Slow Diffusive Motions in a Monolayer of Tetracosane Molecules Adsorbed on Graphite


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Abstract. Monolayers of intermediate-length alkane molecules such as tetracosane (n-C_{24}H_{50} or C24) serve as prototypes for studying the interfacial dynamics of more complex polymers, including bilayer lipid membranes. Using high-resolution quasielastic neutron scattering (QNS) and exfoliated graphite substrates, we have investigated the relatively slow diffusive motion in C24 monolayers on an energy/time scale of ~1-36 μeV (~0.1-4 ns). Upon heating, we first observe QNS in the crystalline phase at ~160 K. From the crystalline-to-smectic phase transition at ~215 K to a temperature of ~230 K, we observe the QNS energy width to be dispersionless, consistent with molecular dynamics simulations showing rotational motion of the molecules about their long axis. At 260 K, the QNS energy width begins to increase with wave vector transfer, suggesting onset of nonuniaxial rotational motion and bounded translational motion. We continue to observe QNS up to the monolayer melting temperature at ~340 K where our simulations indicate that the only motion slow enough to be visible within our energy window results from the creation of gauche defects in the molecules.

INTRODUCTION

Over the past three decades nuclear magnetic resonance (NMR) has been used to investigate the conformational dynamics of lipid molecules in membranes [1]. In particular, measurements of the nuclear quadrupole splitting at selectively deuterated methylene groups along the alkyl chains have yielded the distribution of gauche defects along the chains in the membrane's fluid phase [2].

Recently, we have shown that the melting transition in a monolayer of intermediate-length alkane molecules (n-C_{n}H_{2n+2}), such as tetracosane (n = 24) and dotriacontane (n = 32), adsorbed on a graphite basal-plane surface provides an interesting analog of the gel-to-fluid transition in bilayer lipid membranes [3]. That is, our molecular dynamics (MD) simulations provide evidence of a monolayer melting transition in which intramolecular and translational order are lost simultaneously [3]. Figure 1 shows a top view of the tetracosane (C24) molecules in the MD simulation cell above and below the melting transition at ~340 K. Below the transition, we see translational order manifested by a lamellar structure. We refer to this structure as the "smectic" phase, since both the simulations and quasielastic neutron scattering measurements (see below) indicate some translational diffusive motion within the lamellae. Above the melting transition, the individual molecules have transformed to a more globular shape, i.e. have undergone a "chain-melting," and the monolayer has lost the translational order characterized by its lamellar structure. As in the case of the lipid membrane, the simultaneous "chain" melting and lattice melting of the alkane monolayer result from an abrupt increase in the number of gauche defects within the central region of each molecule.

Here we describe how high-resolution quasielastic neutron scattering (QNS) can be used to investigate the diffusive motion in a C24 monolayer occurring on a time scale of ~0.1-4 ns. We present evidence that
RESULTS AND DISCUSSION

We began our measurements at low temperature where we anticipated the molecular diffusive motion would be simplest to describe. Elastic neutron diffraction measurements on deuterated samples show that, upon cooling, the C24 monolayer undergoes a transition at ~215 K from the “smectic” to a commensurate phase in which the monolayer contracts in a direction parallel to the lamellae boundaries [7]. The nearest-neighbor molecular separation in this direction becomes 4.26 Å = √3 a_0 where a_0 = 2.46 Å is the graphite basal-plane lattice constant compared to ~4.6 Å in the “smectic” phase. This higher-density crystalline phase is similar in structure to that found at room temperature when a C24 monolayer is grown from a heptane (n-C_7H_{16}) solution or when a partial layer of either heptane or C24 is adsorbed above the C24 monolayer [6].

In Fig. 2, we show quasielastic spectra taken on the HFBS at two C24 coverages, θ = 1.0 and 1.15 layers, where unity coverage is defined as a complete monolayer of the “smectic” phase. At all temperatures in Fig. 2, both samples are in the higher-density crystalline phase described above. Due to the compression of the first layer when a partial second layer is added, we estimate that at θ = 1.15 there is about 7% of a second layer present. The solid line in the spectra of Fig. 2 represents the HFBS’ energy resolution function. It is actually the quasielastic spectrum observed from the sample at 100 K where all diffusive motion within the film is frozen out.

For both coverages, we begin to observe QNS having an energy width greater than that of the HFBS’ resolution function at a temperature of ~160 K. MD simulations indicate [8] that this results from motion in the crystalline phase involving a uniaxial rotation of the C24 molecules about their long axis aligned parallel to the graphite basal-plane surface (the molecules are nearly in the all-trans configuration).

Above ~215 K at both coverages, we begin to observe QNS with sufficient intensity to determine the Q dependence of its energy width. At these temperatures, the 1.0-layer sample has transformed to the “smectic” phase, while the 1.15-layer sample remains crystalline. We have fit the quasielastic intensity at each Q to the sum of a Lorentzian and a delta-function convoluted with the instrumental resolution function. As shown in Fig. 3, the half-width at half-maximum (HWHM) of the Lorentzian component is essentially dispersionless at temperatures of 215 K and 230 K for both coverages. Our MD simulations indicate that in the “smectic” phase the
diffusive motion at 215 K and 230 K is still dominated by a uniaxial rotation of the molecules about their long axis aligned parallel to the graphite surface. Such a model is consistent with the dispersionless HWHM of the Lorentzian component of the QNS. Note that at 215 K and 230 K the 1.15-layer sample (right panel) is still in the crystalline phase, and its HWHM's are somewhat smaller than for the 1.0-layer sample.

We see in Fig. 3 that at a temperature of 260 K the energy width of the QNS of the 1.0-layer sample more than triples, and the HWHM also begins to increase with $Q$. Our simulations suggest that this $Q$-dependence results primarily from nonuniaxial rotational motion and, to a lesser degree, translational diffusion.

It is interesting to compare the HWHM's for the higher-coverage sample at 260 K. They have increased very little above their values at 230 K, and they remain dispersionless. In fact, the magnitudes of the HWHM for the 1.15-layer sample at 260 K are about the same as for the 1.0-layer sample at 230 K. These results suggest that both the nonuniaxial rotational motion and translational diffusion are inhibited in the 1.15-layer sample, which remains in the crystalline phase.

At a temperature of 330 K, our measurements support the prediction of the MD simulations [8] that QNS should be observable due to the creation and annihilation of gauche defects in the C24 molecules. In Fig. 4, we have analyzed the $Q$ dependence of the HFBS spectra of the 1.0-layer sample at temperatures of 285 K and 330 K in the same way as for lower temperatures in Fig. 3: (a) the Lorentzian HWHM characterizing the energy width of the observed QNS; and (b) the relative intensity of the Lorentzian component in the spectra. Although the intensity becomes very weak at 330 K, it is clear that there is still QNS within the dynamic range of the HFBS. Comparing with Fig. 3, we see that at each $Q$ the energy width of the QNS continues to increase with temperature above 260 K. There is also a large increase in the slope of the HWHM vs. $Q$ between 260 K and 285 K. For comparison, we have also plotted in Fig. 4 (dashed line) the HWHM of a Lorentzian fitted
\[ \theta = 1.0 \quad \theta = 1.15 \]

**FIGURE 3.** Results of fitting the quasielastic spectra at temperatures \( \leq 260 \) K for the low-coverage sample (left panel) and higher coverage sample (right panel). The energy width (HWHM) of the Lorentzian component is plotted vs. wave vector transfer \( Q \).

to the incoherent inelastic structure factor calculated from the MD simulation at 330 K after removing the center-of-mass motion from the C24 molecules. These calculated HWHM's characterize the contribution to the QNS from both rotational and intramolecular diffusive motions. We conclude that both their magnitude and \( Q \)-dependence are in qualitative agreement with experiment.

In summary, our QNS measurements and MD simulations indicate two principal temperature ranges of interest. The first of these is below 230 K where the diffusive motion primarily involves rotation about the long axis of the molecule aligned parallel to the surface. The second temperature range of interest is above 285 K where we identify the observed QNS with motions involving conformational changes of the molecules as predicted by the simulations.

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