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Simultaneous scanning tunneling microscopy and synchrotron X-ray measurements in a gas environment

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Abstract

A combined X-ray and scanning tunneling microscopy (STM) instrument is presented that enables the local detection of X-ray absorption on surfaces in a gas environment. To suppress the collection of ion currents generated in the gas phase, coaxially shielded STM tips were used. The conductive outer shield of the coaxial tips can be biased to deflect ions away from the tip core. In tunneling contact, the X-ray-induced current is separated from the regular, 'topographic' tunneling current using a novel high-speed separation scheme. We demonstrate the capabilities of the instrument by measuring the local X-ray-induced current on Au(111) in 800 mbar Ar.

Keywords: Scanning tunneling microscopy, Synchrotron radiation, High-pressure surface science, operando catalysis research

Highlights:

- First-time measurement of the local X-ray-induced current on Au(111) at a gas pressure of 800 mbar.
- A coaxially shielded STM tip and mounting scheme that effectively removes the ion current background.
- An electronics scheme to separate X-ray-induced current from tunneling current that enables fast height feedback.
- A quick method for aligning tip and X-ray beam with micrometer precision.

1. Introduction

The synchrotron-based techniques developed over the last decades provide a powerful toolbox for the study of interface phenomena. The weak interaction of X-rays with matter makes these methods particularly suitable for *operando* studies, which are indispensable in research areas such as catalysis [1], environmental science [2], and film growth [3]. However, most X-ray techniques make use of a large beam spot compared to the features studied, thereby averaging out possibly important structural and chemical variations. For techniques that use a highly focused beam on the other hand, finding the feature of interest can be a tedious process as searching based on reciprocal-space images or absorption features is non-trivial.

Scanning probe microscopy can supplement X-ray measurements by providing local structural information in real space. Inspired by this potential, several atomic force microscopes (AFM) [4–9] and scanning tunneling microscopes (STM) [10–14] have been installed on beamline end stations. In the AFM studies, it was shown that the AFM tip can be used to align the X-ray beam with features of interest [6]. Once aligned, the tip can be used for nano-manipulation or the application of local stress [15,16].

Using the combination of STM and X-rays, it was shown that the current collected by the STM tip can be used to measure local X-ray absorption, with a spatial resolution as small as 2 nm [11,17–19]. Thus, local chemical information can be obtained and coupled to the local structure. This combination is highly desirable for the understanding and design of materials and chemical processes.

So far, all synchrotron X-ray assisted STM (SXSTM) studies have been performed in vacuum. Under these conditions, the signal collected by the tip consists of three principal components: the photo-electron current, the regular, ‘topographic’ tunneling current, and an X-ray-induced increase of the tunneling current. While both the photo-electron current and the X-ray-induced increase of the tunneling current are related to X-ray absorption, only the latter provides local information [11,17–19]. To improve the ratio between this local signal and the photo-electron current, most SXSTM experiments employ coated STM tips,

leaving only the tip apex uncoated [17,20–23]. This effectively reduces the number of photo-electrons that the tip collects. Besides this background reduction, the X-ray-induced part of the tunneling current needs to be separated from the topographic tunneling current. Several experimental schemes employing a beam chopper and lock-in amplification have been developed for this purpose [11,24].

To use SXSTM in *operando* studies at elevated pressures, two challenges need to be addressed: i) Inhibitively large ion currents are generated in the gas phase. We show that the insulator-coated tips used in most SXSTM studies are inadequate to suppress these. ii) Most interesting systems for *operando* studies require elevated temperatures and have some degree of roughness in their morphology. To cope with thermal drift and to follow the surface morphology accurately, fast response in the height feedback is necessary. However, the current separation schemes for the topographic and the X-ray-induced tunneling current limit the height feedback in the STM to response frequencies below roughly 1 kHz.

Here, we describe an instrument that can perform SXSTM measurements in a gas environment using hard X-rays, overcoming the barriers for performing *operando* studies. Our methodology includes the following: i) A quick method for aligning tip and beam with micrometer precision. ii) An electronics scheme to separate X-ray-induced current and topographic tunneling current that enables fast height feedback at frequencies up to the chopper frequency. iii) A coaxially shielded STM tip following the approach of ref [17] and a mounting configuration that effectively suppresses the ion current background. As a proof of concept, we have collected the local, X-ray-induced tunneling current on a Au(111) sample at a gas pressure of 800 mbar.

2. Instrument design

2.1 High-pressure cell

The starting point for our experiments was the high-pressure scanning probe and surface X-ray scattering instrument developed earlier in collaboration between our group and the ID03 beamline staff at the ESRF [14], which we modified for the SXSTM measurements. In short, this system consists of an ultrahigh vacuum (UHV) chamber with a high-pressure cell on top (see Figure 1) [25]. The UHV chamber allows for surface preparation by ion bombardment, annealing, controlled gas exposure, and metal evaporation onto the sample. The high-pressure cell can be operated as a flow reactor, providing a controlled gas atmosphere at pressures up to 1.1 bar. The cell is exchangeable, allowing for a versatile combination of experiments.

In our recent work, we fitted a high-pressure scanning probe module on the system [14,26]. Similar to the design of the ReactorSTM and ReactorAFM [27,28], it uses a single piezo tube both for the coarse approach of the tip to the sample and for generating the scanning motion. The coarse approach is achieved by a

stick-slip motion of the tip holder (slider) on two tracks. The piezo tube is separated from the gas environment by a polyimide cap and a viton seal, as depicted in Figure 1b. In our previous design, the reactor wall was a dome-shaped aluminum piece of 1 mm thickness, providing homogeneous and satisfactory X-ray transmission over a large range of angles at photon energies above 18 keV. While this made the system excellent for surface X-ray diffraction measurements, the X-ray transmission was insufficient in the lower energy range relevant to X-ray absorption spectroscopy.

To allow for measurements at energies down to 8 keV, we designed a new cell that uses four thin, exchangeable windows (see Figure 1) sealed on both sides by viton O-rings. In our first test measurements, we used a stack of three 75 μm thick Kapton sheets for each window. In the energy range from 10 keV to 12 keV, the Kapton stack provides an excellent X-ray transparency of 91% to 95% [29]. However, the permeability of Kapton towards gases [30] limits the base pressure of the UHV chamber to the 10^{-6} mbar range. Aluminum foil windows provide a lower transmission (59% for 100 μm thick foil at 11 keV [29]), yet do not suffer from any gas permeability. Beryllium windows were not preferred here for safety reasons, as the reactor needs to be handled frequently for tip exchanges.

The reactor walls and the part of the microscope head that are exposed to gases are gold coated. This prevents undesirable side reactions from occurring on these surfaces. Furthermore, charge collected on the walls from photo-electrons and ions is immediately neutralized. Charging of the native oxide of materials such as aluminum and stainless steel may cause memory effects in the background signal of the SXSTM measurements. This could adversely affect the reproducibility of the data.

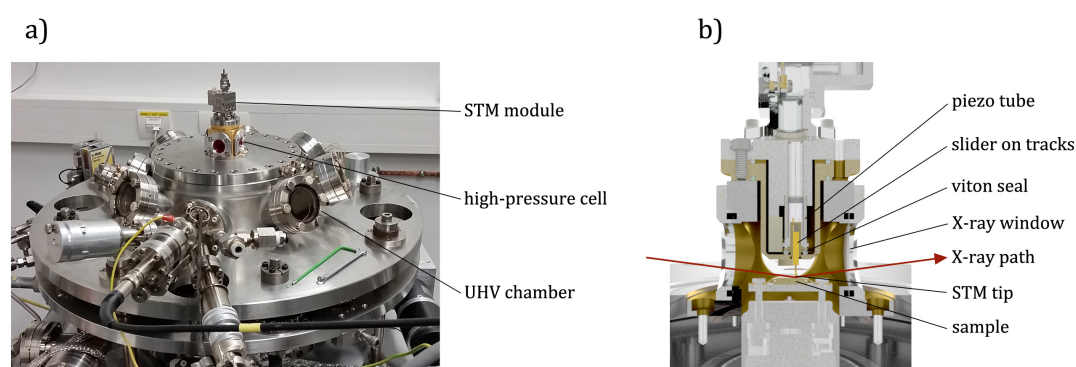


Figure 1: Overview of the set-up. a) Assembly of ID03's ReactorSXR chamber [25], our new high-pressure cell and the recently developed STM module [14]. b) Cross-section of the STM module and high-pressure cell.

2.2 Coaxial tips and mounting scheme

In a gas environment, the X-ray-induced current collected by the STM tip behaves differently as compared to the vacuum situation. Photo-electrons emerging from the sample cause a cascade of ionization processes in the gas.

This effect is particularly pronounced for the high-energy Auger electrons produced during hard X-ray absorption [31,32]. Furthermore, the plasma created in the gas environment helps the charge transport to and from insulating surfaces [33–35]. Hence, when a beam chopper is employed, this may generate oscillating electric fields that can couple into the tunneling current. These effects cause a complex behavior of the background signal in SXSTM measurements, making reproducibility of experiments difficult. It is therefore essential to minimize the background current.

To achieve this reduction in background, we fabricated both insulator-coated and coaxially shielded STM tips. The latter consist of a conductive core, an insulating layer, and a conductive outer layer (shield), as shown in Figure 2. While the core is used for measuring the tunneling current, the shield layer neutralizes ions. Tips are mounted such that the shield potential can be controlled independently with respect to the tip core and the sample, thus creating an electric field that deflects ions away from the tip core.

The insulator-coated tips were fabricated by a combination of SiO₂ sputter deposition and focused ion beam milling similar to that reported in literature [20,22,23]. The fabrication of coaxially shielded tips is less well established [17], hence we explored several preparation methods. As core material, we have used an electrochemically etched Pt_{0.8}Ir_{0.2} wire (\varnothing 0.25mm, Goodfellow). Three different types of insulation layer were tested: i) 100 nm SiO₂ prepared by atomic layer deposition (ALD) through 1100 cycles of bis(diethylamido)silane (Sigma-Aldrich SIBDEA) exposure and an O₂ plasma as co-reactant at 200 °C. ii) 1000 nm SiO₂ prepared by plasma-enhanced chemical vapor deposition (PE-CVD) using a plasma of silane and O₂ at 300 °C. iii) Polymer coating prepared by pushing the tip through a droplet of polyethylene-vinylacetate-based glue gun wax at 80 °C, a method commonly used for electrochemical STM [36].

In all three cases, >2 M Ω core-shield insulation was accomplished with a success rate of over 50%. For the ALD tips, the coating was very homogeneous but too fragile to survive further assembly. For the PE-CVD tips, the insulation thickness was not uniform over the tip due to inhomogeneities in the plasma that are common around conducting three-dimensional objects inserted into the plasma. As a result, the coating at the apex was 7.5 μ m thick while the average was only 1 μ m. For the polymer-coated tips, the coating thickness decreased from 1 mm halfway the tip to a few hundred nanometer at the apex (see Figure 2).

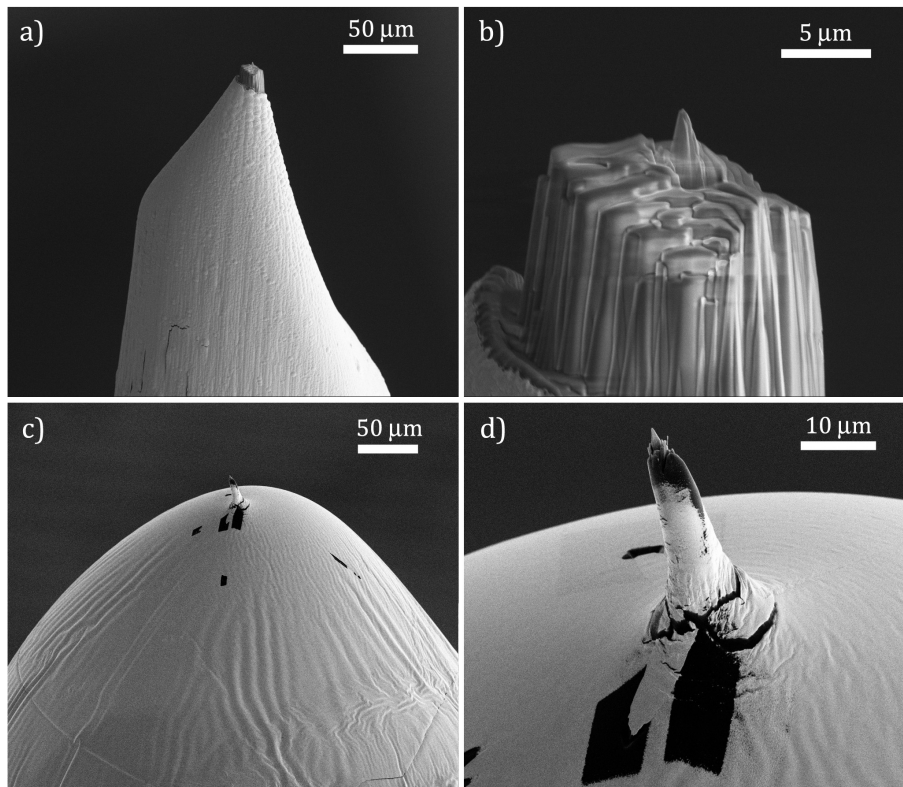


Figure 2: Scanning electron microscopy (SEM) images of coaxially shielded STM tips. a,b) $\text{Pt}_{0.8}\text{Ir}_{0.2}$ with SiO_2/Au coating. c,d) $\text{Pt}_{0.8}\text{Ir}_{0.2}$ with polymer/Au coating.

The shield coating was applied by radiofrequency sputter deposition of a Ti seed layer followed by 500 nm Au for the ALD and PE-CVD tips and 100 nm Au for the polymer tips. Subsequently, the apex was shaped using focused ion beam milling. In this process, the original tip apex was first cut off, followed by a donut-shaped ion milling pattern around the new tip apex to remove the last $\sim 1 \mu\text{m}$ of the isolation and shield layers, so that the core protrudes (see Figure 2). Using a finer donut shape, the apex was sharpened.

The mounting configuration of the coaxial tips is depicted in Figure 3. Essential in the design is that the core and shield of the tip have separate electrical connections. Furthermore, the pick up of ion current by the tip holder needs to be minimized. To achieve this, a gold-coated Kapton foil was attached to the tip using non-conductive glue. Subsequently, the Au coating of the Kapton foil was connected to the shield of the coaxial tip using Agar silver paint. Similarly, a thin Au-coated tungsten wire was contacted to the Au coating of the Kapton through a puncture in the foil. This wire and the tip core were mounted on the tip holder (slider), which consists of two electrically isolated pieces of Au-coated machine steel. Each piece makes contact to one of the tracks on the piezo motor. Shielded wires provide connections from the two tracks to outside of the reactor.

After tip fabrication and mounting, a typical resistance of $2 \text{ G}\Omega$ between tip core and shield could be achieved, allowing for a shield bias of several volt with respect to the core. Thus a strong electrical field between the tip shield and the

sample can be applied, deflecting both positive and negative charges before they can reach the tip core.

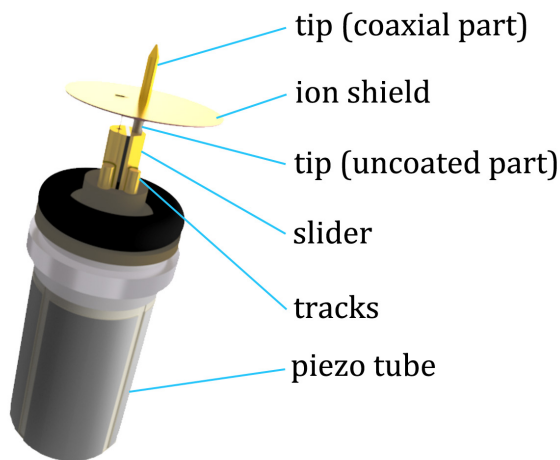


Figure 3: Mounting configuration of the coaxial tip.

2.3 Tip-beam alignment

All experiments were conducted at the ID03 beam line at the European Synchrotron Radiation Facility (ESRF). The beam was focused to $25\ \mu\text{m} \times 25\ \mu\text{m}$ using a toroidal mirror. In the energy window employed, 10 keV to 13 keV, a flux of 10^{13} photons/s was delivered to the sample in the reactor cell with Kapton windows. For the Al dome reactor developed in our previous work [14], the flux on the sample was around 10^{11} photons/s.

In order to align the STM tip with the X-ray beam, the chamber was placed on a hexapod, which in turn stands on a six-circle diffractometer. This allows translation in x (horizontal, perpendicular to the beam), y (parallel to the beam), and z (vertical), as well as tilting and rotation. We used the shadow of the STM tip on the Maxipix X-ray detector [37] as a probe for the alignment. First, the tip was brought to within a few micrometer from the sample surface. With the beam parallel to the sample surface, the chamber height (z) was set such that half of the beam was blocked by the sample. When the chamber is then translated in x , it passes through the configuration in which the beam hits the tip, casting a shadow on the detector. Figure 4a depicts the intensity of the beam spot on the detector recorded during such a chamber translation scan. The minimum of this curve is located at the position where the tip maximally blocks the beam, as is also evidenced by the tip shadow that can be seen in the X-ray detector images (see Figure 4c) and the reflections from the sides of the tip. In order to align in both lateral directions, the chamber was rotated by 90° and the same scan was performed.

To confirm the accuracy of this method, a similar scan was performed with the X-rays incident on the sample at 0.5° . In this case, the beam reflected from the

sample is recorded, which can also be blocked by the tip. Simultaneously, the photo-electron current on the tip core was measured. As shown in Figure 4b, the mismatch between the current maximum and the maximal beam blocking by the tip is only a few micrometer. The origin of this slight mismatch is likely the asymmetry of the tip on the scale of the beam spot, which is also apparent in Figure 4c.

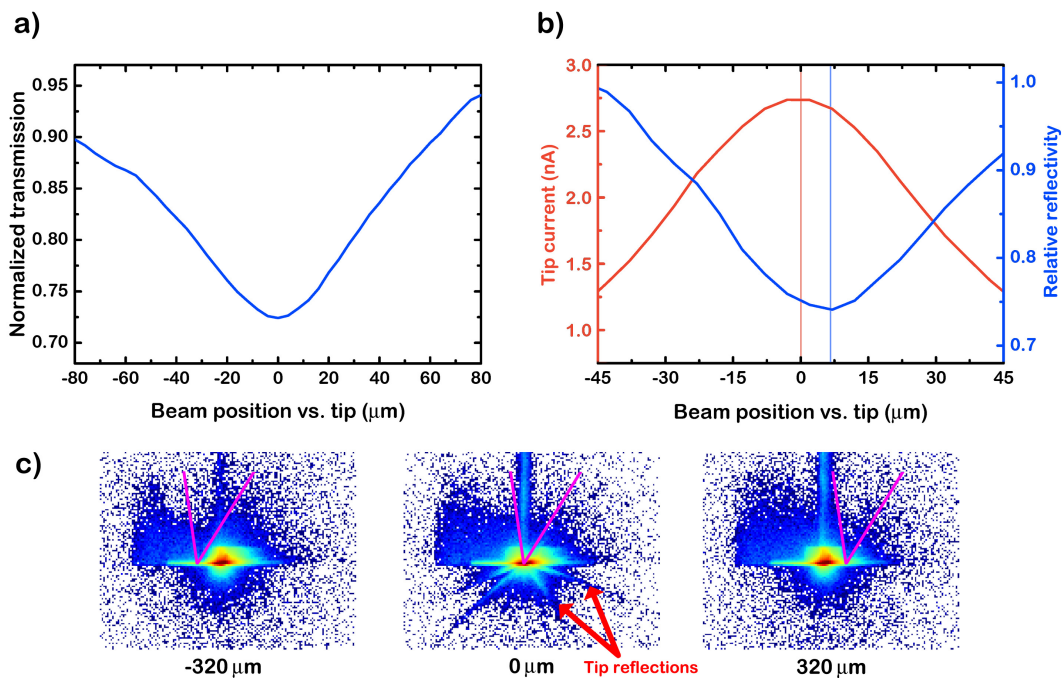


Figure 4: Alignment of STM tip and X-ray beam (tip-sample distance: $\sim 3 \mu\text{m}$). a) Transmission when moving the beam across the tip apex (beam parallel to sample). b) Tip current and X-ray reflection intensity when moving the beam across the sample underneath the tip (0.5° incidence). c) Beam spot on the detector when moving the beam across the tip. The purple lines indicate an estimate of the tip apex profile.

2.4 Electronics scheme for separating the topographic from the X-ray induced tunneling current

To obtain the local X-ray absorption signal, the X-ray beam is chopped [11,24]. Without adjustments to the height-feedback mechanism, which uses the tip current as an input, this would result in an oscillation of the tip height. In principle, the height signal can be lock-in amplified to find the local X-ray absorption signal. However, the height signal has a logarithmic dependence on the tip current, making it relatively insensitive to the X-ray induced signal.

An alternative was proposed by Wang *et al.* [24], who performed height feedback on the tip current only after applying a low-pass filter. In this case the X-ray-induced signal is obtained from the tip current before the low-pass filter using lock-in amplification. In order to completely remove X-ray-induced effects from the height feedback, the X-ray-induced signal obtained from the lock-in amplifier is scaled and subtracted from the low-pass-filtered tunneling current:

$$I_{\text{topo}} = I_{\text{total}}(\text{low pass filtered}) - I_{\text{X-ray/lock-in}}$$

While this scheme allows for stable tunneling, the height feedback is limited to frequencies of maximally $0.1 \times$ chopper frequency in order to reach a stable signal in the lock-in amplifier. The bandwidth of the high-gain preamplifiers used in STM (ranging up to 50 kHz bandwidth for normal operation) restrict the chopper frequency to roughly 10 kHz in order for the chopped X-ray induced signal to be transmitted without distortions. This leads to a maximal feedback frequency of 1 kHz, which is adequate for flat surfaces and slow data acquisition in environments with low vibrational noise levels. However, more flexibility is desired for the nanoparticle systems typically used in catalysis. Furthermore, the challenging boundary conditions resulting from the integration of an STM in a reactor end-station impose limitations to the vibrational noise reduction that can be achieved [14].

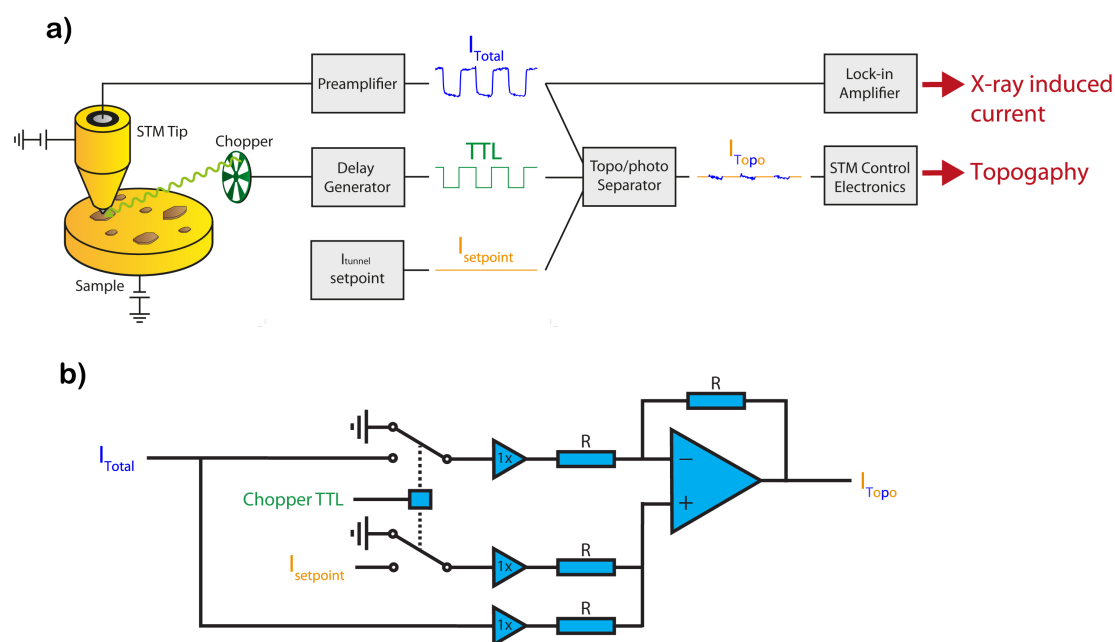


Figure 5: a) High-speed separation scheme for the topographic tunneling current and the X-ray-induced current. b) Electronic circuit for the topo/photo separator, generating the input for the STM control system without X-ray-induced contributions.

In order to perform faster height feedback, we have developed a new scheme to separate the X-ray-induced current from the topographic tunneling current. In essence, our circuit switches the height feedback off during the “beam on” phase of the X-ray chopper. This is accomplished by replacing the measured signal by the tunneling current setpoint during this phase. To do this, the transistor-transistor logic (TTL) signal from the beam chopper is fed into a delay generator, with which the phase and the duty cycle of the TTL signal can be adjusted independent of the phase and duty cycle of the chopper. Thus, the TTL phase can be matched to that observed in the tip signal. Adjusting the TTL duty cycle to prolong the “feedback off” period can be useful if the tip signal is not a perfect square wave, as is the case for our measurements in a gas environment.

The modified TTL signal is used in a purpose-built circuit (Figure 5b), which performs the following operation:

$$I_{\text{topo}} = I_{\text{total}} + \text{TTL} \times (I_{\text{setpoint}} - I_{\text{total}})$$

The output signal can be used for height feedback at frequencies up to the chopper frequency, because the topographic tunneling current is sampled every chopper cycle during the “beam on” phase. Hence, our scheme uses up to 20% of the preamplifier bandwidth, compared to 2% for the current state-of-the-art from Wang *et al.* [24]. In principle, one could construct a similar high-speed circuit for measuring the X-ray-induced signal. However, lock-in amplification of the tip signal provides a better signal-to-noise ratio.

We have first tested the acquisition scheme described above using the X-ray-induced current measured with the tip out of tunneling contact. Thus, we only measured the oscillations in the ion current background induced by chopping of the beam. From the data in Figure 6, it is clear that the X-ray-induced current is effectively removed from the signal. At high chopper frequencies, the signal deviates from the ideal square wave. Nevertheless, Figure 6b shows that the X-ray induced current can still be completely removed by changing the duty cycle of the TTL signal, thus evidencing the robustness of our scheme.

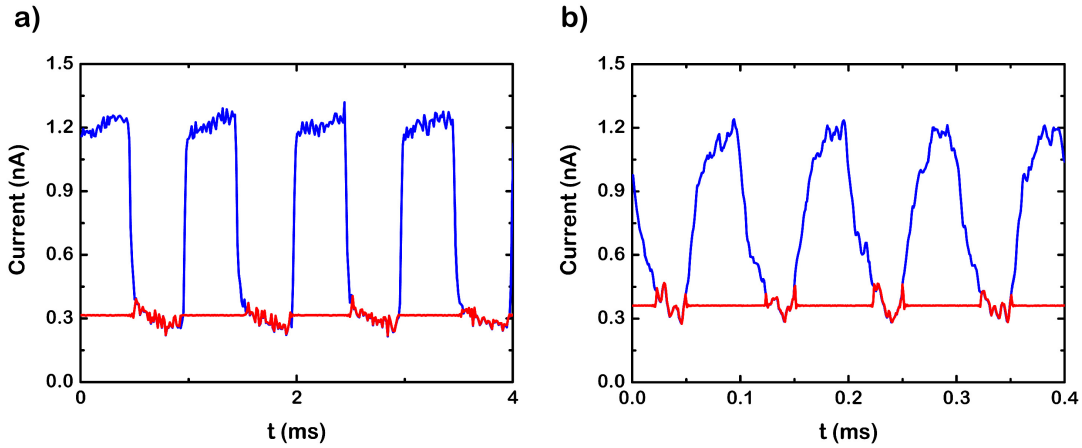


Figure 6: Test of the topo/photo current splitter using the X-ray-induced current recorded with a coaxial Au/polymer coated tip out of tunneling contact in 1 bar O₂. The blue and the red curves show the signal before and after the separation circuit, respectively. a) Chopper at 1 kHz b) Chopper at 10 kHz.

3. Performance

3.1 Current contributions from the gas environment

We have studied the SXSTM background currents generated by photo-electrons and ions by comparing the currents measured with a coaxial tip out of tunneling contact above a Au(111) surface in Ar, O₂ and in a modest vacuum of approximately 1 mbar. Although it is clear that many ionization events occur in the gas phase [31,32], it is not evident how these translate into the detection of a net current. We observed that the background current exhibits complex

behavior, which is strongly dependent on many parameters. For instance, the current measured on the tip core at large tip-sample distances was clearly much higher in Ar than in O₂ (see Figure 7a). In contrast, at a tip-sample distance of only a few micrometers, the current was slightly higher in O₂. In general, the tip current was higher in a gas environment than in vacuum.

From the Au L3 edge absorption spectrum in Figure 7b, it is clear that the tip current follows the absorption coefficient, apart from a difference in background slope. This is expected, as ion current collection is commonly used in conversion electron yield X-ray absorption spectroscopy [32]. Similarly, when changing the beam intensity (data not shown) the current-intensity relation only slightly deviates from linearity.

Figure 7c shows the effects of insulating surfaces in the reactor on the background current. The measurements in this experiment were conducted with the Al dome reactor that naturally has a native insulating oxide layer on all its surfaces in combination with a SiO₂-coated Mo tip *without* a conductive shield. The time-dependent recorded currents on the sample (red) and the tip (blue) show several striking observations: (1) When the X-rays hit the surface (more negative sample currents), electrons are strongly picked up by the tip (positive tip current). However, the tip current quickly decays, although the beam still hits the sample. This clearly indicates charging. A negative charging of the tip insulation by the electrons might explain this effect. (2) When the chopper blocks the X-rays, one observes a significant, negative tip current peak that also decays. Electrons must leave the tip or at least accumulate at the tip surface and insulation interface. This can be explained by the discharging of the tip insulation and the accompanied counter current in the grounded tip wire, which is required to break down the charge separation required for the screening. (3) The sample current also shows charging effects, both when the beam hits as well as stops hitting the sample. Charging of the tip insulation is obviously one reason. However, the magnitude of this effect is probably much smaller than the effect of the insulating Al dome, which forms together with the sample a huge capacitor. (4) Finally, we observed a long time effect, in which the sample current develops a negative offset current while hitting it with the X-rays. As a result, the sample current does not return to 0 even during the “beam off” phase of the chopper. The above described effects are all related to charging and discharging of insulating surfaces in the reactor, which is accelerated in the presence of a gas environment [33–35]. X-ray absorption by the gas was not a major contributor to the background currents, as was shown by measurements with the beam passing through the reactor without hitting the sample.

Figure 7d shows that the background current of the tip depends on the applied sample bias. For non-zero bias, the positively and negatively charged species generated by ionization events in the gas phase are separated by the electric fields between the sample, the tip and the reactor walls. The lack of precise symmetry in the tip current may be related to the fact that a net positive current is injected into the gas phase from the sample. As Figure 7a shows however, the current sign may also be inverted for different gases, again underlining the complexity of the charge dynamics in the background currents.

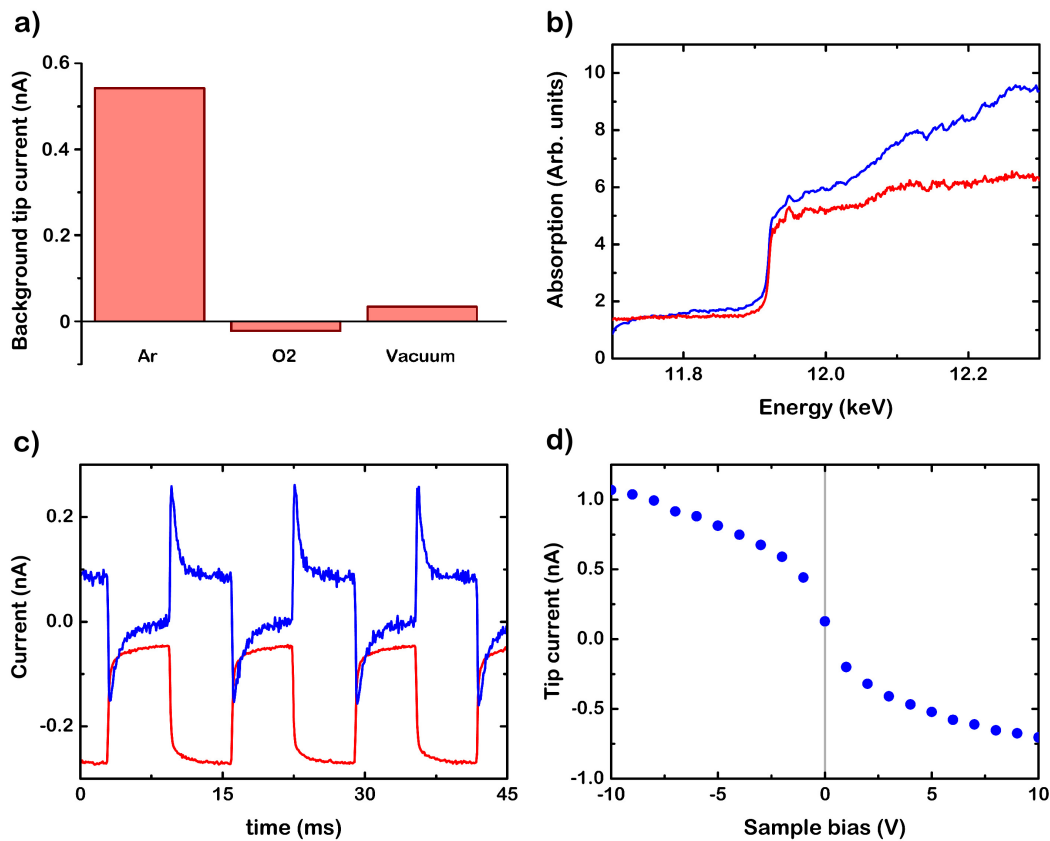


Figure 7: Characteristics of the SXSTM background currents in a gas environment, recorded with the tip out of tunneling contact. a) Background current on the tip in various environments at a photon energy of 12.3 keV for a tip-sample distance of 0.8 mm and a sample bias of -1 V. The Ar or O₂ pressure was 800 mbar, while ‘vacuum’ corresponds to a residual pressure of approximately 1 mbar. b) Au L3 edge absorption spectra from the fluorescence signal on the X-ray detector (red) and the tip current (blue), recorded in 800 mbar Ar. c) Unwanted charging and discharging effects in an uncoated reactor, recorded with a SiO₂/Mo tip *without* Au shield: sample current (red) and tip current (blue) in 500 mbar O₂ with the beam chopped at 80 Hz. d) Dependence of the background signal on the sample bias, recorded in 800 mbar Ar.

3.2 Background reduction using coaxial tips

To test the effectiveness of the coaxial STM tips described in Section 2.2, which consist of a conductive core and shield separated by an insulation layer, we simultaneously measured the current on the tip shield and on the core. As can be seen in Figure 8a, the maximum shield current is two orders of magnitude higher than that of the core. When the beam is placed more than 150 μm away from the tip, the core is completely insensitive towards the X-rays. In this situation, the electric field between tip shield and sample is sufficient to deflect all ions and electrons so that they cannot reach the tip core.

In many SXSTM studies, the STM tips were coated with an insulation layer only, not with a conductive outer shield [20–23]. While this can be adequate in UHV, Figure 7c shows that charging/discharging effects can cause signal distortions in a gas environment. The details of the shape of the distortions depend strongly on the precise beam-tip alignment and the gas environment, which makes it difficult to reproduce results. Hence, it is essential to use STM tips with a conductive outer shield for performing SXSTM measurements in a gas environment.

From Figure 6 it is clear that the coaxial tips provide the required improvement, since the square wave signal is only mildly distorted. It should be noted, however, that the magnitude of the background signal is still in the nA range and varies strongly from tip to tip. The type of insulation layer is not the most important factor, as the polymer-coated tips produced both the best and the worst background. Most likely, the geometrical variation at the tip apex is a major contributor to the observed differences between tips.

In addition to the fast signal distortions of the square wave, some Au/polymer-coated tips showed a slowly increasing offset current, which builds up during several minutes after switching on the beam and can reach values of several hundred pA. The origin of this offset current must be a slow charging effect, which we have not been able to identify further. However, once the offset current has stabilized, it does not interfere with the SXSTM measurements.

To further reduce the background current, we applied a bias voltage to the shield. Figure 8b shows the effect of the bias on the X-ray-induced current received by the tip core. The current was corrected for the leakage current baseline in the absence of X-rays ($2\text{ G}\Omega$ shield-core resistance). Clearly, a small positive bias of $\sim 0.5\text{ V}$ on the shield strongly reduces the background current. Surprisingly, at negative or larger positive shield bias, the background is not effectively reduced or even amplified. This could be due to surface conduction induced by ions landing on the exposed part of the tip insulation layer, as this would increase the leakage current between shield and core.

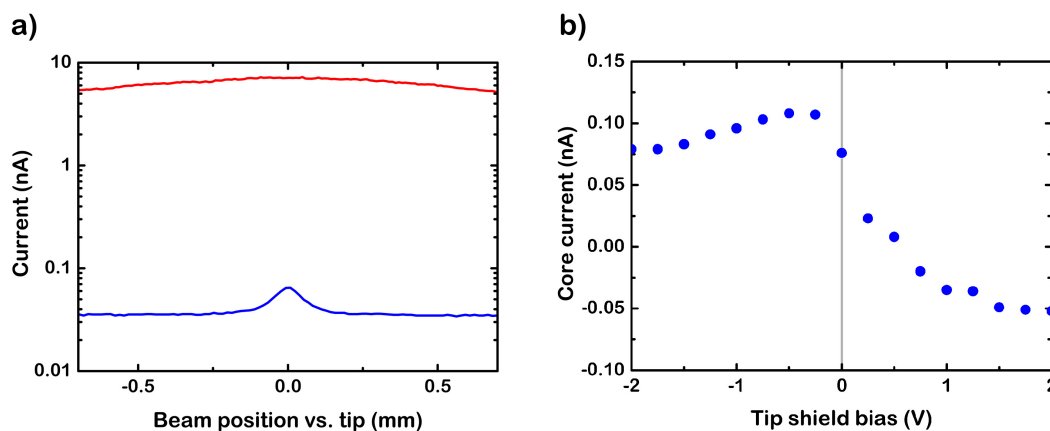


Figure 8: Effectiveness of the background reduction. a) Tip core (blue) and shield current (red) when scanning the beam across the tip in 1 bar O_2 at a tip-sample distance of $10\text{ }\mu\text{m}$. Sample bias: -1 V , tip shield bias: 0 V b) Tip core current versus tip shield bias in 800 mbar Ar.

3.3 Imaging

A requirement in SXSTM measurements is that the tip is sufficiently sharp to properly image the surface: the height resolution should suffice to see atomic height steps and the lateral resolution should be better than 2 nm in order to identify nanoparticles. Figure 9 shows that atomic step resolution on Au(111) was indeed obtained with a Au/polymer coaxial tip in a beam chopped at 10 kHz. Line scans at higher magnification show that the Au(111) steps are imaged with a width of roughly 1 nm. Hence, the tip is sufficiently sharp to meet the lateral resolution criterion.

However, interference is clearly visible in Figure 9, decreasing the practical lateral resolution to approximately 5 nm. Nevertheless, spectral analysis of the tip current signal showed that the 10 kHz chopping frequency was effectively removed by the topo/photo separator described in Section 2.4. Rather, the interference is caused by vibrational and electronic sources coupling into the signal, which are also present in regular STM mode without X-rays and the topo/photo separator. Because the STM was originally designed for X-ray scattering experiments, which require precise control over the incident angle and sample height, the STM could not easily be equipped with a vibrational isolation stage [14]. Improvements to suppress the interference are currently under development.

To get a lower bound estimate of the feedback responsivity, we scanned across a Au step edge with a tip repositioning speed of 5000 nm/s. Because the lateral width of the step edge feature is ~ 1 nm with the Au/polymer tip, the height feedback system needs to respond with a frequency of minimally 5 kHz in order to prevent a collision with the step edge. Since no tip-sample collisions occurred, it is clear that the topo/photo current separator allows for height feedback frequencies that are unfeasible with other separation schemes.

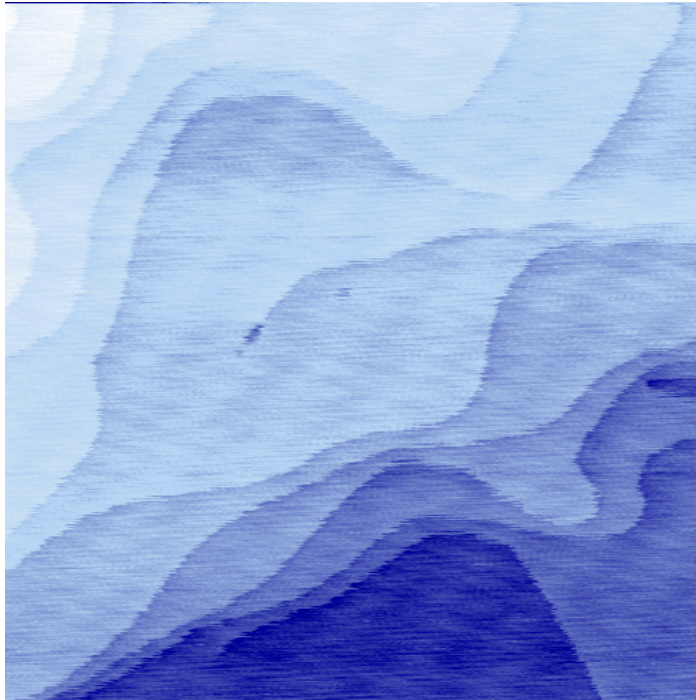


Figure 9: STM image of Au(111) recorded using a Au/polymer coaxial tip in a 12.1 keV beam chopped at 10 kHz in 800 mbar Ar, showing atomic steps. The topo/photo current separator was used to obtain the topographic tunneling current, the input for the STM control system. Scanning parameters: 640 nm × 640 nm, $I_t = 1$ nA, $U_s = -1$ V, tip speed 2500 nm/s.

3.4 X-ray induced tunneling current enhancement

The most essential prerequisite in SXSTM measurements is the ability to measure the X-ray induced increase in the tunneling current, which provides the local X-ray absorption information [17,18]. To provide proof of this ability for our instrument, we measured the X-ray-induced current as a function of the topographic tunneling current. The ion current and other non-local sources of background signal will not depend on the topographic tunneling current set point, because this changes the tip-sample distance within a range of only roughly 1 nm. Hence, a correlation between the topographic tunneling current and the X-ray induced current implies that a local X-ray induced signal is detected: the X-ray induced increase in the tunneling current.

Figure 10 shows the X-ray induced versus the topographic current as recorded on Au(111) in 800 mbar Ar. The X-ray induced signal is shown normalized as (X-ray induced current - ion current background)/ion current background, where the ion current background was measured with the tip out of tunneling contact. Clearly, the magnitude of the X-ray induced signal scales with the topographic tunneling signal, providing a proof-of-principle of our ability to perform SXSTM measurements in a gas environment.

Although the mechanistic details of the processes underlying the X-ray induced contribution to the tunneling current have not been rigorously clarified, the correlation in Figure 10 could be interpreted as follows: the X-ray-induced modification of the tunneling current is assumed to arise from hot secondary

electrons [18,38]. Most of the hot electrons do not have sufficient energy to escape from the sample into the vacuum, but they do have a much higher tunneling probability to the tip than electrons at the Fermi level. Still, this enhanced tunneling probability depends on the tip-sample distance, which changes when the topographic tunneling current setpoint is changed. Thus, the X-ray induced increase in the tunneling current will be higher at smaller tip-sample distance (high topographic tunneling current). Both thermalized and hot electrons have an exponential dependence on the tip-sample distance, each with an exponent that depends on their energy with respect to the vacuum state.

$$I_{t \text{ thermal}} \sim e^{-\alpha d}$$

$$I_{t \text{ hot}} \sim e^{-\beta d}$$

As a result, the relationship between the tunneling current arising from thermal electrons (topographic tunneling current) and the current arising from hot electrons (X-ray induced tunneling current) is given as:

$$I_{t \text{ hot}} \sim I_{t \text{ thermal}} e^{-(\beta-\alpha) d}$$

The linear correlation observed in Figure 10 could imply that the energy of the hot electrons is only marginally higher than the Fermi level. In such a case

$$I_{t \text{ hot}} \approx I_{t \text{ thermal}} \times e^{-(\beta-\alpha) d}$$

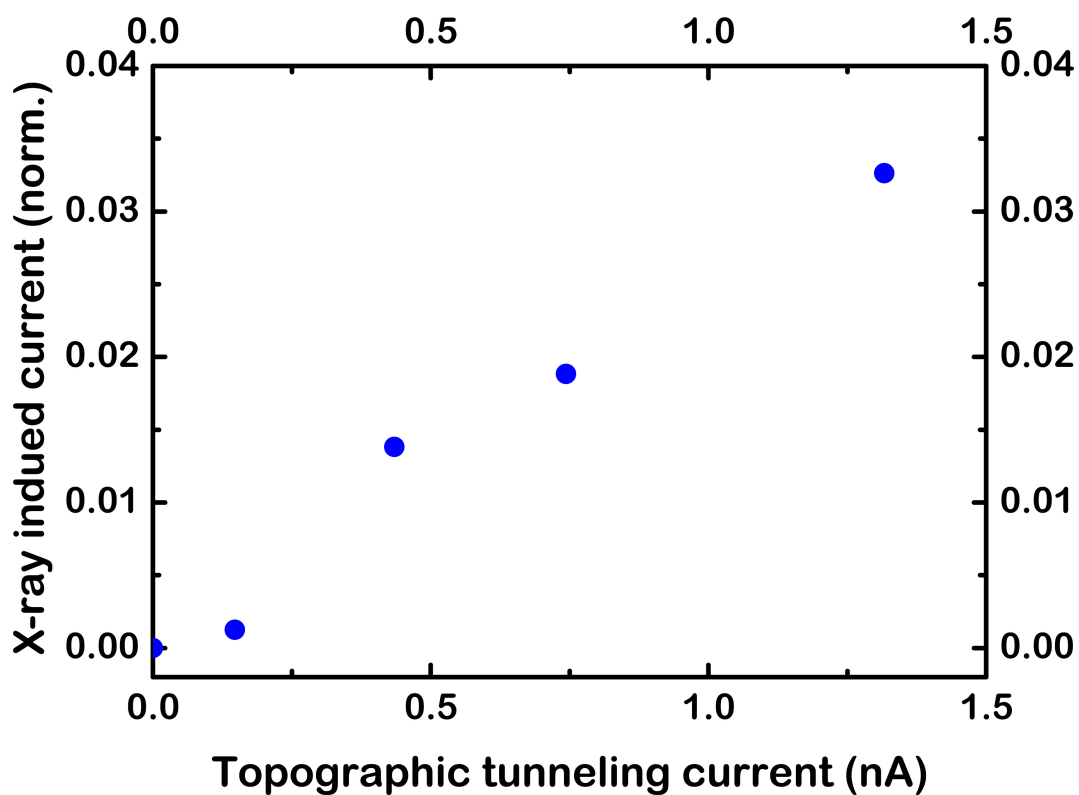


Figure 10: X-ray-induced current versus topographic tunneling current, recorded on Au(111) in 800 mbar Ar using the topo/photo splitter with the 12.1 keV X-rays chopped at 10 kHz. The normalized X-ray induced current was obtained as (X-ray induced current-ion current background)/ion current background, where the ion current

background was measured with the tip out of tunneling contact. Sample bias: -1 V, tip shield bias: 0 V.

Conclusion and outlook

We have developed an instrument suitable for performing local X-ray absorption measurements by modification of a previously developed high-pressure STM and X-ray scattering apparatus [14]. Several essential additions have been made in order to reach the required sensitivity to measure the small X-ray-induced modification of the tunneling current that carries the local X-ray absorption signal [17,18].

First, the STM is mounted on a new high-pressure cell with thin windows, providing high X-ray transparency in the energy range of interest, from 8 keV to 25 keV. Furthermore, we have shown that the beam shadow of the STM tip on a 2D X-ray detector can be used to align tip and beam with a precision of a few micrometer. Our measurements show that background currents from ions and photo-electrons cause much more severe signal distortions in a gas environment than in vacuum, particularly in the presence of insulating surfaces. We have therefore applied a Au coating to the high-pressure cell and employed coaxially shielded STM tips. The conductive outer layer of the coaxial tips can be biased to provide additional ion deflection. To separate the regular, 'topographic' tunneling current from the X-ray-induced current, a new high-speed electronics circuit was developed. Using these technical implementations, the X-ray induced modification of the tunneling current was measured on Au(111) in 800 mbar Ar. Combining our methodology with the knowledge of the spatial and chemical resolution of SXSTM developed in the literature [17,18], *operando* X-ray absorption measurements with nanometer spatial resolution have now come within reach.

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