1,6-Hexanedithiol Monolayers on Au(111): A Multitechnique Structural Study

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Monolayers of 1,6-hexanedithiol \( [HS(CH_2)_6SH] \) deposited on Au(111) from the gas phase were characterized by scanning tunneling microscopy (STM), grazing incidence X-ray diffraction (GIXD), and low-energy atom diffraction (LEAD). Molecular resolution STM images suggest that the molecules lie prone in a striped arrangement with an inter-row spacing of 5 Å. For the films prepared at an elevated temperature, two uniaxial incommensurate phases were found by GIXD. With respect to the surface substrate net, the diffraction patterns of both phases can be described by rectangular \((p \times \sqrt{3})\) nets, where \( p = 4.24 \pm 0.01 \) and \( 4.30 \pm 0.01 \). These values of \( p \) correspond to spacings of \( 12.23 \pm 0.04 \) and \( 12.40 \pm 0.02 \) Å along the nearest-neighbor (NN) direction of the substrate, whereas the spacing along the next-nearest-neighbor direction is 5 Å in both cases. The LEAD patterns can be described by a \( 3 \times 1 \) superlattice with respect to the mesh observed by GIXD. Lattice nonuniformity and angular broadening along the NN direction were observed by GIXD. The structure of the striped phases is consistent with the molecules being fully extended and flat on the surface with their molecular \( C-C-C \) plane parallel to the surface. Using different growth protocols, including liquid-phase deposition, the order of the striped phases was observed to change considerably; however, no evidence of nucleation of other ordered phases was found. Even if denser phases exist, the striped phases may act as effective kinetic traps preventing the transition to other denser phases. The results of both varied growth conditions and performed annealing experiments can be explained by the strong molecular/substrate interaction in the striped phases, which is a consequence of the strong, but not site-specific, interaction of both sulfur atoms with the gold surface.

I. Introduction

Self-assembled monolayers (SAMs) of thiols on gold have been intensively studied because they have shown to be promising materials for microelectronics and biotechnology and can act as a model system for studying more complex membranelike surface layers. In particular, monolayers of \( \omega \)-substituted thiols of the form \( HS-(CH_2)_n-X \) assembled on Au(111) (hereafter denoted as \( HS-C_n-X \)) have been explored for different technological applications.2 Previous work has demonstrated that there are a number of competing interactions present in self-assembled monolayers, including strong headgroup–substrate interaction, endgroup–substrate interaction, chain–chain interaction, and endgroup–endgroup interaction.2 It is the overall balance of these interactions that governs the structure and growth of this class of organic thin films.

The dependence of the structure on the chemical nature of the endgroup has been studied for \( X = OH, COOH, \) \( CH=CH_2, \) and \( CH_3. \) The latter three endgroups resulted in a similar structure,2a,3a whereas \( HS-C_6-X=OH \) was found by STM to adopt a different structure.4 While the interaction of the carboxylic groups is expected to be significant because of the possible dimer formation, the experimental observations for \( HS-C_6-COOH \) suggest that the van der Waals interactions for long alkyl chains may dominate. Consistently, the impact of the olefin groups in \( HS-C_6-CH=CH_2, \) on the system was not found to be large for any chain length.3a
Recent studies have found that throughout its growth process \( HS-C_6-SH \) exhibits a rich variety of structures, before converging to the densely packed high-coverage equilibrium structure in which the molecules pack with their molecular axes inclined by \( \sim 30^\circ \) with respect to the surface normal.\(^5,6\) These findings reflect the complexity of thiol monolayers and the difficulty, shared with Langmuir films, of achieving structural equilibrium. Further understanding of the interplay between the different interactions can be gained through the studies of \( \alpha,\omega \)-dithiol monolayers in which the adsorbates have a thiol functional group at both ends.

Several groups have utilized dithiol monolayers to make molecular devices and multilayers. Brust et al. reported that multilayer structures of copper ions sandwiched between dithiol molecules can be formed by using \( 1,6 \)-hexanediethiol.\(^7\) Ellipsometry measurements suggested that the first monolayer of dithiols adsorbed to the gold substrate with only one of the thiol groups, resulting in concomitant layer-by-layer growth. Other examples are molecular junctions fabricated using 1,4-phenyldithiol as linkers\(^8\) and rigid nanostructures of gold clusters utilizing 1,4-phenyldimethanethiol and 4,4'-biphenyldithiol.\(^9\)

While previous studies have focused on the electronic and photophysical properties of the nanodevices, the structural characterization of these monolayers is incomplete. To exploit the full potential of molecular electronic devices, the structure of the monolayers used to make them needs to be known. In addition, molecular-level information on how the defects affect the quality of the devices is needed. 

There are a few reports, most of them of spectroscopic nature, on the characterization of \( \alpha,\omega \)-dithiol monolayers. Using ellipsometry and X-ray photoelectron spectroscopy (XPS), Tour et al. studied solution-grown monolayers and multilayers on gold films from a series of conjugated organosulfur compounds including 1,4-phenyldithiol and 4,4'-biphenyldithiol.\(^10\) Because multilayer formation of conjugated \( \alpha,\omega \)-dithiols on gold was observed, it was inferred that one thiol end adsorbed to the surface while the other end projected away from it. More recently, Rieley et al. employed XPS to study monolayers of 1,8-octanediethiol on gold.\(^10\) The data were found to be consistent with a structure with an "upright" configuration, which is similar to that suggested by Brust et al.\(^7\) On the basis of STM observations, Nakamura et al.\(^11\) have reported a similar upright configuration for 1,12-dodecanedithiol and a thiophene containing \( \alpha,\omega \)-dithiol monolayers on gold. Using STM and IR spectroscopy, Kobayashi et al.\(^12\) reported a different structure for 1,8-octanediethiol on gold in which the molecules where aligned parallel to the surface.

A recent STM study has obtained information on 1,8-octanediethiol electrochemically adsorbed on Ag(111) from aqueous solution.\(^13\) The adsorption of octanediethiol was found to give rise to close-packed SAMs lying flat on the electrode surface. Two striped structures were suggested with either a head-to-head packing or a "zipperlike" interlocking of sulfur atoms. Both phases were observed to develop domain sizes limited by the number of initial nucleation sites. More importantly, the authors inferred that the octanediethiol molecule adsorbs with only one end attached to the surface while the other forms a silver-mercaptide complex, which is subsequently adsorbed to the surface with the release of a silver ion.

With this situation in mind, we decided to use three complementary structural techniques to study \( 1,6 \)-hexanediethiol \( [HS-C_6-SH] \) adsorbed on Au(111). Grazing incidence X-ray diffraction (GIXD) accurately measures the dimensions, the symmetry, and the average domain size of two-dimensional periodic lattices, and it is sensitive to the in-plane structure of the adsorbates. Low-energy atom diffraction (LEAD) is less precise than GIXD but has a higher surface specificity and is especially sensitive to the corrugation of the outermost surface. Scanning tunneling microscopy (STM) is, in turn, able to study, in real time and real space, the nucleation process occurring on the surface and also to measure with somewhat larger errors the lattice parameters. In particular, STM has the capability of following the structural changes during the growth at surface defects, such as steps and dislocations.

In vacuo gas-phase deposition of \( 1,6 \)-hexanediethiol results in the formation of striped phases (molecules lying down in rows and showing up as stripes in STM images). As will be discussed in detail later, all available experimental evidence strongly suggests that the structure of the striped phases is likely to be composed of fully stretched molecules with their molecular axes parallel to the substrate surface. Because the structure proposed in the present study is different from the upright one commonly assumed in the literature, different growth protocols have been attempted to determine if other denser phases could be assembled in this way. No evidence of other ordered phases was found, and the structural integrity of the striped phases obtained was found to depend strongly on the deposition protocol.

The outline of this paper is as follows. The experimental procedures are explained in section II. In section III, structural studies by STM, GIXD, and LEAD are presented, and a possible structure of the striped phases is proposed. The growth of the monolayers is investigated by GIXD, and the findings are reported in section IV. The results of a GIXD study of thermal behavior of the monolayers are presented in section V, which is followed by the conclusions in section VI.

### II. Experimental Details

#### II.A. Sample Preparation

Au(111) single crystals were used as substrates. Substrates were cleaned by repeated cycles of argon ion sputtering and annealing. The cleanliness of the substrates was signified by the observation of the well-known \( 23 \times \sqrt{3} \) reconstruction.\(^14\) \( 1,6 \)-Hexanediethiol (nominal purity: 96\%) was purchased and used as received. Monolayers were prepared by either the conventional liquid-phase deposition or in vacuo gas-
phase deposition. For the conventional protocol, a clean
substrate was incubated in a millimolar ethanolic solution
of dithiol for 4–16 h. Prior to installation into the
apparatus, the samples were rinsed several times with
solvent and dried with N₂.

For in vacuo gas-phase depositions, the gas manifolds
connected to the thiol source were pumped out for at least
30 min in order to remove any volatile impurities.
Alternatively, several freeze–pump–thaw cycles were
carried out prior to deposition. The purity was checked by
mass spectrometry in the STM studies or by
subsequent mass spectrometry in the GIXD and LEAD
studies. Several in vacuo growth conditions have been
tried: (i) with the substrate at room temperature and
the deposition rate ranging from 0.1 to 10 L/s (1 L = 10⁻⁶
Torr s or 10⁻⁶ Pa s); (ii) with the substrate at 335 K and
the impingement rate kept at ~100 L/s; and (iii) with the
deposition started while the substrate was at 413 K and
continued as the substrate was cooled to 323 K. In this
case the deposition rate ranged from 0.02 to 5 L/s. This
growth protocol is later in the paper referred to as the
“high-temperature growth” protocol.

**II.B. GIXD.** The GIXD experiments were carried out
at the beamline X10B at the National Synchrotron Light
Source. Mounted on a four-circuit diffractometer, a compact
vacuum chamber suitable for in situ studies of organic
thin films (P ~ 10⁻⁸ Torr or 10⁻⁶ Pa) was used for the
measurements. The details of the experimental setup have
been described elsewhere.³ All of the measurements were
taken at a wavelength of 1.130 Å. The resolution was set
by the detector slits. For most of the in-plane diffraction
measurements, Δq_i ranged from 0.006 to 0.01 Å⁻¹ while
Δq_j was 0.015 Å⁻¹.

The scattering geometry used is shown in Figure 1a.
The total momentum transfer q and the perpendicular
momentum transfer q_perp are

\[ q = k_i - k_j \quad \text{with} \quad |q| = 2|k_i| \sin(\theta/2) \]  
(1)

and

\[ |q_perp| = |k_i|(\sin \alpha + \sin \beta) \]  
(2)

where \( k_i \) and \( \alpha \) are respectively the incident wavevector
and angle while \( k_j \) and \( \beta \) respectively the outgoing
wavevector and angle. In addition to the lattice spacing,
determined from the position of the Bragg peaks, the
monolayer domain size \( L_D \) can be determined from their
width. After the instrumental broadening is taken into
account, \( L_D \) can be estimated from the observed width,
Δq_θ, using

\[ L_D \approx 2\pi/\Delta q_\theta \]  
(3)

Noted that if other types of broadening (e.g., inhomogeneous
broadening) exist, they have to be included in the
calculation of the peak width.

A rectangular coordinate system is used with unit
vectors \( \mathbf{a} \) and \( \mathbf{b} \) with |\( \mathbf{a} \)| = 4.997 Å and |\( \mathbf{b} \)| = 8.66 Å. The
surface normal is defined as the \( \{111\} \) direction of the
face-centered-cubic structure of gold. The third unit vector
\( \mathbf{c} \) corresponds to the \( \{111\} \) Bragg point and |\( \mathbf{c} \)| = 2.356 Å.
The reciprocal lattice units are \( \mathbf{a}^* \), \( \mathbf{b}^* \), and \( \mathbf{c}^* \) where |\( \mathbf{a}^* \)|
= 1.257 Å⁻¹, |\( \mathbf{b}^* \)| = 0.726 Å⁻¹, and |\( \mathbf{c}^* \)| = 2.667 Å⁻¹. Diffraction
peaks are indexed as \( (h, k, l) \) and \( \mathbf{q} \) equals \( h\mathbf{a}^* + k\mathbf{b}^* + l\mathbf{c}^* \). Because all of the in-plane measurements
were taken at \( \theta = 0.1 \), the diffraction peaks \( (h, k, l) \)
are abbreviated as \( (h, k) \). The crystallographic directions used
in this paper are indexed according to the face-centered-
cubic structure of gold.

**II.C. LEAD.** The LEAD studies were performed in a
bolometric detection-based atom diffractometer that al-
lows in situ sample preparation. The extensive use of
cryogenics provides for ultrahigh-vacuum conditions in
all of the studies presented here. The diffractometer has
been described in detail elsewhere.³ The monoenergetic
beam of helium (wavenumber \( k_i = |k_i| = 5.27 \text{ Å}⁻¹ \) and
monochromacity \( dk_i/k_i \approx 2\% \)) was produced by supersonic
expansion, and the incident angle \( \theta_i \) was fixed at ~60°
from the surface normal. During the measurements the
substrate temperature was held at 120 K. The angular
distribution of the scattered helium intensity was mea-
sured by the detector for different values of \( \theta_i \). As shown in
Figure 1b, an “in-plane” scattering configuration was
used, i.e., the incident beam, the surface normal \( \mathbf{n} \), and
the detector were all in the same plane. Thus, the parallel
momentum transfer \( |q| \) can be determined by

\[ |q| = |k_i|(|\sin \theta_i - \sin \theta_j|) \]  
(4)

The resolution was limited by the size of the detector. For
the present studies, the \( q \) resolution is ~0.06 Å⁻¹ while the
azimuthal resolution is only ~1 Å⁻¹ because of the
large height-to-width ratio of the slit collimating the
primary beam so chosen to increase detection sensitivity.

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II.D. STM. STM studies were performed in an ultrahigh-vacuum (P \approx 10^{-10} \text{Torr or } 10^{-8} \text{ Pa}) multichamber apparatus, which is equipped with custom-built STM and commercial XPS, quadrupole mass spectrometer probe, and a rapid-entry load-lock. The experimental design allows in situ preparation and chemical characterization of organic thin films. STM tips were prepared by electrochemically etching single-crystal tungsten (111) wires. All images were acquired at room temperature in a high-impedance (\approx 10 \Omega) and constant-current (10^{-3} \text{ pA}) mode. The systematic error in measuring the lattice spacings is \approx 10%.

III. Structural Studies

III.A. Lattice Dimension and Symmetry. (1) STM Results. Monolayers of in vacuo gas-phase grown HS–C$_6$–SH were characterized by STM. Figure 2a is a molecular resolution constant-current STM image of monolayers that were prepared at room temperature. The image shows rows of bright features along three symmetry-equivalent directions. The cross-sectional profiles along the rows (Figure 2b) and across the rows (Figure 2c) show that the lattice dimensions respectively are 13.3 \pm 1.3 and 5.0 \pm 0.5 Å. The interaxial angle \gamma is \approx 95°. With respect to the substrate surface net, the overlayers can be described as a rectangular (9 \times \sqrt{3}) net with two molecules per unit mesh. Schematics of the unit mesh in real space and an idealized reciprocal space map are shown in parts a and b of Figure 3, respectively. Also, the STM data shown in Figure 2a indicate that the bright features are not all aligned in the (110) direction (i.e., the NN direction). Some features are centered along the long axis of the unit mesh (labeled by c), and some are beside the long axis (labeled by s). The implication of this “meandering” of the bright features will be discussed in section III.C.

(2) GIIXD Results. Because of its high resolution and accuracy, GIIXD was employed to measure the lattice spacing, the symmetry, and the interfacial structure. The monolayers were prepared following the “high-temperature growth” protocol, and the observed reciprocal space map is shown schematically in Figure 4a where the diffraction features of the monolayer and the substrate are represented by solid and open circles, respectively. Parts a and b of Figure 5 show (solid squares) the radial scans through the first-order diffraction peaks (0, 0.7) and (1, 0), respectively. The solid lines in Figure 5a,b are least-squares fits to the data assuming a linear background and a Gaussian peak shape. While the intensities will be discussed later, the peak positions are at $q_1 = 0.507 \pm 0.001$ and 1.260 \pm 0.001 Å$^{-1}$. Phase coexistence was also found in the monolayers. Although the coexistence is not clearly shown in the (0, 0.7) Bragg peak, which just shifts upon annealing (solid and open squares in Figure 5a), two diffraction features are clearly present before annealing at the (0, 2.8) Bragg peak (the fourth-order peak; solid squares in Figure 5c). Upon annealing to \approx 423 K, an irreversible transition occurs and only one phase remains (open squares in Figure 5a,c). This phase of larger $q_1$ (i.e., higher density in real space) is denoted as IC1. From a least-squares fit of the data, the first-order diffraction peak along the (110) direction of IC1 is located at 0.514 \pm 0.001 Å$^{-1}$. The observed reciprocal space map implies that the surface net of IC1 is primitive rectangular and the unit mesh dimensions are 12.23 \pm 0.04 and 4.99 \pm 0.02 Å. The statistical error bars represent 3 times the standard deviations resulting from the fit to the data, while systematic errors are estimated to be on the order of 0.01 Å.

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Figure 2. (a) 220 Å \times 220 Å constant-current STM image showing molecular resolution of the striped phase of HS–C$_6$–SH prepared by in vacuo gas-phase deposition. Note that the bright features are “meandering” as explained in the text. (b) "Height" profile along the line trace AB. The dotted lines mark a spacing of 13.3 Å. (c) "Height" profile along the line trace EF. The dotted lines mark a spacing of 5.0 Å. The profiles imply a rectangular surface net with unit mesh dimensions of 13.3 \pm 1.3 and 5.0 \pm 0.5 Å. The meaning of the labels c and s is explained in the text.
The length 12.23 Å equals 4.24\(d\) while the length 4.99 Å is equal to \(x3\)\(d\), where \(d\) = 2.884 Å is the surface lattice parameter of gold. Thus, the suggested net is uniaxially incommensurate with the underlying substrate and the incommensurability is along the \(\langle 11 \bar{2}\rangle\) direction. The real-space dimensions are illustrated in Figure 4b. The real-space long dimension of the unit mesh of the phase, which shows a peak at a smaller value of \(q\) \(| \langle IC2 \rangle\), is 12.40 (0.02 Å or 4.30\(d\). Note that the characteristic lengths of the suggested surface nets are different from those inferred from the STM images but are within the experimental error of the latter. In Figure 5a,c, the arrows mark the expected peak positions for a rectangular \(9 \times \sqrt{3}\) surface net. Notice that the lattice spacing of IC1 is shorter than that of IC2.
by 1.3%, and the difference between IC1 [(4.24 x 2) x \sqrt{3}] and the 9 x \sqrt{3} net is 5.8%.

(3) LEAD Results. The apparent discrepancy between the lattice dimensions measured by STM and GIXD motivated the LEAD study. Monolayers were prepared following the “high-temperature growth” protocol. Parts a–c of Figure 6 show diffraction patterns along the (110) (NN) direction. There is a progression of diffraction peaks with a small separation in momentum transfer space. As indicated by the solid lines, most of the diffraction peaks can be explained by a periodicity of 0.169 Å⁻¹.

Figure 7 shows a diffraction spectrum along the (112) (NNN) direction. The positions of the more intense diffraction peaks show a periodicity of 1.257 Å⁻¹ as indicated by the vertical solid lines. The spacings of 0.169 ± 0.003 and 1.257 ± 0.007 Å⁻¹ correspond to a rectangular surface net with the unit mesh dimensions of 37.18 ± 0.65 and 4.995 ± 0.028 Å. Figure 8 is an idealized reciprocal space map of this net where different symbols represent the three symmetry-equivalent domains. The solid line rectangle outlines the suggested unit mesh.

Evidently, a number of peaks in the diffraction spectrum along the (112) direction (see Figure 7) cannot be explained by a spacing of 1.257 Å⁻¹. This can be understood by considering the instrumental resolution function, which is drawn as a dashed rectangle in Figure 8. As explained in detail elsewhere, the detector was designed to give an optimal \( q \) resolution and signal-to-noise level at the cost of sacrificing the azimuthal resolution. Therefore, off-azimuthal diffraction peaks that fall in the “detection window” are also detected during the scans. The observed \( q \) of these peaks is related to their true \( q \) as

\[
q_{\text{observed}} = q_{\text{true}} \cos \xi
\]

where \( \xi \) is the angle by which the peak is removed from the chosen azimuth. According to this equation, the positions of these peaks are marked as the dashed lines in the spectrum along the (112) direction (Figure 7). The expected off-azimuthal peak positions show an excellent agreement with the data.

The poor azimuthal resolution also explains the intensity distribution of the diffraction spectrum along the (110) direction. The intensities of diffraction features around integral multiples of -1.5 Å⁻¹ were found to be higher than the intensities of other diffraction peaks. By inspecting the reciprocal space map as suggested by the surface net (Figure 8), the high intensities around these positions can be attributed to the large number of diffraction peaks which fall in the detection window. For instance, let us compare the integrated intensity around -1.5 Å⁻¹ with the integrated intensity around -2.03 Å⁻¹, where only one diffraction peak contributes to the observed peak (see the arrows in the middle part of the figure). In Figure 6, the diffraction feature around \( q_{\text{obs}} \) of -4.3 Å⁻¹ appears to be narrower than the diffraction features around \( q_{\text{obs}} \) of -1.5 and -2.9 Å⁻¹. This observation provides further support for the suggested surface net. Because diffraction patterns are superpositions of all of the diffraction peaks that are inside the detection window, the shapes of the diffraction features at -1.5, -2.9, and -4.3 Å⁻¹ can be correlated very well with the distribution of the off-azimuthal peaks at these locations (see Figure 8).

In summary, the helium diffraction data yield a rectangular surface net with unit mesh parameters of 37.18 ± 0.65 and 4.995 ± 0.028 Å, which, within the experimental uncertainties, in one direction is very close to 3 times the spacing of the IC1 and IC2 phases while in

Figure 6. LEAD data taken along the (110) direction. The solid lines mark a spacing of 0.169 Å⁻¹. The asterisk in part c marks a spurious peak which was not observed in all other samples.
the other direction coincides with \( \sqrt{3} \) times the gold atomic spacing. Figure 9 is a schematic of the real-space unit mesh. A uniaxially incommensurate unit mesh was therefore measured by both diffraction probes. Although the absence of long-range periodicity in the STM images suggests a commensurate net, the results are consistent with the diffraction results within the mutual experimental uncertainties. However, it is surprising that no Moiré patterns were observed, since they are expected for incommensurate overlayers. The reasons why the superlattice observed by atom diffraction was not observed by GIXD will be discussed later.

III.B. Peak-Width Analysis of the X-ray Data. High-resolution GIXD can reveal information on the average domain size and the presence of certain types of lattice disorder. When the radial scans through the (0, 0.7) Bragg peak and its higher order peaks are examined, the measurements show that the characteristic peak widths for the monolayers, \( \Delta q_{\text{intrinsic}} \), depend on the in-plane momentum transfer, \( q \), in both the IC1 and IC2 phases.

\[
(\Delta q_{\text{total, observed}} \cos \chi)^2 = (\Delta q_{\text{resolution}} \cos \chi)^2 + \Delta q_{\text{intrinsic}}^2
\]

where \( \Delta q_{\text{total, observed}} \) is the total (observed) peak width, \( \Delta q_{\text{resolution}} \) is the width determined by the slit setting, and the term \( \cos \chi \) is a correction factor for the scattering geometry used in the experiments. Besides the finite domain size, different types of lattice disorder may also contribute to \( \Delta q_{\text{intrinsic}} \). According to the type of disorder, \( \Delta q_{\text{intrinsic}} \) can depend in different ways on \( q \).

In the following we will adopt simple models described in ref 17 because a more sophisticated treatment would require an even more extended data set. On the basis of these models, we will distinguish between two types of disorder. In the first type, the lattice spacing exhibits fluctuations about the ideal spacing, with the long-range order being preserved. Then \( \Delta q_{\text{intrinsic}} \) can be written as

\[
\Delta q_{\text{intrinsic}} = a + b q_i
\]

In the disorder of the second type, the spacing between two successive lattice points varies about an average value but the spacing between two given lattice points is not related to the neighboring spacing, i.e., the fluctuations are random. This leads to

\[
\Delta q_{\text{intrinsic}} = a + b q_i^2
\]

In both cases, the term \( a \) is the width due to the finite average domain size of the monolayers.

Parts a and c of Figure 5 show the radial scans through the (0, 0.7) and (0, 2.8) peaks of the C1 phase, respectively (see the open squares). Clearly, the fourth-order peak is wider than the first-order peak. Least-squares fits of the data with a Gaussian function and a linear background give a peak width of \( 0.0100 \pm 0.0002 \) Å\(^{-1} \) for the (0, 0.7) peak and a peak width of \( 0.014 \pm 0.001 \) Å\(^{-1} \) for the (0, 2.8) peak. Notice that, in the present study, the instrumental broadening \( \Delta q_{\text{resolution}} \) increases with the angle \( \chi \) which is \( \approx 27^\circ \) for the first-order peak and \( \approx 7^\circ \) for the fourth-order peak.\(^{18} \) The \( \Delta q_{\text{resolution}} \) contributions to the total width are thus 0.0099 and 0.0064 Å\(^{-1} \) for the first-order peak and the fourth-order peak, respectively. Because only two data points are available and the (0, 0.7) peak is instrument-resolution-limited, the functional form of \( \Delta q_{\text{intrinsic}} \) cannot be determined. The domain size is found to be substrate-limited at \( \approx 2000 \) Å (see below), and it implies \( a \) to be 0.003 Å\(^{-1} \) (2σ/2000 Å). Assuming a linear \( q \) dependence (i.e., eq 7 to be valid), the mean variation of the lattice parameter can be estimated to be more than 0.4%.

Similarly, the lattice nonuniformity in the IC2 phase was investigated. The radial widths of four peaks were measured. After accounting for the phase coexistence with IC1 and the instrumental effect, \( \Delta q_{\text{intrinsic}} \), the characteristic peak width for the monolayers, were found. The \( \Delta q_{\text{intrinsic}} \) values of the four diffraction peaks of IC2 are plotted as a function of \( q \) (Figure 10a) and \( q^2 \) (Figure 10b). The lines are the least-squares fits to the data. Apparently, it is difficult to discern which model fits better, i.e., whether the disorder is of the first type or the second type. In general, one may even expect that both types of lattice disorder exist in the monolayers and the peak width will behave like:

\[
\Delta q_{\text{intrinsic}} = a + bq^2 + cq^4
\]

While it is hard to determine the coefficients \( a, b, \) and \( c \) with high accuracy, the data put some constraints on the contribution of each type of disorder. From the fit to the data, the linear broadening (disorder of the first type) indicates \( a \) to be close to zero, whereas the quadratic broadening (disorder of the second type) suggests \( a \) to be 0.004 Å\(^{-1} \). Given that the substrate has an average domain size of \( \approx 2000 \) Å, a should be close to 0.003 Å\(^{-1} \). Therefore, the quadratic broadening seems to be dominant, yet it is possible that both types of disorder coexist.

Finally, GIXD was also employed to examine possible orientational disorder in the monolayers. The observed azimuthal width of the diffraction peak \( \Delta q_{\text{azimuthal}} \), the width due to the instrumental resolution \( \Delta q_{\text{inst}} \), and the width due to a finite domain size \( \Delta q_a \) are related by

\[
\Delta q_{\text{azimuthal}}^2 = \Delta q_{\text{inst}}^2 + a^2 + \Delta q_a^2
\]

For the present experimental configuration, \( \Delta q_{\text{azimuthal}} \) is measured by an azimuthal scan through a diffraction feature. Let \( d_\phi \) be the width of the diffraction peak in azimuthal angle \( \phi \); then

\[
d_\phi \approx \Delta q_{\text{azimuthal}}/q_i
\]

Because \( \Delta q_{\text{inst}} \) is very small and \( a \) is independent of \( q_i \), \( \Delta q_{\text{azimuthal}} \) is independent of \( q_i \) if there is no angular broadening. With \( \Delta q_{\text{azimuthal}} \) constant, \( d_\phi \) is thus inversely proportional to \( q_i \). Parts a and b of Figure 11 show the azimuthal scans through the (0, 0.7) peak and the fourth-order peak of the IC1 phase, respectively. The solid lines are the least-squares fits to the data. The data clearly show that the azimuthal width of these peaks is not limited by the domain size and imply that IC1 is orientationally disordered. Notice that, from the (0, 0.7) peak width, the domain size of IC1 is derived to be \( \approx 2200 \) Å, i.e., substrate-limited. Likewise, the IC2 phase is found to be orientationally disordered as well (data not shown).

To conclude, the data suggest a rather complex scenario with respect to the type of disorder present in the system. While more data are needed for a more detailed analysis, the data available appear to be consistent with the STM observation of “meandering” bright features. This is most likely a consequence of the competing chemisorption of the sulphur groups on inequivalent sites of the substrate.

III.C. Proposed Structural Model for the Striped Phases. The lattice dimensions determined by STM, GIXD, and LEAD provide detailed structural information for the striped phases (IC1 and IC2). With respect to the substrate surface net, the striped phases can be described as a primitive rectangular \( p \times \sqrt{3} \) net. The striped phases are similar to the striped phase of monolayers of 1,8-octanediethiol on gold observed in an earlier STM study.\(^{12} \) It is quite possible that the adsorbates are lying flat on...
alkanes on Pt(111). While earlier work already reported to be 5 Å, within experimental uncertainties, which is equal short axis of the unit mesh (along the present study. All of the techniques employed find the periodicity along the HS substrate. Thus, the unit mesh dimensions observed for hydrocarbon chains and/or the corrugation of the gold that the dithiol molecules form a striped phase, the monolayers of substrate. Similar packing has also been observed in the plane containing the carbon atoms parallel to the sub-

Figure 11. Data (solid squares) and the Lorentzian fits (solid lines) of the azimuthal scans through (a) the (0, 0.7) peak and (b) the (0, 2.8) peak of the 1C1 phase.

the surface with the alkyl chains fully extended and the plane containing the carbon atoms parallel to the substrate. Similar packing has also been observed in the monolayers of n-alkanethiols on Au(111) and n-alkanes on Pt(111). While earlier work already reported that the dithiol molecules form a striped phase, the periodicity along the (112) direction (i.e., the spacing across the rows) is, for the first time, accurately measured in the present study. All of the techniques employed find the short axis of the unit mesh along the (112) direction to be 5 Å, within experimental uncertainties, which is equal to \(\sqrt{3}\) times the lattice spacing of Au(111). Notice that this is also the lattice spacing of the commensurate short periodicity of the striped phases of HS\(- C_n - CH_3\), which can be attributed to the packing constraints of the hydrocarbon chains and/or the corrugation of the gold substrate. Thus, the unit mesh dimensions observed for HS\(- C_9 - SH\) indicate that the molecules are lying flat on the surface with their alkyl chains fully extended.

The long side of the unit mesh observed by GIXD (12.40 and 12.23 Å) is close to the length of a dithiol fragment suitably increased by a \(\sim 2.9\) Å S–S distance. Note that while a typical S–S bond is on the order of 2 Å, the van der Waals distance between two S atoms is comprised between 3.6 and 4 Å. The molecular resolution STM image shown in Figure 2a shows that the bright features are elongated. The shape of those bright features is different from some of those from the periodic phase of HS–CH=CH_2, which were found to be round. If the bright features are associated with sulfur atoms, the elongated features observed in the striped phases of HS–C6–SH suggest that there are two sulfur atoms in close proximity of each other.

The long side of these features is parallel to the (110) direction, i.e., the NN direction of the substrate (Figure 2a). Assuming that the features are the sulfur pairs belonging to neighboring molecules, this observation suggests that the sulfur pairs align in rows parallel to the (110) direction. However, as mentioned earlier, while following this general alignment direction, the features may slightly alter their positions perpendicularly to this direction (see features labeled c and s in the figure). Because the domain size in the monolayer at which the image were acquired is rather small, further studies are needed to clarify if this “meandering” of the sulfur pairs is intrinsic to this system or if it disappears after the films have been annealed. Note that the weak intensity of the (1,0) peak seen in X-ray diffraction may be related to the sulfur pairs’ meandering and the consequent presence of partially destructive interference.

Next, it is useful to consider the information provided by the LEAD data which have shown the existence of a \(3 \times 1\) superlattice of the surface net measured by GIXD. While X-rays primarily scatter from the sulfur atoms and STM tunneling takes place possibly through the sulfur atoms, helium is scattered by the corrugation of the surface potential, which is governed by a number of factors including the vertical position of the adsorbates. Two different scenarios can be invoked to explain the existence of the \(3 \times 1\) superlattice detected in the atom diffraction experiments. They are (i) the existence of a height modulation of the substrate and (ii) a height modulation of the dithiol molecules which may be due to a modulation in the binding of the sulfur pairs.

As gold has a much higher electron density than sulfur and carbon, the scattered X-rays are very sensitive to even a slight modulation of the surface gold atoms. As no evidence of modulation of the surface gold atoms was detected in the GIXD experiments, height modulation of the surface gold atoms is unlikely to be present. Because the lattice nets are incommensurate, some molecular height modulation has to be expected. Recently, the formation of sulfur pairs in equilibrium monolayers of HS–CH3 suggested by Fenter et al. has been confirmed using standing wave X-ray photoelectron spectroscopy. In that work, two different positions have been suggested for the sulfur atoms of each pair. It is

(22) The C–C and C–S bond angles are 112°. The C–C bond length is 1.541 Å, and the C–S bond length is 1.81 Å. Without both of the S-bonded H atoms; the dithiol fragment is estimated to be 9.4 Å.
(24) Theoretical studies suggest that sulfur atoms should be imaged as a protrusion on metal surfaces [Lang, N. D. Phys. Rev. Lett. 1986, 56, 1164]. Also, both experimental and theoretical investigations have shown that sulfur atoms are imaged as bright features on Ni(100) surfaces [Partridge, A.; Tatlock, G. J.; Leibsle, F. M.; Flipse, C. F. J.; Hörmandinger, G.; Pendry, J. B. Phys. Rev. B 1993, 48, 8267].
IV. Growth Conditions

The structures of the monolayers grown in vacuo in this study disagree with previous studies, where it is assumed that the hydrocarbon backbones in dithiol monolayers are positioned more or less vertically. In an attempt to clarify this issue, different preparation methodologies to create the “standing-up” phase have been explored. Previous studies in our laboratory and others have demonstrated that by prolonged dosing HS−C₆−CH₃ undergoes a phase transition from the striped phase, in which molecules are lying flat on the substrate, to the “standing-up” phase, in which the hydrocarbon backbones are aligned along the surface normal. The same mechanism was also observed in monolayers of functionalized thiols, such as HS−C₆−OH. In situ GIXD and STM studies were performed to investigate if other phases appear in HS−C₆−SH monolayers after extended dosing. Both STM and GIXD results are in good agreement (data not shown) and suggest that a prolonged dosing is not conducive to the making of other denser phases.

In previous work it has been shown that the self-assembly process of HS−C₁₀−CH₃ monolayers occurs by a low-density striped phase through a complex, intermediate density stage, where little long-range order is present, to the final standing up c(4 × 2) phase which grows when the coverage goes above a critical value. One may then pose the question as to whether the density of domain boundaries of a particular striped phase affects the subsequent growth behavior of that layer. STM was employed to monitor if other phases nucleate for monolayers with a high density of domain boundaries by extended dosing at room temperature. No evidence of other phases was observed. The observations suggest that for the growth conditions of this experiment the density of the domain boundaries is not the limiting factor in producing kinetic traps on the road to the formation of the standing-up phase.

Another growth protocol, in which the deposition was carried out at an elevated temperature and with a very high impingement rate of dithiol, has been tried. The conjecture was that the mobility of the adsorbate increases at higher substrate temperatures allowing the monolayer to escape from possible kinetic traps and thereby succeeding in increasing the surface coverage. In situ GIXD studies were conducted for a monolayer which was grown at 335 K with an impingement rate of 100 L/s. Upon an exposure of ~7 × 10¹³ L, no evidence of ordered phases other than the incommensurate phase c(1 × 2) was observed. The results indicate that at 335 K the mobility is still too low to attain a surface coverage higher than that in the striped phase.

Traditionally, monolayers of thiols on gold have been prepared by immersing a clean gold substrate into a millimolar solution of thiol for hours at room temperature. This raises the question as to whether other phases can be obtained by liquid-phase deposition. Specifically, does the solvent play a crucial role in the intermediate stages of the growth? [Poirier, G. E. Langmuir 1999, 15, 1167].
of this observation is not clear as the existence of an upright phase is not the only hypothesis that could explain the measurements. However, the observation is reported for sake of completeness and objectivity. Finally, we note that in the XPS measurements we detected no features in the region where the oxygen 1s lines normally appear, indicating the absence of sulfonate species.

The growth attempts in this study only cover a limited range of growth parameters (substrate temperature, impingement rate, with or without solvent); nevertheless, the results are very suggestive that the dithiol molecules at the very least do not readily form other denser phases. The difference in growth behavior between HS–C₆–SH and HS–C₉–CH₃ can be rationalized by the energetics involved. In the case of the alkanethiol, the thermal energy is large enough to cause fluctuation in the local density, which allows the incoming molecules to disrupt the ordered striped phase. A coverage-induced phase transition then occurs, and the standing-up phase appears. In the case of α,ω-alkanedithiol, the measurements indicate that the local free energy minimum represented by the striped phase is very deep indeed.

While no denser phases were observed in the monolayers prepared by different growth methods, the quality of the striped phase was found to vary significantly. The monolayers that were grown in vacuo gas-phase deposition at room temperature (Figure 2a) are of poorer quality as compared to those grown at elevated temperatures for which the domain size of the samples is substrate-limited as discussed in section III.B. These results can be tentatively attributed to the low diffusion rate of molecules at room temperature, which is expected from a system with a strong molecule–substrate interaction. This strong molecule–substrate interaction is also manifested by finding a similar domain size on the terraces and around surface heterogeneities (data not shown).

V. Thermal Properties

Apart from the structure and growth behavior, other monolayer properties such as thermal stability may be controlled by the interactions involved. GIXD was used to measure the thermal behavior of HS–C₆–SH monolayers prepared using the “high-temperature growth” protocol. Figure 14a shows a selected set of azimuthal scans through the (0, 0.7) Bragg peak at different substrate temperatures. The radial resolution δq was 0.02 Å⁻¹, and the striped phases IC₁ and IC₂ were not resolvable in this case. The integrated intensities measured are proportional to the total number of molecular monolayer properties such as thermal stability may be measured by the interactions involved. GIXD was used to measure the thermal behavior of HS–C₆–SH monolayers prepared using the “high-temperature growth” protocol. Figure 14a shows a selected set of azimuthal scans through the (0, 0.7) Bragg peak at different substrate temperatures. The radial resolution δq was 0.02 Å⁻¹, and the striped phases IC₁ and IC₂ were not resolvable in this case. The integrated intensities measured are proportional to the total number of molecular monolayer properties such as thermal stability may be argued in light of the energetic differences....
HS and SH are more thermally robust than the striped phase of SH ordered phase is a consequence of the strong molecule dithiol, desorbs at the same. The higher thermal stability of the HS of the bulk 1,6-hexanedithiol and temperature for the striped phase of HS the peaks under part a are plotted as a function of the substrate unit for viewing convenience. (b) The integrated intensities of the one at 463 K) have been translated vertically by one arbitrary different substrate temperatures as indicated. All scans (except f
resulting in larger domains while the IC2 IC1 and IC2. Upon annealing, the molecules diffuse, in this way a standing-up monolayer can be obtained. These results show that the striped phases of HS–C6–SH. IC1 and IC2. Upon annealing, the molecules diffuse, resulting in larger domains while the IC2 → IC1 phase transition takes place. Upon further annealing, IC1 desorbs from the substrate.

These results show that the striped phases of HS–C6–SH are more thermally robust than the striped phase of HS–C9–CH3 which melts at ~373 K and then, as the dithiol, desorbs at ~473 K. Note that the boiling points of the bulk 1,6-hexanediithiol and n-decanethiol are almost the same. The higher thermal stability of the HS–C6–SH ordered phase is a consequence of the strong molecule–substrate interaction in the striped phases and is also consistent with the proposed model in which the molecular backbone is parallel to the surface, and for each molecule, there are two sulfur atoms interacting with the underlying substrate.

VI. Conclusions

The results of a multitechnique study of 1,6-hexanediithiol [HS(CH2)6SH] monolayers on Au(111) are presented. Three structural probes were employed: namely, STM, GIXD, and LEAD. Molecular resolution STM images of vapor-deposited monolayers show rows of bright features and suggest that the molecules lie prone on the surface and arrange in stripes. Two uniaxial incommensurate striped phases IC1 and IC2 were found by high-resolution GIXD. With respect to the surface substrate net, the diffraction patterns of both phases are consistent with a rectangular ($p \times \sqrt{3}$) net, where $p$ is 4.24 in IC1 and 4.30 in IC2. The incommensurability is along the NN direction of the substrate, and the lattice spacing along this direction is 12.23 ± 0.04 Å for IC1 and 12.40 ± 0.02 Å for IC2. Within the experimental uncertainties, the lattice dimensions determined by GIXD and STM are in agreement. LEAD measurements are consistent with a rectangular net with unit mesh dimensions of 37.18 ± 0.06 Å × 4.995 ± 0.028 Å, which can be also described as a $3 \times 1$ superlattice with respect to the surface net observed by GIXD. Lattice nonuniformity along the NN direction of the substrate (in the direction of the chains) and angular broadening of the diffraction features were found by GIXD. From these measurements, it is suggested that the structure of the striped phases is comprised of fully extended molecules lying flat on the surface, with the plane containing the C atoms parallel to the surface. The incommensurability along the NN direction and the existence of the $3 \times 1$ superlattice imply the presence of multiple S/Au binding sites. In the proposed structure, the sulfur endgroups of adjacent molecules are separated by ~2.9 Å and are parallel to the NN direction.

Different growth protocols, including the conventional liquid-phase deposition, have been attempted, but no evidence was obtained for the nucleation of ordered phases other than the striped phases described here. The apparent lack of crystalline high-density standing-up phases in HS–C6–SH can be explained by a strong molecule–substrate interaction in the striped phases, which implies a large energy barrier for the diffusion of both the adsorbates and the domain boundaries. If other phases exist, the observations suggest that the striped phases are effective kinetic traps preventing the phase transition to other denser phases. Finally, the strong molecule–substrate interaction is manifested in the observed thermal behavior of the striped phases.

Recent results by Liu and collaborators31 have shown that the growth of n-octadecanethiol monolayers can be made to proceed much faster and with better results by “nanografting"" the molecules into a preexisting n-octadecanethiol monolayer using the tip of an atomic force microscope. It would therefore be interesting to nanograft dithiol molecules into an alkanethiol monolayer to see if in this way a standing-up monolayer can be obtained.

(30) The boiling point of 1,6-hexanediithiol is 391 K/15 mm from the Aldrich Chemical Co. Catalog. The boiling point of n-decanethiol is 399 K/19 mm from the CRC Handbook of Physics and Chemistry; Lide, D. R. Ed.; CRC Press: Boca Raton, FL, 1992.
The present study highlights the importance of understanding the relationship between the energetics, structure, and growth of thiol monolayers on gold. Also, the results obtained for hexanedithiol on gold show once more the importance of the striped phase in the growth process of these films. The present study demonstrates that a combination of STM, GIXD, and LEAD may be necessary to obtain reliable structural information at a molecular level, without which the utilization of these molecules in the assembly or fabrication of nanomaterials and/or nanodevices may be complicated by the possibly false underlying premises.

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