2 describes the experimental techniques and the instrumentation used to prepare Pc into thin solid films and Langmuir-Blodgett(LB) films. In this thesis, normal Raman scattering, surface enhanced Raman scattering(SERS), infrared, and ultraviolet-visible spectra were used to characterize Pc molecules. The majority of the work was related to the characterization of phthalocyanine LB films using SERS. A small review of SERS theory is given in Chapter 3. It accounts for the enhancing properties of the SERS substrate. Chapter 4 begins with a review of the Raman spectra of Pc. The vibrational characterization is presented for vanadyl phthalocyanine thin films adsorbed onto a halide crystal and reflecting substrates. Infrared and SERS data for vanadyl tetra-tert-butylphthalocyanine and zinc tetra-tert-butylphthalocyanine is also discussed in Chapter 4. The chapter closes with the observation of energy transfer between one LB monolayer of a perylene derivative and one LB layer of a
phthalocyanine complex transferred to a SERS substrate. The effect of surface coverage on surface enhanced resonance Raman scattering of vanadyl tetra-tert-butylphthalocyanine is addressed in chapter 5. Here the LB technique was used to control surface coverage. Chapter 6 is dedicated to the vibrational characterization and electronic spectra of the blue and the green forms of rare earth bisphthalocyanines. An application of SERS is described in Chapter 7, where the high sensitivity of SERS was used to monitor the reversible adsorption of nitrogen dioxide on a single monolayer of various Pc derivatives. The general conclusions of this thesis follow in Chapter 7.
CHAPTER 2: Instrumentation and Experimental Techniques

Vacuum Evaporator Apparatus

Thin solid films of P, substrates for Surface Enhanced Raman Scattering (SERS), and smooth metal films were prepared in traditional and "in house built" vacuum evaporation systems\(^1\)\(^2\) composed of some factory purchased electronic components. The commercial components used to monitor pressure include Balzers IKR 020 cold cathode gauge, Balzers TPR 010 Parani gauge, and Balzer pressure gauge control. The power source used to control the current to the boat for resistive heating was the Balzers BSV 080 glow discharge/evaporation control unit. Edwards E2M2/E2M5 rotary vacuum pumps and an Edwards diffusion pump were employed to obtain high vacuum. Film thickness and the rate of evaporation were monitored by an Inficon XTC single film monitor/controller with a standard 6 MHz gold coated quartz sensor. Film thickness is reported as mass thickness.

The basic working principle behind the apparatus was to convert the material to the gaseous phase by supplying heat of vaporization, and to control the rate at which the gaseous atoms or molecules condense onto the substrate. In the high vacuum regime\((10^{-4}-10^{-3} \text{ Pa})\), the mean free path of particles was large enough to ensure that the evaporant moved to the substrate. High vacuum also reduced the possibility of contamination to the substrate by residual gases. The rate of evaporation and the thickness of the film were carefully monitored in order to reproduce the desired film. These parameters were important to duplicate the SERS substrates with a near constant plasmon absorption profile. For metal films the rate of
evaporation was easily controlled ca. ±0.1 nm/s. The rate of sublimation for Pc molecules was however more difficult. Even after degassing for 10-15 minutes, the best controllable rate was ca. ±0.5 nm/s for Pc derivatives. Thin films of Pc and metal films were prepared in separate systems to avoid contamination to the SERS substrates.

The main components in the evaporator are illustrated in Figure 2.1. The baffle valve separated the working chamber from the high vacuum compartment. The high vacuum chamber was composed of a liquid nitrogen cold trap and a diffusion pump with Santovac 5 diffusion pump oil. Valves(V) were used to isolate and evacuate the working and high vacuum chambers. The pressure gauge control module recorded the pressure at gauges P1 (Parani) and P2 (cold cathode). In our evaporator, high vacuum of 2x10^-4 Pa was obtained within one hour of operation. The sample holder was a modified heater that was attached to the top flange of the evaporator. A VARIAC was used to control the voltage to the heater (maximum temperature 250 °C) when the substrate was to be heated. A type K (NiCr-NiAl) thermocouple was positioned permanently within the heater and near the surface of the substrate. It was used to record the temperature at the substrate. The manually operated shutter masked the substrate from the evaporant but not the crystal oscillator. This arrangement allowed the control of the rate of film growth before opening the shutter and depositing the material on the substrate. A boat was used to contain the material that was to be deposited as a film. It was fastened to a set of electrodes or posts. The posts were also connected by a set of cables to the high
Figure 2.1  Schematic of vacuum evaporator

substrate holder(1), crystal oscillator sensor(2), shutter(3),

boat(4), posts(5), cold trap(6), diffusion pump(7), baffle valve(8)

valves(V), Parani vacuum gauge(P_1), cold cathode gauge(P_2)
power source or transformer, which delivered current to the boat for resistive heating. The transformer delivered high current by stepping down the voltage (4, 8, or 16V) in the secondary. The rate of evaporation/sublimation was controlled by adjusting the amount of current that flowed through the boat. In the preparation of metal films, the XTC Inficon controller was programmed with the constants provided by the manufacturer. In the fabrication of organic thin films of Pc a Z-ratio of unity, and a sample density of 1.5 g/cm³ were selected.

Calibration of film thickness was periodically checked by the procedure outlined in the XTC Technical Manual. The film thickness calibration correction would be in terms of the tooling factor which required the determination of film thickness by an alternative method. In our case, thick films of Ag (ca. 150 nm) or Au were sent to the Xerox Research Centre of Canada where the thickness was determined by the stylus method.

All boats used in the vacuum evaporation apparatus were purchased from R. D. Mathis Company (Long Beach California). Tantalum boats (ME10-.005 Ta) were used for the sublimation of Pc materials. In evaporation of silver and tin, tantalum boats(S9A-.005 Ta) were utilized; however evaporation of gold required tungsten boats(S9A-.005 W). Both silver and tin shots (99.999% purity) were obtained from Johnson and Matthey Aesar Group (Toronto). A gold (99.99%) rod was purchased from Aldrich. These metals were used in the preparation of SERS substrates and smooth metal films.

Thin films of Pc were formed onto Corning 7059 glass slides and halide
crystals suitable for transmission infrared spectroscopy. The glass substrates were used as supplied by Corning (Danville, Virginia). Thin solid films of phthalocyanine were made from i) VOPc, tetra-tert-butylPcM(M = VO, Zn), LnPc₂(Ln = Yb, Lu) and ii) LnPc₃(Ln = Pr, Nd, Sm, Eu, Gd, Tb, Dy, Ho, Er, Tm). Samples denoted i) were supplied by Dr. Kovacs, and Dr. R. O. Loutfy from the Xerox Research Centre of Canada. Those designated ii) were supplied by Dr. A. Y., Tsivadze at the Institute of General and Inorganic Chemistry Academy of Science, Moscow.

**Langmuir Trough and Langmuir-Blodgett Films**

**Introduction**

Irving Langmuir is mostly credited for the work on insoluble monomolecular films at the liquid-gas interface. Among other findings, he showed that the cross-sectional area (~21-25 nm²) of palmitic acid, stearic acid and cerotic acid were very similar regardless of the length of the hydrocarbon chain. His findings confirmed that a monomolecular film was formed on the subphase and that the carboxyl groups were in contact with the subphase, while the hydrocarbon tails were packed vertically above the subphase. Katherine Blodgett, who was Langmuir’s coworker, reported the first successful transfer of a monomolecular film of stearic acid and calcium stearate to a glass slide. The transfer of the Langmuir monolayer at the liquid-gas interface to a solid support is known as the Langmuir-Blodgett or LB technique. A detailed description of Langmuir monolayers and LB films is found in the following sections. A new book devoted to LB films was recently written. It accounts for the renewed interest in the field and includes chapters devoted to monolayer properties and their
molecular structure, LB film deposition, LB characterization and properties, spectroscopy of complex monolayers, multilayers and monolayers of biomolecules, and potential applications of LB films.\textsuperscript{13}

Researchers are attracted to LB films because film thickness and molecular orientation can be precisely controlled. For example Kuhn, studied the dependence of energy transfer between acceptor and donor molecules separated by monolayers of arachidic acid of known thickness.\textsuperscript{14} Others may have interests in LB forming materials because they can be transferred at near ambient temperature and pressure, unlike the conditions used to form films by evaporation or sublimation. It is believed that size reduction offered by the LB technique could lead to an era of organic supermolecular electronics.\textsuperscript{15} Studies on the electrical conductivity in LB films showed that electrical conduction is related to the structure and to the arrangement of molecules.\textsuperscript{15} It is expected that ultra thin films of organic molecules should respond more positively to external stimuli than conventional evaporated thin films. For example, diffusion transport in films is thickness dependent and it is remarkably fast at molecular thicknesses.\textsuperscript{16} The field of LB films is also interdisciplinary and involves elements of physical chemistry, physics, chemical engineering and material science. As an example, our group has collaborations with industrial and academic institutes of organic chemistry who synthesize Pc and perylene derivatives for possible applications in LB film technology. Our current LB research interests are in the preparation of LB films of phthalocyanine and perylene derivatives, and their optical spectroscopic characterization by surface enhanced Raman scattering. Swalen has reviewed various
other optical techniques that can be used to study LB films.\textsuperscript{17,18}

As the number of novel molecules (like Pc) increase in LB technology, further research will be required in the areas of substrate selectivity, interpretation of the isotherm, optimal conditions for good monolayer transfer, and the evaluation of LB structure on the solid support. Some of the latest research in LB films of mostly fatty acids has certainly revealed important results. For example, investigation in LB of cadmium stearate and stearyl alcohol has shown that the transfer ratio was heavily dependent on the contact angle and the surface pressure.\textsuperscript{19} Defects with a diameter of 10 nm were observed in cadmium arachidate LB films by the atomic mass microscope.\textsuperscript{20} Leisieur et al. have noted that defects might be important for diffusion processes.\textsuperscript{21} Another important proposal was that the orientation of docosanoic acid on the subphase was not preserved during transfer to the formvar coated copper grid.\textsuperscript{22}

**LB Instrumentation**

The main components of the LB trough were purchased from Brinkmann Instruments. They include a film balance or Langmuir trough and a central unit that controls the velocity and direction of the moving barrier. The barriers and trough are coated with Teflon. A Lauda Filmlift FL-1 film deposition device is simply a programable stepper motor that allows the substrate to enter and leave smoothly the liquid-gas interface during the transfer of the Langmuir monolayer. The temperature of the subphase was maintained with the Lauda refrigerating model RM6 circulator. In our set up the film balance rests on stone plate placed on the top of a sand filled table (similar to that used to support a micro analytical balance). The heavy table
Figure 2.2 Schematic illustration of the Langmuir film balance and other LB accessories
Figure 2.2
adequately damped the natural vibrations within the room. The entire trough was protected from airborne contaminants by a Plexiglass box. Figure 2.2 illustrates the basic components of the Langmuir-Blodgett trough.

The film balance, control unit, and XY recorder are used to determine experimentally the isotherm of surface pressure(\( \Pi \)) versus molecular area(\( A \)) of the Langmuir film. The moving barrier can move either towards(compression) or away(expansion) from the fixed barrier. The surface pressure was measured by the electronic displacement transducer on the measuring barrier and fed into the Y channel of the HP 7015B XY recorder. The signal generating the area/molecule was fed into the X channel of the recorder. The Lauda Filmlift FL-1 device was used to secure the substrate and move it up and down through the liquid-gas interface in the region of the well. The substrate was fastened to the magnetic arm of the filmlift by a metallic clip.

**Langmuir Monolayers and Langmuir-Blodgett Technique**

In the "classical" formation of a Langmuir monolayer, the *amphiphile* is dissolved in a volatile organic solution and a known number of molecules are spread with a syringe onto a aqueous subphase in the region enclosing the moving and the measuring barrier. The amphiphile is a long chain fatty acid which contains a hydrophilic(carboxylic acid) group and a hydrophobic(saturated hydrocarbon) part. The subphase can affect the properties of amphiphile, and the effect can be observed in the isotherm. For example pH, ions, and temperature dependence were observed
in the II-A measurements of heneicosanic acid\textsuperscript{23}, behenic acid\textsuperscript{24} and pentadecanoic acid\textsuperscript{25}. To the right of the measuring barrier the subphase is clean and to the left of the barrier the subphase contains the floating amphiphiles. The molecules are mechanically compressed by advancing the moving barrier towards the measuring barrier. The rate of compression is described by units of (area/number of molecules)/time). The area(in nm\textsuperscript{2}) and time(in s) quantities are respectively the total area and the time the moving barrier takes to sweep through a designated surface area on the subphase. In our system the moving barrier sweeps through an area of 562 X10\textsuperscript{14} nm\textsuperscript{2} during π-A measurements. The measuring barrier records surface pressure(Π) as the difference in surface tension:

\[ \Pi = \gamma_0 - \gamma \]  \hspace{1cm} (2.1)

where \( \gamma_0 \) is the surface tension of the subphase and \( \gamma \) is the surface tension in the presence of the amphiphile.

At the surface pressure of 0 mN/m(milli-Newton meter\textsuperscript{-1}) the amphiphiles are independent of each other and their orientation is determined by their interaction with the subphase. At low surface pressure the molecules at the liquid-gas interface may be modelled as a two dimensional gas.\textsuperscript{8} As the molecules are compressed together, they form a monomolecular film that passes through phase transitions(i.e., gas→liquid→solid). The liquid phase of fatty acids and phospholipids can be further characterized by liquid-expanded and liquid-condensed phases.\textsuperscript{24} Phase transitions in the isotherm are seen as
plateaus at different surface pressures. Fluorescence probes and fluorescence imaging microscopy have been used to study phase behaviour in monolayers. X-ray reflectivity data for a monolayer of arachidic acid on the subphase has shown that as the monolayer was compressed from the liquid→solid phase, the tail of the fatty acid formed a tilt angle (with respect to the surface normal) which varied from 30° to 0°. In the solid phase the tails were all parallel and vertically upwards.

The monolayer can be compressed until the collapse pressure where the molecules in the solid phase are expelled from the two dimensional arrangement and forced into a three dimensional phase. The collapse pressure is revealed in the isotherm when the surface pressure suddenly drops. Transfer of the Langmuir layer to a solid support is made at constant surface pressure well below the collapse pressure.

The last step in the LB technique is the transfer of the monolayer (held at constant surface pressure) from the liquid-gas interface to a solid substrate. A feedback loop between the surface pressure transducer and the moving barrier maintains the monolayer at constant surface pressure. The transfer cycle may begin with the substrate in air or in the subphase. In the latter case the substrate is lowered into the subphase before monolayer spreading and compression. The monolayer is transferred to a substrate by slowly (a few mm/s) raising and lowering the substrate vertically through the liquid-gas interface. The Langmuir layer results in X, Y, or Z deposition. The notation X, and Z are used to describe the transfer of the Langmuir layer during the downstroke and upstroke of the substrate respectively. In Y deposition, the monolayer is deposited as the substrate is lowered into and raised from the subphase. As each monolayer is
Figure 2.3  Schematic of the Langmuir-Blodgett film processes.
Langmuir-Blodgett (LB) Film Making

![Diagram of molecular transfer and compression]

- **a)** Transfer of Molecules to subphase
- **b)** Compression of Molecules
- **c)** LB Film Transfer
  
  \((\text{X,Y,Z})\) Deposition

*Figure 2.3*
transferred to the substrate the film thickness should increase linearly. This condition can be checked with optical measurements such as ellipsometry, and absorption spectroscopy (UV-Visible, Infrared). Figure 2.3 summarizes the basic steps in LB film making.

**Calibration of the Film Balance for Isotherm Data Acquisition**

With the trough covered with the subphase, the surface pressure or Y-axis of the isotherm is calibrated by simulating a surface pressure ($\Pi = mg/l_{ef}$) along the measuring barrier with a calibration weight of mass ($m$). The quantity $g$ and $l_{ef}$ refer to the acceleration due to gravity and the effective length of the measuring barrier. In our system, the calibration weight is used to simulate a surface pressure of 37.7 mN/m. The F-scale potentiometer on the control unit is used to bring the signal generated by the calibration weight to 37.7 surface pressure units. Zero surface pressure is manually set by taring the signal of the Y-channel. The surface pressure transducer in the Lauda trough is capable of recording surface pressure in the range of 0-100 mN/m.

The calibration of the area per molecule or X-axis of the isotherm is based on the calibration area of $5.62 \times 10^{16}$ nm$^2$ and $8.64 \times 10^{16}$ molecules (i.e., A-scale = 750). The A-scale is used to enter the number of spread molecules on the subphase. When the moving barrier is in the fully expanded position, the largest area per molecule is 0.65 nm$^2$/molecule (i.e., $5.62 \times 10^{16}$ nm$^2$ / $8.64 \times 10^{16}$ molecule). The moving barrier does not go to 0 area since it will damage the measuring barrier. The closest distance between the moving barrier and the measuring barrier is 0.5 mm. At this position the area enclosed
between the two barriers is $1.07 \times 10^{14}$ nm$^2$. Based on $8.64 \times 10^{14}$ molecules this area marks .0124 nm$^2$/molecule (i.e. 0.65 *1.07/562) position. The zero adjustment on the recorder is used to set the .0124 nm$^2$/molecule mark and the A-corrector potentiometer on the control unit is used to set the 0.65 nm$^2$ mark. Lauda conveniently supplies isotherm chart paper whose X-axis is divided into 65 scale units. Trough calibration is only required for Π-A measurements.

When the trough is calibrated, a predetermined number of molecules can be spread onto the subphase for isotherm determination. With the moving barrier in the fully expanded position, the number of molecules spread onto the subphase is observed and fed into the A-scale potentiometer. Lauda supplies tables that list the A-scale setting corresponding to the number of molecules delivered to the subphase. Five scales for area/molecule are supplied by Lauda to accommodate small and large molecules. The possible scales (in units of nm$^2$) are: a) 0→0.26, b) 0→0.65, c) 0→1.625, d) 0→3.25, and e) 0→6.50. In Figure 2.4 the experimental isotherm of arachidic acid is shown. The arachidic acid molecules were spread from a chloroform solution onto the subphase (pH = 5.5) maintained at 20 °C. The three phases are also shown where the gas, liquid and solid states are characterized when the surface pressure is in the region, 0, 0-25, 25-50 mN/m respectively. Extrapolation of solid phase to Π=0 mN/m gave a limiting area of 0.22 nm$^2$ which corresponds to the cross-sectional area of the carboxylic group. The results and the profile of the isotherm were in excellent agreement with other fatty acids like stearic acid for which a limiting area of 0.22 nm$^2$ was reported.$^{27}$
Figure 2.4  Surface pressure versus molecular area isotherm for arachidic acid measured in our laboratory.
Figure 2.4
In House Fabrication of Langmuir-Blodgett Monolayers

This section describes the general method and the materials used to form monolayers and their transfer to solid supports.

Materials

Phthalocyanine, and perylene derivates were spread from chloroform, benzene or toluene. The solvents were HPLC grade and obtained from Aldrich. Underivatized Pcs are insoluble in organic solvents and can not be spread onto the subphase. The solubility of tetra-tert-butyl substituted phthalocyanines and rare earth bisphthalocyanines is, however, adequate and these molecules were transferred as monolayers and multilayers by the LB technique. In this work LB films and spectroscopic studies of i) tetra-tert-butylPcM(M=2H, Zn, VO), LnPc2(Ln=Yb, Lu), N-octyl,N-isobutyl-3,4:9,10-perylene-bis-dicarboximide(PBDC), ii) tetra-tert-butylPcM(M=Cu), LnL2 L=tetra-tert-butylPcLn=(lanthanide element), were considered. Purified compounds designated i)→ii) were supplied by Dr. G. J. Kovacs and Dr. R. O. Loutfy (Xerox Research Centre of Canada), and Dr. E. A. Luk’yamets and Dr. L. Tomilova(Institute of Organic Intermediates and Dyes, Moscow, USSR), respectively. The tetra-tert-butylPc derivatives were easily dissolved in organic solvents, and solutions in the concentration range of 2-3x10^{-4} M were used. The LnPc2 molecules however have limited solubility and solutions of 0.5-1 x10^{-4} M were prepared by sonication. Arachidic acid was purchased from Aldrich.

Trough Cleaning

The trough was thoroughly cleaned prior to work. The water used to clean the trough was of the same grade as that used in monolayer fabrication. The water was double
distilled (the second distillation was from a Barnstead Sybron unit) and filtered by a Milli-Q Plus (Millipore) portable unit. The resistivity of the water measured by the Milli-Q Plus unit was 18 MΩcm. The pH of the water was 5.5 not adjusted.

The cleaning procedure was extensive and involved wiping the Teflon coated trough with a Kimwipe or lens paper wetted with Aldrich HPLC grade acetone. Conrad 70 (Canlab) soap was mixed with cleaning water and poured into the trough. The moving barrier was activated and the soapy solution was forced back and forth to aid in the dissolution of possible contaminants. The cleaning solution was then suctioned off with an electrical vacuum pump. The soap was completely rinsed from the trough by repeating the cycle of adding clean water and removing it with the vacuum pump.

Following cleaning, the subphase was delivered to the trough and the desired subphase temperature (± 0.5°C) was maintained by the refrigerating circulator unit. Prior to monolayer spreading, the moving barrier was swept from the fully compressed limit to the expanded limit. This mechanical movement was used to clean the liquid-gas interface before spreading the monolayer.

**Preparation of Monolayers**

The procedure for monolayer spreading and compression was similar for all molecules. The molecules were dissolved in either chloroform, toluene or benzene and a volume of solution was spread with a 0.1 ml Hamilton syringe. The volume of solution was dependent on the molar concentration and the number of molecules that were to be delivered to the subphase. For isotherm acquisition the exact number of molecules was predetermined. The number of molecules spread to form a monolayer to be transferred
as LB film was, however, based on the dimensions of the substrate and the compressibility of the molecule. This was a trial and error process and the goal was to form a monolayer (at constant surface pressure) with enough surface area that would facilitate transfer as an LB film.

The compression rate ([5.62×10^{16} \text{ nm}^2/\text{N}_0]/\text{time}) during isotherm acquisition was dependent on the number of molecules (N₀) and the time (in s) necessary for the moving barrier to sweep through the area of 562×10^{14} \text{ nm}^2. The time element was dependent of the velocity of the moving barrier. For LB films, the trough was operated in constant surface pressure mode. Here the moving barrier would advance until the desired surface pressure was reached. The monolayer was maintained for approximately 30 minutes at constant surface pressure before it was transferred to the solid support. In all cases the Lauda Filmlift FL-1 device was set to transfer the substrate at a rate of 4.8 \text{ mm/min}. For LB multilayers a pause time of three minutes elapsed before the substrate passed through the upstroke/downstroke cycle.

**Langmuir-Blodgett Substrates**

Monolayers were transferred to Corning 7059 glass slides and Surface Enhanced Raman Scattering (SERS) substrates. The Corning slides were cleaned when monolayers were transferred to glass by the method described by Möbius and Bücher. The glass slides were placed in hot chromosulfuric acid maintained at 110 °C. After two hours in the cleaning solution, the slides were copiously rinsed with double distilled Milli Q water and placed in a NaOH (Aldrich) solution (pH 11) for 12 hours. The resulting slides were
Figure 2.5  SERS substrates, SEM of Ag-Sn spheres(A), TEM of 10 nm Ag(B)
and 4 nm Au(C)
Figure 2.5
hydrophilic. A hydrophobic surface was achieved by further treating the slides in a 5\%(w/v) solution of dichlorodimethylsilane(Aldrich) in chloroform for fifteen minutes. These slides were then rinsed in methanol and finally water.

Three different SERS substrates were formed by evaporating high purity metal onto untreated Corning 7059 glass slides. Silver and gold island films were formed by evaporating 10 nm Ag or 4 nm Au at a rate of 0.1 nm/s onto a glass slide maintained at 220 °C. The SERS substrate was allowed to return to room temperature before removing it from the evaporator. The third SERS substrate is designated as silver coated tin spheres(Ag-Sn spheres), and the preparation involved a two step process. In the first step 100 nm of Sn was evaporated at a rate of 0.5 nm/s onto a glass substrate maintained at 120 °C. Murr has shown that Sn formed hemispherical island films when the temperature of the substrate was kept between 81-131 °C. In the second step, the substrate is allowed to cool to room temperature, and 100 nm of Ag is evaporated at a rate of 0.5 nm/s onto the tin spheres.

Surface morphology of the SERS substrates was observed with the electron microscope. Transmission micrographs of Ag and Au islands were recorded on a JEOL CX100 EM operated at 100 KV. These samples were prepared by securing a type-B carbon substrate(J.B. EM Services Inc., Quebec) onto the glass substrate before metal evaporation. A scanning electron microscope(SEM) was, however, used to record the surface features of Ag-Sn spheres. The substrate in this case was directly observed with the Nanolab7 SEM operating at 15 KV. The electron micrographs from the metal substrates are given in Figure 2.5.
Figure 2.6  Absorption spectra of Au and Ag island films
Figure 2.6
Figure 2.7  Absorption spectrum of Ag-Sn spheres. The spectrum was taken in reflection mode.
Figure 2.7
Figure 2.6 represents typical absorption spectrum taken from the island films. The 488 nm and 514.5 nm laser lines(LL) from the Ar+ laser were in full resonance with the plasmon absorption of the Ag island film. The Au island film, however, had an absorption in the red spectral region and excitation with the 647.1 nm laser line of the Kr+ laser was required for surface plasmon resonance. The plasmon absorption profile for Ag-Sn spheres is shown in Figure 2.7. All three laser lines can be used to excite plasmon resonances in this structure; since the broad plasmon absorption encompasses both blue and red spectral regions.

Raman Instrumentation

Raman data were collected by a Spex 1403 spectrometer interfaced to an IBM computer(XT). The spectrometer was situated on an optical table. The software was written in our laboratory and has been described elsewhere. The program "Raman" was used to move the pair of holographic gratings(1800 grooves/mm) and register the inelastic scattering. Photons were detected by a thermoelectrical and water cooled Hamamatsu R928 multalkali photomultiplier and observed with conventional photon counting electronics. The program "Raman" generated a data file that could be retrieved by a second program "Ramplot" to manipulate and plot the Raman spectrum. Spectra Calc software produced by Galactic Industries is currently being used instead of Ramplot.

The 488 and 514.5 nm radiation from a Spectra-Physics model 164 Ar+ laser and the 647.1 nm and 676.4 nm lines from a Spectra-Physics model 2020 Kr+ were used for Raman excitation. Optical filters supplied from Spectra-Physics were used to remove
plasma lines. The incident laser light was plane polarized by a polarization rotator (Spectra-Physics model 310-21). Scattered light was passed through a scrambler before passing through the entrance slit of the spectrometer to ensure similar sensitivities towards light polarized in different directions. Components of scattered Raman light could also be selected by placing a polarization analyzer in between the collecting optics and the scrambler.

The 90° collection geometry used in the Raman experiments is shown in Figure 2.8. Porto's notation was used to describe the direction and polarization of incident and scattered electric field vectors. In our laboratory, illumination was along the positive Z-axis, observation was along the X-axis and the sample was placed along the Y-axis. The plane of incidence was the ZX plane. Incident light and scattered light was polarized either parallel (P) or perpendicular (S) to the incident plane. The analyzer was rotated to collect S or P scattered light. Four combinations were therefore possible and they are written as:

\[
\begin{align*}
Z(YY)X &= SS \\
Z(YZ)X &= SP \\
Z(XY)X &= PS \\
Z(XZ)X &= PP
\end{align*}
\]  
(2.2)

The symbols outside the parenthesis are the propagation directions for incident light (i.e., Z) and scattered light (i.e., X). The first and second terms in parenthesis refer to the polarization of the electric field of incident (i.e., X, Y) and scattered (i.e., Y, Z) light respectively. The notations SS, SP, PP, PS are equivalent to the Cartesian description of incident polarized light (first letter) and scattered polarized
Figure 2.8 A schematic representation of the 90° scattering geometry.

The film is illuminated at 60° incident angle. Polarization of incident and scattered light leads to four polarization combinations (SS, SP, PP, PS). The four letter notation used by Porto (Bottom)
Porto’s Notation

\[Z(YY)X = SS\]
\[Z(YZ)X = SP\]
\[Z(XY)X = PS\]
\[Z(XZ)X = PP\]
light(second letter). In the special case where the crystal is orientated so that its axes are collinear with the space fixed axis, the element of the polarizability tensor is the quantity in parenthesis described by the Porto notation. As an example, the SS spectrum would be dominated by bands that contain the $a_{yy}$ Raman component. When the analyzer was not used, total(T) polarization was collected and the polarization in the Raman spectra was described as ST or PT.

The depolarization($\rho$) ratio$^{32,33}$ of the 218(E) cm$^{-1}$ or 314(F$_{2}$) cm$^{-1}$ bands of CCl$_{4}$ was used to calibrate S and P electric fields. It can be shown the $\rho = I_{SP}/I_{SS}$, where $I$ is the integrated band intensity in the SS and SP spectra. A value of $\rho = 0.75$ was obtained for the non totally symmetric bands of CCl$_{4}$ with proper calibration. The CCl$_{4}$(Aldrich spectroscopic grade) was contained in a quartz cell and used daily for frequency calibration of the spectrometer. Occasionally the accuracy of the spectrometer was inspected with a capillary of distilled indene(Aldrich).

An XYX-positioner was used to secure and align the sample. A cold cell, or a spinning cell could be attached to the XYX-positioner. The spinning cell was always used for LB samples to prevent thermal decomposition. The typical spectral bandpass and laser power(at the sample) were 5 cm$^{-1}$ and 50 mW respectively.

**Fourier Transform Infrared(FTIR) Instrumentation**

Transmission spectra were collected with a Nicolet 5DX FTIR and Bomem DA3 FTIR spectrophotometer. The Bomem files were imported to Spectra Calc software for spectral manipulations. Infrared spectra were routinely recorded in the region of 400-4000 cm$^{-1}$. 
UV-VIS Absorption Instrumentation

Ultraviolet-visible (uv-vis) absorption spectra were recorded on a HP 8450 diode array and Response spectrophotometer. The files generated by the Response instrument were imported to Spectra Calc for spectral manipulations.

Electrochemistry Instrumentation

Spectroelectrochemical data were obtained with a PAR Model 175 universal programmer and a PAR Model 174A polarographic analyzer. During electrochemical cycling, uv-vis spectra were obtained and stored every 1.2 s on an HP Model 8450A uv-vis spectrometer. This work was conducted by Carol Jennings at Xerox Research Centre of Canada.
CHAPTER 3: Raman Scattering on Metal Surfaces

Introduction

In the classical application of vibrational spectroscopy, both infrared (IR) absorption and Raman scattering (RS) were applied to explore molecular structure in the gas, liquid and solid state through the appearance of group characteristic frequencies.\textsuperscript{1-3} Group theory shows that both methods are complimentary and especially powerful when used together for studies in molecular structure.\textsuperscript{4,5} For example, in centrosymmetric molecules a vibrational mode is either Raman or IR active. The character table contains the components that give rise to infrared (via the change in dipole moment) and Raman (via the polarizability derivative tensor) activity. Raman scattering has always been more challenging than IR absorption because the cross section in RS is about $10^{-10}$ smaller than in IR. In many cases the luminescence from the substrate or from impurities within sample could unfavourably swamp out the Raman signal.

Raman instrumentation has, however, improved with better detection instrumentation and a larger selection of laser lines that could be used to record resonance Raman spectra (RRS). Selecting or "tuning" a laser line into resonance with an electronic transition in the molecule gives rise to resonance Raman scattering (RRS) and enhanced intensity of certain bands. In particular, the Raman modes that are related to the electronic transition are greatly enhanced as much as $10^2 - 10^6$ fold.\textsuperscript{6} The selectivity in RRS is derived from the localization of the electronic transition to one part of the molecule. For example, in the RRS of Fe(II)-α-dimine
complexes the charge is localized in the skeletal region and only the skeletal modes were enhanced.\textsuperscript{7} Topics in RRS have been thoroughly reviewed by Behringer\textsuperscript{8} and Rousseau\textsuperscript{9} et al.

The most recent interest in Raman spectroscopy includes detection and analysis at the sub monolayer\textsuperscript{10} level. Instrumentally, a new development is Fourier transform\textsuperscript{11,12} (FT)\textsuperscript{10} Raman spectroscopy with near IR laser sources (Nd:YAG operating at 1.064 $\mu$m). The current Raman detection systems have significantly advanced over the photomultiplier tubes used in conventional Raman detection.\textsuperscript{10} For example, non resonance Raman spectra were recorded with charge coupled device (CCD) multi-channel detection for a Langmuir layer on the subphase.\textsuperscript{13} Prior to the recent developments of CCD detectors, other experimental Raman techniques were developed and applied to problems in surface science. Greenler was the first to consider a minor enhancement in the Raman intensity for a molecule near a highly reflecting surface.\textsuperscript{14} Raman spectroscopy of adsorbed molecules on surfaces further flourished after the initial report of Fleischmann and coworkers who first observed intense Raman scattering from pyridine adsorbed on a roughened silver electrode.\textsuperscript{15} Their findings led to the development of a new technique that incorporated a roughened surface to enhance the Raman scattering (SERS) of molecules. For molecules adsorbed or near a rough metal surface the SERS intensity was significantly greater than the normal Raman scattering. For carbon monoxide adsorbed on vapour deposited Ag, an enhancement of $10^{6}$ was observed for the adsorbed layer.\textsuperscript{16} Hexter et al. noted an enhancement factor of $10^{4}$ for pyridine on Ag electrodes. Their
calculations were based on the surface area of the roughened electrode and the band intensities of liquid(991 cm⁻¹) and adsorbed(1006 cm⁻¹) pyridine.¹⁷

The current consensus is that Raman scattering and surface enhanced Raman scattering (SERS) are two vibrational techniques, among several others, that may be used to probe the structure at the interface that exists in electrochemical (solid-liquid interface), heterogeneous catalytic (solid-gas interface) and other systems (i.e., solid-solid interface).¹⁸ The vibrational spectrum of the adsorbates is the fingerprint that gives details about the chemical bonds and their geometry at the interface.¹⁹ Chemical reactivity at metal surfaces is also relevant to possible applications of SERS. Ackers and coworkers have applied SERS to identify the surface intermediates in the Ag catalysed reaction of ethylene and oxygen to form ethylene oxide.²⁰ In the same context, the product in the chemical reaction of imidazole (a corrosion inhibitor) on silver sols was identified in the SERS spectra.²¹ The application of SERS to biological systems has been recently reviewed.²² Raman scattering is one of the most important tools for in situ vibrational studies at surfaces.²³

SERS Theory

The observation of intense Raman scattering of pyridine in the electrochemical experiment described by Fleischmann et al. was originally attributed to the increase in the surface area that occurred during the oxidation-reduction cycle.¹⁵ Creighton et al.²³ and Van Duyne et al.,²⁴ however, showed that the growth in surface area by electrochemical cycling could not significantly account for the strong Raman signals
of adsorbed pyridine. They proposed an increase in the Raman scattering cross section for the adsorbed molecules on the electrode surface. Various models were collected and published in the first book dedicated to surface enhanced Raman scattering.\textsuperscript{25} About 572 SERS papers were cited by Seki in 1986.\textsuperscript{26} They include SERS from molecules adsorbed onto colloids(\text{Ag}, \text{Au}, \text{Cu}, \text{Pt}, \text{Rh}, \text{Na}, \text{K}), electrodes(\text{Ag}, \text{Au}, \text{Cu}, \text{Cd}, \text{Hg}, \text{TiO}_2), and island films(\text{Ag}, \text{Au}, \text{Cu}, \text{Ni}, \text{Pd}, \text{Ti}, \text{Li}, \text{Na}, \text{In}, \text{Co}, \text{NiO}). An essential feature of all SERS were the protrusions and bumps at the surface which gives rise to surface roughness. For colloids and island films the metal domains can be approximated as spheres and spheroids respectively. The size of the particles are not homogeneous and smaller or comparable to the wavelength of the exciting laser line. The particle shape is also important, and it can be described by the aspect ratio defined by \(a/b\) where \(a\) and \(b\) are respectively the lengths of the semi-major and semi-minor axis. Prolate spheroids have \(a/b > 1\), while oblate spheroids have \(a/b < 1\). Particle where \(a/b = 1\) are spheres.

The common opinion is that SERS is an electromagnetic effect. The extent of the chemical contribution to the total SERS effect can be examined case by case.\textsuperscript{27} Mechanisms that amplify the electric field at the molecule are considered electromagnetic while those that strengthen the polarizability derivative tensor are termed chemical or electronic. The chemical effect is commonly associated with resonant excitation into the charge transfer(CT) band of the metal-molecule complex.\textsuperscript{27-30} Chemical enhancement is therefore restricted to molecules directly at the surface. For pyridine adsorbed onto silver island the excitation profile for the 1010
cm⁻¹ band confirmed significant chemical enhancement when the laser line was tuned in resonance with charge transfer band which was well outside the surface plasmon resonance.³¹ Otto et al. note that chemical enhancement on coldly deposited metal films originates from adatoms of atomic scale roughness. They support their claim in experiments where no Raman signals were observed upon annealing the film.³² The adatoms were supposedly destroyed upon heating.

The electromagnetic (EM) effect is attributed to resonant excitation of the collective conduction electrons or surface plasmons (SP) modes that are present on rough metal surface. The SP act to concentrate the incident field at the surface of the metal particles.³³ An essential requirement for EM effects is the surface roughness which helps produce radiative SP that can couple to the exciting laser line and participate in the Raman emission.³³,³⁴ For an isolated sphere (with dimensions less than the incident wavelength), the electric field that is produced inside a sphere, by a laser line of electric field $E_0$ and frequency $\omega$ is given by³⁵:

$$E_{x,y,z} = \frac{1}{1 + \left[ \varepsilon(\omega)/\varepsilon_0 - 1 \right] A_{x,y,z}} E_{0x,y,z}$$  \hspace{1cm} (3.1)$$

where $\varepsilon(\omega)$ and $\varepsilon_0$ are the complex dielectric constants of the metal particle and the surrounding medium respectively. The depolarization factor $A$ in 3.1 is an integral and completely shape dependent (i.e., $A = .333$ sphere, $A = .17356$ prolate $(a/b = 2)$ and size independent.³⁵ Resonance is achieved when:
\[ \epsilon_o + [\text{Re}\{\epsilon(w)\} - \epsilon_o]A = 0 \] (3.2)

For a sphere the SP resonance occurs when the real part \( \text{Re}\{\epsilon(w)\} \) of the dielectric constant equals \(-2\epsilon_o\). The imaginary part \( \text{Im}\{\epsilon(w)\} \) of the complex dielectric constant is related to SP damping and prevents it from reaching infinity. A small \( \text{Im}\{\epsilon(w)\} \) component would therefore minimizes damping of the SP.\(^{36}\) In the visible region excitation of the SP are fulfilled by Ag, Cu, Au, alkali metals.\(^{37}\) For this reason most investigation of SERS have been with the coinage metals.

The SP are localized on isolated bumps and do not propagate along the film surface.\(^{35}\) The field \( E_{\text{in}} \) will give rise to a electric field outside (\( E_{\text{out}} \)) the sphere, which is always greater the \( E_o \). The displacement vector \( D \) can be used to relate \( E_{\text{out}} \) to \( E_{\text{in}} \). The boundary conditions require that the normal displacement vector 
\[ D^\perp = \epsilon(w)E^\perp_{\text{in}} = \epsilon_o E^\perp_{\text{out}} \]
and that the tangential displacement electric vector \( E^\parallel_{\text{out}} \)
\[ = E^\parallel_{\text{in}}.\] \(^{35}\)

The Raman emission is also enhanced by the metal particle. The Raman dipole oscillating at \( w \) polarizes the metal particle and uses it as an antenna to amplify its Raman radiation.\(^{35,37,38}\) The overall SERS enhancement therefore results from amplification of both incident and scattered fields. The electromagnetic enhancement in SERS decays as the molecule is separated from the surface. For a molecule located a distance\( (d) \) from a sphere\( (\text{radius } r) \) the SERS intensity\(^{39}\) is proportional to \( [r/(r + d)]^{-12} \). For a monolayer the SERS dependence with distance\(^{39}\) is proportional to \( [r/(r + d)]^{10} \).

A number of SERS experiments have demonstrated good evidence for EM
enhancement. The predicted exponential decay was demonstrated in a series of experiments that used spacer layers of arachidic acid monolayers to control precisely the distance from the metal particle.\textsuperscript{40,41} The experimental dependence of the Raman signal on the aspect ratio (or depolarization factor) and on the dielectric constant of the surrounding media was also in agreement with electromagnetic predictions.\textsuperscript{42} The authors showed that by changing the parameters A and $\varepsilon_s$ the SP resonance moved but that the maximum enhancement coincided with the shifted SP resonance. Further evidence of EM enhancement was observed in the excitation profiles of carbon monoxide on vapour deposited Ag(SP in the red spectral region) and colloidal Ag(SP in the blue spectral region).\textsuperscript{43} For both substrates the excitation profile peaked in the vicinity of the surface plasmon resonance. The interested reader should consult the excellent reviews by Aroca et al.\textsuperscript{34}, Moskovits\textsuperscript{33}, and Chang\textsuperscript{35} for additional details in surface enhanced Raman spectroscopy.

**SERS Spectra: Physisorption, Chemisorption and Depolarization**

The interaction between the metal particle and the adsorbate is revealed in the SERS spectrum. Chemisorption is observed when new bonds are formed between the adsorbate and the surface (i.e., formation of a surface complex). The surface could lower the symmetry of the molecule, which could lead to new band(s) and shifts in band frequency and relative intensity. Chemisorption is also restricted to adsorbates directly in contact with the metal. For example, chemisorption was observed by the presence of a strong band assigned to Ag-N or Ag-O frequency of the metal-molecule complex.\textsuperscript{44} The authors also showed that the surface complex was not formed when
the molecules (deposited as a LB layer) were separated from the surface by a spacer layer of arachidic acid.

For molecules physisorbed onto a SERS substrate, the spectrum shows no change in the vibrational bands or their relative intensity. The SERS spectrum is almost identical to the normal Raman spectrum. The signal to noise ratio is, however, greater in the SERS spectrum. There is no extensive metal-molecule interaction that would alter the molecular electronic distribution and the polarizability derivative tensor.  

Thus for physisorbed species, chemical enhancement can be excluded and the dominate enhancement is from electromagnetic mechanism.

SERS spectra are also depolarized and cannot be used to obtain molecular orientation. The surface roughness features scramble both the incident field into three orthogonal components and the net orientation of the adsorbed molecules. 

Tailoring Metal Island Films in SERS

Electromagnetic enhancement is observed to peak when the laser line is tuned into resonance with surface plasmons absorption band. Several investigations have considered optimal conditions to maximize SERS. Koh et al. used Si particles (whose size could be varied) as an underlayer to roughen the silver surface. They reported an optimal groove depth of 10 nm for the Si layer. Others have controlled the rate of deposition, the mass thickness and the temperature of the substrate during the formation of metal island films. The width and maximum absorption of the SP are related to the shape, size and inter particle spacing and can be varied by changing the deposition conditions. Film preparation is normally controlled to give
reproducible surface morphology and surface plasmon resonances. For example the 647.1 nm laser line was outside the SP absorption (maximum centred at 480 nm) in the 2.5 nm Ag island film but in full resonance with the SP of a 9.5 nm film.\textsuperscript{48}

Recently a substrate (formed by evaporating 20 nm metal film at a rate of 0.1 nm/s to a glass slide maintained at 200 °C) showed SERS activity in the near-infrared.\textsuperscript{32}

The combination of SERS (in resonance with the localized SP) and RRS (in resonance the molecular electronic absorption) gives optimal enhancement and sensitivity. The double resonance effect leads to surface enhanced resonance Raman Scattering (SERRS).

**Minor Raman Enhancement on Reflecting Surfaces**

A minor Raman enhancement (6-10) is predicted for molecules adsorbed onto "smooth surfaces" of highly reflecting metals.\textsuperscript{33} The nomenclature smooth surfaces has been used to describe films that do not support radiative plasmon modes. The original treatment was given by Greenler et al.\textsuperscript{14} and later expanded by Moskovits.\textsuperscript{33}

The surface changes the illuminating field on the molecule and the inelastic scattering from the Raman dipole. As shown in Figure 3.1 the molecule is excited by an incident field \( E_{\text{in}} \) or \( E_{\text{op}} \) and a reflected field \( r_s E_{\text{in}} \) or \( r_p E_{\text{op}} \). The magnitude of the reflected field is calculated from the Fresnel coefficients \( r_s, r_p \) of s and p polarized light. The resultant electric electric field components at the surface are given by\textsuperscript{33}:

\[
\begin{align*}
E_x &= (1 + r_p)E_{\text{in}} \\
E_y &= (r_p - 1)\cos\phi E_{\text{op}} \\
E_z &= (1 + r_p)\sin\phi E_{\text{op}}
\end{align*}
\] (3.3)
where the \( s \) and \( p \) are the electric fields polarized normal and parallel to the plane of incidence (ZY). The Fresnel coefficients relate the amplitude of the reflected beam to the incident beam. In vacuum:\(^5^4\):

\[
\begin{align*}
\mathbf{r}_s &= \frac{(E_s,\text{ ref.})/(E_\text{in})}{\cos \Phi/\epsilon^{1/4} - (1 - \sin^2 \Phi/\epsilon)^{1/4}} \\
&\quad \frac{\cos \Phi/\epsilon^{1/4} + (1 - \sin^2 \Phi/\epsilon)^{1/4}}
\end{align*}
\]

\[\text{(3.4)}\]

\[
\begin{align*}
\mathbf{r}_p &= \frac{(E_p,\text{ ref.})/E_\text{in}}{\cos \Phi - (1/\epsilon - \sin^2 \Phi/\epsilon^{1/4})} \frac{\cos \Phi + (1/\epsilon - \sin^2 \Phi/\epsilon^{1/4})}
\end{align*}
\]

where \( \Phi \) is the angle of incidence to the normal on the surface and \( \epsilon \) is the complex dielectric constant of the metal. On a metal surface, a phase shift of 180\(^\circ\) occurs for the \( s \)-polarized reflected field for all angles of incidence and \( r_s \rightarrow -1 \).\(^5^5\) The \( p \)-polarized reflected field is, however, in phase and changes to 180\(^\circ\) at high angles of incidence. In the limit of a good reflector \( r_p \rightarrow +1 \) and the reflected field reinforces the incident field. The calculations for Ag, Au, and Cu metals show that the normal component (\( E_z \)) of \( p \)-polarized light and an incident angle of 50\(^\circ\)-70\(^\circ\) produces enhanced fields.\(^1^0,^3^4\) Figure 3.2 depicts the square of the electric field on a reflecting surface as a function of the incident angle.

The inelastic light that reaches the detector is also increased by the light that is reflected from the surface as shown in Figure 3.1. The surface selection rules have been developed for polarized incident and scattered light. The predicated intensity patterns for the four polarized Raman spectra are\(^{5^3} \).
$$\begin{align*}
\text{SS} & \quad \alpha \left| a_{xx}(1 + r_p)(1 + r_s) \right|^2 \\
\text{SP} & \quad \alpha \left| a_{yz}(1 + r_p)(1 - r_p) \cos \Phi' + a_{zz}(1 + r_p)(1 + r_p) \sin \Phi' \right|^2 \\
\text{PP} & \quad \alpha \left| \{a_{yy}(1 - r_p) \cos \Phi + a_{yz}(1 + r_p) \sin \Phi' \} (1 - r_p) \cos \Phi' \\
& \quad + \{a_{yy}(r_p - 1) \cos \Phi + a_{zz}(1 + r_p) \sin \Phi' \} (1 + r_p) \sin \Phi' \right| \times 2 \\
\text{PS} & \quad \alpha \left| a_{xy}(r_p - 1)(1 + r_s) \cos \Phi + a_{zz}(1 + r_p)(1 + r_s) \sin \Phi \right|^2
\end{align*}$$

(3.5)

where the primes indicated the quantities for the scattered fields and $a_{ij}$ are the polarizability derivatives tensor components for the normal coordinate. For molecules adsorbed on a good reflecting surface, the PP spectrum is expected to be the most intense and dominated by bands that transform as $a_{xx}$. 
Figure 3.1 Excitation and scattering for a molecule(O) placed above a smooth surface. Incident and reflected beam are represented by 1 and 2 respectively.
EXCITATION

SCATTERING

Figure 3.1
Figure 3.2  The square of the field components as a function of the angle of incidence and with 514.5 nm excitation for In and Ag.

($E_x^2$ - $\Delta$, $E_y^2$ - O, $E_z^2$ - $\bullet$)
Figure 3.2
CHAPTER 4: Vibrational Spectroscopy of Phthalocyanine

Introduction

Routinely vibrational spectroscopy is primarily used to identify functional groups in a molecule through the appearance of "fingerprint" frequencies which are characteristic of the structural units. The normal coordinate analysis (NCA) of molecules is, however, pursued for the assignment of 3N-6 (or 3N-5 for a linear molecule) fundamental vibrational modes to internal coordinates. For small molecules the NCA could be solved on a microcomputer; however, for unsubstituted Pc where \( N \geq 57 \) (as in MPc\(_2\), or H\(_2\)Pc) the problem becomes very complex and is normally solved on a main frame computer. A NCA of copper phthalocyanine was recently published for the totally symmetric modes.\(^1\) The authors noted that the A\(_{1g}\) modes were heavily coupled. Following the theory of characteristic vibrational frequencies, bands of phthalocyanine were identified using the subunits or model compounds that were structurally related to Pc for which the vibrational assignment was known. These would include such molecules as benzene\(^2\), indole\(^3\), pyrrole\(^4\) and porphyrin\(^5\). This strategy has been used by our laboratory.

Jennings\(^6\) has tabulated and compared modes of the H\(_2\)Pc(\(\alpha\), \(\delta\), \(\chi\)), MPc(M= Mg, VO, Cu, Zn, Sn, Pb), and ClMPC(M=Al, Ga, In) to the model compounds. Her data are comprehensive and were used to generate the characteristic frequencies of Pc as shown in Table 1. The Raman spectra and further detailed vibrational assignments of various Pc molecules have been published by our group.\(^7-16\) The macrocycle bands are associated with the inner ring of Pc, which is composed of the sixteen alternating
<table>
<thead>
<tr>
<th>Pc Shift (cm⁻¹)</th>
<th>Pc Assignment</th>
<th>Model Compound: Raman Shift (cm⁻¹) Assignment</th>
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</tr>
<tr>
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<td>423 (indole)</td>
</tr>
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</tr>
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<td>720-730</td>
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<td>CN pyrrole or CC pyrrole</td>
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<td>1520-1550</td>
<td>CN aza group</td>
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<td>1580-1610</td>
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carbon and nitrogen atoms. The Raman spectra of Pcs have also been reported by others.\textsuperscript{17-19} Weak Raman bands due to the coordination of the metal to the pyrrole nitrogens have been observed in the 230-256 cm\textsuperscript{-1} spectral region.\textsuperscript{8,11,13,14} The metal-nitrogen assignment has been more extensively studied in the far infrared.\textsuperscript{20,21} The symmetry assignment of totally and non totally symmetric modes is given by the symmetry species and their activity in the IR, Raman and polarized Raman spectra. For example, for a D_{2h} Pc (i.e., H$_2$Pc) the A$_g$ modes would be IR inactive and the SS Raman polarized spectrum would be dominated by A$_g$ bands. The non totally symmetric modes (i.e., B_{1g}, B_{2g}, B_{3g}) would be observed with large relative intensity in the SP spectrum. The resonance Raman(RR) spectra of Pc are due to Herzberg-Teller\textsuperscript{22} scattering since both totally symmetric and non totally symmetric modes are enhanced and no overtone progressions are observed in the RR spectra.\textsuperscript{12} Herzberg-Teller scattering is normally observed for molecules where the ground and the excited state have similar geometries, as in the porphyrins.\textsuperscript{22} In the excitation profile for metal free phthalocyanine, both the totally and the non totally symmetric bands were enhanced, which confirmed RR scattering by the Herzberg-Teller mechanism.\textsuperscript{23}

**VOPc: Raman and IR of Thin Solid Films, SERS and Symmetry Assignment**

As pointed out in Chapter one Pcs have remarkable physical and chemical properties. Thin solid films and or Langmuir-Blodgett films are two practical methods that can be used to fabricate phthalocyanine into application devices. Debe noted that organic thin films, and especially ordered organic thin films, are becoming important in novel technologies, and will provide specialized surfaces, membranes and optical
Figure 4.1  Infrared of VOPc in KBr pellet (A)

Infrared of a 200 nm thin solid film of VOPc evaporated onto a KBr crystal (B)
Figure 4.1
response functions that cannot be provided by inorganic analogues. In the
development of the film device, it will be necessary to characterize and optimize the
structure and the chemistry of the film. The molecular organization (whether it is
amorphous, polycrystalline, or crystalline) can be non-destructively probed by
infrared (IR) absorption and polarized Raman scattering. The results of a thin film of
vanadyl phthalocyanine (VOPc) prepared at room temperature are presented.

Thin solid films of 200 nm VOPc were grown onto Corning 7059 glass slides
and onto KBr crystals. The Raman and IR spectra can be interpreted as point dipoles
interacting with an external electromagnetic field. The absorption of radiation is
proportional to the square of the scalar product of transition moment and electric
field. For infrared spectroscopy this means that the electric field and the change in the
dipole moment must have a component in the same direction. For VOPc (assuming
C₄ᵥ molecular point group) 23A₁ + 42E IR modes are expected. The change in the
dipole moment for the A₁ and E modes are along the Z and the (X, Y) molecular axis
respectively.

The infrared spectra of VOPc mixed with KBr to form a pellet, and of a 200
nm film are presented in Figure 4.1. The change in relative intensities between the
two spectra was a good indication of partial molecular organization. In the KBr pellet
there exists a random orientation of the VOPc molecules and the relative intensities of
the A₁ and E modes would reflect the extent of the change in the dipole moment
associated with the particular vibration. The decrease in relative intensity of the bands
marked(*) in the thin film sample could be rationalized in terms of face-on
Figure 4.2  RS spectra (SS and SP components) of a 200 nm VOPc film on KBr excited with the 514.5 nm laser line
or edge-on molecular orientation. In the face-on description the molecular plane (the XY plane) is parallel to the substrate and strong E modes are expected in the IR spectrum since the electric field is parallel to the in plane transition moments. The edge-on organization for VOPc would have the molecules standing up with the XY molecular plane perpendicular to the substrate. In this case the totally symmetric bands would be observed with strong relative intensity and the E bands would be absent or weak.

In the IR spectrum of the 200 nm thin film the totally symmetric bands at 728, 779 and 1002 cm\(^{-1}\) decreased in relative intensity; however, the E bands were unaffected. The results clearly showed that there existed a certain degree of face-on molecular orientation for the 200 nm film of VOPc prepared at room temperature. The SS and SP polarized Raman spectra for the VOPc film are shown in Figure 4.2. The SS spectrum was dominated by totally symmetric modes. The SP spectrum however contained mostly B\(_2\) and E modes. The differences in the SS and SP spectra were due to the partial molecular organization. Another important observation was the separation of the Raman bands at \{1338(SS-spectra), 1343(SP-spectra) cm\(^{-1}\}\} and \{1517(SP-spectra), 1525(SS-spectra) cm\(^{-1}\}\}. The doublet resulted from Davydov splitting, which is a solid state effect commonly observed in crystals.\(^{25}\) Each Davydov component has distinct polarization properties that can be probed by polarized Raman spectroscopy. For example in thin solid films of chloroaluminum phthalocyanine the \(A_g\) (at 1339 cm\(^{-1}\)) and \(B_g\) (at 1344 cm\(^{-1}\)) doublet was assigned to Davydov splitting and the bands were well observed in the SS(or PP) and SP(or PS) spectra, respectively.\(^{14}\)
For a molecular crystal with \( m \) molecules per unit cell, factor group symmetry \( u \), and site group \( v \), the number of Davydov components \( N(u,v) \) is given by:

\[
N(u,v) = \frac{m}{h} \sum \chi_u(R) \chi_v(R)
\]  
(4.1)

where \( h \) is the order of the factor group and \( \chi_u(R) \) and \( \chi_v(R) \) are the characters of the \( u \) and the \( v \) irreducible representation for the factor and site groups respectively.\(^{14}\) The possible site groups for each space group can be found in correlation tables. Crystals of VOPc were reported to be triclinic with factor group \( C_1 \) and two molecules per unit cell.\(^{36}\) The observed doublet in the spectra resulted from two molecules per unit cell, \( C_1 \) factor group, and \( C_1 \) site group. The assignment of the symmetry species of the Davydov components is addressed in the next section (Raman studies of VOPc on highly reflecting surfaces).

The SERS of 1 nm VOPc evaporated onto a 10nm Ag island film and on Ag-Sn spheres are shown in Figure 4.3. The SERRS of 1 nm VOPc evaporated onto Ag and Au islands is shown in Figure 4.4. The vibrational frequencies in the SERS and SERRS spectra for all three substrates were observed unshifted with respect to those recorded for the VOPc film on glass or on the KBr crystal. The results suggest that there were no strong metal-molecule interactions that could perturb the vibrational energies of VOPc. The molecules were therefore physisorbed onto the SERS substrates and the Raman enhancement was attributed to the electromagnetic mechanism. The components of the Davydov doublet were observed as one strong
Figure 4.3  Top: SERS(multiplied by 5) for 2 nm VOPc on Ag islands.

Bottom: SERS of 1 nm VOPc on Ag-Sn spheres
SERS of VOPc. LL=514.5nm
Ag Islands
SS- Spectra

Figure 4.3
band in all four polarized SERS and SERRS spectra. The observations confirm that
the doublet was a solid state effect which was not present at the low surface coverage
of 1 nm. The doublets were, however, observed for a 65 nm film of VOPc deposited
on silver coated tin spheres. Another important observation in the SERS and SERRS
spectra was the similarity of all Raman components in the SS, and SP spectra. The
spectra were all depolarized and further discussion is given in the next section (Raman
studies of VOPc on highly reflecting surfaces).

The SERS, SERRS and the infrared data of a 200 nm thin solid film were used
to assign the symmetry of the bands in VOPc. Assuming a C₄ᵥ molecular point group,
the infrared spectra would contain only Aᵥ and E species, whereas the A₁, B₁, B₂ and
E bands were expected in the Raman or surfaced enhanced Raman spectra. Bands that
were observed exclusively in the Raman spectra could be assigned to B₁, and B₂
representations. A strong band in the Raman with a weak counterpart in the IR was
most likely an Aᵥ vibration. A band observed with strong intensity in the infrared and
weak intensity in the Raman was assigned to an E representation. The symmetry
assignment for VOPc is given in Table 4.2.
Figure 4.4  SERRS spectra of 2 nm of VOPc on Ag and Au islands films.

Both spectra are SS polarization.
Figure 4.4
<table>
<thead>
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<td>1610 m</td>
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Raman Studies of VOPc on Reflecting Surfaces

Ultrathin films of VOPc were evaporated on smooth metal surfaces of Ag, Cu, and Au. Metal films (200 nm thick) were formed by evaporating metal at a rate of 0.5 \( \text{nm/s} \) to glass substrates maintained at room temperature. An overcoat of 10 nm of VOPc was then evaporated onto the metal coated substrate which was maintained at 200 °C. Long delay times of 7 s/cm\(^{-1}\) were required to obtain polarized Raman spectra of these samples. All spectra were recorded at an angle of incidence of 60°.

The SS, SP, PP, and PS spectra for 10 nm VOPc on smooth silver are shown in Figure 4.5 and Figure 4.6. It can be seen that there were essentially two different types of spectra since the SS=PP spectrum and the SP=PS spectrum. This was a clear indication that the relative intensity of different irreducible representations was sensitive to the molecular orientation of the molecules on the surface. The spectra could not be entirely considered in the context of surface selection rules for adsorbates on a reflecting surface.\(^{27}\) At 514.5 nm and for Ag, Cu, Au surfaces, Raman modes transforming as \( a_{\text{ax}} \) (z normal to the surface) should give strong Raman signals followed by \( a_{\text{ay}} \) and \( a_{\text{yz}} \), while \( a_{\text{xx}}, a_{\text{yy}}, \) and \( a_{\text{xy}} \) should be weak. The main signals in the SS(or PP) spectra were assigned to the \( A_1 \) irreducible representation. The SP(or PS) however contained mostly \( E \) and \( B_2 \) species.

Similar polarized spectra were also recorded for VOPc evaporated onto glass coated with a 200 nm film of Au and Cu. The SS and SP spectra for VOPc on smooth Au is shown in Figure 4.7. Polarization effects in RRS(excitation at 647.1 nm) were not as pronounced as they were in the RS recorded with the 514.5 nm laser.
Figure 4.5  SS and SP Raman Spectra of 10 nm thin film of VOPc deposited on 200 nm Ag film.
Figure 4.5
Figure 4.6  PP and PS spectra for 10 nm VOPc on 200 nm Ag film
Figure 4.6
line.

Enhanced Raman intensity (minor enhancement due to reflection) was indeed observed for totally symmetric bands with P-polarized incident light at an angle of incidence of 60°. The intensity of the $A_1$ species in the PP spectrum was about 2.5 times stronger than in the SS spectrum. The results were in agreement with the favourable excitation of P-polarized light based on the Fresnel coefficients and the enhanced normal fields. In Raman spectroscopy one would have normally expected to observe the totally symmetric bands with more intensity in the SS spectra than the PP spectra. For a 200 nm VOPc film on glass, the overall intensity of the $A_1$ modes in the SS spectrum was indeed stronger than that of the PP spectrum for a 60° incidence. For adsorbed molecules on mirror surfaces, the total S-polarized field on the molecule was diminished by the out-phase component of the S-polarized reflected light.

Davydov doublets were distinctly observed in the 1340 cm$^{-1}$ and 1520 cm$^{-1}$ regions with components of different irreducible representations. The 1338 and 1525 cm$^{-1}$ bands were assigned to an $A_1$ type in the SERS of VOPc. The second components at 1343 and 1517 cm$^{-1}$ were only observed in the SP and PS spectra and were not observed in the infrared. These bands hence were not E species. The only possible assignment for both frequencies was to the $B_2$ irreducible representation. The symmetry assignment of VOPc in the isoindole spectral region is found in Table 4.3.

The samples [10 nm VOPc/(200 nm metal)] were then overlaid with a rough surface of silver by evaporating 10 nm silver onto the samples maintained at 200 °C. The absorption spectrum for the silver island film on glass showed a plasmon
Figure 4.7  Raman spectra of 10 nm thin film of VOPc on 200 nm Au
Figure 4.7
TABLE 4.3  Symmetry Assignments in the Region of the Isoindole Stretching

Vibrations for VOPc

<table>
<thead>
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Figure 4.8  SERS of 10 nm overlay of Ag evaporated onto the
(10 nm VOPc/200 nm Ag) sample. SS and SP polarized spectra
Figure 4.8
absorption centered at 500 nm. Molecular vibrations in SERS and SERRS were observed unshifted with respect to the values recorded in the thin solid film and indicated that the molecules were physisorbed onto the metal surface. Under similar conditions it was found that the overall intensity in the SERS spectra was less than that in the SERRS spectra. The greater scattering efficiency in SERRS was attributed to the superposition of two enhancements via 1) a large scattering cross section due to excitation in resonance with the molecular electronic absorption of VOPc, and 2) electromagnetic enhancement due to resonance with the localized surface plasmons on the silver island film. Another important observation was that the polarized spectra that were recorded for VOPc on smooth metal were replaced by depolarized spectra when the samples were overlaid with the island film of silver. SERS spectra were depolarized, in the sense that all four polarization combinations (i.e., SS, SP, PP, PS) gave rise to the same general spectrum. For a practising Raman spectroscopist this means that polarization techniques like those used to probe individual symmetry species in crystalline samples cannot effectively be used in a SERS investigation. The depolarization effect in SERS can be seen by comparing Figures 4.4 and 4.8. The spectra in Figure 4.8 also showed that the Davydov doublets were reduced to a single band in the 1340 and 1520 cm\(^{-1}\) region. Evaporating the overlayer of silver had the effect of interfering with the solid state effect that gave rise to Davydov splitting.
IR and SERS of LB Monolayers of VOPcTTb and ZnPcTTb

The vibrational characterization of vanadyl tetra-tert-butylphthalocyanine (VOPcTTb) and zinc tetra-tert-butylphthalocyanine(ZnPcTTb) was attempted by considering the IR, RS and SERRS data. Both molecules have a four-fold principal axis and the tert-butyl groups could produce four isomers. The first step was to separate the frequencies of the parent molecule(i.e., VOPcTT and ZnPcTT) from those of the tert-butyl groups. A direct comparison of the IR and RS spectra of the parent molecule with the tert-butyl substituted molecules allowed a clear separation of the Pc and tert-butyl frequencies. The IR spectra illustrated in Figure 4.9 showed that a number of new bands were present in the spectra of VOPcTTb. These bands were assigned to the tert-butyl moieties and marked(t) in the spectrum. The VOPc vibrations affected by the substitution were the wagging vibrations of the peripheral C-H groups at 728 cm⁻¹ and the C-H bending at 1119 cm⁻¹. The C-H stretching vibrations of the tert-butyl groups were observed with strong relative intensity in the IR. Similar observations were made for ZnPcTTb. The RS and RRS spectra were dominated by the vibrational frequencies of the parent Pc molecule. This was in agreement with the fact that the Raman cross section of molecular vibrations associated with conjugated systems are much larger than those of saturated groups.

SERRS of Langmuir-Blodgett layers of VOPcTTb and ZnPcTTb were also studied. The first step was to obtain the isotherm which would give the stable pressure range at which the Langmuir layer could be transferred to the SERS substrate. The phthalocyanines were dissolved in toluene([Pc]=2.5 x10⁻⁴ M) and
Figure 4.9  Infrared absorption spectra of VOPc and VOPcTTb in KBr pellets.
Figure 4.10  Surface pressure versus area isotherms for VOPcTTb and ZnPcTTb.
Figure 4.10
spread by a syringe onto the subphase which was thermostated at 20 °C. For both molecules, the compression rate during the acquisition of the isotherm was 0.001 nm²/molecule/s. The isotherms were reproducible for both phthalocyanines and are presented in Figure 4.10. The isotherm showed that surface pressures less than ca. 35 mN/m could be used for LB transfer. The surface pressure was measurable at ca. 1 nm²/molecule. At the molecular area ca. 0.5-0.3 nm² the surface pressure increased steadily as expected for the monolayer in a tightly packed crystalline state. The limiting area was found by extrapolating the most linear part (where the surface pressure increased steadily with molecular area) to zero surface pressure. The limiting area for both molecules was about 0.5 nm²/molecule. The isotherm of ZnPcTTb was reported in 1985 by Hann et al.²⁹ who used xylene as the spreading solvent. Their isotherm also showed that the surface pressure was measurable at ca. 1.4 nm² but their limiting area of 1.20 nm²/molecule was much larger than that in our study. Roberts et al.³⁰ noted that the isotherm is dependent on the spreading solvent, yet three²⁹-³¹ groups who all used xylene as the spreading solvent report 0.87nm², 0.96nm², and 0.60nm² for the limiting area for CuPcTTb. The inconsistent data from these groups suggest that serious research in the isotherm of phthalocyanine derivatives is still required. Molecular association as described by Snow et al.³² cannot completely account for the range in the limiting areas reported for CuPcTTb where the bulky tert-butyl groups are expected to reduce or prevent aggregation. Our group, in collaboration with Dr. DeSaja at the Universidad de Valladolid (in Spain) is currently comparing isotherm data collected under similar experimental conditions but
on different film balances.

Langmuir-Blodgett monolayers of ZnPcTTb and VOPcTTb were transferred to clean glass slides, Ag-Sn spheres, and 10 nm Ag island films at a constant surface pressure of 10 mN/m. The substrate was immersed in the subphase prior to spreading the phthalocyanine solution. Compression was done under constant surface pressure mode. The monolayer was maintained at 10 mN/m, for a period of time (ca 30 min.) that it took the moving barrier to completely stabilize and remain at a constant position. The stable Langmuir layer was then transferred to the substrate at a rate of 4.8 mm/min. The entire process was completely repeated in multilayer LB assemblies. The transfer ratio (defined as the ratio of the area removed from the water surface to the area on the substrate coated with the monolayer) was near unity on the glass substrates, 1.7 on Ag-Sn spheres, and 1.2 for Ag island films. Transfer ratios in excess of unity were attributed to the increase in surface area due to the surface roughness in the SERS substrates.

A linear dependence of absorbance was noted for 1, 2 and 3 LB layers of ZnPcTTb and VOPcTTb on glass and suggested uniform deposition of Pc during each transfer cycle. The SERRS spectra (SS, SP, PP, PS) for one LB of ZnPcTTb or one LB VOPcTTb were all identical to the RRS (SS polarization) for 3 LB on glass. Consequently, the SERRS spectra were all depolarized (i.e., the relative intensities of bands in the SS-component were equal to those in the SP, PP or PS spectrum).
Similarly all SERS spectra excited with the 514.5 nm line were all depolarized. Since all the fundamental frequencies were observed unshifted from their values in LB films
on glass or in thin solid films, the molecules were *physisorbed* onto the SERS surfaces. The SERRS of VOPcTTb and ZnPcTTb are presented in Figures 4.11 and 4.12. For both molecules the SERRS spectra was dominated by frequencies of the macrocycle, and the relative intensity of the peripheral groups was almost negligible.

A partial characterization of the vibrational spectra of VoPcTTb and ZnPcTTb is presented in Tables 4.4 and 4.5.
Figure 4.11  SERRS spectra of 1 LB of VOPcTb on Ag-Sn spheres. S and P incident polarization and total collection.
Figure 4.12  SERS of one LB of ZnPcTTb on Ag-Sn spheres.
Figure 4.12
Table 4.4 Observed Vibrational Frequencies of VOPc\textsuperscript{5-}Tb

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Energy Transfer in LB Monolayers and Selective Spectroscopic Tuning

It was recently shown that neat LB layers of N-octyl,N-isobuty1-3,4:9,10-perylene-bis-dicarboximide (PBDC) produced surface enhanced fluorescence with a broad emission at 648-nm which was attributable to excimer formation.\textsuperscript{34} The broad emission was also observed for other perylene derivatives.\textsuperscript{35} In a series of LB reports\textsuperscript{34,36} it was shown that there were at least two processes that could be used to describe fluorescence from molecules adsorbed on rough metal surfaces. There is fluorescence enhancement that results from the electromagnetic field enhancement on the SERS surface and fluorescence quenching due to the nonradiative transfer of the electronic excitation energy to the surface plasmons.\textsuperscript{34-38} In some cases luminescence may be completely removed for molecules in close proximity to a metal surface since the lifetime of the molecular excited state may be considerably shortened in the presence of the metal.\textsuperscript{38} Further studies of energy transfer with LB on rough metal surfaces were considered in our laboratory. In particular our objective was to investigate energy transfer from a sensitizer (or energy donor) LB layer to an activator (or energy acceptor) LB layer on a rough metal surface. The energy transfer between dyes in LB assemblies on glass substrates is well documented in the classical studies by Kuhn and coworkers.\textsuperscript{39-41}

The excimer emission of PBDC centered at 648 nm served as the energy donor and the broad absorption band of ZnPcTTb at 680 nm served as the energy acceptor. Three unique configurations of PBDC and ZnPcTTb were investigated on Ag-Sn spheres. The first sample was made by putting the donor layer adjacent the rough
metal surface\(, i.e., (1\text{LB} \text{ZnPcTTb}/1\text{Lb} \text{PBDC/Ag-Sn})\). In the second assembly the configuration of donor and acceptor layers was reversed \(i.e., (1\text{LB} \text{PBDC}/1\text{LB} \text{ZnPcTTb}/\text{Ag-Sn})\). The third sample consisted of a mixed layer of ZnPcTb and PBDC in the mole ratio 1:1. In all LB preparations, the monolayer was compressed to a constant surface pressure of 10 mN/m and the substrate was withdrawn at a rate of 4.8 mm/min. The 514.5 nm laser line was used to study the energy transfer process. This laser line was in full resonance with molecular absorption band in PBDC and outside the visible absorption band of ZnPcTTb.

Surface enhanced resonance Raman scattering for sample one is shown in the lower trace of Figure 4.13. The intense broad band corresponds to the excimer emission of PBDC and there was no qualitative evidence of energy transfer between donor and acceptor layers. In our experiment, the absence of excimer emission would be indicative of good energy transfer. The upper spectrum where PBDC was separated from the surface by a layer of arachidic acid, was typically observed for enhanced fluorescence and long range electromagnetic Raman enhancement. As expected the fluorescence enhancement increased when the chromophore was separated by a spacer layer of arachidic acid from the rough surface.$^{36}$ The weak Raman intensity of PBDC bands in the upper spectrum can be attributed to the decay in the electromagnetic enhancement as the Raman scatterer was separated from the SERS surface.$^{42,43}$

The spectra shown in Figure 4.14 were taken from a sample that was prepared by first overcoating one half of the SERS substrate with one monolayer of ZnPcTTb
Figure 4.13  Surface enhanced Raman and fluorescence of PBCD in a two layer structure. One LB of PBDC separated from the surface by one LB of arachidic acid (upper trace). The spectrum of the {1LB ZnPcTTb/1LB PBDC/Ag-Sn spheres} assembly is shown in the lower trace.
Figure 4.13
and then adding a second LB of PBDC to the entire substrate. The upper trace contained the typical SERRS and excimer emission of PBDC. The lower spectrum was recorded with the energy acceptor layer directly adjacent to the rough surface. In this case the excimer emission of perylene was completely quenched by the acceptor layer. The spectral evidence in Figures 4.13 and 4.14 clearly shows that the absorption efficiency of the acceptor (i.e., ZnPcTTb) layer was enhanced by the metal surface allowing complete energy transfer from the donor to the acceptor layer. The results are in agreement with enhanced absorption and luminescence of adsorbed molecules on rough metal surfaces.\textsuperscript{38,44}

In the third sample a mixed monolayer of PBDC and ZnPcTTb in a 1:1 mole ratio was transferred to Ag-Sn spheres. The results indicated that there was very little energy transfer in the mixed LB layer since the excimer emission of PBDC was strongly observed.

The accumulative results indicate that the most efficient LB structure for energy transfer was the two layer assembly with the acceptor layer adsorbed directly onto the metal surface. It was also important to note that there was no interference from ZnPcTTb in the spectra recorded with 514.5 nm laser line. Spectra recorded with 647.1 nm (in resonance with red absorption band of ZnPcTTb) excitation gave typical spectra of ZnPcTTb without any interference from PBDC. The choice of laser lines in resonance with one or the other molecular species demonstrated selective spectroscopic tuning in a two component system. Excitation at 514.5 nm produced SERRS of PBDC without any spectral interference of ZnPcTTb. Excitation at 647.1
Figure 4.14  Surface enhanced Raman scattering and fluorescence of 1 LB

of PBDC directly on Ag-Sn spheres (upper trace) and of the assembly

{1LB PBDC/1LB ZnPcTTb/Ag-Sn spheres} (lower trace)
Figure 4.14
Figure 4.15  SERRS of ZnPcTTb for the assembly {1LB PBDC/1LB ZnPcTTb/Ag-Sn spheres}. Only the SERRS of ZnPcTTb is revealed with 647.1 nm excitation.
Figure 4.15
nm however was exclusively sensitive to the ZnPcTTb species and no Raman lines of PBDC were observed. The SERRS of ZnPcTTb was observed without any interference from PBDC as shown in Figure 4.15.
CHAPTER 5: SERRS of Langmuir-Blodgett Monolayers: Coverage Dependence

Early coverage dependence studies of SERS were conducted in ultra high vacuum. The experiments were designed to investigate the intensity relationship of the SERS signal as the surface coverage of the adsorbate increased from 0 monolayers (ML) to a few ML. The most difficult aspect in the early experiment were the uncertainties to which surface coverage was measured. Rowe et al. measured surface coverage by measuring the carbon densities derived from Auger electron spectroscopy. A quartz crystal oscillator was used to monitor pyridine coverage in the work published by Seki et al. Sands et al. used UV-photoemission spectroscopy to calibrate the surface coverage of pyridine. Radioactive tracing was used to measure coverage in $^{14}\text{CN}$ on Ag island films. Surface coverage was determined spectrophotometrically for cobalt phthalocyanine on Ag films. In 1988, Dr. Cotton's laboratory and our group independently submitted reports that used the Langmuir-Blodgett technique (for the first time) to investigate the SERRS intensity versus coverage dependence. Both groups used mixed monolayers composed of dye and fatty acid to control precisely the surface coverage.

The surface coverage was controlled using mixed monolayers of vanadyl tetra-tert-butyphthalocyanine (VOPcTTb) and arachidic acid in various mole ratios of VOPcTTb:arachidic acid. All monolayers were spread from toluene solutions. Stock solutions of VOPcTTb (2.67 x 10^{-4} M) and arachidic acid (1.07 x 10^{-3} M) were used to prepare mixed solutions containing both species. In all cases the appropriate volume of the arachidic acid solution was added to 2 ml of the stock VOPcTTb solution to give the corresponding mole ratio of VOPcTTb:arachidic acid. Mixed monolayers were transferred to Ag-Sn spheres.
and 4 nm Au island films. An internal standard (VOPcTTb:arachidic acid = 10:1) was used to account for differences that may have existed among the SERS substrates. In particular, one LB of the internal standard was transferred to one half of the SERS substrate. The sample of variable composition was then transferred to the remaining portion of the SERS slide. The substrate was immersed into the subphase prior to monolayer compression. The Langmuir layer was compressed to 10 mN/m for ca. 25 min before being transferred at a rate of 4.8 mm/min. In another type of experiment, more than one mixed LB was transferred to the SERS substrate. These experiments were used to investigate the coverage dependence for mixed multilayer assemblies.

All coverage dependence studies for a given batch of SERS slides were run within the same day in attempt to keep the optical path, the power of the laser and other optical parameters constant. The samples were spun in a rotating cell, with the laser probing a circle of radius ca. 0.25 cm on the sample. This spinning sample effectively allowed scattering from an area average and not from a single location. The absorption spectra of 3 LB of VOPcTTb on glass and 4 nm Au island film showed a well defined broad band centred at 706 nm and 650 nm respectively. SERRS with excitation at 647.1 nm was therefore in resonance with both the molecular electronic absorption of VOPcTTb and the localized surface plasmon of Au. SERRS was also measured for VOPcTTb on Ag-Sn spheres.

SERRS was observed to peak at submonolayer coverage for cobalt phthalocyanine vacuum evaporated onto Ag islands. The decrease in SERRS intensity was attributed to damping of the surface plasmon resonance by the adsorbed layer. Zeman et al. used two theoretical models to show that overall Raman intensity factor was heavily dependent on
Submonolayer SERRS coverage dependence on Ag-Sn spheres and Au island films. The curve is for the 685 cm$^{-1}$ band of VOPcTTb normalized to the internal standard (VOPcTTb:arachidic acid = 10:1)
Figure 5.1
Dipole-dipole\textsuperscript{10} interactions between molecules have been also predicted to affect surface coverage dependence. In our case minimal changes were observed between the absorption spectrum of 4 nm Au island film and the absorption spectrum of the composite (VOPcTTb + Au island). The amplitude did, however, vary slightly. In SERRS, overcoating the surface with an absorbing layer, could in the extreme case, eliminate the plasmon resonance and remove the electromagnetic enhancement.\textsuperscript{11} Garoff et al. have calculated the effect of dye concentration on the absorption spectrum of silver spheroids. Their work showed that the dye initially broadens and damps the silver resonance and eventually causes splitting for thicker dye coatings.\textsuperscript{12}

The distribution of the arachidic acid molecules within the VOPcTTb matrix cannot be directly addressed. Fluorescence studies, however, have shown that a good distribution of the fluorescent probe was observed among the host molecules.\textsuperscript{9} In this work we assumed that a homogeneous mixed monolayer was formed. All SERRS showed that the fundamental vibrations of VOPcTTb were not altered by the rough metal surface or the arachidic acid molecules. The SERRS submonolayer coverage dependence for the totally symmetric, macrocycle band at 685 cm\textsuperscript{-1} of VOPcTTb is presented in Figure 5.1. The bars in Figure 5.1 indicate the range of the observed experimental values. The data points were obtained by measuring the relative intensity of the sample (of variable coverage) to the internal standard on the same slide. The SERRS intensity versus coverage on Au island and Ag-Sn spheres showed a maximum at submonolayer coverage. Maximum intensity was observed at 17 and 25 mole percent VOPcTTb on Au island films and Ag-Sn spheres respectively. Zeman et al.\textsuperscript{6} and Kim et al. also observed a maximum of about 0.1 ML in their SERRS study. The
coverage variation (0.1-0.25 ML) between the three groups could be rationalized in terms of the shape of the metal particles which was theoretically shown to affect the field enhancement.\textsuperscript{6} At least for our data, the aspect ratio difference between Ag-Sn spheroids (spheres) and Au (oblate) island was a possibility, but without any detailed theoretical and further experimental work the prediction was not totally conclusive. Self absorption could not account for SERRS decrease with increasing phthalocyanine coverage. In a recent article\textsuperscript{12} it was shown that the resonance Raman scattering from 2 LB of ytterbium bisphthalocyanine on glass was almost twice that for 1 LB, which confirms that self absorption could not be a major contributor in reducing resonance Raman scattering.

Another important feature in our coverage dependence study was that the relative intensities of the vibrational fundamentals for different surface coverage were consistently seen with the same general pattern. The fundamental frequencies and the relative intensities in SERRS were also identical to 3 LB transferred on glass. As shown in Figure 5.2, the relative intensities for VOPcTTb at 20 mole percent and 10 mole percent were identical. The relative intensities in the SERRS spectra recorded by Zeman et al. for 0.1 ML and 36 ML were, however, remarkably different especially in the isoindole region.\textsuperscript{6} We have only observed changes in the relative intensity in the Pc vibrational spectra when the excitation source was tuned in-resonance and off-resonance to the electronic transition.\textsuperscript{14} Kim et al. also observed that the relative intensities were independent of surface coverage.\textsuperscript{7}

Coverage dependence studies of SERRS were also investigated by adding more than 1 LB to the SERS substrate. In this work multilayers of 10 mole percent VOPcTTb were
Figure 5.2  
SERRS of 20 mole percent and 10 mole percent of VOPcTTb on Ag-Sn spheres and Au on island film respectively. Spectra were taken with ST polarization and 50 mW of the 647.1 nm laser line.
Figure 5.2
transferred to sections of the Ag-Sn spheres slide to give 1, 2, 3 and 4 LB layers. The integrated SERRS for the 685 and 1523 cm\(^{-1}\) bands of VOPcTTb were measured. The integrated SERRS intensities(I) for both bands showed a continuous increase in going from 1LB→2LB→3LB→4LB(i.e., \(I_{1LB} < I_{2LB} < I_{3LB} < I_{4LB}\)) as shown in Figure 5.3. The largest marginal increase was observed in going from 2LB to 3LB samples. The increase in intensity with the number of layers could be rationalized in terms of the long range electromagnetic enhancement.\(^{15,16}\) In this respect the second, third, and fourth layers contributed to increase the total SERRS intensity.

SERRS coverage dependence of VOPcTTb evaporated onto Ag-Sn spheres showed the negative dependence with increasing coverage as shown in Figure 5.4. In this work, VOPcTTb was evaporated onto the same substrate in order to keep the enhancing properties of the substrate constant. In Figure 5.4 the decreasing effect of the SERRS signal with increasing coverage was clearly determined, despite the experimental uncertainties in measuring the film thickness of VOPcTTb. The results showed that the emission of the radiative surface plasmons was diminished or quenched with increasing film thickness.
Figure 5.3  SERRS coverage dependence for the 685 and 1524 cm\(^{-1}\) bands of VOPcTTb as function of the number of mixed layers (10 mole percent of VOPcTTb).
Figure 5.3
Figure 5.4

SERRS coverage dependence on the 685 cm$^{-1}$ band of VO$_2$FeTb for vacuum evaporated films on Au-Spheres. Data are normalized to the 10 nm sample. Data was taken from SS and PP polarized spectra.
Normalized SERRS Intensity

VoPcTTb Film Thickness (nm)

Ag-Sn spheres

Figure 5.4
CHAPTER 6: Blue and Green Forms of Rare Earth Bisphthalocyanines

Introduction

Two phthalocyanine rings joined by a central lanthanide(Ln) element form a series of complexes known as the rare earth bisphthalocyanines(LnPc₂). The general molecular structure is shown in Figure 6.1. X-ray diffraction analysis for the neodymium bisphthalocyanine(NdPc₂) complex showed that the Nd atom occupied a central position in between two staggered (ca. 45°) Pc ligands.¹ The metal was eightfold coordinated to the pyrrole nitrogens atoms and Nd-N bonds lengths varied from 2.39-2.49 Å. One ring in the NdPc₂ complex was saucer shaped towards the Nd atom and the other macrocycle was planar. The first report by Kirin et al. pointed out that two forms of NdPc₂ were produced in the reaction of rare earth acetates and o-phthalonitrile.² During product isolation by chromatography, a blue and a green zone was formed on the aluminium oxide column. The blue and green solutions were characterized by intense absorption bands at 634 nm and 672 nm respectively. The optical spectrum of the green material of the Nd complex was similar to the monophthalocyanine complexes and it was concluded by Kirin³ et al. that the green compound was composed of a single Pc ligand. The visible absorption band of the blue Nd complex was however similar to spectra of tin bisphthalocyanine and the blue form of the Nd complex was assigned to contain two Pc rings. The elemental analysis of the blue form of the Nd complex was consistent with the bisphthalocyanine structure.¹ No elemental analysis was however reported for the green complex which decomposed in dimethylformamide to give metal free phthalocyanine.
Figure 6.1  Molecular structure of lanthanide(Ln) bisphthalocyanine
MacKay et al. however isolated stable green and blue forms of gadolinium phthalocyanines. They reported that both compounds were bisphthalocyanine complexes. The latter group also noted that the colour of lanthanide bisphthalocyanine complexes was attributed to changes in the \( \pi \)-electron distribution during the formation of anions, cations, or neutral species rather than the formation of monophthalocyanine and bisphthalocyanine. Kasuga et al. isolated three products in the reaction of neodymium acetate and o-phthalonitrile. They characterized NdPc\(_2\)H (\( \lambda_{\text{max}} \) 345 and 636 nm), NdPc\(_2\) (\( \lambda_{\text{max}} \) 324, 470, and 676 nm) and PcNdCH\(_3\)COO (\( \lambda_{\text{max}} \) 336, and 675 nm). Elemental analysis was used to confirm the structure of the neodymium bisphthalocyanine complexes.

The band at 470 nm is always observed in bisphthalocyanine complexes which are green in colour. This absorption band is also characteristic of the phthalocyanine free radical (Pc\(^-\)) \(^6\) ligand as observed in Pc radicals \(^7\) of H(Pc\(^-\)) and Li(Pc\(^-\)). The green form of NdPc\(_2\) would therefore contain a Q absorption band at 676 nm, a free radical band at 470, a Soret band at 324 nm. The structural formula Pc\(^2\)Nd(III)Pc\(^-\) is used to describe the green form and it takes into account the free radical nature of one Pc ligand. The free radical Pc ligand in lutetium phthalocyanine(LuPc\(_2\)) complexes has been confirmed by ESR where a g-value ca 2 was measured. \(^5\)-\(^10\) The free radical bands in LuPc\(_2\) disappeared by chemical reduction with hydrazine \(^11\) or by electrochemical \(^8\) reduction. In the literature, the structure of blue form of lanthanide bisphthalocyanine complexes has been cited as LnHPC\(_2\) [i.e., GdHPC\(_2\)\(^4\), NdPc\(_2\)H\(^3\), LuHPC\(_2\)\(^12\) The extra hydrogen in the blue form of lutetium bisphthalocyanine(or
LnHPe₂) was never confirmed in the X-ray crystal structure.¹²

The electrochromic properties of rare earth bisphthalocyanines have also been studied.¹³⁻¹⁹ The electrochromic effect is used to describe the faradaic electrochemical reaction that results in a colour change of a material deposited on the surface of an electrode.¹³ The electrochromism of LnPe₂ is attributable to the generation of different π species (i.e., neutral, cations and anions) depending on the applied potential. LB films of LuPe₂ transferred to an indium tin oxide electrode exhibited reversible colour change from purple (−2 V vs. SCE)→blue (−1.7 V)→green (0 V)→brown-red (1.2 V).¹⁴

Our interest in lanthanide bisphthalocyanine complexes was in the potential use of these molecules as ultra thin films and Langmuir-Blodgett monolayers. The additional aim was to use surface enhanced Raman scattering, infrared, and uv-vis spectroscopy to collectively characterize the green (LnPe₂) and blue (LnHPe₂) forms of underivatized lanthanide bisphthalocyanine complexes and octa-tert-butylbisphthalocyanines of lanthanide elements (LnPe₅). The assignment of IR and Raman bands to internal coordinates for the green and blue forms of LnPe₅ has not been addressed in the literature.

Stable green and blue forms of octa-tert-butylbisphthalocyanine of rare earth elements were obtained from Dr. L. Tomilova from the Organic Intermediates and Dyes Institute (Moscow) during her visit (in 1991) to the University of Windsor. The synthesis of these compounds has been described elsewhere.²⁰,²¹ The synthesis of all rare earth bisphthalocyanine complexes begins with the formation of the
monophthalocyanine which converts to the green bisphthalocyanine form. Prolonged heating gives the blue form of the rare earth bisphthalocyanine.

Thin films (ca. 200 nm) of LnPc$_2$ were sublimed to Corning 7059 substrate and halide crystals. Langmuir layers of LnPc$_2^-$ were spread from benzene solutions onto an aqueous subphase (pH = 5.5, temp. = 15 °C). Monolayers were transferred under constant surface pressure (15 mN/m) mode to the glass, and SERS substrates.

Electronic Absorption Spectra of LnPc$_2^-$

In Figure 6.2 the visible electronic spectra of the green and blue form of an early and late lanthanide phthalocyanine complex are shown. The typical bands observed for all green forms of LnPc$_2^-$ were: Soret band (320 nm with a shoulder near 340 nm), radical$^{6,11}$ anion (460-490 nm) and Q-band at 667-687 nm. The spectra presented in Figure 6.2(A,D) were similar to literature$^5,17$ spectra of NdPc$_2$ and LuPc$_2$. In some reports$^4,14,18$, the structural formula of the green form of LnPc$_2$ was incorrectly written as HLnPc$_2$ which does not account for the free radical and charge balance for the neutral molecule.

The blue complexes of LnPc$_2^-$ (or LnHPc$_2$) have the characteristic signature as shown in Figure 6.2(B,D). Unlike the green analogues there was no absorption band in the 460-490 nm spectral window. These spectra were similar to the proposed blue form of GdHPc$_2^4$, NdHPc$_2^3$ and the complexes of tetra(n-butyl)ammoniumLnPc$_2^{22}$. The absorption spectrum of the latter compound would suggest that the absorption spectrum of the blue form of lanthanide bisphthalocyanine could be due to (Pc$^{2+}$ Ln(III)Pc$^{2+}$)$^-$H$^+$. Zirconium and hafnium bisphthalocyanine also form blue solutions.
Figure 6.2  Electronic absorption spectra of the green(A) and blue (B)
form of NdPc$_2$, and green(C) and blue(D) form of LuPc$_2$
Figure 6.2
and their absorption spectra are similar to the blue form of lanthanide bisphthalocyanine complexes.\textsuperscript{23} Since the oxidation state of the metal in ZrPc\textsubscript{2} and HfPc\textsubscript{2} is IV, the blue colour could originate from two Pc\textsuperscript{2-} ligands.

At ca. 700 nm, a weak satellite of the Q band was observed in the spectra of LnHPc\textsubscript{2}\textsuperscript{l}(Ln=Ho, Er, Tm, Yb, Lu). The band was absent in spectra of LnHPc\textsubscript{2}\textsuperscript{s}(Ce, Pr, Nd, Sm, Eu, Gd, Tb, Dy, .). The splitting of the Q band into another band (i.e., satellite or weak shoulder band) has been attributed to exciton interaction between the two Pc rings.\textsuperscript{22}

**Thin Films of LnPc\textsubscript{2}**

Using LnPc\textsubscript{2}(green material) as the starting material, thin films were simultaneously evaporated onto NaCl disks and glass substrates. Green films were obtained for LnPc\textsubscript{2}(Ln=Gd, Tb, Dy, Ho, Er, Tm, Yb, Lu). For these complexes the visible spectra of the thin film and the solution spectra of the thin film dissolved in benzene were identical to the solution spectra of the starting material. The absorption spectra for LuPc\textsubscript{2} are shown in Figure 6.3. Early rare earth elements in the LnPc\textsubscript{2}(Ln= Pr, Nd, Sm, Eu) series however, produced blue coloured films. The absorption spectra for NdPc\textsubscript{2} are also shown in Figure 6.3. When the film was washed with benzene a blue-green solution was collected and an insoluble film remained on the glass slide. The absorption spectrum of the insoluble film was distinctly identified as H\textsubscript{2}Pc by the strong bands at 665 and 700 nm and weaker bands at 602 and 640 nm. The IR spectrum of the thin film of NdPc\textsubscript{2} on NaCl also contained the N-H stretching vibration at 3290 cm\textsuperscript{-1} and another band at 1007 cm\textsuperscript{-1}. 
Figure 6.3 Visible spectra of an evaporated film (200 nm) of NdPc$_2$ and LuPc$_2$
(i), solution of film washed with benzene (ii), and starting material
dissolved in benzene (iii)
which are seen only in H$_2$Pc analogues.$^{14,24-26}$ The absorption spectrum of the blue-green solution contained two strong absorptions at 645 and 679 nm that were observed in the spectra of isolated green and blue forms of the NdPc$_2$ analogues. The starting material, however, gave an absorption spectra typical of the green form of NdPc$_2$. Traces of H$_2$Pc were not found in the IR spectrum of the starting material. Metal free phthalocyanine was therefore produced during sublimation. The sublimation results obtained for NdPc$_2$ were also observed for LnPc$_2$(Ln = Pr, Sm, Eu).

**Chemical Redox and Acid Effects on LnPc$_2$**

In the early studies of lanthanide bisphthalocyanine complexes, MacKay$^4$ et al. proposed an acid base equilibrium to account for the observed blue and green colours. M'Sadak$^{27}$ et al. showed that chemical reduction (with triethylamine) of the green form of the LuPc$_2$ produced the blue form. Addition of acetic acid to the chemically reduced DyPc$_2$(blue complex) produced the original green complex.$^{27}$ Our absorption spectra recorded in the reaction of the green form of PrPc$_2$ and LuPc$_2$ with different acids are presented in Figure 6.4.

The red shifted Q-band to ca. 705 nm was typically observed for the oxidized Pc ligand. The addition of CF$_3$COOH, HNO$_3$ or H$_2$SO$_4$ produced the red complex. The reaction of HCl and CH$_3$COOH with LnPc$_2$ also produced the red complex, although the rate was slower than with HNO$_3$. LnPc$_2$ (Ln = Ce, Pr, Nd) were not stable in CF$_3$COOH or HNO$_3$ and decomposed to give metal free phthalocyanine. The spectral evidence suggests that acids strongly interact with $\tau$ system of the Pc.
Figure 6.4 Visible spectra for the reaction products of LuPc₃ (green material) with with CH₃COOH (i), HNO₃ (ii), hydrazine (iii), PrPc₂ (green material)
CF₂COOH (i), HNO₃ (ii), one after HNO₃ addition (iii)
Figure 6.4
ligand to produce changes in the electronic distribution. These results suggest that under these mild conditions the oxidation state of the rare earth element was not affected in the green-blue inter conversion.

Raman Spectra of Green and Blue Forms of LnPc$_2$

LB monolayers of the green and blue form of LnPc$_2$ were transferred to glass slides, 4 nm gold island(SERRS) and 4nm Ag(SERS) island films. As shown in Figure 6.5 for HoPc$_2$, the spectral features of the green and blue form were maintained in LB assemblies. The SERRS spectra of HoPc$_2$ are shown in Figure 6.6. The characteristic pattern of relative intensity in the SERRS spectra of LnPc$_2$ (Ln = Nd, Sm, Er, Tm, Yb, Lu) was similar to those in Figure 6.6. The main difference between blue and green forms of HoPc$_2$ is shown in the inset of Figure 6.6. The spectral differences observed with the 647.1 nm laser line(full resonance with the blue form) persisted in the SERRS spectra obtained with the 676.4 nm laser line(full resonance with the green form) as shown in Figure 6.7. The band at 729 cm$^{-1}$ was therefore not selectively enhanced by the resonance Raman effect. The strong band at 685 cm$^{-1}$ was characteristic of the macrocycle breathing mode and is normally the most intense band in SERRS or RRS of Pc. In highly symmetric molecules(i.e., MgPc, CuPc, ZnPc, with D$_{4h}$ point group symmetry) two bands at 685 cm$^{-1}$ and 747 cm$^{-1}$ are normally seen in the macrocycle region.$^{28}$ Complexes of lower symmetry(i.e., SnPc$^{29}$ with C$_{4v}$ point group symmetry) however contain three bands at 681 cm$^{-1}$, 733 cm$^{-1}$ and 747 cm$^{-1}$. For the blue form of HoPc$_2$, three bands were observed at 685 cm$^{-1}$, 729 cm$^{-1}$ and 747 cm$^{-1}$ which suggested lower symmetry for the
Figure 6.5  Visible absorption spectra for LB monolayers of the green and blue form of HoPc₂. There were four LB on each side of the glass substrate.
**Figure 6.5**
Figure 6.6 SERRS with the 647.1 nm laser line of one LB of HoPc₂ blue and one LB monolayer of HoPc₂ green on Au island films. The expanded spectral region of the blue (A), and green (B) are shown in the inset. The ST spectra were recorded.
Figure 6.7  SERRS with the 676.1 nm laser line of one LB of HoPc$_2$ blue and one LB monolayer of HoPc$_2$ green on Au island films. The expanded spectral region of the blue (A), and green (B) are shown in the inset. The ST spectra were recorded.
Figure 6.7
blue form. The lower symmetry could result from two unequal Ln-Pc distances or the speculated hydrogen (i.e. LnPc₂H) as proposed by Moussavi et al. and Kasuga et al. A D₄d structure can be envisioned in the totally symmetric case with perfect staggered configuration and equivalent Ln-Pc distances. Differences in the Ln-Pc distance would reduce the symmetry to C₄v or even lower, and the strong band at 729 cm⁻¹, as observed in the blue form of HoPc₂, could derive its intensity from an allowed irreducible representation of a lower symmetry point group. The SERS of one LB of green and blue forms of HoPc₂ on 4 nm Ag, and excitation at 514.5 nm showed that the only difference was in the pattern of relative intensity. The observed SERRS frequencies for HoPc₂ are listed in Table 6.1.

Infrared spectra of Blue and Green Forms of LnPc₂

The pattern of relative intensities for the green and blue forms of HoPc₂ are shown in Figure 6.8. The same trend was also observed for the two forms of LnPc₂ (Ln=Nd, Sm, Er, Tm, Yb, Lu). The intense bands at 728 cm⁻¹ and 1115 cm⁻¹, which are normally observed in the IR spectra of unsubstituted bisphthalocyanines (LnPc₂), were not seen in the spectra of the octa-tert-butyl bisphthalocyanine complexes of rare earth elements. These bands have a large contribution from the C-H bending modes and are no longer observed in the LnPc₂ complexes because of the partial substitution by tert-butyl groups. The IR spectrum of the green form always contained a strong band at 1318 cm⁻¹ which decreased in relative intensity in the spectrum of the blue form. A strong band at about 1079 cm⁻¹ was always present in the blue form of LnPc₂ compounds. Walton et al. posulated
TABLE 6.1
Observed Vibrational Frequencies of the Green and Blue Form of HoPc₄₂. Infrared of a KBr pellet, and surface-enhanced Raman spectra observed with 647.1 nm excitation of one LB on Au.

<table>
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<th>Blue Form</th>
<th>Assignment</th>
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stretch 1463 m 1463 w
1463 m 1482 w 1486 m
1482 w 1510 w 1517 m Pyrrole stretch
1517 m

1525 m
1612 m 1612 m Benzene stretch
2863 w 2863 w C-H stretch
(tb) 2903 w 2902 w C-H stretch
(tb) 2957 m 2957 m C-H stretch
(tb)
Figure 6.8  Transmission infrared spectra in KBr matrix HoPc$_2$ green (A), and HoPc$_2$ blue (B). Spectral resolution was 1 cm$^{-1}$. 
Figure 6.8
that the 1070-1080 cm⁻¹ band may be characteristic of the blue form. We could not
directly compare the diagnostic 1079 cm⁻¹ band of our blue form of LnPc₂ to the
literature value of 1070 cm⁻¹ reported for the blue form of gadolinium⁴
bisphthalocyanine. The authors of the gadolinium complex published no IR spectra
and their tabulated results did not include the pattern of relative intensity. The
observed IR frequencies for HoPc₂ are also presented in Table 6.1.

**Thermal Inter Conversion of Green and Blue Forms**

It was earlier shown that sublimation of the green form of LnPc₂(Ln=Pr, Nd,
Sm, Eu) in the formation of thin films gave three products: H₂Pc and the green and
blue forms of LnPc₂. The resonance Raman spectrum of the contaminated films
contained the 730 cm⁻¹ band (medium relative intensity) characteristic of the blue form.
The 730 cm⁻¹ band was, however, not observed in the SERRS spectrum of one IR
monolayer of the green form of LnPc₂(Ln=Pr, Nd, Sm, Eu). The green→blue inter
conversion could also be observed by IR spectroscopy. The 1079 cm⁻¹ was only
observed as a contaminant when the green form of LnPc₂(Ln=Pr, Nd, Sm, Eu) was
sublimed during thin film fabrication. This band was not present in IR spectrum of
the starting material mixed with KBr to form a pellet.

Pellets of KBr containing the green or blue form of HoPc₂ were also tested for
thermal interconversion. The pellet was heated to 200 °C for four hours in an oven.
The IR spectrum that was recorded after heating the blue form was identical to the
reference spectrum recorded prior to heating. An intense band 1079 cm⁻¹ and
reduced relative intensity were however observed for the green form of HoPc₂ after
heat treatment.

Summary

The purpose of this investigation was to characterize the blue and green form of octa-tert-butylbisphtalocyanines and related compounds. The structure of the green form as LnPc$_2$(free radical) was supported by the visible band at 460-490 nm. The electronic spectra of the blue form of LnPc$_2$ does not contain the free radical band. Its absorption spectrum was similar to complexes of ZrPc$_2$ and HfPc$_2$ where two Pc$^-$ are attached to the central metal. The infrared and Raman data for the green and blue form of HoPc$_2$ were not similar and the difference was lower symmetry in the blue form. The presence of the extra hydrogen in the blue form could account for lower symmetry in the blue form. Chemical reduction of the green form with hydrazine produced the electronic spectrum of the blue form. Under these conditions the reduction was localized to occur on the Pc ligand. Addition of acid also altered the electronic spectrum or rare earth bisphthalocyanines. The green form of the early LnPc$_2$ (Ln=Pr, Nd, Sm, Eu) are not stable during evaporation and give a mixture of H$_2$Pc, blue and green form. Heating a KBr pellet mixed with the green form of HoPc$_2$ produced the characteristic IR band of the blue form.
CHAPTER 7: Chemical Reactivity of Phthalocyanine to NOX Gas

Introduction

The adsorption of small gases on phthalocyanine devices has been reviewed by Wright\(^1\) and Snow et al.\(^2\) Phthalocyanines are easily ionized and are good electron donors. The strongest interaction with oxidizing gases is predicted to occur at centres of high electron density (i.e., the nitrogen atoms of the macrocycle).\(^1\) Thin films of certain phthalocyanines doped with electron accepting gases such as nitrogen oxide and nitrogen dioxide (NO\(_2\)) produced a significant increase (ca. six orders of magnitude) in the dark dc conductivity of the film.\(^2\) In this thesis, the notation NOX is used to describe nitrogen dioxide gas in equilibrium with dinitrogen dioxide gas (i.e., 2NO\(_2\)⇌N\(_2\)O\(_4\)). Electron donor gases like ammonia were, however, used to reverse the enhanced conductivity observed in films doped with NOX gas.\(^3\) The interaction of NO\(_2\) and phthalocyanine has received the most attention in the literature.

Conductivity studies on single crystals of MPc (M=Mn, Co, Ni, Cu, Zn, Pb, H\(_2\)) exposed to NOX showed a reduction in the activation energy.\(^5\) Phthalocyanines are commonly classified as p-type semiconductors. The increase in Pc conductivity was attributed to the increase in the number of holes or charge carriers produced in the charge transfer reaction between NOX and Pc.\(^6\) The toxicity of NOX has attracted the development of Pc chemical sensors for sensitive detection of NOX in the parts per million regime. Phthalocyanine as a general class of organic molecules satisfies the requirements of a good gas sensor.\(^7\) Its conductivity varies with concentration and it can be operated at temperatures in excess of 100 °C to minimize
water condensation effects. Studies have shown that thin film morphology\textsuperscript{8,9} and the
temperature\textsuperscript{10-12} of the sensor device affect the magnitude of the electrical signal
generated by the interaction of NOX and Pc.

The most common Pc-gas conductivity device is the chemiresistor which
consists of Pc evaporated onto an array of interdigital metal electrodes. The
conductivity of the chemiresistor and changes in conductivity due to NOX adsorption
are usually measured in vacuum. A surface acoustic wave (SAW) device was also
reported to respond to changes in electrical conductivity and it was considered for Pc-
NOX sensor application.\textsuperscript{2}

The most recent Pc-NOX gas sensing device uses Langmuir Blodgett
films of phthalocyanine. The LB technique would be advantageous since it offers
reproducibility in film structure and faster response times than evaporated thin films.\textsuperscript{13}
Studies of copper mesoporphyrin IX diol exposed to 10 ppm of NOX showed that
maximum sensitivity was reached with 5-10 LB.\textsuperscript{14} The LB technique together with
molecular engineering could be used to control the architecture of the film. Baker et
al. introduced the first phthalocyanine LB chemiresistor.\textsuperscript{15} Their device consisted of
eight LB layers of an asymmetrically substituted copper phthalocyanine transferred to
a glass substrate containing interdigital aluminium electrodes. This chemiresistor
incorporating LB layers of the CuPc derivative was reported to respond linearly to
0→16 vapour parts per million of NOX. LB films of Pc have also been incorporated
into an optical sensor device which uses surface plasmon (SP) resonances as a mode of
detection.\textsuperscript{16,17} In this work a few LB of Pc were transferred to a silver slide and the
Kretschmann configuration was used to couple P-polarized light to the SP of the silver film. The effect of NOX was to change the optical constants of the composite (Pc + Ag film) and shift the SP resonance curve.

Our interest in phthalocyanine and NOX was to investigate the interaction between the two species. The objective was to apply novel and sensitive techniques that could be used to monitor the adsorption process of NOX and phthalocyanine. To the best of our knowledge there was no thorough and comprehensive spectroscopic study that examined the reactivity of Pc to NOX. There were certainly no Pc + NOX Raman reports. The work by Schoch and Temofonte demonstrated that infrared transmission spectroscopy was not drastically sensitive to the interaction of NOX and Pc.\textsuperscript{18} It should also be pointed out that the interaction between Pc and small molecules is central to the electrocatalytic\textsuperscript{19} activity of Pc complexes and the use of Pc as catalysts\textsuperscript{20}.

**Raman Scattering, SERS, and UV-Vis of Pc + NOX**

**Raman Scattering: Thin Solid Films and NOX**

Thin solid films (200 nm thickness) on Corning 7059 of ytterbium bisphthalocyanine (YbPc\textsubscript{2}) and lutetium bisphthalocyanine (LuPc\textsubscript{2}) were prepared in the evaporator designated for organic materials. Reference Raman spectra were recorded prior to exposing the films to gas. A lecture bottle of NOX was purchased from Matheson or Aldrich. The Pc sample was placed into an in-house built cylindrical glass cell and evacuated to 300 Pa prior to NOX exposure. NOX was admitted until the pressure reached 600 Pa and the film was exposed to gas for one minute before
Figure 7.1  Raman spectra (ST polarization) of 200 nm YbPc₂ before and after NOX. The 514.5 nm laser line was the excitation source.
Figure 7.1
removing the sample. Films of YbPc$_2$ and LuPc$_2$ readily decolorized from green→brown upon NOX exposure.

In Figure 7.1 the normal or off-resonance Raman spectrum of a 200 nm thin film of YbPc$_2$ before and after exposure to NOX is shown. Similar spectra were also observed for LuPc$_2$. The most drastic changes occurred in the pyrrole spectral region. The effect of NOX was to reduce the relative intensity of certain vibrational bands. The gas did not, however, change the frequency of the Raman bands. In particular, NOX strongly affected the macrocycle bands at 1407 cm$^{-1}$ and 1508 cm$^{-1}$. The latter band has been assigned to the C=C pyrrole frequency in bisphthalocyanines complexes.$^{21}$ In Raman spectroscopy a decrease in scattering intensity could be associated with a loss of electron density in the nuclei involved in the vibration. The results were then interpreted in that NO$_2$ reacted with the bisphthalocyanine complex to remove electron density particularly from the C=C pyrrole. The remarkable sensitivity of Raman spectroscopy to the interaction between Pc and NOX led to the direct application of SERS, and SERRS. In particular, the objective was to study the adsorption and desorption processes on a single monolayer of phthalocyanine.

**SERS: Phthalocyanine LB Layers and NOX**

The work began with monolayer formation of LuPc$_2$ and YbPc$_2$. Both bisphthalocyanine complexes were spread from chloroform onto a subphase maintained at 20 °C. The isotherms are shown in Figure 7.2. In the preparation of LB layers, the Langmuir layer was compressed to a constant surface pressure of 10 mN/m, and
Figure 7.2  Surface pressure-area isotherms of LuPe₂ and YbPe₂.
Figure 7.2

LuPc₂

Surface Pressure (mN/m) vs. Area/Molecule (nm²)

YbPc₂

Surface Pressure (mN/m) vs. Area/molecule (nm²)
Figure 7.3  SERS(λ=514.5 nm, ST polarization) spectra of one mixed LB of YbPc₂ before and after exposure to NOX gas
One Mixed LB YbPc$_2$

1 LB + NOX

1 LB

Figure 7.3
transferred at a rate of 4.8 mm/min to Ag-Sn spheres and hydrophillic glass slides. The transfer ratio for neat LB of LuPc₂ and YbPc₂ was on the average .9/layer. Petty et al. found that mixed monolayers of YbPc₂ were better suited for LB films. We found that mixed monolayers in the mole ratio bisphthalocyanine:arachidic acid=3:1 were easy to work with and they gave near unity transfer ratio. The purpose of arachidic acid was to lubricate the otherwise rigid and fragile monolayer on the subphase. The SERS of one mixed LB(3:1) of YbPc₂ on Ag-Sn spheres before and after the reaction with NOX is shown in Figure 7.3. The effect of NOX was similar to that observed for thin film(Figure 7.1). Unfortunately NOX also tarnished and reacted with the Ag in the SERS substrate. The reaction was so severe it caused a 4 nm Ag island film to turn transparent. It was shown by SERS that NOX attacked Ag powder to produce Ag₂O, AgNO₂ and AgNO₃. The SERS of Ag-Sn spheres exposed to NOX is presented in Figure 7.4. The bands at 812(NO₂⁻), 955(Ag₂O), and 1039(NO₃⁻) cm⁻¹ were attributed to the species formed in the reaction between NOX and silver. Though these bands were not present in the SERS spectra of one mixed LB of LuPc₂ or YbPc₂ exposed to NOX, an alternative SERS substrate was considered. Au island films of 4 nm thickness were extremely stable in the presence of NOX. They did not discolour and the uv-vis spectra before and after exposure were almost identical. The Au island film was suitable to monitor the desorption of NOX from Pc. The 647.1 nm laser line was used to excite the localized surface plasmons of the Au island film. Excitation at 647.1 nm for a Au island film covered with Pc produced SERRS.
Figure 7.4  SERS($\lambda=514.5$ nm, ST polarization) spectra of Ag-Sn spheres exposed to NOX gas
NOX + Ag-Sn Spheres

Figure 7.4
Figure 7.5 Surface pressure-area isotherms for CuPcTTb and H₂PcTTb
SERRS was used to monitor the chemical adsorption of NOX on one mixed(Pc:arachidic acid=3:1) monolayer of metal free tetra-tert-butylphthalocyanine(H$_2$PcTTb), copper tetra-tert-butylphthalocyanine(CuPcTTb), LuPc$_2$ and YbPc$_2$. The preparation of H$_2$PcTTb$^{23,26}$, and CuPcTTb$^{17,27,28}$ monolayers was previously reported by other groups. The pressure-area isotherms at 20 °C for neat CuPcTTb and neat H$_2$PcTTb spread from toluene solutions are presented in Figure 7.5. The isotherm of H$_2$PcTTb reported by Kovacs$^{23}$ et al. was similar to the one shown in Figure 7.5. The isotherm for CuPcTTb prepared in our laboratory was however very different than those reported in the literature.$^{17,27}$ Neat monolayers of H$_2$PcTTb, CuPcTTb, YbPc$_2$ and LuPc$_2$ were also transferred to glass slides. All monolayer work in this section was carried out at a constant surface pressure of 15 mN/m and 6 mm/min transfer rate.

Electronic absorption spectroscopy for LB films on glass was also used to monitor the adsorption and the desorption of NOX. The procedure used to treat LB films(on glass and on Au films) with NOX was identical to that described for thin solid films. Desorption studies consisted of evacuating the exposed film at 3 Pa for three hours.

In Figure 7.6 the SERRS for one mixed LB of H$_2$PcTTb before and after exposure to NOX is shown. Frequencies were observed with the same value in both spectra and the relative intensities showed only minor changes. The absorption spectrum for 8 LB on glass is shown in Figure 7.7. The upper spectrum in Figure 7.7 is typically observed in the solution$^{29}$ or the crystal$^{30}$ state of H$_2$Pc derivatives.
Figure 7.6  SERRS(ST polarization) spectra of one mixed LB of H$_2$PcTTb on a 4 nm Au island film (A) and after exposure to NOX gas (B)
Figure 7.6

$H_2PcTTb$

Wavenumber (cm$^{-1}$)
Figure 7.7  Electronic spectra of $\text{H}_2\text{PcTTb}$ in solution (A), eight LB monolayers on glass (B), and eight LB exposed to NOX gas (C)
Figure 7.7
The middle absorption spectrum contained a unique single peak for the Q-band. This band can be attributed to the $\pi$-$\pi$ aromatic interactions in the LB film. The last absorption spectrum was recorded for the LB assembly exposed to NOX. The Q-band was split into two broad bands whose maximum corresponded to the band in the untreated LB sample (near 600 nm) and to the solution band (near 700). The splitting of the Q-band in the LB film suggested that NOX caused a disruption in the $\pi$-$\pi$ aromatic interactions. NOX also self-desorbed at room temperature and normal pressure and the middle spectrum in Figure 7.7 was recovered. The results of UV-vis and SERRS therefore strongly suggested that NOX was physisorbed to H$_2$PcTTb.

The reversible adsorption of NOX on one mixed LB of CuPcTTb is shown in Figure 7.8. SERRS spectra A) and B) were taken before and after exposure to gas respectively. The bottom spectrum was recorded after the sample was put in vacuum. The effect of NOX in spectrum B) was to reduce the relative intensity of the macrocycle breathing at 688 cm$^{-1}$ and the pyrrole stretching band at 1530 cm$^{-1}$. Since Raman intensities originate from changes in the polarizability with the corresponding normal coordinate, an intensity decrease in a Raman band can be attributed to a loss of electronic density in the normal coordinate. The spectral evidence suggested that NOX polarized both the pyrrole and macrocycle groups. The interaction was, however, not strong enough to cause a shift in the vibrational frequencies. The electronic spectra for CuPcTTb and its interaction with NOX is presented in Figure 7.9. In the middle spectrum the Q-band was observed to split when gas was added. The result was similar to that observed in H$_2$PcTTb where NOX acted to only disturb
Figure 7.8  SERRS spectra (ST polarization) of one mixed LB of CuPcTTb on Au film (A), after exposure to NOX gas (B), and after desorption of NOX gas (C)
Figure 7.9  Absorption spectra of 5 LB monolayers of CuPcTTb (A), after NOX adsorption (B), and after gas desorption (C)
Figure 7.9
the $\pi-\pi$ aromatic interactions. The bottom spectrum showed that NOX was desorbed from the multilayer assembly when placed in vacuum for three hours. The observed difference in the SERRS data for $\text{H}_2\text{PcTTb}$ and $\text{CuPcTTb}$ was therefore attributed to the metal. Indirect evidence for the proposed interaction was provided by Electron Paramagnetic Resonance (EPR) measurements in toluene solution. The EPR signal of $\text{CuPcTTb}$ became silent after addition of NOX. The latter results suggested a direct NO$_2$-metal interaction. A diagram of the proposed molecular interaction between NO$_2$ and CuPcTTb is given in Figure 7.10.

The third type of phthalocyanine was a bisphthalocyanine where one atom was coordinated to two Pc macrocycles. The effect of NOX towards LuPc$_2$ or YbPc$_2$ was more drastic than in the previous two cases. The SERRS progress of one mixed LB of YbPc$_2$ exposed to NOX are presented in Figure 7.11. The SERRS spectrum before and after gas adsorption was more pronounced than in CuPcTTb or $\text{H}_2\text{PcTTb}$. The weak band at 1549 cm$^{-1}$ became the most dominant band in NOX exposed sample. In the resonance Raman studies of lanthanide bisphthalocyanines$^{21}$ the band at 1524 cm$^{-1}$ was assigned to C=N aza group. Based on the earlier discussion that NO$_2$ attacked the metal in CuPcTTb, the middle spectrum could be attributed to NO$_2$ coordinating to the lanthanide and decoupling the aza stretching vibration. The polarization effect of NO$_2$ was also sensed by the macrocycle band at 679 cm$^{-1}$. The NO$_2$-YbPc$_2$ adduct could thus be envisioned in terms of NO$_2$ attacking the metal and drawing electronic charge from the inner macrocycle ring. The SERRS in the bottom spectrum showed that the adsorption process was completely reversible. The electronic spectra of YbPc$_2$
Figure 7.10  Molecular diagram of CuPcTTb with one NO$_2$ molecule. The tert-butyl groups are represented by Tb
Figure 7.10
are shown in Figure 7.12. In the middle spectrum the effect of NOX was to shift the Q-band to higher wavelengths. The red shift was characteristic of the oxidized form of rare earth bisphthalocyanine.\textsuperscript{22} For YbPc\textsubscript{2} or LuPc\textsubscript{2} the SERRS and UV-vis absorption spectra showed that desorption of gas was completely reversible.

In summary reversible adsorption of NOX on one mixed LB of CuPcTTb, YbPc\textsubscript{2}, and LuPc\textsubscript{2} was clearly observed in the SERRS spectra. The oxidized macrocycle was identified in the absorption spectrum of rare earth bisphthalocyanine + NOX. Oxidation of the macrocycle resulted in a polarization of the macrocycle where the C=N aza groups were affected. The relative strength of the interaction between NOX and H\textsubscript{2}PcTTb, CuPcTTb and rare earth bisphthalocyanine(LuPc\textsubscript{2}, YbPc\textsubscript{2}) was weak, strong and very strong respectively. The drastic changes in the SERRS and absorption spectra observed for CuPcTTb and YbPc\textsubscript{2}( or LuPc\textsubscript{2}) support that the adsorption site was the metal centres.

The SERS data of one mixed LB of YbPc\textsubscript{2}(Figure 7.3) and the Raman spectrum of the 200 nm film of YbPc\textsubscript{2}(Figure 7.1) showed that the NOX also polarized the C=C in pyrrole. The accumulated data in RS, SERS and SERRS supported the notion that NOX strongly affected the macrocycle bands. In particular the C=C(pyrrole) and C=N(aza group) were mostly influenced by the oxidising gas. This Raman study thus demonstrated selective tuning in the reaction of NOX + Pc. The changes in C=C(pyrrole) were clearly observed with $\lambda=514.5$ nm(off-resonance excitation), while changes to C=N(aza group) were observed with $\lambda=647.1$ nm (resonance Raman scattering).
Figure 7.11  SERRS spectra (ST polarization) of one mixed LB of YbPc$_2$ on Au island (A), after NOX adsorption (B), and after desorption (C)

The inset shows the expanded pyrrole frequency region.
YbPc$_2$

Figure 7.11
Figure 7.12  Absorption spectra of three LB of YbPc$_2$ on glass (A), after NOX adsorption (B), and gas desorption (C)
Figure 7.12
Oxidation and Reduction of Bisphthalocyanine

During the Raman work of Pc + NOX it became obvious that the redox properties of Pc were intrinsically important. We selected to investigate qualitatively the redox properties of a rare earth bisphthalocyanine since they showed the most changes to NOX. The chemical oxidation(with NOX) and reduction(with hydrazine) for a solution of LuPc₂ in chloroform is shown in Figure 7.13. The oxidized(spectrum B) form of LuPc₂ was observed with red shifted Q-band. Reduced(spectrum C) LuPc₂ however showed a blue shifted Q-band. Electrochemical oxidation also produced similar results to the chemical oxidation.

The spectroelectrochemical analysis of a 18 nm evaporated film of LuPc₂ on an indium tin oxide(ITO) working electrode is shown in Figure 7.14. The counter electrode was a Pt wire and the electrolyte was 1.0 M KCl at pH 1.23. The potential difference was measured with respects to the standard calomel reference electrode. At 0 V the Q-band was positioned at ca. 640 nm. At +0.960 V the Q-band clearly showed red shifted to ca. 705 nm. Our spectroelectrochemical results were also similar to those published in the literature.³¹,³²

It can be concluded that the electronic absorption band due to electrochemical oxidation of the macrocycle was the same band as in the chemical oxidation(Figure 7-13B) with NOX. The NO₂ acted as an electron acceptor in the formation of NO₂⁻ LnPc₂ adduct.
Figure 7.13  Chemical oxidation and reduction of LuPc$_2$ in chloroform, green starting material (A), oxidation with NOX gas (B), reduction with hydrazine (C)
Figure 7.13
Figure 7.14 Absorption spectra of a 18 nm LuPc$_2$ film at (I) 0 V, 
(II) +0.488 V, and (III) +0.960 V. The potential are referenced against the standard calomel electrode.
Figure 7.14

LuPc$_2$ Film on ITO
CONCLUSIONS

In the present thesis surface enhanced {resonance} Raman scattering(SE{R}RS) was reported for various phthalocyanine derivatives. SERS and SERRS were reported for VOPc evaporated onto a Ag substrate. The IR spectrum of a thin film of VOPc sublimed onto KBr crystal showed preferred molecular orientation. The Raman data were complemented with the infrared data and the symmetry assignment of bands in VOPc was attempted. Favourable excitation with P-polarized light was observed for ultrathin films of VOPc adsorbed onto highly reflecting surfaces. Davydov splitting was also seen in the Raman spectra of these samples.

SERRS was used to characterize one Langmuir-Blodgett monolayer of VOPcTTb and ZnPcTTb. The SERS and SERRS of VOPcTTb and ZnPcTTb(or any of the Pcs in this thesis) showed no spectral evidence of strong interaction between Pc and the SERS substrate. The SERS spectra showed no shift in the molecular frequencies of Pc and the Pc molecules were therefore considered to be physisorbed onto the substrate. The SERRS spectra were depolarized in that the SS, SP, PP, and PS components gave identical spectra. The vibrational characterization of VOPcTTb and ZnPcTTb was also given.

Enhanced absorption was observed in the energy transfer experiments of one LB of ZnPcTTb(acceptor layer) and one LB of PBDC(donor layer) on Ag-Sn spheres. Only when the acceptor layer was directly adjacent to the Ag-Sn spheres was the excimer emission of PBDC totally quenched(i.e., there was complete energy transfer). Spectroscopic tuning was also identified in the energy transfer experiments. Selecting
a laser line in resonance with either ZnPcTTb or PBDC exclusively produced SERRS of either the donor or acceptor layer.

The LB technique was used to control surface coverage. The coverage dependence on the SERRS intensity using mixed LB films of(VOPcTTB and arachidic acid) showed a maximum at submonolayer coverage. The negative dependence of SERRS with increasing surface coverage was rationalized in terms of de-exication or quenching of the surface plasmons.

SERRS, IR, and uv-vis were used to investigate the spectral features of the green and blue form of rare earth bisphthalocyanines. The differences observed in the IR and SERRS spectra of the green and blue species could be attributed to differences in molecular symmetry.

The reversible adsorption of NO₂ on a single LB of H₂Pc, CuPcTTb and YbPc₂(or LuPc₂) was monitored by the highly sensitive SERRS technique. The changes in the SERRS spectra were attributed to NO₂ interacting with the metal in CuPcTTb, YbPc₂ or LuPc₂ and polarizing the macrocycle vibrations. The data in the electronic spectra showed that NO₂ acted as an oxidant to lanthanide bisphthalocyanine complexes.
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Publications


