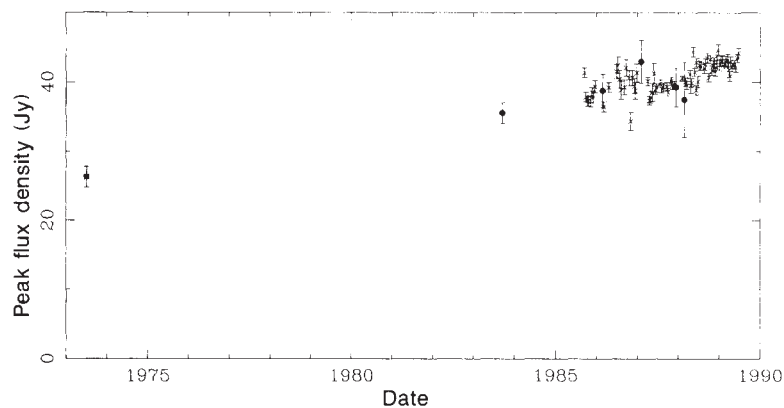


FIG. 4 The peak flux density of the OH 1,612-MHz emission from OH0.9+1.3 plotted as a function of time. Crosses show Hartebeesthoek measurements, circles show Jodrell Bank measurements, and the square is the discovery measurement taken from ref. 21.



expanding-shell model, as shown in Fig. 3. Because the data do not properly constrain a least-squares fit of all parameters, we adopted the stellar velocity of -44 km s^{-1} found from CO measurements¹⁵. This leads to a shell radius of 0.39 ± 0.05 arcsec, an expansion velocity of 18 km s^{-1} (which agrees with the CO data) and a stellar position that agrees with the position of the compact H II region to within the combined errors of ± 0.2 arcsec.

It is well established that the OH shell radii of OH-IR sources increase systematically with mass loss rate^{12,16,17}. This comes about because the OH is produced when H_2O molecules in the outflowing gas are photodissociated by the external ultraviolet radiation. For a spherically symmetric outflow, the OH photodissociation radius is then determined by the mass loss rate and the strength of the ambient ultraviolet radiation field. By a happy coincidence, physical conditions at this photodissociation radius are also right for pumping the OH 1,612-MHz maser¹⁸. Mass loss rates for Vy2-2 and OH0.9+1.3 have been determined as $2.4 \times 10^{-6} M_{\odot} \text{ yr}^{-1}$ and $3.0 \times 10^{-5} M_{\odot} \text{ yr}^{-1}$ for assumed distances of 1 and 8 kpc respectively^{14,15}. These imply OH photodissociation radii of $6.3 \times 10^{13} \text{ m}$ and $3.9 \times 10^{14} \text{ m}$ for an average interstellar ultraviolet flux, which agree well with the values of $5.8 \times 10^{13} \text{ m}$ and $2.9 \times 10^{14} \text{ m}$ derived from our measurements. Thus the OH shells in these two sources are indeed fossil remnants of the OH-IR phase, having the brightness temperatures and shell radii characteristic of OH-IR sources.

Clearly we have been fortunate to catch these two objects in the transition phase: the expansion timescale of the compact H II regions is only 300 years. In fact, the molecular envelopes could evolve even more rapidly, particularly if there is a fast wind, as suggested by recent observations of Herbig-Haro objects and high-velocity outflows in other protoplanetary nebula candidates^{19,20}. Strong changes in the OH maser emission could then occur as the inner parts of the envelope are disrupted. It is possible that such disruption is now being observed in OH0.9+1.3 in the powerful masers at high negative velocity. Single-telescope monitoring has revealed a steady increase in the negative-velocity peak (Fig. 4). The flux density of the emission seems to have increased at the rate of 1 Jy yr^{-1} since the source was discovered in 1973²¹. Such behaviour is unprecedented for an OH-IR source. The implied age of the feature is about 40 yr. It may be possible to resolve the structure of these masers using VLBI. Measurements of the temporal evolution of this remarkable source could be crucial to understanding the evolutionary path from OH-IR sources to planetary nebulae. □

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Positioning single atoms with a scanning tunnelling microscope

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SINCE its invention in the early 1980s by Binnig and Rohrer^{1,2}, the scanning tunnelling microscope (STM) has provided images of surfaces and adsorbed atoms and molecules with unprecedented resolution. The STM has also been used to modify surfaces, for example by locally pinning molecules to a surface³ and by transfer of an atom from the STM tip to the surface⁴. Here we report the use of the STM at low temperatures (4 K) to position individual xenon atoms on a single-crystal nickel surface with atomic precision. This capacity has allowed us to fabricate rudimentary structures of our own design, atom by atom. The processes we describe are in principle applicable to molecules also. In view of the device-like characteristics reported for single atoms on surfaces^{5,6}, the possibilities for perhaps the ultimate in device miniaturization are evident.

The tip of an STM always exerts a finite force on an adsorbate atom. This force contains both Van der Waals and electrostatic contributions. By adjusting the position and the voltage of the tip we may tune both the magnitude and direction of this force. This, taken together with the fact that it generally requires less force to move an atom along a surface than to pull it away from the surface, makes it possible to set these parameters such that the STM tip can pull an atom across a surface while the atom remains bound to the surface. Our decision to study xenon on nickel (110) was dictated by the requirement that the corrugations in the surface potential be sufficiently large for the xenon atoms to be imaged without inadvertently moving them, yet sufficiently small that, when desired, enough lateral force could be exerted to move xenon atoms across the surface.

The experiments were performed using an STM contained in an ultra-high-vacuum system and cooled to 4 K. The entire

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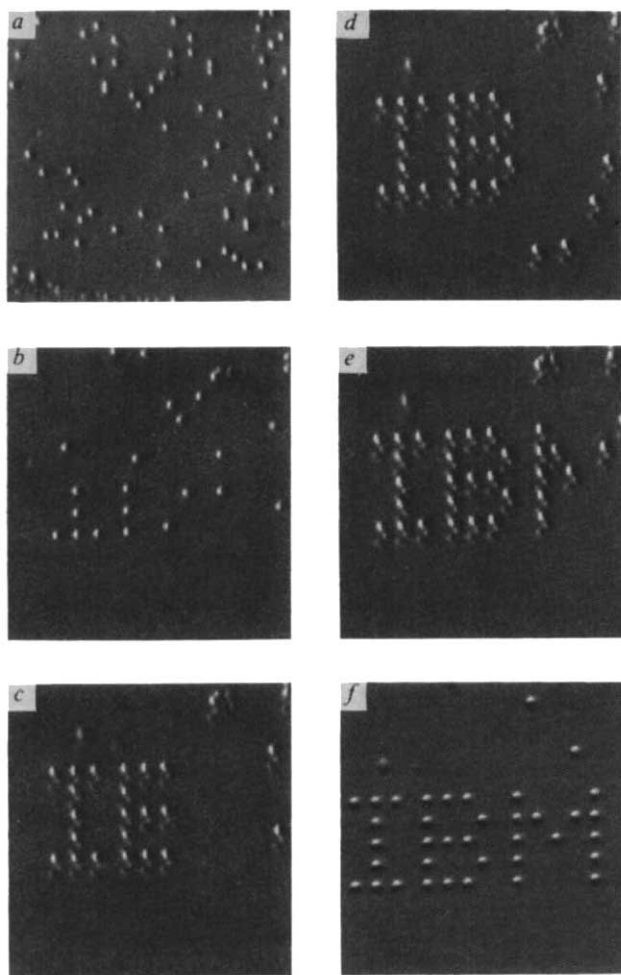


FIG. 1 A sequence of STM images taken during the construction of a patterned array of xenon atoms on a nickel (110) surface. Grey scale is assigned according to the slope of the surface. The atomic structure of the nickel surface is not resolved. The $(\bar{1}10)$ direction runs vertically. *a*, The surface after xenon dosing. *b-f*, Various stages during the construction. Each letter is 50 Å from top to bottom.

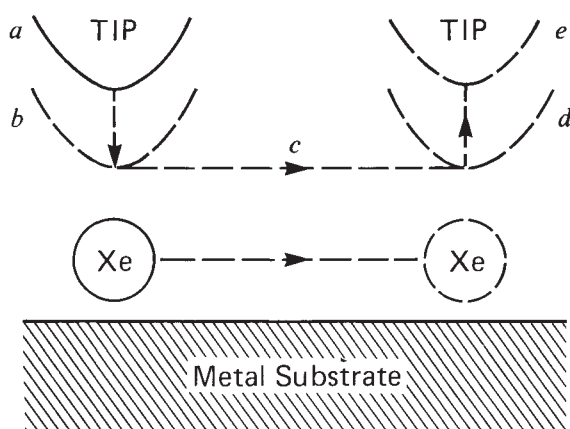


FIG. 2 A schematic illustration of the process for sliding an atom across a surface. The atom is located and the tip is placed directly over it (*a*). The tip is lowered to position (*b*), where the atom-tip attractive force is sufficient to keep the atom located beneath the tip when the tip is subsequently moved across the surface (*c*) to the desired destination (*d*). Finally, the tip is withdrawn to a position (*e*) where the atom-tip interaction is negligible, leaving the atom bound to the surface at a new location.

chamber housing the microscope was cooled to 4 K, which so reduced the contamination rate of the sample surface through adsorption of residual gases that no measurable contamination occurred over weeks. The stability of the sample and of the microscope due to the low temperature are such that one may perform experiments on a single atom for days at a time. This stability proved to be an operational necessity for performing these experiments. This notwithstanding, it is important to realize that the process that we describe for sliding atoms on a surface is fundamentally temperature independent.

The nickel sample was processed by cycles of argon-ion sputtering, annealing in a partial pressure of oxygen to remove surface carbon, and flash annealing. After cooling to 4 K the sample was imaged with the STM and found to be of acceptable quality. Figure 1*a* is an image of the surface taken under constant-current scanning conditions after dosing with xenon to the desired coverage. This image was obtained with a tip bias voltage of 0.010 V relative to the sample, and a tunnel current of 10^{-9} A. Each xenon atom appears as a 1.6-Å-high bump on the surface, apparently at random locations. At this gap impedance the interaction of the xenon with the tip is sufficiently weak to leave the xenon essentially unperturbed during the imaging process.

To move an atom we follow the sequence of steps depicted in Fig. 2. We begin by operating the microscope in the non-perturbative imaging mode described above to locate the atom to be moved and to target its destination. We stop scanning and place the tip directly above the atom to be moved (*a*). We then increase the tip-atom interaction by lowering the tip toward the atom (*b*); this is achieved by changing the required tunnel current to a higher value, typically in the range $1-6 \times 10^{-8}$ A, which causes the tip to move towards the atom until the new tunnel current is reached. We then move the tip under closed-loop conditions across the surface, (*c*), at a speed of 4 Å per second to the desired destination (*d*), dragging the xenon atom with it. The tip is then withdrawn (*e*) by reducing the tunnel current to the value used for imaging. This effectively terminates the attraction between the xenon and the tip, leaving the xenon bound to the surface at the desired location.

Figure 1 is a sequence of images taken during our first construction of a patterned array of atoms, and demonstrates our ability to position atoms with atomic precision. The exact periodicity of the xenon spacing is derived from the crystalline structure of the underlying nickel surface (which is not resolved in Fig. 1). The nickel (110) surface has an unreconstructed rectangular unit cell and is oriented such that the short dimension of the rectangular surface unit cell runs vertically in the image. The xenon atoms are spaced on a rectangular grid which is four nickel unit cells long horizontally and five unit cells long vertically, corresponding to 14×12.5 Å.

Although the STM tip is made from tungsten wire, the true chemical identity and the structure of the outermost atoms of the tip are not known to us. We find that for any given tip and bias voltage, there is a threshold height below which the tip must be located to be able to move xenon atoms parallel to the rows of nickel atoms, and a lower threshold height for movement perpendicular to the rows of nickel atoms. This is consistent with a simple model wherein the xenon interaction with the metal surface is approximated by pairwise interactions with the individual nickel atoms. Simple investigations showed that the magnitude or sign of the applied voltage had no significant effect on the threshold tip height. This suggests that the dominant force between the tip and the xenon atom is due to the Van der Waals interaction, but this tentative conclusion requires further investigation.

In Fig. 3 we show a sequence of images demonstrating how a simple structure, the linear multimer, may be fabricated using this process. First we slide xenon atoms into a chosen row of nickel atoms. We next slide a xenon atom along the row to a position where it will bind with a neighbouring xenon to form

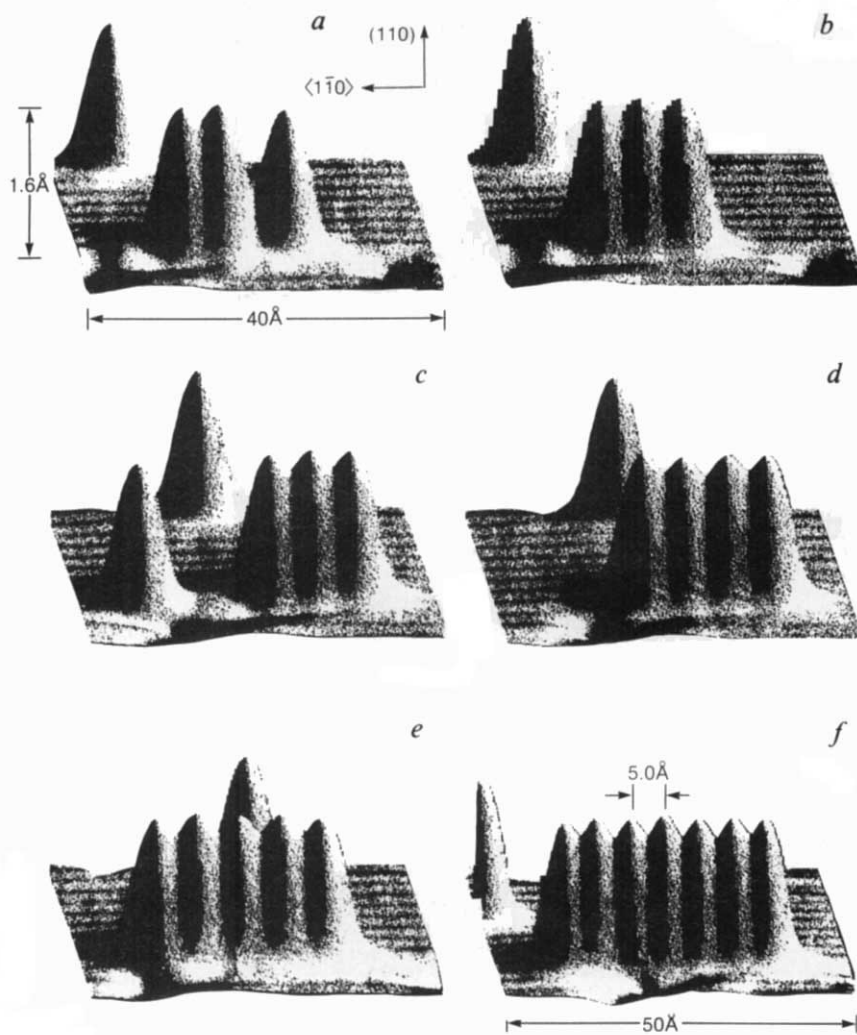


FIG. 3 Various stages in the construction of a linear chain of xenon atoms on the nickel (110) surface. The individual xenon atoms appear as 1.6-Å-high protrusions in these images. The rows of nickel atoms are visible as alternating light and dark stripes. Several point defects are visible in the nickel surface. *a*, The assembled xenon dimer. To the right of the dimer, a xenon atom has been moved into position for forming a xenon trimer. *b*, Formation of the xenon linear trimer. *c-f*, Various stages in construction of the linear heptamer, a process that can be completed in an hour. The xenon atoms are 5 Å apart, occupying every other unit cell of the nickel surface.

a dimer. We repeat the process, forming a linear trimer, then a linear tetramer, and so on. From these images we find that the linear xenon chain along the $\langle \bar{1}10 \rangle$ direction of the nickel (110) surface is stable, the xenon atoms occupying every other surface unit cell along a row of nickel atoms. Attempts to pack the xenon atoms closer were unsuccessful. We find the xenon-xenon spacing along the row to be uniform (excluding end effects) to within 0.2 Å.

We anticipate that there will be a limiting class of adsorbed atoms and molecules that may be positioned by this method. Many new avenues of investigation are open to us. It should be possible to assemble or modify certain molecules in this way. We can build novel structures that would otherwise be unobtainable. This will allow a new class of surface studies that use the STM both to fabricate overlayer structures and to probe their properties. The prospect of atomic-scale logic circuits and other devices is a little less remote. □

A tri-platinum complex containing a coordinatively naked platinum atom

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POLYNUCLEAR clusters of platinum and palladium containing metal-metal bonds may serve as models of the surface of heterogeneous catalysts, and their reactions may mimic those on such surfaces^{1,2}. Ligand-free metal clusters can be created in an ultradispersed condensed phase³, but they are difficult to isolate and characterize because of their high reactivities. Here we report the preparation of a tri-platinum diphosphine complex that contains a coordinatively 'naked' platinum atom. The metal cluster in this complex is electron-deficient relative to other tri-platinum complexes. Bonding of ligands to the naked platinum could reflect that of chemisorbed species on catalytic platinum surfaces.

The potentiostatic electrolysis of $[\text{Pt}(\text{diphos})(\text{RNC})_2](\text{PF}_6)_2$

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