A study of thin solid organic films by Fourier transform surface-enhanced Raman scattering and atomic force microscopy.

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A STUDY OF THIN SOLID ORGANIC FILMS BY FOURIER TRANSFORM SURFACE-ENHANCED RAMAN SCATTERING AND ATOMIC FORCE MICROSCOPY

by

Carol Ann Jennings

A Dissertation Submitted to the Faculty of Graduate Studies and Research 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

Windsor, Ontario, Canada

1994

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ISBN 0-612-01456-8
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ABSTRACT:

Fourier transform surface-enhanced Raman scattering (FT-SERS) in the near-infrared and atomic force microscopy (AFM) have been used to characterize thin solid organic films. The pigments studied were phthalocyanines and perylenes which have applications as charge generation materials in organic photoreceptors. Techniques for film preparation such as Langmuir-Blodgett (LB) deposition and vacuum evaporation are discussed. The literature was reviewed for FT-Raman spectroscopy, surface-enhanced Raman spectroscopy and atomic force microscopy. A model according to Gersten and Nitzan was applied and SERS enhancement factors in the near-IR were calculated for various metal hemispheroids. FT-Raman spectra were presented for thin vacuum evaporated films of titanyl phthalocyanine (TiOPc) and vanadyl phthalocyanine (VOPc) prepared on glass substrates held at a range of temperatures during deposition. The frequencies were assigned and different polymorphic forms were apparent. In order to improve the signal intensity, thin films were deposited on metal substrates which produced SERS in the near-IR. The substrates selected for FT-SERS were vacuum evaporated metal island films of copper and gold of 20 nm mass thickness prepared with the glass substrate held at 200°C. Thin films of 3 nm mass thickness of VOPc and 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA) were vacuum evaporated onto the metal island substrates and produced FT-SERS with an enhancement factor of approximately 100. Single Langmuir-Blodgett monolayers of phthalocyanines and perylenes on the metal island substrates gave FT-SERS. The FT-SERS spectra of vacuum evaporated films and LB monolayers were compared with visible laser excitation. Phthalocyanines and
perylene absorb in the visible region and produced surface-enhanced
resonance Raman scattering (SERRS). The morphology and structure of single
LB monolayers on mica and Si(111) wafers have been determined using atomic
force microscopy. AFM results for a perylene on Si(111) showed good
agreement with the orientation inferred from reflection-absorption infrared
spectroscopy and transmission IR measurements.
ACKNOWLEDGEMENTS

I would like to express sincere thanks to my two supervisors: Dr. R. Aroca from the University of Windsor and Dr. G. J. Kovacs of the Xerox Research Centre of Canada.

Xerox Corporation should be acknowledged for financial support and its commitment to employee training and development. In particular, senior staff and area managers have been very supportive at XRCC. I also thank NSERC of Canada for financial support.

Dr. H. Mizes of Webster Research Center provided technical assistance and suggestions related to atomic force microscopy. Dr. K. Kjoller and Mr. C. Callahan of Digital Instruments are acknowledged for images obtained on the Nanoscope III MultiMode AFM. Mr. P. Gerroir of XRCC with Mr. J. Cole and Mr. D. Edgar of Nissei Sangyo supplied the field emission scanning electron micrograph of a gold film. Dr. C. Tripp of XRCC made the initial proposal for an FT-Raman system and gave useful advice. Mr. C. K. Hsaio of XRCC is acknowledged for his comments and assistance regarding the section on polymorphs of titanyl and vanadyl phthalocyanine. I thank Dr. T. Bluhm of Xerox for the X-ray powder diffraction patterns of titanyl phthalocyanine polymorphs.

Finally, I would like to thank my fiancé, family, friends and colleagues who provided moral support throughout this endeavor.
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<td>AFM</td>
<td>atomic force microscopy</td>
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<tr>
<td>ASR</td>
<td>atomic scale roughness</td>
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<tr>
<td>BTA</td>
<td>benzotriazole</td>
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<td>(BuO)$_2$H$_2$Pc</td>
<td>1,4,8,11,15,18,22,25-octabutoxy-29H,31H-phthalocyanine</td>
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<tr>
<td>CCD</td>
<td>charge-coupled device</td>
</tr>
<tr>
<td>CGL</td>
<td>charge generator layer</td>
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<tr>
<td>CT</td>
<td>charge transfer</td>
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<tr>
<td>CT DNA</td>
<td>calf thymus deoxyribonucleic acid</td>
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<tr>
<td>CTL</td>
<td>charge transport layer</td>
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<tr>
<td>CTM</td>
<td>charge transport molecule</td>
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<tr>
<td>CuTTPc</td>
<td>copper tetra-tertiary-butyl-phthalocyanine</td>
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<tr>
<td>DOX</td>
<td>doxorubium</td>
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<tr>
<td>EF</td>
<td>enhancement factor</td>
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<tr>
<td>EM</td>
<td>electromagnetic</td>
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<tr>
<td>FAD</td>
<td>flavine adenine dinucleotide</td>
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<tr>
<td>FT</td>
<td>fourier transform</td>
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<tr>
<td>HPTCDE</td>
<td>N-hexyl-(N')-ethyl-3,4:9,10-perylenetetracarboxylic diimide</td>
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<td>HPTCO</td>
<td>N-hexyl-3,4:9,10-perylenetetracarboxylic monoimide</td>
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<tr>
<td>HOMO</td>
<td>highest occupied molecular orbital</td>
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<tr>
<td>HOPG</td>
<td>highly oriented pyrolytic graphite</td>
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<tr>
<td>LAM</td>
<td>longitudinal acoustic mode</td>
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<tr>
<td>LB</td>
<td>Langmuir-Blodgett</td>
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<tr>
<td>LUMO</td>
<td>lowest unoccupied molecular orbital</td>
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<tr>
<td>Nc</td>
<td>napthalocyanine</td>
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<tr>
<td>Nd/YAG</td>
<td>neodymium doped yttrium-aluminum garnet</td>
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<tr>
<td>Nd/YLF</td>
<td>neodymium doped lithium fluoride</td>
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<tr>
<td>NEP</td>
<td>noise equivalent power</td>
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<tr>
<td>ORC</td>
<td>oxidation-reduction cycle</td>
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<tr>
<td>Pc</td>
<td>phthalocyanine</td>
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<tr>
<td>PcLuPc$^t$</td>
<td>4,4',4''-tetra-tertiary-butyl lutetium diphthalocyanine</td>
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<tr>
<td>PMT</td>
<td>photomultiplier tube</td>
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<td>PTCDA</td>
<td>3,4:9,10-perylenetetracarboxylic dianhydride</td>
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<tr>
<td>PTCDI</td>
<td>perylenetetracarboxylic diimide</td>
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<tr>
<td>PDCI</td>
<td>(N)-octyl-3,4-pyrene dicarboximide</td>
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<td>Abbreviation</td>
<td>Full Form</td>
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<td>PPTCDE</td>
<td><em>N</em>-pentyl-<em>N’</em>-ethyl-3,4:9,10-perylene tetra-carboxylic diimide</td>
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<td>PTFE</td>
<td>polytetrafluoroethylene</td>
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<tr>
<td>RAIRS</td>
<td>reflection-absorption infrared spectroscopy</td>
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<td>RR</td>
<td>resonance Raman</td>
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<td>SEM</td>
<td>scanning electron microscopy</td>
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<tr>
<td>SERRS</td>
<td>surface-enhanced resonance Raman scattering</td>
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<tr>
<td>SERS</td>
<td>surface-enhanced Raman scattering</td>
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<tr>
<td>STM</td>
<td>scanning tunneling microscopy</td>
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<tr>
<td>TEM</td>
<td>transmission electron microscopy</td>
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<tr>
<td>TTA</td>
<td>tritolyltriazole</td>
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<tr>
<td>(t-bu)₄H₂Pc</td>
<td>tetra-tertiary-butyl metal-free phthalocyanine</td>
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<tr>
<td>UHV</td>
<td>ultrahigh vacuum</td>
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<tr>
<td>XRPD</td>
<td>X-ray powder diffraction</td>
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CHAPTER 1: INTRODUCTORY REMARKS, ELECTROPHOTOGRAPHY, AND OBJECTIVES

1.1. INTRODUCTORY REMARKS

The work described here was made possible through the cooperation of the University of Windsor and Xerox Research Centre of Canada. This arrangement has been particularly attractive allowing the author to experience the best of both worlds. The material in this dissertation does not contain any Xerox proprietary information. The background related to electrophotography has been published in the external literature. The experimental chapters were cleared through the legal department and reviewed by scientific staff.

The first chapter explains why thin solid organic films are of interest scientifically and commercially. The objectives of the research are described later in this chapter. Chapter 2 is a general review of the main techniques used in the experimental section. The preparation of thin solid films by Langmuir-Blodgett (LB) deposition and vacuum evaporation are discussed in general terms. Instrumental considerations and theory are presented for Raman spectroscopy, resonance Raman (RR) spectroscopy and FT-Raman spectroscopy. Chapter 2 also contains information about atomic force microscopy (AFM) and an extensive literature review had been conducted prior to carrying out AFM experiments. Chapter 3 is a literature review and presentation of theories for surface-enhanced Raman scattering (SERS). Some model calculations were carried out in Chapter 3 to predict what sort of metals would be suitable to prepare island films for SERS in the near-IR region. Chapter 4 includes FT-
Raman studies carried out on thin vacuum evaporated films of vanadyl phthalocyanine (VOPc) and titanyl phthalocyanine (TiOPc) at various temperatures. In Chapter 5, vacuum evaporated metal island films of copper and gold are prepared and used as FT-SERS active substrates. Thin solid films of organic pigments were deposited on the metal island films by vacuum evaporation. Langmuir-Blodgett monolayers were also deposited on the metals. FT-SERS experiments were carried out using the 1064.1 nm Nd/YAG laser line and surface-enhanced resonance Raman scattering (SERRS) was obtained using visible laser excitation. Following the trend of moving from larger structures to smaller structures, Chapter 6 describes atomic force microscopy (AFM) of Langmuir-Blodgett monolayers on atomically flat substrates. The AFM results were complemented by infrared spectroscopy providing information on molecular orientation. Chapter 7 is an overall summary of the work and ideas on future areas for study are presented.

A brief overview of electrophotography follows and in particular the role of a photoreceptor. The materials used in the experimental section fall under the general category of phthalocyanines and perylenes. Specific compounds from both of the classes have been used in charge generator layers (CGLs) of photoreceptors.

1.2. ELECTROPHOTOGRAPHY AND ORGANIC PHOTOCONDUCTORS

The inventor of electrophotography was Chester F. Carlson and the original work was described in his patent in 1942. This is an example of persistence as Carlson disclosed and demonstrated the crude process to 20 major corporations. There was a lack of interest but eventually development
was carried out at the Battelle Memorial Institute followed by the Haloid Company which became Xerox Corporation\textsuperscript{2}. The commercialized process became known as "xerography". Electrophotography is the technology underlying most copiers and printers.

The first step in the xerographic process is photoreceptor charging. Either positive or negative charges can be laid down on the surface of a photoreceptor depending on its composition by corona discharge. An opposite charge is present at the conductive substrate producing an electric field across the photoreceptor. The photoreceptor is imagewise exposed. It is important to couple the wavelength(s) of light for imaging with the spectral sensitivity of the photoreceptor. The areas of the photoreceptor which have been exposed to light discharge due to the creation of electron-hole pairs and charge transport. The areas which have not been exposed, retain the surface charge and replicate the optical image. The charge pattern is called a "latent image". The latent image is developed using charged toner. Toner typically contains pigmented particles in a thermoplastic. A two-component developer consists of carrier beads which transport and charge the toner particles. The electrostatic image on the photoreceptor is developed using toner. A piece of paper is transported and covers the developed area. A charge can be applied to the back of the paper during image transfer to attract the toner. The toner is fused to the paper by heat and/or pressure. The photoreceptor can be cleaned using a blade and any residual charge removed by an intense erase exposure.

A printer requires more computing power than a copier. A digital image is dumped from the computer to the printer. A laser light source projects an
image onto the photoreceptor surface. The lasers which are commonly used emit in the red to near-IR region and most xerographic printers use a rotating polygon\textsuperscript{2}. Copier and printer technology draws from many fields; physics, chemistry, engineering and computer science.

One general structure of a photoreceptor is presented in Figure 1-1. A conductive substrate is the starting point in the fabrication. Common substrates are aluminum drums or metallized polyester. A thin layer may be present which blocks charge injection from the conductive layer, smooths out the surface or functions as an adhesive for the next layer. The next layer is the charge generator layer (CGL). A photoactive material is present in a polymeric binder material. A dual-layer photoreceptor has a charge transport layer (CTL) separate from the CGL layer. The charge transport layer commonly contains a charge transport molecule within a polymeric matrix. There could also be a optional overcoat layer to provide abrasion resistance.

Coating techniques for solutions or pigment dispersion have been described\textsuperscript{3}. Photoreceptor coatings are done under highly controlled conditions most commonly on a web or continuous roll format. The coating solution is pumped through a slot and delivered to the web. Important factors to consider are web speed, vibrations, drying conditions, cleanliness and the nature of the coating solution to minimize defects. The charge generator layer can be coated as a dispersion. The pigment particle size must be reduced by milling the pigment with binder in a solvent. There are many devices available to reduce particle size such a ball mills, roller mills, attritors and sand or salt mills\textsuperscript{3}. A less common method for preparing charge generator layers is vacuum evaporation of pigments. Vacuum evaporation allows thin layers of controlled
thickness with few defects to be prepared. The downside is that the evaporated films are usually amorphous and have to be converted to a crystalline state either by heat or solvent treatment\(^3\). Methods for coating drums are dip coating, spraying and ring coating.

A brief review of organic charge generation materials will be presented. Many of these pigments exhibit polymorphism. The polymorphic form affects the physical properties. It is important to monitor particle size and morphology. The classes of materials are: aggregate materials such as thiapyrylium salts, polyazo compounds, phthalocyanines, squaraines, perylenes and other polycyclic aromatics including dibromoanthanthrone. Representative pigments are shown in Figures 1-2 and 1-3. Azo pigments are named according to the number of azo groups: bis, tris, tetrakis ... and are easily synthesized. A general phthalocyanine structure has a central atom(s) represented by M, where M = H\(_2\), Cu, Mg, Zn, Pb, Sn, TiO, VO, Sn, Cl\(_2\)Al, Cl\(_2\)In, (OH)\(_2\)Ge etc. Depending on the central atom(s), phthalocyanines can have absorbance in both the visible and near-IR regions. Perylenes are usually reddish in appearance and are diimides of 3,4:9,10-perylenetetracarboxylic acid. Squaraines that have aggregated in charge generations layers have absorption in the 400-1000 nm region\(^3\). Some further structures of polycyclic charge generation materials appear in Figure 1-3. The crystalline nature of the charge generation material is vital and the tendency to form molecular stacks. It can be seen that the charge generation materials are highly conjugated systems and a number of resonance structures can be drawn.

Two photogeneration mechanisms have been described\(^2\). A singlet-state exciton can be produced by photon absorption. The electron and hole
Figure 1-1: A general structure of a photoreceptor showing the conductive substrate, blocking/adhesive layer, charge generator layer and charge transport layer.
A Photoreceptor Device

- Charge Transport Layer
- Charge Generator Layer (contains pigment)
- Adhesive + Blocking Layer
- Conducting Substrate
Figures 1-2 & 1-3:

Structures of organic charge generation materials³,⁴: a) polyazo compound, b) a phthalocyanine, c) a perylene, d) a squaraine, e) dibromoanthanthrone, f) thioquinacridone, g) dithioketopyrrolopyrrole, h) imidazole perinone, i) bis(benzimidazole)perinone, j) pyranthrone
Figure 1-2

a) A bisazo compound Chlorodiane blue

b) Phthalocyanines

c) Perylenes

d) Squaraines
$X = \text{H, OH or CH}_3$
dissociate by an Onsager-like mechanism. Alternatively, the charge migrates to
the pigment-particle surface and charge exchange occurs with a charge
transport molecule. The charge transport layer (CTL) shown in Figure 1-1
usually consists of a molecular charge transport molecule doped in a polymer.
Charge transport molecules (CTMs) are either hole transporting or electron
transporting. The hole transport molecules are good electron donors while the
electron transport molecules are good electron acceptors. Specific examples of
CTMs can be found in the literature2-4. Some classes of materials used for hole
transport include: arylalkanes, arylamines, hydrazones, poly(phenylene-
vinylene)s, polysilylenes, polygermylenes, poly(N-vinylcarbazole) and
pyrazolines. Examples of electron transport materials are: diphenylquinones,
2,4,7-fluorenone and (4-nitro-butoxycarbonyl-9-fluorenylidene) malonitrile.

Photoreceptors are often described by the energy required to discharge
to 50% of the original voltage, $E_{1/2}$ having units of ergs/cm². Another
parameter often quoted is the dark decay in units of volts/second. The
photoreceptor is charged to a desired voltage, undergoes some dark decay and
is photodischarged by the appropriate wavelength and intensity of light. A
residual voltage remains which is erased by a strong exposure. In practice, a
photoreceptor is cycled over a hundred thousand times. The trend has been to
produce materials which are more robust and with near-IR photosensitivity for
printer applications.

1.3. OBJECTIVES

This author had some previous experience characterizing bulk samples by
FT-Raman spectroscopy. In most cases, the technique was relatively easy and
reduced fluorescence. Good spectra could be obtained from a variety of
materials often with little or no sample preparation. The challenge was to be able to look at small quantities of material. The first objective of the experimental work was to investigate FT-Raman spectroscopy as a technique for obtaining vibrational information from thin solid films. The approach taken was to prepare vacuum evaporated films of titanyl phthalocyanine and vanadyl phthalocyanine 100 nm thick on glass held at various temperatures. The FT-Raman vibrational frequencies could be assigned to functional groups. Both TiOPc and VOPc exist in a number of polymorphic forms and these could be identified using FT-Raman spectroscopy based on the observed frequencies and relative intensities.

The next goal was to move to ultrathin Langmuir-Blodgett monolayers. Raman scattering from a single LB monolayer is extremely weak due to the small number of molecules and small Raman scattering cross section. Single LB monolayers have been measured using normal Raman spectroscopy with visible laser excitation and a charge-coupled device (CCD). Good FT-Raman spectra could not be obtained for single LB monolayers of phthalocyanines and perylenes with the system used here employing the 1064.1 nm Nd/YAG laser line and an InGaAs detector. Therefore, the objective was to find a way of increasing the FT-Raman signal from an LB monolayer. Since the materials of interest did not have any absorption in the near-IR, the resonance Raman effect could not be used to increase the signal intensity. The method selected to increase the signal intensity from single LB monolayers was Fourier transform surface-enhanced Raman scattering (FT-SERS).

In order to observe FT-SERS, it was necessary to find an appropriate substrate to enhance the Raman signal in the near-IR region for subsequent
deposition and detection of thin organic films. At the time this experimental work was initiated, other groups (cited later), were using electrochemically roughened metal electrodes and metal colloids for FT-SERS. It was decided that the path taken would be to prepare metal island films as FT-SERS active substrates. Computer calculations based on one of the theories for SERS were carried out to estimate enhancement factors for various metals in the near-IR. Vacuum evaporated copper and gold metal island films were prepared of various mass thicknesses. What remained was to demonstrate experimentally that FT-SERS could be obtained of thin organic films on the metal island structures. Initially pigments were vacuum evaporated to a mass thickness of 3 nm on the metal island films. Then single LB monolayers were transferred to the metal island substrates. The greatest enhancement was obtained using 20 nm thick metal island films that had been vacuum evaporated onto glass substrates held at 200°C.

A final objective was to learn about the morphology and structure of Langmuir-Blodgett monolayers of organic pigments using atomic force microscopy. Methods are needed to study aging, defects, domains and local structures in order for LB monolayers to have commercial applications. The structure of an extremely thin film does not necessarily correspond to that of the bulk material and it was thought that AFM would be useful for looking at different polymorphic forms. AFM does not require that the sample be conductive and dimensions ranging from microns to nanometers can be imaged. Initially it was believed that molecular resolution would be achieved but this was not the case. A review of the literature revealed many AFM papers about fatty acids and relatively few results reported for LB monolayers of aromatic organic molecules. Substrates had to be selected for AFM
experiments that were atomically flat and allowed LB film transfer. The best conditions had to be determined for imaging so that the LB monolayer was not scraped from the substrate.

At the time the AFM experiments were carried out, the University of Windsor and the Xerox Research Centre of Canada did not have an AFM instrument. Measurements were made on a Park Scientific AFM instrument at the Xerox Webster Research Center. The author also had the opportunity to take samples to the University of Toronto when representatives from Digital Instruments were visiting. As this dissertation neared completion, the Xerox Research Centre of Canada purchased an AFM instrument. Scanning probe microscopy opens up many possibilities not only for characterization but also in the area of building up structures from the bottom up by manipulating single atoms for instance.
Chapter 1: References


CHAPTER 2: EXPERIMENTAL TECHNIQUES
(BACKGROUND): FILM DEPOSITION, RAMAN
SPECTROSCOPY AND ATOMIC FORCE MICROSCOPY

2.1. LANGMUIR-BLODGETT FILMS

An historical perspective has been given concerning the development of
Langmuir-Blodgett (LB) films\textsuperscript{1-3}. It was observed very early on that oil spread
on water very thinly and had a calming effect on waves at sea. Lord Rayleigh
suspected that olive oil on water had molecular thickness. The surface tension
of water was lowered by contaminating oil. Agnes Pockels wrote to Rayleigh
about her methods to measure molecular sizes. The set-up used by Pockels was
very crude but has similarities to modern day LB troughs. A rectangular trough
containing water was used, a tin barrier and a button was the disk to measure
surface tension. Pockels produced the first isotherms which are plots of surface
pressure versus area. Pockels noted that there were two distinct conditions. In
the first situation, the displacement of the barrier had no effect on the surface
tension. In the second one, any small change in the barrier markedly affected
the surface tension. It is now known that this is due to changes in the
orientation of the molecules and packing to form a solid-like state. Others\textsuperscript{1}
confirmed that materials were spreading to monolayer thickness. Non-polar
oils did not behave the same way as polar ones. Irving Langmuir believed the
molecular forces were short-ranged. Langmuir received a nobel prize in 1912
for his many contributions to surface science. Langmuir developed a surface
film balance among other things. Katharine Blodgett was also on the research
staff at General Electric and carried out much of the experimental work.
A monolayer on a liquid surface is a Langmuir film in contrast to built up monolayer assemblies on solid supports known as Langmuir-Blodgett films\(^4\). Six large international conferences involving LB films have taken place, the last was LB6, the 6th International Conference on Organized Molecular Films in 1993 in Trois-Rivières, Québec, Canada and the papers have been published\(^5\). An ACS journal exists which is called *Langmuir* and is dedicated to the study of surfaces and colloids.

An account of the steps to prepare a Langmuir-Blodgett film is given here with appropriate examples. The LB trough used in the experiments was a Fromherz trough\(^6\) shown in Figure 2-1. There are alternate trough designs available commercially which have been described in the literature\(^1\), some of which are rectangular. The Fromherz trough has a circular design and consists of polytetrafluoroethylene (PTFE). Polytetrafluoroethylene is a good material because it can be cleaned using concentrated acid and is resistant to organic solvents. Cleaning of the trough was normally accomplished by soaking the entire PTFE surface with NOCHROMIX\(^\circledR\) (Godax Laboratories) in concentrated sulfuric acid. The acid was suctioned off using an aspirator with a PTFE tip. The trough was rinsed five times with Milli-Q water with one additional distillation (pH \(\sim\)6.3). The cleanliness of the water subphase is extremely important for LB work. A laminar flow cabinet and enclosure were used with the trough. The trough was sitting on a terrazzo vibration-free table.

The Fromherz trough has two movable barriers which allow a film to be transported if necessary. For the experiments reported here, one barrier was moved. There are two calibrations which were performed. The area enclosed
Figure 2-1: Schematic of Fromherz circular Langmuir-Blodgett trough. Molecules spread from syringe and compressed. Surface pressure measured using a Wilhelmy plate. Film lift device not shown.
between the two barriers was easily calibrated with a potentiometer. The total area of the trough was 366 cm\(^2\). If the two barriers are moved to three and nine o'clock, the area should read 183 cm\(^2\). The second calibration was for the surface pressure measurement. Surface tension, \(\gamma\) is defined in equation 2.1 where \(G\) is the surface free energy and \(S\) is the surface area. The temperature, pressure and concentration are kept constant.

\[
\gamma = \left( \frac{\partial G}{\partial S} \right)_{T,P,\eta_i} \quad \text{equation 2.1}
\]

Surface pressure, \(\Pi\) is the difference between the surface tension of the subphase only (\(\gamma\)) and with monolayer present (\(\gamma_o\))\(^1\-\^3\).

\[
\Pi = \gamma - \gamma_o \quad \text{equation 2.2}
\]

There are two techniques used to measure surface pressure, the Langmuir balance and the Wilhelmy method\(^1\). The Langmuir balance has a float which separates an area of the subphase with monolayer present from a region of just water and for this reason is considered a differential method. The surface pressure exerted by the film moves the float until balanced by an equal force applied by a spring. A displacement transducer is used to measure the spring deflection. A Wilhelmy plate was used experimentally with the Fromherz trough. The material used for the plate was a clean piece of filter paper. There
are three forces involved when considering the Wilhelmy plate, gravity and surface tension act downward while buoyancy moves the plate upward. A force balance is given by equation 2.3. The length, width and thickness of the plate are given by l, w and t. The density of the plate material is $\rho_p$ and the density of the subphase which is usually water is $\rho_L$. The gravitational constant is $g$ and the contact angle, $\theta$ is the angle the liquid makes with the Wilhelmy plate. The height of the plate immersed in liquid is $h$.

$$F = \rho_p l w t + 2\gamma (l + w) \cos \theta - \rho_L t w h$$ equation 2.3

The plate was initially wetted by the water only and the pressure is adjusted to read zero with a potentiometer. The surface tension measured was the difference between the pure water and monolayer on the water. In this case a simple equation can be used and it is assumed $t << w$. The presence of a monolayer reduces the surface tension of pure water.

$$\Delta \gamma = \frac{\Delta F}{2w}$$ equation 2.4

The calibration of the surface pressure measured by the Wilhelmy plate was carried out by adding a weight of known mass to the wire support for the plate. The area of the filter paper piece was such that the weight exerted a known force of 100 mN/m.
There are many types of materials that can be deposited as LB films. Fatty acids are one of the most common classes used for LB work. Fatty acids contain a polar head group and a long alkyl chain. On the water surface, fatty acids will orient so that the hydrophilic polar group is in the water with the hydrophobic tail pointing away from the water. LB films however, are not restricted to these classical types of films. The LB material is dissolved in a solvent which is immiscible with the water subphase. The materials which are discussed in later chapters, phthalocyanines and perylenes, are considered pigments in most instances because of their lack of solubility. However, solvents can be selected to provide a small amount of solubility at low concentrations on the order of $10^{-4}$ M. High purity solvents must be used for LB work so that contamination is not introduced. A known volume of LB material in solvent is introduced to the surface of the LB trough using a calibrated syringe. The solvent readily evaporates leaving the LB molecules on the surface of the trough.

As the moving barrier is compressed, surface pressure readings are collected from the Wilhelmy plate or Langmuir balance. A surface pressure versus area isotherm is collected in this manner. There are four regions in the isotherm for a classical LB material. At very high areas, the molecules are spread apart and are in a gaseous state. As compression continues, the tails lift away from the water in the liquid-expanded region, followed by a region of constant slope in which the molecules are in a liquid-condensed state and finally a steeply sloping regime where the film is packed tightly and displays solid behavior. At high surface pressures collapse of the monolayer can occur. A typical surface pressure-area isotherm is shown in Figure 2-2.
Figure 2-2: Surface pressure-area isotherm for arachidic acid. Subphase pH=6.5 and 9.928 x 10^{-4} M CdCl₂. Gaseous phase at large area per molecule not shown. Four distinct phases apparent: liquid-expanded, condensed, solid and collapse.
Figure 2-2: SURFACE PRESSURE - AREA ISOTHERM

Arachidic Acid

Subphase: pH = 6.5, 9.928 x 10^{-4} M CdCl_{2}

Phase transitions:
- Collapse
- Solid
- Condensed
- Liquid-expanded

Area (nm²)/molecule

II (mN/m)
Equation 2.5 was used to determine the area/molecule in Figure 2-2.

\[
\frac{\text{Area}}{\text{molecule}} = \frac{\text{Surface area}}{\text{Avogadro's no. } X \text{ concentration } X \text{ volume}}
\]

 equation 2.5

A line can be extrapolated from the steeply sloping region of the solid part of the isotherm and a value of area/molecule read from the x-axis. The first part of an LB experiment is to collect the surface pressure-area isotherm. Next, deposition onto the substrate must be considered.

Many different substrates can be used for Langmuir-Blodgett films. Glass and quartz are hydrophilic. Atomically flat substrates such as polished silicon wafers and mica can be used as substrates for LB work and later atomic forces microscopy. Glass can be made hydrophobic by using silanizing agents. Langmuir-Blodgett deposition of monolayers onto metal island films is demonstrated in a later chapter for surface-enhanced Raman spectroscopy. Whatever substrate is chosen, it is important that it be clean.

There are a number of deposition techniques which can be used to build up Langmuir-Blodgett films and a few of these are illustrated in Figure 2-3. X-type deposition is carried out by dipping a hydrophobic substrate into the water subphase. The hydrophobic tail is adjacent to the substrate surface. The subsequent film layers have hydrophilic to hydrophobic contact which may not be particularly stable. Z-type deposition involves withdrawing a hydrophilic substrate through the monolayer. The hydrophilic head group is adjacent to
Figure 2-3:  

a) X-type LB deposition onto a hydrophobic substrate.  
b) Z-type LB deposition onto a hydrophilic substrate.  
c) Y-type LB deposition onto a hydrophilic substrate by withdrawal and dipping.
a) X-deposition onto hydrophobic substrate

b) Z-deposition onto hydrophilic substrate

c) Y-deposition onto hydrophilic substrate

d.

e.

etc.
the hydrophilic substrate. Once again, layers of the film have hydrophilic-hydrophobic contact. Y-type deposition involves successive withdrawal and dipping of the substrate. Adjacent layers have hydrophilic-hydrophilic contact and hydrophobic-hydrophobic contact.

A substrate dipper is used for LB deposition with variable speeds and a smooth action. The transfer of a monolayer to a substrate is carried out in a constant pressure mode. A feedback to the control of the moving barrier is necessary so a compensation takes place as material is removed during transfer to the substrate. One measure of the quality of LB deposition is the transfer ratio, $t_R$ given in equation 2.6.

$$t_R = \frac{\text{area removed from trough}}{\text{area transferred to substrate}}$$  \hspace{1cm} \text{equation 2.6}

A consistent transfer ratio is important. A transfer ratio of one suggests good transfer and that the orientation of the monolayer on the substrate is similar to that on the subphase.
2.2 VACUUM EVAPORATED FILMS

A description of high vacuum systems is available in the literature\textsuperscript{10,11}. A Vacuum Generators DPUHV12 model was actually used in the experimental section. Two types of materials were vacuum evaporated, metals and organic pigments. The metals were of high purity and the boat was selected so that an alloy was not formed on heating. The organic pigments were previously purified by vacuum sublimation and were placed in a quartz boat. Pigments were selected which did not decompose before the temperature for evaporation was reached. Current was passed through either the metal boat or metal surrounding the quartz holder. The high vacuum was achieved by the presence of a roughing pump and a diffusion pump. The substrate was Corning 7059 glass and was clipped to a substrate holder which allowed heating or cooling. There are further details in the experimental section.

The vacuum chamber was evacuated to a predetermined pressure. At this point the shutter was closed and current gradually applied to the boat. A quartz crystal thickness monitor was located close to the substrate position. As material started to evaporate, readings were observed on the thickness monitor. Once the evaporation rate had stabilized to the desired rate, the shutter was opened. The current was shut off and shutter closed once the desired thickness was obtained. Figure 2-4 is a schematic representation of the vacuum evaporation system.
Figure 2-4: Schematic of a typical vacuum evaporation experiment. An organic material or metal is heated in a vacuum chamber and deposited on substrate. The mass thickness is followed with a quartz crystal thickness monitor.
2.3 RAMAN SPECTROSCOPY THEORY

2.3.1. Normal Raman

Raman spectroscopy and infrared spectroscopy provide vibrational information which is complementary in nature. The observed frequencies depend on selection rules unique to each technique. The molecular symmetry and crystal form determines which vibrations are active. The relative intensities of bands may be quite different in the Raman spectrum versus the IR spectrum because the mechanisms are very different. For a mode to be Raman active, there must be a change in the molecular polarizability during that vibration. A change in the molecular dipole moment during vibration is a requisite for IR activity.

There are instrumental differences between an FTIR spectrometer and a Raman spectrometer. The source in an IR instrument is very broad in nature spanning a range of 200-4000 cm\(^{-1}\). In comparison, lasers are intense monochromatic sources for Raman work with lines available in the UV, visible and near-IR regions. Molecules undergo absorption of IR radiation during the measurement of an infrared spectrum. The molecules are excited from the ground vibrational state to a higher vibrational state. Modern infrared spectrometers have interferometers which allow all frequencies to be detected simultaneously. The Michelson interferometer has a fixed and a moving mirror. During one scan, the moving mirror travels a distance defined by the desired resolution. The light from two beam paths interact either constructively or destructively. The measured IR intensity is a function of the distance (or velocity x time). A mathematical transformation is carried out to convert from the time domain to a frequency domain, thus the term Fourier transform infrared (FTIR). An infrared spectrum is a plot of intensity
(absorbance or transmittance) versus frequency in wavenumbers (units cm$^{-1}$). An FTIR sample is ratioed against a background to take instrumental factors into account. Two examples of FTIR detectors are the MCT and DTGS detectors.

In a Raman experiment, an intense monochromatic source such as a laser strikes a sample and most of the light is scattered elastically producing Rayleigh scattering. However, a very small proportion of photons will be scattered inelastically giving Raman scattering. There are two possibilities for Raman scattering, Stoke’s and anti-Stoke’s, both causing a shift in the frequency of the light. Figure 2-5 shows the vibrational energy levels. For Stoke’s scattering, the frequency of scattered light is less than the frequency of the incident light. Molecules are excited from a ground vibrational state (v=0) to a higher energy state. The molecules leave this higher energy state and return to an excited vibrational state (eg. v=1). In the anti-Stoke’s situation, molecules start in an excited vibrational state (eg. v=1), are excited to a higher energy state and then return to the ground vibrational state (v=0). The frequency of anti-Stoke’s scattering is greater than the frequency of the incident light. Stoke’s scattering is usually much more intense than anti-Stoke’s because few molecules start out in an excited vibrational energy level.

The laser source used for Raman spectroscopy has an electric field \( E \) and frequency \( \nu_0 \) given by equation 2.7.

\[
E = E_0 \cos(2\pi \nu_0 t)
\]

\text{equation 2.7}
Figure 2-5: Schematic showing vibrational energy levels and photons for elastic Rayleigh scattering, inelastic Stoke’s and anti-Stoke’s normal Raman scattering.
A dipole moment, $\mu$ is induced in the molecule by the electric field of the laser. The molecular polarizability $\alpha$ describes the ease of electron deformation in a molecule.

$$
\mu = \alpha E
$$

The molecular polarizability is a tensor which has nine components, another form of equation 2.8 follows:

$$
\begin{pmatrix}
\mu_x \\
\mu_y \\
\mu_z
\end{pmatrix}
= \begin{pmatrix}
\alpha_{xx} & \alpha_{xy} & \alpha_{xz} \\
\alpha_{yx} & \alpha_{yy} & \alpha_{yz} \\
\alpha_{zx} & \alpha_{zy} & \alpha_{zz}
\end{pmatrix}
\begin{pmatrix}
E_x \\
E_y \\
E_z
\end{pmatrix}
$$

For normal Raman scattering, there are six polarizability components modulated by a molecular vibration because the polarizability tensor is symmetric: $\alpha_{xy} = \alpha_{yx}$, $\alpha_{xz} = \alpha_{zx}$ and $\alpha_{yz} = \alpha_{zy}$.

The molecular polarizability in equation 2.10 consists of the polarizability, at the equilibrium nuclear coordinates ($\alpha_0$) and polarizability which is modulated by the vibrational frequency $\nu_1$:

$$
\alpha = \alpha_0 + \alpha_0 \cos(2\pi \nu_1 t)
$$
The change in polarizability with respect to the normal coordinates, $Q$ can be expressed in the form of equation 2.11:

\[ \alpha = \alpha_0 + \Sigma (\partial \alpha / \partial Q_i)_{Q_i=0} + \ldots \]  

\text{equation 2.11}

It is the second term in equation 2.11, also called the polarizability derivative $\alpha'$ which produces Raman scattering. There must be a change in the molecular polarizability in order for a vibration to be Raman active. From this point on the symbol $\alpha$ will represent the molecular polarizability derivative.

Equation 2.12 is the expression for the intensity of radiation emitted by an oscillating dipole. The intensity is affected by the frequency to the fourth power. The electric field of the laser and the molecular polarizability are included through $\mu = \alpha E$.

\[ I = \left( \frac{16\pi^4 \nu^4}{3c^3} \right) \mu^2 \]  

\text{equation 2.12}

In the case of conventional normal Raman spectroscopy, the scattered light is separated into different wavelengths with gratings. Holographic gratings can be used in double or triple monochromators. One type of detector for Raman scattering is the photomultiplier tube. Spectral sensitivity is an important consideration when selecting a detector for scattered light which can be in the UV, visible or near-IR region. In the conventional Raman
system, the gratings are moved so that different wavelengths are passed over the photomultiplier tube. It can be time consuming to collect an entire spectrum. Alternatively, all frequencies could strike a detector array such as a charge-coupled device (CCD) or photodiode array simultaneously. The time to collect a spectrum in this case is short. There are now FT-Raman spectrometers which have a laser source, interferometer and an appropriate detector. Lasers with near-IR lines are used with infrared detectors. A review of dispersive Raman and FT-Raman systems has been recently published\textsuperscript{13}. Whatever system is used, there must be a way of removing the intense Rayleigh scattering from the weak Raman scattering.

2.3.2 Polarizations of Light

A brief discussion can be made here concerning the notation for different polarizations of laser light that are available for a Raman experiment. The notation being used is that of Porto\textsuperscript{14}. Figure 2-6 shows the four polarizations SS, SP, PS and PP that can be determined in a Raman experiment with a 90 degree geometry between the direction of propagation of the laser and the collection of scattered light. There are two possible polarizations of the laser, perpendicular to the scattering plane, S and parallel, P. The analyzer after the sample also has two positions S and P.

Depolarization ratios\textsuperscript{15} are commonly measured for liquids and have values ranging from 0 to 3/4. In liquid samples, important information is gained about the molecular symmetry of a vibration from the depolarization ratio. A totally symmetric vibration produces a Raman band with a depolarization ratio between 0 and 3/4 while antisymmetric vibrations are depolarized.
Figure 2-6: The four polarizations of light according to Porto\textsuperscript{14} for a ninety degree Raman scattering experiment. The propagation of the laser is along the z-axis. Collection of scattered light is along the x-axis. The four polarizations are SS, SP, PS and PP. The terms inside the brackets describe the electric field vector of the laser and the electric field vector of the light analyzed after the sample.
Figure 2-6

a) \( Z(YY)X = SS \)
b) \( Z(YZ)X = SP \)
c) \( Z(XY)X = PS \)
d) \( Z(XZ)X = PP \)
The experimental work presented in later chapters is for Langmuir-Blodgett films and vacuum evaporated films which exist in the solid state. Polarization ratios SS, SP, PS and PP can provide information about molecular orientation with respect to the substrate if a number of transformations are performed. There are three sets of coordinate systems that must be considered. Firstly, there is the molecular coordinate system. A transformation is carried out to go from molecular coordinates to the second system, the substrate coordinates. The orientation of the molecule with respect to the substrate is important. Thirdly, the laboratory coordinates are defined by the position of the substrate in relation to the incident laser beam and collection of scattered light.

### 2.3.3 Resonance Raman

Resonance Raman (RR) scattering occurs when the laser excitation wavelength coincides with a molecular electronic transition. Resonance Raman scattering is more intense than normal Raman scattering due to a change in the molecular polarizability derivative tensor. \((\alpha_{\rho \sigma})_{GF}\) is the poth component of the Raman scattering tensor and \(\rho, \sigma = x, y\) or \(z\). The ground state is represented by \(G\) and the final state is \(F\). The elements of the polarizability tensor are given by \(16-18\):

\[
(\alpha_{\rho \sigma})_{GF} = \frac{1}{\hbar} \sum_I \left( \frac{<F|r_p|I><I|r_\sigma|G>}{\nu_{GI} - \nu_L - i\Gamma_I} + \frac{<I|r_p|G><F|r_\sigma|I>}{\nu_{IF} - \nu_L - i\Gamma_I} \right)
\]

**equation 2.13**

\(I = \text{vibronic excited states}\)
\( r = \) electron position operator

\( \Gamma_T = \) damping term

From equation 2.13, it can be seen that the first term in the summation becomes very large if \( v_L \) approaches \( v_G \) for an allowed electronic transition and that the damping term related to the bandwidth of the electronic transition prevents the expression from reaching infinity. The polarizability tensor for resonance Raman may become asymmetric, i.e. \( \alpha_{xy} \neq \alpha_{yx} \), etc. A new term is introduced into intensity and depolarization expressions.

\((c_{po})^G\) has been divided into two terms described by Albrecht\(^{16-17}\). The first term is Albrecht’s A term and involves Franck-Condon factors. Only totally symmetric terms are enhanced in the RR spectrum. Albrecht’s B term for RR follows a Herzberg-Teller mechanism where electronic states are mixed and non-totally symmetric modes can be observed.

Resonance Raman scattering can be distinguished from normal Raman scattering in a number of ways. The intensity of normal Raman scattering will follow a \( v^4 \) dependence on excitation frequency whereas RR scattering varies greatly with excitation frequency. The RR time dependence is slower than off-resonance or normal Raman scattering. The relative intensity of Stoke’s to anti-Stoke’s scattering for normal Raman is determined by the Boltzaman distribution. However, this is not true for RR and the intensity of anti-Stoke’s scattering can actually exceed Stoke’s scattering.

**Surface-Enhanced Raman Scattering** is covered in Chapter 3.
2.3.4 **FT-Raman Spectroscopy**

A series of reviews and applications have appeared concerning FT-Raman spectroscopy\(^{19-60}\). There were a number of reasons why FT-Raman systems were developed. Intrinsic fluorescence or fluorescence due to impurities in samples has been a severe problem for Raman spectroscopy using visible laser excitation. Fluorescence is a much more efficient process than Raman scattering and can completely mask Raman signals. The 1064 nm line or in a special case\(^{60}\) the 1339 nm line of a Nd/YAG laser can eliminate or greatly reduce fluorescence. Some materials have an electronic absorption in the visible region and will photodecompose when exposed to visible light. Biomolecules that fall into this category that have been examined by FT-Raman spectroscopy include: cobalamins, bacteriorhodopsin, chlorophylls, cytochrome c and porphyrins\(^{58}\). It may be desirable to obtain normal Raman scattering rather than resonance Raman scattering.

A multiplexing system based on a Michelson interferometer has an internal frequency calibration using a He-Ne laser. The precision and accuracy of frequencies is extremely high. This makes it possible to carry out spectral subtractions in an FT-Raman system. An example of where this is important is for biological assemblies where membrane reorganizations are reflected in small frequency shifts\(^{25}\). Dispersive instruments which have holographic gratings have greater errors in the frequencies and are less predictable\(^{19}\).

Interferometers have a throughput (Jacquinot’s) advantage compared to dispersive spectrometers\(^{20}\). If the Raman experiment is detector noise limited rather than shot noise limited, a multiplexing (Felgett’s) advantage is achieved using the interferometer. It should be mentioned that there is a disadvantage
in using near-IR excitation rather than visible excitation for Raman spectroscopy. The intensity of Raman scattering is proportional to $v^4$ and drops by a factor of sixteen going from 514.5 nm to 1064 nm$^{20}$. The throughput and multiplexing advantages can compensate for the lower Raman scattering cross-section in the near-IR region.

Another advantage of FT-Raman instrumentation is it is relatively cheap compared to a conventional dispersive Raman system operating in the visible with a photomultiplier tube (PMT) and can be operated with minimal training. There are systems which consist of combined FT-Raman and FTIR modules. The information from Raman and IR experiments are complementary since the selection rules are different and both techniques should be applied to a problem. FT-Raman is non-destructive and sample preparation is very simple. Cuvettes, mp capillary tubes, and nmr tubes can be used to hold the sample. The optimal sample cells have been reported and include a spherical cell with a reflecting layer$^{28,38,51}$. Water is highly absorbing and interferes with FTIR spectra. The overtone region of water has some absorption in the near-IR region used for FT-Raman spectroscopy and can reduce the scattering intensity of C-H stretching modes$^{25}$.

Fiber optic probes that were developed for the telecommunications industry have their best performance in the region 1.3 μm to 1.7 μm which is ideal for FT-Raman spectroscopy$^{37}$. There are two ways to carry out remote Raman process monitoring$^{22}$. A small instrument could be used in situ and the spectra shipped to a central location for analysis or a fiber optic bundle could carry the laser beam to the process site and take the scattered light away. The fibers should be of a type which has the lowest water content possible$^{21}$. 
Results have been reported for indene and a polymerization reaction of styrene with initiator at 70°C followed in situ\textsuperscript{37}. A suggested application of FT-Raman with fiber optics is monitoring cure reactions and rates during vulcanization at different depths within a compound or to follow in-service aging of a tire\textsuperscript{35}. Two arrangements of optical fibers have been described for FT-Raman measurements of SERS active colloids and electrodes\textsuperscript{53}. A near-IR system with fiber optics is being developed to monitor contaminants of groundwater in situ that are in wells 100 to 600 feet below the surface of the earth\textsuperscript{42}.

The 1064 nm line of a neodymium-doped yttrium-aluminum garnet (Nd/YAG) solid state laser is the most common excitation source for FT-Raman spectroscopy. Future developments could be based on a neodymium-doped yttrium lithium fluoride (Nd/YLF) diode pumped laser of wavelength 1047 nm which is small and does not require cooling\textsuperscript{21}. The present drawbacks are higher power is needed and lower cost. There could be a move to slightly shorter wavelengths in the range 790-900 nm using a narrow line-width laser with photodiode detection\textsuperscript{22}. A compromise would be made which would cause sample fluorescence to increase but background from thermal emission of hot samples would be moved and less noisy detectors are available\textsuperscript{22}. A near-IR Raman system has been described incorporating a tunable Ti:sapphire laser which can operate at wavelengths between 790-850 nm with charge-coupled device (CCD) detection\textsuperscript{52}.

FT-Raman instruments are available with various scattering geometries such as 180 degree backscattering. It is extremely important that the relatively intense Rayleigh line be eliminated from the scattered radiation entering the
interferometer and ultimately striking the detector. The first problem is the
detector would require an extremely large dynamic range to accommodate
Raman and Rayleigh scattering. It is conceivable the Rayleigh component
would saturate the detector. Secondly, the interferogram would have a large
modulated Rayleigh line on top of a weak Raman spectrum and shot noise
would be distributed over the spectrum\textsuperscript{20,22}. Many approaches including
filtering have been described in the literature to remove the intense Rayleigh
line\textsuperscript{21-23}. Advances in Rayleigh rejection are needed to access longitudinal
acoustic modes (LAMs) close to the Rayleigh line. The scattered light enters a
Michelson interferometer which contains appropriate beamsplitters as well as
moving and stationary mirrors.

The detectors suitable for near-IR radiation include InGaAs, Ge and
PtSi\textsuperscript{19,21,48,59}. Detector response curves are published for Ge and InGaAs\textsuperscript{21}. Both of these materials exhibit lower noise equivalent powers (NEPs) at
cryogenic temperatures, however spectral response is reduced. Thus, the Ge
detector operated at 77K falls off at frequency shifts greater than 2500 cm\textsuperscript{-1}
with a 1064 nm laser line\textsuperscript{19}. This would sacrifice the C-H and N-H stretching
region of the FT-Raman spectrum. The InAs detector is better in terms of
responsivity than other materials but the noise is unacceptable\textsuperscript{21}. Silicon
detectors have a low response above 1 micron and would be useful for anti-
Stokes FT-Raman only\textsuperscript{59}. Platinum silicide extends into the near-IR but one
problem is thermal background and low sensitivity. At present the best choice
for the desired near-IR spectral response and lowest noise is an InGaAs detector
operating somewhere between 77K and room temperature\textsuperscript{48}. 
There have been reports of FT-SERS of pyridine on metal electrodes such as copper, silver and gold using 1064 nm laser excitation\textsuperscript{39-40,42-43}. Electrodes are roughened either ex situ with just the electrolyte present or in situ with the species to be studied also present. This is accomplished by oxidation-reduction cycles (ORCs). The metal electrode is the working electrode, a platinum counter electrode is present and a reference electrode such as the saturated calomel electrode (SCE) is part of the cell. The FT-SERS spectrum depends on the electrode material and the potential applied to the working electrode\textsuperscript{39,43}. Often broad backgrounds are observed due to the metal. The calculated enhancement factors for pyridine are in the range $10^5$-$10^6$ on copper and gold electrodes\textsuperscript{42}. In order for FT-SERS to be useful for chemical sensor applications, the species of interest must be able to be desorbed from the electrode at some potential to allow electrode reconditioning. It has been shown that reconditioning of a copper electrode can take place after pyridine has been adsorbed, but a gold electrode was irreversibly affected by pyridine adsorption\textsuperscript{42}. There is a complex dependence between potential and pH for pyridine adsorbed on a silver electrode\textsuperscript{43}. The positions and relative intensities of bands attributed to ring breathing and deformations indicate there were at least two or three types of pyridine species with at least two orientations. The reproducible splitting of bands was indicative of the existence of multiple environments.

FT-SERS of 3-picoline and 3-chloropyridine has been observed on copper electrodes\textsuperscript{41}. The calculated enhancement factors were $9 \times 10^5$ and $5 \times 10^5$ respectively. This work was part of an initial study of environmental contaminants using FT-SERS. The adsorption of ferro- and ferricyanide ions onto a gold electrode in lithium chloride and cesium chloride has been
discussed. The redox species were adsorbed in ion pairs and were present in a variety of environments which could be due to different metal sites. The corrosion inhibitors benzotriazole (BTA) and tolyltriazole (TTA) give SERS in the near-IR on copper electrodes. These inhibitors are circulated in cooling water to reduce corrosion of copper containing parts. The FT-SERS results indicate that both BTA and TTA are adsorbed through the azole ring and it is this film which protects the copper. Commercial antifreezes which contain water, ethylene glycol and organic inhibitors selectively adsorbed the organic inhibitors on the copper electrode.

2.4 ATOMIC FORCE MICROSCOPY

There are many references available concerning atomic force microscopy (AFM) and scanning tunneling microscopy (STM). This section will give a brief description of AFM. Papers are cited in Chapter 6 for AFM work on phthalocyanines, perylenes, fatty acids and self-assembled monolayers and particular attention is paid to AFM of Langmuir-Blodgett monolayers.

Atomic force microscopy (AFM) has a number of advantages for imaging Langmuir-Blodgett (LB) films and thin solid films compared to other techniques such as electron microscopy and scanning tunneling microscopy (STM). The contact mode of the AFM gives higher resolution in height (z-direction) than electron microscopy. The resolution of the AFM in the x-y plane is also excellent and is affected by the tip geometry and tip composition as well as sample properties. The AFM can image regions on the order of microns to nanometers. LB films for AFM are transferred to atomically flat substrates which include mica, highly oriented pyrolytic graphite (HOPG) and silicon wafers. The AFM can be operated in an aqueous environment in which a non-
conducting sample and tip are submerged and this is of interest for biological studies.

For scanning electron microscopy (SEM), the sample must be conductive, a thin conductive layer applied or replicas prepared. In some instances the sample is not directly imaged. To carry out transmission electron microscopy (TEM), the sample is transferred to a substrate such as a carbon-coated microscope grid. There is insufficient contrast to image a single LB monolayer unless some sort of surface decoration technique is used to increase contrast. Materials can be damaged when subjected to high energy electrons.

Both AFM and STM give local information. Many characterization techniques such as IR, Raman, X-ray or neutron scattering give spatially averaged results over a large number of atoms. AFM and STM are good choices for observing local defects such as pinholes, cracks, grain boundaries or heterogeneous structures such as mixed LB films. In the best case for STM, atomic resolution can be achieved and features observed which are 0.2 nm in diameter. Although STM requires that the sample under investigation be conductive, the AFM can image conductive and non-conductive materials. In STM, electrons tunnel between the tip and the sample. The images reveal aspects of the sample topography and electronic structure. What is observed is the density of electronic states at the Fermi level. Results from AFM can be considered complementary to STM because AFM images are due to energy levels up to the Fermi level.

The contact mode of the AFM senses repulsive forces which are due to an overlap of the electrons in the tip with electrons of the nearby atoms on the
sample surface. One must be careful not to disrupt van der Waals forces between molecules or hydrogen-bonding when imaging LB films. There should be some sort of immobilization of the LB film on the substrate either through covalent bonding or physisorption. Molecular stacking causes strong intermolecular interactions which can help stabilize the position of the LB film. Experimentally, a cantilever with a low spring constant can be used or samples imaged under water. The force applied to an organic surface in a STM experiment is approximately 10^{-7} newtons, orders of magnitude greater than most AFM experiments\textsuperscript{162}. AFM is the preferred technique for imaging thin organic films because sample damage can be minimized.

An atomic force microscope consists of a tip and cantilever to sense the force. The cantilever deflection is detected either electronically or optically. Some sort of mechanical scanning system is required and usually has a piezoelectric component. The tip is held stationary while the sample is moved in the x and y directions. The data which is acquired is a map of forces over points on the surface of the sample. A feedback system is incorporated to control the vertical distance (z) between the tip and sample allowing two modes of operation, constant force and constant height. It is important to minimize any mechanical vibrations of the AFM. A display system is used to convert the data into an image and data processing is performed.

Initial AFM experiments were carried out using metal foil as the cantilever\textsuperscript{161}. Melmed\textsuperscript{128} has reviewed the literature on making sharp tips for field ion microscopy, field electron emission microscopy and STM with an emphasis on electrolytic methods. In addition AFM force sensors have been constructed from carbon and quartz fibers\textsuperscript{126}. The tips are formed by chance
on broken ends. AFM and STM can be performed simultaneously if graphite fibers are used for the probe. Tips for cantilevers have been prepared by attaching diamond fragments by hand to the end of levers\textsuperscript{144,147}. Microfabrication techniques are presently used to provide silicon, silicon dioxide and silicon nitride cantilevers with and without integrated tips\textsuperscript{134,138}.

It is desirable to have a cantilever with a low spring constant in the range 0.1 to 10 N/m with a high resonance frequency\textsuperscript{107}. Rugar and Hansma\textsuperscript{118} calculate a value of 10 N/m for an interatomic spring constant (k) based on vibrational frequencies (ω) on the order of 10\textsuperscript{13} Hz or higher and mass of atoms (m) on the order of 10\textsuperscript{-25} kg using the expression:

\[
k = \omega^2m
\]  

\text{equation 2.14}

The cantilever spring constant and force must be kept much lower than 10 N/m so that van der Waals interactions, hydrogen-bonding and bonds are not disrupted or broken.

There are numerous ways to detect cantilever deflection. Early AFM experiments described by Binnig and Quate\textsuperscript{161} used electron tunneling to measure the deflection of the cantilever. Meyer and Amer\textsuperscript{68} described an optical method to detect cantilever deflection in an AFM by bouncing a laser beam off the back of the cantilever. Deflections were measured with a position sensitive photodiode and the smallest deflection that could be measured was only limited by thermal vibrations. Sarid and Elings\textsuperscript{105} have
reviewed scanning force microscopy and discuss the seven methods to detect cantilever deflection. Additional methods include an electronic method which senses small changes in the capacitance and determines motions of a cylindrical cantilever\textsuperscript{141,142}. Other optical methods are homodyne\textsuperscript{143}, heterodyne\textsuperscript{144}, laser diode feedback\textsuperscript{145} and polarization\textsuperscript{146}. Optical methods are preferred over tunneling because the laser spot averages over a larger area on the cantilever backside and is less sensitive to roughness or contamination\textsuperscript{107}. The detection method should not exert a measurable force on the cantilever.

A mechanical scanning system is an integral part of the AFM which allows data to be collected over the sample surface. Usually the sample is scanned in the x and y directions while the tip is held stationary. A coarse mechanical system brings the tip and the sample into the operating range of a finer system which has piezoelectric components. Piezoelectric materials change dimensions in a known fashion in response to applied voltages.

The imaging resolution of an AFM operating in contact with a LB film has been described in terms of the applied force, tip radius, adhesive force, sample and tip properties\textsuperscript{66}. Contact diameters were calculated using the Young's modulus of the LB film from nanoindentation experiments, the applied force and tip radius. At small forces the behavior was elastic but as the force was increased, a transition to inelastic deformation appeared. At this onset point there is irreversible damage to the LB film and line scans are no longer reproducible. Adhesive forces between the tip and the LB film have the effect of increasing the contact diameter and decreasing the resolution. There were suggestions made to improve imaging resolution\textsuperscript{66}. If the tip is sharpened, the adhesive force is decreased but there is the risk of sample damage by inelastic
deformation. Secondly, a stiffer tip with higher Young's modulus would be beneficial. Adhesion can be minimized by immersing the tip and sample in water. Another alternative would be to image in a non-contact mode.

2.5 SUMMARY

An introduction to Langmuir-Blodgett films was contained in this chapter. A history of the development of LB films was given. The steps to collect a surface pressure-area isotherm and to deposit a LB film on a substrate were described. One way to prepare metal films and thin organic films is by vacuum evaporation which was considered briefly. The next section contained theory and background on Raman spectroscopy including normal Raman, resonance Raman and FT-Raman spectroscopy. Surface-enhanced Raman scattering is covered in Chapter 3. A comparison was made between infrared spectroscopy and Raman spectroscopy. The four polarizations of light SS, SP, PP, and PS have been defined for a 90 degree Raman experiment. A general review of atomic force microscopy was made with more specific references to phthalocyanines, perylenes, fatty acids and self-assembled monolayers reserved for Chapter 6. Atomic force microscopy was compared to scanning tunneling microscopy and electron microscopy. The components of atomic forces microscope have been described.
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CHAPTER 3: SURFACE-ENHANCED RAMAN SCATTERING (SERS) AND MODEL CALCULATIONS OF ENHANCEMENT FACTORS

3.1. INTRODUCTION

During normal Raman scattering only a fraction of the incident light is scattered inelastically. As a further complication, thin solid films which have monolayer or submonolayer thicknesses produce weak Raman scattering due to the small number of molecules in the incident beam. Surface-enhanced Raman scattering (SERS) is one way to improve the sensitivity of the Raman technique. Surface-enhanced Raman scattering occurs when molecules are at or near specially prepared metal substrates. The average of the Raman scattering intensity is typically increased on the order of $10^4$ by SERS. The enhancement factor is the ratio of the scattering intensity in the presence of the SERS active substrate to the intensity that would be observed without the substrate present:

$$\text{EF} = \frac{I_{\text{SERS}}}{I_R}$$  \hspace{1cm} (equation 3.1)

This chapter discusses the substrates that have been used for SERS. A historical perspective is provided which connects such diverse areas as metal colloids, aerosols, light scattering and absorption, color and SERS. SERS theories which fall into two groups are presented. There are selection rules that influence the relative intensities of SERS bands. Model calculations are
carried out based on electromagnetic theory to guide the experimental selection of metal substrates for SERS in the near-IR.

There have been a number of SERS review articles and books in the literature\textsuperscript{1-8}. The first observation of SERS was reported by Fleishmann, Hendra and McQuillan\textsuperscript{9} for pyridine on an electrochemically roughened silver electrode. Many different metals give rise to SERS such as: Ag, Au, Cu, In, Li, Na, K, Ni, Pd, Rh, Ti, Cd, Co, Hg and Al\textsuperscript{3,10-11}. The SERS-active substrates are prepared in various ways and surface roughness is necessary. Metal electrodes are roughened during oxidation-reduction cycles in an electrolyte. Metal island films are produced by vacuum evaporation at elevated substrate temperatures. Cold deposited metal films are prepared with the substrate below 120K\textsuperscript{1}. Repetitive rough structures such as gratings are produced lithographically. Metal colloids which consist of metal spheres or aggregated structures are SERS-active.

It is possible to have surface-enhanced resonance Raman scattering (SERRS) when the laser excitation is in resonance with both an electronic absorption of the adsorbate and the plasmon frequency of the SERS-active substrate. Many dyes and pigments are colored and absorb light in the visible region. Surface-enhanced resonance Raman scattering was used for trace analysis of the fluorescent dye rhodamine 6G on a silver sol\textsuperscript{12}. Raman bands were observed even at a dye concentration of 4 x 10\textsuperscript{-12} M. A thin film of the pigment zinc phthalocyanine (ZnPc) on gold and silver island films produced SERRS with an enhancement on the order of 10\textsuperscript{3} which is above that estimated for SERS alone using the 647.1 nm Kr+ laser line\textsuperscript{13,14}. In Chapter 5, additional examples will be presented of SERRS of phthalocyanines and perylenes.
Connections have been made between a number of seemingly disparate phenomena by Kerker\textsuperscript{15}. Michael Faraday carried out many experiments involving colloidal metal sols. Faraday made particular note of the colors of metal sols and how finely divided metal particles aggregated and changed color when salt was added. John Tyndall was Faraday's assistant and successor. Tyndall was interested in aerosols and made a connection between the blue color of the small aerosol particles, the color of the sky and the fact that light is polarized when observed perpendicular to the incident light from the Sun. The next historical figure was Lord Rayleigh who provided the theory behind Tyndall's observations. Rayleigh derived expressions for scattering and absorption of light and realized the blue color of the sky was due to molecules in the air. The type of factors entering Rayleigh's scattering and absorption expressions included the particle diameter, the wavelength of light, and the dielectric constant. Lord Rayleigh and later Sir C. V. Raman studied sulfur sols. Raman is well known for his discovery of the effect in which a small fraction of scattered light has been shifted in frequency. Surface-enhanced Raman scattering was discovered much later in 1974\textsuperscript{9}. Rough metal surfaces were initially used for SERS and then metal colloids. The colors of silver and gold colloids present in stained glass, Faraday sols and SERS had a common origin according to Alan Creighton\textsuperscript{15}. Kerker offers the explanation that the color of sols and SERS are due to resonances\textsuperscript{15}. The same Irving Langmuir, who worked on Langmuir-Blodgett films discovered that under the right conditions, an electromagnetic wave could cause electrons in metals to vibrate cooperatively leading to strong electrical fields. More will be mentioned about surface plasmons later in the next section. The electron oscillations lead to strong scattering and absorption. Color and SERS are effects which depend on particle size, morphology and dielectric function of the metal.
3.2. ELECTROMAGNETIC THEORY FOR SERS

Theories to explain SERS fall into two categories, electromagnetic (EM) and non-electromagnetic. The electromagnetic theory will be considered first. The electrons in metals such as silver, copper and gold behave like a gas. Metals have collective oscillations of the electrons which can be excited by an incident electric field and localized on the surface of a metal. The collective oscillations are surface plasmons. In order for surface plasmons to radiate, frequency and parallel momentum must be conserved and this condition is fulfilled by metals which have surface roughness features on the order of 10-100 nm. Electric fields at or near the surface of the metal are greatly enhanced and localized. A molecule at or near the metal surface experiences an effective electric field due to the incident electric field from the laser and the electric field arising from the metal surface. Raman shifted radiation is produced which can also be in resonance with the surface plasmon. If the excitation wavelength and Raman scattering wavelength are not too different a double resonance is possible. There are some experimental factors that can be varied in order to obtain the greatest SERS enhancement factor. It is possible to tune into the plasmon resonance by using the appropriate laser wavelength. The dielectric constant can be changed by selecting a different metal. SERS-active substrates can be prepared with different morphologies.

Simple models for surface roughness can be used to calculate SERS enhancements factors. The simplest case is that of an isolated metal sphere. One assumption in the calculations is that the particle size is significantly smaller than the wavelength of the incident light. Campion reports the following expression for the SERS enhancement factor (EF) for a sphere:
\[ EF = \left( \frac{\varepsilon_1(\omega_L) - \varepsilon_2}{\varepsilon_1(\omega_L) - 2\varepsilon_2} \right)^2 \left( \frac{\varepsilon_1(\omega_S) - \varepsilon_2}{\varepsilon_1(\omega_S) - 2\varepsilon_2} \right)^2 \left( \frac{r}{r+d} \right) \]

(equation 3.2)

\( \varepsilon_1 \) is the dielectric function of the sphere and \( \varepsilon_2 \) is the dielectric function of the medium.

\( \omega_L \) and \( \omega_S \) are the incident and Raman scattered frequencies respectively.

\( r \) is the radius of the sphere and \( d \) is the distance of the molecule away from the sphere.

The increase in Raman scattering is roughly proportional to the fourth power of the surface-enhanced electromagnetic field. Some general features can be extracted from expression 3.2. The resonance condition is \( \text{Re} \varepsilon_1 = -2\varepsilon_2 \) at the incident or scattering frequencies. The enhancement is proportional to \( (\text{Im} \varepsilon_1)^{-4} \). Metals such as silver, gold and copper are good enhancers in the visible region while transition metals should work well in the ultraviolet region. The distance dependence term is based on the dipole decay law. Electromagnetic theory can be checked quantitatively by measuring the distance-dependence of the SERS enhancement and this will be discussed later. The enhancement as a function of distance falls off rapidly for small particles or surface roughness features with a high radius of curvature. This equation provides an explanation for the fact that SERS intensities fall off with higher frequency shifts. In practice, incident and scattering fields are not simultaneously in resonance with the surface plasmons. If particles become larger and approach the wavelength of the exciting light, the resonances are red shifted, broadened and the enhancement factor decreases.
Gersten and Nitzan\textsuperscript{16} considered enhancement factors arising from molecules at or near metal spheroids of different aspect ratios. More will be mentioned about this model in the section in this chapter on model calculations. Prior to carrying out SERS experiments in Chapter 5, enhancement factor calculations were completed according to Gersten and Nitzan\textsuperscript{13} for various metals in the visible and near-IR region. The calculations guided the selection of the most appropriate metal(s) for island films and SERS in the near-IR. Whatever simple model is chosen, the experimentally obtained SERS-active substrates are usually more complex. For instance, aggregation takes place with metal colloids and shifts the plasmon resonance to the red. Most rough surfaces will have a distribution of particle sizes. A quantitative comparison between EM theory and experimental results is most appropriate for SERS-active substrates which have well-defined morphologies. An example would be gratings or lithographically produced structures. Electron microscopy and atomic force microscopy are valuable tools to check the substrate morphology.

Other surface-enhanced phenomena in addition to SERS are predicted by electromagnetic theory for rough metal surfaces. The photochemical decomposition of azo sulfonate compounds adsorbed on silver films has been followed\textsuperscript{17}. Excimer laser pulses at 308 nm irradiated the films and surface-enhanced Raman scattering using argon ion laser excitation at 488.0 nm and 514.5 nm was used to follow the reaction products. The monitoring of photolysis in solution was hampered by the production of small amounts of fluorescent products. Fortunately the metal island film quenched the fluorescence by radiationless transfer to the metal while increasing the Raman scattering\textsuperscript{17}. The distance dependence of fluorescence has been reported of a Langmuir-Blodgett monolayer of tetra-tertiary-butyl metal-free
phthalocyanine ((t-bu)₄H₂Pc) on silver island films with arachidic acid LB monolayer spacer layers. The fluorescence spectrum of an LB monolayer of (t-bu)₄H₂Pc with no spacer layer on the Ag film was 200 times higher in intensity than that of an LB monolayer of (t-bu)₄H₂Pc on glass. The fluorescence enhancement factor with five monolayers of spacer was approximately 400. Two processes compete in this system, radiationless decay to the metal and local field enhancement. There are other processes such as absorption, second harmonic generation and luminescence to name a few which are enhanced by rough metal surfaces.

3.3. NON-ELECTROMAGNETIC THEORIES FOR SERS

An overview of the important non-electromagnetic theories for SERS follows. The theories are usually grouped under the umbrella of "chemical" enhancement mechanisms. There are some experimental observations which are difficult to explain in the context of EM theory. There are certain molecules which display a "first layer" effect. The enhancement factor for molecules in the layer adjacent to the metal surface is much higher than predicted by EM theory. Charge transfer (CT) excitations can occur between a metal and an adsorbate for specific systems and would affect the first layer only. An example would be an Ag-C₂H₄ complex which displays SERS in the visible region. An optical absorption band has been observed for this complex at 550 nm which corresponds to a charge transfer excitation of an Ag 5s state to the lowest unoccupied molecular orbital (LUMO), a π* b₂g state. Charge transfer can take place from metal to adsorbate or adsorbate to metal. Molecules such as CH₄ and C₂H₆ do not have affinity levels of the correct energy for SERS in the visible region. One way of thinking of charge transfer excitations is to make an analogy to the resonance Raman effect. The molecular polarizability
derivative tensor is changed and this causes increased Raman scattering. A resonance occurs when the laser excitation coincides with the charge transfer energy.

Molecules such as CO and N₂ have the same Raman scattering cross-section. Experimentally, SERS of CO on cold-deposited silver is fifty times more intense than N₂ on the same substrate. The CO band on Ag is shifted 28 cm⁻¹ from the Raman band of free CO. Nitrogen, however, has a minor shift of 2 cm⁻¹ in frequency when it is adsorbed on cold-deposited Ag. The fifty-fold difference in intensity cannot be accounted for by preferred orientation of CO on Ag alone. Adsorbates on SERS-active substrates can be either physisorbed or chemisorbed. Physisorbed molecules have van der Waals type of interactions with the surface. The Raman frequencies and relative intensities are not expected to differ greatly from the adsorbate in the absence of the surface. On the other hand, chemisorbed molecules have a strong interaction with the metal leading to bond formation. The frequencies and relative intensities can be quite different from the molecule alone. The molecular electronic distribution has been changed and the molecular polarizability derivative tensor is that of the surface-adsorbate complex. The electromagnetic mechanism would operate on the surface-adsorbate complex. A second possibility is that there are two SERS mechanisms operating simultaneously. One which causes a fifty-fold increase over the other electromagnetic enhancement.

Excitation profiles are a plot of SERS intensity against excitation wavelength. Usually the intensities are calibrated with an internal standard. There have been some unusual results obtained for adsorbates on roughened
electrodes at different potentials\textsuperscript{21,22}. The intensity maxima occur at different potentials for different excitation wavelengths. One explanation is that the potential affects the gap between the energy levels of the metal and adsorbate\textsuperscript{2}. If the charge transfer is from the metal to the adsorbate, a negative potential will push the Fermi level up and lead to a red shift in the CT excitation wavelength\textsuperscript{7}. Conversely, for CT from an adsorbate to metal, positive potentials push the Fermi level down and lead to a blue shift in the CT excitation\textsuperscript{7}. The Raman molecular polarizability derivative, $\alpha$ can be expressed in terms of the Kramers-Heisenberg equation\textsuperscript{7}:

$$
\alpha = \frac{\sum_i \langle f | H | i \rangle \langle i | H | g \rangle}{(E_i - E_g) - h\nu + i\Gamma_i} + \frac{\sum_i \langle f | H | i \rangle \langle i | H | g \rangle}{(E_i - E_g) + h\nu}
$$

[equation 3.3]

$g$, $i$ and $f$ are ground, intermediate and final states of the molecule. $H$ is the Hamiltonian operator for molecule-radiation interaction. $\Gamma_i$ is the natural linewidth of the intermediate state. $\nu$ is the frequency of the incident light.

The denominator becomes small and $\alpha$ large in equation 3.3 when $\text{CT} = h\nu = E_i - E_g$. A large molecular polarizability derivative causes increased Raman scattering and equation 3.3 is the condition for the resonance Raman effect. There would also be increased scattering due to the presence of the rough metal.
The CT (excited model) has been described in which a resonance Raman effect leads to increased Raman scattering. There are a couple of ground state charge transfer models 23-27. In the ground state, there is some amount of charge transfer between the metal and adsorbate which is modulated by molecular vibrations4,23,24. The total polarizability is affected by this electron transfer and vibrations which change the metal-adsorbate distance are influenced the most. The second ground state CT model involves vibrations that push and pull electrons to and from the metal surface and vibrationally modulate the reflectivity of the surface or Mie scattering of metal particles5,25-27.

Three classes of biomolecules were deposited on silver island films in an attempt to determine short-range and classical electromagnetic contributions to SERS28. The first class of adsorbates were biomolecules such as adenine and calf thymus deoxyribonucleic acid (CT-DNA) which have electronic transitions in the UV region only. Secondly, flavine adenine dinucleotide (FAD) and doxorubicin (DOX) were compounds with low coefficient of extinction electronic transitions in the visible region. The third category included β-carotene which has a high coefficient of extinction electronic transition in the visible region. The silver island films had extinction spectra with maxima from 450-660 nm depending on the thickness of the metal. The term "extinction" was used because the spectra were not corrected for scattering and reflectance. Excitation profiles were obtained for different bands of each molecule on the metal and enhancement factors calculated. The first two classes of molecules had chemical interactions with the silver surface and the Raman enhancement was short-ranged. The β-carotene on the other hand, displayed long-range enhancement characteristic of a classical electromagnetic
enhancement. The type of enhancement mechanism which is operative may depend not only on the chemical nature of the adsorbate, but also its electronic absorption\textsuperscript{28}. For the \(\beta\)-carotene, the maximum SERS enhancement occurred in the region of maximum overlap between the molecular electronic absorption and the plasmon absorption of the metal island film\textsuperscript{28}.

Otto et al. have described a search for SERS-active sites\textsuperscript{2}. The adatom hypothesis involves strong electron-photon coupling at regions of atomic scale roughness leading to electron tunneling from the metal via the adatoms to the electronic states of the adsorbate\textsuperscript{29,30}. Adatoms are clusters of metal atoms much smaller than the surface roughness features 10-100 nm normally discussed in regards to EM theory. Regions of atomic scale roughness (ASR) are believed to consist of clusters of five or six atoms\textsuperscript{1}. Otto et al.\textsuperscript{2} also described adsorption sites of lowered affinity called "E-sites" which have extra lines compared to normal lines of the adsorbates. There are frequency shifts at the E-sites and the adsorbate is chemisorbed.

Campion\textsuperscript{7} has taken smooth metal surfaces with very low enhancement factors and created defects of well-known structure. Defects were created on high index faces of single crystals of silver by cutting, thermal evaporation of adatoms or by underpotential deposition. The observations were that on Ag(521), Ag(987) stepped and kinked surfaces, the Raman spectrum of physisorbed pyridine is unenhanced\textsuperscript{7}. On Ag(540) the pyridine was chemisorbed but the Raman spectrum was not enhanced. Therefore, chemisorption alone is not sufficient to produce SERS\textsuperscript{7}. 
Moskovits has addressed challenges to the electromagnetic model. For instance, most EM calculations assume a molecule is physisorbed to the metal. There is no reason why one cannot consider chemisorption. The only adjustment that is necessary is to use the polarizability tensor of the metal-adsorbate complex and assume the EM field is acting on this complex. Adatoms are not necessary for SERS for the CT-model, only chemisorption sites. The intensity of adsorbates on cold deposited porous films usually increases with increasing substrate temperature and then decreases. Rather than invoking adatoms, the results can be explained by the idea that the surface structures of the films are being changed and tuned to allow EM enhancement. The weakness or absence of SERS for water on electrodes in aqueous systems can be explained by the fact that water is inherently a weak Raman scatterer and secondly the high vibrational frequency does not allow the simultaneous resonance between the incident and Raman scattered excitation and the surface plasmon. The saturation of the SERS effect with coverage can often be explained by the effect metal curvature has on the distance dependence of SERS. Features of low curvature produce the longest-ranged enhancement. The experiments with CO and N\textsubscript{2} on SERS-active surfaces have been mentioned and differences between ethylene and ethane. In the later case, there appears to be another mechanism operating on top of the EM enhancement. SERS spectra do not have to be the enhanced version of the normal Raman spectrum. This will be discussed in the section of this chapter on surface-selection rules which allow for different relative intensities of bands. Under a couple of circumstances, "forbidden" bands are also possible.

3.4 DISTANCE AND COVERAGE DEPENDENCE OF SERS ENHANCEMENT FACTOR.
The distance dependence of the SERS enhancement factor should provide evidence for either an electromagnetic mechanism or chemical mechanism. A chemical mechanism requires immediate contact with the SERS-active substrate. The electromagnetic mechanism can be relatively long-ranged depending on the substrate geometry. Murray et al.\textsuperscript{31-33} carried out the earliest SERS distance dependence studies. The sample geometry was complex consisting of the following layers: CaF\textsubscript{2} underlayer / vacuum evaporated Al\textsubscript{2}O\textsubscript{3} / dip coated adsorbate / spin coated spacer / top vacuum evaporated Ag. The adsorbate to silver distance was not well-defined because of the protrusions in the CaF\textsubscript{2} layer influencing subsequent layers and enhancement could have also been due to the Al-Al\textsubscript{2}O\textsubscript{3}.

SERS distance dependence results for Langmuir-Blodgett films on metal islands were reported independently by two groups\textsuperscript{34,35}. Cotton et al.\textsuperscript{35} spaced dyes from silver island films. Kovacs et al.\textsuperscript{34} used LB spacer layers of arachidic acid of well known thicknesses to separate tetra-tertiary-butyl metal-free phthalocyanine ((t-bu)\textsubscript{4}H\textsubscript{2}Pc) from silver and indium island films. Calculation of an absolute enhancement factor was possible by comparison with the intensity of an unenhanced spectrum of one or more LB monolayers of (t-bu)\textsubscript{4}H\textsubscript{2}Pc. Weak enhancement was observed even at 15 nm from the surface. The data in both cases\textsuperscript{34,35} supported the notion of an electromagnetic mechanism.

Silver island films have been prepared in UHV on sapphire substrates\textsuperscript{36}. At low temperatures, cyclohexane molecules were condensed on top of the metal and were used as spacer molecules for subsequent deposition of aromatic molecules such as benzene. The intensity of the Raman band assigned to C-C stretching of benzene reached a maximum at 1.3-1.7
monolayer coverage with an enhancement factor of 10^4. When a clean silver island surface was exposed to oxygen prior to aromatic molecules, the 10^4 enhancement was lost and the long-range classical electromagnetic enhancement was observed. The explanation provided was that atomic oxygen passivates the surface by adsorbing on regions of atomic scale roughness or defects^{36}.

There have been many studies related to distance-dependence and coverage dependence using pyridine adsorbed on metal surfaces^{3}. If chemisorption sites are present in low concentrations and are necessary, the SERS signal should have a maximum value at a low surface coverage. Adsorption of additional physisorbed species can lower the local field^{3}. The coverage dependence of SERS of cobalt phthalocyanine (CoPc) on roughened films has been measured and the maximum intensity was observed for 0.07 monolayer^{37}. An electromagnetic model could explain the results and damping of the plasmon resonance above 0.1 monolayers. If there are no special adsorption sites then the normal EM enhancement is in effect. If molecules are adsorbed, the effect on the dielectric constant of the medium must be considered^{3}.

3.5 SURFACE-SELECTION RULES

The frequencies and relative intensities of Raman bands of a molecule are influenced by the presence of a surface and are governed by surface selection rules^{38}. The model begins with a molecule above a smooth metal surface. The orthogonal coordinate system consists of the z-axis normal to the surface and the plane of incidence is yz. There are two possible polarizations of incident linearly polarized light, P with the electric field parallel to the plane of
incidence and $S$ with the electric field vector perpendicular to the plane of incidence. The electric field at the surface is due to incident and reflected beams. The angle the incident beam makes with the surface normal is $\phi$. The electric field components have been expressed in terms of Fresnel coefficients$^3$:

$$E_x = (1 + r_s)E_S \quad \text{[equation 3.4]}$$

$$E_y = (r_p-1)\cos\phi E_p \quad \text{[equation 3.5]}$$

$$E_z = (1 + r_p)\sin\phi E_p \quad \text{[equation 3.6]}$$

$$r_s = \frac{\cos\phi - (\varepsilon - \sin^2\phi)^{1/2}}{\cos\phi + (\varepsilon - \sin^2\phi)^{1/2}} \quad \text{[equation 3.7]}$$

$$r_p = \frac{\varepsilon \cos\phi - (\varepsilon - \sin^2\phi)^{1/2}}{\varepsilon \cos\phi + (\varepsilon - \sin^2\phi)^{1/2}} \quad \text{[equation 3.8]}$$

The Fresnel coefficients $r_s$ and $r_p$ are defined by equations 3.7 to 3.8$^1$. $\varepsilon$ is the dielectric function of the metal.
The squared amplitude of the electric fields ($E_{zp}^2$, $E_{yp}^2$, $E_{xs}^2$) have been calculated for different angles of incidence (0 to 90°) with 514.5 nm excitation of silver and indium surfaces. The conclusions are that the field parallel to the surface is small for good reflectors, the perpendicular component is strong due to the superposition of incident and reflected fields and best Raman enhancement should occur for P-polarized light at a calculated 60° angle of incidence.

Four combinations of polarizations are possible experimentally based on the polarization of the incident light and analyzed light after a 90° Raman scattering experiment: SS, PS, SP and PP. Expressions similar to 3.4 to 3.8 can be generated for scattered light. The primes ('s) in subsequent expressions indicate scattered radiation. To define this model further, the geometry of the molecule must be considered relative to the surface. The simplest case has the coordinates of the molecule and surface in common. The expressions for the four polarizations according to Moskovits are given in equations 3.9 to 3.12. The molecular polarizability derivative tensor component elements are $\alpha_{ij}$.

$$SS \propto |\alpha_{xx} (1+r_s)(1+r'_s)|^2$$  \hspace{1cm} (equation 3.9)

$$SP \propto |\alpha_{yx} (1+r_s)(1-r'_p) \cos \phi' + \alpha_{zx} (1+r_s)(1+r'_p) \sin \phi'|^2$$  \hspace{1cm} (equation 3.10)
\[ PP \propto \left\{ \alpha_{yy} (r_p - 1) \cos \phi + \alpha_{yx} (1 + r_p) \sin \phi \right\} (1 - r_p') \cos \phi' + \left\{ \alpha_{zy} (r_p - 1) \cos \phi + \alpha_{zz} (1 + r_p) \sin \phi \right\} (1 + r_p') \sin \phi' \mid^2 \]

(equation 3.11)

\[ PS \propto \mid \alpha_{xy} (r_p - 1) (1 + r_s') \cos \phi + \alpha_{xz} (1 + r_p) (1 + r_s') \sin \phi \mid^2 \]

(equation 3.12)

Returning to the Fresnel coefficient expressions in 3.7 and 3.8, as \( | \epsilon | \) becomes very large, which is true to the red\(^1\), \( r_s \to -1 \) and \( r_p \to +1 \). The tangential field component of the surface becomes close to zero and the normal component of the incident field is increased. The term in PP which is related to \( \alpha_{zz} \) remains. Raman-active modes of the molecule are observed which have a non-zero \( \alpha_{zz} \) component. This is an example of how the relative intensities of Raman bands can be quite different for a molecule near a metal surface compared to the free molecule.

The condition necessary for the tangential field component being stronger than the normal component is that \( 1 / | \epsilon | \) becomes very large\(^1\), making \( r_s \to +1 \) and \( r_p \to -1 \). Raman modes with non-vanishing \( \alpha_{xx}, \alpha_{yy}, \alpha_{xy} \) are expected. Raman bands of molecules on smooth surfaces fall into three classes\(^1\): those with \( \alpha_{zz} \), those with one or all of \( \alpha_{xx}, \alpha_{yy}, \alpha_{xy} \) and one or both of \( \alpha_{xz} \) and \( \alpha_{yz} \). The excitation profile (intensity versus wavelength) for each class of bands is different.
The next step in the development of surface selection rules is to consider molecules adsorbed on metal spheres\textsuperscript{3}. If the same coordinate system exists for the molecule and the sphere, the $\alpha_{zz}$ components are enhanced when the excitation frequency is to the red of the plasmon frequency, followed by $\alpha_{xz}$ and $\alpha_{yz}$ components. If SERS experiments are carried out just to the blue of the plasmon frequency, all types of modes are equally enhanced appearing to violate the surface selection rule\textsuperscript{1} in which modes which are normal to the metal surface are strong.

The surface selection rules affect the relative intensities of "allowed bands" in SERS spectrum. There are two ways to have "forbidden modes" appear in SERS spectra. Firstly, the symmetry of the adsorbate can be lowered by bonding to the surface. Secondly, a steep field gradient near the surface makes it necessary to include additional terms in the induced dipole moment expression.

The orientation of adsorbed molecules on SERS-active substrates can be determined by applying surface selection rules\textsuperscript{39-43}. The orientation of low symmetry molecules on metal was calculated based on the relative intensities of $v_{\text{sym}}$(C-H) and $v_{\text{asym}}$(C-H) vibrations of methyl groups\textsuperscript{43}. Two assumptions must be made. The polarizability tensor should not change when a molecule is adsorbed to the surface. Therefore, relative intensities of adsorbate bands should not be due to a resonance Raman effect. The total radiation experienced by the adsorbate must be a vector sum of the normal and tangential electric field components. In general, this approach could be used for two modes of known spatial relation that have discrete frequencies. The polarizability ellipsoid changes along the adjacent carbon to methyl carbon for
the $v_{\text{sym}}$(C-H) mode. The $v_{\text{asym}}$(C-H) mode has polarizability changes in a perpendicular direction.

### 3.6 MODEL CALCULATIONS OF SERS ENHANCEMENT FACTORS

A computer program\textsuperscript{44} written in FORTRAN for a VAX system was used to calculate the enhancement factor of a molecule above a prolate metal hemispheroid according to Nitzan and Gersten\textsuperscript{16}. The identity of the metal hemispheroid was varied and also the excitation wavelength. Literature values of dielectric constants of metals at the excitation and scattering wavelengths were collected and input to the program. The surrounding medium was assumed to have $\varepsilon_0=1$. The excitation wavelengths used as input were 488.0 nm (Ar$^+$ laser), 514.5 nm (Ar$^+$ laser), 647.1 nm (Kr$^+$ laser) and 1064.1 nm (Nd/YAG laser).

The molecule selected for the calculations was a phthalocyanine derivative, tetra-tertiary-butyl metal-free phthalocyanine ((t-bu)$_4$H$_2$Pc). In the past, Langmuir-Blodgett monolayers of this material were deposited on vacuum evaporated metal island films\textsuperscript{45-47}. The enhancement factors were calculated for a band which appears at a frequency shift of 686 cm$^{-1}$ corresponding to a macrocycle breathing mode. For the purpose of these calculations, the geometry of the hemi-spheroid was fixed with a semi-major axis of 38 nm and a semi-minor axis of 33 nm. Different values could also be input based on geometries that are possible experimentally. The enhancement factors were calculated at the top of the spheroid and to a distance 24.50 nm using 0.50 nm increments. The enhancement factor (EF) expression according to Nitzan and Gersten was\textsuperscript{16}: 

\begin{equation}
\text{EF} = \frac{(\varepsilon_2 - \varepsilon_1) - \frac{1}{2} (\varepsilon_2 + \varepsilon_1)}{(\varepsilon_0 - \varepsilon_1) - \frac{1}{2} (\varepsilon_0 + \varepsilon_1)}
\end{equation}
EF = |A(ω_l)|^2 |A(ω_s)|^2 \quad [\text{equation 3.13}]

ω_l is the laser excitation wavelength
ω_s is the wavelength of scattered radiation

\[
A(\omega) = 1 + \frac{\varepsilon_0 - \varepsilon(\omega)}{\varepsilon(\omega) - \varepsilon_0 P} \quad [\text{equation 3.14}]
\]

\varepsilon_0 is the dielectric constant of the surrounding medium
\varepsilon(\omega) is the wavelength dependent dielectric constant for the metal
L and P are geometrical factors

\[
L = \frac{\xi_0 Q_1'(\xi_1)}{Q_1(\xi_0)} \quad [\text{equation 3.15}]
\]

\[
P = \frac{\xi_0 Q_1'(\xi_0)}{Q_1(\xi_0)} \quad [\text{equation 3.16}]
\]

Q_1 is a Legendre function of the second kind

The geometry of the prolate hemi-spheroid used for the calculations is shown in Figure 3-1. The focus point f, is related to the semi-major axis b and semi-minor axis a through equation 3.17. The height above the tip of the
spheroid is H. The geometrical terms $\xi_0$ and $\xi_1$ are related to $a$, $H$ and $f$ through equations 3.18 and 3.19.

\[
\begin{align*}
f &= (a^2 - b^2)^{1/2} & \text{[equation 3.17]} \\
\xi_0 &= a/f & \text{[equation 3.18]} \\
\xi_1 &= (a + H)/f & \text{[equation 3.19]}
\end{align*}
\]

An example of the prolate enhancement factor output for a gold hemi-spheroid with semi-major axis=38 nm and semi-minor axis=33 nm and a laser excitation wavelength of 1064.1 nm appears in Appendix I. The enhancement factor drops rapidly from 155 for a molecule at the top of the metal to 4.48 at a distance of 24.50 nm from the metal. The first set of enhancement factor calculations were completed for (t-bu)$_4$H$_2$Pc on silver, gold and copper metal hemi-spheroids using various laser excitation wavelengths. The input parameters and calculated enhancement factors for a molecule at the top of the Ag, Au and Cu hemi-spheroids appear in Table 3-1. In the case of silver, the enhancement factor decreases with increasing wavelength. The EF was modeled to be 403 using 488.0 nm laser excitation and 151 for 1064.1 nm excitation. Gold and copper display a slightly different behavior from silver with the highest enhancement calculated for 514.5 nm excitation with EF values of 690 and 272 respectively.
Figure 3-1: Prolate hemi-spheroid geometry^16 used for enhancement factor model calculation.
Table 3-1: Calculated Enhancement Factors for Ag, Au, Cu and Ni.

<table>
<thead>
<tr>
<th>METAL</th>
<th>Reference for ε values</th>
<th>EF laser line = 488.0 nm</th>
<th>EF laser line = 514.5 nm</th>
<th>EF laser line = 647.1 nm</th>
<th>EF laser line = 1064.1 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ag</td>
<td>Johnson &amp; Cristy⁴⁸</td>
<td>403</td>
<td>332</td>
<td>213</td>
<td>151</td>
</tr>
<tr>
<td>Au</td>
<td>&quot;</td>
<td>270</td>
<td>690</td>
<td>274</td>
<td>155</td>
</tr>
<tr>
<td>Cu</td>
<td>&quot;</td>
<td>253</td>
<td>272</td>
<td>268</td>
<td>154</td>
</tr>
<tr>
<td>Ni</td>
<td>Johnson &amp; Cristy⁴⁹</td>
<td>179</td>
<td>177</td>
<td>169</td>
<td>148</td>
</tr>
</tbody>
</table>

Input parameters: EF for band at shift 687 cm⁻¹, semimajor axis = 38 nm, semiminor axis = 33 nm.
Table 3-2: Calculated Enhancement Factors for Various Metals at 1064 nm.

<table>
<thead>
<tr>
<th>METAL</th>
<th>Reference for $\varepsilon$ values</th>
<th>$\varepsilon_{\text{Re}}$ (1064 nm)</th>
<th>$\varepsilon_{\text{Im}}$ (1064 nm)</th>
<th>$\varepsilon_{\text{Re}}$ (1148 nm)</th>
<th>$\varepsilon_{\text{Im}}$ (1148 nm)</th>
<th>ENHANCEMENT FACTOR (top of spheroid)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>Johnson &amp; Cristy$^{50}$</td>
<td>-10.7</td>
<td>25.5</td>
<td>-12.9</td>
<td>27.6</td>
<td>140</td>
</tr>
<tr>
<td>Cr</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-0.235</td>
<td>25.4</td>
<td>0.254</td>
<td>26.0</td>
<td>127</td>
</tr>
<tr>
<td>Co</td>
<td></td>
<td>-25.2</td>
<td>33.7</td>
<td>-27.8</td>
<td>37.3</td>
<td>147</td>
</tr>
<tr>
<td>Cu</td>
<td>Johnson &amp; Cristy$^{48}$</td>
<td>-49.7</td>
<td>4.97</td>
<td>-59.9</td>
<td>6.56</td>
<td>154</td>
</tr>
<tr>
<td>Au</td>
<td></td>
<td>-48.8</td>
<td>3.64</td>
<td>-58.6</td>
<td>4.78</td>
<td>155</td>
</tr>
<tr>
<td>Ir</td>
<td>Kirillova et al.$^{51,52}$</td>
<td>-34.2</td>
<td>39.2</td>
<td>-38.7</td>
<td>42.5</td>
<td>145</td>
</tr>
<tr>
<td>Fe</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-7.28</td>
<td>23.7</td>
<td>-8.88</td>
<td>25.4</td>
<td>143</td>
</tr>
<tr>
<td>Mn</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-9.56</td>
<td>29.3</td>
<td>-9.99</td>
<td>31.4</td>
<td>140</td>
</tr>
<tr>
<td>Mo</td>
<td>Moritani et al.$^{53}$</td>
<td>-14.3</td>
<td>20.7</td>
<td>-20.6</td>
<td>21.1</td>
<td>160</td>
</tr>
<tr>
<td>Ni</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-18.7</td>
<td>30.1</td>
<td>-20.8</td>
<td>32.5</td>
<td>148</td>
</tr>
<tr>
<td>Os</td>
<td>Lynch et al.$^{54}$</td>
<td>-6.56</td>
<td>14.5</td>
<td>-11.0</td>
<td>16.9</td>
<td>163</td>
</tr>
<tr>
<td>Pd</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-32.8</td>
<td>31.0</td>
<td>-37.1</td>
<td>34.3</td>
<td>150</td>
</tr>
<tr>
<td>Pt</td>
<td>Weaver$^{55}$</td>
<td>-23.0</td>
<td>44.2</td>
<td>-24.5</td>
<td>-49.7</td>
<td>140</td>
</tr>
<tr>
<td>Rh</td>
<td>Weaver et al.$^{56}$</td>
<td>-52.0</td>
<td>57.5</td>
<td>-55.8</td>
<td>61.5</td>
<td>140</td>
</tr>
<tr>
<td>Ag</td>
<td>Johnson &amp; Cristy$^{48}$</td>
<td>-58.3</td>
<td>0.610</td>
<td>-69.3</td>
<td>1.11</td>
<td>151</td>
</tr>
<tr>
<td>Ti</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-4.04</td>
<td>27.8</td>
<td>-4.01</td>
<td>29.1</td>
<td>134</td>
</tr>
<tr>
<td>W</td>
<td>Weaver et al.$^{57}$</td>
<td>-5.15</td>
<td>22.7</td>
<td>-7.37</td>
<td>25.2</td>
<td>140</td>
</tr>
<tr>
<td>V</td>
<td>Johnson &amp; Cristy$^{49}$</td>
<td>-5.54</td>
<td>21.5</td>
<td>-8.76</td>
<td>22.5</td>
<td>144</td>
</tr>
</tbody>
</table>

Input parameters: laser 1064.1 nm, EF for band at shift 687 cm$^{-1}$ (1148 nm), semimajor axis = 38 nm, semiminor axis = 33 nm.
High SERS enhancement factors are expected for prolate hemi-spheroids with large aspect ratios, a/b. This can be accomplished for instance by depositing an underlayer of rough calcium fluoride followed by metal. There has been a report on the fabrication of tailored needle substrates for SERS\textsuperscript{58}. A base layer of latex microspheres was deposited on a glass slide and a metal was evaporated at a grazing angle to produce needles 50-75 nm in diameter with aspect ratios 5:1 or greater. It was possible to obtain good SERS spectra from ca. 100 pg of benzoic acid illuminated on the needles\textsuperscript{58}.

Table 3-2 displays the calculated enhancement factors for a series of metals excited at 1064.1 nm. The modeling indicated that a wide range of metals should give FT-SERS enhancement on the order of 10\textsuperscript{2}. One objective of the later experimental work is to prepare vacuum evaporated metal island films as SERS-active substrates in the near-IR. The SERS-active substrates would allow signals to be obtained from thin solid organic materials which could be vacuum evaporated or deposited as Langmuir-Blodgett monolayers on the metal island film. There have been previous reports in the literature of FT-SERS on metal electrodes and colloids\textsuperscript{59-68}. However, there had not been any reports of FT-SERS using metal island films. Gold and copper were selected for the preparation of metal island substrates for two reasons. Firstly, FT-SERS on Au and Cu had been demonstrated on electrodes and colloids\textsuperscript{60-69}. Secondly, these materials were readily obtainable in a pure form and could be vacuum evaporated. These two metals would demonstrate the feasibility of FT-SERS on vacuum evaporated metal island films knowing that it was likely that the technique could be extended to other metals.
3.7 SUMMARY

Surface-enhanced Raman scattering (SERS) has been observed for many specially prepared rough metal surfaces. Electromagnetic theory requires collective oscillations of electrons in metals which are surface plasmons. Roughness is required to have radiative plasmons. The electric fields at or near rough metal surfaces are greatly enhanced and localized. An adsorbate experiences an effective electric field due to the incident light, Raman scattered radiation and the surface plasmons. There are chemical models which require immediate contact between the adsorbate and metal. The SERS enhancement factor has a distance and coverage dependence. Surface-selection rules have been described which affect the relative intensities of the observed SERS bands. The presence of forbidden frequencies is explained by either a lowering of adsorbate symmetry on the surface or extremely high electric field gradients. Model calculations of SERS enhancement factors were completed for various metals at visible and near-IR frequencies to guide the later experimental work.
Chapter 3: References


44. FORTRAN program written by G. J. Kovacs, Xerox Research Centre of Canada and modified by C. A. Jennings.


68. Wentrup-Byrne, E.; Sarinas, S.; Fredericks, P. M. *Applied Spectroscopy* 1993, 47(8), 1192.
CHAPTER 4: FT-RAMAN SPECTROSCOPY OF THIN FILMS OF TITANYL PHTHALOCYANINE AND VANADYL PHTHALOCYANINE

4.1. INTRODUCTION

The FT-Raman spectra of titanyl phthalocyanine (TiOPc) and vanadyl phthalocyanine (VOPc) polymorphs are presented. The characterization of polymorphs was corroborated by X-ray data. Both materials were vacuum evaporated onto a substrate whose temperature was varied during vacuum evaporation. The FT-Raman spectra of the films formed at different temperatures are discussed. It is shown that FT-Raman is an appropriate analytical technique to distinguish between polymorphic forms.

Phthalocyanines (Pcs) have a wide range of applications. Some Pcs exhibit near-IR photosensitivity and could be used in conjunction with laser diodes in high-speed printers. A typical photoreceptor structure consisting of a conductive substrate, a charge generator layer and a charge transport layer has been given in Chapter 1. Optional layers in the photoreceptor include an adhesive layer, blocking layer and an overcoat. The phthalocyanine is part of the charge generator layer and is normally in the form of a dispersion in a polymeric binder. The types of phthalocyanines considered for this application are: X-metal-free phthalocyanine (X-H₂Pc)¹⁻³; trivalent metal-Pcs such as chloroaluminum phthalocyanine (AlPcCl)⁴, chloroindium phthalocyanine (InPcCl)⁵,⁶; and tetravalent metal-Pcs such as vanadyl phthalocyanine (VOPc)⁷ and titanyl phthalocyanine (TiOPc)⁸. Third-order non-linear optical susceptibilities have been measured for three polymorphs of TiOPc⁹.
Candidates for non-linear optical waveguides include VOPc, substituted tetra-tertiary-butylphthalocyanines and TiOPc\textsuperscript{10}. Properties such as photosensitivity and non-linear optical effects are influenced by molecular packing and orientation.

Figure 4-1 gives the molecular structure of a tetravalent metal-phthalocyanine such as TiOPc and VOPc. Both TiOPc and VOPc belong to the $C_{4v}$ symmetry point group.

Several crystalline polymorphic forms of titanyl phthalocyanine exist and the nomenclature will be introduced here. Polymorphs can be identified on the basis of their X-ray powder diffraction patterns shown in Figure 4-2 for type I, type II, type III and type IV TiOPc. Single crystal structures of type I and type II TiOPc have been reported by Hiller and others\textsuperscript{11}. Type I is monoclinic with four molecules per unit cell and space group $P2_1/c$ while type II is triclinic with two molecules per unit cell and space group P-1. Single crystals of type III and type IV have not been obtained, therefore no single crystal structure has been reported. Rietveld refinement was used to solve crystal structures of TiOPc from X-ray powder diffraction data\textsuperscript{12}. The results for type I and type II TiOPc were obtained by comparing the data with known crystal structures\textsuperscript{11}. The most probable crystal structure for type IV TiOPc has been proposed\textsuperscript{12}. The best fit structure by Rietveld refinement for type IV TiOPc was a triclinic unit cell with two molecules per unit cell and space group P-1\textsuperscript{12}. Enokida has described the preparation and properties of five polymorphs of TiOPc: $\alpha$, $\beta$, $\gamma$, m and an amorphous form\textsuperscript{8}.
Figure 4-1: Molecular structure of TiOPc and VOPc.
Figure 4-1

\[ M = \text{Ti or V} \]
Figure 4-2: X-ray powder diffraction patterns of TiOPc polymorphs (types I, II, III, & IV).
Figure 4-2

[Graph showing data for types I, II, III, and IV with counts on the x-axis and another value on the y-axis]
Three polymorphic forms (I, II and III) have been observed for vanadyl phthalocyanine\textsuperscript{13}. Type I occurs if VOPc is vacuum deposited onto fused quartz at room temperature. Type II is prepared by crystal growth from solvents such as quinoline or 1-chloronaphthalene, crystal growth from vapor or by heating phase I to 200°C and above. Phase III occurs if a melt at 610°C is quenched. Type II VOPc is triclinic with \textit{P\overline{1}} symmetry and two molecules per unit cell. A single crystal X-ray structure is not available for type I VOPc.

In the following sections of Chapter 4, the Raman results for a series of thin solid films of TiOPc and VOPc deposited onto a substrate whose temperature was varied during vacuum evaporation are discussed. The technique selected for the study was Fourier-transform Raman spectroscopy (FT-Raman) that provides spontaneous Raman spectra for pigments strongly absorbing in the visible spectral region. The objective of the work was to provide an analytical method to distinguish the different polymorphic forms of these important materials.

4.2. EXPERIMENTAL

4.2.1. Preparation of TiOPc, VOPc and Vacuum Evaporation of Films.

Titanyl phthalocyanine was synthesized by refluxing 1,3-diiminoisoindoline and titanium butoxide in chloronaphthalene followed by washing with boiling \( N, N \)-dimethylformamide and methanol\textsuperscript{14}. The pigment was further purified by a train sublimation process\textsuperscript{15}. An amorphous TiOPc reference film was prepared by vacuum evaporation with the substrate at room temperature. A type II reference film was prepared by vacuum
evaporation with the substrate held at 95°C and subsequent exposure to cyclohexanone vapor for two minutes outside the vacuum chamber. The polymorphic conversion to obtain type IV TiOPc has been described in the patent literature\textsuperscript{16}. The procedure for preparing VOPc from vanadium pentoxide and phthalonitrile has also been described in a patent\textsuperscript{17}.

Corning 7059, 1" X 3" glass slides were coated on one-side with vacuum grease (Dow Corning, \# 4 Compound) and placed against the substrate holder. The vacuum evaporations were carried out in a Vacuum Generators (VG) UHV system from a quartz boat containing either the TiOPc or VOPc powder. The background pressure was typically $9 \times 10^{-5}$ to $1 \times 10^{-4}$ Pa. The sample holder was temperature controlled using either liquid nitrogen for cooling or electrical heating to achieve substrate temperatures from -30°C to 240°C. The film deposition rate was 0.8 to 1.0 nm / second. The film thickness was monitored with a Kronos, Inc., Model QM311 thickness monitor during the deposition. The final thickness was 100 nm for films in the TiOPc and VOPc temperature series and 600 nm for the TiOPc reference films.

4.2.2. X-Ray Powder Diffraction.

Thin film X-ray powder diffraction patterns (XRPD) were collected by a Philips 1710 X-ray powder diffraction system modified with parallel beam optics on the diffraction side. The system used CuK$_\alpha$ radiation ($\lambda=0.1542$ nm) and contained a pulse light analyzer and graphite monochromator. The data were collected at an incidence beam angle of 1.0 degree using a step size of 0.1 degrees and count time of 10 seconds at each step.
4.2.3. **FT-Raman Spectroscopy.**

A Bomem Ramspec 150 spectrophotometer with a Nd/YAG laser emitting at 1064.1 nm and an InGaAs detector were used to acquire FT-Raman spectra. The system had a Michelson interferometer and was interfaced to a NEC Powermate computer. Bomem Easy Software was used to collect the data and files were imported to Spectra Calc (Galactic Industries Corp.). All Raman files were ratioed against an instrument response curve. A back-scattering geometry was used with resolution set at 4 cm\(^{-1}\). Typically 525 scans were averaged requiring 50 minutes to collect the data. The laser power at the sample was 0.75 watt for the 600 nm amorphous TiOPc reference film, 0.025 watt for the type I TiOPc solid sample, 0.10 watt for the 600 nm type II TiOPc reference film, and 0.025 watt for the type IV TiOPc solid sample. The laser power for 100 nm vacuum evaporated films of both TiOPc and VOPc was 1.82 watts.

4.3. **RESULTS AND DISCUSSION**

The FT-Raman spectra of a 600 nm film of amorphous TiOPc on glass (evaporated at room temperature), type I TiOPc solid in a melting point capillary tube, a 600 nm film of type II TiOPc on glass (evaporated at 95°C and subsequently exposed to cyclohexanone vapor for 2 minutes), and type IV TiOPc solid in a melting point capillary tube appear in Figure 4-3 and can be considered reference materials. Table 4-1 lists the frequencies and intensities of these polymorphs of TiOPc and band assignments are given. The Raman bands have been assigned using model compounds such as benzene\(^{18}\), pyrrole\(^{19}\), indole\(^{20}\) and porphyrin\(^{21}\). Titanyl phthalocyanine belongs to the point group C\(_{4v}\) and since there are 58 atoms, 168 vibrations are expected. The total irreducible representation consists of the following species: \(\Gamma = 23a_1 + \)
19a_2 + 21b_1 + 21b_2 + 42e (doubly degenerate). Based on C_{4v} symmetry, the a_1 and e species are both Raman and IR active, while b_1 and b_2 species are Raman active only. The a_2 species are neither Raman nor IR active.

Raman depolarization ratios in thin solid films can be used to interpret the symmetry type of observed bands. The theory has been developed in a previous paper\textsuperscript{22} for thin solid films. Three sets of cartesian coordinates are required, one each for the laboratory, substrate and molecule. The laboratory coordinates describe the laser and collection of Raman scattered light. The substrate position is defined by an angle β between the incident laser beam and substrate normal. Finally the molecular orientation with respect to the substrate is defined by three Euler angles, φ, Θ, and Ψ. Transformations are carried out to go from molecular to substrate coordinates and then from substrate to laboratory coordinates. Four polarizations of light SS, SP, PS and PP are available experimentally by adjusting a polarizer at the output of the laser and an analyzer after the sample. A computer program can be used to generate theoretical curves for the value of the depolarization ratio SS/SP as a function of the experimental angle β and molecular stacking angle Θ for a particular symmetry type, a_1, b_1, b_2 or e. The experimentally measured depolarization ratio is then compared with theoretical curves to assist in the assignment of the symmetry type. The SS/SP depolarization ratio has been experimentally measured for TiOPc thin solid films on glass using the 647.1 nm Kr\textsuperscript{+} laser line and was compared with theoretical values\textsuperscript{23}. The symmetry types listed in Table 4-1 were based on the values of the SS/SP depolarization ratios and on the activity of the vibrations in the Raman and IR spectra.
Figure 4-3: FT-Raman spectra of various polymorphs of TiOPc.  
a) 600 nm amorphous film  
b) solid type I  
c) 600 nm type II film  
d) solid type IV
Figure 4-3

TIOPc
678
678
749

a) 600 nm amorphous film

b) solid type I

1511
1505

678
750

678

1337

1143

1509

(600 nm type II film)

1400

Wavenumbers (cm⁻¹)

Intensity (arbitrary units)
Table 4-1: FT-Raman Frequencies of Selected TiOPc Polymorphs

<table>
<thead>
<tr>
<th>600 nm RT amorphous film (cm⁻¹)</th>
<th>Type I solid in capillary tube (cm⁻¹)</th>
<th>600 nm film type II (cm⁻¹)</th>
<th>Type IV solid in capillary tube (cm⁻¹)</th>
<th>Assignments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1524</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>C-N pyrrole stretch, a₁</td>
</tr>
<tr>
<td>1511</td>
<td>1505</td>
<td>1513</td>
<td>1509</td>
<td>benzene (v₁₉₉), a₁</td>
</tr>
<tr>
<td>1449</td>
<td>1453</td>
<td>1450</td>
<td>1453</td>
<td>----</td>
</tr>
<tr>
<td>1432</td>
<td>1428</td>
<td>1429</td>
<td>----</td>
<td>isoindole stretch, a₁</td>
</tr>
<tr>
<td>1337</td>
<td>1343</td>
<td>1340</td>
<td>----</td>
<td>pyrrole stretch, a₁</td>
</tr>
<tr>
<td>1305</td>
<td>1312</td>
<td>1303</td>
<td>1316</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1206</td>
<td>1213</td>
<td>----</td>
<td>----</td>
<td>C-H bend</td>
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<td>1187</td>
<td>1192</td>
<td>1191</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1164</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1143</td>
<td>1147</td>
<td>1141</td>
<td>1148</td>
<td>pyrrole ring breathing</td>
</tr>
<tr>
<td>1104</td>
<td>1108</td>
<td>1105</td>
<td>1108</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1027</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1006</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>C-H bend, a₁</td>
</tr>
<tr>
<td>938</td>
<td>----</td>
<td>----</td>
<td>----</td>
<td>----</td>
</tr>
<tr>
<td>----</td>
<td>----</td>
<td>836</td>
<td>834</td>
<td>macrocycle stretch, a₁</td>
</tr>
<tr>
<td>789</td>
<td>----</td>
<td>----</td>
<td>788</td>
<td>macrocycle stretch</td>
</tr>
<tr>
<td>----</td>
<td>----</td>
<td>----</td>
<td>767</td>
<td>----</td>
</tr>
<tr>
<td>749</td>
<td>751</td>
<td>750</td>
<td>750</td>
<td>benzene (v₁), e</td>
</tr>
<tr>
<td>678</td>
<td>680</td>
<td>678</td>
<td>678</td>
<td>macrocycle breathing, a₁</td>
</tr>
<tr>
<td>589</td>
<td>----</td>
<td>590</td>
<td>----</td>
<td>benzene radical</td>
</tr>
<tr>
<td>483</td>
<td>----</td>
<td>483</td>
<td>----</td>
<td>isoindole ring def.</td>
</tr>
</tbody>
</table>
Titanyl phthalocyanine polymorphs display subtle differences in the FT-Raman spectra in the region around 1500 cm\(^{-1}\). The amorphous film has a very strong band at 1511 cm\(^{-1}\) and a shoulder at 1524 cm\(^{-1}\). Types I, II and IV have a single strong band in the range 1505 cm\(^{-1}\) to 1513 cm\(^{-1}\). The CN stretching vibration in pyrrole\(^{19}\) occurs at 1530 cm\(^{-1}\). Vibrations normally associated with carbon-carbon stretching in benzene rings from 1565 cm\(^{-1}\) to 1620 cm\(^{-1}\) were extremely weak or absent. Carbon-hydrogen stretching bands normally not seen in the Raman spectrum also were not observed in the FT-Raman spectra because they do not cause significant changes in the polarizability of the large TiOPc molecule. The macrocycle refers to the inner core of the phthalocyanine which consists of sixteen alternating carbon and nitrogen atoms which are conjugated. Titanyl phthalocyanine polymorphs have slightly different infrared spectra\(^8\). The most sensitive regions to polymorphic form\(^8\) were 700-800 cm\(^{-1}\) and 1600-2000 cm\(^{-1}\).

FT-Raman spectra of 100 nm TiOPc films covering a substrate temperature range of -30\(^\circ\)C to 240\(^\circ\)C during vacuum evaporation appear in Figure 4-4. The Raman scattering of these films was quite weak. In the region of 470 cm\(^{-1}\), a large background was present due to the Corning glass slide. Substrate temperatures of -30\(^\circ\)C to room temperature produced the same polymorphic form with a visible absorption band at 720 nm and X-ray diffraction results indicating amorphous TiOPc. As the temperature was increased, the stable type II TiOPc was formed with a typical X-ray peak at 2\(\theta\)=7.5\(^\circ\). Pure type II TiOPc at 102\(^\circ\)C has a characteristic Raman band at 1513 cm\(^{-1}\) which is not split as observed for the amorphous form.
Figure 4-4: FT-Raman spectra of 100 nm TiOPc films deposited with various substrate temperatures.
Vanadyl phthalocyanine also belongs to the \( C_{4v} \) point group and 168 vibrations are expected. Again the \( a_1, e, b_1 \) and \( b_2 \) species are Raman active. FT-Raman spectra of 100 nm vacuum evaporated VOPc films on glass at room temperature, 90°C, 210°C and 240°C are shown in Figure 4-5. Assignments of observed FT-Raman bands are shown in Table 4-2. Vanadyl phthalocyanine films have a pyrrole C-N stretch at 1522 cm\(^{-1}\) to 1525 cm\(^{-1}\) while type I, II and IV TiOPc polymorphs exhibit the pyrrole C-N stretch at 1505 cm\(^{-1}\) to 1513 cm\(^{-1}\). However, amorphous TiOPc films have a band at 1511 cm\(^{-1}\) with a shoulder at 1524 cm\(^{-1}\).

At room temperature, the VOPc film was amorphous and above 200°C the type II polymorphic form was present. The relative intensities of FT-Raman bands for the room temperature VOPc film and the 240°C VOPc film were quite different. At room temperature, the 1522 cm\(^{-1}\) peak (C-N pyrrole stretch) was more intense than the 680 cm\(^{-1}\) one (macrocycle breathing). At 240°C, the intensities were reversed. The intensity of the 753 cm\(^{-1}\) band (benzene breathing) decreased with increasing temperature. The peak at 785 cm\(^{-1}\) (room temperature) gave way to a peak at 778 cm\(^{-1}\) (240°C). Minor changes with temperature occurred in the C-H bands in the 1200 cm\(^{-1}\) region.
Figure 4-5: FT-Raman spectra of 100 nm VOPc films deposited with various substrate temperatures.
Table 4-2: FT-Raman Frequencies and Relative Intensities of VOPc films.

<table>
<thead>
<tr>
<th>VOPc film room temperature</th>
<th>Assignment</th>
<th>VOPc film deposited on 240°C substrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency (cm⁻¹)</td>
<td>Relative Intensity</td>
<td>Frequency (cm⁻¹)</td>
</tr>
<tr>
<td>1522</td>
<td>(100)</td>
<td>C=C stretch benzene</td>
</tr>
<tr>
<td>1451</td>
<td>(19)</td>
<td>C-N pyrrole stretch</td>
</tr>
<tr>
<td>1432</td>
<td>(2)</td>
<td>isoindole stretch</td>
</tr>
<tr>
<td>1341</td>
<td>(48)</td>
<td>isoindole stretch</td>
</tr>
<tr>
<td>1307</td>
<td>(19)</td>
<td>pyrrole ring stretch</td>
</tr>
<tr>
<td>1210</td>
<td>(6)</td>
<td>C-H</td>
</tr>
<tr>
<td>1197</td>
<td>(7)</td>
<td>C-H</td>
</tr>
<tr>
<td>1187</td>
<td>(13)</td>
<td>C-H</td>
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<td>1162</td>
<td>(5)</td>
<td>C-H</td>
</tr>
<tr>
<td>1143</td>
<td>(21)</td>
<td>pyrrole ring breathing</td>
</tr>
<tr>
<td>1106</td>
<td>(23)</td>
<td>C-H</td>
</tr>
<tr>
<td>1033</td>
<td>(8)</td>
<td>C-H</td>
</tr>
<tr>
<td>1008</td>
<td>(7)</td>
<td></td>
</tr>
<tr>
<td>948</td>
<td>(13)</td>
<td></td>
</tr>
<tr>
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<td>(11)</td>
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<td>785</td>
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<td>benzene (ν₁)</td>
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<td>(7)</td>
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<tr>
<td>724</td>
<td>(7)</td>
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<tr>
<td>591</td>
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</tr>
<tr>
<td>485</td>
<td>(41)</td>
<td>isoindole ring deformation</td>
</tr>
</tbody>
</table>
4.4 CONCLUSIONS

FT-Raman spectra have been obtained for 100 nm vacuum evaporated films of TiOPc and VOPc with substrate temperatures ranging from -30°C to 240°C. The observed frequencies have been assigned to specific vibrations for both of these phthalocyanines. Titanyl phthalocyanine forms in an amorphous form when evaporated onto substrates held at -30°C to room temperature. Pure type II TiOPc forms with substrate temperatures from 102°C to 240°C. Titanyl phthalocyanine type I, II and IV polymorphs can be distinguished on the basis of slight changes in the FT-Raman spectrum, most notably in the C-N pyrrole stretching region. Vanadyl phthalocyanine forms in an amorphous form on room temperature substrates and as type II on substrates above 200°C. There were changes in band frequencies and relative intensities between the VOPc FT-Raman spectra obtained from room temperature films and from films produced at 240°C. TiOPc or VOPc can be identified on the basis of the FT-Raman spectrum. The FT-Raman scattering intensity was weak in general for these thin solid vacuum evaporated films. To improve the signal intensity a thin film could be deposited on a substrate which gives surface-enhanced Raman scattering in the near-IR.24,25.
Chapter 4: References


CHAPTER 5: FOURIER TRANSFORM SURFACE-ENHANCED RAMAN SCATTERING OF EVAPORATED FILMS AND MONOLAYERS

5.1. INTRODUCTION

Surface-enhanced Raman scattering (SERS) of molecular species on metal island films of Au, and Au films roughened with underlayers of CaF₂, was obtained using the near-infrared excitation line of the Nd:YAG laser at 1064.1 nm. The molecular species were vanadyl phthalocyanine (VOPc) and 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA), which were coated onto the Au surfaces as thin vacuum evaporated layers. An enhancement factor of about 100 was obtained for the electromagnetic enhancement. Demonstration of the infrared SERS effect for the two macrocycle molecules is presented and comparisons are made with previous SERS results obtained with visible light excitation. This is the first reported observation of SERS from metal island films with excitation in the infrared region.

Fourier transform surface-enhanced Raman scattering for a single Langmuir-Blodgett (LB) monolayer on a gold island film and on a copper island film using near-IR excitation at 1064.1 nm is reported here for the first time. FT-SERS was also observed for a thin vacuum evaporated film of vanadyl phthalocyanine on a copper substrate. Similar attempts to obtain FT-SERS from vacuum evaporated VOPc on GaAs and Ga₇Al₃As semiconductor substrates were not successful. The types of LB monolayers used were copper tetra-tertiary-butyl-phthalocyanine (CuTTPc), a mixed monolayer of arachidic acid / asymmetrically substituted diphthalocyanine (PcLuPct), and N-pentyl-N'
ethyl-3,4:9,10-perylenetetracarboxylic diimide (PPTCDE). For comparison, surface-enhanced resonance Raman scattering (SERRS) spectra were obtained using visible laser excitation in resonance with both an electronic absorption of the LB material and the surface plasmons of either the copper or gold metal island films. In all cases, the LB material was physisorbed on the metal with the electromagnetic mechanism leading to FT-SERS. The results illustrate the FT-Raman application to submicron thin solid films.

Fourier transform (FT) Raman spectroscopy is a recent addition to the long list of Raman spectroscopic techniques. The FT-Raman system consisted of a Michelson interferometer and an appropriate detector for inelastic light scattered from a sample excited with a Nd:YAG laser operating at 1064.1 nm. A major advantage of the technique for practical applications is the fact it is "fluorescence-free". The intensity of non-resonant scattered light is very weak for thin solid vacuum evaporated films or Langmuir-Blodgett (LB) monolayers due to the small amount of material present. Very few compounds have absorption in the near-infrared which rules out the use of the resonance Raman (RR) effect to increase the intensity of scattering in this region. One approach to increase the scattering is to deposit thin films on a metal island substrate to obtain Fourier transform surface-enhanced Raman scattering (FT-SERS).

Chapter 5 discusses vacuum evaporated vanadyl phthalocyanine (VOPc) and 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA) films on gold island substrates. The recently published results for VOPc on a copper island film as well as FT-SERS of single Langmuir-Blodgett monolayers of phthalocyanine and perylene derivatives on copper and gold island films appear here also.
was the first of a number of new techniques, encompassed under the general name of surface-enhanced spectroscopy. In the visible region of the spectrum, SERS is now a common analytical technique routinely used, for instance, in biochemistry and electrochemistry. Experiments are generally carried out on rough surfaces with the appropriate dielectric constant and shape, as required by the electromagnetic enhancement; Ag, Au and Cu are the surfaces largely used. Infrared FT-SERS of pyridine on Cu and Au electrodes and 3-picoline and 3-chloropyridine on a copper electrode were demonstrated. Very large enhancement factors of $10^5$-$10^6$ and $9 \times 10^6$, respectively, were estimated. These are two of the very few reports on FT-SERS, and there were no previous reports of FT-SERS on metal island films. Other reports on FT-SERS include investigations of pyridine, ferricyanide and ferrocyanide ions on Cu, Ag and Au surfaces and electrodes, of pyridine on Ag and Au electrodes, of dyes and pyridyls on Au and Cu colloids, and of pyridine and ruthenium complexes on Cu and Au colloids. The electromagnetic model used as a predictive theory for planning infrared SERS experiments, indicated that metal and semiconductor surfaces could be used for Raman enhancement.

There have been cases in the literature of SERS experiments on non-metallic surfaces using visible laser excitation. Roughness-induced SERS appeared for surface carbon on PbTe and was thought to involve interband transitions of the PbTe. SERS has been observed for copper phthalocyanine (CuPc) vacuum evaporated onto GaP particles. Lines due to CuPc and GaP were enhanced, and the enhancement factor depended on the particle size. SERS occurred at a semiconductor-electrolyte interface for I$_2$ adsorbed on a TiO$_2$ electrode. Ueba presented a theory for Raman scattering from an adsorbed molecule on a semiconductor. Excitonic or interband excitations of
the semiconductor can be coupled to the electronic excited states of an adsorbed molecule\textsuperscript{19}. A measurement of the excitation profile of the Raman scattering intensity can be used to determine which mechanism is operating\textsuperscript{19}. In this chapter, a thin layer of VOPc was vacuum evaporated on GaAs and Ga\textsubscript{0.7}Al\textsubscript{0.3}As surfaces in order to explore possible enhancement of the Raman signal with the near-IR excitation frequency.

The three general classes of LB monolayers transferred to FT-SERS substrates included a substituted metal phthalocyanine, a substituted lanthanide diphthalocyanine, and a perylene derivative. The substituted metal phthalocyanine was copper tetra-tert-butylphthalocyanine (CuTTPc). The preparation of a lightly substituted metal-free phthalocyanine has been discussed by Kovacs et al\textsuperscript{20}. Similarly, electrical and structural properties of an LB film of CuTTPc have been reported by Hann et al\textsuperscript{21}. A mixed monolayer of CuTTPc reversibly adsorbs NO\textsubscript{2}/N\textsubscript{2}O\textsubscript{4} and is a candidate for a phthalocyanine-based sensor\textsuperscript{22}.

Lanthanide diphthalocyanines have interesting semiconductive and electrochromic properties\textsuperscript{23-25}. The second type of LB monolayer used for FT-SERS is an asymmetrically substituted lutetium diphthalocyanine. Four tert-butyl groups are present at the 4, 4', 4'', 4''' positions of one of the Pc rings (PcLuPc\textsuperscript{t}). The vibrational characterization of lutetium, ytterbium, dysprosium and holmium diphthalocyanines (LuPc\textsubscript{2}, YbPc\textsubscript{2}, DyPc\textsubscript{2}, HoPc\textsubscript{2}) was reported previously for vacuum evaporated films and LB monolayers\textsuperscript{26,27}. The interaction of diphthalocyanine LB films with NO\textsubscript{2}/N\textsubscript{2}O\textsubscript{4} gas mixtures has also been documented\textsuperscript{27-29}. 
Langmuir-Blodgett monolayers of N-pentyl-N'-ethyl-3,4:9,10-peryleneetetracarboxylic diimide (PPTCDE) and other perylene derivatives were candidates for FT-SERS on metal islands. Akers has reported vibrational spectra of vacuum evaporated perylenetetracarboxylic dianhydride (PTCDA) and perylenetetracarboxylic diimide (PTCDI)\textsuperscript{30,31}. SERS and SERRS results for PTCDA and a diimide were presented in a separate report\textsuperscript{32}. SERS results were also published for LB films of N-octyl-N'-isobutyl-3,4:9,10-peryleneetetracarboxylic diimide\textsuperscript{33}. Electronic and vibrational spectra of LB monolayers of N-octyl-3,4-peryleneedicarboximide (PDCI) appear in the literature\textsuperscript{34}. LB monolayers of N-hexyl-3,4:9,10-peryleneetetracarboxylic monoimide (HPTCO) and N-hexyl-N'-ethyl-3,4:9,10-peryleneetetracarboxylic diimide (HPTCDE) display surface-enhanced fluorescence on silver islands\textsuperscript{35}. Energy transfer has been shown to take place between LB multilayers consisting of perylene derivatives as donors and LuPc\textsubscript{2} as the acceptor\textsuperscript{36}.

5.2 EXPERIMENTAL

5.2.1. Vacuum Evaporation of Metal Island Films and Thin Organic Films.

Evaporated thin films were prepared in a Vacuum Generators DPUHV12 vacuum system. Film thicknesses were monitored using a Kronos Inc. Model QM-311 quartz crystal oscillator. Gold was obtained as fine casting grains of 99.99\% purity from Johnson Matthey and Co. Ltd., and the copper was Balzers of 99.9\% purity with a grain size of 0.2-0.7 mm. The copper and gold films were evaporated at 0.1 nm/s under a pressure of 1.8 \times 10^{-4} \text{ Pa} from a tungsten boat to a thickness of 20 nm onto Corning 7059 glass slides held at 200°C. Dow Corning No. 4 compound, a high-vacuum silicone grease, was applied to the back of the slide to ensure good thermal contact with the substrate heater.
After evaporation of the copper or gold, the vacuum grease was carefully removed from the back of the slide with a swab soaked in methanol.

A sample of a 20 nm gold film evaporated onto Corning 7059 glass held at 200°C was examined without further treatment using a Hitachi Model S-4000 field emission scanning electron microscope (FESEM). An accelerating voltage of 3 kV was used with the specimen at 0° angle of tilt.

A 3 nm film of vanadyl phthalocyanine was vacuum evaporated onto a 20 nm gold film at 0.1 nm/s under a pressure of $3.2 \times 10^{-4}$ Pa from a quartz crucible. The synthesis of the VOPc has already been described in Chapter 4. A 3 nm vacuum evaporated film of VOPc on a 20 nm copper island film was prepared in a similar manner. A 3 nm film of 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA) was vacuum evaporated onto a 20 nm gold film at 0.1 nm/s under a pressure of $3.7 \times 10^{-4}$ Pa from a quartz crucible. A sample of PTCDA was provided by Professor Nagao and the synthesis has been reported.

A substrate with surface roughness was prepared by first depositing calcium fluoride, 20 nm Au and then finally 3 nm VOPc. Optical grade calcium fluoride (Alfa Products) was vacuum evaporated to a 250 nm thickness at 0.3 nm/s at $1.4 \times 10^{-4}$ Pa initial pressure onto Corning 7059 glass slides held at room temperature. A layer of 20 nm Au was vacuum evaporated onto the CaF$_2$ at a rate of 0.1 nm/s under a pressure $3.0 \times 10^{-4}$ Pa with the substrate at room temperature. Finally a top layer of 3 nm of VOPc was vacuum evaporated under a pressure of $2.4 \times 10^{-4}$ Pa again with the substrate at room temperature.
5.2.2. Preparation of Nonmetallic Substrates.

Non-metallic surfaces were prepared for FT-SERS. A gallium arsenide (GaAs) substrate of {100} orientation from Sumitomo Electric Industries Ltd. was used with a surface finish consisting of both sides lapped followed by one side polished, the other etched. A 3 nm film of VOPc was vacuum evaporated at a rate of 0.1 nm/s under a pressure of $2 \times 10^{-4}$ Pa from a quartz crucible onto each side of the GaAs separately. A second FT-SERS non-metallic surface consisted of a similar GaAs substrate as above with metallorganic chemical vapor deposited (MOCVD) epitaxial layers on top. The subsequent layers were a 1.7 µm thick GaAs buffer layer followed by a 4.65 µm thick Ga$_7$Al$_3$As layer. A 3 nm film of VOPc was vacuum evaporated on the top Ga$_7$Al$_3$As layer with conditions as stated previously.

5.2.3. Preparation of Langmuir-Blodgett Films.

The copper tetra-tertiary-butylphthalocyanine (CuTTPc) was kindly provided by Dr. Tomilova from the Organic Intermediates and Dyes Institute of the Academy of Sciences of the USSR in Moscow. A solution of $4.40 \times 10^{-4}$ M CuTTPc in toluene (Eastman Kodak ACS Spectro grade) was prepared and 100 µL spread onto the subphase of a Fromherz circular type Langmuir-Blodgett trough$^{38}$. For the subphase Milli-Q water with one additional distillation was used at ambient temperature ($\sim 23^\circ$C) and without further treatment (pH $\sim 6.3$). A Fromherz trough and dipper, housed in a glove box on a terrazzo vibration-free table were used for film compression and transfer to substrates. The monolayer was compressed at $4 \times 10^{-3}$ nm$^2$/molecule/s and the film was transferred at 6 mm/min onto the gold film on Corning 7059 glass by Z-type deposition at a constant pressure of 10 mN/m.
The synthesis of an asymmetrically substituted bisphthalocyanine PcLuPc has been reported\textsuperscript{39}. The LB film solution consisted of a 1:10 molar ratio (PcLuPc:arachidic acid) in chloroform. It was necessary to add arachidic acid to obtain good transfer to the substrate. The subphase was maintained at 15°C and transfer was carried out at 3 mm/min with a constant pressure of 15 mN/m to the 20 nm thick gold and copper island films on Corning 7059 glass by Z type deposition. These LB films were prepared in a Lauda Langmuir film balance equipped with the Lauda Film lift FL-1 electronically controlled dipping device.

The \textit{N}-penty1-\textit{N}'-ethyl-3,4:9,10-perylenetetracarboxylic diimide (PPTCDE) was synthesized and purified by Professor Nagao\textsuperscript{37}. The PPTCDE solution was 3.6 x 10\textsuperscript{-5} M in chloroform and 4 mL were spread on the Fromherz trough. The PPTCDE was compressed at a rate of 1 X 10\textsuperscript{-3} nm\textsuperscript{2}/molecule/second and transferred to metal island substrates at 35 mN/m constant pressure. The slide was removed from the trough at 3.6 mm/min and gave Z type deposition.

5.2.4 Raman Systems and UV-visible Spectrophotometer.

A Bomem Ramspec 150 spectrophotometer with a Nd/YAG laser emitting at 1064.1 nm and an InGaAs detector were used to acquire the FT-Raman spectra. The system had a Michelson interferometer and was interfaced to a NEC Powermate 286 Plus computer. Bomem Easy (version 1.45) software was used to collect the data and files were imported to Spectra Calc\textsuperscript{TM} (Galactic Industries Corp.). All Raman files were ratioed against an instrument response curve. A backscattering geometry was used with resolution set at 4 cm\textsuperscript{-1}.

The incident power of the Nd/YAG laser onto the samples were (a) 1.82 W onto the 200 nm VOPc film, (b) 3.5 W onto the 200 nm PTCDA film, (c) 0.61 W
onto the 3 nm PTCDA on 20 nm Au (200°C) film, 0.290 W onto the 3 nm VOPc on 20 nm Au (200°C), 0.290 W onto the 3 nm VOPc on glass film, (f) 0.117 W onto the 3 nm VOPc on 20 nm Au on 250 nm CaF₂ film, (g) 0.070 W onto the 3 nm VOPc on 20 nm Cu (200°C), (h) 0.440 W onto the single LB monolayer of CuTTPc on 20 nm Au (200°C), (i) 0.017 W onto the mixed LB monolayer of 1:10 PcLuPc / arachidic acid on 20 nm Au (200°C), and (j) 0.343 W onto the single LB monolayer of PPTCDE on 20 nm Au (200°C). Typically 525 scans were averaged, requiring 50 minutes to collect the data.

Samples were compared where possible by putting single and multiple layers all on the same slide. For example, on one slide there was a region with VOPc on Au and another region of VOPc on glass only. The slide was aligned with the laser beam focussed onto the VOPc on Au region, and a spectrum was collected. Next, the slide was translated horizontally so the laser beam focussed on the VOPc on glass only region, and scans were collected. It was necessary to collect 2500 scans from a 3 nm VOPc film on glass to acquire the signal needed to calculate the SERS enhancement factor. To calculate the enhancement factors, the signal-to-noise ratios (S/N) were calculated as the square roots of the number of scans both for the VOPc film on glass and for the VOPc film on the roughened Au substrate. The enhancement factor is then given as the ratio of the band heights corrected for the S/N ratios and for the incident laser powers.

A Spectra Physics model 2020 Kr+ laser provided the 647.1 nm laser line with 50 mW of s-polarized light at the sample. A rotating sample cell was used to prevent laser damage to LB films. The typical spectral band pass was 4 cm⁻¹ and Raman shifts were measured with a Spex-1403 double monochromator.
The 514.5 nm line of a Spectra Physics Ar+ laser was used for the PPTCDE compound on gold.

Electronic spectra were measured in the region 200-1100 nm using a Shimadzu UV-160 spectrophotometer with a glass slide as a reference.

5.3 RESULTS AND DISCUSSION

5.3.1. SERS-active Surfaces for the Near-IR.

It is widely accepted that an electromagnetic enhancement of the Raman signal may be achieved on rough surfaces of certain metals and semiconductors\(^{10}\). The role of the surface roughness is to couple the incident field of the exciting radiation to the surface plasmons, enhancing the local optical field and efficiency of light scattering, absorption, and fluorescence. Experiments have shown that signal enhancement of 3 and 4 orders of magnitude could be obtained from this effect in the visible region\(^{40}\). Calculations of enhancement factors also give an optimum value of \(~10^4\) for the average enhancement on Ag\(^{41}\). Metal island films used for experiments with visible light excitation were approximated as spheroidal particles, and those with dimensions in the range 10-100 nm produced the strongest enhancement\(^{42}\).

In the quasi-static limit, \(r/\lambda < < 1\), where \(r\) is the radius of a metal sphere, only the dipole surface modes are excited by the plane-wave field, and the enhancement of the field is due to the accumulation of electromagnetic energy in the local plasma oscillations. For most metals used in SERS the maximum enhancement lies in the visible region. Using the same metals (Ag, Au and Cu), the design of a SERS active metal surface for the infrared region
requires an increase in the particle size in order to tune the film plasmon frequency to the exciting laser line. The question arises as to the effect of particle size and particle interactions on the enhancing properties of the film. In general, increasing the particle size and packing of metal particles would also increase the damping of local plasma oscillations, inhibiting the enhancing ability of the film. The latter discussion implies that only a very modest enhancement should be expected from metal island films of Ag, Cu, and Au evaporated onto glass substrates (held at an elevated temperature of ~200°C) with a mass thickness sufficient to provide a measurable value of light absorption in the near-infrared region.

Gold has a dielectric constant value that gives plasmon absorption in the red region of the visible spectrum for a mass thickness of only 4 nm. Therefore, metal island films of Au of 15-25 nm mass thickness give considerable absorption in the 1000 nm region as shown in Figure 5-1. Au films deposited onto glass substrates held at 200°C produced an enhanced Raman signal. An electron micrograph of the 20 nm Au film is shown in Figure 5-2. The rough Au surface contains metal islands that are on average larger than 100 nm, with a high packing density.

SERS spectra were obtained with several Au island films, but the 20 nm film produced the largest enhancement factor. Copper films were also deposited onto glass substrates held at 200°C and the best enhancement was observed with 20 nm mass thickness films. In an attempt to increase the enhancement factor observed for FT-SERS from Au, 20 nm of Au was evaporated onto CaF₂ underlayers. This surface improved the enhancement by a factor of 2 over that of the evaporated Au island film.
Figure 5-1: Absorption spectrum of a 20 nm Au island film on glass.
Figure 5-2: Field emission scanning electron micrograph of a 20 nm Au island film on glass.
The island films of Au consist largely of oblate islands. However, it is known from previous experimental results that CaF$_2$ underlayers are very jagged structures which provide a more prolate character to the bumps (roughness) of the covering metal film. Murray and Allara used two different sample configurations consisting of Si/CaF$_2$/Al+oxide/Raman scatterer molecule/optional polymer spacer/Ag and Si/Al+oxide/Raman scatterer molecule/optional polymer spacer/Ag, for visible SERS experiments. Our sample configuration consisted of Corning 7059 glass/CaF$_2$/Au/organic film. FT-SERS was obtained using near-IR laser excitation at 1064.1 nm.

5.3.2 UV-Vis absorption spectra of VOPc and PTCDA. FT-SERS on gold island films.

The molecular structure of VOPc appeared in Figure 4-1. The structure of 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA) is given in Figure 5-3. Absorption spectra of thin solid films of the molecular pigments used in the SERS experiments are given in Figures 5-4 and 5-5. It can be seen that there was no molecular electronic absorption beyond 850 nm; therefore, the FT-Raman results correspond to spontaneous Raman spectra. Both VOPc and PTCDA have been characterized spectroscopically, and for which SERS and (SERRS) visible spectra were previously reported. The FT-SERS spectrum of a 3 nm VOPc film on a 20 nm Au island film is given in Figure 5-6, where for comparison the FT-Raman spectrum of a 200 nm solid film of VOPc is included. An enhancement factor (EF) of ~50 was estimated by comparison of the FT-SERS signal with the signal obtained from the reference film. The EF found for the 20 nm Au film on CaF$_2$ was ~100. The FT-SERS spectrum is the enhanced version of the FT-Raman spectrum of the film, and there are no frequency shifts or noticeable intensity changes.
Figure 5-3: Molecular structure of 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA).
Figure 5-4: Electronic absorption spectrum of a 100 nm evaporated film of VOPc on glass.
Figure 5-5: Electronic absorption spectrum of a 100 nm evaporated film of PTCDA on glass.
The FT-Raman and FT-SERS spectra for PTCDA followed closely the relative intensity patterns and frequency changes observed for SERRS and SERS in the visible region. The FT-Raman spectrum of a 200 nm thin solid film of PTCDA is shown in Figure 5-7 along with an FT-SERS spectrum of a 3 nm film of PTCDA on a 20 nm Au island film. The frequencies observed in the FT-Raman spectrum of PTCDA include 1052 cm\(^{-1}\), 1307 cm\(^{-1}\), 1384 cm\(^{-1}\), 1574 cm\(^{-1}\) and 1592 cm\(^{-1}\). In the same region, the frequencies observed in the FT-SERS are 1293 cm\(^{-1}\), 1376 cm\(^{-1}\), 1569 cm\(^{-1}\) and 1578 cm\(^{-1}\).

5.3.3 Comparison of Near-Infrared FT-SERS and Visible SERS for VOPc and PTCDA on gold films.

Vanadyl phthalocyanine has strong absorptions in the visible region as can be seen in Figure 5-4. However, there is a transparent window between 400 and 550 nm, and excitation with the 514.5 nm laser line was considered to give SERS, while excitation with the 647.1 nm line gave surface-enhanced resonant Raman scattering (SERRS). Comparing the SERS spectra obtained on Ag-coated Sn spheres with the FT-SERS results may help to understand the influence of metal-molecule interaction and/or molecular orientation on the observed frequencies and relative intensity patterns. The symmetries of the fundamental isoindole stretching vibrations of VOPc have been studied using polarized Raman spectroscopy of submicron thin films on smooth gold surfaces. In general, there is good agreement between the two spectra, and the most intense Raman frequencies observed in SERS at 1521, 1338, 1001, 836, 678, 590 and 484 cm\(^{-1}\) are also intense in the FT-SERS spectrum shown in Figure 5-6. It can be concluded that both spectra correspond to molecular species physisorbed onto the Ag and Au surfaces, respectively. The FT-Raman and visible RS are similar in one other important behavior. The thick films (bulk) of
Figure 5-6: (a) FT-Raman spectrum of 200 nm film of VOPc and (b) FT-SERS of a 3 nm film of VOPc on 20 nm Au island film.
Figure 5-7: (a) FT-Raman spectrum of 200 nm PTCDA and (b) FT-SERS of 3 nm film of PTCDA on a 20 nm Au island film.
VOPc show a strong band at 1343 cm\(^{-1}\) in both the FT-Raman and visible Raman scattering (RS). However, the SS spectrum\(^{46}\) (s-polarized light incident and s-polarized light collected) of the visible RS of a VOPc film showed a very intense band at 1337 cm\(^{-1}\), which was the only one observed in SERS in this wavenumber region. It was assumed\(^{44}\) that the doublet (1343-1337 cm\(^{-1}\)) was due to correlation field splitting. Identical results are obtained in the FT-Raman experiments: a band at 1343 cm\(^{-1}\) is observed in bulk with a side band at 1338 cm\(^{-1}\), but the FT-SERS shows only one band at 1338 cm\(^{-1}\).

The SERS of PTCDA in the visible region was shown to be different from the Raman spectrum of a solid sample, which was similar to both the corresponding resonant Raman spectrum (RRS) and SERRS\(^{32}\). For example, a doublet was observed in the RRS of the thin solid film at 1574 and 1592 cm\(^{-1}\), which was also seen in the RS and SERRS, and was seen as well in the FT-Raman spectrum. Indeed, the RRS and SERRS (488 nm excitation) and the spontaneous RS obtained with the 1064.1 nm line are similar not only in frequency values but also in relative intensities. In the PTCDA SERS report\(^{32}\), it was shown that the 1574 cm\(^{-1}\) line was preferentially enhanced in the SERS spectrum excited with the 647.1 nm laser line on Ag-coated Sn spheres, while both frequencies were seen in the SERRS spectrum (488 nm excitation on Ag-coated Sn spheres). The FT-SERS results are consistent with previous findings, and of the doublet (1574 and 1592 cm\(^{-1}\)) observed in the FT-Raman spectrum shown in Figure 5-7, only the 1574 cm\(^{-1}\) band is preferentially enhanced in the FT-S\(\text{\textregistered}\)RS. The preferentially enhanced band at 1574 cm\(^{-1}\) is, in turn, split into a doublet at 1569 and 1578 cm\(^{-1}\). Also in agreement with the visible RS and SERS is the shift of the band centered at 1307 cm\(^{-1}\) in the FT-Raman spectrum of the thin solid film to 1293 cm\(^{-1}\) in the FT-SERS. In summary, FT-SERS spectra observed on gold
for both molecular systems are consistent with the previous observations of SERS in the visible spectral region, and unique intensity patterns found in the SERS spectra are also observed in the FT-SERS.

5.3.4 FT-SERS of Thin Vacuum Evaporated VOPc Films on Cu Islands and Semiconductors

FT-SERS was obtained for a 3 nm vacuum evaporated film of vanadyl phthalocyanine on a 20 nm copper island film, the Raman spectra are shown in Figure 5-8. Vanadyl phthalocyanine does not absorb significantly at the 1064.1 nm Nd/YAG laser line but does absorb strongly at the 647.1 nm line of the Kr+ laser. The 20 nm copper island film prepared at 200°C displayed a broad absorption spectrum with some absorption at both 647.1 and 1064.1 nm. The spontaneous Raman scattering of a 200 nm thick VOPc film evaporated at room temperature onto glass is included in Figure 5-8 (a) as a reference. Figure 5-8 (b) shows surface-enhanced resonance Raman scattering of a VOPc film on a 20 nm copper island film. Electromagnetic SERS is combined with the resonance Raman effect. Figure 8-5 (c) displays the FT-SERS spectrum of a 3 nm VOPc film on copper islands. The frequency values are practically the same for the FT-Raman of thick VOPc (top) and the FT-SERS (bottom). This indicates that the VOPc is physisorbed rather than chemisorbed onto the metal surface.

Table 5-1 gives the assignments for the spontaneous Raman scattering of VOPc, SERRS of VOPc on Cu, and FT-SERS of VOPc on Cu. The isoindole and benzene radial deformations occur at 484 cm⁻¹ and 591 cm⁻¹ respectively. A macrocycle can be defined by the sixteen inner alternating carbon and nitrogen atoms. Macrocycle breathing and stretching modes are observed in the 650 cm⁻¹ to 1000 cm⁻¹ region while carbon-hydrogen bending modes are
Figure 5-8: (a) FT-Raman spectrum of 200 nm VOPc on glass. Laser line 1064.1 nm. (b) SERRS spectrum of 3 nm VOPc on 20 nm Cu. Laser line 647.1 nm. (c) FT-SERS spectrum of 3 nm VOPc on 20 nm Cu. Laser line 1064.1 nm.
Figure 5-8

(a) FT-Raman of 200 nm VOPc film

(b) SERS of 3 nm VOPc on 20 nm Cu

(c) FT-SERS of 3 nm VOPc on 20 nm Cu

WAVENUMBERS (cm\(^{-1}\))

500 1000 1500

INTENSITY (arbitrary units)
seen from 1000 cm\(^{-1}\) to 1300 cm\(^{-1}\). The resonance Raman scattering of phthalocyanine has contributions mainly from the Herzberg-Teller term. Excitation at 647.1 nm enhances both totally and non-totally symmetric vibrations of VOPc. The SERRS spectrum in Figure 5-8 (b) has contributions from both the resonance Raman effect and surface-enhanced scattering.

SERRS and FT-SERS for 3 nm VOPc on copper islands are remarkably similar in relative intensities as can be seen in Figure 5-8 (b) & (c). The observation is simply a consequence of comparable relative intensities found in the correlation of the resonant Raman spectrum (laser frequency in resonance with the Q-band of the VOPc) with the spontaneous FT-Raman spectrum. The SERS spectrum of this particular molecule seems to be reduced to the original spectrum (RRS or FT-Raman) times the enhancement factor, a textbook case of physisorption. FT-SERS was not observed for a 3 nm thick film of VOPc on either GaAs or Ga\(_{0.7}\)Al\(_{0.3}\)As semiconductor substrates.

5.3.5 **FT-SERS of Single LB Monolayers on Copper and Gold Island Films.**

FT-SERS is reported here for the first time for a single Langmuir-Blodgett monolayer on vacuum evaporated metal island films. Figure 5-9 shows the FT-SERS spectrum and SERRS spectrum of a single LB monolayer of copper tetra-tertiary-butyl-phthalocyanine (CuTTPc) on a 20 nm gold film. Molecular modeling by Hahn et al. suggested that the bulky tertiary-butyl groups of CuTTPc are interleaved with neighboring molecules on water and transmission electron micrographs of CuTTPc on supported grids indicated the molecules form columns in the plane of the film with 1.9 nm between columns and an in-
Table 5-1: Raman Frequencies and Intensities for vacuum evaporated VOPc and VOPc on Cu

<table>
<thead>
<tr>
<th>200 nm VOPc LL=1064 nm (RS) Wavenumbers (relative intensity)</th>
<th>3 nm VOPc on 20 nm Cu (200°C) LL=647nm (SERS) Wavenumbers (relative intensity)</th>
<th>3 nm VOPc on 20 nm Cu (200°C) LL=1064 nm (FT-SERS) Wavenumbers (relative intensity)</th>
<th>Assignments</th>
</tr>
</thead>
<tbody>
<tr>
<td>485 cm(^{-1}) (44)</td>
<td>484 cm(^{-1}) (26)</td>
<td>484 cm(^{-1}) (36)</td>
<td>isoindole ring def.</td>
</tr>
<tr>
<td>592 cm(^{-1}) (24)</td>
<td>591 cm(^{-1}) (19)</td>
<td>591 cm(^{-1}) (18)</td>
<td>benzene radial</td>
</tr>
<tr>
<td>680 cm(^{-1}) (79)</td>
<td>679 cm(^{-1}) (91)</td>
<td>680 cm(^{-1}) (68)</td>
<td>macrocycle breathing</td>
</tr>
<tr>
<td>753 cm(^{-1}) (50)</td>
<td>751 cm(^{-1}) (22)</td>
<td>752 cm(^{-1}) (38)</td>
<td>benzene (v(_1))</td>
</tr>
<tr>
<td>787 cm(^{-1}) (18)</td>
<td>---</td>
<td>787 cm(^{-1}) (13)</td>
<td>macrocycle stretching</td>
</tr>
<tr>
<td>836 cm(^{-1}) (10)</td>
<td>835 cm(^{-1}) (29)</td>
<td>836 cm(^{-1}) (18)</td>
<td>macrocycle stretching</td>
</tr>
<tr>
<td>947 cm(^{-1}) (15)</td>
<td>946 cm(^{-1}) (29)</td>
<td>947 cm(^{-1}) (11)</td>
<td>macrocycle stretching</td>
</tr>
<tr>
<td>1008 cm(^{-1}) (7)</td>
<td>1007 cm(^{-1}) (18)</td>
<td>1010 cm(^{-1}) (9)</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1032 cm(^{-1}) (7)</td>
<td>---</td>
<td>---</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1107 cm(^{-1}) (22)</td>
<td>1107 cm(^{-1}) (18)</td>
<td>1106 cm(^{-1}) (16)</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1143 cm(^{-1}) (19)</td>
<td>1143 cm(^{-1}) (26)</td>
<td>1142 cm(^{-1}) (21)</td>
<td>pyrrole ring breathing</td>
</tr>
<tr>
<td>1162 cm(^{-1}) (6)</td>
<td>---</td>
<td>1162 cm(^{-1}) (9)</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1187 cm(^{-1}) (22)</td>
<td>---</td>
<td>1186 cm(^{-1}) (18)</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1204 cm(^{-1}) (22)</td>
<td>1196 cm(^{-1}) (20)</td>
<td>1194 cm(^{-1}) sh.</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1306 cm(^{-1}) (15)</td>
<td>1304 cm(^{-1}) (27)</td>
<td>1306 cm(^{-1}) sh.</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1304 cm(^{-1}) (15)</td>
<td>---</td>
<td>1321 cm(^{-1}) sh.</td>
<td>C-H bend</td>
</tr>
<tr>
<td>1341 cm(^{-1}) (44)</td>
<td>1342 cm(^{-1}) (75)</td>
<td>1340 cm(^{-1}) (59)</td>
<td>pyrrole ring stretch</td>
</tr>
<tr>
<td>1450 cm(^{-1}) (18)</td>
<td>1446 cm(^{-1}) (41)</td>
<td>1450 cm(^{-1}) (20)</td>
<td>isoindole ring stretch</td>
</tr>
<tr>
<td>1522 cm(^{-1}) (100)</td>
<td>1524 cm(^{-1}) (100)</td>
<td>1523 cm(^{-1}) (100)</td>
<td>C-N pyrrole stretch</td>
</tr>
</tbody>
</table>
stack distance of 0.33 nm. Excitation at 647.1 nm is in resonance with an electronic absorption of the CuTTPc molecule and gives surface-enhanced resonance Raman scattering on the gold island surface. The 1064.1 nm Nd/YAG laser line is in resonance with the plasmons in the gold film only and gives rise to FT-SERS. Relative intensities in the region of the macrocycle breathing vibration (691 cm\(^{-1}\)) are different due to the specific intensity enhancement of the chromophore vibrations by the resonance Raman effect. The band at 1332 cm\(^{-1}\) in SERRS is centered at 1330 cm\(^{-1}\) in FT-SERS and similarly the 1532 cm\(^{-1}\) band in SERRS is measured at 1526 cm\(^{-1}\) in FT-SERS. The latter differences could be due to the contribution of two fundamentals to the envelope of the observed Raman bands in the FT-SERS. One of the components could be selectively enhanced by the resonance Raman effect. The band near 1330 cm\(^{-1}\) is a pyrrole ring stretch while C-N pyrrole stretching and \(C_\beta-C_\beta\) pyrrole stretching account for the bands at 1526 cm\(^{-1}\) and 1532 cm\(^{-1}\).

The FT-SERS spectrum of a mixed monolayer (1:10) of the asymmetrically substituted diphthalocyanine PcluPc and arachidic acid on a 20 nm film are presented in Figure 5-10 (a). Surface-enhanced resonance Raman results for the mixed monolayer on copper and gold island films appear in Figures 5-10 (b) and 5-10 (c). The purpose of the arachidic acid was to act as a lubricant on the water and improve the transfer to the substrate. A 20 nm gold island film absorbs light more strongly at the 647.1 nm laser line than the 20 nm copper island film. The observed frequencies for the mixed LB monolayer on gold and copper are similar. The FT-SERS signals were very weak since the PcluPc is diluted in arachidic acid. Considering the FT-SERS results first, the 560 cm\(^{-1}\), 760 cm\(^{-1}\) and 913 cm\(^{-1}\) peaks can be assigned respectively to an isoindole deformation, and two macrocycle stretching modes. It was interesting to note
Figure 5-9: (a) FT-Raman spectrum of a single LB monolayer of CuTTPc on 20 nm Au. Laser line 1064.1 nm. (b) SERRS spectrum of a single LB monolayer of CuTTPc on 20 nm Au. Laser line 647.1 nm.
a) FT-SERS of single LB monolayer of CuTTPc on 20nm gold

b) SERRS of single LB monolayer of CuTTPc on 20nm gold

INTENSITY (arbitrary units)

WAVENUMBERS (cm$^{-1}$)

LL = 1064 nm

LL = 647 nm
that there were no Raman bands from the arachidic acid. The most intense band in the SERRS spectra at 681 cm\(^{-1}\) arises from macrocycle breathing and macrocycle stretches appear at 744 cm\(^{-1}\) and 819 cm\(^{-1}\). In the low frequency region, an isoindole deformation was observed at 479 cm\(^{-1}\) with the benzene radial deformation at 577 cm\(^{-1}\). The PCluPct is physisorbed because known frequencies of diphthalocyanines were observed and not shifted as expected for chemisorption. There is a specific intensity enhancement of chromophore vibrations by the resonant Raman effect in Figures 5-10 (b) and 5-10 (c) which explains the changes in relative intensities between FT-SERS and SERRS.

The FT-SERS and SERRS spectra of a single LB monolayer of \(N\)-pentylyl-\(N'\)-ethyl-3,4:9,10-perylenetetracarboxylic diimide (PPTCDE) on a 20 nm gold island film are shown in Figure 5-11. The PPTCDE does not have any absorbance in the near-IR at 1064.1 nm while the gold island film absorbs in this region allowing surface-enhanced Raman scattering. The 514.5 nm laser line is in resonance with an electronic absorption in the perylene and the 20 nm gold island film also absorbs at this wavelength producing SERRS. An upper limit of (ca. 10\(^3\)) for the enhancement of a resonance Raman signal has been suggested\(^{47}\). Earlier work\(^{48}\) on the perylene chromophore reported a SERRS enhancement factor of at least 10\(^4\). The resonance Raman in tetracarboxylic derivatives is due mainly to Franck-Condon scattering and overtones have been observed\(^{48}\).

The \(a_{1g}\) modes for perylene have been reported\(^{49}\) at 550, 1104, 1298, 1369, 1380, 1574 and 1581 cm\(^{-1}\). The FT-SERS frequencies of PPTCDE on gold were 541, 1073, 1292, 1375, 1450 and 1576 cm\(^{-1}\). The 541 cm\(^{-1}\) and 1073 cm\(^{-1}\) bands correspond to a benzene ring deformation and benzene breathing
Figure 5-10: A mixed monolayer of 1:10 PcuPc / arachidic acid. (a) FT-Raman spectrum on 20 nm Au. Laser line 1064.1 nm. (b) SERRS spectrum on 20 nm Au. Laser line 647.1 nm. (c) SERRS spectrum on 20 nm Cu. Laser line 647.1 nm.
Figure 5-10

(a) FT-Raman of single mixed LB on 20nm Au
(b) SERRS of single LB on 20nm Au
(c) SERRS of single mixed LB on 20nm Cu

Wavenumbers (cm⁻¹)

Intensity (arbitrary units)
Figure 5-11:  
(a) FT-SERS spectrum of a single LB monolayer of PPTCDE on 20 nm Au. Laser line 1064.1 nm. (b) SERRS spectrum of a single LB monolayer of PPTCDE on 20 nm Au. Laser line 514.5 nm.
modes respectively. Characteristic group frequencies for the perylene moiety include 1375 cm$^{-1}$ and 1576 cm$^{-1}$ vibrations which involve stretches of the naphthalene group$^{49}$. There are two different naphthalene groups in PPTCDE due to asymmetric alkyl substitution and shoulders appear on the sides of these naphthalene bands. The peak at 1292 cm$^{-1}$ is also a vibration of the parent perylene moiety$^{50}$. The PPTCDE LB monolayer is physisorbed on the gold island film since frequency shifts remain unchanged from PPTCDE alone. In general, a weak continuum inelastic scattering background appeared towards higher wavenumbers for all films which showed FT-SERS which is due to the metal film. The resonance Raman contribution to the SERRS scattering for PPTCDE selectively enhances vibrations of different symmetry type than those seen in the FT-SERS, differences seen in Figure 5-11.
5.4 CONCLUSIONS

FT-SERS on copper and gold island films is reported for the first time. Observed enhancement factors are below $10^3$, an upper limit that would be predicted from the electromagnetic model for metal films with a high density of large metal islands on a glass surface. Recorded FT-SERS spectral frequencies and relative intensities for both VOPc and PTCDA are in agreement with previous SERS findings and support a model of physisorbed molecules on the Au surface. FT-SERS was also observed for a 3 nm vacuum evaporated film of VOPc on a 20 nm copper island film.

FT-SERS is reported for the first time for single LB monolayers on metal island films. The CuTTPc, PcLuPct, and PPTCDE LB monolayers were physisorbed to the metals and the electromagnetic mechanism caused the surface-enhanced Raman scattering. The resonance Raman effect has been shown to selectively enhance chromophore vibrations in SERRS spectra. Gold island films absorb more in the red than copper island films. This results in a greater efficiency of surface plasmon generation which in turn results in greater SERRS enhancement. FT-SERS is a viable technique for obtaining more intense signals for the study Langmuir-Blodgett films or thin vacuum evaporated films in the near-IR.
Chapter 5 References


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CHAPTER 6: SPECTROSCOPY AND ATOMIC FORCE MICROSCOPY OF LANGMUIR-BLODGETT MONOLAYERS

6.1. INTRODUCTION

Thin solid films of perylenes and phthalocyanines have been prepared using the Langmuir-Blodgett (LB) technique. The orientation of LB monolayers of a series of perylenes on metal surfaces has been determined using reflection-absorption infrared spectroscopy. The atomic force microscopy (AFM) instrument was used to study LB monolayers on atomically flat substrates such as mica and silicon wafers. The observed thickness of a monolayer of N-hexyl-N'-ethyl-3,4:9,10-perylenetetracarboxylic diimide (HPTCDE) was 1.83 nm by AFM which agrees with the orientation calculated from the reflection-absorption IR measurements. The importance of AFM as an analytical method for characterization of LB film structures and defects is demonstrated. In addition to HPTCDE, AFM results are also included for LB monolayers of N-hexyl-3,4:9,10-perylenetetracarboxylic monoimide (HPTCO); 1,4,8,11,15, 18,22,25-octabutoxy-29H,31H-phthalocyanine ((BuO)8H2Pc) and tetra-tertiary-butyl metal-free phthalocyanine, (t-bu)4H2Pc.

Thin solid films which differ from bulk material, can be prepared by the Langmuir-Blodgett technique. A single LB monolayer has two interfaces which affect its structure, the first being the substrate/monolayer interface and the second is the monolayer/air interface. The orientation of the monolayer on the substrate is influenced by whether the substrate is hydrophobic or hydrophilic and the manner in which transfer to the substrate is completed. The monolayer is either physisorbed or chemisorbed to the substrate and
intermolecular forces between the molecules in the monolayer play an important role in determining structure. The most thermodynamically stable form of a monolayer may not even be a uniform film.

Mixed monolayers consist of at least two components within a given layer. Segregation or formation of domains could be observed rather than a homogeneous mixing of materials. It is possible to build up a number of successive LB monolayers giving highly organized structures in three dimensions. Strong intermolecular forces between adjacent monolayers is a stabilizing feature and leads to rearrangement after transfer in stearic acid LB films\textsuperscript{3,4}. Two techniques to experimentally explore LB film structure are infrared spectroscopy and atomic force microscopy (AFM).

Reflection-absorption infrared spectroscopy (RAIRS) using different polarizations of light provides information on the orientation of molecules in an LB monolayer. In fact, molecular orientation in Langmuir-Blodgett multilayers can be determined from the combined infrared spectral data obtained in a transmission geometry and in a reflection-absorption geometry. The approach has been illustrated by Umemura et al.\textsuperscript{5a} using LB monolayers of cadmium stearate. The quantity of interest that can be evaluated with this method is the tilt angle formed between the direction of a dynamic dipole moment and the normal to the surface of the film. The information so obtained corresponds to an average molecular orientation of molecules on the surface.

If molecular resolution was possible with local probe methods such as AFM, specific molecular information in LB films, rather than data which is
averaged over a large sample region as is the case with infrared spectroscopy, would be found. An AFM instrument can be used to study surface coverage, local defects such as pinholes, cracks, grain boundaries and heterogeneous structures in mixed LB films. Scanning tunneling microscopy (STM) typically requires that the sample under investigation be conductive while AFM can be applied to conductive and non-conductive materials. The force exerted by the tip of the AFM is on the order of $10^{-9}$ N, and is preferred for imaging thin organic films such as LB monolayers to minimize sample damage. It is now known that the force on an organic monolayer can be reduced by imaging under water in a liquid cell. The AFM also has advantages compared to electron microscopy because samples can be imaged directly. For scanning electron microscopy, a conductive overcoat is applied to non-conductive films or a replica is made. A problem with transmission electron microscopy is low contrast for a single LB monolayer and the need to transfer the monolayer to a microscope grid.

In this chapter, atomic force microscopy was used to characterize single LB monolayers of perylenes and phthalocyanine derivatives on mica and silicon wafers. The pigment structures appear in Figure 6-1.

There have been STM reports in the literature concerning vacuum evaporated phthalocyanine films. Copper phthalocyanine was sublimed onto Cu(100) at room temperature and high resolution images were obtained for submonolayer coverages in an ultrahigh vacuum (UHV) STM system. The CuPc molecules were flat on the substrate with two different molecular orientations. There was excellent agreement between the model of the charge
Figure 6-1: Structures of Langmuir-Blodgett materials.
(a) $N$-hexyl-$N'$-ethyl-3,4:9,10-perylenetetracarboxylic diimide (HPTCDE).
(b) $N$-hexyl-3,4:9,10-perylenetetracarboxylic monoimide (HPTCO).
(c) 1,4,8,11,15,18,22,25-octabutoxy-29H,31H-phthalocyanine ((BuO)$_8$H$_2$Pc).
(d) tetra-tertiary-butyl metal-free phthalocyanine, ((t-bu)$_4$H$_2$Pc).
densities of the highest occupied molecular orbitals (HOMOs) and the experimental data. Lead phthalocyanine films $^8$ were vacuum evaporated onto a substrate of silicon wafer-SiO$_x$-Au composition with imaging in air. Different polymorphic forms of PbPc existed depending on the substrate temperature. Crystallites of 100 nm size were observed as well as 15 nm substructures but no molecular resolution was possible. Scanning tunneling microscopy results were presented for vanadyl naphthalocyanine $^9$ (VONc) on highly oriented pyrolytic graphite (HOPG) and zinc naphthalocyanine (ZnNc) on MoS$_2$ substrate $^{10}$, both imaged under ambient conditions. Molecular resolution was obtained of VONc and indicated a slipped stack orientation with the oxygen atom pointing towards the HOPG. There was a lattice matching of the VONc with the graphite substrate. In the case of ZnNc, the molecules were stacked and canted with respect to the MoS$_2$ substrate. In general, it is important to be able to determine the polymorphic form and orientation of phthalocyanine molecules on substrates because this influences the optical and electronic properties. The stacking and molecular orientation is temperature and substrate dependent.

Literature about AFM or STM of Langmuir-Blodgett monolayers of phthalocyanines has been scarce $^{11}$. A derivatized phthalocyanato-poly-siloxane was transferred to HOPG and gold by the LB technique, then imaged by STM. The polymer rods were parallel to the substrate and aligned in the dipping direction. However, fine structure was not observed and movement was apparent. Porphyrins are biologically important, large organic molecules with some similarities to phthalocyanines. Porphyrin-based molecules with pendant isocyano groups were reacted with epitaxially grown Au(111) on mica and imaged in air using STM $^{12}$. The porphyrin molecules were anchored by covalent bonding to the substrate.
The pigment 3,4:9,10-perylenetetracarboxylic dianhydride (PTCDA) was investigated on graphite using video STM$^{13}$. The PTCDA was vacuum evaporated very slowly and formed a monolayer of flat molecules. Initially the PTCDA was coincident with the graphite but approached bulk α-PTCDA as the thickness was increased. High resolution images displayed a herringbone structure with two molecules per unit cell. Crystals of a perylene radical cation salt were prepared electrochemically and were imaged using STM and AFM$^{14}$.

There have been numerous AFM and STM experiments performed on fatty acid LB films$^{15-20}$. STM images of behenic acid on HOPG showed two types of molecular arrangements parallel to the substrate$^{16}$. The type II form, with a 6 nm periodicity indicated a rearrangement had taken place on the HOPG following LB deposition. Reorganization and crystallite formation was seen by AFM for LB films of cadmium arachidate on substrates that had been submerged under water for different times$^{18}$. As the submersion time increased, bilayers peeled off the substrate and flipped onto existing covered areas to form patches and eventually crystallites. Two LB monolayers of barium arachidate were deposited onto a silanated silicon wafer so that alkyl chains were imaged by AFM$^{15}$. Molecular resolution occurred at higher pH values where a large number of barium ions were incorporated into the bilayer giving additional stability to the film. Annealing of the LB films led to LB film dewetting and domains being formed that coexisted with bilayer. In summary, STM and AFM are tools to study the quality of LB films with respect to preparation method, pH, aging and annealing.

Monolayers can also be self-assembled rather than prepared by LB deposition. The types of molecules that form self-assembled monolayers
include trichlorosilanes on silicon\textsuperscript{21}, alkanethiols on gold\textsuperscript{22,23} and decanol at the air/gold interface\textsuperscript{24}. The self-assembled monolayers were securely anchored to the gold making them stable to the forces exerted by the STM and AFM procedures.

Phthalocyanines (Pcs) and perylenes are commercially important with a wide range of applications. Both of these classes of materials can be used as photoreceptor pigments in copiers and laser printers for instance. Diphthalocyanines in particular exhibit electrochromic properties changing color when different voltages are applied\textsuperscript{25-27}. Perylene and phthalocyanine derivatives are available which are water soluble and can be used as colorants in inks. Some phthalocyanines can reversibly bind gases such as nitrogen dioxide with applications as gas sensors\textsuperscript{28-32}.

6.2. EXPERIMENTAL


A Fromherz\textsuperscript{33} trough and dipper housed in a glove box on a terrazzo vibration-free table were used for film compression and transfer. Generally milli-Q water with one additional distillation was used without further treatment (pH $\sim$ 6.3) at ambient temperature ($\sim$ 23°C). Two types of substrates, muscovite mica and silicon wafers were used for LB deposition and subsequent atomic force microscopy.

Muscovite mica (#4V-2 grade) was purchased from the Asheville-Schoonmaker Mica Company, Newport News, Virginia. Tweezers and a needle
were cleaned with Aldrich denatured HPLC grade ethyl alcohol (0.2 μm filtered). A sheet of mica was gripped using the tweezers and the needle was used to strip off an upper layer of contaminated mica. Successive layers of mica were peeled in a similar manner until a sheet was obtained with a large area showing a single visible color (thickness) on reflection in visible light. The mica had to be thick enough to support itself and typically gave a purple or yellow color on reflection. A smaller area was cut using a continuously heated platinum wire. The mica piece was clamped into the LB sample holder and immediately immersed in the Mega pure water subphase prior to the spreading of the monolayer.

Polished silicon test wafers of (111) orientation, p-boron doped and 3" diameter were purchased from Wafer Net, San Jose, CA. A diamond-tipped scribe was used to cut the wafers into 2.5 cm X 7.5 cm pieces suitable for our holder and LB trough. Any debris was rinsed off with the Mega pure water. The substrate was immediately immersed in the Mega pure water subphase prior to the spreading of the monolayer.

The two perylenes, \(N\)-hexyl-\(N'\)-ethyl-3,4:9,10-perylenetetracarboxylic diimide (HPTCDE) and \(N\)-hexyl-3,4:9,10-perylenetetracarboxylic monoimide (HPTCO) were synthesized and purified by Professor Nagao\(^\text{34}\). For spreading onto the water surface, the HPTCDE was dissolved (\(\sim 1 \times 10^{-4}\) M) in Burdick and Jackson high purity chloroform. Similarly, the HPTCO was dissolved (\(\sim 3 \times 10^{-5}\) M) in B&D high purity chloroform. The compression speed of the trough was (1.9 \(\times\) 10\(^{-3}\) nm\(^2\)/molecule)/s for HPTCDE and (1.7 \(\times\) 10\(^{-3}\) nm\(^2\)/molecule)/s for HPTCO. Film transfer was carried out at 3.6 mm/min with a constant pressure of 15 mN/m by vertical substrate withdrawal for both HPTCDE and HPTCO on
mica and Si(111). Multilayers were transferred to IR transparent substrates (i.e., ZnS, Ge) and smooth silver films. The latter substrate was used for reflection-absorption infrared spectroscopy. Smooth silver films were prepared in high vacuum by evaporating 110 nm of silver onto Corning 7059 glass slides kept at 200°C. Silver thickness was monitored by an XTC Inficon quartz crystal oscillator. LB films were transferred at a constant pressure of 25 mN/m. The substrate was moved vertically through the gas-liquid interface at a rate of 3.6 mm/min. Up to 4 monolayers were transferred to smooth Ag substrates. Fifteen to thirty LB monolayers were transferred on withdrawal to the IR transmission substrates. Z-type deposition was used for all multilayer transfers with the water surface being cleaned each time before reinsertion of the substrate followed by spreading of a new film.

The 1,4,8,11,15,18,22,25-octabutoxy-29H,31H-phthalocyanine, ((BuO)₈-H₂Pc) was obtained from Aldrich and was used without further purification. For spreading onto the water surface, (BuO)₈H₂Pc was dissolved (∼ 6 x 10⁻⁴ M) in Baker Analyzed reagent 1,1,1-trichloroethane. The compression speed of the trough was (3.5 x 10⁻³ nm²/molecule)/s. Film transfer was carried out at 3.6 mm/min with a constant pressure of 15 mN/m by vertical substrate withdrawal.

The tetra-tertiary-butyl metal-free phthalocyanine, ((t-bu)₄H₂Pc) was synthesized in-house from 4-t-butyl phthalonitrile in refluxing amyl alcohol, the reaction being promoted by sodium amylate. Purification was by acid pasting and contains all of the possible isomers which are related by the different relative positions of the t-butyl groups. For spreading onto the water surface, the (t-bu)₄H₂Pc was dissolved (∼ 5 x 10⁻⁴ M) in Eastman Kodak ACS spectro grade toluene. The compression speed of the trough was (∼ 1.1 x 10⁻³
nm²/molecule)/s. Film transfer to the substrate was done at 3.6 mm/min at a constant pressure of 8 mN/m by vertical substrate withdrawal.

6.2.2. FTIR spectrometer and Reflection-Absorption Measurements.

A Bomem DA3 FTIR spectrometer with a MCT liquid nitrogen cooled detector was used to record IR spectra. A Spectra Tech spectral reflectance variable angle accessory was used to obtain RAIR spectra. An incidence angle of 75° to the surface normal was used in all cases. For data analysis, all files were imported to Spectra Calc™ software available from Galactic Industries Corp.

6.2.3. Park Scientific Atomic Force Microscope.

The AFM studies of HPTCDE on mica were carried out using a Park Scientific Instrument scanning force microscope with integrated Si₃N₄ tips. The force constant of the cantilever was determined from geometry and using Young's modulus. Scanning electron micrographs of the tips indicate that the Si₃N₄ tips have a radius of curvature on the order of 100 nm at the end of the tip. A high resolution image of the muscovite mica substrate was taken in the constant height mode. In the first attempt to image HPTCDE on mica, too large a force of 100 nN was applied which removed the HPTCDE from the substrate. A lower force of 10 nN was used to image a different region of the HPTCDE on mica. The micron sized images were taken in the constant force mode. All of the AFM images were collected in air. The height and distances were calibrated using a displacement meter. The scans of the mica substrate also provided a small-scale calibration since the dimensions of the mica unit cell are well known.
6.2.4. Digital Instruments Nanoscope III SPM.

All images were collected in the tapping mode with a 125 μm silicon probe. A "J" (125 μm) scanner was used in a Nanoscope III MultiMode AFM.

6.3. RESULTS AND DISCUSSION

Figure 6-2 is a high resolution AFM image of bare muscovite mica [KAl₂(Si₃Al)O₁₀(OH)₂] displaying atomic scale resolution. A Fourier transform of the image had the two largest peaks at \( k₁ = 12.6 \text{ nm}^{-1} \) and \( k₂ = 11.5 \text{ nm}^{-1} \) separated by an angle of 129 degrees. Using the \( k₁ \) and \( k₂ \) values, the calculated unit cell area was 0.21 nm². From the literature³⁹,⁴⁰, it is known that the surface of mica is a triangular lattice with a spacing of 0.52 nm between the vertices and a surface unit cell area of 0.23 nm².

It was difficult to accurately determine the transfer ratio of HPTCDE to mica as the substrate piece was very small and irregular in shape. The transfer ratio is the area of material removed from the trough divided by the area of substrate used for the deposition. The HPTCDE had a transfer ratio of 1.0 to the Si(111) substrate using z-deposition. At the transfer pressure of 15 mNm⁻¹, the area per molecule was 0.48 nm² taken from the surface pressure-area isotherm. As shown in Figure 6-3, if too large a force is applied to the monolayer of HPTCDE on mica, the AFM tip can scrape away the film. One application of the AFM would be as a modifying tool to write features on a submicron scale. The AFM or STM could also be used for nanotechnology⁴¹ to build structures from the bottom up by manipulation and selective placement of atoms.
Figure 6-2: High resolution image of muscovite mica obtained on a Park Scientific AFM.
Figure 6-2

Mica surface
Another region of the LB monolayer of HPTCDE on mica was imaged with a smaller force of 10 nN and the large area image appears in Figure 6-4. Molecular resolution was not observed. Three different height levels were apparent as black, gray and white. The height difference between the black and gray regions was $1.83 \pm 0.34$ nm measured for seven cross-sections and between the gray and white features was $1.86 \pm 0.28$ nm using six cross-sections. The HPTCDE does not appear to be coating the mica uniformly. The black areas are the underlying mica substrate, the gray regions HPTCDE monolayer, and the white features are two layers of HPTCDE. The HPTCDE monolayer thickness of $1.83 \text{ nm} \pm 0.34 \text{ nm}$ is consistent with reflection-absorption infrared measurements. The molecular orientation in the LB film has been elucidated through infrared spectroscopy\textsuperscript{42}. Using the absorbance ratio between the transmission and reflection-absorption infrared spectra, mean molecular tilt angles of the long and short axes of the chromophore in the LB layers have been determined.

The FTIR transmission and reflection-absorption spectra of HPTCDE are shown in Figure 6-5. HPTCDE dispersed in a KBr pellet corresponds to isotropic molecular orientation. The bands in the region around 1600 cm\textsuperscript{-1} are due to C=C stretching vibrations in the benzene rings. Peaks at 1656 cm\textsuperscript{-1} and 1659 cm\textsuperscript{-1} have been assigned to anti-symmetric carbonyl stretching and symmetric carbonyl stretching respectively\textsuperscript{42}. Anti-symmetric carbonyl stretching causes changes in the dynamic dipole moment along the short axis of HPTCDE while symmetric carbonyl stretching changes the dynamic dipole moment along the long axis of HPTCDE. The tilt angle $\phi$ is defined as the angle between the dynamic dipole moment and the normal to the substrate surface. The tilt angle can be determined using equation 6.142-44.
Figure 6-3: HPTCDE Langmuir-Blodgett monolayer scraped from the mica substrate when the applied force is too large. Park Scientific AFM image.
HPTCDE perylene on mica

1 \mu m
Figure 6-4: HPTCDE Langmuir-Blodgett monolayer on mica imaged with force of 10 nN. Three different heights were observed using the Park Scientific AFM.
Figure 6-4

HPTCDE perylene on mica

0.5 µm
Figure 6-5: FTIR spectra of HPTCDE in the carbonyl stretching region. The transmission and reflection spectra show the change in relative intensity of the symmetric and anti-symmetric C=O stretches.
\[
\frac{A_T}{A_R} = \frac{\sin^2\phi}{2m_z\cos^2\phi + m_x\sin^2\phi} \quad \text{equation 6.1}
\]

\(A_T\) and \(A_R\) are the measured IR absorbance for transmission and reflectance and are normalized to the number of LB layers. \(m_z\) and \(m_x\) are enhancement factors for light which is polarized normal (z) and parallel (x or y) to the surface. In the RAIRS experiment, \(m_x < m_y\) yielding equation 6.2.

\[
\frac{A_T}{A_R} = \frac{\tan^2\phi}{2m_z\cos^2\phi} \quad \text{equation 6.2}
\]

A method for estimating \(m_z\) based on a hypothetical ratio of \(A_T/A_R\) for isotropic films has been reported\(^{42,43}\). For HPTCO, the tilt angles with respect to the surface normal were 36 degrees and 64 degrees for the long and short axis respectively. For HPTCDE, the values for the same tilt angles were 60 and 80 degrees respectively. These head-on molecules would form a monolayer about 2 nm in height. The image of the LB monolayer of HPTCDE on Si(111) appears in Figure 6-6. The 1.0 \(\mu\)m scan shows that the film is not homogeneous and there are some voids present.

The transfer ratio of HPTCO to Si(111) by z-deposition was 1.1. At the transfer pressure of 15 mNm\(^{-1}\), the area per molecule was 0.31 nm\(^2\) taken from the surface pressure-area isotherm. The limiting area\(^{42}\) obtained for this molecule using the low compressibility region of the isotherm was 0.24 nm\(^2\).
Figure 6-7 is the 1.0 μm image of HPTCO. An island structure of HPTCO occurs rather than a continuous film. Assuming that the film formed on Ag is also an island film, it can be concluded that the molecular orientation occurs within the islands of the LB film.

The transfer ratio of (BuO)$_8$H$_2$Pc to Si(111) by z-deposition was 0.71. At the transfer pressure of 15 mN/m$^{-1}$, the area per molecule was 0.87 nm$^2$ from the surface pressure-area isotherm. Ordered LB multilayers of octa-$n$-alkoxy phthalocyanines such as (BuO)$_8$H$_2$Pc were fabricated on glass and described by others previously$^{45}$. A value of 0.71 nm$^2$ per molecule was reported$^{45}$ at a surface pressure of 30 mN/m$^{-1}$. X-ray diffraction patterns for some octa-$n$-alkoxy phthalocyanines corresponded to the maximum molecular dimension with the plane of the Pc molecule orthogonal to the substrate$^{45}$. Figure 6-8 is the AFM image of the (BuO)$_8$H$_2$Pc monolayer on Si(111). The mean surface roughness was low at 0.180 nm.

The transfer ratio of (t-bu)$_4$H$_2$Pc to Si(111) was 1.3 by z-deposition. The area per molecule at 8 mN/m$^{-1}$ was 0.44 nm$^2$ extrapolated from the surface pressure-area isotherm. Baker and co-workers reported the preparation and properties of tetra-tertiary-butyl metal-free phthalocyanine$^{46}$. Kovacs et al.$^{47}$ provided further evidence of ordered true monolayers with a perpendicular distance between adjacent molecules of 0.334 nm, 0.547 nm between molecules in the stacking direction and 1.715 nm as the overall in-plane length of the Pc molecule. The structure and morphology of a related material, copper tetra-tertiary-butyl phthalocyanine has been investigated using transmission electron microscopy, transmission electron diffraction, X-ray
Figure 6-6: AFM image of HPTCDE Langmuir-Blodgett monolayer on Si(111) wafer. Nanoscope III SPM.
Figure 6-7: AFM image HPTCO Langmuir-Blodgett monolayer on Si(111) wafer. Nanoscope III SPM.
diffraction and light microscopy\textsuperscript{48}. The transmission electron diffraction patterns of Cu(t-bu)$_4$Pc had two perpendicular periods corresponding to 0.332 nm and 1.8 nm\textsuperscript{48}. Computer-based molecular modeling and a simulation suggested a zig-zag packing for the most frequent isomers\textsuperscript{48}. The AFM image of a (t-bu)$_4$H$_2$Pc Langmuir-Blodgett monolayer appears in Figure 6-9. Some voids are present but there are regions where the (t-bu)$_4$H$_2$Pc is continuous.

The factors affecting lateral resolution of the AFM for LB films have been considered by others\textsuperscript{49,50}. The contact diameter for repulsive mode AFM was calculated taking into account the applied force, tip radius, Young’s modulus and Poisson’s ratio\textsuperscript{49}. Adhesive forces between the tip and the sample increase the contact diameter and decrease the resolution when operating in the contact mode\textsuperscript{49}. The best AFM resolution for soft organic molecules in air may be 2-3 nm with small forces applied\textsuperscript{49}. Surfaces that are imaged in air are usually covered with an adsorbed layer of water and the adhesive forces between the tip and sample may be greater than the force desired for imaging\textsuperscript{6}. One approach to reduce this adhesive force would be to image with the tip and sample immersed in liquid\textsuperscript{6}. In this study of LB monolayers of perylenes and phthalocyanines, the fact that molecular resolution was not observed, could be due to a large contact diameter.
Figure 6-8: AFM image of (BuO)$_3$H$_2$Pc Langmuir-Blodgett monolayer on Si(111) wafer. Nanoscope III SPM.
Figure 6-9: AFM image of (t-bu)$_4$H$_2$Pc Langmuir-Blodgett monolayer on Si(111) wafer. Nanoscope III SPM.
6.4. CONCLUSIONS

The AFM technique has been used to investigate the quality and surface coverage of Langmuir-Blodgett monolayers of important electroactive organic molecules on mica and Si(111) surfaces. With the AFM contact mode, it was possible to determine the monolayer height by indentation. The AFM tapping mode revealed film topography, surface coverage and voids in the nanometer regime. However, molecular resolution was not achieved. Transmission and reflection-absorption FTIR spectroscopy provided an average molecular orientation of HPTCDE with an LB monolayer height in close agreement with the value found using the AFM contact mode. It is important to point out that together, spectroscopy and AFM provide the ideal tool for structural and physical studies of LB films.
Chapter 6 References


CHAPTER 7: OVERALL SUMMARY, CONCLUSIONS AND FUTURE OUTLOOK

7.1 Overall Summary and Conclusions

A discussion and conclusions have been presented in the experimental sections: Chapter 4, Chapter 5 and Chapter 6. The key points will be summarized here.

Phthalocyanines and perylenes are important charge generation materials which are used in organic photoreceptors. Several crystalline polymorphs exist for titanyl phthalocyanine and vanadyl phthalocyanine. The polymorphic form is affected by heating, solvents and processing conditions. X-ray powder diffraction is commonly used to identify polymorphs and it has been shown that FT-Raman spectra can be used to distinguish polymorphs. Polymorphs exhibit different physical properties and photosensitivity. FT-Raman spectra were presented for four TiOPc reference materials that had previously been identified by X-ray powder diffraction. Films of titanyl phthalocyanine were vacuum evaporated onto glass held at -30°C to 240°C. The TiOPc was amorphous from -30°C to room temperature and pure type II TiOPc was present from 102°C to 240°C. FT-Raman spectra were collected of vacuum evaporated vanadyl phthalocyanine deposited on glass substrates held from -30°C to 240°C. The vanadyl phthalocyanine is amorphous at room temperature and at 240°C is type II. FT-Raman frequencies of both TiOPc and VOPc were assigned based on information from model compounds and other phthalocyanines. It was therefore, demonstrated that FT-Raman spectroscopy could be applied to thin solid vacuum evaporated films.
Since the Raman scattering intensity is weak for thin solid films, surface-enhanced Raman scattering was used to increase the intensity in the near-IR. Molecular modeling suggested a number of metals should give SERS using 1064.1 nm excitation. Copper and gold metal island films were prepared and characterized as FT-SERS substrates. The enhancement factor was approximately 100. It was initially shown that FT-SERS could be obtained for 3 nm vacuum evaporated phthalocyanine and perylene films on metal island structures. FT-SERS was then demonstrated for single Langmuir-Blodgett monolayers on copper and gold island films. The experimental results suggested the organic materials were physisorbed on the metal island films with electromagnetic enhancement. Surface-enhanced resonance Raman spectra were presented for comparison using visible laser excitation.

The trend in the experimental work was to study thin solid films on smaller and smaller scales. Thin vacuum evaporated films were studied, then Langmuir-Blodgett monolayers and finally the atomic force microscope was used to characterize the structure of the LB monolayers. Langmuir-Blodgett monolayers of two perylenes, HPTCDE and HPTCO were transferred to atomically flat substrates of mica and Si(111) wafers. For comparison, a transmission infrared spectrum and a reflection-absorption infrared spectrum were measured of a monolayer of HPTCDE on IR substrates. The HPTCDE film on mica had three regions of different thicknesses measured using the AFM with the step height 1.83 ± .34 nm in agreement with the orientation inferred from transmission IR and RAIRS measurements. If the force on the sample is too great the LB monolayer can be scraped from the film. Four different LB monolayers on Si(111) were examined using a tapping mode AFM. The materials considered were HPTCDE, HPTCO, CuTTPc and (BuO)₈H₂Pc.
Molecular resolution was not obtained. The AFM was an excellent probe for local defects in the LB films. In contrast, techniques such as X-ray diffraction and infrared spectroscopy give results averaged over an area of sample. An advantage of AFM over STM is that lower force can be exerted on the sample and a conductive sample is not necessary.

7.2 Future Prospects

Returning to the model calculations of SERS enhancement factors at 1064.1 nm for various metals reported in Table 3-2, it can be seen that other elements than copper and gold should produce FT-SERS of a similar magnitude. The calculations were carried out for a specified set of input parameters including morphology. Future modeling could be done with different morphologies. An optimum morphology for FT-SERS enhancement could be selected and the challenge would be to experimentally produce these structures. This author carried out some limited experiments to check the morphology of 20 nm mass thickness vacuum evaporated films of indium, vanadium, chromium and aluminum on glass that had been held at 200°C during deposition. Electron micrographs revealed closely packed spheres of less than 0.15 microns diameter for indium. The 20 nm mass thickness chromium consisted of isolated somewhat spherical structures on glass with a typical diameter of about 0.3 microns. There was a lot of space between the centers of the spherical structures. The 20 nm mass thickness vanadium on glass had a distribution of shapes ranging from spherical to cubic with a lot of space between the centers. The aluminum film consisted of closely packed spheres of less than 0.15 microns diameter. It was assumed that an aluminum oxide layer was formed on removal from the vacuum chamber.
The future of FT-Raman spectroscopy can be considered. It is expected that Raman spectroscopy in general will be used more routinely as a complementary technique to infrared spectroscopy. The major advantage of FT-Raman spectroscopy compared to normal Raman spectroscopy is reduced fluorescence. Also since some materials absorb in the visible region, it may be desirable to collect data away from resonance. There presently are many infrared digital libraries available for compound identification. In the near future it is expected Raman digital libraries will be common. Computer-assisted spectral interpretation and intelligent systems have been applied to infrared data bases and should be useful for Raman data sets. FT-Raman results reported in the experimental section were obtained using the 1064.1 nm Nd/YAG laser line. If materials still exhibit fluorescence at this wavelength there is an opportunity to move to an even longer wavelength with a Nd/YLF laser. Reduced fluorescence can be compromised for a gain in scattering intensity if the shorter wavelengths of a tunable Ti-sapphire laser are used. FT-Raman will have applications for on-line measurements or remote sensing.

Atomic force microscopy and scanning tunneling microscopy have become popular for examining local structures on the micron to nanometer scale. Whenever an LB film is prepared, its properties must be characterized before it can be used in an application. AFM could be applied to mixed monolayers to determine if aggregates are formed or a homogeneous mixture. There appears to be some interest in the literature towards self-assembled monolayers and organized structures1. In practice, these materials may be easier to fabricate than Langmuir-Blodgett films and fit in with the idea of nanotechnology and building structures from the bottom-up. Two examples of self-assembled structures are self-assembled vesicles to deliver drugs and a
self-assembled sandwich of electronically active materials\textsuperscript{1}. The STM can also be used to move molecules and atoms and could be used to fabricate structures from the bottom up.

Chapter 7: References

Appendix I: Sample output of the prolate enhancement factor for (t-bu)$_4$H$_2$Pc on a gold hemi-spheroid

The value of $\varepsilon_0$ is 1.00.
The wavelength of laser excitation is 1064.1 nm.
The wavelength of scattered light is 1148.0 nm.
The spectral shift is 686.8 wavenumbers.
The dielectric constant at the laser wavelength is $-48.787 + 3.637i$.
The dielectric constant at the scattering wavelength is $-58.634 + 4.781i$.
The length of the semi-major axis is 38.0 nm.
The length of the semi-minor axis is 33.0 nm.
The aspect ratio of the prolate spheroid is 1.15.

<table>
<thead>
<tr>
<th>Distance from top of spheroid (nm)</th>
<th>Enhancement factor</th>
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</thead>
<tbody>
<tr>
<td>0.00</td>
<td>0.155E+03</td>
</tr>
<tr>
<td>0.50</td>
<td>0.135E+03</td>
</tr>
<tr>
<td>1.00</td>
<td>0.118E+03</td>
</tr>
<tr>
<td>1.50</td>
<td>0.103E+03</td>
</tr>
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<td>0.911E+02</td>
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<td>0.716E+02</td>
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<td>3.50</td>
<td>0.639E+02</td>
</tr>
<tr>
<td>4.00</td>
<td>0.572E+02</td>
</tr>
<tr>
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Appendix I continued:

Sample output of the prolate enhancement factor for \((t\text{-}bu)_{4}\text{H}_2\text{Pc}\) on a gold hemi-spheroid

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<th>DISTANCE FROM TOP OF SPHEROID (NM)</th>
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<td>0.463E+01</td>
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<tr>
<td>24.50</td>
<td>0.448E+01</td>
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</tbody>
</table>
VITA AUCTORIS

Carol Ann Jennings was born in 1958 in Toronto, Ontario. She graduated from Alderwood Collegiate Institute in 1977. From there she went to the University of Toronto where she obtained a B.Sc. in Chemistry in 1981. From 1982-1984 she held a position at the University of Toronto. During 1985-1986 she completed a M.Sc. in Chemistry at the University of Windsor. She became a member of the research staff at Xerox Research Centre of Canada in 1986. During 1989 to 1990 she successfully completed three graduate courses at the University of Waterloo in polymer chemistry as a part-time special student. In 1990 she entered a Ph.D. program as a part-time student at the University of Windsor. In May of 1994 Carol transferred from the Xerox Research Centre of Canada to the Research and Technology group of Xerox Graphics Systems. She expects to complete all the requirements for a Ph.D. in 1994.