Field Emission Enhancements in Carbon-implanted Group 5B Metals

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by

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Abstract:

Vanadium, Niobium, and Tantalum foil samples were implanted with high-energy carbon ions by plasma source ion implantation. Carbide formation was confirmed in some cases by x-ray photoelectron spectroscopy. The relative work functions of the samples were measured with a Kelvin probe, and work function calculations were calculated from current versus voltage measurements performed by scanning tunneling microscopy. Kelvin probe and STM measurements were performed on unimplanted samples of the same metals. Work function enhancements are discussed in terms of nano-scale surface roughness, as measured by high-resolution tapping-mode atomic force microscopy.
I. Introduction

In recent years, there has been much research into field emission enhancement from carbon-containing cold cathodes, including carbon, boron, and sodium implants into diamond produced by chemical vapor deposition (CVD), CVD diamond thin films, carbon nanotubes (CNT), and implanted carbon layers on the surface of transition metals and dielectrics. Research into diamond in particular has shown it to be an excellent emitter at low voltages, possibly due to the negative electron affinity (NEA) of certain hydrogenated diamond crystal planes. On the other hand, CNT field emitter arrays have received recent experimental attention thanks to the high aspect ratio and subsequent magnified local electric field of the nanotube structure, despite its low robustness. Carbon ion-implanted layers are often also marked by this enhanced local electric field largely due to physical grain boundaries and surface defects and electrical inhomogeneities inherent in the implantation process.

It is well known that the physical surface features – especially those with a high aspect ratio – and the electrical inhomogeneities introduced during implantation of
carbon ions produce regions of local electric field enhancement, a critical condition to electron emission. Field emission, also known as Fowler-Nordheim tunneling, occurs when a high electric field incident on the surface of a metal lowers its potential barrier to the point that electron emission is possible via quantum mechanical tunneling. This variety of electron emission is thus distinct from photo-emission and thermionic emission, in which electrons are given sufficient energy to escape over the work function, otherwise known as the local barrier height (LBH). For this reason, the utility of field emission devices is their activity at low temperatures and their minimal activation times, making field emission widely usable as a current source across thin barriers in metal-semiconductor junctions and in field emission microscopy, femtosecond cameras, and field emission displays (FED’s). Field emitters also find use in electron beam microlithography and scanning electron microscopy because they are point sources to first approximation, with a radius on the order of an angstrom.\textsuperscript{7} The greatest hurdle researchers must overcome with current field emission technologies is coping with a lack of robustness due to the large aspect ratio of field emitter arrays (FEAs), which are easily damaged by mass electron transport, especially at high voltages.

A. Field Emission from Transition Metal Carbides

There is a substantial body of work on the subject of the role of carbides in enhancing the field emission properties of transition metals. Mackie et al. measured work functions of group 4B carbides, specifically ZrC and HfC, to be in the range of 3.5 eV and observed field emission current densities on the order of $10^8$ A/cm\textsuperscript{2} on single tips.\textsuperscript{8,9,10} By depositing ZrC films onto Mo and W field single field emitter cathodes, they were able to repeatably and consistently lower the work function of the cathode.
material and improve current stability and beam confinement of the array. To determine whether the subsequent field enhancement was a result of a change in surface work function or in the enhancement factor $\beta$, a family of curves with constant $\phi$ was plotted against a family of curves with constant $\beta$. The plot’s domain was the FN intercept, and the range was the FN slope. For the case of ZrC films deposited onto a W tip, the graph suggested a small decrease in the $\beta$ factor, corresponding to an increase in the tip radius, and a more substantial decrease in the surface work function, as shown in figure 1.

![Figure 1](reprinted with permission from Mackie et al.)

The relation between the tip radius and enhancement factor was assumed to be:
\[
\beta = \left[ 0.5r \ln \left( \frac{D}{r} \right) \right]^{-1}
\]

where \(D\) is the tip to screen distance in an FEM tube.\(^{13}\)

Then by assuming only a small increase in tip radius, the change in work function due to deposition is given by the FN ratio:

\[
\text{Ratio}_{FN} = \left( \frac{m_{\text{deposited}}}{m_{\text{clean}}} \right)^{2/3}
\]

where \(m\) is the slope of the FN curve for the emitter before and after ZrC deposition.\(^{11}\)

Using a similar procedure, Xie et al. were able to obtain a work function decrease of 0.6 eV after depositing ZrC on single crystal Si Spindt-type single emitters and a decrease on the order of 1 eV for Mo polycrystalline wire. Again, the decrease in field enhancement factor was sufficiently small to calculate FN ratios that corresponded with the change in work function due to deposition.\(^{14}\)

Carbides of hafnium, one period below Zr in group 4B, have been shown to have lower work functions than ZrC by several tenths of an eV.\(^{15,12}\) Similarly, Mackie et al.\(^{13}\) have demonstrated that deposition of HfC films on W single tips, Mo single tips, and Mo FEA's reduces surface work function below that of experiments with ZrC deposition on the same substrates. In addition to the success of ZrC and HfC in lowering work functions beyond those of uncoated tips, these carbides boast some of the highest melting points of all materials and thus can maintain functionality at high temperatures and fields without becoming subject to atom migration. ZrC and HfC usually exist in an fcc crystal structure but are found with stoichiometries as low as \((C/Zr) = 0.84\). Nonetheless, Mackie et al.\(^{16}\) have shown optimum current stability, optimal resistance to surface
contamination, and minimal noise levels with (C/Zr) = 1. Using x-ray photoelectron spectroscopy (XPS) to measure carbon content of implanted silicon, Chen et al.\textsuperscript{17} found the most preferable emission characteristics to come from the more stoichiometric films also in the case of deposition of SiC on to an Si tip surface.

Tsang et al. also studied SiC deposition, characterized by high dose carbon implants into silicon and showed that both surface morphology and electrical inhomogeneities lent a hand in producing local electric field enhancement.\textsuperscript{18} Using conducting atomic force microscopy (CAFM), he found localized graphitic clusters in samples that were not annealed. Thermal annealing, however, produced a more uniform SiC layer, destroying the clusters. Subsequent XPS experiments revealed the presence of graphitic sp\textsuperscript{2} carbon in the as-implanted samples and its absence after annealing, prompting the conclusion that electrical inhomogeneities were eliminated by complete carbonization during the first hour of thermal annealing at 1200 °C. After longer annealing times, small protrusion structures became detectable on AFM, leading to local electric field enhancement.

In this work, plasma-sourced ion-implantation (PSII) of Group V transition metals was performed to produce a thin carbon-implanted layer. In cases where implantation yields a stable carbide, experimenters expected reductions in the work function beyond that of the unimplanted metals, as seen in previous studies of other groups. In cases where carbide formation is minimal or inexistent, the effect on field emission is entirely unknown. Experimental work on the field emission enhancing properties of carbon largely surpasses theoretical work, but several groups have put forth explanations on why carbon implants and films enhance field emission.\textsuperscript{19,20} The fact that carbon has very low or negative electron affinity -- and the conduction band lies close to or even above the
vacuum level -- makes it energetically favorable for an electron in that band to break from the atom. Burden\textsuperscript{20} also suggests that the coexistence of carbon hybridized to $sp^2$ and $sp^3$ orbitals in the same region of a carbon-containing compound produces inclusions and grain boundaries and hence regions of locally amplified electric field, thereby making electron emission easier.
II. Theory

A. Fowler-Nordheim tunneling

Fowler and Nordheim first described the dependence of electron emission on a high electric field as a simple barrier penetration situation, in which an incident electric field so deforms a potential barrier that low-energy electrons may tunnel through the classically forbidden region. The Heisenberg principle requires a correlation between uncertainty in position and momentum of a surface electron. Supposing that an electron near the surface of the material has a positional uncertainty on the order of the width of the potential barrier, tunneling through the barrier is possible, as predicted by the Wentzel-Kramers-Brillouin (WKB) method for one-dimensional potentials. The current is predicted as:

\[ I = \int_{0}^{eV} \rho_{s}(r,E)\rho_{s}(r,-eV + E)\Gamma(E,eV,r)dE, \]
where \( \rho_s \) and \( \rho_t \) are the density of states of the sample and tip, respectively, \( eV \) is the sample bias, \( E \) is the electron energy, \( r \) is located directly under the tip, and \( T \) is the transmission probability, given by:

\[
T(E, eV) = \exp \left[ -\frac{2z\sqrt{2m}}{h} \sqrt{\phi_s + \phi_t} + \frac{eV}{2} - E \right],
\]

for work functions \( \phi \) and tip-sample distance \( z \).\(^{21}\)

In the case of an incident electric field on a one-dimensional metallic surface, the WKB method describes a triangular potential. Multiplying the WKB probability by the differential rate of electron arrival at the surface, and integrating the electron kinetic energy from zero to the Fermi level, Fowler and Nordheim arrived at an expression for current density, valid for an instantaneous potential barrier.\(^7\)

![Figure 2](reprinted from Gomer\(^7\))
For a more realistic case, use of a classical image potential correction is necessary because no accurate model for the potential of electrons at a surface exists. The image potential only slightly alters the potential step, rounding it off at the corners, as shown in figure 2. This more general Fowler and Nordheim equation provides for a rather accurate prediction of tunneling electron current for field emitters at voltages greater than the potential barrier:

\[
    J = \frac{1.54 \times 10^{-6} E^2}{\phi} \exp \left[ -\frac{6.83 \times 10^9 \phi^{3/2}}{E} \right],
\]

where \( J \) is the current density of emitted electrons in A/m\(^2\), \( \phi \) is the work function in electron volts, and \( E \) is the local electric field strength in V/m.\(^{22} \) Numerical factors are derived from geometrical considerations, including the assumption of a point-like tip and a field emitting area in the shape of a perfectly flat infinite plane, eliminating dependence on surface topography. Thus, from the Fowler-Nordheim equation, one may derive the explicit electric field dependence of the work function:

\[
    \Delta \phi = \frac{eE}{4\pi \varepsilon_0},
\]

valid for fundamental charge \( e \), electric field \( E \), and permittivity of free space, \( \varepsilon_0 \).\(^{23} \)

In the presence of surface adsorbates, Miller predicted a secondary potential well a distance \( z \) outside of the initial potential step, making it possible for an electron to tunnel through two smaller gaps, which may significantly increase electron tunneling probability.\(^{24} \) Gomer\(^7 \) presents a work function adjustment due to adsorbed surface double layers, in which the dipole moments (to first approximation) of the adsorbed species alter the surface charge distribution. Under the assumption that the layers are planar and parallel, consequent enhancement of the work function rests solely on the
number of dipoles and their strengths, aside from a solid angle factor of $4\pi$. Plausibly assuming the dipole is centered on the electrical midpoint in the double layer, such that an electron need only do work against half of the layer, Gomer concludes:

$$\Delta \phi = 2\pi P_i N_s \theta,$$

where $P_i$ is the dipole moment of an adsorbed particle, $N_s$ is the total number of adsorption sites per unit area, and $\theta$ is the fraction of filled sites.\(^7\) The surface dipole assumption is also valid for surfaces without adsorbed layers and is likewise dependent on the number of dipole sites. More loosely packed surfaces lead to atomic scale corrugations on the surface and the subsequent formation of dipoles oriented in the opposite direction of the surface dipole, as seen in figure 3. The result of this phenomenon, known as the Smoluchowski effect, is an overall lowering of the macroscopic work function.\(^{25}\)

![Figure 3](image-reprint.png)
B. Plasma Source Ion Implantation

During the PSII, the target to be implanted is submerged in an ionized plasma and is then pulsed with a high negative voltage, in this case -35 kV at 25 Hz. The target and mount are electrically isolated from the chamber walls by a quartz rod. The pulse repels free electrons throughout the plasma while attracting the ions, which are subsequently implanted into the surface of the target. Anders\(^{26}\) presents a planar model of a sheath, assuming collisionless ion flow, zero electron mass, an instantaneous voltage pulse, the instantaneous formation of the ion sheath, singly charged ions, and that the implant current wholly provides the charge of the expanding sheath. By equating the current density as derived by Child’s law to the charge per unit time crossing the boundary of the ion sheath, Anders arrives at the sheath velocity:

\[
\frac{ds}{dt} = \frac{2 s_0^2 u_0}{9 s^2} ,
\]

for

\[
s_0 = \left( \frac{2 \varepsilon_0 V_0}{en_0} \right)^{\frac{1}{2}} \quad \text{and} \quad u_0 = \frac{2eV_0}{M}
\]

where, \( s \) is the sheath position, \( s_0 \) is the thickness of the ion matrix sheath, \( \varepsilon_0 \) is the permittivity of free space, \( V_0 \) is the pulse voltage, \( e \) is the fundamental charge, and \( M \) is the ion mass.\(^{27}\) Analytically integrating the differential equation produces

\[
s = s_0 \left[ \frac{2}{3} \omega_{pi} t + 1 \right]^{\frac{1}{3}} ,
\]

where \( \omega_{pi} \) is the ion plasma frequency.\(^{28}\) Assuming that the target current is only supplied by ionization during sheath expansion, this solution allows us the average ion current per pulse as
\[ I_{ion} = \left( \frac{enA}{I_{pulse}} \right)^{s_f}, \]

with \( s_f \) the final sheath extent as calculated by the above analytical solution to the sheath velocity differential equation.\(^{25}\)

For a low-pressure plasma at high pulse voltage and small pulse-width, these assumptions are good approximations. The plasma in this work, however, consists of multiple ions of different masses and charges. The sheath velocity of a plasma of this consistency must be modified by replacing the ion mass with an effective mass, given by:

\[ \sqrt{M} = \sum_j k_j \sqrt{Z_j M_j} \]

with \( Z_j \) and \( M_j \) signifying atomic number and mass for each ion \( j \) and \( k_j \) defined as the fractional density of ion \( j \) as follows: \(^{24}\)

\[ k_j = \frac{n_j}{\sum_z Z_n_z}. \]

As the voltage pulse continues, the sheath of ions expands radially outward from the pulsed sample. When the pulse ends, however, the sheath collapses as the charge from the sample and holder is discharged through the electronics, and the neutralized target is again ready for the next pulse. The plasma, on the other hand, may still be nonuniform by the time of the next pulse, depending on repetition rate and sheath thickness. This situation causes the next pulse to expand the ion sheath further, and if left unchecked, creates the possibility of arcing with the chamber walls on subsequent pulses. This effect is more pronounced in low density plasmas because of the wide sheath produced with a single pulse. Anders \(^{26}\) is quick to point out that the pulse expansion does eventually reach a steady state in much less time than is required for the total implant.
PSII is a low-temperature surface processing technique that causes minimal thermal distortion of the sample with no macroscopic change in dimensions and produces no degradation in the surface finish. Between pulses, the target to be implanted loses heat energy by radiation and conductive contact with the sample stage. The temperature of the implant is determined by four major factors: heating by the pulses, heat loss through conduction with the stage, radiation loss, and convection with the implant plasma. Thus, the time dependence of the temperature is given by:

\[
mc_p \frac{dT}{dt} = Aq - A\varepsilon \sigma (T^4 - T_0^4) - \frac{K_q A_q (T - T_b)}{L_q},
\]

where \( A \) is the target’s surface area, \( q \) is the time-average heat from the pulses, \( \varepsilon \) is the emissivity of the target surface, \( T_0 \) is the temperature of the plasma, \( K_q \) is the thermal conductivity of the quartz rod that holds the sample stage, \( A_q \) is the cross-sectional area of the rod, \( L_q \) is its length, and \( T_b \) is the temperature of the bottom wall of the chamber, with which the rod is in contact. Heat loss by radiation from the rod is assumed to be minimal and may be ignored.

After the implantation procedure, electrical activation of doped materials, usually done by high-temperature annealing, is necessary to increase the substitutional fraction of the dopant species and to repair the crystal lattice damage due to the implantation process that otherwise would degrade the electrical properties of the material. The samples in this work, however, were not annealed so as to maintain the high-aspect surface features and electrical inhomogeneities that are crucial to localize high-density electric fields and hence electron emission at low temperatures. The surface of a crystalline solid does not exhibit the three-dimensional periodicity of the bulk material, and surface atoms relax to eliminate dangling bonds. A common example is the (001) face of silicon, in which
structure relaxation results in the formation of Si-Si dimers. In compound semiconductors, charge transfer between dangling bonds on cations and anions produces local regions of high electric field, which may enhance field emission characteristics.

C. X-ray Photoelectron Spectroscopy

Siegbahn et al.\textsuperscript{30} were the first to use the high-resolution energy spectra of x-ray stimulated electrons to study atomic energy orbitals. This technique, known as X-ray Photoelectron Spectroscopy (XPS) or Electron Spectroscopy for Chemical Analysis (ESCA), uses soft x-ray photons -- often from MgK\textalpha -- to prompt electron emission from a target material.\textsuperscript{31} All electrons with binding energy $E_b < h\nu$ may be excited by incident x-rays, but the probability for certain emissions is higher, producing sharp peaks in the energy spectrum. If the electron kinetic energy is measured by a detector, the binding energy is calculated as:

$$E_p = h\nu - KE + \phi_s,$$

where $\phi_s$ is the work function of the detector, which must be calibrated.\textsuperscript{32} The effective sample depth measured by XPS is based on the electron mean free path (MFP) and is a function of both sample composition and electron kinetic energy and thus in general is not constant throughout the spectrum. As a rule of thumb penetration depth is limited to 5-20 Å in base metals and 15-40 Å in oxides but can reach as deep as 40-100 Å in polymeric materials. In the case of metals, the minimum depth measurement occurs around electron energies of 50 eV and increases linearly to first approximation at higher energies, reaching 20 Å near 2000 eV, as shown in figure 4.\textsuperscript{32} Relative intensities of the photo-peaks may be used to calculate abundance ratios of elements within the material. A complete analysis would involve MFP variations due to crystallographic structure and
electron kinetic energy, but a simpler analysis can be performed by simply dividing relative peak heights by the photoelectric cross-sections of the element. Photoelectric cross sections for the C1s level for different elements when subjected to MgKα can be found in reference 32.

Figure 4 (reprinted from Riggs and Parker32)

A depth distribution of the form

\[ P(z) = \frac{\sqrt{b}}{\sqrt{\pi}} \exp \left[ -b(z - z_{eq})^2 \right] \]

is expected, for distance from surface z, and a Gaussian distribution centered at an equilibrium distance \( z_{eq} \) and of width \( \frac{1}{\sqrt{b}} \). \(^{33}\)

D. Scanning Tunneling Microscopy

The theory of STM tunneling current was first described with perturbation theory by Bardeen in 1961. \(^{34}\) In the limit of small bias voltage and a relatively large sample-tip
distance, the perturbation technique provides a sufficient analysis of the tunneling mechanism for a single-atom STM tip.

\[ j = \int_{0}^{\infty} dE_z D(E_z) N(E_z) \]

in which \( E_z \) is the energy component normal to the metal surfaces. The normal energy distribution is given by:

\[ N(E_z) = \frac{4\pi ne}{\hbar^3} \int [f(E) - f(E + eV)] dE_{\text{parallel}} , \]
where \( f(E) \) is the Fermi-Dirac distribution function, and the transmission probability is approximated by:

\[
D(E) \approx \exp(-2\kappa s),
\]

which holds for small voltages and energies near the Fermi level. In this equation, \( s \) is the tip-sample spacing, and \( \kappa \) is the vacuum decay constant:

\[
\kappa = \sqrt{\frac{2m\phi}{\hbar^2}}
\]

By examination of the exponential dependence of the tunneling probability on tip-sample distance, Sasaki and Tsukada point out that in the case of a small protrusion in the sample surface, almost all tunneling current originates from that area.\(^{36}\)

Assuming the tip is a mathematical point source of current, Tersoff treats STM tunneling as a first-order perturbation situation and arrives at:

\[
\frac{dI}{dV} \propto \rho(R, E_F - eV)
\]

where \( \rho \) gives the local density of states (LDOS) at the surface of the protrusion, assuming that the LDOS of the tip is smooth relative to that of the surface, or else higher-order derivatives will play a measurable role.\(^{37}\) In this case, a more complete theoretical simulation is necessary; one can be found in van Loenen et al., but it is well beyond the scope of this paper.\(^{38}\) Stroscio avoided this problem by plotting the ratio of differential to total conductivity measurements as a function of energy, thereby removing the dependence of the tunneling current on the voltage gap and separation distance and obtaining relatively direct measurement of the LDOS. Then, to ensure that the plot contained no LDOS features of a tungsten probe tip, they scanned both silicon surfaces
and nickel films and finding no common peaks, concluded that tungsten probe tips contribute no significant features to the spectra.\textsuperscript{39}

In any case, the dynamic inductance (dI/dV) can be measured using STM in spectroscopy mode by holding the tip-sample distance constant and varying the bias voltage over a single point on the surface of the sample. I(V) characteristics may be compared to a \textit{modified} form of the FN equation:

\[ I = 1.4 \times 10^{-6} A \frac{1}{\phi} e^{9.86\phi^{\frac{1}{2}}} \beta^2 V^2 e^{-6.53 \times 10^3 \phi^{\frac{1}{2}}/\beta} \]

for current I and emitting area A and where $\beta$ is the field enhancement factor.\textsuperscript{12} For smooth emitters with tip to sample distance of less than about one micron, $\beta$ may be expressed under a hyperbolic model as a function of the tip radius:

\[ \beta_H = \frac{2}{r_t \ln \left( \frac{4R}{r_t} \right)} \]

where $r_t$ is the tip radius and R is the tip-sample distance.\textsuperscript{40}

In current vs. distance plots in STS of metal surfaces, an exponential dependence of tunneling current on the tip-sample separation is well known. Kuk plots tunneling current as a function of gap distance on semi-logarithmic axes, obtaining the work function as:

\[ \phi = 0.952 \left( \frac{d \ln I}{dS} \right)^2 \]

for a Na single-atom tip, where S is the tip-sample separation.\textsuperscript{41} This equation holds only for local barrier heights (LBH’s) when the tip-sample distance is greater than \(~5\ \text{Å}, where the barrier can be treated as a small perturbation between two independent metallic states, corresponding to the tip and the sample surface. For closer distance, the
perturbation term becomes larger, and the slope may vary over this region. Unfortunately, the size and shape of the probe tip and the detailed topology of the sample surface play a large role in measuring the barrier height. Kuk also presents a more reliable measure of work function for gap voltages below the work function, where tunneling current is not described by the Fowler-Nordheim equation but depends exponentially on the work function: \[ I \approx Ve^{-1.05\phi}, \]

where \( \phi \) is in eV.

Citing Tersoff-Hamann theory, Hamers and Padowitz also give an equation for current that is linear in voltage for small gap biases, about 10 mV:

\[ I = 32\pi^3 h^{-1} e^2 \phi^2 R^2 \kappa^{-4} e^{2\phi E_d} D_1(E_F) D_2(r_0, E_F) V, \]

where \( R \) is the tip radius, \( D_1 \) is the tip density of states at the Fermi level, \( D_2 \) is the surface density of states at the Fermi level directly under the tip, and \( \kappa \) is the vacuum decay constant. \( ^{21} \)
III. Experimental Procedure:

Square foils (7 mm x 7 mm) of Group V transition metals – 99.8% vanadium, 99.98% niobium, and 99.95% tantalum – were implanted with carbon ions at ~35 kV for seven hours at a pulse rate of 125 Hz. The vanadium foil was 0.127 mm thick, the niobium was 1.0 mm, and the tantalum was 0.5 mm. The implantation chamber was pumped down by a CIT-Alcatel Type 2033 mechanical pump and a Seiko Seiki STPH-1000 turbo-maglev, backed by an Edwards GVSP 30A scroll pump, all of which were dry pumps. Pump speed was throttled by an iris valve. The implant plasma was held at a pressure of 2.4 × 10⁻³ Torr and was composed of 2% methane gas with a mixture of 4% hydrogen and 94% argon to minimize carburization on the walls of the implantation chamber. Use of methane (CH₄) as carbon source, however, decreased carbon implantation energies by a factor of 25% because the total mass of a methane molecule, 16 amu, derives only 12 amu of that mass from carbon. Therefore during a 35 kV implantation, carbon ions implanted at only 26.25 kV. The plasma was driven by an RF power supply capable of an output of 2.00 kW and operated at 850 W at 13.6 MHz.
through three variable capacitors to allow for fine tuning to match the impedance load of the power supply to that of the plasma. It was inductively coupled to the chamber, providing a higher plasma density than a capacitively coupled arrangement would have, and thus allowing a greater ion flux per pulse as well as a longer pulse length without danger of the ion sheath extending to the wall, minimizing the possibility of arcing and damage to the samples. Figure 6 is a schematic of the PSII system.

![Figure 6](image)

**Figure 6** (reprinted with permission from Theodore, Manos

The delivered ion dose during implantation may be calculated by

\[ \phi = \frac{w \nu n I_i t}{qA} \]

for pulse width \( w \), pulse frequency \( \nu \), average atoms to be implanted per ion \( n \), ion current \( I_i \), total implantation time \( t \), fundamental charge \( q \), and sample area \( A \). While most of these quantities may be directly measured, \( n \) and \( I_i \) must be calculated. For a
methane plasma, the relative concentrations of CH$_3^+$ and CH$_4^+$ are difficult to determine experimentally. Fortunately, in each case $n_{\text{carbon}} = 1$, and no calculation is necessary. Calculating ion current, however, is more involved but was done by our group$^{43}$, giving:

$$I_i = \frac{I_t - I_d}{1 + \gamma},$$

where $I_t$ is the total current as measured by an ammeter at the output of the transformer, $I_d$ is the displacement current, which can be measured by pulsing the sample with no plasma in the chamber, and $\gamma$ is the secondary electron emission coefficient. By substitution, the dose rate may be calculated as:

$$\phi = \frac{\nu \int \gamma}{\gamma(qA(I_t - I_d))}.$$

Previous experimentation with this chamber allowed samples to be placed in a region of highly uniform plasma density.$^{27}$ High negative voltage pulses were created by a pulse-forming network (PFN) in which a thyratron, driven by the generated pulse, determined the pulse rate by grounding out for a certain time, thereby triggering a 10X step-up transformer. The PFN was capable of producing 100 kV, 30 A pulses at 200 Hz. A network of capacitors and inductors removed “ripple” from the pulse shape in order to achieve a more uniform voltage. After implantation, the samples underwent no annealing. For comparison, similar samples of vanadium, niobium, and tantalum metals were cut and examined without being submitted to carbon implantation.

XPS measurements were carried out on the implanted and bare metals to determine whether carbides were formed during implantation. The preparation chamber of the XPS system was pumped down to $10^{-3}$ Torr with a roughing pump, followed by a turbo molecular pump and a diffusion pump, which decreased the pressure to $10^{-8}$ Torr. After movement to the analysis chamber, an ion pump dropped the pressure to $\leq 10^{-9}$
Torr. For analysis, a twin anode MgKα source was used to produce an emission current of 10 mA at 10 kV for a power output of 200 W. The detector was a channeltron, a three-centimeter-long horn-shaped continuous photomultiplier. The channel was made of evacuated, semiconductive glass. When an incident electron strikes this surface, it expels more electrons, which are then accelerated towards the anode, hitting the walls several more times and each time generating more secondary electrons. Figure 7 is a photograph of the XPS system. The bulbous top-most portion is the hemispherical mirror analyzer with the channeltron located in an appendage in the foreground below. The window under it looks into the analysis chamber, and the preparation chamber sticks out on the right side.
The change in contact potential, which is related to the work function, due to implantation was measured with a Kelvin Probe apparatus, capable of measuring the relative work function with respect to a nondestructive, vibrating stainless steel probe tip positioned parallel to the sample. In this configuration, the steel probe tip and sample produce a parallel plate capacitor with a capacitance that oscillates with the probe tip. Kelvin probes operate by adjusting an offset DC voltage between the two surfaces to equalize their Fermi energies and thus produce an equal and opposite charge across the capacitor gap. The zero value of the current flow can then be correlated with the value of the bias voltage, thereby giving the relative work function or contact potential. Unfortunately an irreparable pinhole in the bellows of the Kelvin probe vacuum system prevented measurements at pressures below 6.0 x 10^{-2} Torr. Some 101 measurements of each foil sample were taken to maximize the signal to noise ratio at a frequency of 108 Hz and a bias voltage of 2.000 V. Figure 8 is a photograph of the Kelvin Probe apparatus. The coil on the top of the chamber provides oscillations, while the window looks in at the interface of the oscillating probe tip with the sample.
Kelvin probe contact potential measurements were then compared to current versus voltage curves produced by STM, which measures the variation of tunneling current as a function of voltage. For STM imaging, the samples were fixed to magnetic sample mounts with a conducting silver paste. The mounts were then attached to a stainless steel cylinder which ran along guide rails for motorized travel in the z direction. Tunneling currents were measured with a Pt/Ir 0.25 mm wire pulled by hand to a tip radius of ~ 250 nm. Maximum lateral resolution was 0.015 nm, and set point currents could be adjusted between ±100 nA in increments of 25 pA. Over an area of 7.32 nm x 7.32 nm, some 16,384 measurements were taken to produce of a visual image of the surface features. Then by adding a sample-and-hold circuit to the feedback loop of the microscope, the piezo-voltage was prevented from changing during the “hold” phase, maintaining a constant tunneling gap between the sample and the microscope probe, aside from the effects of thermal drift. When the bias voltage across the gap was changed, the current was measured with an analog to digital converter. This process was completed by averaging 128 I-V spectra for each measurement to maximize the signal to noise ratio. Figure 9 is a photograph the easyscan STM system. Atop the mottled gray base, a glass cylinder houses the sample and the probe tip to minimize convective effects and air movements.

Figure 9
To determine relative tip-sample height, a plot of current vs. Z distance was constructed over the point of the I-V measurements; then several I-V curves were produced with varying setpoint currents at that location, and the I-Z curve was used to calculate the height for the given setpoint. The absolute tip-sample distance was determined by assuming that the current goes to infinity at Z = 0, a valid assumption because the 100 nA current limit prevents measurements with the tip nearer than ~ 0.3 nm to the sample. Knowledge of the tip to sample height is required in to obtain the $\beta$ factor for calculating work functions via FN curves. Tip radius is also necessary for calculating the $\beta$ function and was measured with a high-resolution Axioskop 2 light microscope at 400x with additional digital magnification, as seen in figure 10, and under a scanning electron microscope (SEM) at even higher magnification in figures 11 through 13.

Figure 10
In an attempt to correlate surface morphology with reduced work function measured by Kelvin probe and STM, an atomic force microscope (AFM) was used to measure surface roughness. The AFM utilized a Tap 300 “tapping” cantilever with an n-type (100) Si tip with radius of 10 nm and a resonant frequency of 300 kHz ± 100 kHz. A piezo-strip oscillated the cantilever near its resonance while a 1.0 mW laser diode (670 nm) measured the vertical height of the oscillations to a resolution of better than 1.0 Å. Lateral resolution in the plane of the sample was dependent upon the number of pixels provided by the software, which was able to resolve features of ~10 nm diameter. A feedback loop regulated the force of the cantilever tip on the sample, minimizing surface destruction. Five surfaces of size 5 \( \mu \text{m} \times 5 \mu \text{m} \) were scanned on each bare metal and each implanted metal. Deviation from mean height, or roughness, measurements were
calculated by arithmetic averaging, $R_a$, and the root mean squared method, $R_q$, over the scanned areas. A photograph of the AFM appears in figure 14. The samples were mounted on the silver disk and rotated to a position under the cantilever, which is located directly under the matte silver laser housing. The optics also point to this location.

Figure 14
IV. Data Analysis

A. STM Data

STM imaging was used to evaluate topographically smooth locations for obtaining I-V and I-Z curves. Figure 15 is a map of tunneling current taken at constant height across an area of 500 nm x 500 nm. The procedure for obtaining stable I-V curves in small areas was to scan a large sample size, choose the smoothest area -- in this case perhaps in the upper left-hand corner, and then zoom in to choose the smoothest area again. This technique minimized the role large-scale (greater than a few nanometers) surface features had in effecting work function measurements.
Figure 15

Figure 16 is a plot of the STM tunneling current from bare vanadium measured at 128 equally-spaced tip-sample distances made at 0.100 s intervals. The gap voltage was set at a constant 0.05 V. Error bars are indicative of surface roughness; even though the I-Z curve was collected over a single point, thermal effects produce a lateral drift between the sample and tip, making it necessary to take into account surface features in the area of the measurement. In this case, maximum surface features reached a distance of 0.4 nm above the height of the sample, making the error bars 0.2 nm in either
direction. The stepwise appearance of this curve is a result of the limited resolution of current measurement for this STM, 25 pA.

![Tunneling Current as a Function of Gap Distance on Bare V at 0.05 V](image)

**Figure 16**

I-Z curves were taken in order to calculate tip to sample distances on I-V curves. Once the regressions had been performed to obtain an equation of the exponential fall-off of the current with distance at a set voltage, several I-V curves at different setpoints were produced. By interpolating the magnitude of the tunneling current at 0.05 V on an I-V curve, that current could be plugged into the equation of the I-Z curve to obtain a distance...
for that particular I-V spectrum. This procedure could be completed with the following formula:

\[
Z = \frac{1}{B} \ln \left( \frac{F(V)}{A} \right),
\]

where A and B are constants given by the exponentials for I-V and I-Z curves, as follows:

\[
i = A \exp(Bz)
\]

for current i, voltage V, and separation z, and where F(V) is linear in V for small voltages (less than ~ 10 mV) and becomes exponential at higher gap biases. Such I-Z curves were measured twice for each metal, once on the bare specimen and once on the carbon-implanted metal. Each is similar in shape to that of bare vanadium and is closely mapped by an exponential decline, as expected theoretically. These I-Z curves appear in Appendix I: “Tunneling Currents Measured with respect to Gap Distance.”

Figure 17 is an overlay of I-V curves of bare vanadium taken at increasing tip-sample separations. As is well known, measurements taken a greater distance away from the sample require exponentially higher voltages before they become significant current sources.
For voltages greater than \( \sim 0.4 \) V, curves are neatly approximated by exponentials of the form:

\[
i = a \exp(bV),
\]

for constant \( a \) and \( b \).

As the tip-sample distance closes, however, the curves become increasingly more linear and are better fit by equations of the form:

\[
i = aV \exp(bV),
\]

and thus contain a linear and exponential portion. Although the researcher did not reach currents low enough to see completely linear behavior on bare vanadium, he was able to do so on bare niobium and tantalum samples. Figure 18 contains six plots of tunneling
current against voltage on bare niobium, taken at different tip-sample separations with the intent of making the linearization transition more visible. In the data sets that are best fit by pure exponentials, the points are colored red, those best fit by straight lines are colored blue, and those best fit by a multiplicative hybrid are green. Curve fit R values are above 0.993 for all regressions, despite the stepwise appearance of the curves due to limited current resolution. Numerically, one may see the linearization trend as an increase in the scaling parameter a (or m1) and a decrease in the exponential parameter b (or m2) as defined in the equation:

\[ i = aV \exp(bV) \]

Note that the exponential approaches unity as (bV) becomes smaller, and voltages in this region are usually one to two orders of magnitude below unity. Overlays of I-V curves of unimplanted and implanted samples of each metal are available in Appendix II: “Tunneling Currents Measured with respect to Bias Voltage.” The researcher has been unable to explain a distance-scaling error of ~ 4 in the plot of bare Ta. It is likely that an error was made with a setting during data collection.
Figure 18
In the case of voltages greater than the work function, it is possible to calculate the barrier potential using the Fowler-Nordheim method:

\[ I = 1.4 \times 10^{-6} A \frac{1}{\phi} e^{9.86\phi \frac{1}{2}} \beta^2 V^2 e^{-6.53 \times 10^{-7} \phi \frac{3}{2} \beta} \]

for emitting area A and field enhancement factor \( \beta \).\textsuperscript{12} Traditionally, this procedure is done by plotting \( \ln \left( \frac{I}{V^2} \right) \) against \( \frac{1}{V} \), from which a straight line is obtained with slope:

\[ m = 6.53 \times 10^7 \frac{\phi \frac{3}{2}}{\beta} \]

and intercept:

\[ b = \ln \left( 1.40 \times 10^{-6} A \beta^2 \frac{e^{9.86\phi \frac{1}{2}}}{\phi} \right) \]

In the case of this work, however, the 100 nA limit on the STM required voltages to remain well below the work function, and attempts to produce an FN plot resulted in logarithmic curves, such as Figure 19. On the high voltage end of this plot, it appears as though the behavior may be approaching linearity, but current limit prevents any further progression. The discontinuities at the high-voltage end (small values of \( 1/V \)) are a product of current overloads causing the STM software to abort measurements and to average values of zero at that location for some of the 128 measurements that were taken.
For lower voltages, however, Kuk presents another method of calculating the approximate work function, as follows:

\[ I \approx Ve^{-0.05\phi}, \]

where the constant has been adjusted so that the voltage is measured in volts, the current in nanoamps, and the work function in electron volts.\(^{40}\) By this method, Table 1 is calculated. Accepted values for bare materials are 4.3 eV for vanadium and niobium and 4.25 eV for tantalum.\(^{46}\) Error values listed in this table arise only from discrepancies with the curve fitting. Additional errors – most notably thermal drift – have not been accounted for. Deviations from the accepted value may be an effect of the thermal drift, or may be a result of any contradictions with Kuk assumptions. It is worthy to note that
vanadium foils used in this experiment were by far the most reflective and the least rough, characteristics necessary for STM. Niobium foils in particular were visibly more “matte” than vanadium and contained large-scale (~ 1-2 µm) surface features. Kuk’s paper provides no detailed justification for this linear approximation, nor does he detail under what circumstances it holds. It appears, however, to be appropriate for the cases we analyzed.

**Work Functions, Calculated by Kuk Linear Approximation**

<table>
<thead>
<tr>
<th>Material</th>
<th>Work Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bare V</td>
<td>4.33 ± 0.03 eV</td>
</tr>
<tr>
<td>C-imp V</td>
<td>4.28 ± 0.03 eV</td>
</tr>
<tr>
<td>Bare Nb</td>
<td>5.22 ± 0.01 eV</td>
</tr>
<tr>
<td>C-imp Nb</td>
<td>4.71 ± 0.01 eV</td>
</tr>
<tr>
<td>Bare Ta</td>
<td>4.45 ± 0.01 eV</td>
</tr>
<tr>
<td>C-imp Ta</td>
<td>4.48 ± 0.01 eV</td>
</tr>
</tbody>
</table>

*Table 1*

Figure 20 is a comparison of graphs of DOS measurements of bare and implanted vanadium, niobium, and tantalum measured at a tip-sample distance of about 2 nm. The noise on the low voltage end is due again to finite current resolution but in this case is magnified as a derivative over a discontinuous region. For most of these plots, any changes due to implantation are obscured by the noise. Qualitatively, it may be possible to discern a slightly lower increase in the density of states with voltage after implantation, but with such a low signal-noise ratio, it is impossible to say without better measurements.
B. Kelvin Probe Data

Table 2 contains average contact potential measurements made on implanted and unimplanted samples by KP. The stainless steel measurement tip should have a work function of 4.1 eV, but a calibration run on a stainless steel sample yielded an average CPD of -0.4 eV, indicating the need for recalibration.

<table>
<thead>
<tr>
<th></th>
<th>Bare</th>
<th>Bare</th>
<th>C-Implanted</th>
<th>C-Implanted</th>
<th>∆CPD</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Avg. CPD</td>
<td>Std. Dev.</td>
<td>Avg. CPD</td>
<td>Std. Dev.</td>
<td></td>
</tr>
<tr>
<td>Vanadium</td>
<td>-0.771 eV</td>
<td>0.020 eV</td>
<td>-0.808 eV</td>
<td>0.159 eV</td>
<td>-0.037 eV</td>
</tr>
<tr>
<td>Niobium</td>
<td>-0.007 eV</td>
<td>0.015 eV</td>
<td>-0.555 eV</td>
<td>0.018 eV</td>
<td>-0.548 eV</td>
</tr>
<tr>
<td>Tantalum</td>
<td>-0.172 eV</td>
<td>0.023 eV</td>
<td>-0.819 eV</td>
<td>0.026 eV</td>
<td>-0.645 eV</td>
</tr>
</tbody>
</table>

Table 2

The column ∆CPD indicates the change in contact potential difference before and after implantation, which is equal to the work function enhancement, assuming the contact potential of the stainless steel tip remains constant. The ∆CPD for vanadium, measured to be about -0.04 eV correlates well with the work function enhancement of 0.05 eV measured by I-V fitting in Table 1. Similarly the ∆CPD for Niobium, measured here to be -0.56 eV is not far from the work function enhancement of 0.51 eV as measured by STM I-V. Tantalum samples, however, are more puzzling. STS curves indicate that the work function actually slightly increases from 4.45 ± 0.01 eV to 4.48 ± 0.01 eV after implantation. KP measurements, however, show a decrease of -0.65 ± 0.03 eV in contact potential in the implanted samples, more in line with results obtained by other groups listed in the introductory section of this paper. This apparent contradiction may be explained by an error in estimating the slope of the tantalum I-V lines at small
tip-sample separations due to too few measurements. At small tip-sample distances, electron emissions from both bare and implanted tantalum quickly reach values above the STM current limit, and the measurements cannot be used.

C. XPS Data

Figure 21

Figure 21 is a comparison of scans of the carbon C1s peak in bare V and C-implanted V. In the unimplanted plot, the high-energy shoulder of the peak is characteristic of carbon peaks in metals. In the implanted plot, a shoulder at a lower binding energy (BE) has appeared, indicating a C-metal bond and the formation of a carbide. Fujihana et al. specified binding energies for C-O bonds to be 281.6 eV, C-C bonds as 284.2 eV, and C-metal bonds to range from 281.0 eV to 283.3 eV for transition metal carbides. A constant shift of ~7 eV from book values indicates that the XPS needs to be recalibrated. Because peak height is proportional to the implanted concentration, only a proportionally small amount of carbide was formed.
Figure 22

Figure 22 is a comparison of scans of the V2p metal peak in bare and implanted vanadium. Two metal peaks are visible in the bare spectrum: a sharp peak at 523 eV, corresponding to the 2p\(^{3/2}\) state; a lower peak at 531 eV, the 2p\(^{1/2}\) state; and the O1s peak is just visible at 536 eV.\(^{48}\) In the implanted spectrum, a third peak has appeared at higher BE than the 2p\(^{3/2}\) state, as has a high BE shoulder developed on the 2p\(^{1/2}\) state. These peaks provide further evidence of either carbon or oxygen, and likely both, bonding to the metal as mixed oxides and carbides.\(^{49}\)

Figure 23
Similarly, figure 23 is a comparison of the C1s peak on bare and implanted Nb. The bare Nb scan shows only the C1s peak, complete with its high BE shoulder. In the implanted scan, a slight low BE hump has appeared at the foot of the C1s peak, corresponding to C-metal bonding.\textsuperscript{47} The small height of this peak indicates that only a relatively small amount of carbon is engaged in C-metal bonds or that only the top surface is carbide. In the unimplanted Nb3d metal scan, figure 24, the two large peaks at 214 eV and 217 eV correspond to the 3d\textsuperscript{5/2} and 3d\textsuperscript{3/2} states, in order of increasing BE.\textsuperscript{48} The small peaks are also pictured in bare metal scans in Moulder et al.\textsuperscript{48} but are unlabeled. The implanted scan shows a shift of the 3d\textsuperscript{5/2} state to a higher BE of 215 eV, indicating that that state has lost electrons to a bond. There is also a new peak at 213 eV, which may correspond to either C-metal bonding or O-metal bonding.

![Bare Niobium and C-implanted Niobium](image)

**Figure 24**

Figure 25 is likewise a comparison of scans of the C1s peak on bare and C-implanted Ta. Just as seen in the V and Nb comparisons, only a high BE shoulder appears on the bare peak, but a low BE peak is located at the foot of the C1s peak in the implanted case, indicating some carbide formation.
Figure 26 is again a comparison of the metal Ta4f region for bare and implanted samples. The bare sample shows four peaks, two of which are the 4f⁷/₂ state at 35 eV and the 4f⁵/₂ state at 37 eV as identified by Moulder et al.\textsuperscript{48} The other two are identified by Riggs and Parker\textsuperscript{31} as tantalum(V) oxide. In the implanted case, the Ta4f peaks shift up to 37 eV and 39 eV, the tighter hold indicating a loss of electrons from that orbital, and a third peak at 34 eV, likely a C-metal peak. The metal-oxide peaks have contracted, perhaps due to displacement by carbon during implantation.
D. AFM Data

<table>
<thead>
<tr>
<th>Material</th>
<th>Bare</th>
<th>Std Deviation</th>
<th>C-implanted</th>
<th>Std Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>V</td>
<td>12.2 nm</td>
<td>2.0 nm</td>
<td>24.1 nm</td>
<td>2.5 nm</td>
</tr>
<tr>
<td>V</td>
<td>16.0 nm</td>
<td>1.8 nm</td>
<td>29.8 nm</td>
<td>2.6 nm</td>
</tr>
<tr>
<td>Nb</td>
<td>114.4 nm</td>
<td>26.6 nm</td>
<td>144.3 nm</td>
<td>25.3 nm</td>
</tr>
<tr>
<td>Nb</td>
<td>100.9 nm</td>
<td>22.3 nm</td>
<td>125.6 nm</td>
<td>24.8 nm</td>
</tr>
<tr>
<td>Ta</td>
<td>129.0 nm</td>
<td>53.8 nm</td>
<td>159.4 nm</td>
<td>68.8 nm</td>
</tr>
<tr>
<td>Ta</td>
<td>117.06 nm</td>
<td>49.6 nm</td>
<td>145.13 nm</td>
<td>68.8 nm</td>
</tr>
</tbody>
</table>

Table 3

Table 3 is a table of average surface roughness measurements on bare and C-implanted group 5B metals. Measurements were taken over five areas of 5 µm x 5 µm for each sample and computed by arithmetic averaging for $R_a$ and the root mean squared method for $R_q$. According to these data, implantation approximately doubled the roughness of the vanadium foil but had little effect on the rougher niobium and tantalum foil. Perhaps the effect of carbon deposition onto rougher surfaces is a small factor when compared with the surface features that existed before implantation. In these measurements, the standard deviation values also provide an indirect measurement of surface roughness. Naturally, a greater degree of variation is expected from surfaces with more prominent features. Figures 26 through 28 are AFM images of surfaces of vanadium, niobium, and tantalum, respectively.

It is difficult to directly relate these measurements of surface roughness to effective work function calculations that appear in table 1. While virtually no enhancement in work functions of vanadium and tantalum due to implantation was calculated, the roughness of vanadium increased on the order of twofold. Moreover, even though the work function of niobium was increased by ~ 0.5 eV, the roughness remained
constant within one standard deviation. Effective work functions for niobium, however are \(~ 0.9\) eV higher than accepted values, which seems inconsistent with prominent surface features on the sample, just as the \(~ 0.2\) eV difference of observed over accepted work function values on bare Tantalum is not explainable by surface roughness.

Figure 27: Vanadium
Figure 28: Niobium

Figure 29: Tantalum
V. Conclusion

Because of current limits of the STM, all I-V curves produced within the linearization region were not analyzable using the Fowler-Nordheim equation. At larger tip-sample distances (greater than ~1.5 nm), more voltage may be applied before reaching the current ceiling and traditional exponential behavior is observed. Below this range, linearization can be traced by modeling data as the product of a voltage with an exponential, and for tip-surface separations of less than ~0.7 nm the exponential approaches unity. Based on linear approximation models of STM I-V curves at low voltages, no significant work function enhancements were seen on C-implanted V and Ta outside of the uncertainty in measurement. On niobium samples, a work function enhancement of 0.51 ± 0.02 eV was observed after implantation. The discrepancy between observed and accepted values of the work function of bare niobium may be explained by the low reflectivity of the surface; STS requires a conductive and very smooth surface for stable electron currents. Little information can be gleaned from
density of states measurements plotted as \( \left( \frac{I}{V} \right)^{-1} \frac{dI}{dV} \) against V, which are dominated by noise due to integration over discontinuous current resolution at lower voltages. Kelvin probe measurements taken on a the sample foils indicate similar field emission enhancements for vanadium and niobium samples – a small enhancement of 0.04 eV for vanadium and a significant decrease of 0.56 eV for niobium. KP measurements of the tantalum foil, however, show a \( \Delta \text{CPD} \) of 0.65 eV, even though STS measurements showed no significant change. A possible explanation is that the relatively small amount of data points below the STM current limit prevented an accurate calculation of the I-V slope. XPS BE scans of the bare and implanted metals revealed a low energy shoulder or subpeak 2 to 3 eV below the C1s peak in each of the implanted specimens, corresponding to carbide formation, even if it is well below stoichiometric levels. Scans of the metal peaks also exhibited signs that can be interpreted as carburization or oxidation. AFM surface roughness measurements did not correlate well with work function enhancements. Only V samples displayed substantial signs of increased roughness after implantation, although the more irregular Ta and Nb foils may have experienced a similar increase that was masked by their rougher initial surfaces. Further research with a more powerful STM is necessary to accurately describe FN behavior of Group 5B in the 1 to 3 V region and thus calculate work function enhancement and particularly to view DOS plots, which typically become interesting in the region of 1 to 4 V.
Appendix I: Tunneling Currents Measured with respect to Gap Distance

Tunneling Current as a Function of Gap Distance on Bare V at 0.05 V

Tunneling Current as a Function of Gap Distance on C-implanted V at 0.05 V
Appendix II: Tunneling Currents Measured with respect to Bias Voltage

Tunneling Current as a Function of Bias Voltage on Bare V at Several Tip-Sample Distances

Tunneling Current as a Function of Bias Voltage on C-implanted V at Several Tip-Sample Distances
Tunneling Current as a Function of Bias Voltage on Bare Ta at Several Tip-Sample Distances

Tunneling Current as a Function of Bias Voltage on C-implanted Ta at Several Tip-Sample Distances
VI. Acknowledgements

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