Fatty Acid Langmuir Films on Liquid Mercury: X-ray and Surface Tension Studies

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The structure and phase behavior of liquid-mercury-supported molecular films of fatty acids ($\text{CH}_n(\text{CH}_2)_m\text{COOH}$, denoted $\text{C}_{n\text{C}_{m\text{H}}}$) were studied for molecular lengths $7 \leq n \leq 24$, by surfacetensitometry and X-ray methods. Two qualitatively different film structures were found, depending on coverage. For high coverage, the film consists of a monolayer of roughly surface-normal molecules, showing a pressure-dependent sequence of structures similar, though not identical, to that of the corresponding water-supported Langmuir films. At low coverage, phases consisting of surface-parallel molecules are found, not observed on the aqueous subphases employed to date. In this range, a two-dimensional (2D) gas followed by a single and, for $14 \leq n \leq 24$, also by a double layer of surface-parallel molecules is found as coverage is increased. Depending on chain length, the flat-living phases have a crystalline 2D-ordered, a smectic-like 1D-ordered, or a disordered in-plane structure consisting of molecular dimers. The structure and thermodynamics of the films are discussed.

I. Introduction

Langmuir monolayers of fatty acids on water served as the model of choice for studies of Langmuir films since the first quantitative measurements of Agnes Pockels on stearic acid over a century ago. Naturally, they have been also among the first three Langmuir monolayers studies by X-ray methods, published contemporaneously in 1987. Langmuir films were, and are, extensively studied in physics as a model for two-dimensional matter, in biology as a model for the cell membrane, and in chemistry as templates for oriented growth of crystals from solutions, etc. They are being currently studied also as means for constructing few-molecule electronic devices.

The vast majority of all experimental studies of Langmuir films over the last century employed a variety of macroscopic methods. Surface tensiometry, the most prominent of these methods, provides a description of the surface thermodynamics through surface pressure ($\pi$)–molecular area (A) isotherms. With the advent of synchrotron-based liquid surface X-ray scattering techniques, it became possible to study the structure of these films in situ with Ångström scale resolution. Over the last two decades, these X-ray methods have been employed in studies of many Langmuir films. The structure of fatty acid Langmuir films on aqueous subphases, in particular, was investigated in detail as a function of chain length and temperature. Peterson and co-workers were able to show that as far as the phase diagram is concerned, a decrease in temperature is equivalent to an increase in chain length in these films. Once this effect is accounted for quantitatively, the phase diagrams of fatty acids of all chain lengths were found to merge into one "universal" phase diagram. The rich phase diagram which emerged from these experimental studies supported a detailed theoretical description and interpretation within a Landau-type mean-field theory by Kaganer and Loginov. Despite this great progress, new phases, such as a phase with an intermediate tilt between nearest- and next-nearest neighbor directions, and novel effects, like a shift in the phase boundaries upon the tuning of the aqueous subphase, have been observed.


of the interactions between the headgroups,\textsuperscript{16} are still being discovered. Further details of the plethora of results obtained for fatty acids on aqueous solutions can be found in several excellent reviews published over the past decade.\textsuperscript{3,4,8,11}

Until recently, all X-ray studies of Langmuir films were carried out on aqueous subphases. On such subphases, the strong hydrophobic repulsion of the fatty acid’s tail from the surface invariably orients the molecules roughly along, or slightly tilted from, the surface normal. To the best of our knowledge, the only nonaqueous subphase employed in X-ray studies of a Langmuir film is formamide,\textsuperscript{11,17,18} which is also polar and induces the same surface-normal molecular orientation. To investigate the role of the molecules’ interaction with the subphase in the determination of the Langmuir film’s structure, we have undertaken to study the structure of Langmuir films of organic molecules on mercury. Mercury does not induce a hydrophobic repulsion of the chains from the surface but rather provides an attractive interaction with the chains. It also does not provide a hydration shell for ionic headgroups. It has several technical advantages over water. The high surface energy of mercury, ~500 mJ/m\textsuperscript{2}, as compared to that of water, 72 mJ/m\textsuperscript{2}, enhances the spreadability of films and allows spreading of molecules that are not readily, or not at all, spreadable on water. The practically zero solubilities of all organic compounds, and most inorganic ones, in mercury allows investigation of Langmuir films of compounds which cannot be studied on water due to their high water solubility. As shown below, the higher surface tension of mercury also allows extension of the X-ray reflectivity measurements to 4-fold larger scattering vectors, yielding a commensurately higher resolution as compared to those achievable on water. The important differences in both tail and head-group interactions with the subphase enables investigations over a considerably broader scope of properties of Langmuir films than those available in Langmuir films on aqueous subphases only. This should provide for a deeper insight into the role of the subphase in forming the structure and phase diagram of Langmuir films.

Knowing the surface structure of bare mercury is a prerequisite for any study of mercury-supported Langmuir films. We have determined this structure with subångström resolution a few years ago,\textsuperscript{19} using surface X-ray techniques. This was followed by a study of dense self-assembled alkanethiol monolayers\textsuperscript{20} and multilayers\textsuperscript{21} on the surface of mercury. Very recently we presented results for stearic acid on mercury,\textsuperscript{22} and a systematic study of the chain length variation of the structure and phase diagram of Langmuir films of alkanes on mercury,\textsuperscript{23} for both fatty acids and alkanes, the attractive interaction of the alkyl tail with the subphase induces, at low coverages, new phases comprising surface-parallel molecules, not observed on water. For alkanes only surface-parallel phases (of up to three layers) are found for all chain lengths. In contrast, stearic acid on mercury exhibits phases with surface parallel molecules at low coverages and phases with surface-normal molecules at high coverages.\textsuperscript{24} The surface-normal phases have structures very similar to those of stearic acid on water. A schematic view of the real-space structure of the various surface-normal and surface-parallel phases found for C18OOH for different coverages is shown in Figure 1. Two surface-parallel phases, a single layer (SL) and a double layer (DL), are shown as well as two surface normal phases, one tilted (Ov) and one untitled (LS).

We present here a systematic study of the chain length dependence of the structure and phase diagram of fatty acids on mercury for a range of chain lengths and surface coverages, as well as first results on their temperature dependence.

Finally, self-assembled layers (SAMs) of organics on solid substrates, mostly alkanethiols on gold, have been investigated extensively by X-ray, and other, methods.\textsuperscript{24–27} and surface-parallel phases were found at certain coverage ranges. These SAMs should be, however, clearly distinguished from Langmuir films on a liquid substrate. While on a solid substrate the SAMs structure is largely determined by its epitaxy to the structure of the underlying solid, on a liquid mercury subphase the film’s molecules are not confined to lattice sites. Thus, varying the coverage is considerably simpler, and more accurate, on a liquid surface than on a solid one. More importantly, in the most extensively studied SAMs by far, alkanethiols on gold, the crystalline structure of the gold surface was found to determine the SAMs structure by epitaxy, both in the surface-normal\textsuperscript{26} and the surface-parallel\textsuperscript{27} phases. Since no static corrugation potential exists for a liquid surface like mercury, such epitaxy of the organic mono-

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure1.png}
\caption{Schematic real space model of a Langmuir film of stearic acid on mercury. With increasing coverage stearic acid shows successively a single (SL) and a double (DL) layer of lying-down molecules, a tilted phase of standing-up molecules (Ov) and an untitled phase of standing-up molecules (LS).}
\end{figure}

layer to the structure of the subphase does not occur, resulting in less constraints for the emergence of different types of lateral order in the film.

II. Experiment

The experimental methods are discussed in detail in a previous publication. Thus, only a summary is given here.

A. The Langmuir Trough. A specially designed Langmuir trough, suitable for simultaneous film balance and X-ray studies, was used. It consisted of a $175 \times 65 \times 3.5$ mm$^3$ KeF trough, enclosed in a hermetically sealed aluminum box. The box was filled with either helium (in X-ray measurements) or nitrogen (for surface tension measurements) to minimize mercury oxidation and air scattering, and was also equipped with an automatic shutter, which was opened for counting, and kept closed during spectrometer movement.

B. Materials. Triple distilled, 99.999% mercury was purchased from Merck. Fatty acids were purchased from Fluka or Aldrich with molarities in the range of $(3-8) \times 10^{-4}$ using HPLC grade, 99.9% pure chloroform.

C. Measurement Methods. 1. Surface Pressure–Molecular Area Isotherms. The surface pressure, $\pi = \sigma_0 - \sigma$, the difference between the surface tension of the bare ($\sigma_0$) and the film-covered ($\sigma$) mercury varies with the surface coverage, $A$, the area of the molecular representation. Features in the $\pi$–$A$ isotherm curve (e.g., kinks, plateaus, etc.) reveal the occurrence of structural changes in the film as the coverage $A$ is varied. Since a well-sealing barrier is notoriously difficult to construct for mercury, the coverage was increased not by barrier compression but rather by a stepwise addition of standard solution, employing a calibrated micropipet. In each step, material was added only after a full pressure equilibration of the previous step was reached.

2. X-ray Measurements. The structure of the deposited films was studied at several coverages using surface-specific X-ray techniques. These were done using the Harvard/BNL liquid surface spectrometer at beamline X22B, NSLS, Brookhaven National Laboratory, U.S.A., at wavelengths $\lambda = 1.55 - 1.58$ Å. The trough was mounted on an active vibration isolation unit, attached to the spectrometer. This arrangement effectively eliminated pickup of vibrations from the environment, which plagued early X-ray surface studies of mercury. The X-ray methods used here have been discussed in detail.

The surface-normal structure was studied by X-ray reflectivity measurements, wait times, etc. The X-ray methods used here have been discussed in detail. The Parrat matrix method was employed to fit the measured $R(q)$ data. When we had two phases in coexistence with each other, e.g., a phase of standing-up molecules in coexistence with a phase of lying-down molecules, we fitted the measured XR by a sum of the model intensities reflected from each phase. As demonstrated by the results discussed below, all the previous studies using the same method, the model and strategy employed here are well good fits to the measurements. Consistent trends are obtained in the fitted parameter values with coverage. At the same time, the number of fit parameters is kept down to a manageable number and the interparameter correlations to a minimum. This lends confidence in the results and conclusions discussed below.

Further details on the trough, the measurement procedures, and data analysis methods are given in ref 23.

III. Results: Surface Pressure–Area Isotherms

The measured surface pressure–area isotherms for fatty acids of several chain lengths, $7 \leq n \leq 24$, are shown in Figure 2. The various parameter values derived from the isotherms are listed in Table 1. The isotherms were all measured at a temperature $T = 23^\circ$C. A smooth solid line connects the measured points (open circles). The dashed lines at large molecular areas are fits by the Volmer equation, $\pi(A - A_1) = k_0T$, the ideal gas law in two dimensions for molecules of a nonzero area $A_1$. Although the fit is not perfect, the agreement of all isotherms with the Volmer equation is reasonable, in the sense that the exclusion areas $A_1$ obtained from the fits (shown in Figure 3 and listed in Table 1) are close, within their admittedly rather large experimental uncertainties, to the calculated area occupied by a flat lying molecule, $A_{1calc} = 4.8(1.265n + 1.86)\AA^2$/molecule, obtained from the width, $w \approx 4.8\AA$, and length, $l \approx 1.265n + 1.86\AA$ of a CnOCH molecule. A linear fit, shown in a solid line in Figure 3, to the measured $A_1$ values yields $A_1 = [(65.5 \pm 0.4)n + (0.1 \pm 6)]\AA^2$/molecule, again, very close to the $A_{1calc}$ values, shown below.


(33) Kjaer, K.; et al. J. Phys. Chem. 1989, 93, 3200. Petrov, J.; et al. J. Phys. Chem B 1999, 103, 3417. However, the headgroup sizes derived in these studies correspond to hydrated carboxylic headgroups and were reduced here, somewhat arbitrarily, by the radius of a water molecule, $\approx 1.4\AA$.
measurements, are required to elucidate the structure of associated with transitions between such phases. Further a single "gas" phase, our surface pressure and X-ray dependent phases, quasi-2D gas and liquid, rather than the "gas" phase actually consists of at least two A

results to be described below. While it is possible, and agreement is interpreted as indicating that the Langmuir film in the low coverage regime consists of a "gas" of flat-lying molecules, the second plateau is observed for n ≥ 14 only. The decrease in A1 below the lines for n ≤ 8 is discussed in the text.

as a dash line in Figure 3. A similar agreement of the Volmer equation with the measured isotherms, and of A1, with A1calc, was also observed for alkanes on mercury and for C1800H on mercury. Here, as here, this agreement is interpreted as indicating that the Langmuir film in the low coverage regime consists of a "gas" of flat-lying molecules. This conclusion is supported by the X-ray results to be described below. While it is possible, and even plausible, that in the low coverage regime, A ≫ A1, the "gas" phase actually consists of at least two A-dependent phases, quasi-2D gas and liquid, rather than a single "gas" phase, our surface pressure and X-ray measurements did not reveal any features that could be associated with transitions between such phases. Further measurements, in particular high-resolution isotherm measurements, are required to elucidate the structure of the Langmuir film in this regime.

 Fits to the two-dimensional van der Waals equation (35) (τ + aA2) (A − Avdw) = kBT, with the interaction parameter a and the exclusion area Avdw, were only marginally better and resulted in practically the same exclusion area as for the simpler Volmer equation. Thus, both Volmer and van der Waals equations yield here an exclusion area A1 ≈ A1calc rather than the A1 ≈ 2A1calc expected for a hard-disk liquid. This effect, found also for Langmuir films on water, was addressed in a number of theoretical and experimental studies. The a values obtained from the fit exhibit a large scatter, as neither the van der Waals nor the Volmer equations of state follow closely enough the measured isotherms. The force constant of the long-distance molecular pair interaction should be related, in principle, to a. However, a reliable extraction of the force constant of any pair potential from the isotherms requires a better-fitting state equation, a more realistic effective pair potential (which will account for the chain—chain, the chain—headgroup, and the headgroup—headgroup interactions), and the derivation of an expression relating the force constant and the interaction parameter for the specific choices of potential and state equation.

As we show below, the X-ray results reveal that the 2D gas at large A consists, at least for the longer molecules, of molecular dimers rather than single molecules. We have therefore carried out also fits assuming dimers rather than monomers as the basic structural units. These fits yielded slightly larger molecular exclusion areas, Avdw = [(6.6 ± 0.3)n + (5 ± 6)] Å2/molecule (see Table 1). These larger areas are more consistent with the lengths of the dimers derived from the X-ray measurements discussed below.

For coverages higher than the bend in the isotherm, i.e., A ≪ A1, three qualitatively different types of behavior are observed in the isotherms, depending on the molecular length n. For 14 ≤ n ≤ 24, we find the same behavior which was observed in our study of stearic acid on mercury. As A is reduced from A1, we first obtain a

Table 1. Exclusion Areas for Fatty Acids of Chain Lengths n As Obtained from Fits of the Measured Isotherms by the Volmer Equation Assuming Monomers, A1, and Dimers, Avdw, and by the van der Waals Equation, A1vdw

<table>
<thead>
<tr>
<th>n</th>
<th>A1</th>
<th>A1vdw</th>
<th>A2</th>
<th>x1</th>
<th>x2</th>
<th>x3</th>
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<tbody>
<tr>
<td>7</td>
<td>23(5)</td>
<td>27</td>
<td>32</td>
<td>51.4</td>
<td>11</td>
<td>38</td>
</tr>
<tr>
<td>8</td>
<td>41(6)</td>
<td>46</td>
<td>49</td>
<td>57.5</td>
<td>20</td>
<td>42</td>
</tr>
<tr>
<td>10</td>
<td>68(7)</td>
<td>69</td>
<td>74</td>
<td>69.6</td>
<td>26</td>
<td>37</td>
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<tr>
<td>12</td>
<td>80(7)</td>
<td>80</td>
<td>86</td>
<td>81.8</td>
<td>33</td>
<td>52</td>
</tr>
<tr>
<td>14</td>
<td>83(7)</td>
<td>85</td>
<td>91</td>
<td>93.9</td>
<td>45</td>
<td>39</td>
</tr>
<tr>
<td>16</td>
<td>115(8)</td>
<td>118</td>
<td>124</td>
<td>118.2</td>
<td>62</td>
<td>47</td>
</tr>
<tr>
<td>18</td>
<td>115(8)</td>
<td>118</td>
<td>124</td>
<td>118.2</td>
<td>62</td>
<td>47</td>
</tr>
<tr>
<td>20</td>
<td>130(10)</td>
<td>132</td>
<td>141</td>
<td>130.4</td>
<td>71</td>
<td>45</td>
</tr>
<tr>
<td>24</td>
<td>157(10)</td>
<td>157</td>
<td>162</td>
<td>154.7</td>
<td>76</td>
<td>46</td>
</tr>
</tbody>
</table>

A The experimental uncertainty is given in parentheses. The area occupied by a flat-lying molecule as calculated from its geometrical dimensions, A1calc, and the area at the beginning of the second plateau A2 (as observed in the isotherm) are also listed. The surface pressures, x1 and x2 at A1 and A2, respectively, and the collapse pressure, xc, are also given.

Figure 2. Surface pressure (τ)—area per molecule (A) isotherms for fatty acids of different chain lengths 7 ≤ n ≤ 24 (solid lines + points), shifted for clarity by τ = 5 mN/m each. The fit to the Volmer equation is shown in dashed lines.

Figure 3. The Volmer-fitted exclusion area, A1 (open circles), and starting point of the second plateau, A2 (open circles), with linear fits (lines). The dash line shows the area A1calc of a single molecule laying flat on the surface, as calculated from the molecular dimensions. The second plateau is observed for n ≥ 14 only. The decrease in A1 below the lines for n ≤ 8 is discussed in the text.

References:

(36) Israelachvili, J. Langmuir 1994, 10, 3774. See also the extensive review by Fainerman, V. B.; Vollhardt, D. J. Phys. Chem. B 2003, 107, 3098, and in ref 38, and several references therein. Note that despite the graphic similarity with our isotherms, our A1 does not correspond to A1 in these studies, but rather to A2 and ω, respectively, i.e., the area occupied by a single molecule at the monolayer’s most compressed state.
plateau (in most cases with a small slope), then a step to a second, higher, plateau. When a molecular area of $A \sim 30 \text{ Å}^2$/molecule is reached, a faster rise in the surface pressure is observed with decreasing $A$ until the film collapses at $A \leq 20 \text{ Å}^2$/molecule, when we start observing small crystallites on the otherwise clean mercury surface. The “two step” structure suggests the existence of both a single- and a double-layer phase of flat-lying molecules. As found in the stearic acid study$^{22}$ the precollapse rise in the isotherm is due to phases of standing up molecules, rendering the second, higher--$\pi$, plateau a coexistence region between lying down and standing up phases.

For $8 \leq n \leq 12$ no second plateau and a rather sloping first plateau are observed. The bend at $A_1$ becomes less pronounced and practically disappears for C7OOH. The surface pressure at the bend also decreases with decreasing $n$, yielding a shorter existence range for, and perhaps incomplete coverage of the surface by, a single layer of surface-parallel molecules. This phase seems to disappear completely for C7OOH and shorter chain lengths. This suggested trend is supported also by the observed n-dependence of the fitted exclusion areas. For $10 \leq n \leq 24$ the measured $A_1$ values are well approximated by $A_1 = [(6.5 \pm 0.4)n + (0 \pm 6)] \text{ Å}^2$/molecule, but for $n \leq 8$ the values fall below the line, indicating that at $A_1$ a molecule occupies a surface area less than that required for a surface-parallel orientation. The exclusion area fitted for C7OOH, $A_1 = 23 \text{ Å}^2$/molecule, is, in fact, close to the $\sim 20 \text{ Å}^2$/molecule occupied by a standing-up molecule. This suggests that the shorter-chain molecules tend to stand up already in the higher $A$ parts of the isotherm and thus show a phase sequence closer to that of Langmuir films of fatty acids on water.

The areas at the beginning of the second plateau also grow linearly with $n$: $A_2 = [(3.3 \pm 0.4)n + (1 \pm 8)] \text{ Å}^2$/molecule, very close to $A_2/2$ and $A_1^{\text{calc}}/2$. This, and our earlier results for C18OOH on mercury,$^{22}$ suggests that the step at $A_2$ occurs on the completion of a double layer of flat-lying molecules. The isotherms show a clear trend of sharper transitions between the regimes, and flatter plateaus, with increasing molecular length. This, in turn, indicates a better ordering of the molecules in the condensed layers for longer molecules. The structural suggestions made in this section, based on the isotherms, are further examined below in the light of the X-ray measurements described in the next section.

IV. Results: X-ray Measurements

X-ray reflectivity and GID measurements were generally done at the high $A$ end and the middle of each plateau and close to the low $A$ ends of the isotherms, where a strong rise in the surface pressure is observed. We now describe the X-ray results for two fatty acids, C14OOH and C24OOH, showing a different phase behavior and sequence for the standing-up but similar behavior for the lying-down phases. For completion, a very short summary of our previously published results for C18OOH$^{31}$ will also be given.

We discuss first the surface-normal structure at all coverages, and the in-plane structure of the more familiar monolayer phases of standing-up molecules, at $A \leq 30 \text{ Å}^2$/molecule. The more novel in-plane structure of the single and double layers of flat-lying molecules observed for $A \geq 40 \text{ Å}^2$/molecule and, in particular, their variation with chain length and temperature will be discussed separately at the end of this section, to point out n-dependent trends in these novel phases.

A. C14OOH. 1. Surface Normal Structure. Figure 4a shows a set of reflectivities measured (points) at room temperature (RT) for C14OOH at the areas per molecule listed and shown by arrows in Figure 4c. The curves are normalized to the Fresnel reflectivity, $R_f(q)$, of an ideally flat and smooth mercury surface. The box model employed yields good fits (lines). The corresponding electron density profiles derived from the fit. The curves marked SL and DL denote the thickness of the single and double layer of lying-down molecules and ML the thickness of a monolayer of standing-up molecules. (c) The isotherm, with arrows marking the positions where X-ray data were measured. The XR in (a) show from bottom to top the successive growth of two layers of flat-lying molecules, a coexistence between a standing-up phase and a two-layer lying-down phase, and a phase consisting of standing-up molecules only. The curves in both parts of the figure are shifted for clarity.

Table 2. Nominal Coverage (A) According to the Isotherms, Layer Thickness (d), Phase, Fractional Coverage of the Topmost Layer in % (Top), and the Roughness of the Fatty Acid--Air ($\sigma_{\text{fat}}$) and the Mercury--Fatty Acid ($\sigma_{\text{Hg}}$) Interfaces, As Obtained from Fits to the XR Curves in Figure 4 for a C14OOH Layer at $T = 25 ^\circ C$

<table>
<thead>
<tr>
<th>A, Å$^2$</th>
<th>d, Å</th>
<th>phase</th>
<th>top, %</th>
<th>$\sigma_{\text{fat}}, \AA$</th>
<th>$\sigma_{\text{Hg}}, \AA$</th>
</tr>
</thead>
<tbody>
<tr>
<td>88</td>
<td>4.8</td>
<td>SL</td>
<td>100</td>
<td>0.9</td>
<td>1.1</td>
</tr>
<tr>
<td>64</td>
<td>9.6</td>
<td>DL</td>
<td>40</td>
<td>1.2</td>
<td>1.3</td>
</tr>
<tr>
<td>48</td>
<td>9.6</td>
<td>DL</td>
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<tr>
<td>30</td>
<td>18.5</td>
<td>ML</td>
<td>30</td>
<td>1.0</td>
<td>1.2</td>
</tr>
<tr>
<td>20</td>
<td>18.6</td>
<td>ML</td>
<td>100</td>
<td>1.5</td>
<td>1.2</td>
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</table>

Figure 4. (a) Measured, Fresnel-normalized X-ray reflectivities (points) for C14OOH on mercury and their model fits (lines) for the listed area per molecule, taken at RT. (b) The electron density profiles derived from the fit. The curves marked SL and DL denote the thicknesses of the single and double layer of lying-down molecules and ML the thickness of a monolayer of standing-up molecules. (c) The isotherm, with arrows marking the positions where X-ray data were measured. The XR in (a) show from bottom to top the successive growth of two layers of flat-lying molecules, a coexistence between a standing-up phase and a two-layer lying-down phase, and a phase consisting of standing-up molecules only. The curves in both parts of the figure are shifted for clarity.
respectively, indicates a larger unit cell with $A$ Å$^2$/molecule for 30 Å$^2$/molecule, as compared with the molecule, except for the bottom curve that was taken at $T = 27.0 \, ^\circ C$. The single peak found indicates an hexagonal in-plane structure. (b) The temperature dependence of the GID peak’s position. (c) BR for $A = 20$ Å$^2$/molecule at $T = 27.0 \, ^\circ C$ showing a monolayer of upright molecules. There were no measurable differences in the BR for the different temperatures.

are shown in Figure 5 for the same coverage of 20 Å$^2$/molecule, except for the bottom curve that was taken at $T = 27.0 \, ^\circ C$ and a coverage of 30 Å$^2$/molecule.

For all measured temperatures $T$ and coverages $A \leq 30$ Å$^2$/molecule, i.e., for the standing-up phases, a single GID peak was observed, indicating an undistorted hexagonal unit cell. While the width of the GID peaks does not change significantly with temperature, the peak positions clearly change linearly with $T$, as shown in Figure 5b. Assuming the peak shift to result only from thermal expansion, the expansion coefficient obtained from a linear fit to the peak positions at Figure 5b is $(d/dT) \approx (6.5 \pm 0.6) \times 10^{-4}$ K$^{-1}$. This is very close to the linear expansion coefficients of bulk hexagonal rotator phase $R_{II}$ of alkanes, $6.5 \times 10^{-4}$ K$^{-1}$ and of the solid surface-frozen monolayer in alkane melts, $9 \times 10^{-4}$ K$^{-1}$. The BR shown in Figure 5c does not change with temperature. The fact that its peak is at $q_0 = 0$ indicates that the molecules are upright, to within $\theta \leq 3^\circ$, from the surface normal. This structure is the same as that observed for a Langmuir film of C14OOH on water at similar temperatures and at the highest coverage. No tilted phases were observed for C14OOH on mercury for any other coverage $A$, as was found also for C14OOH on water. The GID peaks at coverages of 20 and 30 Å$^2$/molecule were both measured at $T = 27.0 \, ^\circ C$. The peak positions, $q_1 = 1.502 \, \AA^{-1}$ and $q_2 = 1.486 \, \AA^{-1}$, respectively, indicates a larger unit cell with $A = 20.6$ Å$^2$/molecule for 30 Å$^2$/molecule, as compared with the $A = 20.2$ Å$^2$/molecule for 20 Å$^2$/molecule. The peaks’ full width at half-maximum for the lower coverage is significantly larger than that of the high coverage: $\Delta q_1 = 0.036$ Å$^{-1}$ at $A = 30 \, \AA^2$/molecule, as compared to $\Delta q_2 = 0.016$ Å$^{-1}$ at $A = 20 \, \AA^2$/molecule. This, in turn, yields a shorter crystalline coherence length of $L = 0.95/\Delta q_2 \approx 160 \, \AA$ at the lower coverage, as compared to $L \approx 350 \, \AA$ for the high coverage. This reduction in coherence length may result from the coexistence of two phases on the surface at this coverage: regions of hexagonally packed, standing-up molecules, enclosed within a double layer of lying-down molecules. At the highest coverage, all results (the hexagonal symmetry, the linear expansion coefficient and the coherence lengths) strongly support the conclusion that the standing-up phase is a hexatic-like LS rotator phase, similar to that of Langmuir films on water.

For the lying-down phases no in-plane GID peaks were detected at room temperature, indicating that no long-range in-plane order exists.

**B. C18OOH.** The results for the standing-up phases of C18OOH have been already published$^{31}$ and will be summarized here only briefly. Similar to C14OOH, for the standing-up monolayer phase of C18OOH, we observe an LS phase$^3$ at the highest coverage with lattice constants and coherence lengths similar to those of C14OOH. However, unlike C14OOH, tilted phases were found in this regime for C18OOH. We find an Overbeck–Möbius phase (Ov)$^3$ which has a next-nearest-neighbor tilt of the molecules in the standing-up phase, with a distorted hexagonal packing in the plane perpendicular to the molecules’ long axis. As is typical for these phases, increasing the coverage causes a continuous decrease in the tilt angle and a continuous shift of the Ov phase into an upright LS phase. For low coverages, lying-down phases, GID measurements at room temperature reveal here a one-dimensional in-plane order, quite unlike the in-plane-disordered lying-down phase of C14OOH. These results are further discussed at the end of this section.

**C. C24OOH.** 1. Surface-Normal Structure. The XR measurements carried out for C24OOH at several different coverages are shown in Figure 6. The fit parameters are summarized in Table 3. The same fitting procedures were used as for C14OOH.

Beginning with the 161, 116, and 76 Å$^2$/molecule curves, we observe the formation of a single layer, coexistence of a single layer and a double layer, and a 100% complete double layer, respectively. The two scans at high coverage, 27 and 21 Å$^2$/molecule, were taken for the same sample for each pair. However, the upper curve of each pair was taken 1 h later than the lower curve of the pair. For 27 Å$^2$/molecule, the fits of both scans result in 60% of the area being covered by a monolayer of standing-up molecules and 40% being covered by a double layer of lying-down molecules. Despite the same relative coverage, a comparison of the XR curves reveals a clearly visible decrease in the oscillation period. As shown in Table 3, this

![Figure 5](image1.png)

**Figure 5.** GID scans for the standing-up phase of C14OOH. (a) GID scans at $A = 20$ Å$^2$/molecule for the listed temperatures and for $A = 30$ Å$^2$/molecule for $T = 27.0 \, ^\circ C$. The single peak found indicates an hexagonal in-plane structure. (b) The temperature dependence of the GID peak’s position. (c) BR for $A = 20$ Å$^2$/molecule at $T = 27.0 \, ^\circ C$. The single peak found indicates an hexagonal in-plane structure. (d) BR for $A = 20$ Å$^2$/molecule at $T = 27.0 \, ^\circ C$ showing a monolayer of upright molecules. There were no measurable differences in the BR for the different temperatures.

![Figure 6](image2.png)

**Figure 6.** Same as Figure 4, but for C24OOH. Note the formation of a single and double layer of flat-lying molecules. The tilted and untilted standing-up phases observed at 21 and 27 Å$^2$/molecule are discussed in the text.
We show here equal-intensity contour plots of the time-dependent GID measurements for C24OOH at a coverage of $A = 27 \, \text{Å}^2$ molecule. The scans from (a) to (d) were taken 15, 50, 60, and 90 min after film deposition.

**Table 3. Nominal Coverage (A) According to the Isotherms, Layer Thickness (d), Phase, Fractional Coverage of the Topmost Layer in % (Top), and the Roughness of the Fatty Acid–Air ($\sigma_{\text{res}}$) and the Mercury–Fatty Acid ($\sigma_{\text{fatty}}$) Interfaces, As Obtained from Fits to the XR Curves in Figure 6 for a C24OOH Layer at $T = 25 \, ^\circ\text{C}$**

<table>
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<th>A, $\text{Å}^2$</th>
<th>d, Å</th>
<th>phase</th>
<th>top, %</th>
<th>$\sigma_{\text{at}}, \text{Å}$</th>
<th>$\sigma_{\text{res}}, \text{Å}$</th>
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<td>161</td>
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<tr>
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<td>DL</td>
<td>40</td>
<td>1.2</td>
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<tr>
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<tr>
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<td>ML</td>
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<tr>
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<td>ML</td>
<td>60</td>
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<td>ML</td>
<td>100</td>
<td>1.1</td>
<td>1.1</td>
</tr>
</tbody>
</table>

corresponds to an increase in the thickness of the standing-up part of the monolayer from 28.5 Å for the first scan to 32.3 Å for the later scan. The second of these values equals the extended length of a C24OOH molecule, $l \approx 32.2 \, \text{Å}$. The same behavior occurs for the higher coverage. However, in this case both curves yield a 100% coverage by a monolayer of standing-up molecule, without a trace of a contribution from a lying-down phase. This suggests that upon increasing the coverage above that of a complete double layer of lying-down molecules, some of the molecules start forming a standing-up tilted phase with a tilt of $\theta \approx \arccos(28.7/32.2) = 27^\circ$ from the surface normal. For rigid chains this corresponds to a shift, due to tilt, of two carbons along the chain in adjacent molecules, as expected from the requirement of space filling in closely packed chains, and as indeed found in SAMs of alkane thiols on gold substrates. The change with time indicates that this phase is metastable and transforms with time to an untilted monolayer phase.

2. **Surface-Parallel Structure.** To determine the in-plane structure of the tilted and untilted standing-up monolayer phases, we carried out GID measurements at different coverages and temperatures. To follow possible changes with time of the in-plane structure that might accompany the variations found in the XR results, we also carried out time-dependent GID scans at a few coverages. A typical time sequence is shown in Figure 7. We show here equal-intensity contour plots of the GID peaks for a coverage of $27 \, \text{Å}^2$/molecule. The scans were taken 15, 50, 60, and 90 min after film deposition. We observe at least three different phases and often also a coexistence of some of the phases. The sequence of metastable phases terminates after slightly over 1 h in a crystalline CS phase with a body-centered rectangular unit cell and a herringbone packing of the molecules.

We focus first on the transient phases and return below to discuss in detail the stable CS phase and its temperature dependence. Figure 7a shows two peaks at $q_1 = 1.316 \, \text{Å}^{-1}$, $q_2 = 0.45 \, \text{Å}^{-1}$ and at $q_3 = 1.512 \, \text{Å}^{-1}$, $q_4 = 0.38 \, \text{Å}^{-1}$. These two peaks often appear together immediately after deposition of new material on the mercury surface. Attempts to associate them with a single phase result in an unphysical unit cell. Moreover, since they disappear rather quickly (a few minutes after deposition), it is not clear whether they are the only first-order peaks of this particular phase or whether other peaks may exist somewhere else in $q$-space. This pattern is replaced within a few minutes by the pattern shown in Figure 7b. Here we find two GID peaks as well, which, contrary to those in Figure 7a, exist for up to a few tens of minutes and can be measured with better reproducibility and accuracy. Their peak positions are $q_1 = 1.300 \, \text{Å}^{-1}$, $q_2 = 0.66 \, \text{Å}^{-1}$ and at $q_3 = 1.447 \, \text{Å}^{-1}$, $q_4 = 0.38 \, \text{Å}^{-1}$. They can be assigned to the (02) and (11) diffraction peaks of a body-centered rectangular cell with dimensions $a = 9.66 \, \text{Å}$ and $b = 4.86 \, \text{Å}$ and two molecules per unit cell. A detailed fit to the corresponding GID pattern and Bragg rods, shown in Figure 8, yields a tilt angle of $\theta = 27^\circ$ in a direction rotated azimuthally by $\phi = 25^\circ$ from the nearest-neighbor direction. This yields a unit cell in the plane perpendicular to the molecular long axis, with the lattice parameters $a = 4.86 \, \text{Å}$, $b = 8.62 \, \text{Å}$ and with a molecular area of $A = ab/2 = 20.9 \, \text{Å}^2$/molecule. The ratio of $b/a = 1.77$ is close to $3^{1/2} = 1.73$, which indicates a hexagonal hexitic-like rotator phase. This phase is very similar to the OV phase observed for stearic acid (C18OOH) on mercury at high coverages (see ref 22 and summary above) except that for lignoceric acid (C24OOH) the peak positions and tilt angles do not change with coverage. Another difference is the significantly sharper GID peaks, which are close to the resolution limit, $\Delta q_{\text{res}} = 0.0056 \, \text{Å}^{-1}$. The full width at half-maximum (fwhm) of the peaks, $\Delta q_{\text{fwhm}} \approx 0.009 \, \text{Å}^{-1}$, indicates a large crystalline coherence length, of the order of 80 Å. This can be rationalized by the stronger chain–chain interaction of longer chains, which drives the system toward a closer packing and longer-range order in the crystalline layer.
The tilted Ov-like phase, discussed in the last paragraph, converts within about 1 h into a crystalline CS phase, the GID pattern of which is shown in Figure 7d. Similar slow relaxation, going through a complicated phase sequence, was also found for Langmuir films of tetradecanoic acid on water. During the conversion, a coexistence of the two phases is observed, with the GID patterns showing the peaks of both phases (Figure 7c). The CS phase is crystalline with a herringbone ordering. At room temperature, its two lowest peaks are observed at $q_a = 0$ Å$^{-1}$ and $q(11) = 1.512$ Å$^{-1}$ and $q(02) = 1.688$ Å$^{-1}$. This corresponds to two untilted molecules in a body-centered rectangular unit cell of dimensions 5.02 × 7.44 Å². The resultant molecular area, 18.6 Å²/molecule, equals the closest packing of crystalline alkyl chains of, e.g., Langmuir films on water, the untilted phase of the crystalline fatty acids and alkanes, and surface-frozen monolayers on alkane melts.

The temperature dependence of the herringbone CS phase is shown in Figure 9 at 10, 19, and 25 °C. The peaks move from $q(11) = 1.512$ Å$^{-1}$ and $q(02) = 1.688$ Å$^{-1}$ at $T = 25$ °C to $q(11) = 1.516$ Å$^{-1}$ and $q(02) = 1.712$ Å$^{-1}$ at $T = 10$ °C. The corresponding unit cell vectors change from $a = 5.01$ to 5.02 Å and from $b = 7.44$ to 7.34 Å upon decreasing the temperature. Thus, the thermal expansion of this phase is highly anisotropic. While the length of the unit cell hardly changes with the decrease of the temperature, the corresponding vectors change significantly, showing a coexistence of the two phases. The corresponding unit cell vectors change from $a = 5.01$ to 5.02 Å and from $b = 7.44$ to 7.34 Å upon decreasing the temperature. Thus, the thermal expansion of this phase is highly anisotropic. While the length of the unit cell hardly changes with the decrease of the temperature, the corresponding vectors change significantly, showing a coexistence of the two phases.

The packing of the molecules in the unit cell is very dense, and the area per molecule shrinks from 18.6 to 18.4 Å²/molecule over this temperature range. The peak width for the herringbone packing is always resolution limited, indicating a crystalline coherence length of more than 1000 Å, at all temperatures. It is interesting to note that upon prolonged exposure to the beam, the peak width is observed to increase, and the peak moves to lower $q$ values. These symptoms, well-known from conventional x-ray diffraction, indicate the proliferation of X-ray-induced defects in the (two- and three-dimensional) crystalline structure.

In conclusion, for C24OOH, although an Ov-like phase of tilted molecules (~27°) is observed on short timescales following film deposition, the final, equilibrium phase at room temperature and below is always a crystalline CS phase with a high density of 18.5 Å²/molecule. This is in contrast with C18OOH, where an equilibrium Ov phase was observed, and with C14OOH, where a hexagonal untilted rotator phase is the equilibrium phase at the highest coverages.

**D. The Structure of the Lying-Down Phases.** We now discuss our GID and BR measurements of the lying-down phases of fatty acids on mercury. Figure 10 shows the GID patterns measured at low $q$ values for the fatty acids indicated. The patterns were recorded each for a coverage corresponding to two layers of lying-down molecules. Scans for a coverage of a single layer of lying-down molecules yield the same patterns, but beam damage by the intense synchrotron radiation becomes a more serious problem and prevents employing the long measuring times required for achieving good statistics. No variation is observed in the patterns upon varying the coverage within the lying-down phase. In view of the fact that in-plane GID peaks appear for C14OOH only below $T = 18$°, the GID pattern shown was recorded at $T = 5$ °C. The GID peaks of C18OOH and C24OOH were observed for all temperatures measured, 5 °C ≤ $T$ ≤ 30 °C. All peaks are resolution limited, suggesting a crystalline coherence length in excess of 1000 Å. All scans show four diffraction peaks at positions which are multiples of the lowest-q peak position in each curve. These are $q_a = 0.1477$ Å$^{-1}$, $q_e = 0.1201$ Å$^{-1}$, and $q_i = 0.0932$ Å$^{-1}$, corresponding to repeat distances of $d = 42.5$ Å, $d = 52.3$ Å, and $d = 67.4$ Å, for C14OOH, C18OOH, and C24OOH, respectively.

The respective calculated fully extended molecular lengths are 19.6, 24.6, and 32.2 Å for C14OOH, C18OOH, and C24OOH, rendering the repeat distances longer by 3.0–3.3 Å than twice the molecular length. This, in turn,
strongly suggests that the order is formed by molecular dimers rather than individual molecules. The two carboxylic head groups reside, most likely, at the center of the dimer, and their interaction stabilizes the dimer. Since the excess \( \sim 3 \text{ Å} \) length renders the distance between the carboxylic groups most probably too large for hydrogen bonding, it is likely that at least one H-gatom participates in the bonding at the center of the dimer. This suggestion is supported, albeit circumstantially, by the 2.8–3.0 Å diameter of the mercury atom (or ion)\(^{(47)}\) and by the \( \sim 3.2 \text{ Å} \) interatomic distance at the freeman surface of mercury, taken from the liquid peak position in our previous reflectivity and GID measurements on bare mercury.\(^{(47)}\) It is also supported by the exceptionally strong GID peaks which suggest that the ordered phase comprises scatterers of high electron density.

All chain lengths studied exhibit the low-\( q \) peaks indicating ordering in the direction of the molecular axis. The peaks show similar characteristics for all chain lengths: position commensurate with twice the molecular length, sharp peaks with resolution-limited widths, several diffraction orders, etc. By contrast, the very existence of GID peaks at \( q_z \sim 1.3 \text{ Å}^{-1} \), which corresponds to order in the direction perpendicular to the molecular axis, is very different for different chain lengths. The \( q_z \)-range studied was 1.15 \( \leq q_z \leq 1.35 \text{ Å}^{-1} \). This corresponds to repeat distances 4.65 Å \( \leq l \leq 5.46 \text{ Å} \), which can be expected from close, or even loose, packing of molecules of widths \( \sim 4.8 \text{ Å} \), as discussed above. For C1400H no GID peaks could be observed in this range for all temperatures studied, 5 °C \( \leq T \leq 27 °C \). For C1800H, while no GID peaks were found in this \( q_z \)-range at room temperature, below \( T = 20 °C \) a single GID peak, shown in Figure 11a, was observed at \( q_z = 1.302 \text{ Å}^{-1} \). This corresponds to a repeat distance of \( l = 4.83 \text{ Å} \), which is typical of interchain distances of alky chains in several close-packed two- and three-dimensional systems. The intensity of this peak is, however, rather low and the points of the corresponding BR, shown in Figure 11b in open circles, are even more so. Unfortunately, beam damage does not allow collecting data with better statistics. Thus, the quality of the available BR is insufficient for a detailed modeling with a reasonable level of confidence. Nevertheless, the data are reasonably consistent with a model, shown by the solid line in Figure 11b, of a double layer of flat-lying cylindrical molecules, where the molecules in the second layer lie in the hollows of the molecules in the first layer. Connecting the centers of the cross sections of two adjacent cylinders, one in the first layer and the other in the second layer, yields a line tilted by 30° from the surface normal. Thus, the BR should resemble that of a tilted molecular phase, as indeed it does, see, e.g., part b or c of Figure 8, making allowances for the much narrower BR peak there. The BR's peak here is at \( q_z \approx 0.7 \text{ Å}^{-1} \), yielding a tilt of \( \phi = \arctan(0.7/1.302) = 28^° \), which agrees well with the expected value. It is interesting to note that despite the low intensity of the GID peak, its width has the same resolution-limited value as that of the longitudinal GID peaks, indicating an ordering coherence length in excess of 1000 Å.

In contrast with the shorter chain length fatty acids, C1400H and C1800H, for the lying-down phases of C2400H at room temperature, we could readily observe two diffraction peaks, shown in Figure 12, at \( q_z = 1.262 \text{ Å}^{-1} \) and 1.312 Å\(^{-1} \). The corresponding repeat distances are 4.98 and 4.79 Å and hence belong to ordering perpendicular to the molecular axis, as discussed above. As can be seen in Figure 12b for the \( q_z = 1.262 \text{ Å}^{-1} \) GID peak, the BR peaks at \( q_{z1} = 0 \text{ Å}^{-1} \) and a fit (line) yields a width corresponding to \( d \approx 5 \text{ Å} \), i.e., that of a single lying-down layer. For the \( q_z = 1.312 \text{ Å}^{-1} \) GID peak the BR peaks at \( q_z \approx 0.66 \text{ Å}^{-1} \) and can be fitted (line) by a \( \sim 27^° \)-tilted layer of 10 Å thickness, which corresponds to a double layer of lying-down molecules, where the molecules of the second layer reside in the hollows of the first layer. The rather large, \( \sim 0.2 \text{ Å} \), shrinking of the repeat distance from the single to the double layer is a manifestation of the much stronger van der Waals intermolecular interaction within the layers, due to the existence of a second layer, which causes a tighter packing.

We now compare the molecular areas of the lying-down molecules obtained from the X-ray measurements and from fits to the exclusion areas of the isotherms. Assuming a molecular width of 4.8 Å, obtained from the GID and also the XR measurements on the lying-down phases, the unit cell lengths obtained from the GID measurements yield molecular areas of 102, 126, and 162 Å\(^2\)/molecule for C1400H, C1800H, and C2400H, respectively. From the relation \( A_{eq} = (6.6n + 5) \text{ Å}^2 \) obtained above for the exclusion area from isotherm fits assuming dimers, we

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obtain 97, 124, and 163 Å²/molecule for C14OOH, C18OOH, and C24OOH, respectively. The two sets of numbers are obviously in very good agreement.

To summarize the structure of the lying-down phases, the basic building block of the structure in these phases consists of extended molecular dimers, bound at their center by the two carboxylic headgroups and perhaps incorporating at least one mercury atom. At room temperature, C14OOH shows no in-plane order at all. C18OOH shows a one-dimensional ordering along the dimer’s long axis, but no order in the direction perpendicular to the dimer’s long axis. This can be regarded as a one-dimensional, smectic-like order in a quasi-two-dimensional layer. C24OOH exhibits a two-dimensional crystalline structure, with order both along, and perpendicular to, the dimers’ long axis. In C14OOH a 1D order along the dimer’s axis can be induced by lowering the temperature to \( T = 18 \) °C. The 1D order in C18OOH is converted to a 2D order, like that of C24OOH at room temperature, by lowering the temperature below \( T = 20 \) °C. The dimers in the top layer of the double-layer phase are found in all cases to reside in the hollows of the dimers of the first layer.

**E. The Phase Diagram.** From the combined surface tensiometry and X-ray measurements, we can construct a preliminary phase diagram. Note, however, that although isotherms were measured for a broad range of chain lengths, the X-ray data were measured only for C14OOH, C18OOH, and C24OOH.

The room-temperature projection of the three-dimensional (\( A, n, \pi \)) phase diagram on the (\( n, \pi \)) plane is shown in Figure 13. When looking at this plot, one must bear in mind that as \( A \) is decreased from a large value of a few hundred Å², \( \pi \) increases monotonically, but certainly not linearly: there are \( A \) ranges where \( \pi \) increases hyperbolically, whereby it stays constant, and where it increases with a large and varying slope. Film collapse, at \( A \lesssim 20 \) mN/m, marks the transition from a quasi-2D monolayer to 3D crystallites. This is marked in a bold solid line in the figure.

For C70OH and shorter chain lengths, as the molecular area \( A \) is reduced the surface pressure \( \pi \) starts increasing from near-zero only when the \( A \) values become very close to that of standing-up molecules, a few tens of mN/m. Thus, for \( n \leq 7 \), no lying-down phases seem to exist, and the only phase is a monolayer of standing-up molecules. For \( 8 \leq n \leq 12 \) a single layer of lying down molecules, SL, followed by a monolayer of standing up molecules, ML, are most likely formed upon decreasing \( A \). Conclusive X-ray measurements in these ranges are difficult to carry out due to the relatively small aspect ratio of the molecules, which make it difficult to distinguish standing-up from lying-down phases. Moreover, hints from corresponding films on water are not available because of the larger solubility of these short chain fatty acids in water. Also, the 3-fold higher surface roughness of water, as compared to mercury, severely restricts the measurable \( \pi \) range and, hence, the correspond resolution.

For \( n = 14 \) the X-ray and isotherm data show disordered, liquidlike SL and DL phases of lying-down molecules followed, at higher \( \pi \) values, by a standing-up LS rotator phase. For \( n = 18 \) the DL phase shows a smectic-like one-dimensional in-plane order of dimers. At about the same n-range the standing-up monolayer starts showing an additional hexatic phase: untitled LS and tilted Ov. Somewhere between \( n = 18 \) and \( n = 24 \) both the lying-down and standing-up phases become more ordered. The lying-down phases show a two-dimensional crystalline order of dimers, and the standing-up phases show a fully crystalline herringbone CS structure, although a transient Ov structure is often also observed for short periods following the deposition. In general, therefore, the order grows with increasing chain length and decreasing temperature. For example, at room temperature the DL phase of C14OOH is not ordered, C18OOH is smectic-like and C24OOH is crystalline. This trend is in line with the behavior of fatty acid Langmuir films on water, where an increase in the chain length was found to be equivalent to a decrease in temperature. Once this effect, properly quantified, was taken into account, the \( (n, \pi) \) phase diagrams of all fatty acids on water, regardless of their chain length, collapsed onto one “universal” phase diagram. Quantitative establishment of, or ruling out, a similar behavior for films of fatty acids on mercury will have to await the availability of a more complete set of data on the temperature dependence of the phase diagram than the few measurements carried out in this study.

In general, the high-coverage standing-up phases of fatty acids on mercury are very similar to the phases of fatty acids on water, despite the fact that they cover a very small part of the full \( (n, \pi) \) phase diagram. The high pressure at which these standing-up phases appear on mercury seems to suppress almost everywhere on the phase diagram the tilted phases observed for fatty acids on water. For C14OOH we did not observe tilted phases at all. For C24OOH we observed tilted phases, but with different symmetries than those of C24OOH on water. Moreover, the observed tilted phases were found to be unstable at all coverages and convert with time to the crystalline untitled CS phase. Only C18OOH exhibits here the same tilted and untitled phases found for C18OOH monolayers on water. For the highest coverage the phase sequence at room temperature and below seems to be the same as that of Langmuir films of fatty acids on water, but additional temperature and chain length dependent measurements are needed to quantify this assumption confidently.

The discussion above demonstrates that many details of the phase diagram can be derived from the isotherms and the (somewhat sparse) X-ray data presented here. Thus, although the general features of the phase diagram, as shown in Figure 13, are well determined, the exact assignments of phase boundaries in the \( (n, \pi) \) plane, and the determination of their exact temperature variation, will have to await more detailed X-ray measurements.

**Figure 13.** Surface pressure (\( \pi \))–chain length (\( n \)) phase diagram for fatty acid films on mercury. For discussion, see text.
The phase diagram in Figure 13 can be used to derive a rough estimate for the adsorption energies, through a comparison of the heat of vaporization with the surface pressure at the transition from the single-layer lying-down phase to the standing-up monolayer, marked by the bold dashed line. The details of the method are discussed in our previous publication on Langmuir films of alkanes on mercury. In a simplified view, the surface pressure \( \pi \) should be equal to the difference between the adsorption energy \( E_{\text{ads}} \), and the heat of vaporization \( \Delta H_{\text{vap}} \). A fit of the simple theoretical expression derived in ref 23 to the experimental data, shown in a dash line in Figure 13, yields adsorption energies of 5.3 kJ/(CH2 mol) for a methyl group and \( E_{\text{ads}} \approx 28 \text{ kJ}/(\text{CHOOH mol}) \) for the carboxylic headgroup. The result for a methyl group agrees very well with the 5.4 kJ/(CH2 mol) obtained from alkanes on mercury. The adsorption energy of the carboxylic headgroup is five times larger, making the headgroup adsorption energy comparable to that of the full alkyl chain in short molecules. Vertical orientation of the molecule relative to the surface maximizes the headgroup–subphase contact, while surface-parallel orientation maximizes the alkyl–subphase contact. Thus, the relative magnitude of these two interactions will determine the molecular orientation at the surface at low surface pressures. When the former is larger, at short chain length, the molecules stand up spontaneously at low pressures. When the latter is larger, at longer chains, the molecules tend to lie down and can be induced to stand up only by increasing the pressure. This explains our observation of only a standing-up phase at \( n \leq 7 \) and a sequence of a lying down phase at low \( \pi \) followed by a standing up phase at higher \( \pi \) at higher \( n \). The large adsorption energy of the headgroup also explains why short-chain fatty acids form Langmuir films on mercury, while the same-length alkanes do not form Langmuir films on mercury, and rather prefer to form 3D crystallites. The transition among the various phases found, and the very existence of a the lying-down phases, can be rationalized, in principle, within a balance among several competing interactions. A major factor is the Hamaker constant of the liquid metal–organic film–vapor system which is large and positive in contrast with the small and negative Hamaker constant found for Langmuir films on water. This favors here a thin layer of lying-down molecules over a thick layer of standing-up molecules (albeit consisting of islands), as found on water. Other factors are the bulk cohesive energy, which favors standing-up molecules with maximal chain contact, as in bulk, the interaction of the acidic headgroup with the mercury (attractive) and the chains (repulsive) and the attractive chain–mercury interaction. Work on these issues is in progress.

### F. Conclusion

The major features of the phase behavior of fatty acid films on mercury were determined in the present study. Unlike fatty acid molecules on water, which show only standing-up phases, on one hand, and alkanes on mercury, which show only lying-down phases, on the other hand, we observed here both lying-down and standing-up phases. These differences demonstrate the importance of the subphase as well as the headgroup in determining the structure of the Langmuir film.

The lying down phases differ from those of alkanes on mercury by showing disorder, 1D-smectic-like order, and 2D-order of molecular dimers, depending on chain length, while for alkanes no in-plane order was found for any chain length. The standing-up phases are found here to be similar to those of fatty acids on water, with increasingly higher ordering with chain length. However, due to the much higher pressures required to generate these phases, untilted stable phases are mostly observed, albeit transient tilted phases do appear for a short time following film deposition.

Finally, the present results, in conjunction with similar studies of Langmuir films of alkanes and alcohols on mercury should provide for a deeper understanding of the relative importance and role of the subphase–headgroup and subphase–alkyl chain interactions in determining the structure of, and molecular orientations in, these films. Clearly, the standing-up phases appear only when a strong headgroup–subphase interaction exists. For alkanes, lacking such headgroups, only lying-down phases are found. However, more subtle differences in structure and phase sequence also depend on the relative strength of these two interactions. These issues will be addressed in future publications.

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