Calorimetric, X-Ray Diffraction, and Spectroscopic Studies of the Thermotropic Phase Behavior and Organization of Tetramyristoyl Cardiolipin Membranes

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ABSTRACT The thermotropic phase behavior and organization of aqueous dispersions of the quadruple-chained, anionic phospholipid tetramyristoyl diphosphatidylglycerol or tetramyristoyl cardiolipin (TMCL) was studied by differential scanning calorimetry, x-ray diffraction, 31P NMR, and Fourier-transform infrared (FTIR) spectroscopy. At physiological pH and ionic strength, our calorimetric studies indicate that fully equilibrated aqueous dispersions of TMCL exhibit two thermotropic phase transitions upon heating. The lower temperature transition is much less cooperative but of relatively high enthalpy and exhibits marked cooling hysteresis, whereas the higher temperature transition is much more cooperative and also exhibits a relatively high enthalpy but with no appreciable cooling hysteresis. Also, the properties of these two-phase transitions are sensitive to the ionic strength of the dispersing buffer. Our spectroscopic and x-ray diffraction data indicate that the lower temperature transition corresponds to a lamellar subgel (Lc) to gel (Lα) phase transition and the higher temperature endotherm to a Lα to lamellar liquid-crystalline (Lc) phase transition. At the Lc/Lα phase transition, there is a fivefold increase of the thickness of the interlamellar aqueous space from ~11 Å to ~50 Å, and this value decreases slightly at the Lα/Lβ phase transition. The bilayer thickness (i.e., the mean phosphate-phosphate distance across the bilayer) increases from 42.8 Å to 43.5 Å at the Lc/Lα phase transition, consistent with the loss of the hydrocarbon chain tilt of ~12°, and decreases to 37.8 Å at the Lα/Lβ phase transition. The calculated cross-sectional areas of the TMCL molecules are ~79 Å² and ~83 Å² in the Lc and Lα phases, respectively, and we estimate a value of ~100 Å² in the Lβ phase. The combination of x-ray and FTIR spectroscopic data indicate that in the Lc phase, TMCL molecules possess tilted all-trans hydrocarbon chains packed into an orthorhombic subcell in which the zig-zag planes of the chains are parallel, while in the Lα phase the untitled, all-trans hydrocarbon chains possess rotational mobility and are packed into a hexagonal subcell, as are the conformationally disordered hydrocarbon chains in the Lβ phase. Our FTIR spectroscopic results demonstrate that the four carbonyl groups of the TMCL molecule become progressively more hydrated as one proceeds from the Lc to the Lα and then to the Lβ phase, while the two phosphate moieties of the polar headgroup are comparably well hydrated in all three phases. Our 31P-NMR results indicate that although the polar headgroup retains some mobility in the Lc phase, its motion is much more restricted in the Lα and especially in the Lβ phase than that of other phospholipids. We can explain most of our experimental results on the basis of the weakly disordered nature of the polar headgroup of TMCL relative to other phospholipids and the covalent attachment of the two phosphate moieties to a single glycerol moiety, which results in a partially immobilized polar headgroup that is more exposed to the solvent than in other glycerophospholipids. Finally, we discuss the biological relevance of the unique properties of TMCL to the structure and function of cardiolipin-containing biological membranes.

INTRODUCTION

Cardiolipin (CL) or diphosphatidylglycerol is an anionic phospholipid occurring in the plasma membranes of many types of Gram-negative and Gram-positive bacteria and in the mitochondrial and chloroplast inner membranes of eukaryotes (1–5). CL is usually a relatively small component (≤10 mol %) of the total membrane lipid in such membranes and its primary role appears to be supporting the function of key membrane proteins (5). Specifically, CL plays an important role in the structural stabilization and activation of many mitochondrial enzymes, especially those involved in ATP synthesis and energy transduction (6–12), and in maintaining the structure and function of the Type II photoreaction center of photosynthetic bacteria and plants (13). In higher eukaryotes, CL is an effector of the cytochrome P-450-dependent cholesterol side-chain cleavage enzyme and activates cytochrome c oxidase and the mitochondrial phosphate carrier protein (14–16). It has been suggested that the capacity for structurally specific interactions with membrane CL may actually be highly conserved in such proteins (17). CL may also be found in substantially higher quantities (~55–60 mol %) in the membranes of some microbial organisms, wherein elevated levels of CL and its derivatives appear to enhance the tolerance of these organisms to halophilic and resource-depletion stress (5,18–21). However, it is not clear whether elevated CL levels reflect a requirement for the enhancement of the structural integrity of the cell membrane, for the support of stress-related increases in energy procurement and transduction, or both (12,18).

Unlike other naturally occurring glycerophospholipids, CL is a quadruple-chained amphiphile composed of two
phosphatidate moieties esterified to the 1- and 3-hydroxyl groups of a single glycerol molecule (referred to here as the headgroup glycerol). Consequently, one would expect the CL polar headgroup to be a relatively rigid, mobility-restricted entity, the cross-sectional area of which would be small relative to the combined cross-sectional areas of the four hydrocarbon chains. Thus, CL should be more prone to form inverted nonlamellar phases than phospholipids with larger headgroups such as phosphatidylcholine (PC) and phosphatidyglycerol (PG), although the formation of such inverted nonlamellar phases would be expected to be inhibited by the high surface charge of the CL bilayer (see (22)). Indeed, the lamellar phase of CL membranes only converts to the inverted hexagonal phase when the negatively charged phosphate groups at the surface of such bilayers are protonated at low pH or screened by dispersal in solutions of high ionic strength, or if divalent metal cations are added (23–26). It has been suggested that these properties of CL may be an important aspect of the capacity of certain microorganisms to regulate the lamellar/nonlamellar phase propensity of their membrane lipids (27,28).

Most of the studies that have been performed on CL model membranes have concentrated on characterizing its role in supporting the function of enzymes such as cytochrome c oxidase (6–10), and on a characterization of its pH, divalent metal, and salt-induced lamellar-nonlamellar phase behavior (23–26). As a result, relatively little is known about the overall thermotropic phase behavior, or about the detailed structure and organization, of CL bilayer membranes. However, some differential scanning calorimetry (DSC) studies have shown that the L_{\alpha}/L_{\beta} phase transition temperatures of CL bilayers are higher than those of the corresponding diacyl PGs (24,29,30), and are more sensitive to the nature and amounts of mono- and divalent cations present in the dispersal media (29,30). Also, Fourier transform infrared (FTIR) spectroscopic studies suggest that the headgroup glycerol of CL may be involved in strong intramolecular interactions, presumably with the phosphate entities of the polar headgroup (31). In this study, we present a rigorous and systematic DSC, x-ray diffraction, and FTIR and 31P-NMR spectroscopic characterization of the overall thermotropic phase behavior of tetramyristoyl cardiolipin (TMCL), and of the structure and organization of the various lamellar phases that TMCL can form at physiological pH and ionic strength.

MATERIALS AND METHODS

TMCL was obtained from Avanti Polar Lipids (Alabaster, AL) and was used without further purification. Samples were prepared for DSC under the following set of conditions. In one set of measurements, 4–5 mg of TMCL were dispersed in 600 \mu l of a buffer composed of 50 mM Tris, 150 mM NaCl, 5 mM EDTA, and 1 mM NaN_3, pH 7.4, by vigorous agitation at temperatures near 60°C, and a 500 \mu l aliquot was introduced into the Hastelloy capsule of a Calorimetry Sciences (Spanish Fork, UT) Multi-Cell DSC. Before initial data acquisition, the sample was preconditioned by heating to 50°C followed by cooling to −7°C at scanning rates near 60°C h⁻¹. Data acquisition scans were then performed at scanning rates near 10°C h⁻¹. Typically, the process first involves three cycles of heating and cooling between −7°C and 60°C. Subsequently, the samples were cooled to −30°C and incubated at that temperature for 12 h, then heated to 8°C and incubated at that temperature to ensure that all ice had completely melted. This low-temperature incubation protocol was repeated twice and the sample was then cooled to −7°C and a further set of data-acquisition runs, consisting of two cycles of heating and cooling between −7°C and 60°C, were performed. In another set of DSC measurements, −1 mg of lipid was dispersed in 1 ml of buffer composed of 100 mM sodium phosphate, 150 mM NaCl, and 5mM EDTA, pH 7.4, by methods similar to those described above. Subsequently, the sample was incubated at −20°C overnight and then allowed to thaw slowly at temperatures near 4°C. Subsequently, 0.5 ml of sample was loaded into a precooled sample cell of a Microlab (Northampton, MA) VP-DSC instrument and data were acquired between 1 and 55°C at heating and cooling rates near 10°C h⁻¹ and 60°C h⁻¹. The phosphate buffer was utilized to facilitate a direct comparison of our DSC and x-ray diffraction results with previous studies, but this buffer is not suitable for FTIR and 31P-NMR spectroscopic studies, because the presence of large amounts of phosphate in the buffer swamps the lipid phosphate signals, so the Tris buffer was utilized instead. Moreover, the use of two different buffers allowed us to investigate the effect of buffer composition and ionic strength on TMCL phase behavior.

FTIR spectroscopy was performed on samples containing 2–3 mg of TMCL. Samples were dispersed as described above in 100 \mu l of a D,O-based buffer containing 100 mM sodium phosphate and 150 mM NaCl, pD 7.4. The dispersion was squeezed between the CaF_2 windows of a heatable, demountable liquid cell (NSG Precision Cells, Farmingdale, NY) equipped with a 25 \mu M Teflon spacer. Once mounted in the sample holder of the spectrometer, the sample could be heated between −20°C and 90°C by an external, computer-controlled water bath. Before initial data acquisition, samples were subject to the same low-temperature incubation protocol described above for the DSC samples. Infrared spectra were then acquired as a function of temperature with a Digilab FTS-40 Fourier-transform Spectrometer (Bio-Rad, Digilab Division, Cambridge, MA) using data acquisition parameters similar to those described by Mantsch et al. (32). The experiment involved a sequential series of 2°C temperature ramps with a 20 min interramp delay for thermal equilibration, and was equivalent to an effective scanning rate of 4°C per hour. The data obtained were analyzed using computer programs obtained from the instrument manufacturer and from the National Research Council of Canada. In cases where absorption bands appeared to be a summation of components, a combination of Fourier deconvolution and curve-fitting procedures were used to obtain estimates of the position of the component bands and to reconstruct the contours of the original band envelope.

For the 31P-NMR spectroscopic measurements, 10–15 mg of TMCL were dispersed in 0.6 ml of a buffer composed of 100 mM Tris, 150 mM NaCl, 5 mM EDTA, and 1 mM NaN_3, pH 7.4. Before initial data acquisition, samples were subjected to the same low-temperature incubation protocol described above for the DSC samples. Spectra were acquired as a function of temperature with a Varian Unity-300 spectrometer (Varian, Palo Alto, CA) operating at 121.41 MHz for 31P. Spectra were acquired using previously published single-pulse data acquisition techniques and other data acquisition parameters (33).

Samples for x-ray diffraction measurements were also prepared by dispersing 5 mg of TMCL in 0.1 ml of both buffers (100 mM sodium phosphate, 150 mM NaCl, and 5 mM EDTA, pH 7.4; and 50 mM Tris, 150 mM NaCl, 5 mM EDTA, and 1 mM NaN_3, pH 7.4, respectively), by vigorous vortex mixing at temperatures near 60°C. Before initial data acquisition, samples were subjected to the same low-temperature incubation protocol described above for the DSC samples. X-ray diffractograms were recorded with a SWAX-camera (HECUS X-Ray Systems, Graz, Austria), which was mounted on a sealed-tube Seifert generator (Ahrensburg, Germany) operating at 2 kW (50 kV and 40 mA). The x-ray beam was filtered for CuKα radiation (\( \lambda = 1.54 \text { Å} \)) using an Ni foil and a pulse-height discriminator, built into the detection system. The scattered intensity was recorded with two linear
position-sensitive detectors (HECUS X-Ray Systems) simultaneously in the small- and wide-angle regimes of $10^{-3} \text{A}^{-1} < q < 1 \text{A}^{-1}$ (SAXS) and 1.2 $\text{A}^{-1} < q < 2.7 \text{A}^{-1}$ (WAXS), where $q = 4 \pi \sin(\theta)/\lambda$ is the modulus of the scattering wave vector. Calibration in the small-angle region was performed with silver stearate and in the wide-angle region with a p-bromo-benzoic acid standards, respectively. Samples were loaded into thin-walled quartz capillaries that were thermally equilibrated for 10 min before initiating data acquisition. Temperature was controlled with an accuracy of 0.1°C with a programmable Peltier unit. Diffraction patterns were recorded at each temperature for 2700 s.

Background-corrected SAXS patterns were analyzed in terms of a global analysis technique described previously (34–36). Briefly, the scattered intensity of randomly oriented bilayers

$$I(q) = \frac{S(q)|F(q)|^2}{q^4}$$

is separated into the contributions from the lamellar lattice, given by the structure factor $S(q)$ and the modulation of the electron density across the lipid bilayer, expressed by the form factor $F(q)$. The electron density profile, $\rho(z)$, is modeled as a simple summation of Gaussian peaks, where two Gaussians are located at the position of the phosphate group on each side of the bilayer and a third Gaussian of negative amplitude at the center of the bilayer. $F(q)$ is the Fourier transform of $\rho(z)$ and contains an adjustable parameter, $d_{PP}$, as a measure for the membrane thickness. From $d_{PP}$, the phosphate-phosphate distance across the bilayer, we further calculate the hydrocarbon chain length by the equation

$$d_C = (d_{PP}/2) - d_{H1},$$

where $d_{H1}$ is the distance from the phosphate group to the polar/apolar interface (37). With PC bilayers, Nagle and co-workers have typically reported values of $\approx 4.95 \text{Å}$ for $d_{H1}$ (38). However, Fabst et al. (37) have shown that a $d_{H1}$ value of $\approx 4 \text{Å}$ gives a good overall agreement between the $d_C$ values obtained by Nagle’s group and the values derived from the global data analysis technique. This is because the global analysis technique models the lipid headgroup with a single peak, whereas Nagle and co-workers model the PC headgroup with two contributions, one for the phosphate group and one for the choline group. The latter reports slightly larger $d_{PP}$ values than those derived by the global analysis technique used here, which has compensated for by a smaller $d_{H1}$ value. The $d_{H1}$ value used for these studies (4 Å) assumes that the distance between the phosphate group and the polar/apolar interface of TMCL bilayers is comparable to those of PC bilayers, which seems to be justified by our results (see below). The hydrocarbon-chain tilt angle ($\theta_t$) was estimated from the $d_C$ values by the equation

$$d_C = d_C^0 \cos(\theta_t),$$

where $d_C^0$ refers to nontilted hydrocarbon chain length, as approximated by the product of the distance between two C-bonds (1.27 Å, (39)) and the number of hydrocarbons per chain, which yields a value of $\approx 17.8 \text{Å}$. The projected lateral area ($A$) per TMCL molecule was estimated from the hydrocarbon-chain tilt angle and the per chain area ($A_C$) by the following equation:

$$A = \frac{4A_C}{\cos \theta_t}.$$
of TMCL exhibit two major heating endothermic transitions. The lower-temperature transition is less cooperative than the higher temperature transition but the energetics of the two processes are comparable in magnitude. However, the thermodynamic properties of these two transitions differ in the two buffer systems and with the instrument used. Thus, for example, at a scanning rate of 10° C h⁻¹ the temperatures, enthalpies, and widths (at half-height) of the lower-temperature transition obtained using the Tris buffer in the Multi-Cell DSC are 24.9°C, 14.8 kcal/mol, and 1.2°C. In the case of the higher-temperature transition, the temperatures, enthalpies, and widths recorded with the Tris buffer and Multi-Cell DSC are 38.9°C, 12.9 kcal/mol, and 1.5°C, whereas the corresponding values obtained with phosphate buffer in the VP-DSC are 41.2°C, 13.5 kcal/mol, and 0.3°C, respectively. Nevertheless, when examined in the high-sensitivity power-compensation instrument (i.e., the Microcal VP-DSC), the properties of the latter transition are relatively insensitive to the scanning rate in the range 10°C–60°C h⁻¹ (data not shown). Using data obtained in our x-ray-diffraction and spectroscopic studies (see below), we have assigned the lower-temperature event to a subgel-to-gel (L_c/L_g) transition and the higher-temperature event to a gel-to-liquid-crystalline (L_β/L_α) phase transition involving the melting of lipid hydrocarbon chains.

Figs. 1 and 2 also show that upon cooling, the conversion of the L_α phase of TMCL to the L_β phase may be associated with either one (phosphate buffer) or two (Tris buffer) cooling exotherms, depending on the nature of the dispersal buffer. These exothermic events are observed at temperatures only slightly below those observed upon heating, and have been identified as interconversions between the L_α and L_β phases of this lipid on the basis of our FTIR spectroscopic and x-ray diffraction results (data not shown). Multiple cooling exotherms at the L_α/L_β phase transitions have been observed with some mixed-chain PCs (42) and cationic phospholipids (43). This phenomenon usually arises from heterogeneity in the domain sizes from which the L_β is nucleated (42,44). Thus, the differences between the cooling exotherms observed in Tris and phosphate buffer is probably reflecting ionic strength-induced variations in the sizes of the domains formed in the L_α phase. Unlike the L_α/L_β phase transition of TMCL, conversion of the L_β phase to the L_c phase exhibits a pronounced cooling hysteresis, being initiated at temperatures some 10°C below the onset of the heating endotherm (Figs. 1 B and 2 B) and, as indicated by the considerably smaller enthalpy change observed (~2.5 kcal/mol), this phase transition is not completed during the cooling scan. Interestingly, when aqueous dispersions of TMCL are reheated within a few hours after cooling to temperatures near 0°C, one usually observes two low-temperature endotherms centered near 14–18°C and 25–30°C (see Figs. 1 C and 2 C). Moreover, the combined enthalpy of these two peaks (~6–8 kcal/mol) is usually smaller than that of the single L_c/L_β single phase transition observed upon heating the fully equilibrated samples. Moreover, the size of the peak near 25–30°C seems to increase at the expense of the lower-temperature peak upon incubation at temperatures near 0°C. The latter observation suggests that the formation of the stable L_c phase formed using the low-temperature incubation protocols described in the Materials and Methods could be proceeding via the formation of an intermediate subgel phase. However, the results of our time-resolved FTIR spectroscopic studies (not shown here) showed no evidence for the formation of such an intermediate L_c phase.

X-ray diffraction

Illustrated in Fig. 3 A are the SAXS profiles exhibited by TMCL in the sodium phosphate buffer system at temperatures bracketing the calorimetrically-resolved L_c/L_β and L_β/L_α phase transitions. The bilayer properties calculated from these x-ray-diffraction data are summarized in Table 1 and Fig. 4. At temperatures near 2°C, TMCL exhibits small-angle diffractionograms containing a sharp, resolution-limited, first-order Bragg peak. Aside from the fourth-order peak, the higher-order diffraction peaks are barely visible, which is
analogous to diffraction patterns displayed by phosphatidylethanolamines (PEs) (45). The global data analysis gives a good overall fit to these data except for the fourth-order peak. This can be attributed to simplifications involved in polycrystalline theory (36), but has no significant affect on the structural parameters obtained. As reported in Table 1, we find that at 2°C, TMCL forms a multibilayer system with a lamellar repeat distance \( d \) of 54.2 Å, where the individual layers of thickness \( d_{\text{PP}} \); 42.8 Å are separated by a small water layer \( d_W \); 11 Å, again very much like the PEs (45). At temperatures near 35°C, TMCL exhibits two SAXS peaks that can be indexed as the first and second lamellar diffraction order. The peaks appear to be considerably broader than at 2°C, which is due to an increase of the stacking fluctuations from \( \sigma_{\text{PT}} \) (2°C) = 3.7 Å to \( \sigma_{\text{PT}} \) (35°C) = 10.3 Å. The central peak width given by the scattering domain size remains unaffected. The approximately three-fold increase of \( \sigma_{\text{PT}} \) manifests itself also in a larger \( d \) of \( \approx 93.3 \) Å. The observed increase of \( d_{\text{PP}} \) of \( \approx 43.5 \) Å is much less pronounced, and hence the increase of \( d \) is largely due to a swelling of the water layer by \( \approx 40 \) Å. Finally, at temperatures above the higher-temperature phase transition, aqueous TMCL exhibits Bragg peaks that index on a lamellar lattice with a \( d \) of \( \approx 83 \) Å at 55°C. The membrane thickness \( d_{\text{PP}} \); 37.8 Å is now significantly smaller than observed at lower temperatures, as expected for the melting of the hydrocarbon chains. The bilayer separation has decreased slightly to \( \approx 45 \) Å, but is still significantly larger than in uncharged lipid/water systems, which is also reflected in the fluctuation parameter \( \eta \) (Table 1), which is approximately three times the value exhibited in phosphatidycholine (PC) dispersions (36). Comparable x-ray diffraction experiments were also performed with TMCL in the lower ionic strength Tris buffer (not shown). Our analyses of the data obtained in those experiments indicate that the structural parameters such as bilayer thickness and hydrocarbon chain packing were the same as those obtained in phosphate buffer. However, the positional correlation of the bilayers decreased in the Tris buffer, resulting in broader Bragg peaks and an increase in the thickness of the water layer between the

![FIGURE 3 Scattered x-ray intensity of TMCL in small-angle (A) and wide-angle regime (B) at three temperatures in the L_c (2°C), L_p (35°C), and L_a (55°C) phase. Data shown were acquired during heating using samples that were initially fully equilibrated at low temperatures. Arrows in panel A indicate the position of the first-order Bragg peak except for the pattern at 2°C, which has an additional arrow marking the fourth diffraction order. Solid lines show the best fits to the SAXS data applying a global analysis technique (see Materials and Methods). Bracketed numbers in panel B give the Miller indices of the corresponding Bragg rods.](image1)

![FIGURE 4 Low-resolution, one-dimensional electron density profiles along the bilayer normal calculated from the small-angle scattering diffraction patterns exhibited by aqueous dispersions of TMCL. Electron density profiles were calculated for (A) L_c phase at 2°C; (B) L_p phase at 35°C; and (C) L_a phase at 55°C.](image2)

### TABLE 1 Bilayer properties calculated for the three polymorphic phases exhibited by aqueous dispersions of TMCL

<table>
<thead>
<tr>
<th>( T^\circ ) (°C)</th>
<th>( d ) (Å)</th>
<th>( d_{\text{PP}} ) (Å)</th>
<th>( d_c ) (Å)</th>
<th>( d_W ) (Å)</th>
<th>( \sigma_{\text{PT}} ) (Å)</th>
<th>( \eta )</th>
<th>( A_c ) (Å)</th>
<th>( \Delta ) (Å)*</th>
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<td>0.305</td>
<td>—</td>
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*For abbreviations, see text.

TMCL: 100 mM sodium phosphate, 150 mM NaCl, and 5 mM EDTA, pH 7.4.
bilayers (i.e., larger \(d\)-values). The latter is consistent with the expected effects of increased electrostatic repulsion under those experimental conditions.

Illustrated in Fig. 3B are the patterns of WAXS reflections exhibited by the three polymorphic phases of TMCL. The pattern of diffraction peaks observed in this region arises from the two-dimensional packing of the lipid fatty acyl chains. At temperatures near 2°C, the WAXS diffractogram contains a relatively sharp Bragg peak centered near 4.6 Å and a broader Bragg reflection centered near 3.8 Å, suggesting that the latter arises from an off-equatorial reflection typical for hydrocarbon chains being tilted with respect to the bilayer plane (46). These peaks were assigned to the (2 0) and (1 1) spacings of an orthorhombic subcell with lattice parameters \(a = 9.2\) Å, \(b = 4.2\) Å, and a per-chain area of 19.3 Å² (see Table 1). From the SAXS data we estimate a hydrocarbon-chain tilt angle of some 12°, from which we obtain a projected lateral area per TMCL molecule of some 83.2 Å² (46,47). For this phase the projected lateral area (283.2 Å²) is simply four times the \(A_c\) value and is only slightly larger than obtained for the \(L_c^9\) phase at 2°C (Table 1). We also note that the \(d_c\) value obtained from the analysis of the SAXS data (~17.8 Å) is in excellent agreement with our estimates of the untilted length of TMCL all-trans hydrocarbon chains, consistent with the conclusion that TMCL hydrocarbon chains are untilted at temperatures between the two phase transitions. The increase of the membrane thickness across the lower-temperature transition can thus be entirely explained by the removal of the chain tilt. Finally, at temperatures above the higher-temperature transition, TMCL exhibits a broad diffuse WAXS peak, consistent with the low degree of lateral packing order, which occurs in highly mobile melted hydrocarbon chains (i.e., the \(L_a\) phase). However, in the absence of dilatometric measurements we were unable to determine the lateral area per molecule in the \(L_a\) phase. Nevertheless, if the area expansion across the hydrocarbon chain-melting phase transition of TMCL is of comparable magnitude to that observed with PCs (~25%, see (48)), the projected lateral area should be near 100 Å².

Fourier-transform infrared spectroscopy

Illustrated in Fig. 5 are the regions of the C–H stretching, C–O stretching, CH₂ scissoring, and asymmetric phosphate O–P–O stretching bands of the FTIR spectra exhibited by aqueous dispersions of TMCL at temperatures bracketing the two thermotropic phase transitions reported by DSC. These regions of the spectra encode information about hydrocarbon chain conformation, bilayer interfacial conformation and hydration, hydrocarbon chain packing interactions, and the polarity of the phosphate polar headgroups, respectively. It is clear from even a cursory examination of these spectra that the two phase transitions detected by DSC are structurally distinct thermotropic events. Thus, for example, in the C–H stretching region (2800–3000 cm⁻¹), the changes accompanying the lower-temperature transition centered near 25–30°C involve relatively small decreases in the intensities of all absorption bands, whereas the changes accompanying the

![Figures A-D](https://example.com/figures.png)

**Figure 5:** FTIR spectra observed upon heating fully equilibrated aqueous dispersions of TMCL. The absorbance spectra shown were acquired at the temperatures indicated and typify the C–H stretching (A), the C–O stretching (B), the CH₂ scissoring (C), and the asymmetric phosphate stretching (D) regions of the IR spectra of this compound at temperatures bracketing the two calorimetrically resolved thermotropic phase transitions. The vertical dashed line (A) marks the position of the maxima of CH₂ symmetric stretching band observed in the in the three polymorphic forms of this lipid. The dashed lines (B) represent the subcomponents of the observed absorption band. The asterisks (D) mark the positions of the peaks of the CH₂ wagging band progression.
higher-temperature transition near 39–42°C are considerably larger and involve significant increases in both band frequency and bandwidth (Fig. 5 A), indicating that the major changes in hydrocarbon chain conformation coincide with the higher-temperature event. In contrast, the C=O stretching, CH$_2$ scissoring, and asymmetric phosphate stretching regions of the FTIR spectrum exhibit major changes in band frequency, band shape, and band width at the lower-temperature transition and considerably smaller changes at the higher-temperature transition (Fig. 5, B–D), indicating that the major changes in interfacial conformation and hydration, hydrocarbon chain packing, and polar headgroup polarity occur at the lower-temperature transition. A more detailed examination and structural interpretation of the spectroscopic changes associated with these two phase transitions is presented below.

In the C-H stretching region of the FTIR spectrum, the dominant features are the absorption bands centered near 2850 cm$^{-1}$ and 2915 cm$^{-1}$ (Fig. 5 A). These bands arise from the symmetric and asymmetric stretching vibrations of CH$_2$ groups of the lipid hydrocarbon chains, respectively (49,50). In particular, the frequency of the symmetric stretching band is determined by the trans/gauche rotamer content of the lipid hydrocarbon chains and is therefore an excellent spectroscopic marker for changes in the conformation of these chains (49,50). At temperatures below 20°C, fully equilibrated TMCL samples exhibit sharp symmetric CH$_2$ stretching bands at frequencies centered near 2850 cm$^{-1}$ (Fig. 1 A), values consistent with the existence of predominantly all-trans polyethylene chains. Upon heating to temperatures near the L$_c$/L$_b$ phase transition of this lipid, this band broadens slightly without any significant change in its peak frequency, indicating that no gauche conformers are induced at the lower-temperature transition. However, at the L$_b$/L$_a$ phase transition temperature, we see discontinuous increases in its frequency and width, indicating significant increases in the gauche rotamer content and overall mobility of the lipid hydrocarbon chains. These results clearly indicate that the lower-temperature transition involves a solid phase rearrangement of the TMCL molecules with all-trans lipid hydrocarbon chains, whereas the higher-temperature transition involves the melting of the TMCL hydrocarbon chains.

In the 1400–1500 cm$^{-1}$ region of the IR spectrum, the dominant feature is the absorption band between 1466 cm$^{-1}$ and 1473 cm$^{-1}$, which arises from the scissoring vibrations of the hydrocarbon chain methylene groups (see Fig. 5 C). The properties of this band are sensitive to the lateral packing interactions between hydrocarbon chains (49,50). At low temperatures, the CH$_2$ scissoring band of TMCL appears as a single, sharp absorption peak centered near 1473 cm$^{-1}$, suggesting that the all-trans lipid chains are packed in a subcell with their zigzag planes parallel. When heated to temperatures in between the L$_c$/L$_b$ and L$_b$/L$_a$ phase transitions, this CH$_2$ scissoring band broadens and its peak frequency shifts to values near 1468 cm$^{-1}$ (Fig. 5 C), suggesting that the hydrocarbon chains are more mobile and packed in a hexagonal subcell. Moreover, since the spectroscopic properties exhibited in the C-H stretching region indicate that the hydrocarbon chains are still in the all-trans conformation (see previous paragraph), we can also conclude that the hydrocarbon chains are rotationally disordered, most probably because of an increase in the rates and amplitude of the reorientational fluctuations of the all-trans polyethylene chains at the L$_c$/L$_b$ phase transition. Finally, when heated to temperatures above the L$_b$/L$_a$ phase transition, the CH$_2$ scissoring band of TMCL broadens considerably without a significant change in its peak frequency (see Fig. 5 C). Since the contours of the C-H stretching regions indicate that the hydrocarbon chains are melted at these temperatures (see previous paragraph), we can conclude that the liquid-crystalline phase contains hexagonally arranged, conformationally and rotationally disordered hydrocarbon chains.

The spectra shown in Fig. 5 B also indicate that the major changes in the C=O stretching region between 1650 cm$^{-1}$ and 1800 cm$^{-1}$ also occur mainly at the L$_c$/L$_b$ phase transition. Since the infrared absorption in this region arises exclusively from the stretching vibrations of the ester carbonyl groups of TMCL, these bands can provide valuable information about the structure and properties of the polar/apolar interfacial regions located near the backbone glycerol moiety of the three polymorphic forms of this lipid (49,50). At temperatures below the onset of the L$_c$/L$_b$ phase transition, the C=O stretching band seems to be a summation of a relatively sharp band centered near 1735 cm$^{-1}$ and a broader band of lesser intensity centered near 1716 cm$^{-1}$ (see dashed curves in Fig. 5 B). These bands arise from the stretching vibrations of subpopulations of free and hydrogen-bonded ester carbonyl groups, respectively (51,52). At temperatures between the L$_c$/L$_b$ and L$_b$/L$_a$ phase transitions, the C=O ester stretching band is also resolvable into two components (see dashed curves in Fig. 5 B). However, at these temperatures the component bands are of comparable intensity, are broader, and are centered at frequencies near 1743 cm$^{-1}$ and 1728 cm$^{-1}$, respectively. These latter two components are also found at temperatures above the L$_b$/L$_a$ phase transition of the lipid, but at those temperatures the lower-frequency component is even broader and is the major contributor to the integrated intensity of the observed C=O stretching band (see dashed curves in Fig. 5 B). These results indicate that there is a progressive increase in the degree of hydrogen bonding of the fatty acid carbonyl groups, resulting from an increased penetration of water into the glycerol backbone region of the TMCL molecules, as one proceeds from the L$_c$ through the L$_b$ into the L$_a$ phase.

An examination of the region encompassing the phosphate O-P-O asymmetric stretching band also indicates that fairly drastic changes occur at the L$_c$/L$_b$ phase transition with relatively minor changes occurring at hydrocarbon chain-melting L$_b$/L$_a$ phase transition temperature (Fig. 5 D). At
low temperatures, the broad contours of the phosphate O-P-O asymmetric stretching band are largely obscured by several sharp, relatively weak absorption bands arising from the CH₂ wagging band progression (see marked bands in Fig. 5 D). These bands arise from the coupled CH₂ wagging vibrations of methylene groups on all-trans polymethylene chains, and are usually quite sharp when the rates and amplitudes of chain reorientational fluctuations are low (49,50). However, despite the interference from these bands, the presence of the broad underlying absorption of the phosphate O-P-O asymmetric band is obvious and its peak seems to be centered near 1216 cm⁻¹. Upon heating to temperatures in between the Lₐ/Lₐ phase and Lₐ/Lₐ phase transitions, the CH₂ wagging band progression largely disappears and the peak of the phosphate O-P-O asymmetric stretching is more easily resolved at progression largely disappears and the peak of the phosphate O-P-O asymmetric stretching vibrations, consistent with the phosphate O-P-O asymmetric stretching band narrows slightly, but its peak frequency remains near 1216 cm⁻¹. This frequency is near the lower end of the range expected of phosphate O-P-O asymmetric stretching vibrations, consistent with the phosphate groups being well hydrated and/or bonded to hydrogen-bonding donor groups.

**3¹P-Nuclear magnetic resonance spectroscopy**

Illustrated in Fig. 6 are the 3¹P-NMR powder patterns acquired at temperatures bracketing the two thermotropic phase transitions observed by DSC upon heating after prolonged sample incubation at low temperatures. At temperatures below the onset of the lower-temperature transition, fully equilibrated samples of aqueous TMCL exhibit broad 3¹P-NMR powder patterns with basal line widths near 250 ppm (spectrum A), characteristic of the so-called rigid-limit spectra expected of the crystalline forms of phospholipids (53,54). This suggests that TMCL forms an Lᶜ phase in which reorientation of the phosphate moieties of the TMCL headgroup is extremely slow on the 3¹P-NMR timescale. We also note that despite the effective “immobilization” of the phosphate headgroups, 3¹P-NMR spectra of the Lᶜ phase of TMCL were easily acquired using simple single-pulse data acquisition techniques. This situation differs markedly from what occurs when similar techniques are used to record the 3¹P-NMR spectra of the subgel phases of phospholipids such as dimyristoyl phosphatidylethanolamine and dimyristoyl phosphatidylglycerol, for which the extremely long longitudinal relaxation times of the 3¹P nucleus makes the recording of their Lᶜ phase spectra impractical (55,56). Given that such slow longitudinal relaxation of the 3¹P nucleus have been attributed to long-lived H-bonding interactions between the phosphate and appropriate H-bond donor groups (55,56), it appears that the phosphate moieties of the polar headgroup of TMCL may not be involved in long-lived H-bonding interactions with either solvent molecules of other potential H-bonding donors in their Lᶜ phase.

When heated to temperatures between the completion and onset temperatures of its lower-temperature and higher-temperature thermotropic phase transitions, significant narrowing of the 3¹P-NMR powder patterns of TMCL occurs (Fig. 6, spectrum B). However, the 3¹P-NMR powder patterns observed in this phase are themselves quite broad (basal line widths ~140 ppm), and their widths and contours are atypical of those normally exhibited by the Lₐ-like lamellar gel phases of other 1,2-diacyl glycerophospholipids (for examples, see (32,33,55,56)). This observation suggests that in the Lₐ phase of this lipid, reorientation of its headgroup phosphate moieties is either significantly slower than that of the Lₐ phases of most phospholipid bilayers, or that the range and amplitude of reorientational motion accessible to phosphate groups in the TMCL Lₐ phase are much more restricted than is typical of phospholipid Lₐ phases, or both.

Finally, at temperatures above its Lₐ/Lₐ phase transition temperature, TMCL exhibits 3¹P-NMR powder patterns that show some of the features normally observed in the Lₐ phases of most other phospholipids (Fig. 4, spectrum C). However, the apparent chemical shift anisotropy (~25 ppm) of fluid TMCL dispersions is considerably smaller than that of the Lₐ phases of phospholipids such as PC, PE, and PG, for which values near 35–40 ppm are commonly observed (see (32,33,55,56)). This observation suggests that even in its Lₐ phase, the phosphate moieties of TMCL may be subject to motional restrictions that are not normally observed in most other phospholipids.
DISCUSSION

A number of new and important conclusions about the thermotropic phase behavior and organization of TMCL bilayer membranes can be drawn from the calorimetric, x-ray diffraction, and spectroscopic data presented here. For example, we demonstrate that the overall thermotropic behavior of TMCL is strongly influenced by the nature of the dispersal medium, especially the temperature at which its thermotropic phase transitions are observed. Such behavior is commonly observed with anionic phospholipid membranes (see (57–59)) and was thus generally expected. However, the data presented here and elsewhere (see (29,30,60)) indicates that CL membranes are more sensitive to such effects than membranes composed of other anionic phospholipids such as PG and phosphatidylserine. This observation can probably be attributed to the reduced shielding of the negatively charged phosphate groups on the surface of CL membranes by the single headgroup glycerol moiety to which both phosphates are covalently linked. This is because the relative rigidity and restricted mobility of the CL headgroup effectively reduces its capacity for intra- and intermolecular interaction with other CL phosphate groups, and as a result negative charges at the surfaces of CL bilayers will not be as shielded from interaction with the solvent and its dissolved ions, as would be the case with anionic lipids such as PG and phosphatidylserine. In this respect, the properties of CL bilayers may actually be more comparable to those of bilayers composed of the anionic glycerophospholipid phosphatic acid (PA), which is also very sensitive to the properties of the dispersal medium, presumably because its negatively charged phosphate headgroups are not shielded from interaction with the water and ions.

The calorimetric data presented here is in general agreement with previous calorimetric studies (29,30) reporting that TMCL exhibits a L_β/L α phase transition temperature (~40°C) that is markedly higher than that of the corresponding diacyl PG (~23°C (61)). Given the low capacity for shielding of the phosphate groups by the CL headgroup glycerol (see above), the relatively high L_β/L α phase transition temperature of TMCL can probably be rationalized by the combined effects of the relatively small size of its polar headgroup and the existence of a H-bonding and/or a cation interaction network, as postulated for PA bilayers (62). Our calorimetric results also indicate an enthalpy for the TMCL L_β/L α phase transition (12.9 kcal/mol) that is slightly higher but generally compatible with the range of values (~11–12.5 kcal/mol) reported in the literature under comparable conditions (29,30). Also, the per chain normalized enthalpy (~3.2 kcal/mol) is compatible with those observed at the L_β/L α type phase transitions of most phospho- and glycolipids that have been examined so far. Our calorimetric measurements are thus consistent with the assignment of the TMCL L_β/L α phase transition as a simple hydrocarbon chain-melting event like those observed in most naturally occurring glycerolipids.

These studies have also shown that TMCL readily forms a highly ordered lamellar crystal-like (L_c) phase when cooled to temperatures below 10°C. To our knowledge, this aspect of the polymorphic phase behavior of CL membranes has not been reported previously. Our spectroscopic and x-ray diffraction data also indicate that although the temperature at which the L_β/L α phase transition occurs varies with the nature of the dispersal medium, the structural properties of the L_c phase are essentially the same under all conditions explored here. The structural picture of the TMCL L_c phase that emerges from these studies is as follows. In the hydrophobic domain, the closely packed all-trans hydrocarbon chains of TMCL are slightly tilted (~12°) to the bilayer surface and seem to be arranged in an orthorhombic subcell (x-ray diffraction data) with their zigzag planes parallel to each other (FTIR spectroscopic data). In the bilayer polar/apolar interfacial region near the backbone glycerol, there appears to be two populations of ester carbonyl groups (FTIR spectroscopic data). One of these gives rise to a sharp, higher-frequency C=O stretching band consistent with its being fairly rigid and located in a relatively nonpolar environment free of H-bonding interactions with either water or other H-bonding donor groups. The other population gives rise to a broad, lower-frequency C=O stretching band, and is probably located in a more polar and more heterogeneous environment where it is interacting via H-bonding with water or other donor groups. Given the overall low mobility of CL molecules in the L_c phase, the heterogeneity underlying the broad lower-frequency ester C=O band probably arises from heterogeneity in the orientation of H-bonding donor groups. This suggests that the source of the H-bonding donor groups is external to the CL molecule itself and is most probably water molecules in the dispersal medium. Our studies also indicate that the two phosphate moieties of the TMCL molecule are essentially immobilized on the 31P-NMR timescale in the L_c phase. However, the frequency of the phosphate O-P-O asymmetric stretching band (FTIR spectroscopic data) also suggests that these phosphate moieties are well hydrated. This latter conclusion is supported by the x-ray diffraction results, which show that the L_c phase lamellar repeat distance (~54 Å) is significantly greater than the phosphate-phosphate bilayer thickness (~43 Å), consistent with the presence of an aqueous interlamellar space of ~11 Å. Also, given the relative ease with which 31P-NMR spectra of the L_c phase were acquired, it is unlikely that the phosphate moieties are involved in strong, long-lived H-bonding interactions in this phase. Thus, interactions involving these negatively charged phosphate groups at the surface of the L_c phase of TMCL are probably better defined by hydrogen bonding to water and by Gouy-Chapman type interactions with counterions (for references on the latter, see (55)).

The phosphate-phosphate distance (~43 Å) and the hydrocarbon chain length (~17. 4 Å) calculated for the L_c phase of TMCL also provide some insight into the possible
conformations of the lipid molecules in the bilayer. There are two extreme conformations for the alignment of the backbone glycerols of TMCL, one along the bilayer normal as observed in single-crystal x-ray studies of PC and PE (63,64), and one roughly parallel to the bilayer surface as observed in single-crystal x-ray studies of PG and PA (62,65). The latter conformation can be excluded because the length of two opposed untitled hydrocarbon chains (~35 Å, see Table 1) and the mean thickness of the headgroup and backbone moieties relative to the bilayer normal (~2.5 Å, our estimates) are together too short to account for the observed P-P distance of ~43 Å. However, our experimental data can accommodate a conformation in which the TMCL backbone glycerols are oriented similar to PE and PC, with a slight tilting of the hydrocarbon chains. Such a conformation is obviously favorable for an optimal intramolecular packing of the hydrocarbon chains in the $L'_c$ phase and is also compatible with the results of our x-ray data obtained in the $L_b$ phase.

Our studies have also provided a fairly clear picture of the structures and organization of the $L_b$ and $L_a$ phases of TMCL and of the structural changes occurring at the calorimetrically resolved thermotropic phase transitions. When the $L_b$ gel phase of TMCL is cooled to low temperatures, the formation of the $L'_c$ phase is probably initiated by a marked but progressive reduction in the amount of water present in the interlamellar space and bilayer polar/apolar interfaces, as indicated by the decrease of the lamellar repeat distance upon cooling (data not shown). Initially, this process probably involves the formation of relatively small domains of the $L'_c$ phase that reconvert to the $L_b$ gel phase at temperatures between 10 and 20°C. This assumption is supported by WAXS data, which showed only hardly discernible Bragg peaks immediately after cooling below the $L'_c/L_b$ phase transition (data not shown). However, with prolonged low-temperature annealing, the growth of $L'_c$ phase continues and the small domains eventually coalesce into the extensive lattices that exhibit the structural properties described above. Upon heating, these structures reconvert to the $L_b$ gel phase at temperatures between 25 and 30°C. This process is accompanied by a modest increase in the hydration of the bilayer polar/apolar interfaces but by a considerable swelling of the interlamellar spaces. We suggest that such a large increase of the bilayer separation can only occur through a change of the membrane surface charge density, probably through the release of counterions. There is also an increase in the rates and amplitudes of reorientational fluctuations of the hydrocarbon chain segments, which are reflected by a decrease within close-contact, coupled interactions between vibrating CH$_2$ groups (FTIR data), caused by a distortion of the orthorhombic lattices formed at low temperatures (x-ray diffraction data). The $^{31}$P-NMR spectroscopic data also suggest that although there is some increase in the mobility of the phosphate moieties when the $L'_c$ phase converts to the $L_b$ phase, the rates of such motions remain slow on the $^{31}$P-NMR timescale, presumably because of the restrictions imposed by the tethering of the phosphate moieties to the headgroup glycerol. Finally, upon conversion to the $L_a$ phase, there is a cooperative melting of the hydrocarbon chains which is accompanied by an increase in the hydration of the bilayer polar/apolar interfaces, a small decrease in the thickness of the lipid bilayer and the interlamellar spaces, and a considerable increase in the rates and amplitudes of motion in all regions of the lipid bilayer. In this phase, the motions of phosphate moieties are also fast on the $^{31}$P-NMR timescale, but the range and amplitudes of such motions are restricted, presumably because of the tethering effect alluded to above.

In conclusion, we note that virtually all of the physical characteristics of CL bilayers as described in this and other studies can be directly or indirectly attributed to the restrictions placed on the flexibility and mobility of its polar headgroup by the tethering of the two phosphatidate moieties to a single glycerol molecule. As noted above, the low flexibility and impaired reorientational mobility of the CL headgroup diminishes its capacity for self-shielding of its negatively charged phosphate moieties and should therefore make CL-containing membranes more sensitive to interactions with the buffer. This property could be of considerable biological importance. Also, because both the intrinsic and effective sizes of the CL headgroup are small relative to the cross-sectional area of its hydrocarbon chains, CL can form inverted nonlamellar phases when its headgroup charges are either neutralized or effectively screened (22–26). This property may be biologically important because charge neutralization seems to be an important aspect of the interaction between CL and many of the CL-requiring membrane enzymes (9–10). These structural properties also seem to increase the structural integrity of CL membranes (note its relatively high $L_b$/$L_a$ phase transition temperature), presumably in part by imparting greater cohesion to the headgroup and polar/apolar interfacial regions of the membrane. In principle, these properties could explain why some organisms actually increase the CL content of their membranes under conditions of chronic resource depletion and halophilic stress (5,18–21). Moreover, these same properties may also explain why the penetration into the polar/apolar interface of CL membranes by the antibiotic peptide gramicidin S is not as efficacious as observed with membranes composed of other anionic lipids (66), and why CL-containing membranes are more resistant to the effects of some antimicrobial peptides than are membranes containing other negatively charged lipids (67–69). Thus, aside from its specific roles in supporting the function of key membrane enzymes, it also appears that CL confers upon cell membranes various physical properties that may be physiologically important.

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