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Probing Single Molecules with a Tunable Femtosecond Laser Coupled RF-STM

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Probing Single Molecules with a Tunable Femtosecond Laser Coupled RF-STM

DISSERTATION

submitted in partial satisfaction of the requirements
for the degree of

DOCTOR OF PHILOSOPHY

in Materials Science and Engineering

by

Weicai Cao

Dissertation Committee:
Professor Wilson Ho, Research Advisor
Professor Martha Mecartney, Committee Chair
Professor Regina Ragan

2015
DEDICATION

To

Annie and our family
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Professional Skills

- Designing, constructing, maintaining and improving Scanning Tunneling Microscope (STM) along with related scientific instruments.
- Setting up and maintaining complicated tunable femtosecond laser system which covers wide wavelength range.
- Probing single molecules, single atoms and nanoclusters with a homemade tunable femtosecond laser (100fs, 210~1040nm) coupled low temperature (10 kelvin) ultrahigh vacuum (4×10^{-11} torr) Radio Frequency (80MHz) STM.
- Characterizing microstructure of materials with scanning electron microscope (SEM), atomic force microscope (AFM), transmission electron microscope (TEM) and x-ray diffraction (XRD); characterizing mechanical, optical and electrical properties of materials.
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Graduate Research Assistant of Dr. Wilson Ho 07/2009-Present

- Moved a homemade STM system to a new lab; designed and assembled a molecular interlock system for exchange of evaporators; designed, machined and assembled atom and molecule evaporators; designed and built a mobile pump station (leak detector) to leak check and detect current of evaporators; designed and built a station to capture the helium (He) gas used during LHe LT experiments, co- designed and built systems for storage, purification, and liquefaction of He.
• Set up a new tunable femtosecond laser system that can cover whole solar spectrum from 210nm to 1040nm with second, third and fourth harmonic generations; overlapped two ultrafast laser beams (MaiTai fundamental+fundamental, fundamental+2<sup>nd</sup> harmonic, fundamental+3<sup>rd</sup> harmonic) spatially and temporally on the optical table and in the STM junction; co-designed, modified and built new microscopes to detect weak laser induced signal in the tunneling regime.

• Performed STM experiments of molecules, atoms and nanoclusters on oxide and metal surfaces (properties of these systems were probed at the single molecule level with sub-Å resolution; light molecule interactions mediated by plasmons at the nanoscale were also studied by combining laser with STM). Investigated molecules are C<sub>60</sub>, alizarin, Th-Th-BT-Th-Th, azulene, and PDI, investigated atoms and nanoclusters included Ag, Au, Fe, K and Mg.

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Rotation Graduate Researcher 10/2008-07/2009

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• Synthesized and characterized microstructure and mechanical properties of Al<sub>2</sub>O<sub>3</sub>-YAG-ZrO<sub>2</sub> three phase ceramic; Characterized microstructure and catalytic activities of Au nanoparticles.

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ABSTRACT OF THE DISSERTATION

Probing Single Molecules with a Tunable Femtosecond Laser Coupled RF-STM

By

Weicai Cao

Doctor of Philosophy in Materials Science and Engineering

University of California, Irvine, 2015

Professor Wilson Ho, Research Advisor

Professor Martha Mecartney, co-Advisor and Chair

Scanning Tunneling Microscope (STM) has become a powerful tool in nanoscience for imaging, manipulation and electronic spectroscopy. STM inelastic electron tunneling spectroscopy (IETS) first achieved chemical identification of molecular species by characterizing vibrational energies. Recently, with the STM itProbe and H₂ rotational spectromicroscopy, molecular structure and chemical bonds are observed with the STM. Despite these successes in spatial resolution, various efforts have been made to combine fs laser with STM to overcome the temporal resolution limitation of STM, there is so far no clear evidence of simultaneous fs and Å resolution.

Electronic properties of organic molecules are of central importance to applications such as molecular electronics, organic LEDs, and solar cells. Properties of these molecules can be probed by the scanning tunneling microscope (STM) at the single molecule level and with sub-Å spatial resolution. The molecular orbital of 4, 7-Di ([2, 20-bithiophen]-5-yl) benzo[c] [1, 2, 5] thiadiazole (4T-BTD) with intramolecular donor-acceptor-donor sites is probed with the electronic state dI/dV imaging and H₂ rotational and vibrational spectromicroscopy. 1, 4-Phenylene Diisocyanide (PDI) is probed by imaging with a CO-terminated tip and H₂. PDI can
self-assemble on noble metal surfaces to form nanostructures, which could have potential applications in molecular electronics and catalysis.

Further combination of a RF-STM with a tunable femtosecond laser enables the investigation of light-molecule interactions. In this dissertation, efforts are spent to setup a new tunable fs laser (220 nm~1040 nm) to couple with the RF-STM. The effects of the femtosecond laser are followed by detecting photo induced electron emission and photochemistry. A new double lock-in technique is applied to detect the weak laser-induced signal in the tunneling regime. To sharpen the energy width and increase the lifetime of the excited states of molecules, thin aluminum oxide and copper oxide are grown on metal surfaces to provide electronic isolation of the metal substrate and adsorbed molecules. Metal nanoclusters are grown on metal and oxide to improve laser-induced signal through plasmonic enhancements.
Chapter 1

Introduction

1.1 Background of STM

Scanning tunneling microscopy (STM) has been a powerful tool to study surface sciences for imaging, spectroscopy and manipulation [1] after its invention in 1980’s [2]. By modifying various experimental conditions such as temperature and pressure, coupling magnetic fields or electromagnetic radiation, and functionalizing the STM tip with single molecules, STM has demonstrated its versatility to probe chemistry and physics of various materials at the atomic scale.

The heart of STM is the quantum mechanical effect of tunneling through a potential barrier applied between two metallic leads. An applied potential difference between the sample and tip causes a tunneling electron current when the distance between the tip and sample is close, typically 5Å to 10Å. This current depends exponentially on the gap distance of the tip and sample surface, typically if the tip and surface get 1 Å closer, the tunneling current will become 10 times bigger. The tunneling current can be kept constant with STM electronics feedback on, which applies offset voltage to the STM piezos to keep a preset tunneling current. STM images obtained in this constant current mode are obtained by scanning the STM tip across the surface and adjusting the tip position required to maintain a constant current. These applied piezo voltages are used to reconstruct STM topographic images which are a convolution of the geometric and electronic structure of the surface [1, 3].

The tunneling current can be expressed by using first-order perturbation theory as:

\[ I = \frac{2eV}{\hbar} \frac{\exp(-\frac{\Delta}{\lambda})}{1 + \exp(-\frac{\Delta}{\lambda})} \]
\[ I_T = \frac{4\pi e}{\hbar} \int_{-\infty}^{\infty} \rho_t (\epsilon - \epsilon V_T) \rho_s (\epsilon) (f_t (\epsilon - \epsilon V_T) - f_s (\epsilon)) |M(\epsilon - \epsilon V_T)|^2 d\epsilon \]  

(1.1)

where \( \rho_t \) and \( \rho_s \) represent the density of states (DOS) of the tip and sample respectively, \( f_t(\epsilon) \) and \( f_s(\epsilon) \) are temperature dependent Fermi distribution functions of the tip and sample respectively, and \( |M(\epsilon)|^2 \) is the tunneling matrix element taking into account the coupling between the electron wavefunctions of the tip and sample. The difficulty in evaluating equation (1.1) lies in determining the tunneling matrix element which depends strongly on the local density of states of both the tip and surface. However, a simple approximation has been performed by J. Tersoff and D. R. Hamman [3, 4] to simplify the expression for the tunneling current.

In the Tersoff-Hamman approximation the STM tip is treated as atomically sharp where only the last atom in the tip contributes to the tunneling process. In addition, the only contributions of wavefunction states of the tip are those that have spherical \( s \)-like orbitals. Using these approximations and taking a zero temperature and low bias limit the tunneling current expression can be simplified to:

\[ I_T \propto \rho_t \int_{0}^{eV_T} \rho_s (\epsilon) d\epsilon \]  

(1.2)

and the derivative of this expression with respect to voltage will give the differential conductance:

\[ \frac{\partial I_T}{\partial V} \propto \rho_s (\epsilon) \]  

(1.3)

which is direct measure of the local density of states (LDOS) of the sample surface or atoms and molecules adsorbed on the surface [1, 3 and 4].

STM electronic spectroscopy is performed in the constant height, the STM feedback electronics are off after the tunneling gap is set; the tunneling current is recorded as the sample bias is ramped in certain bias range. For STM, all the spectroscopy information is recorded in the I-V curve. I-V curve is mostly linear since the tunneling is predominantly a linear process. If
there are electronic states on the surface or molecules adsorbs on the surface, a step will appear in the I-V curve. Such a change is more apparent in the dI/dV curve, with a peak at the bias of the electronic states. Another reason to record dI/dV signal is expressed in equation 1.3, dI/dV is directly related to density of states on the surface or molecules adsorbs on the surface. The dI/dV signal can be readily recorded with lock-in detection by applying a small modulating bias to the tunneling bias.

Despite being a powerful tool in imaging, manipulation and spectroscopy, STM has its limitations, the first one being lack of chemical sensitivity. In 1998, STM inelastic electron tunneling spectroscopy (IETS) first achieved chemical identification of molecular species by characterizing vibrational energies [1, 5]. The operation of STM can be treated as an equivalent circuit consisting of a power supply and a resistor (STM junction, typical 1MΩ to 1GΩ); the resistor is of atomic dimensions, if the molecule in the STM junction moves in the form of vibration, rotation, etc, the resistance will change. Such a resistance change is reflected in the tunneling current change; however, such a small change is too small to be measured in the I-V curve. A few times a change in the dI/dV could be observed, most of time d²I/dV² needs to be recorded and the junction resistance change due to the molecule motion is associated with a positive peak on the positive bias side and a negative peak on the negative bias side [1, 5].

1.2 Imaging with H₂ and CO

Even though STM can routinely achieve high spatial resolution of surfaces and image single molecules, it can also provide electronic and vibrational signatures of single molecules, STM imaging fails to resolve complex chemical structure until recently. Scanning tunneling hydrogen
microscopy (STHM) [6], inelastic tunneling probe (itProbe) [7] and hydrogen rotational spectromicroscopy [8, 9] achieved chemical and structural imaging of single molecules with unprecedented details. Such great-detail chemical structures have also been observed with a noncontact atomic force microscope (NC-AFM) [10].

In all these techniques, small molecules like H\(_2\) and CO are attached to the tip or trapped in scanning probe junction, acting as a sensor of the chemical structure. In the itProbe and NC-AFM, a CO terminated tip is attached to the tip. As the tip scanned across the surface, either the shift in the resonance frequency or inelastic tunneling signal is recorded. Skeletal structure of molecule and chemical bonds can be imaged. Especially STM itProbe broadens STM’s applications to study electronic properties, vibrational properties, molecular structure and chemical bonding of the same molecules combing with STM imaging, electronic spectroscopy and IETS.

Recently it has been demonstrated that by trapping H\(_2\) in the STM junction at low temperature, the chemical structure of organic molecules can be imaged with high resolution revealing geometrical structures and intermolecular hydrogen interactions between monolayers of organic molecules, this way of imaging is called Scanning Tunneling Hydrogen Microscopy (STHM) [6]. More recently, the rotational excitations of hydrogen molecule weakly adsorbed on surfaces have been measured with the STM [11, 12]. It was also shown that similar to the vibrations of a CO molecule attached to the tip [7], changes in the adsorption potential well induced by varying the tip-sample distance shift the energies in both vibrational and rotational modes for the hydrogen temporally trapped in the tunnel junction [13]. The sensitivity of the H\(_2\) modes to different chemical environments has been applied to achieve sub-Å structural and molecular orbital imaging of MgP on metal and oxide surfaces [8, 9].
1.3 Laser STM

Shortly after the invention of STM in 1982 [1], the idea of combining optical techniques with STM has been pursued [13]. One way to couple optical techniques with STM is the STM induced light emission; this technique now has achieved submolecular resolution [14]. Due to the thermal effects induced by laser and some other experimental challenges, for long time, little success has been achieved toward the combination of high spectral and temporal resolution of laser with the spatial resolution of STM [13]. Submolecular Raman spectral imaging has been achieved by coupling the Raman process to tunneling [15].

More importantly, fs laser techniques like 2 Photon Photoemission (2PPE) has achieved fs temporal resolution [16], by combining fs laser with STM, simultaneous temporal fs resolution and spatial sub-Å resolution could be achieved. However, despite the successes in spatial and spectral resolution, although lots of efforts have been spent, there is so far no clear evidence of simultaneous fs and Å resolution, achieving this great goal has been the “holy grail” in the field [17].

Besides STM induced light emission, there were also efforts in the Ho Group to achieve simultaneous fs- Å resolution [18, 19]. In these efforts, the Ti: Sapphire laser beam fundamental beams are used for the pump-probe time resolved experiments. The problem by using two fundamental beams is they will cause optical interference when the two beams spatially and temporally overlapped. One way to solve this is using two different polarization beams although the two beams will still slightly interfere [18, 19], a better way to avoid interference is by using two different color laser beams.
In this dissertation, efforts are spent to setup a tunable wavelength (220nm~1040nm) femtosecond laser system by using a harmonic generator to extend the wavelength from a MaiTai Deepsee laser. Different color laser beams (fundamental and 2\textsuperscript{nd} harmonic, fundamental and 3\textsuperscript{rd} harmonic) are made overlapped on the diagnostic path. However, it’s hard to overlap the two color beams in the junction given there are lenses inside the UHV chamber to focus the beams and the refractive index is different for different colors and will change the beam path. The excited states of molecules are usually in the visible to the UV, another reason that the wavelength of the laser needs to be tunable in this range is in order to be able to excite the tunneling electrons into resonance with an excited state of the molecule.

To reach sub-Ångström spatial limit in optical experiments of single molecules, previous studies have been shown that it is necessary to couple light to the tunneling process [19]. The difficulty in such experiments is that the rate of production of photoinduced tunneling electrons is very low in relation to the background of bias-induced tunneling electrons in the operation of the STM. We are using a new method by detecting the electrons produced at the repetition rate of the femtosecond laser (~80 MHz), which provides an intrinsic modulation of the photon emitted or photon-induced tunneling electrons. An external bias tee was used to separate the DC and RF signal, a 50Ω terminal resistor was used as the RF current to voltage converter. The RF signal will go to the 200MHz RF lock-in (or spectrum analyzer) through the RF amplifier for detection. Further discrimination is achieved by connecting the output of 200 MHz lock-in amplifier to a second lock-in amplifier that detects a lower frequency signal generated by modulating the sample bias, for dI/dV and d^2I/dV^2 spectroscopy. To improve, a new cryogenic amplifier can be installed to get better signal/noise ratio, a fabricated plasmonic tip or nanoclusters on surface can be used for better plasmonic enhancement.
1.4 Summary of Contents

In the introduction, methods to extend both the spatial and temporal resolution of STM were addressed. In the remaining part of this chapter a brief summary of the following chapters is presented of the work conducted in this thesis.

Chapter 2: “A Tunable fs Laser Coupled RF-STM” describes the effort to set up the new 220 nm~1040 tunable fs laser system, including alignment, characterization and simulation on the optical table. And then this chapter describes the alignment of different color beams to the STM junction, especially the effort to align two different color beams overlap spatially and temporally in the STM junction. Lastly, this chapter describes the detection of laser induced current, including the DC rectified current and the RF current with the double lock in technique.

Chapter 3: “Imaging Molecular Orbitals of 4, 7-Di ([2, 20-bithiophen]-5-yl) benzo[c] [1, 2, 5] thiadiazole (4T-BTD) with a STM” by W. Cao, C. J. Patel, G. C. Welch, G. C. Bazan and W. Ho, prepared to submitted to Nano Lett. 4T-BTD is an organic molecule with intramolecular donor-acceptor-donor sites. This paper first describes the STM topography of the 4T-BTD molecule on the NiAl(110) surface, which shows bias dependence, the electron poor benzothiodiazole fragment in the acceptor site can only be clearly imaged at bias near the LUMO peak. LUMO dI/dV spectra clearly show that the center acceptor unit has higher peaks, LUMO dI/dV imaging more clearly highlights the center acceptor part. H2 rotational IETS imaging is performed on the molecule, LUMO structure of the molecule are more clearly imaged at sub-Å resolution, which is the first time observe sub- Å details of the molecular orbital on the metal surface by using H2.
Chapter 4: “Imaging Metal-Molecule Chain and Single Molecule of 1, 4-Phenylene Diisocyanide (PDI) on Noble Metal Surfaces and Their Interaction with Small Molecules” by W. Cao, C.J. Patel, M. Feng, H. Petek and W. Ho, prepared to be submitted to Journal of Chemical Physics. This paper first describes the chains on noble metal surfaces and the IETS of CO\textsubscript{2} trapped on the chain, and then focuses on the imaging of single PDI molecules. Single PDI molecules are studied on Cu (110) surface with H\textsubscript{2} imaging. At low H\textsubscript{2} coverage, while the STM topography image only shows two dimples which may be the N-C groups, the STM H\textsubscript{2} IETS image clearly shows the two N sites. At high H\textsubscript{2} coverage, the topography image already shows three parts of the molecule with interactions with the surface, the IETS image shows more details of the molecule and its interaction with the substrate, with better contrast and sharper transitions.

Chapter 5: “Exploration of New Oxide Surfaces and Study of Molecules in Various Chemical Environments” describes the STM work of molecules on oxide surfaces. Large aluminum oxide islands are formed on NiAl (110) surface by modifying previous recipes, CuO is formed on Cu (110) surface with well-defined structure. STM imaging and spectroscopy are performed on molecules (including C\textsubscript{60}, azulene, alizarin and CO) adsorbed on Al\textsubscript{2}O\textsubscript{3} and CuO surfaces. The thin oxide provides electronic decoupling from the underlying metal substrate that sharpens the electronic states energy width of the molecule, the oxide can also increase the lifetime of excited states of adsorbed molecules.

Chapter 6: “Imaging and Electronic Spectrum of Fe Nanoclusters on Al\textsubscript{2}O\textsubscript{3}/NiAl (110) at Various Temperatures with a Laser-STM” describes the imaging and electronic properties of Fe nanoclusters that are formed on Al\textsubscript{2}O\textsubscript{3}/NiAl (110) surface with STM. Fe clusters are formed on oxide at room temperature and 84K. Bigger Fe cluster can be formed by annealing the sample to
high temperature. Electronic properties are studied at 84K and 21K, DC and RF dI/dV spectra are also studied with 220nm fs laser in junction.

The concluding chapter (Chapter 7) summarizes the dissertation and significance. New directions for further experimental directions are proposed that should elucidate the experiments conducted in this thesis.

Appendix A includes the catalog of the STM4 laser system and describes the alignment and the key parts and homemade parts of the system. Appendix B describes the evaporator loadlock of the STM4 system and new evaporators.


Chapter 2

A Tunable fs Laser Coupled RF-STM

2.1 Introduction/Overview

Shortly after the invention of STM in 1982 [1], the idea of combining optical techniques with STM has been pursued [2]. One way to couple optical techniques with STM is the STM induced light emission [3]; this technique now has achieved submolecular resolution [4]. Due to the thermal effects induced by laser and some other experimental challenges, for long time, little success has been achieved toward the combination of high spectral and temporal resolution of laser with the spatial resolution of STM [2]. Submolecular Raman spectral imaging has been achieved by coupling the Raman process to tunneling [5]. Regarding temporal resolution, although lots of efforts have been spent, there is so far no clear evidence of simultaneous fs and Å resolution [6-15].

In the Ho group, the first STM built in 1998 was to study photo dissociation of O2 molecules on Pt [6]. Later the group pushed STM induced light emission to submolecular resolution [4]. There were also efforts to achieve simultaneous fs- Å resolution [7, 8]. In this thesis, a tunable femtosecond laser was assembled; different colors of laser beams are aligned to overlap temporally and spatially; photoemission current is detected in the STM junction with a RF-STM.
2.2 STM4 Laser System Setup

In order to have the femtosecond laser interacting resonantly with the single molecule, the wavelength of the laser needs to be tunable in order to be able to excite the tunneling electrons into resonance with an excited state of the molecule. Since excited states are usually in the visible to the UV, we have coupled the near-IR output of the Mai Tai Deepsee Ti:sapphire laser to nonlinear crystals in Photop Harmonic Generator to generate second, third, and fourth harmonics, thus achieving continuous tunable femtosecond light output from 1040 nm down to 210 nm, with a gap between 520 and 690 nm. A delay stage and several other linear optical stages are used to get the temporal and spatial overlap of fundamental-fundamental, fundamental-2\textsuperscript{nd} harmonic and fundamental-3\textsuperscript{rd} harmonic. A BEOC-LPC is used to stabilize the laser power.

Fig2.1 is a diagram of the STM4 laser system, Fig2.2 is a photo of the STM 4 laser table, the inset shows the diagnostic path, Fig2.3 shows the powers at different wavelengths; all parts are listed in Appendix of this dissertation. As shown in fig2.1 and fig2.2, after the laser beam comes out from the Mai Tai, a periscope is used to change the beam polarization from horizontal to vertical, then a mirror pair is used to steer the beam into the LPC. Since the diameter of the incoming beam is required to be smaller than 4mm for the LPC to stabilize laser power, a lens pair is used to slightly reduce the beam size. After the LPC, a 30/70 beam splitter is used to divide the laser power, ~30\% of the beam will go through the fundamental path and 70\% of the beam will go into the harmonic generator to generate 2\textsuperscript{nd}, 3\textsuperscript{rd} and 4\textsuperscript{th} harmonics.

On the fundamental path, three mirrors are used to steer the beam into a Michelson Interferometer, a 50/50 beam splitter is used to get the same power of the two arms. One arm is a
Newport UBBR1-1S retroreflector mounted on a homemade piezo shake attached to a Newport UTS100CC delay stage, ensuring parallel of the outgoing beam to the incoming beam; the other arm is a mirror pair sits on a plate attached to a 3-direction stage. For the fundamental pump-probe experiments, we need both arms of fundamental beam. For the time resolved experiments using two different colors (like 2PPE), we only need the retroreflector-shaker-delay stage fundamental beam. Different from the former STM2 laser system setup, we have the delay stage and shaker in the same arm. The reason is that we need both fundamental+2nd and fundamental+3rd for time resolved experiments, thus not convenient to put either the delay stage or shaker on one of the 2nd or 3rd harmonic paths.

70% of the fundamental power goes into the harmonic generator to generate 2nd, 3rd and 4th harmonics. The horizontal-polarized 2nd harmonic is generated through second harmonic generation (SHG) by focusing the vertical-polarized fundamental beam through a 2nd harmonic BBO crystal; since we need vertical beam (polarized along the tip direction) to couple with STM to get maximum photon-induced signal, we use two periscopes to change the polarization, the 1st periscope changes the beam height to 3 inches and the 2nd periscope change the polarization (to vertical) and change the height back. From the beam splitter to the dichroic mirror (d1) which combines the fundamental and 2nd harmonic, we need to make sure the travel distance is the same so they can temporally overlap; first we need to roughly measure the distance, then we uses linear stages for fine adjustments. The vertical-polarized 3rd harmonic is generated through sum frequency generation (SFG) by focusing the horizontal polarized fundamental and 2nd harmonic beams through the 3rd harmonic BBO crystal, a wave plate is used to rotate the fundamental beam while keep the 2nd harmonic beam horizontal. The 3rd harmonic beam also travels the same distance as the fundamental to get the temporal overlap; same as 2nd harmonic path, a linear stage
is used to do fine adjustment. The vertical polarized 4th harmonic is generated through a SHG process by focusing 2nd harmonic beam through a 4th harmonic crystal, the 4th harmonic travels shorter than the other 3 beams since 4th harmonic is used only for one-photon photoemission experiments. The diameter of the harmonic beams is too big so we used a lens pair in each path to reduce the beam size.

2.3. Characterization and Simulation on the Optical Table

With the development of ultrafast lasers, different lab techniques, most notably the time-resolved two photon photoemission (2PPE) have been successfully studied the ultrafast charge carrier dynamics [15, 16]. However, 2PPE and other time-resolved techniques do not have the spatial resolution to study single molecules and atoms. Thus, combining laser techniques like 2PPE with STM could achieve the simultaneous temporal and spatial resolution in principle to study the ultrafast dynamics at single molecule level.

Before shining the laser beams into the junction, the beams can be characterized on the optical table. Especially for the pump probe experiments; it’s desirable to make sure the beams overlap spatially and temporally before shining into the junction. A flip mirror mount is used to direct the beam to the diagnostic path; the distance between the flip mirror mount and the nonlinear crystal on the optical path is about the same as the distance from the distance between the flip mirror mount and the STM junction. A 3.38” fused silica viewport and two fused silica windows are used to simulate the path to the STM junction. Same fused silica lens is used to focus the beam to the non-linear crystal.

After the 2 arms of fundamental beams combine on the dichroic mirror of the Michelson Interferometer, they travel to the chamber to shine into the STM junction; we flip m30 on the
optical path to direct the beams towards the diagnostic path to characterize the beam. The same 3” focal length lens is used to focus down the beams at the BBO crystal, which is mounted on the 3-direction linear stage for fine adjustments. 2nd harmonic beam (usually with green color) is generated through the BBO crystal, a color filter is used to filter out the fundamental residual. When the 2 arms of beams are spatially and temporally overlapped, we could see the fundamental beam blinking; the blinking of the 2nd harmonic after the BBO crystal is much more obvious. To quantitatively characterize the beam, a photodiode (from Thorlabs) is used to covert the 2nd harmonic laser signal to current, then voltage signal is obtained after the current goes through a terminal resistor (with different resistances, also from Thorlabs). The voltage signal can be fed into an oscilloscope or a DAQ, a LabVIEW program is used to record the voltage signal as the delay stage moves. An example of the autocorrelation data is shown in fig6; as could be seen on the top figure, if the polarization of both beams is vertical to the table, there will be significant interference. Such interference will cause tip crash in the tunneling regime even with very small laser power, however, photons need to couple with tunneling process to study the ultrafast dynamics at the single molecule level [7, 8]. One way to minimize the interference is to use one vertical polarized beam and one horizontal polarized beam (to optical table); the horizontal beam is got by using a waveplate and a polarizer (fig6). The interference gets much smaller, though there is still some especially around time zero.

Another better way to minimize the interference is to use different wavelength of laser beams. We can use fundamental and 2nd harmonic or fundamental and 3rd harmonic. Similar to fundamental-fundamental, we use the same fused silica lens to focus the beams down on a different nonlinear crystal. The orientation of the nonlinear crystal will be quite different from using 2nd harmonic to 3rd harmonic. 3rd harmonic beam is generated by focusing the fundamental
and 2\textsuperscript{nd} harmonic beam on the nonlinear crystal ("2+1"); we use a Schott glass filter to filter fundamental and 2\textsuperscript{nd} harmonic residual before the beam reaches the photodiode. 2\textsuperscript{nd} harmonic beam is generated by focusing the fundamental and 3\textsuperscript{rd} harmonic on the nonlinear crystal ("3-1"); the same Chroma filter for fundamental overlap is used to filter fundamental and 3\textsuperscript{rd} harmonic before the beam reaches the photodiode. The 2\textsuperscript{nd} harmonic and 3\textsuperscript{rd} harmonic are weaker than the fundamental; sometimes a CCD camera is needed to detect the even weaker beam after the nonlinear crystal and filter.

2.4. Alignment of Laser Beams into STM Junction

After characterization on the optical table, the laser beams are ready to shine into the junction. To align the laser to the junction, a pair of plano-convex lenses was mounted on two separate translation-and-tilt stages inside the UHV chamber. The laser incident and exit angles are 45\degree from the surface normal. One of the lenses was used to focus the laser into the junction; the other was used to image the STM junction onto a CMOS or CCD camera. When the laser was aligned to the junction, the STM tip was shown on the bottom in and its mirror image appeared on the top due to the high reflectivity of the sample. The focal spot in the junction was estimated to be around 50 \mu m in diameter. The laser electric field was linearly polarized in the plane of incidence (p polarized) to maximize the laser-induced signal [17]; except for changing polarization of one arm of fundamental to minimize the interference when 2 fundamental beams are used. Fig2.7 (B) is a CMOS tip-mirror image with the fundamental in the junction while fig2.7(C) is a CMOS tip-mirror image with the 4\textsuperscript{th} harmonic in the junction, both were taken on the laser outgoing side.
To set up the new system, the initial alignment to the junction could be very tricky. First tip-mirror image should be clearly seen on the bottom mirrors of the periscope on both the beam incoming and outgoing side by shining white night from the 8” front window, and the image should be focused and in the center of view, it also depends on which angle we view from (fig2.8 A). Then the tip-mirror image could be got on a CCD/CMOS at the laser outgoing side, but due to the 2D nature of our two lens system, even when the beam seems at the junction as shown in fig2.7 (B) and (C), it may hit somewhere on the sample and then reflects out (eg, point e in fig2.9A). We can see the sample from the pyrometer viewport with the help of a finder scope, the CMOS tip-mirror image got from here is 3D, and when the beam seems at the junction on the image, it is actually in the junction. Since no commercial finder scope can see 3rd or 4th harmonic (220nm~330nm), a homemade finder scope made of fused silica lenses was assembled to help seeing the UV beams from the pyro viewport (fig2.8 D).

If the two p-polarized fundamental is in the junction or along the tip mirror axis, we can clearly see the STM Z feedback voltage oscillation, both in and out of tunneling regime due to the interference (fig2.9), multiphoton current can also be detected, as shown in fig2.10 A. Even if the polarization of the fundamental beams is different, we can still clearly see more multiphoton signals around time 0 if the two arms of fundamental overlap on the tip-mirror axis.

Although we can easily and reproducibly overlap two fundamental beams temporally and spatially in the STM tip-sample junction; it is very hard for us to align two different colors (fundamental + 2nd harmonic “1+2” or fundamental+3rd harmonic “1+3”) to the junction. The main reason is that we have a lens in the UHV chamber to focus the laser beam to the STM junction (fig2.7A); the refractive index for different wavelengths is different and it will change the beam overlap unless the beams hit the center of the lens. The beam path is very hard to check.
or simulate given that the lens is in UHV chamber. It’s also hard to overlap the 2 different color beams with CCD/CMOS from the laser outgoing viewport or pyro viewport, given that the lens of the finder scope will change the overlap. For the 2PPE setup, a curved mirror outside the UHV chamber is used to focus the beam on the sample, so as long as the 2 beams overlap on the diagnostic path, they overlap on the sample in the UHV chamber [15, 16]. 2PPE also has the hemispherical energy analyzer to collect the photoemitted electrons [15, 16]; it’s probably more sensitive than the STM tip/sample to collect electrons.

### 2.5. Laser Induced Current Detection

To reach sub-Ångström spatial limit in optical experiments of single molecules, previous studies have been shown that it is necessary to couple light to the tunneling process [8]. The difficulty in such experiments is that the rate of production of photoinduced tunneling electrons is very low in relation to the background of bias-induced tunneling electrons in the operation of the STM. One way to detect the laser-induced current is through the DC rectified current. The multiphoton current shown in figure.10 is the DC current amplified by the DC preamplifier used for DC bias induced tunneling current, such current could not be detected in the tunneling regime given that certain laser power is needed to detect the current, and such a laser power will make the tunneling junction unstable.

We are using a new method by detecting the electrons produced at the repetition rate of the femtosecond laser (~80 MHz), which provides an intrinsic modulation of the photon emitted or photon-induced tunneling electrons (fig2.11). We were using an external bias tee to separate the DC and RF signal, a 50Ω terminal resistor was used as the RF current to voltage converter. The RF signal will go to the 200MHz RF lock-in (or spectrum analyzer) through the RF amplifier for detection. Fig2.12 shows the power spectrum with 220nm in the STM junction of
Ag tip and Cu (110) surface. The 79.7MHz peak is at the repetition rate of the Mai Tai Deepsee laser, illustrating the feasibility of detecting the electron signal created by each laser pulse by lock-in detection at the repetition rate of the oscillator. Further discrimination is achieved by connecting the output of 200 MHz lock-in amplifier to a second lock-in amplifier that detects a lower frequency signal generated by modulating the sample bias, for $dI/dV$ and $d^2I/dV^2$ spectroscopy. This double lock-in approach is shown in Figure 2.11 (A), the RF current and $dI_{RF}/dV$ detected is shown in fig2.13. However, these measurements are performed with tip retracted. While the bandwidth of the terminal resistor is essentially infinite, the gain is greatly compromised, making it not sensitive enough to go to tunneling to get detectable laser-induced RF or DC signal.

To improve, a new cryogenic amplifier can be installed to get better signal/noise ratio, a fabricated tip can be used for better plasmonic enhancement. We can also use two different color beams for pump-probe. Schematic layout of the 80 MHz Laser-STM showing the detection of the 80 MHz component of the $dI/dV$ and $d^2I/dV^2$ signals due to changes in the STM junction induced by time delayed pulse pairs.

### 2.6 Conclusions

A tunable fs laser (210nm~1040nm with a gap from 520nm~690nm) system has been set up by using a Mai Tai Deepsse laser (690~1040nm, ~100fs) and a Photop Haramonic Generator to generate 2$^{nd}$, 3$^{rd}$ and 4$^{th}$ harmonics. 4$^{th}$ harmonic (~220nm) can be used in 1-photon photoemission and deep UV laser-induced chemistry. The fundamental, 2$^{nd}$ harmonic and 3$^{rd}$ harmonic can be used for plasmon enhanced chemistry and time-resolved studies. Fundamental and fundamental (“1+1”), fundamental and 2$^{nd}$ harmonic (“1+2”), fundamental and 3$^{rd}$ harmonic
(“1+3”) are temporally and spatially overlapped on the optical table diagnostic path. Fundamental and fundamental can be reproducibly aligned to overlap in the STM junction although causing significant optical interference. Fundamental and 2\textsuperscript{nd}, fundamental and 3\textsuperscript{rd} could not be reproducibly aligned into the STM junction due to the lens inside the UHV chamber. The laser-induced signal is still too noisy even the double lock-in technique is used to detect the weak laser-induced signal in the tunneling regime. To improve the signal to noise ratio, a cryogenic amplifier can be used. A fabricated Ag, Au or Cu tip can also be used to improve the plasmonic enhancement.

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Fig 2.1. STM 4 fs laser system layout (m stands for mirror, d stands for dichroic mirror, TS stands for translational stage, LPC stands for laser power controller; for a complete list of parts, refer to A.1 catalog of Appendix A in this dissertation).
Fig2.2. Photo of the STM4 laser system layout, (A) photo of the whole laser system, with the key parts labeled, (B) the diagnostic path.
Fig2.3. STM 4 fs laser power spectrum
Fig 2.4. autocorrelation data obtained on the diagnostic path, the laser wavelength is 800 nm. (A) without wave plate or polarizer, (B) with wave plate and polarizer, so the polarization of the two fundamental beams is 90 degrees away and the interference is minimum.
Fig 2.5. “1+2” autocorrelation data obtained of on the diagnostic path, the fundamental is 840nm and the 2\textsuperscript{nd} harmonic is 420nm.
Fig 2.6. “3-1” autocorrelation data obtained on the diagnostic path, the fundamental is 840nm and the 3rd harmonic is 280nm.
Fig2.7. (A) optical path through the STM junction, (B) CMOS camera tip mirror image from outgoing viewport with the fundamental (780nm) shining into the junction, (C) CMOS camera tip mirror image from outgoing viewport with the 4\textsuperscript{th} harmonic (220nm) shining into the junction
Fig2.8. (A) tip-mirror image on the bottom mirror of the periscope, (B) sample and tip from pyro viewport, (C) tip mirror image from pyro viewport with the CMOS camera, the beam is 2\textsuperscript{nd} harmonic 420nm, (D) homemade finderscope made of fused silica lens to see 3\textsuperscript{rd} and 4\textsuperscript{th} harmonic beams in junction.
Fig 2.9. (A), different beam position around the junction, Z-oscillation or multiphoton current can be detected when the beam is along the tip-mirror axis (eg, a, b and c), the oscillation or current won’t be detected when the beam hit other place (eg, c and d); (B) Z oscillation out of tunneling, (C) Z oscillation when tunneling.
Fig 2.10. (A) multiphoton current detected in the STM junction, 2 fundamental beams are both p-polarized, (B) multiphoton current detected in the STM junction, one fundamental beam is p-polarized the other one is s-polarized.
Fig2.11 (A) Schematic of experimental arrangement combining a tunable femtosecond laser system with a low temperature RF-STM. The uniqueness of the instrument lies in the lock-in detection of laser induced current due to each laser pulse of a repetitive pulse train at 80 MHz. Laser induced RF current was detected with a bias tee and terminal resistor; (B) laser induced RF current detection with a UHV low temperature ~80MHz amplifier with Q=100 and sensitivity of 20,000V/A.
Fig2.12 (A) Noise spectrum with the center frequency set at 79.5 MHz of the spectrum analyzer. Spectrum obtained by blocking the femtosecond laser. (B) Power spectrum of the photoemitted electron signal induced by 230 nm femtosecond laser irradiating the junction. The peak of photoemitted electron pulses are seen at 79.7 MHz, with the spectrum analyzer’s center frequency set to this frequency, the repetition rate of the femtosecond oscillator.
Fig 2.13 Laser-induced DC and RF current detection with the double lock in technique, the junction is formed by a Ag tip and Cu (110) surface, 220 FHG beam shines into the junction at 2.5mW power out of tunneling, (A) photo-induced electron from tip to sample at positive sample bias, (B) $I_{DC}$ detected by the DC lock-in (black), $I_{RF}$ detected by the RF lock-in (blue), $dI_{RF}/dV$ by the double lock-in technique (red).
(A)

vacuum
Ag tip Cu(110)

(B)

Cu(110) FHG (220nm) P=800mW Vmod=420mV T=84K 030312

$V = 1.12V$
Bibliography:


Chapter 3

Imaging Molecular Orbitals of 4, 7-Di ([2, 20-bithiophen]-5-yl) benzo[c] [1, 2, 5] thiadiazole (4T-BTD) with a STM

3.1 Abstract

STM imaging and spectroscopy are performed to study an organic semiconductor molecule 4, 7-Di ([2, 20-bithiophen]-5-yl) benzo[c] [1, 2, 5] thiadiazole (4T-BTD) on NiAl (110) surface. The STM topography image shows bias dependence, the benzo[c][1, 2, 5] thiadiazole could only be clearly observed at positive biases near lowest unoccupied molecular orbital (LUMO) peak. LUMO and HOMO (highest occupied molecular orbital) dI/dV spectroscopy and imaging show the structure and electronic properties of the molecule at the single molecule level. Inelastic electron tunneling spectroscopy (IETS) imaging at the hydrogen rotational energy shows the sub-Å details of the LUMO orbital of the molecule.

3.2 Introduction

The scanning tunneling microscope (STM) has been a powerful tool to study single-molecule chemistry by imaging and electronic spectroscopy [1]. Despite these successes, various efforts have been made to extend the structural and chemical sensitivities of STM. In 1998, Seventeen years after the invention of STM, STM inelastic electron tunneling spectroscopy (STM-IETS) first achieved chemical identification of molecular species by characterizing vibrational energies [2]. Only recently, scanning tunneling hydrogen microscopy (STHM) [3], inelastic tunneling probe (itProbe) [4] and hydrogen rotational spectromicroscopy [5, 6] achieved chemical and
structural imaging of single molecules with unprecedented details. Such great-detail chemical structures have also been observed with a noncontact atomic force microscope (NC-AFM) [7].

4, 7-Di ([2, 20-bithiophen]-5-yl) benzo[c] [1, 2, 5] thiadiazole (4T-BTD) is a semiconductor benzothiadiazole molecule which has potential applications in organic light emitting diodes (LEDs), organic field effect transistors (FETs) and solar cells [8, 9]. 4T-BTD has donor-acceptor-donor sites inside a single molecule, enabling intramolecular electron transfer; 4T-BTD and its small molecular semiconductor relatives have received great attention due to their advantages versus polymer semiconductors: ease of purification, higher charge mobility and overall better performance [10, 11]. Electronic properties of such organic molecules are of central importance to applications, understanding the properties of the conjugated building blocks will help building blocks to extended semiconducting molecules with predictable properties, and ultimately understanding and controlling the collective behavior of multiple components in functional devices [12]. Structure and electronic properties of these molecules can be probed by the STM at the single molecule level and with sub-Å spatial resolution [4, 5, and 6].

The rotational excitations of hydrogen molecule weakly adsorbed on surfaces have been measured with the STM [13, 14]. It was also shown that similar to the vibrations of a CO molecule attached to the tip [4], changes in the adsorption potential well induced by varying the tip-sample distance shift the energies in both vibrational and rotational modes for the hydrogen temporally trapped in the tunnel junction [13]. The sensitivity of the H$_2$ modes to different chemical environments has been applied to achieve sub-Å structural and molecular orbital imaging of MgP on metal and oxide surfaces [5, 6].

3.3 Experiment
The experiments were performed using a home-built STM operating at 12.5 K and a base pressure of $3 \times 10^{-11}$ Torr [15]. The preparation of NiAl (110) follows previously reported procedures [16]; the NiAl sample was sputtered with Ne ions and annealed at ~1350K. The Ag tip was electrochemically etched and in situ cleaned by a few sputter-anneal cycles. The clean surface at 12.5 K was dosed in situ with 4T-BTD. As hydrogen molecules have limited residence time on the NiAl(110) surface at 12.5 K, a background pressure of H$_2$ is kept at $2 \times 10^{-10}$ Torr throughout the experiment to maintain a constant population of adsorbed molecules on the surface following an initial dose at $5 \times 10^{-10}$ Torr for 5 min. A bias voltage is applied to the sample with the tip at virtual ground.

### 3.4 Results and Discussion

STM topography and spectroscopy are performed to study the 4T-BTD molecule on the NiAl (110) surface in the absence of H$_2$. The molecules show bias dependence in the topography image. The electron poor benzothiodiazole fragment, which acts as the acceptor unit, can only be clearly imaged at positive biases close to the LUMO energy. Fig 2.1 A is taken at sample bias 1V and the benzothiodiazole fragment is clearly seen, while the benzothiodiazole fragment could not be cleanly imaged in fig 2.1E at sample bias 0.5V. Electronic dI/dV spectrum are performed on a few points of the molecule from 0.8V to 1.8V, the background spectrum on NiAl (110) is flat at this bias range (fig 2.1D, 2.2D and 2.3D BKGD spectrum). There are LUMO peaks on different points of the molecule, although the peak energies of the spectra at different points of the molecules are the same around 1.25V, they show different intensities, with the benzothiodiazole fragment has the strongest peak and two electron rich bithiophen units (donor parts) have the weakest peak. During the spectrum on point D of fig 2.1A, the molecule rotates to a different orientation as shown if fig 2.1E, showing the STM-induced rotational motion of this big 4T-BTD
molecule and the possibility of building blocks to extended semiconducting molecules by STM [12].

Spectroscopy LUMO dI/dV imaging is performed to further study the molecular orbitals in the absence of H₂. The dI/dV imaging is taken at 1.2V, near the LUMO peak, dI/dV signal is recorded at each point during the scan. The image is taken at constant current mode (feedback is on during moving the tip and is off during taken the dI/dV signal, fig 2.2B, 2.3B and 2.4B) or constant height mode (tunneling gap is set on the first point and feedback is off during the scan, fig 2.2C, 2.3C and 2.4C). Electronic dI/dV spectra are also taken on various points on the molecules. From the spectroscopy imaging; there are two kinds of LUMO geometries of molecules (fig 2.2 and 2.3). Both kinds have much stronger LUMO dI/dV signal in the center acceptor unit than in the two side donor units (fig 2.2 B, C and fig 2.3B, C). The first LUMO geometry is shown if fig 2.2, there is a dip (around point B in the topography image fig2.2A) in the benzothiodiazole fragment between the high dI/dV signal [1, 2, 5] thiadiazole and benzene ring parts, both in the constant current and constant height dI/dV image. This dip is confirmed by the almost flat dI/dV spectrum on point B of fig 2.2A (fig 2.2D). In the constant current mode, the thiodiazole unit has the strongest signal, there are also side features in the donor parts; while in the constant height mode, the benzene ring part has the strongest dI/dV signal and there are no side features. There is no dip in the second LUMO geometry (fig 2.3B and C), suggesting more hybridization in these molecules, there are no obvious side features in the constant current dI/dV imaging (fig 2.3B). Same as in the first LUMO geometry, the thiodiazole unit has the strongest signal in the constant current mode dI/dV imaging (fig 2.3B) and the benzene ring part has the strongest dI/dV signal in the constant height mode (fig 2.3C). Point dI/dV spectra are also performed on this molecule, different from the molecule in fig 2.2, the points near the center of
this molecule (4, 5, 6, 7 a, b and c) all have strong peak near 1.2V, with points a and 5 have the strongest dI/dV signal.

Calculations show that 4T-BTD molecule has a HOMO-LUMO gap of 2.31eV [11]. The LUMO peak is about 1.2V (fig 2.1, 2.2 and 2.3), thus the HOMO peak should be around -1.1V. Since the NiAl (110) background has some electronic states at the negative bias range -1.4V~-0.5V (fig 2.4D BKGD spectrum), the HOMO dI/dV spectra and images are not as clear as the LUMO ones. However, the electric properties and HOMO geometry can still be studied by dI/dV spectra and HOMO dI/dV imaging (fig 2.4). The molecule shown in fig2.4 is the same one as in fig2.3; the benzothiodiazole unit is not clearly imaged in the topography fig 2.4A at V_B=-1.1V. The constant current dI/dV image highlights the donor units on the two sides of the molecule, especially the top unit (fig 2.4B). The donor units are not so obvious in the HOMO constant height image (fig 2.4C). In both the constant current and constant height dI/dV images, the electron poor [1, 2, 5] thiadiazole are not observed. As can be seen from the point spectra in fig 2.4D, the -1.1V peak is only obvious in the top parts of the molecule (points 1 and 2), spectra on the other points are very similar to the background spectrum.

The STM tip and surface (or molecules absorbed on the surface) form a nanocavity and molecular H_2 can be trapped, the rotational and vibrational modes of the trapped H_2 can be characterized by STM IETS [13, 14]. After introducing H_2 into the UHV chamber, IETS spectra are performed on different points of the molecule to study the interaction between H_2 molecule and the 4T-BTD molecule. While the H_2 IETS spectrum on the background is nearly featureless, the H_2 IETS spectra on different points of the molecule clearly show strong features. H_2 is an electron donor [5], 4T-BTD molecule is an electron acceptor here and the LUMO orbital of the molecule readily accepts electron from H_2. Different from the IETS spectra of H_2 trapped
between tip and MgP molecule on Au (111) and NiAl (110) surfaces, on which both H₂ vibrational and rotational modes could be clearly observed [5, 6]; the IETS spectra of H₂ molecule trapped between tip and the 4T-BTD molecule shows very strong \( j=0 \rightarrow 2 \) rotational mode but the \( v=0 \rightarrow 1 \) vibrational mode signal is much weaker compared to the rotational mode (fig 2.5). The rotational mode position varies from about 37mV to 42mV on the eight points of the molecule; the intensities on the points are also different. These differences suggest the trapped H₂ experience spatially dependent interactions with the underlying 4T-BTD molecule.

Constant current H₂ rotational IETS images (feedback is on during moving the tip and is off during taken the \( d²I/dV² \) signal) are taken based on the sensitivity of H₂ interacting with the 4T-BTD molecule. Surprisingly, LUMO of 4T-BTD are clearly imaged on the NiAl (110) surface at \( V_B=37.1 \)mV, \( I_T=0.4 \)nA, despite the strong electronic interaction of the molecule and metal substrate. As shown in fig 2.6 B and C, the H₂ rotational IETS imaging matches really well with the calculation from ref.11, except the IETS image is slightly not symmetric probably due to the molecule is slightly tilted as shown in the topography image fig 2.6A. In contrast, the H₂ rotational imaging of MgP can only observe the submolecular electronic structure on the oxide surface [5, 6].

Since the spectra show different peak positions on different points of the molecule (fig 2.5C), H₂ rotational IETS images are taken at different biases; these images highlight different parts of the molecule, fig 2.6D highlights the LUMO geometry on the left side of the molecule and fig 2.6E highlights the LUMO geometry of the right side of the molecule. In contrast, the IETS image taken at \( V_B=60.1 \)mV, \( I_T=0.65 \)nA (far away from H₂ rotational mode) is featureless (fig 2.6F).
3.5 Conclusions

STM imaging and spectroscopy are performed on 4T-BTD molecule, a molecule that has intramolecular donor-acceptor-donor sites. The electron poor benzothiodiazole fragment in the acceptor site can only be clearly imaged at bias near the LUMO peak. LUMO dI/dV spectra clearly show that the center acceptor unit has higher peaks, LUMO dI/dV imaging more clearly highlights the center acceptor part. In contrast, HOMO dI/dV spectra weakly shows the two side donor units have stronger peak and the HOMO imaging confirms this. H₂ rotational IETS imaging is performed on the molecule, LUMO structure of the molecule are more clearly imaged at sub-Å resolution, which is the first time observe sub-Å details of the molecular orbital on the metal surface by using H₂. Such an understanding of this kind of molecules will help building blocks to extended semiconducting molecules with predictable properties.

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Fig 3.1 (A) 29Å ×29Å constant current STM topographic image of a 4T-BTD molecule on NiAl (110) surface, the tunneling gap is set with $V_B = 1.0V$ and $I_T = 0.1nA$ and the temperature is at 12.5K, (B) diagram of this 4T-BTD molecule, (C) ball-stick model of the molecule, (D) Tunneling spectra taken on different points of the 4T-BTD molecule in (A), the tunneling gap is set with $V_B = 1V$ and $I_T = 0.1nA$, (E) 29Å ×29Å constant current STM topographic image of a 4T-BTD molecule on NiAl (110) surface, the tunneling gap is set with $V_B = 0.05V$ and $I_T = 0.1nA$, the molecule rotates after doing spectroscopy on point D in panel (A), also the “benzothiazole fragment” is not so obvious as with $V_B = 1V$. 
Fig 3.2: (A) 29Å ×29Å STM topography image taken at $V_B=1V$, $I_T=60pA$, (B) constant current $dI/dV$ image taken at $V_B=1.2V$, the tunneling gap is set at $V_B=1V$, $I_T=60pA$, (C) constant height $dI/dV$ image taken at $V_B=1.2V$, the tunneling gap is set at $V_B=1V$, $I_T=150pA$, (D) $dI/dV$ spectra taken at different points of the molecule, the tunneling gap is set at $V_B=1V$, $I_T=60pA$. 
Fig 3.3: (A) 29Å × 29Å STM topography image taken at $V_B=1.2\, \text{V}$, $I_T=50\, \text{pA}$, (B) constant current $dI/dV$ image taken at $V_B=1.2\, \text{V}$, $I_T=50\, \text{pA}$, (C) 29Å × 29Å constant height $dI/dV$ image taken at $V_B=1.2\, \text{V}$, $I_T=50\, \text{pA}$, (D) $dI/dV$ spectra taken at different points of the molecule, the tunneling gap is set at $V_B=1.2\, \text{V}$, $I_T=50\, \text{pA}$. 
Fig 3.4: (A) 29Å ×29Å STM topography image taken at $V_B=-1.1V$, $I_T=60pA$, (B) constant current $dI/dV$ image taken at $V_B=-1.1V$, $I_T=60pA$, (C) 29Å ×29Å 29Å ×29Å constant height $dI/dV$ image taken at $V_B=-1.1V$, $I_T=60pA$, (D) $dI/dV$ spectra taken at different points of the molecule, the tunneling gap is set at $V_B=-1.07V$, $I_T=100pA$. 
Fig 3.5: (A) 30.5Å ×30.5Å STM topography image taken at $V_B=37.1\text{mV}$, $I_T=0.4\text{nA}$, (B) IETS image at $V_B=37.1\text{mV}$, $I_T=0.4\text{nA}$, (C) IETS spectra taken at different points of the molecule, the tunneling gap is set at $V_B=38\text{mV}$, $I_T=400\text{pA}$.
Fig 3.6: (A) 30.5Å ×30.5Å STM topography image taken at $V_B=37.1$ mV, $I_T=0.4$ nA, (B) IETS image at $V_B=37.1$ mV, $I_T=0.4$ nA, (C) 30.5Å ×30.5Å IETS image taken at $V_B=44.9$ mV, $I_T=0.48$ nA, (D) 29Å ×29Å IETS image at $V_B=-39.1$ mV, $I_T=0.65$ nA, (E) 30.5Å ×30.5Å IETS image at $V_B=34.9$ mV, $I_T=0.45$ nA, (F) 29Å ×29Å IETS image at $V_B=60.1$ mV, $I_T=0.65$ nA.
Bibliography


Chapter 4

Imaging Metal-Molecule Chain and Single Molecule of 1, 4-Phenylene Diisocyanide on Noble Metal Surfaces and Their Interaction with Small Molecules

4.1 Abstract

1, 4-Phenylene Diisocyanide (PDI) forms 1-D nanochains on Au (111) surface, IETS spectra are performed on the CO$_2$ adsorbed on the chain. PDI forms chains on Cu (110) surface at room temperature, CO$_2$ also prefers to adsorb on the chains even the Cu-PDI chains break after cooling down to 12.5K. The single PDI molecule, which can capture small molecules like H$_2$, is also studied on Cu (110) and Ag (100) surfaces. Imaging of the single molecule with H$_2$ and CO shows sub-Å details of the molecule. On the Cu (110) surface, at low H$_2$ coverage, the H$_2$ IETS image highlights the two N atoms in the molecule; at high H$_2$ coverage, the topography image when the tip is close shows the molecule structure and the interaction of molecule with the surface, the more detailed ring structure and molecule interaction with the substrate are observed in the IETS image. H$_2$ IETS spectra are taken on different points of the molecule, showing H$_2$ sensitivity of interaction with the molecule. Moreover, CO adsorbed on the Ag (100) surface can be picked up by a Ag tip and high resolution images of single PDI molecule is also got with the Ag-CO tip.

4.2 Introduction

The scanning tunneling microscope (STM) has been a powerful tool to study single-molecule chemistry by imaging and electronic spectroscopy [1]. Despite these successes, various efforts
have been made to extend the structural and chemical sensitivities of STM. STM inelastic electron tunneling spectroscopy (STM-IETS) first achieved chemical identification of molecular species by characterizing vibrational energies [2]. Only recently, scanning tunneling hydrogen microscopy (STHM) [3], inelastic tunneling probe (itProbe) [4] and hydrogen rotational spectromicroscopy [5, 6] achieved chemical and structural imaging of single molecules with unprecedented details. Such great-detail chemical structures have also been observed with a noncontact atomic force microscope (NC-AFM) by using a CO-terminated tip [7].

1, 4-Phenylene Diisocyanide (PDI) is a unique molecule that can self-assemble into nanostructures on noble metal Au [8-11], Cu [12] and Ag [13] surfaces. The negatively charged carbon atom in the two N-C groups readily to donate electrons to noble metal atoms on the surface and form nanostructures like 1D nanochains [9, 10], which are 1-D conductor. Moreover, the nanochains can chemically and physically absorb small molecules like carbon dioxide (CO₂) [11, 12]. When linked between nanoparticles [14, 15] or formed layers on surface [16], PDI molecules show interesting electrical and plasmonic properties, making it a potential candidate for molecular electronics. Structure and electronic properties of the PDI molecule can be probed by the STM at the single molecule level and with sub-Å spatial resolution [4, 5, and 6].

The vibrational and rotational excitations of hydrogen molecule weakly adsorbed on surfaces have been measured with the STM [17, 18]. It was also shown that similar to the vibrations of a CO molecule attached to the tip [4], changes in the adsorption potential well induced by varying the tip-sample distance shift the energies in both vibrational and rotational modes for the hydrogen temporally trapped in the tunnel junction [17]. The sensitivity of the H₂ modes to different chemical environments has been applied to achieve sub-Å structural and molecular orbital imaging of MgP on metal and oxide surfaces [5, 6]. With a CO terminated tip,
even higher resolution STM itProbe [4] and NC-AFM images [7] have been achieved. Here we focus on the sub-Å imaging and spectroscopy of single PDI molecule with H₂ and CO, the understanding of single PDI sub-molecular should help the understanding the self-assembled structure, its electronic properties and the interactions with nanoparticle and small molecules.

4.3 Experiment

The experiments were performed using a home-built STM operating at 12.5 K and a base pressure of 3×10⁻¹¹ Torr [19]. The Au (111), Cu (110), Ag (100) sample sputtered and annealed to about 520°C. The Ag tip was electrochemically etched and in situ cleaned by a few sputter-anneal cycles. PDI molecules are dosed onto Au (111) and Cu (110) at room temperature to form chains and then cooled down to 12.5K for low temperature study. PDI molecules are dosed onto the clean Cu (110) and Ag (100) surfaces at 12.5 K for single-mole studies. H₂ molecules were introduced into UHV chamber for imaging on Cu (110) surface, as hydrogen molecules have limited residence time on the Cu (110) surface at 12.5 K, for the low H₂ coverage imaging, a background pressure of H₂ is kept at 5×10⁻¹⁰ Torr throughout the experiment to maintain a constant population of adsorbed molecules on the surface following an initial dose at 1×10⁻⁹ Torr for 5 min. For the high H₂ coverage imaging, a background pressure of H₂ is kept at 2×10⁻⁹ Torr. CO is evaporated onto the Ag (100) surface and picked up by the Ag tip to image the single PDI molecule. For all these experiments, a bias voltage is applied to the sample with the tip at virtual ground.

4.4 Results and Discussion

At room temperature, PDI molecules are evaporated onto Au (111) surface following reported recipes [8, 10, and 11]; Au-PDI chains are formed on the surface. Then the chains are cooled
down to 12.5K for imaging and absorb CO₂. The chains are consisted by alternating PDI molecule and Au atoms, the bigger dots are PDI molecules and the smaller ones are Au atoms (fig 4.1 A and B). Both PDI and Au has a ~2V peak in the dl/dV spectrum, showing the chain is a 1D conductor (fig 2.1C). CO₂ is evaporated onto the chain and IETS are performed on the CO₂, showing two modes at ~20mV and ~50mV. PDI molecules are also evaporated onto the Cu (110) surface at room temperature to form chains, the chains break after cooling down to 12.5K, possibly because Cu atom takes off the CN groups and sink into the Cu surface (fig 4.2 A and B). CO₂ still prefers the chains even the Cu atoms are missing (fig 4.2C). IETS are also performed on the CO₂, showing two modes at ~20mV and ~2.5mV (fig4.2D).

Single PDI molecules are evaporated onto the Cu (110) surface at 12.5K, at this temperature, the molecules freeze and chain structures are not formed. The STM tip and surface (or molecules absorbed on the surface) form a nanocavity and molecular H₂ can be trapped, the rotational and vibrational modes of the trapped H₂ can be characterized by STM IETS [17, 18]. After introducing 5×10⁻¹⁰ Torr H₂ into the UHV chamber, IETS spectra are performed on different points of the molecule to study the interaction between H₂ molecule and the PDI molecule. While the H₂ IETS spectrum on the background is nearly featureless, the H₂ IETS spectra on different points of the molecule clearly show strong features (fig 4.4C). Single molecules are imaged on the Cu (110) surface by H₂ topography and IETS imaging at this lower H₂ coverage. The topography image only show two dimples near the two N-C groups, same as the topography image without H₂(Fig 4.3A and B), but the H₂ IETS imaging clearly resolve the submolecular structure of the PDI molecule (fig 4.3 C). The two bright spots in fig4.3C (more clearly 3 and 5 in fig 4.4B, which is the same IETS image) are ~6Å apart, suggesting they are two N atoms in the PDI molecule, which are about 6Å away, Making the two weak spots near N
(1 and 6 in fig 4.4B) probably C atoms. From the point spectra in fig 4.4C, the two N (3 and 5) have very similar spectra; the two C (1 and 6) also have similar spectra, except the peak intensities are different. The spectrum on transition point 2 shows transitional features from the spectra on point 1(C) to point 3(N). H₂ IETS spectrum on the center of the molecule (point 4) is almost the same as on the BKGD, showing the center of the ring does not interact with H₂ at this coverage. Six spots nearby (7 and 9 are listed as examples here) can also trap H₂, the spectra on 7 and 9 are almost the same and they are similar to the spectra on the N sites (3 and 5). There are some gaps between N and the trapped state (point 8 with spectrum shown is an example here) which do not attract H₂. All the H₂ IETS spectra (except on BKGD, points 4 and 8, which do not attract H₂) show the ~40mV \( j=0 \rightarrow 2 \) rotational mode, although the peak position and intensity are different. Different from the H₂ IETS spectra on MgP [5, 6] and 4T-BTD molecules [20], the \( \nu=0 \rightarrow 1 \) vibrational peaks on the PDI molecule are strong. H₂ rotational and vibrational IETS images are taken at different biases, the image shown here (fig 4.3C or fig 4.4B) is taken at the 5mV vibrational mode and shows most features of the molecule.

After the imaging and spectra at low H₂ coverage, more H₂ is introduced to the UHV chamber to do Scanning Tunneling Hydrogen Microscopy (STHM) [3] and H₂ IETS image. At this high H₂ coverage, three parts of the molecule (the benzene ring and two N-C groups) can be clearly resolved in the topography image when the tip is close (fig 4.5B), such submolecular details could not be resolved when the tip is far away (fig 4.5A). The tip-close topography image (fig 4.5B) also shows there is interaction between the PDI molecule and Cu (110) substrate. The H₂ IETS image at the -8.8mV H₂ vibrational mode shows more clearly the ring structure in the center, two N-C groups, and the interaction with the substrate (fig 4.5C and 4.6B). H₂ IETS spectra are also performed on different points of the molecule at this high H₂ coverage, generally
at the same tunneling set point the signals are stronger than the spectra on molecule at low H$_2$ coverage (fig4.4C), spectra on only one N-C group is shown (1, 2, 3 and 4 on fig 4.6B) since the spectra on the other group are symmetric. As seen in fig 4.C, spectrum on the Cu (110) background already shows strong IETS features at such high coverage, including the rotational modes and vibrational modes, showing the Cu surface starts to attract H$_2$ at high coverage. The H$_2$ rotational and vibrational modes are also observed in the center benzene ring of the molecule, although the signal is weaker than most points on the molecule and even the background. The points above and below the molecule (6 and 7) also have weaker H$_2$ signals than the background. The points on or near the N-C groups (1, 2 3 and 4) all show strong H$_2$ IETS signals, although the peak position and intensity are very different, showing strong H$_2$ sensitivity over these N-C groups. Such high sensitivity is confirmed in the IETS images (fig 4.5C or 4.6B); very sharp transitions could be imaged, most notably around point 3 in fig4.6B. IETS images at different biases highlight different parts of the molecule (fig 4.7); while vibrational modes images highlight (fig 4.7A and C) the CN groups of the molecule, rotational modes also highlight the interactions around the molecule, especially between the CH groups and substrate (fig 4.7B and D).

Single PDI molecules are studied also studied with a CO terminated tip on the Ag (100) surface, CO is picked up by the Ag tip from substrate by scanning close and is confirmed by IETS spectra showing CO vibrational modes at ~4mV and ~19mV on the background. CO in the topography image looks like a protrusion instead of a dip (fig 4.8A); also indicating the CO on surface is imaged with a CO tip [4]. STM topography image of the PDI molecule when the CO tip is close show similar structure with H$_2$ imaging, the dip on the two N-C groups and the interaction between molecule and substrate (fig 4.8 B and D). These CO tip topography images
suggesting that the interaction features around CH groups (fig 4.8B, D and fig 4.5B) are not from the interaction with CO or H₂, but from the CH groups on the molecule and substrate, probably H bonds.

4.5 Conclusions

1, 4-Phenylene Diisocyanide (PDI) forms 1-D nanochains on Au (111) surface, STM-IETS is performed on the adsorbed CO₂. PDI also forms chains on Cu (110) surface, even though the chains break down at 12.5K, CO₂ still prefers to go to the chains and STM-IETS is performed. Single PDI molecules are studied on Cu (110) surface with H₂ imaging. At low H₂ coverage, while the STM topography image only shows two dimples which may be the N-C groups, the STM H₂ IETS image clearly shows the two N sites. At high H₂ coverage, the topography image already shows three parts of the molecule with interactions with the surface, the IETS image shows more details of the molecule and its interaction with the substrate, with better contrast and sharper transitions.

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Fig 4.1 Au-PDI chains on the Au (111) surface, (A) 112Å ×112Å STM topography image taken at $V_B=1V$, $I_T=0.1nA$, (B) 56Å ×56Å STM topography image taken at $V_B=1V$, $I_T=0.1nA$, (C) $dI/dV$ spectra over the PDI and Au in the chain, the tunneling gap is set at $V_B=2V$, $I_T=0.1nA$, (D) IETS spectrum of the CO$_2$ adsorbs on the chain, the tunneling gap is set at $V_B=-65mV$, $I_T=0.3nA$. 
Fig 4.2 Broken Cu-PDI chains on the Cu (110) surface. (A) 58Å ×58Å STM topography image taken at $V_B=0.65\,\text{V}$, $I_T=0.15\,\text{nA}$, (B) 29Å ×29Å STM topography image taken at $V_B=-0.5\,\text{V}$, $I_T=0.20\,\text{nA}$, (C) 58Å ×58Å STM topography image taken at $V_B=65\,\text{mV}$, $I_T=0.10\,\text{nA}$, CO$_2$ adsorbs on broken Cu-PDI chains on the Cu (110) surface, (D) IETS spectrum of the CO$_2$ adsorbs on the broken chain, the tunneling gap is set at $V_B=65\,\text{mV}$, $I_T=0.20\,\text{nA}$. 
Fig 4.3 STM topographic images and H2 IETS image at low H2 coverage, (A) 20Å×20 Å STM topography image taken at $V_B=23.4\,\text{mV}$, $I_T=0.2\,\text{nA}$, (B) 21.75Å×21.75 Å STM topography image taken at $V_B=5\,\text{mV}$, $I_T=0.2\,\text{nA}$, (C) 21.75Å×21.75 Å IETS image taken at $V_B=5\,\text{mV}$, $I_T=0.2\,\text{nA}$, (D) ball-stick model of single PDI molecule.
Fig 4.4 H$_2$ IETS spectra on different points of the molecule at low H$_2$ coverage, (A)

21.75 Å×21.75 Å STM topography image taken at $V_B$=5mV, $I_T$=0.2nA, (B) 21.75 Å×21.75 Å IETS image taken at $V_B$=5mV, $I_T$=0.2nA, (C) IETS spectra on different points of the molecule, the tunneling set point is at $V_B$=65mV, $I_T$=2.5nA.
Fig 4.5 STM topography images and H₂ IETS image at high H₂ coverage, (A) 17.5Å×17.5 Å STM topography image taken at V_B=65mV, I_T=0.15nA, (B) 17.5Å×17.5Å STM topography image taken at V_B=5mV, I_T=0.2nA, (C) 21.75Å×21.75 Å IETS image taken at V_B=-8.8mV, I_T=0.2nA, (D) ball-stick model of single PDI molecule.
Fig 4.6 \( H_2 \) IETS spectra on different points of the molecule, (A) \( 21.75\text{Å} \times 21.75 \text{ Å} \) STM topography image taken at \( V_B=-8.8\text{mV}, I_T=0.2\text{nA} \) (B) \( 21.75\text{Å} \times 21.75 \text{ Å} \) IETS image taken at \( V_B=-8.8\text{mV}, I_T=0.2\text{nA} \), (C) IETS spectra on different points of the molecule, the tunneling set point is at \( V_B=65\text{mV}, I_T=2.5\text{nA} \).
Fig 4.7: H₂ IETS image at different setpoints, (A) 21.75Åx21.75 Å IETS image taken at $V_B=5.1\text{mV}$, $I_T=0.2\text{nA}$, (B) 21.75Åx21.75 Å IETS image taken at $V_B=48.1\text{mV}$, $I_T=1.92\text{nA}$, (C) 21.75Åx21.75 Å IETS image taken at $V_B=-8.8\text{mV}$, $I_T=0.2\text{nA}$, (D) 21.75Åx21.75 Å IETS image taken at $V_B=39.1\text{mV}$, $I_T=1.5\text{nA}$. 
Fig 4.8: (A) 58Å×58 Å STM topography image taken at $V_B=50\text{mV}$, $I_T=0.18\text{nA}$, (B) 20Å×20Å IETS image taken at $V_B=10\text{mV}$, $I_T=1.75\text{nA}$, (C) 17Å×17Å IETS image taken at $V_B=42.5\text{mV}$, $I_T=0.05\text{nA}$, (D) 17Å×17 Å IETS image taken at $V_B=-4.2\text{mV}$, $I_T=0.25\text{nA}$. 
Bibliography


Chapter 5

Exploration of New Oxide Surfaces and Study of Molecules in Various Chemical Environments

5.1 Abstract

Molecules on oxide surfaces are particularly challenging to characterize due to limitation on the amount of current that can be used due to the weak adsorption energy. If too much tunneling current is used or the tunneling gap is too small, the molecule can be perturbed and either move or desorb from the surface. For STM, the oxide needs to be thin (< 10 Å) and is grown on a metal substrate by oxidation. Still the thin oxide provides electronic decoupling from the underlying metal substrate that sharpens the electronic states of the molecule. Thus it is possible to detect molecular orbitals and image their spatial distribution. An effort was spent to get partially oxidized (Al$_2$O$_3$) on the clean NiAl (110) surface, with about half oxide coverage and large Al$_2$O$_3$ islands, C$_{60}$, azulene, alizarin molecules were studied on this surface. A monolayer of CuO has been grown on Cu (110) surface, Cu/CuO stripe structure is also formed, allow the simultaneous investigation of molecules on both metal and oxide surfaces, C$_{60}$ and CO molecules are studied on this surface.

5.2 Introduction

Many metals found in nature are covered with an oxide. Oxides are known to exhibit different chemical and physical properties, as well as different composition, stoichiometry, and structure.
In interfacial electron transfer, oxides provide electronic isolation of the metal substrate and lowest lying conduction band that can accept photogenerated electrons from the molecules. For these reasons, it is desirable to investigate different oxides.

5.2.1 Al$_2$O$_3$/NiAl (110)

NiAl has simple cubic crystal structure, with Ni sits in the corners and Al sits in the center (CsCl prototype). It has good physical properties (low density: 5.85g cm$^{-3}$, high melting point: 1638C, high thermal conductivity, etc.) to be a material in aerospace industry. But its mechanical properties (lack of high fracture toughness and high temperature strength) limited its use now [1, 2]. Since the first report of the formation of ultrathin well-ordered Al$_2$O$_3$ layer (5Å) by oxidation of NiAl [3], it plays a key role in many technologies, like serving as stable supports for metal clusters [4] and an STM decoupling layer [5]. However, the microscopic structure of this thin film is still not clear. In the first paper of Freund, they use different electronic spectroscopy technologies to investigate of the structure of the thin film. Their LEED data shows that Al$_2$O$_3$ thin film contains distorted hexagonal oxygen layers and the oxygen-oxygen spacing is larger than that of bulk Al$_2$O$_3$. Their XPS and AES data show that the thickness of the film is 5Å; the thickness is compatible with two bilayers consisting of an aluminum layer and a quasi-hexagonal oxygen layer. Their EELS data suggest the structure is mostly likely similar to α-Al$_2$O$_3$ (0001) or γ-Al$_2$O$_3$ (111) [3]. There is no evidence showing the Ni-O interactions (NiO or NiAl$_2$O$_4$) or Ni existing in the Al$_2$O$_3$–NiAl interface, the Ni atoms dissolved into the bulk NiAl during annealing process at around 1250K (the diffusion coefficient is high enough at this temperature for Ni diffuse into bulk). The fact that the Al$_2$O$_3$ layer is chemically inert to most molecules up to 800K leads to the presumption that the thin film is oxygen terminated since electronic structure of O$^{2-}$ is near to that of rare gas, so that the sequence of the layer is NiAl-Al-
O-Al-O. Later, this group took out the Al₂O₃/NiAl sample from the UHV chamber and studied with TEM, the data supported the γ-Al₂O₃ (111) structure [6].

Stierle et al. reported their detailed study of the Al₂O₃/NiAl structure using an extended surface x-ray diffraction (SXRD), which allows them to derive an atomically resolved structure model. The distorted double layer of hexagonal O ions act as building blocks for the film and the Al ions host on distorted octahedral or tetrahedral sites. Uncoordinated Al ions are located in the oxide/metal interface to fulfill the stoichiometry. The resulting structure is closely related to bulk κ- Al₂O₃, which was shown both in theory and experiment before. They also show the atomic positions within the structure [7]. Later, Kresse et al. found that the short bond lengths (Al-Al 2.08 Å, Al-O 1.51 Å) in the structure that Stierle reported are unstable in their ab initio calculations. Since they observed square features in STM image, they explained them by a square arrangement of O atoms. The topmost Al atoms are pyramidally and tetrahedrally coordinated. The stacking consequence of the thin film is 4(Al₄O₆Al₆O₇) and the stoichiometry is Al₁₀O₁₃, different from commonly Al₂O₃. The oxide unit cell covers 16 NiAl substrate unit cell. They propose the same building blocks can be found on surfaces of bulk oxides [8].

Researchers are also interested in the grain boundary structure of the thin Al₂O₃ film. The Freund group studied the structure with STM to determine the atomic structure of the antiphase domain boundaries of the thin Al₂O₃ film, APDB result from the insertion of an oxygen row but do not induce change in the geometric height, the apparent corrugation of APDB in STM images at certain biases must arises from the electronic properties of these dislocations [9]. Combining STM images and DFT calculations, Kresse et al. derived a model of APDB of Al₂O₃/NiAl. It indicates that the structure is O deficient; two Al 3p electrons fail to find the corresponding empty O conduction band states, the O-deficient domain boundary acts as a native electron
donor. By transferring these electrons to the NiAl support, an insulating surface layer will be restored [10].

5.2.2 CuO/Cu (110)

Molecular Oxygen is long-time known to dissociatively chemisorb on the Cu (110) surface and forms the 2×1 structure at O coverage of 0.5 monolayer [11]. Different techniques including EELS, LEIS, photoemission, XRD, He diffraction, etc. have been used to study the structure, all these studies conclude that O chemisorbed on the long bridge site (along the <100> direction), but the detailed atomic geometry remains controversial for 20 years, based on different experiments, “buckled-row model” and “missing-row model” have been suggested [12, 14]. In early 1990’s, with the STM experiments of nucleation, structure and growth of p(2×1)O-Cu(110), it’s clear that this new phase is formed by reaction of chemisorbed O atoms with the Cu atoms evaporated from the step edges and diffused across terraces, the structure is an “added-row” type [12, 13, 14]. The oxygen-induced p (2×1) O-Cu (110) has been attracted much interest for the past two decades after the structure gets clear. One great advantage of this surface over Al₂O₃/NiAl(110) is the much smaller unit cell and the surface atoms can be imaged, thus the clusters, molecules and atoms on this surface has a more uniform and well-defined chemical environment surrounded. For STM experiments, it’s also served as a magnetic insulator [15] and one dimensional substrate [16].

Besides the 2×1 structure, there are some other kinds of chemisorbed O induced reconstruction, most notably c (6×2) O [17].

5.3 Experiments and Discussion

The aluminum oxide and CuO are formed and different molecules are studied on the oxide surfaces with the homemade STM.
The preparation of NiAl (110) surface and aluminum oxide patches is modified from previous procedures [5]. The sample holders are modified, adding sapphire hemispheres and rings between sample holder and NiAl sample [18], enabling the NiAl sample annealed at higher temperature \( \sim 1200 ^\circ C \), at such a high temperature, sputtering is not necessary to get a clean metal surface. Partial crystalized oxide patches are formed on a clean NiAl(110) surface by 240L oxidation (120s at \( 2 \times 10^{-6} \) torr) at \( \sim 530 ^\circ C \) followed by annealing at \( 1050 ^\circ C \) for 7 mins. Furthermore, full crystalized aluminum oxide surface can be formed by 550L oxidation at \( \sim 530 ^\circ C \) (275s at \( 2 \times 10^{-6} \) torr) followed by annealing \( 1050 ^\circ C \) for 7 mins (fig 5.1).

The Cu (110) surface is cleaned by sputter (0.5kV Ne beam) anneal (480°C) cycles, some defects still remain on the surface (fig5.5 A). The \( p(2\times1) \) O-Cu (110) is formed by 3L oxidation (150s at \( 2 \times 10^{-8} \) torr) at \( \sim 300 ^\circ C \). The oxide looks uniform without any defects (fig 5.5 B and fig 5.5 C). Cu/CuO stripes can also be formed by smaller amount \( O_2 \) oxidation or annealing the full oxide surface, but the metal looks kind of amorphous. Also, the Cu stripe + CuO stripe width is limited to 6~14nm [19], without the big terraces, molecules prefer to absorb on metal or edge between Cu and CuO (fig 5.7 B).

Electrochemically etched Ag tips which are in-situ cleaned before the experiments are used in these experiments.

### 5.3.1 Molecules on \( \text{Al}_2\text{O}_3/\text{NiAl} \) (110)

\( \text{C}_60 \) is a well-studied molecule with vibronic features [20] and can be superconductor with alkali metal doped [22]. In our experiments, \( \text{C}_60 \) is evaporated onto the \( \text{Al}_2\text{O}_3/\text{NiAl} \) (110) surface at room temperature, all molecules adsorb on NiAl at this temperature even though the molecules also go to oxide at \( 10K \) [20, 21]. A full oxide surface is formed to get the molecules onto the oxide (fig 5.1). The \( \text{C}_60 \) layer looks different on metal and on the oxide (fig 5.1 A, B and C), the
plan was to study the life time of the vibronic states of the C$_{60}$ layer by shining fs laser [20]. K was also doped for study of superconductivity [22]. For all these experiments, L He is needed to cool the molecules down to 10K; however, they were not possible given the shortage of L He at the time these experiments were performed.

Azulene is a molecule with unique electronic and photoelectric properties [23]. With 193 nm photon absorption, the azulene molecules will first convert to its isomer naphthanlene and then dissociate to smaller molecules [24]. Here, we shine 220nm fs laser into the STM junction, multiphoton at this energy can also covert the molecule to naphthanlene and dissociate. Fig 5.3 (A) shows the azulene molecules look uniform before shining the 220nm fs laser beam, after shining the laser, smaller species appears as shown in fig 5.3(B), probably due to photo isomerization and photo dissociation. Smaller species also show up on the oxide after shining fs laser as shown in fig 5.3(C) and fig 5.3(D). The goal next will be studying the photochemistry at single molecule level and characterizing the dissociated products with a CO tip at 10K.

Alizarin (1, 2-dihydroxyanthraquinone) is an organic red dye that is used for dying textile fabrics and is an abundant material for dye sensitized solar cells. The molecules are thermally evaporated onto the Al$_2$O$_3$/NiAl (110) surface at 10K. Fig 5.4 shows the STM topography images and dI/dV spectra of three molecules on the oxide, the topography images show the LUMO geometry when scanned near the LUMO peak, the dI/dV spectrum from 0 to 1.6V show the LUMO peak(s) around 1V, however, only the molecule on the grain boundary shows the HOMO peak. Molecule 2 disappears after taking the spectrum. More molecules need to be studied to confirm these data. The ultimate goal of this project is to study the electron transfer between the molecule and TiO$_2$. Also the alizarin molecule may stand up on the surface for fs laser simulated Raman experiments [25].
5.3.2 Molecules on CuO/Cu (110)

CO is evaporated onto the CuO/Cu (110) surface at 13.5K, CO prefers to go to the metal surface, given the metal surface is kind of amorphous, it’s hard to tell individual CO molecules apart and some CO molecules do not look round (fig 5.7 A and B). However CO looks big protrusions on CuO surface and they form 1D chain (fig 5.5 B and D). The hindered rotational mode of CO on CuO has slight blue shift [26]; it’s also possible to pick up CO from the oxide. However, it’s easy to see H₂ signal on the CuO surface without introducing any H₂ into the UHV chamber. With such strong H₂ IETS background, it’s hard to see IETS modes from other molecules like CO, NO or O₂.

C₆₀ is also thermally evaporated onto the CuO/Cu (110) surface at room temperature, different from the layers that C₆₀ forms on Al₂O₃/NiAl (110) (fig 5.2), they form clusters on CuO instead (fig 5.6A). Mg is thermally evaporated onto the C₆₀/CuO surface; Mg prefers the C₆₀ clusters than the CuO. Individual Mg atoms could be imaged on the C60 clusters if the coverage is low (fig 5.7 B and C). With more Mg evaporated onto the surface, the Mg fully covers around the C₆₀ clusters instead of adsorbing on CuO. MgC₆₀ is a semiconductor used in solar cell [27]; investigating the number and site dependence of dosed Mg on C₆₀ upon electronic structure may help improve the structure, furthermore, time-resolved study of charge transfer could be performed by combining fs laser with STM.

4T-BTD and alizarin are also evaporated onto the Cu/CuO stripes, the molecules prefers the metal or the edge.
5.4 Conclusions

Large aluminum oxide islands are formed on NiAl (110) surface by modifying previous recipes. CuO is formed on Cu (110) surface with well-defined structure. STM imaging and spectroscopy are performed on molecules (including C\textsubscript{60}, azulene, alizarin and CO) adsorbed on Al\textsubscript{2}O\textsubscript{3} and CuO surfaces. The thin oxide provides electronic decoupling from the underlying metal substrate that sharpens the electronic states of the molecule, the oxide can also elongate the lifetime of excited states of adsorbed molecules.

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Fig 5.1: partial and full oxide at room temperature, (A) 2000Å×2000Å STM topography images of partial oxide on NiAl (110) surface, \( V_B=2V, I_T=0.10nA \), oxide is on the left and metal is on the right, the lines are grain boundaries, (B) 2000Å×2000Å STM topography images of partial oxide on NiAl (110) surface, \( V_B=2V, I_T=0.10nA \), oxide is in the middle and the two sides are metal, (C) 2000Å×2000Å STM topography images of full oxide on NiAl (110) surface, \( V_B=3V, I_T=0.10nA \), (D) 2000Å×2000Å STM topography images of full oxide on NiAl (110) surface, \( V_B=3V, I_T=0.10nA \).
Fig 5.2: STM topography images of C$_{60}$ adsorbed on Al$_2$O$_3$/NiAl (110), the C$_{60}$ is thermally evaporated onto the surface at room temperature and the experiments are performed at 84K, all images are 125Å×125Å at $V_B=2.5$V, $I_T=0.10$nA, (A) C$_{60}$ layer on metal, (B) C$_{60}$ layer on oxide (C) irregular layer on oxide, (D) K on C$_{60}$ layer on oxide.
Fig 5.3: STM topography images of azulene on Al$_2$O$_3$ and NiAl(110) before and after 220nm fs laser irradiation, all the images are 250Å×250Å, (A) on NiAl before laser irradiation, $V_B=1.00$V, $I_T=0.10$nA, (B) on NiAl after laser irradiation, $V_B=1.50$V, $I_T=0.10$nA, (C) on oxide before laser irradiation, $V_B=1.50$V, $I_T=0.10$nA, (D) on oxide after laser irradiation, $V_B=1.50$V, $I_T=0.10$nA.
Fig 5.4: STM topography images and spectroscopy of alizarin on Al$_2$O$_3$ at 13.5K, (A) 87Å×87Å STM topography images of three alizarin molecules on Al$_2$O$_3$/NiAl, $V_B$=1.5V, $I_T$= 0.10nA, (B) 87Å×87Å STM topography images of two alizarin molecules on Al$_2$O$_3$/NiAl, $V_B$=1.5V, $I_T$= 0.10nA, molecule 2 disappear during spectroscopy, (C) dI/dV spectroscopy of the molecules from 0V to 1.6V, the tunneling gap is set at $V_B$=1.5V, $I_T$= 0.10nA, (D) dI/dV spectroscopy of the molecules from -1.6V to 0V, the tunneling gap is set at $V_B$=1.5V, $I_T$= 0.10nA.
Fig 5.5: (A) 500Å ×500Å STM topography image taken at $V_B = -3V$, $I_T = 0.12nA$ on Cu (110) at room temperature, (B) 58Å ×58Å STM topography image taken at $V_B = 0.1V$, $I_T = 0.1nA$ on CuO at 12.5K, with a CO on the surface, (C) 17.5Å ×17.5Å STM topography image taken at $V_B = 0.5V$, $I_T = 0.8nA$ on CuO at 12.5K, (D) 58Å ×58Å STM topography image taken at $V_B = -0.5V$, $I_T = 0.1nA$ of CO chains on CuO at 12.5K.
Fig 5.6: STM topography image of C$_{60}$ and MgC$_{60}$ on CuO/Cu(110) surface, (A) 500Å ×500Å STM topography image taken at $V_B=2.05V$, $I_T=0.12nA$ on CuO at room temperature, the clusters are formed by C$_{60}$, which are thermally evaporated onto the surface, (B) 1000Å ×1000Å STM topography image taken at $V_B=2.05V$, $I_T=0.12nA$ on CuO at room temperature, with Mg evaporated on C$_{60}$/CuO, the individual dots are Mg, (C) 500Å ×500Å STM topography image taken at $V_B=2.05V$, $I_T=0.13nA$ with Mg on C$_{60}$/CuO at room temperature, (D) 250Å ×250Å STM topography image taken at $V_B=1.08V$, $I_T=0.12nA$ with more Mg on C$_{60}$/CuO.
Fig 5.7: (A) 464Å ×464Å STM topography image taken at $V_B=0.6\,V$, $I_T=0.1\,nA$ on CuO/Cu at 13.5K, (B) 116Å ×116Å STM topography image taken at $V_B=0.6\,V$, $I_T=0.1\,nA$ on CuO/Cu at 13.5K, (C) 81Å ×81Å STM topography image taken at $V_B=37\,mV$, $I_T=0.5\,nA$ on CuO/Cu at 13.5K, (D) 81Å ×81Å STM topography image taken at $V_B=37\,mV$, $I_T=0.5\,nA$ on CuO/Cu at 13.5K.
Bibliography


25. Original data from Peter Lu’s Group at Bowling Green State University, OH.


Chapter 6

Imaging and Electronic Spectrum of Fe Nanoclusters on Al₂O₃/NiAl (110) at Various Temperatures with a Laser-STM

6.1 Abstract

Fe nanoclusters are grown by thermally evaporating Fe onto Al₂O₃/NiAl (110) surface at room temperature and ~84K. At room temperature, Fe prefers to absorb on the NiAl (110) surface and can grow into islands/films with enough coverage on metal, sparse Fe nanoclusters are grown on the oxide. At 84K, Fe grows into size-limited nanoclusters on both Al₂O₃ and NiAl (110) surface; still with less coverage on oxide. The nanoclusters on oxides are bigger than the ones on metal, with increasing Fe coverage, the number of nanoclusters on both metal and oxide will increase instead of forming islands/thin films on NiAl (110), DC and RF spectroscopy are performed on the nanoclusters on Al₂O₃ and NiAl (110) at 84K. Furthermore, the nanoclusters are annealed to form larger ones, the larger clusters are cooled down to ~21K and dI/dV spectroscopy is performed on the clusters.

6.2 Introduction

Metal films and particles on oxide surfaces are of great importance in many technological applications including heterogeneous catalysis [1, 2], spintronic devices [3] and photovoltaic devices [4]. In this work we focus on the creation of small iron clusters on a thin oxide film serving as a model system to investigate the catalytic, magnetic and optical properties of
nanosized metallic particles on an insulating substrate. Fe nanoclusters on oxide have been proposed to have potential applications such as catalysis [3] and spintronics [4].

The Scanning Tunneling Microscope (STM) has been a powerful tool to study nanoscience by imaging and electronic spectroscopy [5]. Different metal nanoclusters adsorb on thin oxide film grown on metal substrates can be studied with a STM [1]. Fe clusters adsorb on MgO thin film grown on Ag (100) [6] and alumina thin film grown on Ni₃Al (111) [7, 8] have been characterized with STM imaging. Theoretical studies are also carried out to study the magnetic properties [9] and catalytic behaviors [2] of Fe clusters supported on oxide. However, electronic properties of the Fe nanoclusters have only been studied on the metal Au (111) surface [10], but not on the oxide surfaces. Nanoclusters on oxide surfaces are particularly challenging to characterize the electronic properties due to limitation on the amount of current. Electronic properties of single atoms and molecules have been characterized on 5Å alumina thin film grown on NiAl (110) surface [11, 12]. In this work, electronic properties of Fe nanoclusters on oxide Al₂O₃/NiAl (110) are studied at 84K and 21K.

6.3 Experiment

The experiments were performed using a home-built STM operating at various temperatures from 21K to room temperature and a base pressure of 3×10⁻¹¹ Torr [13]. The preparation of NiAl (110) surface and aluminum oxide patches is modified from previous procedures [11, 12], the NiAl sample was annealed at higher temperature ~1200°C, the partial crystalized oxide patches are formed on a clean NiAl(110) surface by 240L oxidation (120s at 2×10⁻⁶ torr) at ~530°C followed by annealing at 1050°C for 7 minutes. The Ag tip was electrochemically etched and in-situ cleaned before experiments. The Fe is thermally evaporated onto the surface at room
temperature or 84K and forms nanoclusters on the oxide. Electronic properties are studied by performing dI/dV spectrum on the nanoclusters. Moreover, with 220nm laser shining into the junction, DC dI/dV spectroscopy and RF dI/dV spectroscopy with double lock-in technique are also performed. The nanoclusters grown on the surface at room temperature are annealed to ~600°C to get bigger size nanoclusters; the bigger nanoclusters are cooled to 21K to perform electronic properties studies.

6.4 Results and Discussion

For comparison, Fe clusters and thin films on NiAl (110) metal surface are also studied. Fe is evaporated onto the clean NiAl (110) surface (fig 6.1 A) at room temperature. At small coverage, the Fe will first saturate on the step edges, and then small clusters are grown on the terraces (fig 6.1 B). With enough coverage, layers of Fe films are grown from step edges (fig 6.1 C and D). For the partial 5Å oxide crystalized patches covered Al₂O₃/NiAl (110) surface, Fe prefers to absorb on the NiAl (110) surface and grown into islands, sparse Fe nanocluster could be found on the oxide with enough dosage.

With the Al₂O₃/NiAl (110) surface cooled down to 84K, Fe grows into size-limited nanoclusters on both oxide and metal; more clusters could be found on the metal. The size of the Fe nanoclusters are different on the metal (~2nm) from the ones on oxide (~3nm) (fig 6.3 A and B). Different from the layers or islands of Fe film grown on NiAl (110) at room temperature, more Fe nanoclusters are grown on both oxide and metal if more Fe is dosed (fig 6.3C). A dimer of Fe clusters on oxide is shown if fig 6.3 D, the clusters are even more crowded on the metal.

Electronic properties of the clusters are measured by performing dI/dV spectroscopy with the lock-in technique. From the dI/dV spectrum on the Fe nanocluster in fig 6.4A on NiAl (110),

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the cluster has a peak at 2.2V (fig 6.4B). With 220nm fs laser shining into the junction, DC dI/dV spectroscopy is performed, the 2.2V peak slightly shifts to ~2.3V. On the negative bias side, the dI/dV signal is dominated by the photoemission from sample to tip. The dI/dV signal in and out of tunneling on both Fe nanocluster and NiAl (110) background is measured (fig 6.4C). All the spectra have two peaks around ~-0.8V and 0.5V, with slightly different peak position and intensity especially for the -0.8V peak. Furthermore, the DC and ~80MHz laser induced RF signal can be separated by using an external bias tee, the RF signal is fed into a 200MHz lock-in, the output of 200 MHz lock-in amplifier is fed into a second lock-in amplifier that detects a lower frequency signal generated by modulating the sample bias, for dI/dV and d²I/dV² spectroscopy. The dI_{RF}/dV spectrum on the Fe nanocluster at tunneling has ~1V peak, very similar to the background dI_{RF}/dV at tunneling. While the noise spectrum without laser is without any peaks and the ~1V peak on dI_{RF}/dV spectra out of tunneling (on cluster and background) are less obvious.

Same measurements are performed on nanoclusters deposited on oxide (fig 6.5), the dI/dV spectrum on the oxide background has no peaks from -2V to 2V since the oxide film has a band gap at this range (fig 6.5B). Besides the peak at 2.2V, two other peaks at 1.2V and -1.1V on the nanocluster are also observed (fig 6.5B and fig 6.5D). With 220nm fs laser in the junction, the 2.2V peak shifts to 2.3V, suggesting possible laser induced effects on the electronic states of the nanocluster. The dI_{RF}/dV spectra on the nanocluster and the oxide background are noisier than the ones on metal NiAl (110), the DC dI/dV with laser on is also dominated by photoemission, the peak position is similar to NiAl, but not so sharp.

To get bigger size nanoclusters, the clusters deposited at RT are annealed to 600°C. Fe islands are grown on metal even at small coverage, the clusters on oxide are grown bigger to
~4nm on oxide, during annealing, some clusters diffuse to grain boundaries so more clusters can be found there (fig 6.6). The annealed sample is cooled to 21K and electronic properties of Fe clusters on oxide are studied (fig 6.7). Fig 6.7A shows one cluster has peaks at ~2.2V, ~1.2V and ~1.1V; while there are no peaks on the other cluster. Fig 6.7B shows the electronic spectra on different points of the dimer, the peak energy and intensity vary on different points.

6.5 Conclusions

In conclusion, Fe clusters are formed and investigated on Al2O3/NiAl (110) surface with STM. Fe thin films are formed on metal surface at room temperature while sparse cluster can be formed on oxide. At 84K, clusters are both formed on oxide and metal surfaces. Bigger Fe cluster can be formed by annealing the sample to high temperature. Electronic properties are studied at 84K and 21K, DC and RF dI/dV spectra are also studied with 220nm fs laser in junction. For most of the Fe clusters formed and investigated at 84K, there is a DC dI/dV peak around ~2V. For most of the bigger clusters investigated at ~20K, other states besides the 2V are observed. With laser shining, a small shift of 2V peak in DC dI/dV spectra is detected; suggesting some laser induced effect on electronic states. DC dI/dV spectra on the negative bias with laser shining are dominated by the photoemission from sample to tip. RF dI/dV spectra with the double lock-in technique are noisy although some differences on nanocluster and background are recorded.
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Fig 6.1: Fe nanocluster and films on NiAl(110) at room temperature, (A) 1000Å×1000Å STM topography image taken at $V_B=2.0\text{V}$, $I_T=0.11\text{nA}$, (B) 2000Å×2000Å STM topography image taken at $V_B=2.05\text{V}$, $I_T=0.11\text{nA}$, (C) 1000Å×1000Å STM topography image taken at $V_B=2.05\text{V}$, $I_T=0.11\text{nA}$, (D) 250Å×250Å STM topography image taken at $V_B=1.08\text{V}$, $I_T=0.13\text{nA}$. 
Fig 6.2: Fe clusters and islands on Al$_2$O$_3$/NiAl (110) surface at room temperature, (A)
4000Å×4000Å STM topography image taken at $V_B=2.0$V, $I_T=0.10$nA, (B) 2000Å×2000Å STM
topography image taken at $V_B=2.05$V, $I_T=0.10$nA, (C) 1000Å×1000Å STM topography image
taken at $V_B=2.05$V, $I_T=0.10$nA, (D) 1000Å×1000Å STM topography image taken at $V_B=2.05$V,
$I_T=0.10$nA.
Fig 6.3: Fe clusters on Al₂O₃/NiAl (110) at 84K, (A) 250Å×250Å STM topography image taken at $V_B=1.08$V, $I_T=0.10$nA, (B) 250Å×250Å STM topography image taken at $V_B=2.00$V, $I_T=0.10$nA, (C) 250Å×250Å STM topography image taken at $V_B=2.50$V, $I_T=0.12$nA, (D) 62Å×62Å STM topography image taken at $V_B=2.00$V, $I_T=0.10$nA.
Fig 6.4: Spectroscopy of Fe cluster on NiAl (110) at 84K, (A) 60Å×60Å STM topography image taken at $V_B=2.50\,V$, $I_T=0.10\,nA$ of a 2~3nm Fe cluster on NiAl, (B) $dI/dV$ of the NiAl BKGD and the Fe nanocluster, the red curve shows the $dI/dV$ with 220nm fs laser in junction, the negative bias side is dominated by photoemission, (C) $dI_{DC}/dV$ on the nanocluster and NiAl at tunneling and out of tunneling, the photoemission at negative bias dominates the signal, with double peaks around 0V, the peak position slightly varies, (D) $dI_{RF}/dV$ with the double lock-in technique. The tunneling set point for all the spectrum is at (2.5V, 0.1nA).
Fig 6.5: Spectroscopy of Fe cluster on Al$_2$O$_3$ at 84K, (A) 125Å×125Å STM topography image taken at $V_B=3.00\,\text{V}$, $I_T=0.10\,\text{nA}$ of a 3~4nm Fe cluster on Al$_2$O$_3$/NiAl, (B) dI/dV of the Al$_2$O$_3$ BKGD and the Fe nanocluster, the red curve shows the dI/dV with 220nm fs laser in junction, the negative bias side is dominated by photoemission, (C) d$I_{\text{DC}}$/dV on the nanocluster with and without the 220nm in junction, showing the peak shifts from 2.2V to 2.3V with laser, (D) d$I_{\text{RF}}$/dV with the double lock-in technique, the whole d$I_{\text{DC}}$/dV on the nanocluster with laser is also shown for comparison. The tunneling set point for all the spectrum is at (3V, 0.1nA).
Fig 6.6: Fe nanoclusters and islands after annealing to 600°C. (A) 2000Å×2000Å STM topography image taken at $V_B=3.01\,\text{V}$, $I_T=0.10\,\text{nA}$, (B) 1000Å×1000Å STM topography image taken at $V_B=3.01\,\text{V}$, $I_T=0.10\,\text{nA}$, (C) 500Å×500Å STM topography image taken at $V_B=3.01\,\text{V}$, $I_T=0.10\,\text{nA}$, (D) 250Å×250Å STM topography image taken at $V_B=3.01\,\text{V}$, $I_T=0.10\,\text{nA}$.
Fig 6.7: Spectroscopy of the annealed clusters at 21K, (A) 140Å×140Å STM topography image taken at \( V_B = 3.00V \), \( I_T = 0.10nA \) of two Fe cluster on Al\(_2\)O\(_3\), (B) dI/dV of the Al\(_2\)O\(_3\) BKGD and the two Fe nanoclusters in (A), the tunneling set point is at \( V_B = 3.00V \), \( I_T = 0.10nA \), (C) 70Å×70Å STM topography image taken at \( V_B = 2.50V \), \( I_T = 0.12nA \) of Fe cluster dimer on Al\(_2\)O\(_3\), (D) dI/dV of different points on the dimer in (C), the tunneling set point is at \( V_B = 2.50V \), \( I_T = 0.12nA \).


Chapter 7

Concluding Remarks and Future Prospects

7.1 Concluding Remarks

Two important molecules are extensively studied in this dissertation with STM at sub Å resolution by probing with H\(_2\) or a CO terminated tip. The molecular orbital of 4, 7-Di ([2, 20-bithiophen]-5-yl) benzo[c] [1, 2, 5] thiadiazole (4T-BTD) with intramolecular donor-acceptor-donor sites is probed with the electronic state dI/dV imaging and H\(_2\) rotational and vibrational spectromicroscopy. H\(_2\) rotational spectromicroscopy on the 4T-BTD molecule shows the sub Å detailed LUMO structure of the molecule. Understanding the properties of the conjugated building blocks like 4T-BTD will help building blocks to extended semiconducting molecules with predictable properties. 1, 4-Phenylene Diisocyanide (PDI) is probed by imaging with a CO-terminated tip and H\(_2\). At low H\(_2\) coverage the STM H\(_2\) vibrational spectromicroscopy clearly shows the two N sites; at high H\(_2\) coverage, the topography image already shows three parts of the molecule with interactions with the surface, the H\(_2\) vibrational spectromicroscopy shows more details of the molecule and its interaction with the substrate. Topography image with a CO-tip also shows different parts of the molecule and the interaction with the substrate. PDI can self-assemble on noble metal surfaces to form nanostructures, which could have potential applications in molecular electronics and catalysis.

Further combination of a RF-STM with a tunable femtosecond laser enables the investigation of light-molecule interactions. In this dissertation, efforts are spent to setup a new tunable fs laser (220 nm–1040 nm) to couple with the RF-STM. The wavelength of a MaiTai
Deepsee (690~1040) is extended by a harmonic generator to generate the 2\textsuperscript{nd}, 3\textsuperscript{rd} and 4\textsuperscript{th} harmonics. The 2\textsuperscript{nd}, 3\textsuperscript{rd} and 4\textsuperscript{th} harmonic can be applied for photon induced chemistry or photoemission, the 2\textsuperscript{nd} harmonic or 3\textsuperscript{rd} harmonic can also be used to perform pump-probe time resolved study by overlapping with the fundamental. The fundamental and fundamental (1+1), fundamental and 2\textsuperscript{nd} harmonic (1+2), fundamental and 3\textsuperscript{rd} harmonic (1+3) are aligned to overlap on the diagnostic path on the optical table. Only fundamental and fundamental can be routinely aligned to overlap in the STM junction, the UHV in-situ lens change the spatial and temporal overlap of different color beams.

The effects of the femtosecond laser are followed by detecting photo induced electron emission and photochemistry. 4\textsuperscript{th} harmonic UV laser beam induced DC and RF photocurrent has been detected out of tunneling, multiphoton current induced by two overlapped fundamental beams is also detected. The signal is too noisy to be detected at tunneling; a new double lock-in technique is applied to detect the weak laser-induced signal in the tunneling regime. To sharpen the energy width and increase the lifetime of the excited states of molecules, thin aluminum oxide and copper oxide are grown on metal surfaces to provide electronic isolation of the metal substrate and adsorbed molecules. Different molecules are studied on the oxide. Metal nanoclusters are grown on metal and oxide to improve laser-induced signal through plasmonic enhancements.

### 7.2 Future Prospects

#### 7.2.1 Fabricated Tip

To detect the laser induced signal in the tunneling regime, the current STM4 configuration with the in situ UHV cryogenic RF amplifier is not sensitive enough. It has been shown by using the
fabricated grating on a Au tip, much greater plasmonic enhancement are achieved to get the photoinduced current in the nA range [1]. For more plasmonic enhancements, a fabricated Ag, Au or Cu tip can be used in the laser-STM experiments. Combining the fabricated tip, cryogenic RF amplifier and the double-lock in technique, there is a good chance to detect the photoinduced current in the tunneling regime and successfully perform the long time pursued spatially resolved lifetime measurement with a laser STM. To get the two different color laser beams aligned in the STM junction for time resolved experiments, one way is to use parabolic mirror outside of the UHV chamber for better simulation on the optical table, although the beam size in the junction will become bigger given the longer focal length [2]. An electron analyzer could also enhance the sensitivity to detect photoinduced electrons [2].

### 7.2.2 Designed Molecules

Preliminary data in this dissertation shows the possible photoisomerization and photodissociation of the azulene molecules with big laser power shining into the junction out of tunneling. Coworkers in the Ho Group already got sub-Å structural and molecular orbital images with H$_2$ and CO terminated tip [3]. With the new grating tips, plasmonic enhancement would induce photoisomerization and dissociation, one molecule at a time. With the H$_2$ and CO terminated tip imaging, one should be able to image the sub-Å structure of the naphthalene after isomerization and the smaller molecules after dissociation. The molecular orbital of 4T-BTD molecule is extensively investigated in this dissertation; the life time of the excited state of this molecule could be further investigated with the laser STM. Alizarin molecules may stand up on metal and oxide surfaces, which could be used for the proposed fs-stimulated Raman experiments [4]. CO$_2$ and CO$_2^-$ adsorbed on Au-PDI chains could be further investigated by STM-IETS.
7.2.3 Oxide Surfaces

Aluminum oxide on NiAl(110) and copper oxide on Cu(110) are investigated in this dissertation. Aluminum oxide has a large band gap, could be used as an insulator to study single molecules. Although copper oxide on Cu(110) is not a good electric insulator, it has been proved to be a magnetic insulator [5], the magnetic properties of small molecules like NO and large magnetic organic molecules can be probed on this well-defined oxide, a Cu tip instead of Ag tip might be more stable on the CuO surface. Other insulating surfaces can also be investigated, especially the ones with various thicknesses of insulating layers, like MgO on Ag(100) [6]. Such thicker insulating layers will further increase the life time of the excited states of adsorbed molecules.

7.2.4 Ag and Au Nanoclusters

Fe nanoclusters are grown on oxide and metal surfaces and studied with the laser STM in this dissertation, besides DC dI/dV spectroscopy, RF dI/dV spectroscopy is also recorded with the double lock-in technique. The catalytic properties of Fe nanoclusters can be further investigated. With the new amplifier and grating tips, RF signal may be detectable in the tunneling regime. Ag and Au nanoclusters which have better plasmonic properties can now be also grown on the surface as the Au and Ag evaporators are ready. The plasmonic enhancement of these nanoclusters can help detect the photoinduced signal in the tunneling regime. Moreover, the plasmons induced by the laser and greatly enhanced by the grating tip and nanoclusters could be used in photocatalysis [7].
Bibliography


3. S. Li, A. Ye and W. Ho, to be published.

4. Original data from Peter Lu’s Group at Bowling Green State University, OH; private communication with H. Kumar Wickramasinghe.


Appendix A

STM4 Optics Setup

A.1 Catalog

Below is the catalog of the optical devices in STM4, the optical path and setup photo is shown as in fig2.1 and fig2.2 of chapter 2.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Device Purpose</th>
<th>Company</th>
<th>Product # &amp; Characteristics</th>
<th>Price</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ti: Sapphire Laser</td>
<td>MaiTai Deepsee</td>
<td>Spectra Physics</td>
<td>Mai Tai EHP DS + Warrenty supplement +Gold Level Service Tunable from 690-1040, 70fs-100fs pulse width, 80MHz, 2.5W at 800nm</td>
<td>160,000</td>
</tr>
<tr>
<td>Periscope 1</td>
<td>Change the beam from p to s, change the height of the beam from 4.38&quot; to 5.5&quot;</td>
<td>Newport</td>
<td>9151-M 12” Periscope kit and m1, m2</td>
<td>434.88</td>
</tr>
<tr>
<td>m1</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 45 degrees AOI, reflect over 99.6% from 680 to 1060 nm, optimized for s</td>
<td>499</td>
</tr>
<tr>
<td>m2</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35P 45 degrees AOI, reflect over 99.6% from 680 to 1060 nm, optimized for p</td>
<td>499</td>
</tr>
<tr>
<td>m3</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S</td>
<td>499</td>
</tr>
<tr>
<td>m4</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S</td>
<td>499</td>
</tr>
<tr>
<td>lens1</td>
<td>Combine with lens 2 as a telescope to reduce beam size f=75mm</td>
<td>CVI</td>
<td>PLCX-25.4-38.6-UV</td>
<td>140</td>
</tr>
<tr>
<td>lens2</td>
<td>f=65mm</td>
<td>CVI</td>
<td>PLCX-25.4-33.7-UV</td>
<td>300</td>
</tr>
<tr>
<td>TS 1</td>
<td>Adjust the distances between lenses</td>
<td>Newport?</td>
<td>From Stock</td>
<td></td>
</tr>
<tr>
<td>LPC</td>
<td>Stabilize laser power, 700-1100nm</td>
<td>BEOC</td>
<td>MI - 10450 – NIR, 5450</td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>-----------------------------------</td>
<td>-------</td>
<td>------------------------</td>
<td></td>
</tr>
<tr>
<td>BS 1</td>
<td>20% power goes to fundamental path, 80% goes to harmonic generator</td>
<td>Chroma</td>
<td>20-80bs-NIR 20/80 beam splitter, 20R, 80T. 600-1200nm wavelength range, laser-grade substrate AR-coating. Actually 30-70 beam splitter 200</td>
<td></td>
</tr>
<tr>
<td>m5</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 499</td>
<td></td>
</tr>
<tr>
<td>m6</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 499</td>
<td></td>
</tr>
<tr>
<td>m7</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 464.07 w/discount</td>
<td></td>
</tr>
<tr>
<td>m8</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 464.07</td>
<td></td>
</tr>
<tr>
<td>m9</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 464.07</td>
<td></td>
</tr>
<tr>
<td>BS 2</td>
<td>50% power goes to delay stage, 50% goes to retroreflector</td>
<td>Chroma</td>
<td>50/50bs-NIR 600-1200nm range, 25mm diameter, laser-grade substrate, AR coating 200</td>
<td></td>
</tr>
<tr>
<td>TS 2</td>
<td>Translational Stage, adjust the two beams to overlap</td>
<td>Newport</td>
<td>Can be adjusted in 3 directions, same as TS8 From Stock</td>
<td></td>
</tr>
<tr>
<td>m10</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 464.07</td>
<td></td>
</tr>
<tr>
<td>m11</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S 464.07</td>
<td></td>
</tr>
<tr>
<td>Retroreflector</td>
<td>Make sure reflected beam and incoming beam parallel</td>
<td>Newport</td>
<td>UBBR1-1S 1343</td>
<td></td>
</tr>
<tr>
<td>Shaker</td>
<td>With the PZ-70 power supply, modulate the laser beam at low frequency</td>
<td>Homemade</td>
<td>Refer to shaker manual for more detail</td>
<td></td>
</tr>
<tr>
<td>Delay Stage</td>
<td>Vary the delay time between pump and probe beams</td>
<td>Newport</td>
<td>UTS100CC 3460.53</td>
<td></td>
</tr>
<tr>
<td>m12</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45 182</td>
<td></td>
</tr>
<tr>
<td>lens3</td>
<td>Combine with lens4</td>
<td>CVI</td>
<td>PLCX-25.4-25.8-UV-355- 266</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Description</td>
<td>Manufacturer</td>
<td>Part Number</td>
<td>Cost</td>
</tr>
<tr>
<td>---</td>
<td>-----------------------------------------------------------------------------</td>
<td>--------------</td>
<td>----------------------------------------------------------------------------</td>
<td>--------</td>
</tr>
<tr>
<td>1</td>
<td>as a telescope to reduce the 2nd harmonic beam size f=50mm</td>
<td></td>
<td></td>
<td>532</td>
</tr>
<tr>
<td>2</td>
<td>Laser grade UV fused silica lens, with 355-532nm AR coating f=250mm</td>
<td>CVI</td>
<td>PLCX-25.4-128.8-UV-355-532</td>
<td>225.00</td>
</tr>
<tr>
<td>3</td>
<td>Adjust the lens position for different wavelengths</td>
<td>Thorlabs</td>
<td>LNR25M</td>
<td>620</td>
</tr>
<tr>
<td>4</td>
<td>Adjust the 2nd harmonic path distance to make it overlap with the fundamental</td>
<td>Thorlabs</td>
<td>LNR25D (for laser path direction) LNR25M(for aligning purpose)</td>
<td>710</td>
</tr>
<tr>
<td>5</td>
<td>Adjust the lens position for different wavelengths</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Adjust the 2nd harmonic path distance to make it overlap with the fundamental</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45</td>
<td>182</td>
</tr>
<tr>
<td>8</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45</td>
<td>182</td>
</tr>
<tr>
<td>9</td>
<td>Change the beam height of 2nd harmonic</td>
<td>Newport</td>
<td>BSD-2A</td>
<td>401.75</td>
</tr>
<tr>
<td>10</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45</td>
<td>182</td>
</tr>
<tr>
<td>11</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45</td>
<td>182</td>
</tr>
<tr>
<td>12</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45</td>
<td>182</td>
</tr>
<tr>
<td>13</td>
<td>Mirror for 2nd harmonic</td>
<td>Photop</td>
<td>HR340/550-45</td>
<td>182</td>
</tr>
<tr>
<td>14</td>
<td>Dichroic mirror Reflect 2nd harmonic, transmit fundamental</td>
<td>Chroma</td>
<td>600dcoxru lot#239437</td>
<td>275</td>
</tr>
<tr>
<td>15</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S</td>
<td>464.07</td>
</tr>
<tr>
<td>16</td>
<td>Mirror for 3rd harmonic, interchangeable with m26 short for below 880nm</td>
<td>Photop</td>
<td>m20L: HR288/340-45 m20S: HR230/300-45</td>
<td></td>
</tr>
<tr>
<td>17</td>
<td>Mirror for 3rd harmonic, short for below 880nm</td>
<td>Photop</td>
<td>m20L: HR288/340-45 m20S: HR230/300-45</td>
<td></td>
</tr>
<tr>
<td>18</td>
<td>Mirror for 3rd harmonic</td>
<td>Photop</td>
<td>HR230/350-45</td>
<td>382</td>
</tr>
<tr>
<td>19</td>
<td>Mirror for 3rd harmonic</td>
<td>Photop</td>
<td>HR230/350-45</td>
<td>382</td>
</tr>
<tr>
<td></td>
<td>Description</td>
<td>Manufacturer</td>
<td>Model Number</td>
<td>Notes</td>
</tr>
<tr>
<td>---</td>
<td>-----------------------------------------------------------------------------</td>
<td>--------------</td>
<td>--------------</td>
<td>-------</td>
</tr>
<tr>
<td>lens5</td>
<td>Combine with lens 6 as telescope to reduce 3rd harmonic beam size f=50mm</td>
<td>CVI</td>
<td>PLCX-25.4-23.6-CFUV-248-355</td>
<td>530</td>
</tr>
<tr>
<td>lens6</td>
<td>CaF2 lenses with AR coating f=200mm</td>
<td>CVI</td>
<td>PLCX-25.4-93.2-CFUV-248-355</td>
<td>375</td>
</tr>
<tr>
<td>TS 5</td>
<td>Adjust distance between lens 5 and lens 6</td>
<td>Thorlabs</td>
<td>LNR25M</td>
<td>535</td>
</tr>
<tr>
<td>TS 6</td>
<td>Adjust the 3rd harmonic path distance to make it overlap with the fundamental</td>
<td>Thorlabs</td>
<td>LNR25D (for laser path direction) LNR25M(for aligning purpose)</td>
<td>620 535</td>
</tr>
<tr>
<td>m24</td>
<td>Mirror for 3rd harmonic</td>
<td>Photop</td>
<td>HR230/350-45</td>
<td>382</td>
</tr>
<tr>
<td>m25</td>
<td>Mirror for 3rd harmonic</td>
<td>Photop</td>
<td>HR230/350-45</td>
<td>382</td>
</tr>
<tr>
<td>d2</td>
<td>Reflect 3rd harmonic, fundamental and 2nd harmonic transmitted</td>
<td>Photop</td>
<td>HR230/350-45</td>
<td>382</td>
</tr>
<tr>
<td>m26</td>
<td>4th harmonic mirror, interchangeable with m20 30 degrees AOI</td>
<td>Photop</td>
<td>HR210/235-30</td>
<td>Come with the HG</td>
</tr>
<tr>
<td>m27</td>
<td>4th harmonic mirror, 30 degrees AOI</td>
<td>Photop</td>
<td>HR210/235-30</td>
<td>Come with the HG</td>
</tr>
<tr>
<td>lens7</td>
<td>Combine with lens 6 as telescope to reduce 4th harmonic beam size f=50mm</td>
<td>CVI</td>
<td>PLCX-25.4-23.6-CFUV-193-248</td>
<td>550</td>
</tr>
<tr>
<td>lens8</td>
<td>CaF2 lenses with AR coating f=150mm</td>
<td>CVI</td>
<td>PLCX-25.4-70.6-CFUV-193-248</td>
<td>395</td>
</tr>
<tr>
<td>TS7</td>
<td>Adjust distance between lens 7 and lens 8</td>
<td>Newport</td>
<td>Al made, not fancy</td>
<td>From Stock</td>
</tr>
<tr>
<td>m28</td>
<td>4th harmonic mirror, 45 degrees AOI</td>
<td>Photop</td>
<td>HR210/235-45</td>
<td>290</td>
</tr>
<tr>
<td>m29</td>
<td>Mirror for fundamental</td>
<td>Newport</td>
<td>10Q20UF.35S</td>
<td>464.07</td>
</tr>
<tr>
<td>m30</td>
<td>Mount with a flip mount, reflect the beam for diagnosis</td>
<td>CVI</td>
<td>DUVA-PM-1025-UV</td>
<td>220.8</td>
</tr>
</tbody>
</table>
In August, 2013, we found the 2\textsuperscript{nd} harmonic beam was not optimized; the beam is too broad when it meets lens 3 and lens 4. So we revised the 2\textsuperscript{nd} harmonic path, lenses 3 and 4 are now in the later position, the 2\textsuperscript{nd} harmonic beam now travels at 3 inches height for long distance. A few mirrors are added as below.
A.2 MaiTai-DeepSee

To connect the MaiTai or DeepSee controller to our PC, we can use either RS232 cable or USB-RS232 adaptor. In case of using the latter, we have to use the National Instrument RS-232 USB adaptor, otherwise there is software crashed frequently somehow.

The laser beam coming out of the MaiTai has a horizontal polarization. We used the periscope with s- and p-polarized dielectric mirrors for femtosecond pulse laser to change the polarization to vertical since LPC and the harmonic generator require the vertical polarization for input beam. The repetition rate actually depends on the wavelength.

<table>
<thead>
<tr>
<th>Table A.2 Mai Tai Frequency Dependence of Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \lambda ) (nm)</td>
</tr>
<tr>
<td>-------------------</td>
</tr>
<tr>
<td>700</td>
</tr>
<tr>
<td>800</td>
</tr>
<tr>
<td>900</td>
</tr>
<tr>
<td>940</td>
</tr>
<tr>
<td>950</td>
</tr>
<tr>
<td>1000</td>
</tr>
</tbody>
</table>

There is a huge RF interference peak at 80MHz picked up by our RF detection circuit, so the wavelength near 940nm might not be used.

DeepSee unit is attached to the end of MaiTai output inside the laser box and can provide the
computer-controlled group velocity dispersion (GVD) compensation in order to achieve the shortest pulses possible at the sample. This compensation is done by moving the motor that moves optical materials on a stage. DeepSee GUI screen shows the range of the motor which are determined by upper and lower “soft limits”. The smaller percent values correspond to a lower absolute value of GVD. We found the shortest pulse width at each wavelength was achieved when the motor is at the lower limit by examining interference signal from the autocorrelation stage. Since the pulse width is minimized, the output power from the harmonic generator described in detail below is also maximized, more than twice with the DeepSee as that without it.

A.3 LPC

After turning on the power of LPC controller, LPC shows the minimum power and transmission state, which means that the polarization of the incoming beam is vertical according to BEOC. The glass mounted on the circuit board is the LC cell and the cubic crystal behind that is output polarizer, there is NO input polarizer in our LPC (figA.1).

The laser power will fluctuate if the input beam size is bigger than the input aperture which is 4mm. Power threshold of LPC is 4W/4mm, while MaiTai has the max power of 2.75W (2.6W with DeepSee) @ 800nm. We reduced the beam size from 4mm to 3.5mm by using the pair of Fused Silica lens with $f=65$mm and 75mm, and now the power threshold for 3.5mm is 3W.

A.4 Harmonic generator (TP 2000B Tripler, Photop Technologies)

2\textsuperscript{nd}, 3\textsuperscript{rd} and 4\textsuperscript{th} harmonic beams are generated here. The 1\textsuperscript{st} waveplate depicted on the manual has been removed from our setup since our input beams has a vertical polarization. Read the manual carefully (figA.2). Sheng gave us a set of crystals and here is the table. The crystals with boldfaced name are being used now.
### Table A.3 Nonlinear Crystals for Harmonic Generator and Diagnosis

<table>
<thead>
<tr>
<th>Harmonic</th>
<th>Material (BBO or LBO)</th>
<th>Thickness</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>60fs 2\textsuperscript{nd} harmonic</td>
<td>5x5x1.1mm LBO</td>
<td>1.1mm</td>
<td>680-1,100nm</td>
</tr>
<tr>
<td>100fs 2\textsuperscript{nd} harmonic</td>
<td>5x5x2mm LBO</td>
<td>2mm</td>
<td>680-1,100nm</td>
</tr>
<tr>
<td>60fs 3\textsuperscript{rd} harmonic</td>
<td>7x7x0.6mm BBO</td>
<td>0.6mm</td>
<td>680-1,100nm</td>
</tr>
<tr>
<td>100fs 3\textsuperscript{rd} harmonic</td>
<td>7x7x1mm BBO</td>
<td>1mm</td>
<td>680-1,100nm</td>
</tr>
<tr>
<td>60fs 4\textsuperscript{th} harmonic</td>
<td>7x7x0.6mm BBO</td>
<td>0.6mm</td>
<td>680-1,100nm</td>
</tr>
<tr>
<td>100fs 4\textsuperscript{th} harmonic</td>
<td>7x7x1mm BBO</td>
<td>1mm</td>
<td>680-1,100nm</td>
</tr>
<tr>
<td>2\textsuperscript{nd} auto crystal</td>
<td>8x5x0.5mm BBO</td>
<td>0.5mm</td>
<td>600-1,100nm</td>
</tr>
<tr>
<td>3-1 or 1+2 auto crystal</td>
<td>10x10x0.4mm</td>
<td>0.4mm</td>
<td>720-900nm</td>
</tr>
</tbody>
</table>

Here is some note we’d better keep in mind when optimizing THG and FHG in the harmonic generator (You can refer the layout of the optics inside HG and the generated laser path in the above photos).

1. Make sure the incoming fundamental beam go through near the center of the SHG crystal since the beam will slightly diverts as $\lambda$ is changed.
2. Find “absolute zero” before calibration so that actuators can rotate the crystals with max range. First, using axis positioner window of TPI, send large negative value to each axis until – limit is lit on. Then turn off the motor and manually turn the actuator toward the positive direction until “-limit” was off. Finally press zero to reset. Now the positioner is at zero position.

3. Make sure SHG spot can be optimized within the range of the angle of the SHG crystal for all wavelengths.

4. Make sure arrows of time plate (TP) and SHG crystal are always antiparallel otherwise building up. Currently the crystal orientation of TP is Arrow Downward, while that of SHG crystal is Arrow upward.

5. For THG, we need to use THG crystal as well as the time plate (TP) with M3 and M4 mirrors labeled “S(hort)” for $\lambda < 900$nm. M3 and M4 mirrors must be replaced by those labeled “L(ong)” for $\lambda \geq 900$nm.

6. For FHG, take out TP and swap the crystal from THG to FHG. There are thick green line on the crystal holder and scribed mark on the post holder so that we can align them. We must use M3 and M4 mirrors for FHG, but keep in mind that incident of angle of FHG mirror is $30^\circ$.

7. To optimized THG(FHG) signal, need to adjust F1, F2 and both angles of WP2 ($\theta,\phi$)

Another problem we found was that the beam size of higher harmonics got broader with length. We tried reducing the beam size with a pair of plano-convex lens in the telescope configuration.
(d₂/d₁=f₂/f₁). The resulting beam size, however, was not reduced and collimated at all. Instead, we could happen to reduce the size by swapping the lens so that lens with the shorter focal length is placed in front of lens with the longer focal length. It’s also recommended that plano-convex UV grade CaF₂ lens with BBAR coating would better for shorter wavelength such as THG and FHG. The combinations of lenses we chose are

THG: CaF₂ lens, f₁=50mm, f₂=200mm, BBAR 248 to 355nm coating for the UV region
FHG: CaF₂ lens, f₁=50mm, f₂=150mm, BBAR 193-248nm coating for the UV region

According to Sheng, we need a faster lens to focus the beam first since the FHG (THG) beam is diverging a lot before lens.

With these pair of lens, the THG beam is reduced to 1.5mm x 1mm at far field (115” away from the lens with f₂) where 1.5mm is measured along the direction vertical to the table. The FHG beam size is ~2mm x 2mm. SHG beam coming out of the harmonic generator has the horizontal polarization. We change the polarization by using two periscopes. The beam size of SHG is again reduced with the telescope configuration of a pair of lenses:

SHG: Fused Silica lens, f₁=50mm, f₂=200mm, BBAR 355-532nm coating for the UV region

A.5 Autocorrelation stage

Our autocorrelation (AC) stage is the Michaelson interference setup consisting of two arms, one fixed and another movable by the linear translation stage (ESP 301-2G Newport) controlled by Labview program through GPIB cable. We use the delay stage UTS100CC from Newport (with a mounting plate UTS-TP from the same company) in the fundamental beam to get the delay τ between the 2 beams. The temporal resolution of the stepping motor is 1 en (0.1 micron), so the temporal resolution is 0.667fs/en. The fundamental beam is guided to a Michelson Setup as in the Figure A.3. One arm has the delay stage, retroreflector and shaker. The other arm has a mirror
pair mounted on a plate, which is connected to a 3-direction translation stage. Initially we also want to use the mirror pair on the delay stage arm since the mirrors are specially made for MaiTai laser and have higher reflectivity. But it turns out with the machine shop made mirror-mount; it’s very hard to keep the beam constant horizontally when the stage moves. It’s much easier to make the two fundamental beams overlap after we use the retroreflector UBBR1-1S from Newport.

Different from the previous fs laser setup in STM2 (which has the delay stage and shaker in different arms), we have the delay stage and shaker in the same arm. The main reason is that we want to do pump-probe experiments with 2 different color beams (fundamental and 2nd harmonic, fundamental and 3rd harmonic), to make things easier, the delay stage and shaker are in the same arm. However, it’s still valuable to have 2 fundamental beams overlap; we can characterize the beam with autocorrelation.

The shaker to modulate the delay is home built, since PZ-80 from Burleigh has been obsolete for long time, we used a piezo tube (EBL# 2 Piezoceramic Tube 1.00” OD x 0.060” Wall x 2.00” Length, gold over nickel electrodes) mounted in a homemade pieces designed, a detailed drawing is shown in Fig A.4~FigA.6. The housing is made of aluminum, electrically isolated from the piezo with two macor pieces; all these pieces are designed and machined as light as possible and glued together by Varian torr seal. The retroreflector is connected to the front macor piece through a homemade connector. The controller of the piezo is a Burleigh PZ-70 (we got a manual from company). We use Viking Thorkom connector from Source Research Inc. to apply the high voltage from PZ-70 to the piezo. Parts list is as follows:
Table A.4 Homemade shaker commercial electric parts

<table>
<thead>
<tr>
<th>Socket Contact</th>
<th>TS-100-AU</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plug</td>
<td>TNP07-101</td>
</tr>
<tr>
<td>Dummy Filler</td>
<td>TD-200</td>
</tr>
<tr>
<td>Bulkhead Receptacle</td>
<td>TBR07-101-P</td>
</tr>
<tr>
<td>Backshell .150-185 Cable Dia</td>
<td>TSR07-201</td>
</tr>
<tr>
<td>Pin Contact</td>
<td>TP-100-AU</td>
</tr>
</tbody>
</table>

There are 7 pins in the TKR-07 receptacle, among them 3 are HV and 4 are ground. Modulation output from EG&G 7265 lock-in amplifier was fed into PZ-70 as an input sine wave. It’s necessary to tune the DC bias voltage to a few hundred volts such that the amplified sine wave signal is always positive, otherwise PZ-70 will shut down. The modulation frequency is 220 Hz.

For example, the amplitude from the lock-in is 1.05V (rms) and turn the Gain knob to increase the voltage until the output/100 BNC connected with the oscilloscope shows 5 V (p-to-p), which corresponds to ~500V applied to the piezo shaker. 4V p-p voltage on the monitor output (about 400V on the HV output) gave us the clearest dS/dτ. Other parameters of the lock-in for dS/dτ and d^2S/dτ^2 are TC 100ms, sensitivity 50mV for 1st harmonics and 1mV for 2nd harmonics.

We did the piezo shaker calibration as follows. The first calibration is to obtain the relationship between the HV output and the monitor output,

It is: \( HV_{output}(V) = 100.61 \times MonitorOutput(V) + 0.2842. \)

A second calibration is the relationship between the piezo motion with the monitor output, it’s done by comparing the time zero change in the autocorrelation spectra with tuning the bias voltage.

The relationship is \( piezo(\text{microns}) \times 2 = -1.13 \times MonitorOutput(V) - 0.08. \)

Note that the piezo shrinks at the positive voltage applied.

**A.6 Simulation Path**
We have two BBO crystals, one for 1+1 AC (crystal orientation is vertical to the table) and another for 3-1 or 1+2 AC. For 3-1 AC, the fundamental needs to be perpendicular to THG, so we use WP1 of the harmonic generator which hasn’t been used to change the polarization. The black lines on the crystal holder should be horizontal to the optical table surface. Since each of the fundamental and THG generates SHG at BBO, we need minimize these signals to distinguish 3-1 SHG signal from them. Empirically SHG spot generated from the fundamental can be minimized by adjusting the BBO crystal position and angle. We usually monitor the beams after BBO with CCD camera to check the spatial overlap and check the temporal overlap by moving the translation stage of THG. Once you see the 3-1 SHG spot, you can maximize the signal by adjusting the crystal position, angle and fine tuning the optical path. A picture below shows the BBO crystal orientation for $\omega=880\text{nm}$ for the 3-1 autocorrelation. Note the polar angle of the crystal with respect to the laser path is rotated to maximize the AC signal. For the 2+1 autocorrelation, the crystal must be rotated along the azimuthal angle with respect to the laser path.

As for the in-plane crystal orientation, the black lines marked on the rotational crystal mount, should be parallel to the SHG and the fundamental input polarizations for 2+1. In this case, it should be vertical to table, i.e., "both fundamental and the second harmonic beams are vertical to the table as well as the BBO crystal orientation". For 3-1 case, the crystal orientation must be parallel to the fundamental beam polarization, which was already rotated to the horizontal polarization by the half waveplate, i.e., parallel to the table.

To detect AC signals for 3-1 and 2+1, we need a color filter such as one used for 1+1 AC measurements. We used the same color filter as 1+1 AC for 3-1 and UG11 color filter by Schott
for 2+1 AC measurement. Optical density data is on the STM4 manual and also could be got through the company website.

**A.7 Beams into Junction**

To focus the beams into the junction, we use a lens inside the UHV chamber (focal length 3 inches); the lens is mounted on the lens holder assembly for adjusting. The assembly drawings can be found in the STM4 folder. To get a 3-D view of the junction through the pyro viewport, we need to use a finder scope. We can only see fundamental and 2\textsuperscript{nd} harmonic with the commercial finder scope. To see 3\textsuperscript{rd} and 4\textsuperscript{th} harmonics in the junction, we need a finder scope with fused silica lenses, which is not commercially available. We assembled a home-made finder scope with the fused silica lenses and cage assembly from Thorlabs, POs of these parts could be found in the STM4 laser system POs. The home-made finder scope works though it could be improved to a better cage system or lenses.

![Inside of LPC](image)

**Fig A.1 Inside of LPC**
Fig A.2 Inside of the Harmonic Generator

FigA.3 (A) Schematic of the Michaelson interference setup, (B) Photo of the delay stage and shaker.
FigA.4 (A) homemade Al housing body for the shaker piezo, (B) Al cover for the housing.
Fig A.5 (A) macor piece attached to the piezo and Al mating piece, (B) macor piece attached to the piezo and Al housing.
Fig A.6 Al mating piece to connect the macor piece and retroreflector.
FigA.7 (A) BBO crystal position for “2+1” simulation at 840nm and 420nm, (B) crystal position for “3-1” simulation at 880nm and 293nm.
Appendix B

STM4 Doser Loadlock and Doser

B.1 New Lab

To couple to the new femtosecond laser described in this thesis, we need a new lab with darkroom capabilities. So the first major task for me was to move STM4 (which was in the main lab between STM1 and STM3) to the previous storage room with my partners, fig B.1 shows the initial stage and later stage of the moving, (A) shows the lab after we cleaned up and relocate the optical table, (B) shows the lab after we moved the STM and gas system. It took a few months to move the system and debug, making it back to work.

B.2 New Loadlock and Other Ports Change

Before as shown in figB.2 (A), every time we want to study a new molecule or metal atom, we need to vent the whole chamber, it usually takes ~4 weeks to bake and degas until the system is ready to run experiments again. So it’s desirable to have an evaporator interlock system to enable rapid turnaround of metal and molecular dosers that can be made without venting the vacuum chamber, thus allowing fast screening of molecules and metal atoms. The modified system with the new loadlock is shown in fig B.2 (B), a zoomed in image of the loadlock is shown if fig B.3 (A). To shine laser beams into the junction, two lens holder assemblies are mounted on the two sides of the chamber (3.38 ports, through hole), before dosers were mounted there. Before the gas doser was mounted on the side and a molecule doser was mounted in the front port, later we mounted Mg or C\textsubscript{60} dosers on the side to form MgO on Ag (100) or C\textsubscript{60} layer on Al\textsubscript{2}O\textsubscript{3} at RT.
The gas doser now is mounted in the front with a linear stage; the gas tube can get really close to the scanner door, enabling a more effective gas or molecule dose from the gas line.

The main parts of the loadlock are a small ion pump, a getter pump, a linear translator, two gate valves, a gold seal valve and tees (crosses, reducer and nipple). The flange on the port which loadlock was assembled has a tap hole, so a short nipple is used before the gate valve, which also has tap holes. After the gate valve is a tee with a viewport, the linear translator for the dosers is attached to the straight side of the tee while the pumps connect through the other side. A six-way cross is connected to the tee first on the pumps side, one side the six-way cross is connected to the gold seal valve, the other side connects to the ion pump and getter pump through a gate valve. Three ports are blanked on the six-way cross now for later improvements like gauge or RGA. The ion pump and getter are mounted on a 4.62 inches tee, enabling bigger volume for the getter. The process to change doser is:

1 close both gate valves,
2 vent the load lock by opening gold seal valve,
3 take the old doser out,
4 put the new doser in,
5 connect the load lock to pumps,
6 start pumping, wait till the pressure reaches lowest,
7 open the valve near the pumps, activate getter pump if necessary, start baking.

B.3 New Dosers

Before the Ho group dosers use a 3.38 flange, same as the lens holder ports, the other ports are 2.75” flange, so we need a reducer to mount the dosers. Now with a 2.75 inches linear translator, new dosers with 2.75 inches are designed, machined and assembled; the vacuum length of these
dosers is 12.5”~13 inches. Fig B.3 (B) shows a doser with water cooling. On a 2.75” flange, three holes are drilled, the center hole is for water cooling; one of the side holes is for electrical feedthrough and the other one for rotational feedthrough. The watercooling system contains a 0.375” out tube, a 0.25” inner tube and a machined water tank top, it’s very important to bend the out tube and inner tube same time before welding. It’s also very important to weld the tube concentric to the flange, usually, we can cut several tubes and roll on the flat table in the machine shop and choose the straightest one. The electrical feedthrough is from CeramTec (electrode diameter 0.094 inches); the rotational feedthrough is from MDC (less expensive ones may be found at Huttington). Home-made adaptors are used to connect the electrical feedthrough and rotational feedthrough to the 2.75 inch flange. One more hole could be made on the flange to mount the thermocouple if needed. The electrical feedthrough is connected to 30mil or 20mil W wire (which wound around a ceramic crucible) through Cu rod and Ta rod, a shaft collar is needed to fix the rods. The ceramic crucible is made by making threads on a one-end-sealed ceramic tube (0.313 inch OD, 0.188 inch ID) from Ortech. Fig B.4 (A) shows the drawing of the doser after machining and welding, fig B.4 (B) shows the dimension of the welding holes on the 2.75” flange. Fig B.5 (A) shows the dimension of the one-end sealed water tank; fig.5 (B) shows the dimension of the adapter to weld to the 2.75 inches flange to connect the electrical feedthrough.

Fig B.3 (C) shows a dual-doser with two crucibles mounted, enabling co-evaporation of two molecules same time onto the surface, the inset shows the top view of the two crucibles. It needs two smaller crucibles to fit into the space and a good isolation between the two crucible housings; two pairs of electrodes are needed to heat up the crucibles. Fig B.6 (A) is the drawing after the doser is assembled. The top shield and crucible structure are supported by two 6-32
screw rods, given that the top structure is heavy and the doser is 13” long, stainless steel tubes are needed to outfit the screw rods to stabilize. A home-made shaft collar is needed to clamp in the middle. Fig B.6 (B) shows the dimension of the welding holes and tap holes on the flange and fig B.6 (C) shows the dimension of the shaft collar. Fig B.7 shows the housing structure for the two crucibles, it’s made up of three pieces, the body structure as shown in (A), the separating piece as shown in (B) and the cover as shown in (C). To make smaller crucibles for the new dual doser, new jig to cut the ceramic tubes is needed, the design is shown if figB.8, (A) shows the mail piece, (B) shows the female piece and (C) shows the clamping piece.

Below is a table of different dosers for STM4:

<table>
<thead>
<tr>
<th>Doser Material</th>
<th>Doser Type</th>
<th>Crucible and Current</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg</td>
<td>3.38 flange, to chamber, face junction or heater, w/ water cooling</td>
<td>Old fat crucible, 28~32A</td>
</tr>
<tr>
<td>K</td>
<td>2.75 flange, load lock, strip from SAES getter, face junction, no need of water cooling</td>
<td>The strip is spot welded to the two Ta rods, 5~6A</td>
</tr>
<tr>
<td>Fe</td>
<td>3.38 flange, to chamber, face junction, w/ water cooling</td>
<td>Ceramic Crucible wound with 30 mil W wires, 16~18A</td>
</tr>
<tr>
<td>C60</td>
<td>3.38 flange, to chamber, face heater, no water cooling</td>
<td>Ceramic Crucible wound with 20 mil W wires, 3.4A~3.8A</td>
</tr>
<tr>
<td>Ag</td>
<td>3.38 flange, to chamber, face heater, w/ water cooling</td>
<td>Ceramic Crucible wound with 30 mil W wires, 11~12A</td>
</tr>
<tr>
<td>Au</td>
<td>3.38 flange, to chamber, face junction, w/ water cooling</td>
<td>Ceramic Crucible wound with 30 mil W wires, 14~15A</td>
</tr>
<tr>
<td>Th-Th-BT-Th-Th</td>
<td>2.75 flange, load lock, face STM junction, w/ water cooling</td>
<td>Ceramic Crucible wound with 30 mil W wires, around 2.75A</td>
</tr>
</tbody>
</table>

Table 1 Evaporators of STM4
Fig B.1 (A) Initial stage of the new STM4 lab (which used to be the storage room) after cleaning and setting up new optical table, (B) latter stage of the new STM4 lab after setting up the STM table and pumping systems.
FigB.2 (A) Former STM4 setup, with no optical setup or load lock, it should be also noted that the gas doser is on the side and a molecule doser is in the front; (B) new STM4 setup, with load lock and optical setup, the gas doser is now in the front with a linear translator and a molecule or metal doser is mounted on the side facing the heater.
Fig B.3(A) STM4 doser loadlock system, (B) a load lock doser with water cooling, (C) a load lock doser with two crucible (inset) mounted.
Fig B.4 (A) Load lock doser with water cooling after welding, this one has a third miniflange for thermocouple, (B) two possible flange welding holes dimension, the shutter hole can be small.
FigB.5 (A) water tank top, bottom is welded to the water flowing outer tube, top attached to the Ta shield, (B) welding adapter, top welded the 2.75” flange and bottom is welded to the minflange.
Fig B.6 (A) Dual molecule evaporator with 2 crucibles, (B) flange holes dimension, (C) shaft collar dimension.
FigB.7 (A) housing body dimension, (B) housing middle piece dimension, (C) housing top piece dimension.
FigB.8 (A) mail piece for the jig, (B) female piece for the jig, (C) top clamping piece for the jig.