Effect of Variations in the Structure of a Polyleucine-Based α -Helical Transmembrane Peptide on Its Interaction with Phosphatidylethanolamine Bilayers

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ABSTRACT High-sensitivity differential scanning calorimetry and Fourier transform infrared spectroscopy were used to study the interaction of a cationic α -helical transmembrane peptide, acetyl-Lys₂-Leu₂₄-Lys₂-amide (L₂₄), and members of the homologous series of zwitterionic n-saturated diacyl phosphatidylethanolamines (PEs). Analogs of L_{24} , in which the lysine residues were replaced by 2,3-diaminopropionic acid (acetyl-DAP₂-Leu₂₄-DAP₂-amide (L₂₄DAP)) or in which a leucine residue at each end of the polyleucine sequence was replaced by a tryptophan $(Ac-K_2-W-L_{22}-W-K_2-amide (WL_{22}W))$, were also studied to investigate the roles of lysine side-chain snorkeling and aromatic side-chain interactions with the interfacial region of phospholipid bilayers. The gel/liquid-crystalline phase transition temperature of the PE bilayers is altered by these peptides in a hydrophobic mismatch-independent manner, in contrast to the hydrophobic mismatch-dependent manner observed previously with zwitterionic phosphatidylcholine (PC) and anionic phosphatidylglycerol (PG) bilayers. Moreover, all three peptides reduce the phase transition temperature to a greater extent in PE bilayers than in PC and PG bilayers, indicating a greater disruption of PE gel-phase bilayer organization. Moreover, the lysine-anchored L_{24} reduces the phase transition temperature, enthalpy, and the cooperativity of PE bilayers to a much greater extent than DAP-anchored $L_{24}DAP$, whereas replacement of the terminal leucines by tryptophan residues (Ac-K₂-W-L₂₂-W-K₂-amide) only slightly attenuates the effects of this peptide on the chain-melting phase transition of the host PE bilayers. All three peptides form very stable α -helices in PE bilayers, but small conformational changes occur in response to mismatch between peptide hydrophobic length and gel-state lipid bilayer hydrophobic thickness. These results suggest that the lysine snorkeling plays a significant role in the peptide-PE interactions and that cation- π -interactions between lysine and tryptophan residues may modulate these interactions. Altogether, these results suggest that the lipid-peptide interactions are affected not only by the hydrophobic mismatch between these peptides and the host lipid bilayer but also by the electrostatic and hydrogen-bonding interactions between the positively charged lysine residues at the termini of these peptides and the polar headgroups of PE bilayers.

INTRODUCTION

All biological membranes consist of lipid and protein molecules held together by noncovalent interactions (Yeagle, 1993). The fundamental structural element of all biological membranes is a lipid bilayer, which also functions as the major permeability barrier. The zwitterionic phospholipids phosphatidylcholine (PC) and phosphatidylethanolamine (PE) are the major lipid components in animal cell membranes, and PE is often a major component of bacterial cell membranes as well (Daum, 1985; Devaux and Seigneuret, 1985). In addition, eukaryotic cell membranes exhibit an asymmetry in lipid distribution across the lipid bilayer. For example, in human erythrocyte membranes, PC and sphingomyelin are mainly found in the outer leaflet, whereas PE and phosphatidylserine (PS) are mainly found in the inner leaflet of the bilayer (Verkleij et al., 1973). The basic functions of biological membranes are carried out by membrane proteins that may associate loosely with the membrane surface

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(peripheral proteins) or traverse the lipid bilayer (integral membrane proteins). Most of the integral membrane proteins are composed of tightly packed bundles of transmembrane α -helical segments that contain \sim 20 hydrophobic residues (Kleinschmidt, 2003). Recent studies have shown that PE can assist in the correct folding of integral membrane proteins such as the lactose permease and the high affinity phenylalanine permease of Escherichia coli (Bogdanov and Dowhan, 1998; Zhang et al., 2003). A number of x-ray crystal structures also indicate that integral membrane proteins such as cytochrome c oxidase from Rhodobacter sphaeroides (Svensson-Ek et al., 2002) and the purple bacterial reaction center from Thermochromatium tepidum (Jones et al., 2002) may contain bound PE or other phospholipids. It was also found that aminophospholipids such as PE in the cytoplasmic leaflet of Golgi, endosome, and plasma membranes contribute to the protein-mediated vesicle budding competence of these membrane systems (Pomorski et al., 2003).

Although the mutual interactions of lipids and proteins are fundamentally important to both the structure and the function of all biological membranes, our understanding of the physical principles underlying lipid-protein interactions remains incomplete, and the actual molecular mechanisms whereby associated lipids alter the activity, and presumably

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also the structure and dynamics, of integral membrane proteins are largely unknown. This is due in part to the fact that most transmembrane proteins are relatively large, multidomain macromolecules with complex and often unknown three-dimensional structure and topology. Despite the fact that about one-third of all proteins are integral membrane proteins, less than 100 unique membrane protein structures have been reported to date, compared with the huge number (over 14,000) of soluble proteins in the Protein Data Bank. Moreover, membrane proteins can interact with lipid bilayers in complex, multifaceted ways (Gennis, 1989; Selinsky, 1992; Marsh and Horváth, 1998). To overcome this problem, a number of workers have designed and synthesized peptide models of specific regions of natural membrane proteins and have studied their interactions with model membranes of defined lipid composition (Davis et al., 1983; Ren et al., 1997; de Planque and Killian, 2003).

The synthetic peptide acetyl- K_2 -G-L₂₄-K₂-A-amide (P₂₄) and its analogs such as acetyl-Lys-2-Leu-24-Lys-2-amide (L_{24}) have been successfully utilized as a model of the hydrophobic transmembrane α -helical segments of integral membrane proteins (Davis et al., 1983). These peptides contain a long sequence of hydrophobic leucine residues capped at both the N- and C-termini with two positively charged, relatively polar lysine residues. Moreover, the normally positively charged N-terminus and the negatively charged C-terminus are blocked to provide a symmetrical tetracationic peptide that will more faithfully mimic the transbilayer region of natural membrane proteins. The central polyleucine region of these peptides was designed to form a maximally stable α -helix, particularly in the hydrophobic environment of the lipid bilayer core, whereas the dilysine caps were designed to anchor the ends of these peptides to the polar surface of the lipid bilayer and to inhibit the lateral aggregation of these peptides. In fact, circular dichroism (CD) (Davis et al., 1983) and Fourier transform infrared (FTIR) (Zhang et al., 1992a,b; Axelsen et al., 1995) spectroscopic studies of P_{24} have shown that it adopts a very stable α -helical conformation both in solution and in lipid bilayers, and x-ray diffraction (Huschilt et al., 1989), fluorescence quenching (Bolen and Holloway, 1990), and FTIR spectroscopic (Zhang et al., 1992a,b; Axelsen et al., 1995) studies have confirmed that P_{24} and its analogs assume a transbilayer orientation with the N- and C-termini exposed to the aqueous environment and the hydrophobic polyleucine core embedded in the hydrocarbon core of the lipid bilayer when reconstituted with various PCs. DSC (Huschilt et al., 1985; Morrow et al., 1985; Zhang et al., 1992b) and ²H NMR spectroscopy (Huschilt et al., 1985; Morrow et al., 1985; Paré et al., 2001) studies have shown that P_{24} and L_{24} broaden the gel/liquid-crystalline phase transition of the host phospholipid bilayer and reduces its enthalpy. The phase transition temperatures $(T_m s)$ of PC and phosphatidylglycerol (PG) bilayers are shifted either upward or downward by P_{24} or L_{24} , mainly depending on the degree of mismatch

between the peptide hydrophobic length and the lipid hydrophobic thickness (Zhang et al., 1992b; Liu et al., 2002, 2004). In contrast, the phase transition temperatures of PE bilayers are substantially decreased by the presence of P_{24} in a manner essentially independent on the peptide-lipid hydrophobic mismatch (Zhang et al., 1995a). ²H NMR (Pauls et al., 1985) and electron spin resonance (ESR) (Subczynski et al., 1998, 2003) spectroscopic studies have shown that the rotational diffusion of P_{24} about its long axis perpendicular to the membrane plane is rapid in the liquidcrystalline state of the bilayer and that the closely related peptides L_{24} and acetyl-Lys-2-(Leu-Ala)-12-Lys-2-amide $((LA)_{12})$ exist at least primarily as monomers in liquidcrystalline POPC bilayers, even at relatively high peptide concentrations.

Statistical analysis shows that the hydrophobic core of the transmembrane α -helices of many integral membrane proteins are normally flanked by aromatic residues like trytophan or tyrosine and that positively charged lysine or arginine residues are found adjacent to these aromatic residues (Killian and von Heijne, 2000). To study the roles of tryptophan and lysine residues in lipid-protein interactions, we have previously synthesized three analogs of P_{24} , Ac-K₂- L_{24} -K₂-amide (L₂₄), Ac-DAP₂-L₂₄-DAP₂-amide (L₂₄DAP; DAP is diaminopropionic acid), and Ac-K₂-W-L₂₂-W-K₂amide ($WL_{22}W$), and studied their interactions with zwitterionic PC and anionic PG bilayers (Liu et al., 2002, 2004). First, in the peptide $L_{24}DAP$, the two pairs of capping lysine residues at the N- and C-termini of L_{24} have been replaced with the lysine analogs DAP, in which three of the four side-chain methylene groups have been removed. This peptide was used to test the so-called snorkel model first suggested by Segrest et al. (1990) to explain the behavior of positively charged residues in the amphipathic helices present at the surfaces of blood lipoproteins and later extended to transmembrane α -helices by von Heijne et al. (Monne et al., 1998). According to the transmembrane peptide version of the snorkel model, the long, flexible hydrophobic side chains of lysine or arginine residues could extend along the transmembrane helix so that the terminal charged moiety can reside in the lipid polar headgroup region whereas the α -carbon of the amino acid residue remains well below (or possibly above) the polar/apolar interfacial region of the phospholipids molecule, even when the hydrophobic length of the peptide is considerably different from that of the host lipid bilayer. Because of the shorter spacer arms between the charged group and the α -carbon in DAP, L₂₄-DAP is expected to be less accommodating to the peptide-lipid hydrophobic mismatch than L_{24} , and any effects of such mismatch on the thermotropic phase behavior of the host lipid bilayer should thus be exaggerated. Second, in the peptide $WL_{22}W$, the Leu-3 and Leu-26 residues of L_{24} are replaced with tryptophans. The preference of aromatic tryptophan or tyrosine residues for the polar/apolar interfacial region of the host lipid bilayer has been found to be one of the common features of natural membrane proteins (von Heijne, 1994). Since tryptophan residues have been proposed to anchor the ends of α -helical transmembrane peptides to the polar/apolar interface of the lipid bilayer, we also expect that $WL₂₂W$ would be less accommodating to hydrophobic mismatch between the peptides and host lipid bilayer than would L_{24} .

To further clarify the role of interfacially localized tryptophan and lysine residues in lipid-protein interactions, we have used DSC and FTIR spectroscopy to study the interactions of L_{24} , $L_{24}DAP$ and $WL_{22}W$ with a series of zwitterionic PEs with different hydrocarbon chain lengths and have compared these results with those obtained previously for a homologous series of zwitterionic PCs (Liu et al., 2002) and anionic PGs (Liu et al., 2004). Our results demonstrate that the snorkeling of the terminal lysine residues at the end of the model peptides is required for optimization of the hydrogen-bonding and/or electrostatic interactions between the lysine residues at the termini of these peptides and the polar headgroups of the PE. Our results also indicate that the presence of interfacially localized tryptophan residues only slightly affects the interactions of these model peptides with the zwitterionic phospholipids PE and PC (Liu et al., 2002), although tyryptophan residues strongly modulate such interactions in anionic PG bilayers (Liu et al., 2004).

MATERIALS AND METHODS

The phospholipids used in this study were obtained from Avanti Polar Lipids (Alabaster, AL) and were used without further purification. Commercially supplied solvents of at least analytical grade quality were redistilled before use. Peptides were synthesized and purified as trifluoroacetic acid (TFA) salts using previously published solid-phase synthesis and reversed phase high-performance liquid chromatographic procedures (Mantsch et al., 1985).

Samples were prepared for DSC as follows. Appropriate quantities of peptides and PE were codissolved in methanol to obtain the desired lipid/ peptide ratio, and the solvent was removed in a stream of nitrogen at temperatures near $60-70^{\circ}$ C to ensure sample homogeneity. Then the sample was redissolved in chloroform/methanol (1:1), and the solvent was again removed in a stream of nitrogen at temperatures near 60–70C. Later the sample was redissolved in benzene and lyophilized overnight to form a white powder. Samples containing 0.5 mg lipid were then hydrated by vigorous vortexing with water at temperatures some $10-15^{\circ}$ C above the gel/liquidcrystalline phase transition temperature of the lipid. DSC thermograms were obtained from 0.5-ml samples with a high sensitivity Microcal VP-DSC instrument (Microcal, Northampton, MA), operating at heating and cooling rates of 10^oC per hour. The data were analyzed and plotted with the Origin software package (OriginLab, Northampton, MA).

Peptide samples to be used in FTIR spectroscopic experiments were converted to the hydrochloride salt by two cycles of lyophilization from 10 mM hydrochloric acid. This procedure was necessary because the trifluoroacetate ion gives rise to a strong absorption band (\sim 1670 cm $^{-1}$) that partially overlaps the amide I absorption band of the peptide (Zhang et al., 1992a,b). Typically, samples were prepared by codissolving lipid and peptide in methanol at a lipid/peptide ratio of near 30:1 (mol/mol). After removal of the solvent and drying of the film (see above), samples containing 2–3 mg lipid were hydrated by vigorous mixing with 75 μ l D₂O-

based buffer (50 mM Tris, 150 mM NaCl, 1mM NaN $_3$, pD 7.4). The paste obtained was then squeezed between the $CaF₂$ 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 temperature could be varied between 20° C and 90°C by an external, computer-controlled water bath. Infrared spectra were acquired as a function of temperature with a Digilab FTS-40 Fouriertransform spectrometer (Bio-Rad, Digilab Division, Cambridge, MA) using data acquisition parameters similar to those described by Mantsch et al. (1985). The experiment involved a sequential series of 2° C temperature ramps with a 20-min interramp delay for thermal equilibration and was equivalent to a scanning rate of $4^{\circ}C$ per hour. Spectra were analyzed with software supplied by the instrument manufacturers and other programs obtained from the National Research Council of Canada.

RESULTS

To evaluate the DSC and FTIR spectroscopic data presented below, we need to compare the intrinsic hydrophobic length of the model peptides L_{24} , L_{24} DAP, and WL₂₂W with the intrinsic hydrophobic thickness of the various PE bilayers used in this study (see Table 1). On the basis of measurements of a molecular model of L_{24} in an ideal α -helical conformation, we estimate that the mean hydrophobic length of L_{24} (the average length of the leucine sequence measured at any point on the surface of the helix) is \sim 30.6 Å. Note that the effective hydrophobic length measured in this manner is two helical half-turns shorter at each end of the hydrophobic helical core than the maximal hydrophobic length of this peptide, which would be 36.0 Å . Thus the mean hydrophobic length of L_{24} is intermediate between the mean hydrophobic thickness of 14:0 PE and 16:0 PE bilayers (where the number before the colon represents the total number of carbon atoms in the hydrocarbon chains and the number after the colon the number of double bonds). For the shorter chain 12:0 PE bilayers, L_{24} hydrophobic length exceeds bilayer hydrophobic thickness in both the gel and the liquid-crystalline phases, whereas for other three PEs, the hydrophobic length of L_{24} will be less than that of gel state but more than that of the liquid-crystalline bilayer. The same considerations also apply to L_{24} DAP and WL₂₂W, which are taken to have the same effective hydrophobic length as L_{24} .

TABLE 1 Hydrophobic thicknesses of the bilayer formed by various phosphatidylethanolamines

PЕ	Hydrophobic thickness $(A)^*$		
	Gel phase	Liquid-crystalline phase	Mean [†]
12:0	29.3	19.7	24.5
14:0	34.2	22.8	28.5
16:0	39.5	26.3	32.9
18:0	44.7	29.8	37.3

*Hydrophobic thicknesses were estimated using the equations used by Sperotto and Mouritsen (1988) to calculate the hydrophobic thicknesses of PC bilayers.

[†]Mean of the hydrophobic thicknesses of the gel and liquid-crystalline phases.

We stress here that the pattern of matching of lipid bilayer thickness and peptide hydrophobic length just described applies only if these model peptides adopt an ideal α -helical conformation that is not influenced by lipid bilayer thickness. In fact, our spectroscopic studies show that in PE bilayers these model peptides do adopt a predominantly α -helical conformation that is, however, affected by gel-state PE bilayer thickness (see below). Moreover, the hydrocarbon chains of the PE molecules do change their degree of conformational order, and thus the effective hydrophobic thickness of the PE bilayer, in response to the presence of these model peptides. Thus the actual degree of hydrophobic mismatch between these model peptides and the various PE bilayers, in both the gel and the liquid-crystalline states, may be somewhat different than indicated by the intrinsic effective hydrophobic length of this peptide and the hydrophobic thickness of PE bilayers in the absence of peptide.

Thermotropic phase behavior of PE bilayers in the absence of peptides

As illustrated in Fig. 1, in the absence of peptides the DSC thermograms of unannealed aqueous dispersions of each of four n-saturated diacyl PEs studied here exhibit a single asymmetric fairly energetic and highly cooperative lamellar gel/lamellar liquid-crystalline phase transition on heating. This transition is freely reversible, as shown by the absence of significant cooling hysteresis (data not shown). In addition, the temperatures of the phase transitions of these PEs increase progressively but nonlinearly, and the transition enthalpy increases linearly, with increases in the hydrocarbon chain length, as expected. For a thorough discussion of the thermotropic phase behavior of the complete homologous series of *n*-saturated diacyl PEs, see Lewis and McElhaney (1993) and references cited therein.

FIGURE 1 Effect of L_{24} on the DSC thermograms of a series of n-saturated diacyl-PEs. Thermograms are shown as a function of the acyl chain length $(N:0)$ of the lipids, at the peptide concentrations (mol %) indicated.

Thermotropic phase behavior of various PE bilayers in the presence of L_{24}

The effects of the incorporation of L_{24} on the main phase transition of PE bilayers are also illustrated in Fig. 1. In all cases the incorporation of increasing quantities of L_{24} produces an apparently two-component DSC endotherm, consisting of a broad, lower-temperature component and a sharp, higher-temperature component. The relative contribution of the sharp component of the DSC endotherm, which initially possesses a phase transition temperature, enthalpy, and cooperatively similar to that of the PE alone, decreases in magnitude as the concentration of L_{24} increases. In fact, the sharp component of the DSC endotherm is barely detectable at a peptide concentration of 3.3 mol % in 12:0 PE bilayers and a concentration of 6.7 mol % in the three longer chain PE bilayers studied. In contrast, the relative contribution of the broad component increases as the peptide concentration increases, and it is the only or at least the major component that persists at the highest peptide concentrations tested. Moreover, our FTIR spectroscopic studies of the temperature dependence of the methylene stretching frequency changes indicate that both the sharp and broad components of the DSC endotherms are accompanied by increases in the rotational conformational disorder of the PE hydrocarbon chains (data not shown). Using the rationale provided in our previous DSC studies of the interaction of P_{24} and related peptides with PC, PE, and PG bilayers (Zhang et al., 1992b, 1995a; Liu et al., 2002), we thus assign the sharp component of our DSC endotherms to the hydrocarbon chain-melting phase transition of peptide-poor PE domains and the broad component to the melting of peptide-rich PE domains. Note that these characteristic effects of the incorporation of L_{24} and its analogs on the sharp component of the DSC endotherm are noted in all the PEs studied and are essentially hydrocarbon chain length independent.

The effects of L_{24} incorporation on the temperatures of the sharp and broad components of the DSC heating endotherms for all four PEs are illustrated in Fig. 2. As noted above, the temperature of the sharp component changes little with increases in L_{24} concentration in each of the four PE bilayers examined. However, in all cases there is a gradual decrease in the temperature of the broad component with increasing peptide concentration, resulting in a gradual increase in ΔT (the downward shift of the phase transition temperature of the broad component relative to that of the sharp component). Moreover, at all peptide concentrations, the phase transition temperature of the broad component of the DSC endotherm always occurs at a lower temperature than that of the sharp component, regardless of PE hydrocarbon chain length. At 6.7 mol % L_{24} concentration, the ΔT is largest in 12:0 PE and smallest in 18:0 PE bilayers, but the variation with hydrocarbon chain length is small $(<1°C)$. Qualitatively similar results were observed when L_{24} DAP and WL₂₂W are incorporated into PE bilayers.

FIGURE 2 Effect of L_{24} concentration on the peak temperatures of the two components of the DSC thermograms exhibited by the mixtures of L_{24} and the n-saturated diacyl-PEs. Data are presented for the sharp (\triangle) and broad (\blacktriangledown) components of the DSC endotherms.

The effects of the lipid acyl chain length on the ΔT values observed with PE-peptide mixtures containing 3.3 mol $\%$ L₂₄ are illustrated in Fig. 3. With all of the PEs examined, the ΔT values obtained are all negative and only weakly dependent on lipid hydrocarbon chain length, with the absolute ΔT values diminishing only slightly with increasing hydrocarbon chain length (from -2.6° C for 12:0 PE to -2.0° C for 18:0 PE). As illustrated in Fig. 3, these observations contrast sharply from those that occur when L_{24} is incorporated into PC bilayers. With the latter, the sign and magnitude of ΔT is

FIGURE 3 Hydrocarbon chain length dependence of the differences (ΔT) between the transition temperatures of the peptide-rich and the peptide-poor PE components of DSC thermograms exhibited by L_{24}/PE mixtures at a peptide concentration of 3.3 mol $\%$ (\blacksquare). To facilate comparison, comparable data obtained from studies of L_{24} with PC bilayers (\bullet) are also shown.

strongly hydrocarbon chain length dependent in a manner consistent with the sign and degree of the hydrophobic mismatch between peptide hydrophobic length and lipid bilayer hydrophobic thickness. With the shorter chain PC bilayers (\leq 14:0 PC), the temperature of the broad component of the DSC endotherm is progressively increased relative to that of the sharp component as hydrocarbon chain length decreases, and vice versa in the longer chain PC bilayers (\geq 16:0 PC), whereas the ΔT value is near zero for 15:0 PC bilayers. As noted previously (Liu et al., 2002), the effective hydrophobic length of P_{24} (or L_{24}) match the mean thickness of 15:0 PC bilayers. Thus the ΔT value becomes increasing positive in PC bilayers with mean hydrophobic thicknesses that are progressively less than the effective hydrophobic length of the peptide and vice versa. Qualitatively similar results are also obtained when L_{24} is incorporated into in anionic PG bilayers (Liu et al., 2004). The possible molecular basis for these results will be discussed later.

The variation in the enthalpy of the overall phase transition as a function of L_{24} concentration in each of the four PE bilayers studied is shown in Fig. 4. As noted earlier, in all cases the contribution of the sharp component to the total enthalpy change measured decreases steeply with increasing peptide concentration with little dependence on PE hydrocarbon chain length (see Fig. 1). However, the variation in the enthalpy of the overall and broad components of the DSC endotherms depends strongly on PE hydrocarbon chain length. With the shorter chain PEs (i.e., 12:0 and 14:0 PE), the transition enthalpies of both components decrease as L_{24} concentration increases. However, in the longer chain PE bilayers, these transition

FIGURE 4 Peptide-concentration dependent changes in the enthalpy of the thermotropic transitions exhibited by L_{24} -containing PE bilayers. The data presented represents the total enthalpy change measured for mixtures of L₂₄ with 12:0 PE ($-\blacksquare$), 14:0 PE($-\triangle$), 16:0 PE ($-\blacktriangle$), and 18:0 PE $(-\bullet -).$

enthalpies first decrease and then increase again as peptide concentration increases, with this effect becoming more pronounced as PE hydrocarbon chain length increases. A similar pattern of behavior was observed in our previous studies of P_{24}/PE systems (Zhang et al., 1995a), in which case the apparent increase in the overall phase transition enthalpy, and that of the broad component of the DSC endotherm, was ascribed to the release of heat due to the disaggregation of peptide multimers in the gel states of the longer chain PE bilayers as the gel/liquid-crystalline phase transition is approached on heating. It is noteworthy, however, that similar behavior is not observed when L_{24} is incorporated into PC and PG bilayers (Liu et al., 2002, 2004). With the latter lipids the overall phase transition enthalpy decreases linearly with increasing L_{24} concentration over a similar range of hydrocarbon chain lengths, indicating that significant peptide aggregation does not occur in the more loosely packed gel states of these phospholipids.

A closer examination of Fig. 1 also shows that the width of the broad component of the DSC endotherms increases modestly with increases in peptide concentration but is almost independent of lipid hydrocarbon chain length. For example, in 12:0 PE bilayers, the width of the broad component (as measured from the starting to completion temperatures) exhibits an increase from 8° C to 11° C as peptide concentrations increases from 1.7 to 6.7 mol %. Similar behavior occurs when the peptide P_{24} is incorporated into the same PE bilayers (Zhang et al., 1995a). However, markedly different behavior occurs when L_{24} is incorporated into either PC or PG bilayers (Liu et al., 2002, 2004). With either L_{24} -containing PC or L_{24} -containing PG bilayers, the widths of the broad component in the DSC thermograms are

always considerably larger at comparable peptide concentrations and decrease markedly with increases in lipid hydrocarbon chain length. The possible basis of these experimental observations will be explored in the Discussion.

Thermotropic phase behavior of PE bilayers in the presence of various peptides

To examine the effect of truncating the lysine side chains of L_{24} or of replacing the terminal leucines with aromatic tryptophan residues on the peptide-PE interactions, we have also studied the effects of L_{24} DAP and WL₂₂W incorporation on the thermotropic phase behavior of various PE bilayers by high-sensitivity DSC. A comparison of the effects of the incorporation of various amounts of L_{24} , L_{24} DAP, and $WL_{22}W$ on the phase behavior of 16:0 PE bilayers is shown in Fig. 5. The patterns of peptide concentration-dependent changes in PE thermotropic phase behavior shown therein are typical of our observations of the interactions of these peptides with all of the other PEs examined. We will therefore present the overall effects of these peptides on the thermotropic phase behavior of 16:0 PE bilayers as a qualitative example of what occurs with all of the PE bilayers studied. As noted earlier, the DSC thermograms exhibited by the $L_{24}/16:0$ PE mixtures (*left* panel) are composed of overlapping short and broad components with the former decreasing and the latter increasing with increases in L_{24} concentration. At the 6.7 mol % peptide, the DSC thermogram exhibited by the $L_{24}/16:0$ PE mixture is mainly composed of a broad component centered near 57.8°C with only traces of the sharp component centered near 63.2° C. The DSC thermogram exhibited by the corresponding $WL_{22}W/16:0$ PE mixtures (*right* panel) is also composed of overlapping sharp and broad components, but in this case the sharp component of the DSC endotherm predominates

FIGURE 5 A comparison of the effects of L_{24} , $L_{24}DAP$, and $WL_{22}W$ on the DSC thermograms of 16:0 PE. The thermograms shown were obtained at the peptide concentrations (mol %) indicated.

over the broad component, even at the highest peptide concentration tested. In contrast to L_{24} and $WL_{22}W$, the incorporation of L_{24} DAP has only a small effect on the gel/ liquid-crystalline phase transition of 16:0 PE bilayers and does not appear to induce clearly resolvable sharp and broad components in the DSC endotherm. At 6.7 mol % peptide concentration, the DSC thermogram exhibited by the $L_{24}DAP/16:0$ PE mixture is only slightly broader than that of the pure 16:0 PE. This marked difference between L_{24} DAP and L_{24} or $WL_{22}W$ is also observed when these peptides are incorporated into 12:0, 14:0, and 18:0 PE bilayers and suggests that the $L_{24}DAP$ is not interacting extensively with the host PE bilayers.

The plot of ΔT versus the hydrocarbon chain length of the PE bilayers for L_{24} , L_{24} DAP, and WL₂₂W are shown in Fig. 6. Although ΔT remains negative in all of the peptide/PE mixtures studied, the magnitude of the decrease in the temperature of the broad component is largest for L_{24} , moderate for $WL_{24}W$, and smallest for $L_{24}DAP$, especially in the longer chain PEs. For all the three model peptides, only a very small increase in ΔT is observed when the hydrocarbon chain length of PE bilayers is increased from 12 to 18 carbons. This is markedly different from the results of the same three model peptides in PC and PG bilayers, in which the ΔT initially decreases significantly with an increase of the lipid hydrocarbon chain length and is strongly positive in very short chain phospholipids. It is also interesting to note that in all four hydrocarbon chain lengths studied, the ΔT values exhibited by L₂₄/PE and WL₂₂W/PE mixtures are \sim 1.5°C and 1.0°C lower, respectively, than that the ΔT exhibited by L₂₄DAP/PE mixtures, again indicating less extensive interactions between the latter peptide its host PE bilayer.

FIGURE 6 Hydrocarbon chain length dependence of the differences (ΔT) between the transition temperatures of the peptide-rich and the peptide-poor PE components of DSC thermograms exhibited by peptide/PE mixtures at a peptide concentration of 3.3 mol %. Data are presented for mixtures of PE with $L_{24} (\triangle)$, $WL_{22}W (\triangle)$, and $L_{24}DAP (\blacktriangledown)$.

Illustrated in Fig. 7 is a comparison of the peptide concentration-dependent changes in the enthalpy of the thermotropic phase transitions observed when the peptides L_{24} , L_{24} DAP, and W L_{22} W are incorporated into the various PE bilayers examined. With the shorter chain lipids 12:0 PE and 14:0 PE, incremental increases in peptide content are accompanied by a progressive decrease in the enthalpy values throughout the peptide concentration range examined. Also, the magnitude of this effect varies with the nature of the peptide and decreases in the order L_{24} > $WL_{22}W$ > L_{24} DAP. With the longer chain PEs, the measured enthalpy values also decrease at low peptide concentrations, in a manner and magnitude comparable to what occurs with the shorter chain lipids. Unlike the shorter chain PEs, however, at higher peptide concentrations a marked increase in enthalpy values is observed. Similar behavior has been observed previously in studies of P_{24} - and $(LA)_{12}$ -containing PE membranes (Zhang et al., 1995a, 2001) and have been ascribed to additional energy arising from increased dispersal of the aggregated peptide at temperatures below the onset of the lipid chain-melting phase transition. Here, we also find that the magnitude of this phenomenon is dependent on the nature of the peptide and decreases in the same order described above (i.e., $L_{24} > W L_{22} W >$ L_{24} DAP). We also note that across the range of PE hydrocarbon chain lengths examined, the effects of L_{24} -DAP on the enthalpy of the lipid phase transition are considerably smaller than observed with peptides L_{24} and $WL_{22}W$. These results suggest either that L_{24} DAP is intrinsically less disordering to PE bilayers than are L_{24} and $WL_{22}W$, or that $PE-L_{24}$ DAP interactions are being minimized by the lateral aggregation of L_{24} DAP in PE bilayers.

A comparison of the effects of L_{24} , L_{24} DAP, and $WL_{22}W$ on the transition widths of PE bilayers is shown in Fig. 8. The transition widths were measured from the starting temperature to the ending temperature of the broad component of the DSC endotherms at 3.3 mol % peptide concentration. For all three model peptides, there is only slight change in the transition widths of the broad component of the peptide/PE mixtures when the acyl chain length of PE bilayers is increased from 12 to 18 carbons. Moreover, in all four PEs examined, the transition widths of the broad component of L_{24} DAP/PE mixtures (~5.2°C) is much smaller than that of L_{24}/PE mixtures and $WL_{22}W/PE$ mixtures $(-9.2^{\circ}$ C and 8.4°C, respectively). These results again show that the incorporation of L_{24} or $WL_{22}W$ has much larger effects on the organization of PE bilayers than does the incorporation of L_{24} DAP.

Fourier transform infrared spectroscopic studies of peptide-containing PE bilayers

In these studies, infrared spectra of mixtures of the peptide with each of the four PEs studied were recorded as a function of temperature and as a function of the mol fraction of the

FIGURE 7 Effects of L_{24} (∇), WL₂₂W (\bullet), and $L_{24}DAP$ (∇) concentration on the total transition enthalpies of 12:0. 14:0, 16:0, and 18:0 PE bilayers.

peptide. The use of FTIR spectroscopy permits a noninvasive monitoring of both the structural organization of the lipid bilayer and the conformation of the incorporated peptide (for a more detailed description of this application of IR spectroscopy, see Lewis and McElhaney, 1996, 2002; Tamm and Tatulian, 1997; and references cited therein). Here, changes in the degree of hydrocarbon chain rotational isomeric disorder coincident with the lipid gel/liquidcrystalline phase transition were monitored by an examination of the methylene symmetric stretching band near 2850 cm^{-1} , and the conformational stability of the embedded peptide was monitored by an examination of the contours of

FIGURE 8 Plot of the transition width versus the hydrophobic chain length of the lipid bilayer at a peptide concentration of 3.3 mol %. Data are presented for mixtures of PE with $L_{24} (\triangle)$, $WL_{22}W (\triangle)$, and $L_{24}DAP (\blacktriangledown)$.

the amide I band centered near 1650 cm^{-1} . Our studies (data not presented) showed that the incorporation of these peptides into PE bilayers had no discernable effect on the ester carbonyl stretching band near 1735 cm^{-1} , nor on the O-P-O asymmetrical stretching bands near 1230 cm^{-1} , suggesting that peptide incorporation at the levels studied here $(\sim 3.3 \text{ mol } \%)$ did not have a major effect on the hydration of either the phosphate polar headgroups or the polar/apolar interfacial regions of these lipid bilayers.

Illustrated in Fig. 9 are the amide I and $C=O$ stretching regions of the FTIR spectra exhibited by mixtures of L_{24} with the four PEs examined here. The spectra shown were acquired at temperatures where the host lipids are in their gel (L_β) and liquid-crystalline (L_α) phases. Under all conditions examined, the amide I absorption band of L_{24} is dominated by sharp absorption components centered near 1655–1657 cm^{-1} and near 1647–1650 cm^{-1} . These absorption bands can be ascribed to the amide I vibrations of unexchangedand deuterium-exchanged α -helical peptide domains (Chirgadze and Brazhnikov, 1974; Rabolt et al., 1977), and their prominence is therefore consistent with L_{24} maintaining a predominantly α -helical conformation under all conditions examined. However, Fig. 9 also shows that the gel to liquidcrystalline phase transition of the host PE bilayer is accompanied by a small downward shift in the frequency of the absorption maximum of the amide I band envelope. This frequency shift is minimal $(< 2$ cm⁻¹) with L24-containing 12:0 PE mixtures but increases with increases in lipid hydrocarbon chain length and approaches values near 5–6 cm-¹ with 18:0 PE-based mixtures. A similar pattern of behavior has been reported in our studies of mixtures of L_{24} with PC and PG bilayers (Liu et al., 2002, 2004) and in

FIGURE 9 The $C=O$ stretching and amide I bands of the FTIR spectra exhibited by L_{24} -containing PE bilayers. The absorbance spectra shown were acquired with mixtures containing 3.3 mol $\%$ L₂₄ under conditions where the lipid bilayer hosts are in the L_B-phase and L_{α}-phase states. The dashed lines mark the absorption maxima (1656 cm^{-1}) of the amide I absorption bands arising from fully proteated α -helical peptide domains. The small absorption bands centered near 1670 cm^{-1} arise from small amounts of trifluoroacetate counterions that were not completely removed during sample preparation.

studies of the interaction of the peptides P_{24} and $(LA)_{12}$ with PC and PE bilayers (Zhang et al., 1992b, 1995a,b, 2001).

A comparison of the lipid phase state-induced changes in the amide I band maxima observed with mixtures of peptides L_{24} , L_{24} DAP, and W L_{22} W with 16:0 PE is shown in Fig. 10. These peptides all exhibit a pattern of incremental hydrocarbon chain length-dependent decreases in amide I band maxima that is qualitatively similar to that described for L_{24} above. However, as is evident from Fig. 10, the magnitude of this effect is markedly smaller with the peptide $L_{24}DAP$, for which the maximal chain length-dependent frequency shifts observed never exceeds 2 cm^{-1} , indicating that the coupling between the conformation of $L_{24}DAP$ and changes in the hydrophobic thickness of these PE bilayers is attenuated compared with the other two peptides. Interestingly, our studies of the interactions of these same peptides with PC and PG bilayers (Liu et al., 2002, 2004) also show a similar pattern in which lipid phase state-induced decreases in the amide I band maxima of L_{24} DAP are smaller than in L_{24} and $WL₂₂W$, although in those cases this affect is smaller in magnitude than observed here with PE bilayers. The molecular basis of these experimental observations are explored in the Discussion below.

DISCUSSION

In this study we have shown that interactions of model peptides that mimic the hydrophobic transmembrane α -helical segments of integral membrane proteins with zwitterionic PE bilayers are quite different from their interactions with zwitterionic PC or anionic PG bilayers. Specifically, we demonstrate that all three model peptides reduce the $T_{\rm m}$ s of PE bilayers to a grater extent than in PC and PG bilayers and

that the peptide-induced reduction of the $T_{\rm m}$ s of these PE bilayers is largely independent of the lipid bilayer thickness. The latter observations contrast sharply from those reported in our previous work, in which the direction and magnitude of the peptide-induced shift in the $T_{\rm m}$ s of PC and PG bilayers was shown to be dependent on the thickness of the lipid bilayer (Liu et al., 2002, 2004). Evidently the nature and strength of the interactions of the somewhat polar and charged amino acid residues at the ends of these model peptides with the polar headgroups of zwitterionic PE and anionic PG or zwitterionic PC bilayers are quite different. We provide a possible molecular interpretation of these experimental observations below.

The $T_{\rm m}$ s of saturated PCs are comparable to those of saturated PG bilayers and are $20-30^{\circ}$ C lower than those of saturated PE bilayers of comparable hydrocarbon chain length (see Lewis et al., 1987; Zhang et al., 1997; and references cited therein). This observation can be rationalized on the basis of stronger electrostatic and H-bonding interactions between the polar headgroups at the surfaces of PE bilayers (see Boggs, 1980, 1986, 1987, and references cited therein). Under physiologically relevant conditions, PE amino groups are fully protonated and, being positively charged, are capable of both net attractive electrostatic interactions with negatively charged groups and H-bonding interactions with H-bonding acceptor groups. Thus, the intermolecular forces that constitute the basis of the relatively high transition temperatures of PEs are, in effect, a summation of contributions arising from electrostatic attraction between positively charged amino groups and negatively charged phosphate moieties, electrostatic repulsion arising from both amino-amino and phosphatephosphate contacts, as well as H-bonding interactions

FIGURE 10 The C=O stretching and amide I bands of the FTIR spectra exhibited by peptide-containing 16:0 PE bilayers. The absorbance spectra shown were acquired with mixtures containing 3.3 mol % of the peptides indicated under conditions where the lipid bilayer hosts are in the L_{β} -phase and L_{α} -phase states. The dashed lines mark the absorption maxima (1656) cm⁻¹) of the amide I absorption bands arising from fully proteated α -helical peptide domains. The small absorption bands centered near 1670 cm^{-1} arise from small amounts of trifluoroacetate counterions that were not completely removed during sample preparation.

between the headgroup amino protons and H-bonding acceptor groups in the headgroup and polar/apolar interfacial regions of the lipid bilayer. With anionic PG bilayers, their relatively low transition temperatures are largely attributable to electrostatic repulsion between the negatively charged headgroup phosphate moieties, though this effect will be partially mitigated by H-bonding interactions between the exchangeable protons of the headgroup glycerol moiety and H-bonding acceptor groups in the headgroup and polar/ apolar interfacial regions of the lipid bilayer. Finally, with PC bilayers, there are no H-bonding donor groups on their polar headgroups, and as a result electrostatic attraction between positively charged choline groups and negatively charged phosphate moieties and electrostatic repulsion arising from both choline-choline and phosphate-phosphate contacts will predominate. However, the steric bulk of the Nmethyl groups will markedly reduce the frequency of close contacts interactions between the positively charged choline nitrogen and the negatively charged phosphate groups, and thus the electrostatic attraction component will be markedly attenuated in magnitude. Thus, despite the zwitterionic character of PC polar headgroups, electrostatic repulsion between phosphate moieties at the surfaces of PC bilayers may well have as large an effect on bilayer T_m as occurs with PG bilayers. Largely because of these effects, the forces favoring the gel phase over liquid-crystalline phases are expected to decrease in the order $PE \gg PG \sim PC$, and the incorporation of peptides such as L_{24} into PE, PC, and PG bilayers may be expected to affect interactions between lipid headgroups in two main ways. First, the presence of these peptides will reduce the access of adjacent phospholipid molecules to potential hydrogen-bonding partners, thus leading to the disruption of part of the hydrogen-bonding networks of PG and PE bilayers. Second, the positively charged side-chain amino groups at the N- and C-termini of these peptides will compete with positively charged and Hbonding donor groups on adjacent lipid polar headgroups for electrostatic and/or H-bonding acceptor sites, thereby disrupting the pattern of lipid-lipid interactions at the surface of these bilayers. As noted above, the electrostatic and Hbonding network at the surface of a PE bilayer is quite strong and is largely responsible for the relatively high transition temperature. Consequently, any significant peptide-induced disruption of the surface electrostatic and H-bonding network of a PE bilayer will have a considerably greater effect on its T_m than will occur with PC and PG bilayers, where the corresponding surface networks are considerably weaker. Moreover, given the magnitude of the changes in T_m that are likely to occur upon disruption of the electrostatic and H-bonding network at the surface of a PE bilayer, such changes may well mask the additional changes arising from hydrophobic mismatch effects, which are thus expected to be reduced in magnitude.

A major finding of this study is that the replacement of lysine by DAP residues at the ends of the model peptides markedly affects their interactions with zwitterionic PE bilayers. Specifically, our studies show that $L_{24}DAP$ is much less effective than L_{24} at reducing the $T_{\rm m}$ and enthalpy and broadening the phase transition of PE bilayers. These results indicate that the capacity of L_{24} DAP to disrupt lipid-lipid interactions in PE bilayers is either intrinsically smaller than that of L_{24} , or that this peptide may be more prone to lateral aggregation in PE bilayers than L_{24} , or both. Our experimental observations can be attributed to the different lengths of the methylene spacers between the α -carbons and sidechain amino groups of DAP (one $CH₂$ spacer) and lysine (four $CH₂$ groups). Thus, when lysine-capped peptides such as L_{24} are incorporated into lipid bilayers, their lysine side chains can extend away from their helical surfaces and the side-chain amino groups can thus compete effectively with positively charged and H-bonding donor groups on adjacent lipids for electrostatic and/or H-bonding acceptor sites. Also,

because their lysine side chains are longer than those of the leucine residues of the peptide, lateral contacts between L_{24} molecules in lipid bilayers can result in intermolecular contacts between the positively side-chain amino groups at the peptide termini and, in turn, charge repulsion between these side-chain amino groups will reduce the probability of forming peptide aggregates under such conditions. However, with L_{24} DAP, the DAP side chains do not extend beyond those of the leucine residues on the peptide surface. Thus, when such peptides are incorporated into lipid bilayers, the side chain amino groups cannot compete with positively charged and H-bonding donor groups on adjacent lipids for electrostatic and/or H-bonding acceptor sites as effectively as can those of L_{24} . Moreover, because the DAP side chains are so short, lateral contacts between $L_{24}DAP$ molecules in a lipid bilayer will not result in the type of intermolecular close contacts between the positively side-chain amino groups at the peptide termini that are likely to occur with a lysinecapped peptide like L_{24} . Consequently, the energetic cost of peptide lateral aggregation in L_{24} DAP-containing lipid bilayers will be smaller than that of comparable L_{24} -containing lipid bilayers. We therefore suggest that, in general, $L_{24}DAP$ will not be as perturbing of lipid thermotropic phase behavior as L_{24} because of a combination of its smaller capacity to disrupt the electrostatic and H-bonding network at the lipid bilayer surfaces and the smaller energetic penalties against its aggregating in lipid bilayers. Moreover, these factors may be especially important in PE bilayers, because the electrostatic and H-bonding networks at the surfaces of PE bilayers are quite strong and are thus more likely to favor PE-PE interactions over PE-peptide interactions unless peptide inclusion induces a significant disruption of this network. Thus L_{24} DAP should be inherently less perturbing of PE bilayers than L₂₄. Interestingly, our previous studies indicated no major differences between L_{24} and L_{24} DAP on the thermotropic phase behavior of either or PC of PG bilayers (Liu et al., 2002, 2004). Most probably this is because the electrostatic and H-bonding networks at the surfaces of PC and PG bilayers are considerably weaker than in PE bilayers and the differential effects of lysine and DAP in modulating these forces are not so obvious as in PE bilayers.

Our studies also show that the effect of $WL_{22}W$ on the thermotropic phase behavior of PE bilayers is somewhat smaller than that of L_{24} . Given the arguments presented above (see previous paragraph), it appears that the replacement of a leucine residue at each end of the hydrophobic core of L_{24} by tryptophan residues attenuates the capacity of the terminal lysine residues to interact with or otherwise perturb the electrostatic and H-bonding network at the surfaces of PE bilayers. This could be an indication that the tryptophan residues are forming cation- π -interactions with adjacent lysine residues, as suggested in our previous studies of the interaction of $WL_{22}W$ with anionic PG bilayers (Liu et al., 2004). Indeed, if these cation- π -interactions do occur between lysine and tryptophan residues, then the lysine

residues may not be able to compete as effectively as those of L_{24} for positively charged and H-bonding donor groups on adjacent lipids for electrostatic and/or H-bonding acceptor sites. However, despite its having a smaller effect on the thermotropic phase behavior of PE bilayers than L_{24} , it is clear that the effects of both peptides are of comparable magnitude and are both much greater than that of $L_{24}DAP$. Thus, the effect of the cation- π -interactions between lysine and tryptophan residues is considerably less effective than the lysine snorkeling effect in regulating lipid-peptide interactions in lipid bilayers.

In these studies we also observe small decreases in the frequency of peptide amide I band maxima at the gel/liquidcrystalline phase transitions of their lipid bilayer hosts, and that the magnitude of these frequency shifts decrease with increases in lipid hydrocarbon chain length. Similar results have been reported in studies of these peptides with PC and PG bilayers (Liu et al., 2002, 2004) and in studies of the interaction of P_{24} with PC and PE bilayers (Zhang et al., 1992b, 1995a). As noted previously, these changes seem to arise from a lipid phase state-induced decline in populations giving rise to amide I absorption centered near 1660–1665 cm^{-1} and a concomitant increase in populations giving rise to amide I absorptions centered near $1645-1650$ cm⁻¹ and have been assigned to conformational changes in the deuterium-exchanged N- and C-termini of these peptides (Liu et al., 2002). However, we also find that these lipid phase state-induced decreases in peptide amide I band maxima are consistently smaller in $L_{24}DAP$ -containing PE membranes than observed with the either L_{24} -containing or WL22W-containing membranes. This observation is consistent with the idea that interactions of $L_{24}DAP$ with PE bilayers are less extensive than those of the peptides L_{24} and WL₂₂W and with the suggestion that, unlike L_{24} - and W L_{22} W-containing PE bilayers, significant peptide lateral aggregation occurs in $L_{24}DAP$ -containing PE bilayers.

In conclusion, our studies highlight the importance of the H-bond interactions and electrostatic attractions between the lipid polar headgroups and the terminal lysine and/or tryptophan residues in peptide-lipid interactions. Our studies also indicate an important role of the length and flexibility of the terminal lysine residues on the interactions of these model peptides with zwitterionic PE bilayers and extend our understanding of the role of interfacially localized lysine and tryptophan residues in natural membrane proteins. Further structural and computer modeling investigations are required to elucidate the detailed mechanism of how the terminal lysine and/or tryptophan residues may interact with adjacent lipid headgroups at the membrane polar/apolar interface.

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