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The position of Loki-B in March is constrained in one dimension by its occultation egress time; the constraint in the other dimension is based on its position in the images relative to Kanehekili and the assumption that Kanehekili was fixed. The resulting error box includes Loki Patera, a prominent dark lake-like feature seen by Voyager, lending some support to this reasoning.

A bright hotspot, presumably Loki-A, was also seen in speckle observations by Howell (personal communication), polarization measurements made by some of us in November and December (Goguen and Sinton), and in eclipse observations on 8 November 1989 (Spencer and Sinton). If Loki-A was actually in the same location as Loki-B, at or near Loki Patera, its 3.8-µm flux dropped by a factor of 10 between December 1989 and March 1990. But if our tentative position for Loki-A is correct, it faded below detectability by March 1990. Our tentative location places it close to Amaterasu Patera, identified as a hotspot by Voyager in 1979*. In December it may have masked the much fainter lower-latitude emission from Loki-B. Loki-A’s 3.8-µm December flux is comparable with previous observations of Loki at its brightest1. A similar dramatic fading of a hotspot in the Loki region occurred between December 1985 and September 198610.

In contrast, Kanehekili’s 3.8-µm flux was virtually the same in December and March, indicating a different type of volcanic activity. The December error box, but not the position assuming no motion between December and March, includes a conspicuous unnamed dark rectangular area of flows (but not scanned in the infrared) by Voyager.

The third, very faint hotspot seen in the March occultation egress is farther west, in a region seen only at low resolution by Voyager (Fig. 3). Because its position is only constrained in one dimension it is not yet possible to correlate it with any specific feature seen by Voyager. Its error ‘box’ just excludes the location of the short-lived source Poliiahu seen on 10 July 198511,12.

The 24 December 1989 temperature of Loki-A is constrained to be \(\geq 370\) K by its 3.8-µm flux and the fact that its radius is \(\approx 120\) km, if circular. Because all occultation observations are at 3.8 µm, the only direct temperature measurements come from post-occultation images taken in March in which Loki and Kanehekili are resolved at both 3.8 µm (Fig. 1d) and 4.8 µm. Colour temperatures (Table 1) are within the normal range for Io hotspots13. Kanehekili is probably some hotter than Loki-B. The 2.2-µm emission images taken in March do not resolve individual hotspots but indicate small areas with temperatures over 600 K somewhere on the disk, as did disk-integrated eclipse measurements at this wavelength in 197910.

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**Imaging C_{60} clusters on a surface using a scanning tunnelling microscope**


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RECENT developments in the synthesis of macroscopic amounts of the C_{60} and C_{70} carbon clusters\(^1\) have made possible spectroscopic\(^2-4\) and vibrational Raman\(^5\) studies of these species. Infrared\(^2\) and vibrational Raman\(^4\) spectra strongly support the proposed ‘soccer ball’ structure for the C_{60} cluster\(^2\), and NMR spectroscopy directly confirms its icosahedral symmetry\(^5\). Here we present images of C_{60} clusters obtained with the scanning tunnelling microscope. These species are roughly spherical and form mobile hexagonal arrays on a Au(111) surface with an intercluster spacing of 11.0 Å. Brighter features evident in the arrays are tentatively identified as C_{70} clusters.

We prepared samples of C_{60} with some C_{70} by resistively heating graphite in 100 torr of helium and subliming these clusters out of the resulting soot onto a gold foil. Mass spectrometry, vibrational Raman and NMR spectroscopy indicate that this method produces clean mixtures of C_{60} and C_{70} clusters\(^4\). The C_{70} NMR spectrum had five lines with intensity ratios 1:2:1:2:1 and chemical shifts 130.8, 145.3, 147.3, 148.1 and 150.6 p.p.m. (internally referenced to the C1 resonance of fully deuterated toluene at 137.5 p.p.m.), in agreement with measurements of Taylor et al.\(^6\). This five-line spectrum supports a proposed elongated ‘soccer ball’ shape for C_{70} (ref. 9). The scanning tunnelling microscopy (STM) measurements were made in a ultra-high-vacuum system which has been described previously\(^9\). The gold foil with the fullerene deposit was transferred into this system through an airlock and left for several hours. A gold(111) single-crystal face was prepared by cycles of sputtering and annealing at 600 °C, as in previous studies\(^11\).

The clusters were transferred onto the crystal by placing the fullerene-coated foil 3 cm away and flashing it to 800 °C. The coated crystal had roughly equal Auger intensities for the 272-eV carbon peak and the 255-eV Au peak, indicating submonolayer fullerene coverage.

The crystal was then transferred (under ultra-high-vacuum conditions) to the STM chamber and slow-scan (1 line per second) images were recorded at a tunnel current of 0.3 nA. Figure 1 shows a series of images recorded a few minutes apart at a tip bias of -0.1 V. In Fig. 1a the slow upward scan sweeps the tip across an ordered array of numerous similar dots which we believe are C_{60} (‘fullerene’) molecules. As the tip moves into the upper region of the frame, the surface steps down and the image becomes unstable. Some of the clusters appear distinctly brighter. In Fig. 1b, the initial tip position was translated away from this unstable region, but now instability and a large asperity appear in the right portion of the frame. A triad of brighter dots in the centre of this frame serves as a landmark. Figure 1c shows that the instabilities reappear if we translate back toward the top of the image. In Fig. 1d instrumental drift has subsided and the hexagonal character of the cluster arrays is apparent. In these arrays we find an intercluster spacing of 11.0±0.5 Å (calibrated using the Au(111) atomic spacing\(^11\)), in reasonable agreement with an estimate obtained by adding the 7.1-A diameter calculated for fullerene\(^12\) to the 3.35-Å layer separation of graphite. The straight rows of clusters are nearly parallel to the

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Au atomic rows, so the array may be commensurate. In Fig. 1d, the vacancy above and to the left of the three bright dots has been filled in, and the bright dot at the lower right has moved.

Our data showed ordered arrays of equivalent mobile species on the Au(111) surface with a spacing consistent with fullerene diameter. It is highly unlikely that the chemically unreactive gold surface can dissociate fullerene, especially if the fragments must be equivalent subunits that form an array with the proper spacing. We therefore conclude that these images represent intact fullerene clusters on the gold surface.

The observation of distinctly brighter dots in the ordered arrays (with an apparent height 2 Å greater than for the C_{60} clusters) demands an explanation. Conceivably the taller features are second-layer fullerene clusters. We would not, however, expect on-top registration for van der Waals' spheres, particularly as the STM tip would probably dislodge molecules in thicker layers. Here the taller features seem to be quite stable, even at the island edges. The most likely explanation is that the brighter dots are C_{70} clusters. The elongation of the C_{60} molecule, together with differences in its electronic structure relative to C_{60} (ref. 18), could easily account for the brighter clusters. Systematic work on samples with varying proportions of C_{60} and C_{70} is required to substantiate this assignment.

It may seem surprising that STM can image such a thick molecule which, by virtue of the 1.5-2.0 eV gap between its highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO)\(^{19}\), might be thought of as insulating. But the C_{60} ionization potential is in the range 7.5-7.72 eV (ref. 19), so the LUMO is not far from the metal Fermi level (\(E_F\)). Furthermore, interactions with the Au surface and neighbouring clusters will shift, split and broaden the cluster orbitals thus giving some density of states at \(E_F\), \(\rho(\mathcal{E}_F)\) (ref. 16). STM is quite sensitive to \(\rho(\mathcal{E}_F)\) on the topmost atoms in the cluster, because the tunnel current varies roughly exponentially as \(\rho(\mathcal{E}_F)\) \(\approx 10^4\) eV/Å, so orbitals with \(z\) the elevation of the top of the cluster in angstroms\(^{20}\). For STM on metal surfaces in vacuum, \(\phi\) is typically 1 eV, so a reduction of \(\rho(\mathcal{E}_F)\) by a factor of 10 relative to the metal only reduces the apparent height of the molecule by 2 Å. Experimentally the molecules are easily detected, as the corrugation in close-packed regions is \(\sim 1.5\) Å and the peak-to-vacancy corrugation, or apparent height, is \(\sim 4\) Å for fullerene and 6 Å for the taller clusters. The same ordered arrays and two heights of peaks are observed at bias voltages between \(-1\) and 1 V, so orbitals with sharply defined energies do not seem to be involved.

A potential difficulty for STM is the fact that the tip, which moves roughly 10 Å above the bare metal surface, withdraws only another 4 Å when traversing the molecules. Given the cluster diameter of \(\sim 10\) Å, the tip approaches them very closely and may cause them to move. Indeed, in this system the observed image noise in areas bordering close-packed islands may indicate the presence of STM-induced \(^{21}\) or thermal motion of molecules not constrained by their neighbours. This behaviour may be relevant to earlier unsuccessful attempts to image these clusters using STM\(^{22}\).

We have not yet observed internal structural description of individual fullerene. These nearly spherical C_{60} clusters, which interact with the surface and each other only through van der Waals' interactions, may occupy rotational states which might smear out the atomic detail. Reducing the temperature or using a more reactive substrate could hinder such rotations. The ease with which images of these carbon clusters can be made suggests that STM studies of these species and their derivatives\(^{23}\) should be interesting and revealing.

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