Structure of Monolayers Formed by Coadsorption of Two n-Alkanethiols of Different Chain Lengths on Gold and Its Relation to Wetting

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We have characterized and correlated the structure and wetting properties of self-assembled monolayers (SAMs) on gold derived from different two mixtures of n-alkanethiols (C₁₂SH and C₁₆SH; C₁₆SH and C₂₂SH); in each of these SAMs, one thiol had a perdeuterated alkyl chain and one a perprotonated chain to allow the two n-alkanethiols to be distinguished. The SAM by polarized infrared external reflection spectroscopy is compatible with values obtained by diffraction methods.‡12-13 Monolayers composed of more than one component would lead to domains in which each component has a near-surface region of the shorter chains may, for appropriate concentrations of coadsorbates, contain densities of gauche conformations comparable to those in liquid phases (Figure 1).33

In this study, we examined mixed SAMs comprising two n-alkanethiols having different chain lengths; one had a perdeuterated alkyl chain to allow the two components to be distinguished spectroscopically. Since wetting is due to short-range interactions (<5 Å),10-12 we hypothesized that the structure of these mixed SAMs could be thought of as being a disordered hydrocarbon layer supported on a crystalline or liquid-crystalline underlayer.27-29 That is, we expect that the polymethylene chains projecting beyond the region of the shorter chains may, for appropriate concentrations of coadsorbates, contain densities of gauche conformations comparable to those in liquid phases (Figure 1).33

In earlier studies, we have shown that the alkyl chains of thiolate monolayers supported on gold formed a canted phase which can be described well by an average orientation of a single, all-trans chain having a tilt of ~27° relative to the surface normal.11b,20 The value determined by infrared spectroscopy is compatible with values obtained by diffraction methods.12,13,34 This value originates from the formation of a commensurate lattice of thiophanes on the gold surface, (√3×√3)R30°, that yields a greater area per thiophane (21.4 Å²)34 than the cross-sectional area of an alkyl chain (18.4 Å²).35 As a result, the chains tilt to maximize their van der Waals interaction.16 For the mixed systems studied here, macroscopic phase separation of the two components would lead to domains in which each component has the average structure found in pure SAMs (Figure 1a). Smaller degrees of mixing should yield SAMs where the tails of the longer n-alkanethiolate must adopt more complex conformations (and perhaps tilt further away from the surface normal) to achieve van der Waals interaction, and the chain groups in the near-surface region should be less ordered than the alkyl chains are in pure SAMs (Figure 1b,c). In general, we expect that the highest gauche densities per long chain will exist in those phases in which the value of its mole fraction is lowest and, most likely, highest gauche densities per long chain will exist in those phases in which the value of its mole fraction is lowest and, most likely.
The d+ and d- modes of the spectral references. Mode assignments are given in Table of n-CpD25SH and n-C22Ha5SH, respectively, were used as the references. Monocomponent SAMs derived from pure samples were obtained; monocomponent SAMs derived from end of the alkyl chain, \parallel = parallel, \perp = perpendicular. MacPhail et al. \cite{ref18} have reported a frequency for 2915 cm\textsuperscript{-1} for an orthorhombic crystalline sample of n-C\textsubscript{12}H\textsubscript{25} at T < 10 K.

Results

SAMs Prepared from Ethanolic Solutions Containing Mixtures of C\textsubscript{12}D\textsubscript{3}SH and C\textsubscript{22}H\textsubscript{4}SH. Figure 2 displays data from ellipsometry, X-ray photoelectron spectroscopy (XPS), and wetting by water and hexadecane (HD) for SAMs derived from ethanolic solutions containing mixtures of n-C\textsubscript{12}H\textsubscript{25}SH and n-C\textsubscript{22}H\textsubscript{45}SH (total thiol concentration = 1 mM); similar data on related systems have been discussed in detail elsewhere. \cite{ref29,ref28} The vertical line in Figure 2 marks the solution composition required to form a SAM having an equimolar composition of the two adsorbates. This SAM exhibited the lowest receding value (\(\theta_{\text{r}}(\text{H}_2\text{O})\)) and the highest hysteresis (\(\Delta \cos \theta = \cos \theta_{\text{r}} - \cos \theta_{\text{a}}\)) in the contact angles of water.

SAMs derived from these solutions were characterized by infrared spectroscopy (Figure 3). Spectra of the C-H and C-D stretching regions (2800-3050 and 2000-2300 cm\textsuperscript{-1}, respectively) were obtained; monocomponent SAMs derived from pure samples of n-C\textsubscript{12}D\textsubscript{3}SH and n-C\textsubscript{22}H\textsubscript{45}SH, respectively, were used as the spectral references. Mode assignments are given in Table 1. \cite{ref17,ref29,ref28,ref37,ref38}

The position and width of peaks in the C-H stretching region change with surface composition. The d* and d-r modes exhibit asymmetry (beyond the asymmetry in d- due to lower frequencies and become narrower. For SAMs of in-plane CH\textsubscript{2} groups located at either end of the alkyl chain, \parallel = parallel, \perp = perpendicular. MacPhail et al. \cite{ref38} have reported a frequency for 2915 cm\textsuperscript{-1} for an orthorhombic crystalline sample of n-C\textsubscript{12}H\textsubscript{25} at T < 10 K.
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The C-H stretching region. (b) The C-D stretching region. Total thiol and d- modes, assigned as the symmetric and asymmetric CH2 (alkanes) is observed in these spectra. The line shapes of the d+ and d- modes in these spectra also require that the spectral contributions of the methylene of the gauche segments be, to a similar degree, "abnormally intense". The key insight here is that the structure is not conserved. The "disordered" phases have chain architectures that are distinctly different from the canted phases formed by single-component SAMs. The projections of the transition dipole moments of the C-H stretching vibrations along the surface normal direction are likely to be significantly different (and presumably larger) for many relevant long chain concentrations.

The methyl modes (r+ and r-) in the C-H region exhibit no change in frequency with composition. The ratio of the intensities of these two modes (r+/r-) in the spectra is approximately the same in the spectra, although our ability to measure the intensity of these modes is limited. This trend is consistent with the belief that the thermal population of chain-end gauche conformers yields a disordered methyl surface projection at room temperature. It is unclear from the spectra if the weighting of the r+ and r- modes changes with composition.

In Figure 3b, the peaks associated with the C-D stretching modes change in intensity with changes in the ratios of concentrations of thiols in the contacting solutions (and, thus, of the mixed SAMs): the widths and positions of the peaks, however, remain relatively unchanged. Given the quality of our data and the complexity of interpreting C-D stretching frequencies, we are hesitant to analyze these data further. The spectra do, however, also provide a useful comparison with spectra given later in the paper.

Figure 4 displays the wetting and IR data of the mixed SAMs relative to their surface composition. The lowest values of θ(0) (HD) and θ(0) (HD) occur over a range of surface compositions where the longer thiolate is the minor component. Over this range of compositions, the positions of the d+ and d- modes suggest that the portions of the longer component contacting the solvent are characterized by considerable disorder. In SAMs that contain higher surface compositions of the longer thiolate, this component is present in a simpler form and the wetting properties of the mixed SAMs by HD resemble those of a low-energy methyl surface.

As we have noted above, the integrated intensities of the d+ and d- modes cannot be directly related to the composition of the longer alkanethiolate in the SAM (Figure 4c): at intermediate compositions, the intensity is greater than expected for a simple statistical weighting of the integrated intensity of a pure C22 SAM. The intensities of these modes are affected by, among other factors, the details of the mode couplings as well as the orientation of the polymethylene chain. The simplest case arises for orientations of an all-trans chain for which cants of the main chain axis further away from the surface normal become more intense. The deviations in d+ and d- evidenced in Figure 4 suggest that the average orientation of the chain in the mixed SAMs is greater than the tilt for a pure SAM (~27° from the surface normal); however,
Figure 4. Wetting properties and C-H stretching mode data of SAMs on gold derived from ethanolic solutions containing mixtures of CD₇(CH₂)₇SH and CH₃(CH₂)₁₁SH. The infrared data are for the longer n-alkanethiolates in the mixed SAMs, namely, those elements of the SAM derived from CH₃(CH₂)₁₁SH. The compositions of the SAMs were determined by XPS (see Experimental Section); we estimate the error in surface compositions to be ±5% of the x-axis. The dashed lines in the lowest panel represent the intensities expected for these modes if the longer alkanethiolate were present in large-domain-size islands (Figure 1a). AU = arbitrary units.

Figure 5. Ellipsometry, XPS, and wetting data for SAMs on gold derived from ethanolic solutions containing mixtures of HS(CH₂)₇CH₃ and HS(CH₂)₁₁SH. Data are plotted as in Figure 2. Thicknesses determined from XPS were calibrated to the ellipsometric thickness of a SAM derived from HS(CH₂)₇CH₃ (see Experimental Section).

Figure 6. PIFERS spectra of SAMs on gold derived from ethanolic solutions containing mixtures of HS(CH₂)₇CH₃ and HS(CH₂)₁₁SH. Total thiol concentration = [C₇₃] + [C₁₁₃] = 1 mM. The ratios of the two thiols on the surface ([C₇₃]SAM + [C₁₁₃]SAM) were estimated from XPS data; we estimate the error in surface compositions to be ±5% in the value of [C₇₃]SAM ([C₇₃]SAM + [C₁₁₃]SAM). The dashed and dotted vertical lines correspond to methyl and methylene vibrational modes, respectively. Mode assignments are given in Table I.

because the changes in intensity occur concomitant with changes in the line width and peak frequency of these modes (Figure 3a), the data do not allow a simple picture to be drawn for the longer alkanethiolate. What is clear from these data is that the detailed phase behaviors of the longer chains are responding to the short-chain constituents of the SAM as if the presence of the shorter alkanethiolate constitutes a significant and varying perturbation of the local molecular environment of the longer alkanethiolate.

SAMs Prepared from Ethanolic Solutions Containing Mixtures of n-C₇₃H₂₇SH and n-C₁₁₃D₁₇SH. Figure 5 displays the ellipsometric, XPS, and wetting data on SAMs derived from ethanolic solutions containing mixtures of C₇₃H₂₇SH and C₁₁₃D₁₇SH (total thiol concentration = 1 mM). These experiments are complementary to those in the preceding section. Here, the shorter chains are protonated and thus are the more easily analyzed by PIFERS. The data are qualitatively similar to those in Figure 2. The differences in wetting between the mixed and pure SAMs are less dramatic than those data summarized in Figure 2c and probably reflect the smaller difference between the chain lengths of the two alkanethiols used (difference = 6 vs 10 carbon atoms).³⁵

SAMs derived from these solutions were characterized by infrared spectroscopy (Figure 6). The C-H spectra were obtained using a SAM derived from C₁₁₃D₁₇SH as a reference; the low intensity of C-D stretching modes relative to C-H modes precluded obtaining spectra of the longer alkanethiolate having useful intensity.⁴¹

The IR spectra in Figure 6 exhibit trends opposite those observed in Figure 3a. The methylene C-H stretching modes (d⁺ and d⁻) exhibit little change in position or peak width with decreasing surface composition of the alkanethiol; the methyl C-H stretching modes (r⁺ and r⁻) change both in width and position. The lack of weighted intensity for the d⁺ and d⁻ modes at higher frequencies (or even large band asymmetries) suggests that the density of gauche conformers in the shorter component of the
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Figure 7. Wetting properties and C–H stretching mode data of SAMs on gold derived from ethanolic solutions containing mixtures of CH3(CH2)1SH and CH3(CH2)2SH. The infrared data are for the shorter n-alkanethiolate in the mixed SAMs, namely, those elements of the SAM derived from CH3(CH2)1SH. The compositions of the SAMs were determined by XPS (see Experimental Section); we estimate the error in surface compositions to be ±5% of the x-axis. The dashed lines in the lowest panel are linear least-squares analyses of the data and illustrate that the intensity of methylene modes for the shorter thiolate are linearly related to its composition in the SAM. AU = arbitrary units.

Discussion

Kinetic and Thermodynamic Considerations Important to Multicomponent SAMs. It is well-known that n-alkanes of dissimilar chain lengths (lengths comparable to those employed here) do not form thermodynamically stable solid solutions.61–63 Phase separation occurs very rapidly in samples quenched from the melt and involves, in all likelihood, a combination of molecular transport mechanisms. (These mechanisms find only incomplete analogies in SAMs, since longitudinal and lateral diffusion of chains along growing lamellae occur.) We are inclined to believe that the equilibrium state of mixed SAMs comprised of adsorbates of dissimilar chain lengths would consist of phase-separated domains. This conclusion follows from the likely inability of the favorable entropy of dilution to compensate effectively the increased van der Waals interactions experienced by chains in a crystalline domain. We can estimate the enthalpic contribution that drives this phase separation. A first-order approximation, for a phase transition of the sort shown in Figure 1a, would be equal to the weight-differentially the group constituent binding enthalpies of the CH3 groups according to the local characteristics of the environment in which they exist (either crystalline or disordered domains). A reasonable estimate would be to equate this enthalpic difference to a group constituent heat of fusion. As judged from the melting points of the n-alkanes, this enthalpy would be of the order of ~0.4 kcal/mol of CH3.4,4 For the case of a SAM composed of C12 and C21 chains, an approximate driving force of 4 kcal/mol would be available to offset a composition dependent entropy of mixing. We therefore conclude that the structures of the mixed SAMs discussed below are in part metastable, kinetically trapped structures: that is, they are mixtures of components that would tend to phase separate, given appropriate conditions. There exists increasing experimental evidence from our and others’ laboratories (which we do not present in detail here) that suggests that thiols adsorbed on gold do yield approximately kinetic structures.27,28 It is only for the simplest adsorbates (e.g., CH3SSCH3 adsorbed on Au(111) in UHV) or for very long exposures that the true thermodynamic end points of the assembly are likely to be approached. For single-component SAMs the main point of concern is the time that is required to achieve rigorously the limiting coverage defined by the preferred (\sqrt{3}×3\sqrt{3})R30° habit of the sulfur ligands on the (111) surface. The significant steric repulsions expected for a chain trying to penetrate a high coverage surface suggests that the approach to equilibrium kinetics could be very slow. Strong pinning of adsorbates at surface defects is also likely to affect the rate at which equilibrium is reached. The mixed SAMs implicitly raise another interesting and complex notion which has only been casually addressed in earlier work. It is clear that the adsorption process is dynamic; in the mixed SAM is low and remains constant over all compositions of the SAM. The integrated intensities of the methylene C–H stretching modes (d* and d−) in this experiment exhibit a linear relation with the concentration of the shorter component in the mixed SAM (Figure 7c). This linear relation suggests that the orientation of this component, when in a mixed SAM, is the same as in a pure SAM. The IR data, taken in total, determine that the dilution by the longer chained component does not constitute a major perturbation of the local molecular environment of the shorter alkanethiolate. As in Figure 4a, the lowest values of θ (HD) and θ (HD) in the C12/C4 system occur when the longer alkanethiolate is a minor component of the mixed SAM (Figure 7a). The spectral characteristics of the shorter thiolate in these SAMs are similar to those of a pure SAM although the position of the r+ and r− modes are ~2 cm−1 lower than they are in pure SAMs. At lower surface concentrations of the shorter alkanethiolate, the methyl C–H stretching modes broaden and shift further in position (Figures 6 and 7b). Stole and Porter have observed similar shifts and broadening (albeit of greater magnitude) in the methyl C–H stretching frequencies of a methyl-terminated SAM on gold upon transforming it from contact with air to a condensed phase (in their case, water, methanol, and CCl4).42 These differences between the spectra of mixed and pure SAMs suggest that the methyl groups of the shorter component in these mixed SAMs are not present at the monolayer/air interface, but are instead in contact with a condensed phase,42 presumably the hydrocarbon chains of the longer alkanethiol. The detailed structural arguments advanced by Stole and Porter involved perturbations beyond those of simple dielectric screening (i.e., media) effects on band intensities (although these are no doubt important). The consequence of solvent contact in their view was to enhance surface disorder. This interpretation is doubtful, given the now well-accepted view that the ambient surface of these structures presents considerable conformational disorder at room temperature even in the absence of solvent.11,13,40,44,45 Indeed, what is more striking about these latter trends in the data is that the “band shifts” of the asymmetric methyl C–H stretching modes more correctly indicate, at least to a considerable degree, a reweighting of the relative intensities of the r+ and r− contributors.46 These trends are indistinguishable in both magnitude and direction from those occurring on cooling a single-component SAM below 200 K.11,13 To this extent then, the coadsorbed perdeuterated long chain does not constitute significant perturbation of the local molecular environment felt by the shorter chained constituent. This perturbation is one, at least for a significant range of mole fractions in the SAM, that the thermal population of chain end gauche conformers in the shorter chain are quenched by the presence of the longer constituent, a notion consistent with its being “buried” in the matrix formed by the latter. That is, in brief, the chain ends of the shorter chains become more ordered by contact with the disordered, projecting ends of the longer chains than they would be in a pure single-component phase. Similarly, in the work of Stole and Porter, the observed effects may reflect increased order rather than decreased order for chain ends in contact with solvent (relative to the order characterizing the monolayer/air interface).
presence of excess adsorbate, molecules both adsorb and do not order.

The latter exchange process is incompletely understood at present but is now thought to involve kinetic exchange processes that somehow couple to defects in the overlay. It is unlikely that mixed SAMs could express quantitative structural differences that depend very strongly on the exact details of the sample history.

Structure of the Mixed SAM. The combined weight of the IR data suggests that the shorter alkanethiolate is present in the mixed SAMs, over all surface compositions, in a highly orientated form similar to but differing in some subtle regards from that of a pure SAM (Figures 3b and 6), and that portions of the longer alkanethiolate contain gauche conformers when present in the mixed SAMs (Figure 3a).6 The strong influence of the coadsorbate on the phase state of the longer chains implicitly argues against the occurrence of a macroscopic phase segregation of the constituents under the preparative conditions employed here. The hydrocarbon chain of the shorter thiolate in the mixed SAMs is like that in a pure C₆H₁₃SAM on gold; namely, it is trans-extended (Figure 7b) and tilted ~27° from the surface normal (Figure 7c). At concentrations of short chains in the SAMs less than ~10%, the perturbation of the methyl mode intensities suggests that the population of chain-end gauche conformers in the shorter chain constituent is in fact lower than that found in the pure SAM at this temperature. The longer thiolate contains high densities of gauche conformers that appear to be localized in specific regions of the adsorbate molecule; their density is a function of the concentration of the longer thiolate in the mixed SAMs (Figure 4b). Given the structure of the shorter alkanethiolate in the mixed SAMs, we infer that the region of the longer chain near the gold must be structurally similar to that of a pure SAM (i.e., trans-extended and tilted ~27° from the surface normal) over all surface concentrations, and that the tail (i.e., the portion of the longer alkanethiolate that extends beyond the end of the shorter alkanethiolate) must contain most of the gauche conformers. The band shifts present in the d⁻ and d⁺ modes of crystalline (2918 and 2850 cm⁻¹, respectively) and liquidlike regions (~2928 and ~2858 cm⁻¹, respectively) of the longer thiolate, when present as a minor component of the mixed SAM (Figure 3a; [% C₂H₅SAM < 10%]), suggest that much of the tail region of the longer thiolate may be in contact with the methyl surface of the shorter alkanethiolate. The spectral changes observed in the r⁺ and r⁻ modes in Figure 6 support the presence of an interaction between the methyl groups of the shorter adsorbate and a condensed phase, presumably the disordered tail of the longer thiolate. At higher surface concentrations of the longer thiolate, the line shapes of the d⁻ and d⁺ modes suggest that the longer thiolate contains a much lower density of gauche conformers than that described immediately above. This decrease may be due to the presence of small clusters of longer thiolates themselves providing for domains of largely trans-extended longer chains; these clusters need not be oriented ~27° from the surface normal as would be obtained in a pure SAM. Figure 8 schematically summarizes these structural conclusions.

The general structure that emerges from the IR spectra is one containing a disordered hydrocarbon layer supported on a hydrocarbon underlayer whose structure is similar to and contains comparable molecular orientations as that found in a pure SAM. The IR and wetting data of the mixed SAMs are incompatible with the formation of macroscopic domains each containing one component (Figure 1a); the mixed SAMs are not composed of a crystalline and liquidlike phases are both present in the longer thiolate of our mixed SAMs.

Relation of Structure to Wetting. The wetting properties of the mixed and pure SAMs are different and must be related to the different structural features present in the mixed SAMs. The interfaces of the mixed SAMs expose a mixture of CH₃ and CH₂ groups, the latter contributing to a higher energy surface, and contain a higher density of gauche conformers than do the pure SAMs; these conformers are concentrated near the monolayer–air (liquid) interface. Wetting is a measure of the interfacial free energy, and differences in it between surfaces must reflect differences in the enthalpic and/or entropic interactions of a liquid with the interfaces.

Figures 3 and 6 allow comparison between a macroscopic property of the mixed SAMs, wetting, and microscopic details of their structure. The minimum values of θ(HD) occur when the longer alkanethiolate is the minor component of the SAM. Over the range of surface compositions where θ(HD) is lowest, the IR spectra consist of the surface of the mixed SAM to be composed of disordered hydrocarbon chains supported on a crystalline-like hydrocarbon underlayer (Figure 8). For SAMs composed of ~20–60% of the longer thiolate, cos θ(HD) and cos θ(HD) change little. The small change in cos θ(HD) and cos θ(HD) suggests that the interfacial free energies of these surfaces (γSV) are similar even though the amount of disorder present in the mixed SAMs, as manifested in the positions of the d⁻ and d⁺ modes, is different. Bain et al. have shown that wetting by HD can be sensitive to the 3-A region near the surface. The small similarity in wetting by HD suggests that this 3-A region near the surface of the mixed SAMs is similar in composition and degree of disorder over this range of surface compositions. The shifts in the d⁻ and d⁺ modes thus must represent the different amounts of crystalline hydrocarbon domains that are present below this 3-A region.

The contact angles of water (θW and θW) are relatively insensitive to the disorder present in the SAMs. The surface of the mixed SAMs should be composed of various amounts of CH₃ and CH₂ groups. The equilibrium (or mean) contact angles of water on the different structural features present in the mixed SAMs. The interfaces of the mixed SAMs expose a mixture of CH₃ and CH₂ groups, the latter contributing to a higher energy surface, and contain a higher density of gauche conformers than do the pure SAMs; these conformers are concentrated near the monolayer–air (liquid) interface. Wetting is a measure of the interfacial free energy, and differences in it between surfaces must reflect differences in the enthalpic and/or entropic interactions of a liquid with the interfaces.

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in the absence of other effects, would be slightly lower on the mixed SAMs than on the pure SAMs (±6-10%). The only change observed in the contact angles of water is a decrease in θ, in the C₁₄/C₁₂₂ system (≈6°). As this system also exhibits an increased hysteresis in the contact angles of water (∆cosθ = cosθ₁ - cosθ₂), it is difficult to assess whether any of the changes (or lack of changes) in the contact angles of water are controlled by changes in the composition,27 morphology,65 or degree of disorder present in the SAMs.

Relation between Hysteresis and Structure. While macroscopic theories exist to explain contact angle hysteresis,61,65,67 no microscopic theory is currently available to explain the contact angle hysteresis that have been observed in mixed monolayer systems of the type characterized here. Although we realize that the structure of the mixed SAMs may be different when contacted with a condensed liquid from that determined (by IR) for the SAM in contact with air (Figures 8 and 9a), some similarities no doubt do exist. Our discussion here is intended to summarize the various data concerning hysteresis in related systems in the literature.68-72 and compare them with the structural elements determined in this paper and in related systems.

As in previous studies,28,29 we observed increased hysteresis in the contact angles of water on mixed SAMs on gold derived from C₁₂SH and C₁₄SH relative to the pure SAMs (Figure 2). Significant increases in hysteresis have been observed28 with mixed SAMs composed of adsorbates having smaller differences in the chain lengths of the two adsorbates and are consistent with our observation here that little change in hysteresis occurred in the C₁₂/C₁₈ system (Figure 5). Bain et al.28 have observed similar mixed systems that, while the hystereses in the contact angles of n-alkanes (HD and decane) do not change with surface composition, the hysteresis in the contact angles of bicyclohexyl and water do. The maximum hystereses of water and bicyclohexyl consistently occurred when the SAMs had an equimolar composition of the longer and shorter thiolates.

From the data presented here, SAMs having equimolar compositions consist of shorter thiolates that are present in an oriented, largely trans-extended form, and of longer thiolates that contain gauche conformers. The apparent weight per chain density of gauche conformers in the longer thiolate, as manifested in the positions of the peak maxima of the d- and d* modes, is lower in the SAMs that contain a 1:1 composition of the two thiolates than in those that contain lower concentrations of the longer thiolate (Figure 4b). In Figure 4c, ratios of the two components ranging from 2:1 to 1:2, the largest deviations in the intensities of these modes from those expected for thiolates in a crystalline form oriented 27° from the surface normal are observed. As the 27° cant angle results from dense packing of the hydrocarbon chains on the (3×3)/3R30° lattice on gold,26 these deviations suggest that many of the hydrocarbon chains at the surface are not as densely packed. The range of surface compositions where a maximum amount of hydrocarbon chains is not present in a densely packed fashion occurs over the same range where the hystereses in the contact angles of water and bicyclohexyl are at a maximum. This correspondence suggests that hysteresis between disimilar phases may be determined by amounts of molecular-level structural heterogeneity and possibly roughness.66 Although the exact details of the structure of mixed SAMs when contacting these liquids are unclear, we expect that, based on the immiscibility of alkane and water, contact with water would only modestly perturb the structure of the mixed SAMs from that determined for them by IR (Figure 9a).

Over this intermediate range of surface compositions (2:1 to 1:2), the hystereses in the contact angles of HD and decane exhibit little change (Figures 2 and 5, and refs 28 and 29) and we can only presume that the similarities between the structure and chemical nature of the SAM and of the contacting liquids are responsible. The exact structure of the mixed SAMs, when in contact with HD or decane, is one probably restructured to some degree by these liquids (Figure 9b). The exterior region of the SAMs, when in contact with these liquids, may more bear little resemblance to that pictured in Figure 8. The internal, crystalline region of the SAM, however, should be affected little by solvent and is probably similar in structure to Figure 8.

In a limited way, these mixed SAMs are analogous to binary mixtures of shorter and longer n-alkanes. In these systems, Snyder et al. have observed that the conformational disorder of the longer n-alkane reached a maximum for mixtures of ~1:1 composition.48 The level of conformational disorder in the longer n-alkane was greater when the difference in chain length between the two alkanes was greater.49 The observations noted here that the hystereses in the contact angles of water and bicyclohexyl reach maxima when the surface composition is ~1:1.48 and that the observed hystereses are lower when the difference between the chain lengths of the adsorbates is reduced strongly suggest that the hystereses in the contact angles of water and bicyclohexyl are sensitive to the presence of conformational disorder. By analogy, hexadecane and decane are not sensitive to this disorder.

Conclusions

Mixed monolayers obtained by adsorption of alkanethiols having different chain lengths on gold contain specific regions of order and disorder; the region closest to the gold contains a much lower density of gauche conformers than the region closer to the monolayer/air interface. The highest weight per chain density of gauche conformers is observed in the longer alkanethiolate when its surface composition is low; the greatest amount of disorder in the longer alkanethiolate is observed in SAMs having approximately equimolar composition. The interplay between entropic and enthalpic contributions to the interfacial free energy of the SAMs is manifested differently in the wetting properties of water and hexadecane; the contact angles of hexadecane are more sensitive to the density of conformational disorder present in the longer n-alkanethiolate than those of water; the hysteresis in (HD, O) is more sensitive to the amount of conformational disorder present in the longer n-alkanethiolate than the hysteresis in (θ, HD).

Experimental Section

Materials. Dodecane- and docosanethiol were available from previous studies.23 1-Bromooctadecane-d₄ and 1-bromo-dodecaned-₂₄ were obtained from Cambridge Isotopes Laboratories (98% D); diacetic acid was from Aldrich. Dioxygen was removed from absolute ETOH (Quantum Chemical Corp., Tuscola, IL) by bubbling N₂ through it prior to use. Si(100) wafers (test grade; 50 and 100 mm), Au (99.999%) and Cr (99.997%) were obtained from Silicon Sense (Nashua, NH), Materials Research Corp. (Orangeburg, NY), and Aesar, respectively.

n-C₄₋₁₂SH and n-C₁₀₋₁₈SH. The thiols were prepared from the corresponding bromides by displacement with sodium thioacetate and subsequent acid methanolysis.23 The alkyl thioacetates and thiols were purified by column chromatography (silica gel) and exhibited the same values of 2R as authentic samples not containing deuteriums. SAMs derived from adsorption of these thiols had indistinguishable properties by wetting, ellipsometry, and XPS from those prepared from pure samples of C₄₋₁₂SH and C₁₀₋₁₈SH, respectively.

Preparation and Characterization of Monolayers. Evaporated films of Au (2000 Å) were prepared onto Cr-primed Si(100) wafers as previously described.25 SAMs were formed by exposing the Au films to ethanolic solutions containing mixtures of the two alkanethiols for 24-48 h. The SAMs were removed from solution and rinsed with ETOH prior to characterization. Ellipsometric
Acknowledgment. We are grateful to Colin D. Bain and John P. Folkers for valuable discussions and suggestions.

Registry No. C2H2, 112-55-0; C2SH, 2880-00-9; C22SH, 1773-83-3; Au, 7440-57-5; hexadecane, 544-76-3.

References and Notes
(1) This research was supported in part by the Office of Naval Research and by the National Science Foundation (Grant CHE-88-12794). XPS spectra were obtained using instrumental facilities purchased under the DARPA/URIP program and maintained by the Harvard University Materials Research Laboratory. R.G.N. acknowledges support from the Department of Energy through the UIUC-MRL (DEFG02-91ER-45439).


(41) From Figures 3b and 6, the intensity of the C-D stretching modes (v' and v") of a perdeuterated C16 SAM is ~40% the intensity of the C-H stretching modes for a perprotonated C16 SAM. This reduced intensity limits our ability to assess reliably the magnitude of band shifts that occur at low mole fractions of the perdeuterated C16 component given the baseline artifacts that we observe in this spectral region.


(44) Diffraction studies have shown that the density of gauche conformers at the methyl/vapor/vacuum interface, even at much lower temperatures (~35 K), is greater than the density present in the bulk of a SAM at room temperature: domain sizes for the methyl surfaces of various SAMs are 26-44 Å at ~35 K; the domain size for the SAM is 60-70 Å at ~300 K. 12,14


(46) The reweighting of the r1 and r5 contributors is most evident in the IR spectra in Figure 6 for mixed SAMs derived from solutions having ratios of the two components ([C16H31]/[C16H30]) of 4 and 6. For these mixed SAMs, the r5 spectral envelopes are significantly broader than for those of pure SAMs and their "peaks" occur at wavenumbers lower than that for the pure SAM. These features can be readily explained by an increase in the relative intensity of the r5 contributor (see ref 47). The result of this reweighting is that the r5 spectral envelope in these spectra appear broader than that of a pure SAM (Table I).

(47) As the thermal population of gauche conformers are quenched in an a-alkane, the concentration of gauche conformations in the ends of the longer a-alkane monotonically decreased as its concentration in the mixture was increased; the concentration of gauche conformers in the ends of the shorter a-alkane remained constant as its concentration in the mixture decreased. 16


