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Molecular dynamics simulations of hexane on graphite at various coverages: the difference explicit hydrogens make M.W. ROTH, M.J. CONNOLLY, University of Northern Iowa, Department of Physics, PAUL A. GRAY, University of Northern Iowa, Department of Computer Science, CARLOS WEXLER, University of Missouri Columbia, Department of Physics and Astronomy — Molecular Dynamics simulations of hexane (C_6H_{14}) adlayers on graphite are carried out for coverages of $0.5 \le \rho \le 1$ monolayers. The hexanes have explicit hydrogens and the graphite is modeled as an all – atom, six – layer structure. Above $\rho \cong 0.9$ the herringbone solid loses orientational order at $T_1 = 140 \text{ K} \pm 3 \text{ K}$. At $\rho = 0.878$ the system presents vacancy patches and T_1 decreases to ca. 100 K. As coverage decreases further, the vacancy patches become larger and by $\rho = 0.614$ the solid is a connected network of randomly oriented domains. All cases show a weak nematic mespohase. The melting temperature is $T_2 = 160 \text{ K} \pm 3 \text{ K}$ and falls to ca. 145 K by $\rho = 0.614$. The dynamics and energetics observed demonstrate that the explicit-hydrogen model of hexane is substantially more realistic than the UA approximation.

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