A new scanning tunneling microscope reactor used for high-pressure and high-temperature catalysis studies

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We present the design and performance of a homebuilt high-pressure and high-temperature reactor equipped with a high-resolution scanning tunneling microscope (STM) for catalytic studies. In this design, the STM body, sample, and tip are placed in a small high pressure reactor (~19 cm³) located within an ultrahigh vacuum (UHV) chamber. A sealable port on the wall of the reactor separates the high pressure environment in the reactor from the vacuum environment of the STM chamber and permits sample transfer and tip change in UHV. A combination of a sample transfer arm, wobble stick, and sample load-lock system allows fast transfer of samples and tips between the preparation chamber, high pressure reactor, and ambient environment. This STM reactor can work as a batch or flowing reactor at a pressure range of 10⁻¹³ to several bars and a temperature range of 300–700 K. Experiments performed on two samples both in vacuum and in high pressure conditions demonstrate the capability of in situ investigations of heterogeneous catalysis and surface chemistry at atomic resolution at a wide pressure range from UHV to a pressure higher than 1 atm.

I. INTRODUCTION

The understanding of heterogeneous catalysis at a molecular level has been one of the central issues of physical chemistry for the past half century.1 Single crystal surfaces have served as valuable model catalysts providing insights into heterogeneous catalysis under vacuum conditions.2 This vacuum surface science approach of catalysis studies has revealed a tremendous amount of information for a great number of catalytic systems. However, industrial heterogeneous catalytic reactions are usually carried out at high pressure and high temperature. There are considerable experimental challenges in the high pressure studies, which are necessary in order to understand molecular behavior under realistic conditions. The potential difference in adsorption, surface structures, and catalytic mechanisms between the model studies at low pressure and industrial reactions at high pressure is often referred as pressure gap.2,3

A key component in studying the pressure gap is to characterize the adsorbed layer of the reactant gases at high pressure during catalytic processes. A simple extrapolation of the insights into the adsorption structure obtained at low pressure and low temperature is not necessarily applicable to high pressure and high-temperature conditions which could have different energetic pathways. Thus, to obtain a complete understanding of catalysis, it is necessary to perform studies of surface catalytic reactions under high pressure of reactants.

Scanning tunneling microscope (STM) has the unique capability of studying catalyst surfaces atom by atom, which is invaluable for elucidating the adsorption structure and the mobility of reactant molecules during catalysis. This technique can be applied in a pressure range from UHV to atmospheric or higher pressure since the tunneling process between the sample and tip only occurs in a very close range of 5–50 Å. It has been applied to catalytic studies under a condition of relatively high pressure by a few groups4–9 since the first demonstration.4 In our laboratory we performed high pressure studies of STM by filling reactant gases into STM chamber connected to the UHV preparation chamber.4,5 However, this method has disadvantages such as large volume of reactant gases and limits in sample heating, reactant gas pressure, and spatial resolution. The new high pressure and high-temperature reactor STM and UHV system presented here overcome these limitations and allows for catalytic studies under a wide range of pressure (from 10⁻¹³ to several bars) and temperature (from 300 to 700 K). All parts of this high pressure high-temperature STM reactor/UHV system including STM body, high pressure high-temperature reactor, in situ heating system, tip and sample change system, sample transporter, and wobble stick, sample annealing stage were newly designed and homebuilt in the two years of 2006 and 2007.10 The high pressure reactor is a small cylinder chamber with a volume of approximately 19 cm³, placed inside the vacuum environment of the UHV chamber by a special docking scaffold and mounting framework. It is vibrationally isolated from the UHV chamber with three springs, offering the capability of imaging surface with atomic resolution. The sample can be heated in situ to 700 K by an external heating lamp installed under the high pressure reactor in the STM chamber. Both the sample and tip can be

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conveniently placed and transferred using a transfer arm and wobble stick.

II. APPARATUS

This section presents the details of our high pressure and high-temperature reactor STM. The whole STM system includes sample preparation chamber, STM chamber, and sample/tip loading system (Fig. 1). We will discuss sample preparation and STM chambers, high pressure reactor and gas introduction system, STM body and sample heating, and sample transfer and tip change in Secs. II A–II D, respectively.

A. Sample preparation and STM chambers

The sample preparation chamber is equipped with an Auger electron spectrometer for analysis of surface composition. It is pumped by a sputtering ion pump and a turbomolecular pump with a base pressure of $7 \times 10^{-14}$ bar. Sample cleaning and preparation are carried out by an argon ion gun for sputtering and an electron bombardment gun for annealing. Sample temperature is monitored by using an ex situ infrared thermometer (pyrometer). During cleaning, the sample holder is placed in a four-finger sample stage fabricated out of alumina for thermal isolation. The sample can be heated to 1000 °C with an increase in background pressure less than $2 \times 10^{-12}$ bar. The four-finger sample stage matches a three-finger stainless steel fork on a transfer rod allowing the sample holder to be transferred to the STM chamber after sample preparation. A universal sample holder and tip exchanger on a wobble stick were designed for engaging with the sample stage, storage slots on a docking disk, and sample stage in the high pressure reactor for rapid transfer of samples and tips.

STM chamber is also a stainless steel chamber pumped by its own sputtering ion pump and turbomolecular pump. It has a base pressure of $1.2 \times 10^{-13}$ bar. This chamber is separated from the sample preparation chamber by an 8 in. gate valve. A quadrupole mass spectrometer is installed in this chamber for monitoring reactants and products leaked from Kalrez o-rings of high pressure reactor during STM scanning. The STM chamber contains the high pressure reactor Fig. 1 (5), a docking scaffold assembled on a custom-designed sample manipulator, and a wobble stick for sample and STM tip transfer between the transfer rod, high pressure reactor, and docking disk.

The sample heating source is located outside the high pressure reactor to avoid heating elements in the high pressure environment. It consists of a halogen lamp with an elliptical reflector that focuses the radiation onto the sample through a sapphire window welded at the bottom of the reactor (Fig. 2). The distance between the lamp and the reactor can be adjusted to focus the light on the back of the sample for efficient heating. The heating rate can be controlled by adjusting the power supplied to the lamp.

B. High pressure reactor and gas introduction system

The high pressure reactor is a special vessel designed for assembling the STM body and sample, and providing a high pressure environment of reactant gases. It houses a home-built STM body. A sample assembly stage is mounted at the end of the STM body. The sample is thermally isolated and electrically insulated from the STM body. Figure 3(a) is a picture of the high pressure reactor with a volume of $\sim 19$ cm$^3$. The reactor consists of cell lid, cell neck, middle stage, and bottom stage. The cell lid has a set of precisely
designed holes to glue a set of pin-socket contacts [set II in Fig. 3(a)] for assembling a set of male contacts (set I) from the docking scaffold and a set of male contacts [set III in Fig. 3(c)] for the wiring connections from the shear piezoelectric plates and the scanning tube. These pin-socket contacts provide convenient detachable wiring connections for the high pressure reactor. The contacts (set I) from the docking scaffold can be inserted to the vacuum side of the interfacial contacts (set II) glued to the cell lid. Another set of contacts (set III) is glued on a set of holes of the STM body [Fig. 3(b)] which have the exact same size and arrangement as the holes on the lid [Fig. 3(a)]. This pin-socket wiring structure [Fig. 3(c)] makes dismantling and assembling the STM convenient when maintenance is required on the high pressure reactor and STM body.

For sample transfer and tip change, a port is opened on the wall of the bottom stage of the high pressure reactor [Fig. 3(a) (9)]. A sapphire window is welded at the center of the bottom of the reactor to transmit light to heat the sample. Recesses are fabricated on the cell neck, cell top stage, cell bottom stage, and port of the reactor, hosting Kalrez o-rings forming gas-tight seals. The sealing of Kalrez o-rings allows pressurization of the reactor while maintaining a high vacuum in the surrounding chamber. All four sections of the reactor are assembled together by four venting screws and sealed by these o-rings. A bayonet seal [inset of Fig. 3(a)] is fabricated to seal the port on the reactor. The reactor was plated with a layer of gold (thickness: ~5 μm) to avoid possible reactions between the materials of the high pressure reactor and reactant gas.

Figure 4 schematically shows the setup of gas introduc-
tion for the high pressure reactor. For gas introduction, the male part of a Swagelok fitting ([4] in Fig. 4) is welded on a 1/8 in. tube of the cell lid ([3] in Fig. 4). A 1/8 in. PEEK tube ([7] in Fig. 4) capable of supporting high pressure was glued on a 1/8 in. stainless steel tube ([6] in Fig. 4) assembled into a female part of the Swagelok fitting ([5] in Fig. 4). Another end of the PEEK tube is also glued and assembled to another Swagelok fitting welded to a hole of a double side conflat (CF) flange. This design isolates the reactant gases in the high pressure reactor from the vacuum environment of the STM chamber. The high pressure gases in the reactor can be quickly pumped down by a turbomolecular pump to obtain a UHV environment after completion of a high pressure experiment. Thus, this high pressure reaction system can work under both UHV and high pressure, offering the capability of studying catalysts over a wide pressure range from $1.2 \times 10^{-13}$ to several bars. In addition, the reactions can be carried out with batch or flowing mode.

C. STM body and sample heating

The STM body [Fig. 3(b)] is the key component of the high pressure reactor. As mentioned above, it is screwed onto the cell neck stage. The STM body includes a coarse approaching system, a scanning tube, a receiver of the tip holder, and wire connections to these parts. The coarse approach is carried out by six sets of shear piezoelectric plates [three of them are schematically shown in a bottom view of STM body in Fig. 3(d)] located between a hexagonal sapphire [Fig. 3(b) (2)] and the wall of the STM body. One side of each shear piezoelectric set is glued on the internal wall of the STM body while the other end contacts the surface of the hexagonal sapphire [Figs. 3(d) and 3(e)]. By applying negative or positive voltages to the first/third and the second/fourth piezoelectric plates, respectively, the lateral force moves the hexagonal sapphire forward and backward [Fig. 3(e)]. A single piezoelectric scanning tube is glued to an alumina disk which is, in turn, glued to one end of the hexagonal sapphire. Five Kapton wires are glued to the five components ($+x$, $-x$, $+y$, $-y$, and $z$) of the scanning tube through holes on the alumina disk. Another alumina disk is glued to the other end of the scanning tube onto which a small SmCo magnet and a bowl-shaped tip receiver is glued [Fig. 3(b) (4)]. The central part of the receiver of tip holder is a SmCo magnet. The tip change mechanism is described below. A flexible coaxial wire is glued to this tip receiver for transmitting the tunneling current.

At the front of the STM body one CuBe plate spring [Fig. 3(b) (3)] is used to hold two sets of shear piezoelectric plates. The pressure applied to the hexagonal sapphire by the shear piezoelectric sets can be fine tuned by a screw [Fig. 3(b) (5)] in the spring plate. This pressure can change the speed of the coarse approach. A K-type thermocouple is spot welded to the sample stage for both sample bias and temperature measurements. A second thermocouple is attached to the STM body to monitor the temperature of the shear piezoelectric plates during sample heating and reaction at high temperature. Thus, thermal diffusion and possible increases in the temperature of the STM body can be simultane-ously monitored when the sample in the high pressure reactor is heated.

A sample assembly stage is screwed to the end of the STM body [Fig. 3(b) (6)], which is thermally and electrically insulated from it by three precisely aligned sapphire balls and insulating washers. Thus, our experiments show that the shear piezoelectric plates, scanning tube, and SmCo magnet do not degrade when the sample is at a temperature of 700 K and the pressure of reactant gases is 1 bar.

D. Sample transfer and tip change

The sample transfer between STM chamber and sample preparation chamber is carried out by a magnetic transfer rod with a three-finger fork. A wobble stick can transfer the sample and tip between the three-finger fork, the high pressure reactor, and the slots in the docking disk. It can accept and release the sample holder conveniently. In addition, the sample holder and tip exchanger can be introduced or removed from the system through a load-lock system.

Replacement of the STM tip is accomplished by a magnetic tip exchanger with the same geometry as the sample holder [Fig. 5(a)] and a tip holder [Fig. 5(b)]. The tip exchanger can be easily transferred to and from the high pressure reactor [Fig. 5(c)], the storage disk, and the load-lock system.

III. PERFORMANCE

In this section we illustrate the performance of the instrument with two examples of experiments carried out under UHV and high pressure conditions. These include highly ordered pyrolytic graphite (HOPG) and hex-Pt(100) single crystal. In Sec. III A, the results of a clean HOPG surface and a self-assembled organic monolayer on HOPG under ambient and UHV conditions will be discussed. In Sec. III B,
STM images of a clean hex-Pt\(_{100}\) surface with atomic resolution will be presented along with the roughening of the surface upon reaction with high pressure CO.

A. Clean HOPG and self-assembled monolayer on HOPG under both ambient and UHV conditions

Atomically resolved images can be routinely obtained on HOPG samples, both under UHV and in ambient condition at a tunneling current of 1.0–2.0 nA and sample bias of 0.1–0.5 V [Fig. 6(a)].

To test the behavior of the STM body under high pressure conditions, a HOPG sample deposited with a self-assembled monolayer of hexadecanedioic acid is assembled in the reactor. Then, 1 bar nitrogen was introduced into the reactor while the STM chamber is pumped down to high vacuum. Images of the self-assembled monolayer with atomic resolution, as shown in Fig. 6(b), can be obtained under an environment of 1 bar nitrogen. This image has five lamellae. Each lamella in terms of the section between two adjacent blue lines consists of parallel packed molecule. Similar to the self-assembly of other carboxylic acids on HOPG,\(^{11}\) the ordered self-assembled structure is formed through intermolecular hydrogen bonds between two adjacent lamellae. This demonstrates the satisfactory performance of the homebuilt STM body and high pressure reactor.

B. Reactive surfaces and CO induced roughness under high pressure

CO oxidation is an extremely important industrial catalytic process. Platinum is an active catalyst for this reaction. Here hex-Pt\(_{100}\) is selected as a highly reactive surface. The CO adsorption on hex-Pt\(_{100}\) under a wide range of CO pressures was studied with this instrument. As is well known, the top layer of a clean hex-Pt\(_{100}\) is a quasihexagonal layer with \(\sim 20\%\) extra Pt atoms in contrast to the underlying layer with a \(1 \times 1\) structure. The clean hex-Pt\(_{100}\) was prepared by the procedure reported in literature.\(^{12}\) After it is cleaned by a combined Ar\(^+\) sputtering, annealing in oxygen environment, and a final annealing to 1150 K for several minutes in UHV followed by a slow cooling to room temperature, a clean hex-Pt\(_{100}\) surface is formed. Our STM gives images of the clean hex-Pt\(_{100}\) with atomic resolution at room temperature [Fig. 7(b)]. It clearly demonstrates the reliability of the STM body, sample heating, sample sputtering, sample transfer, and tip exchange mechanism of this homebuilt system. Upon exposed to an environment of \(\sim 0.9\) bar CO, the clean hex-Pt\(_{100}\) significantly restructures and presents as a surface covered with clusters with sizes ranging from 2 to 5 nm. Figure 8 is one image of the highly roughed surface formed in an environment of high pressure CO. A more extensive account of these results will be reported elsewhere.\(^{13}\)

IV. SUMMARY

A new high pressure and high-temperature STM reactor was designed and homebuilt with the purpose of simulating industrial catalysis reaction conditions. The STM body is housed in a high pressure reactor equipped with \textit{in situ} heating and fast sample transfer and tip change. We have demonstrated the good performance of this instrument with ex-
amples that include HOPG, both clean and with adsorbed hexadecanedioic acid. We have also shown results of the CO induced reconstruction of hex-Pt\textsubscript{100} over a wide range of CO pressures, demonstrating the capability of studying catalytic reactions at atomic resolution in a high pressure environment. This STM will serve as an important tool in the effort to overcome the pressure gap of catalysis studies and for surface science studies at a condition of high pressure.

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10 In 2004, a few parts of a STM body were made with an attempt of building a STM reactor by a student. But those parts made in 2004 did not work.