Relationship between CYP1A2 Localization and Lipid Microdomain Formation as a Function of Lipid Composition

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Running Title: CYP1A2 Localization in Microdomains of Lipid Vesicles

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Text pages – 28
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Figures – 10
References – 45
Abstract – 250 words
Introduction – 700 words
Discussion – 1440 words

Abbreviations – Cytochrome P450 (P450), NADPH-cytochrome P450 reductase (CPR), phosphatidylcholine vesicles (V-PC), liquid-ordered domains (lo), liquid-disordered domains (l_d), detergent-resistant membranes (DRMs), ER lipids vesicles (V-ER), DRM lipid vesicles (V-DRM), mixed phospholipid vesicles (V-Mix PL), endoplasmic reticulum (ER), N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (NBD-PE), Lissamine™ rhodamine B 1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (Rh-PE), and 6-dodecanoyl-2-dimethylaminonaphthalene (laurdan), General polarization (GP), 1,2-di-(9,10-dibromo)stearoyl-sn-glycero-3-phosphocholine (BrPC), phosphatidylcholine (PC), phosphatidylethanolamine (PE), Phosphatidylinositol (PI), Phosphatidylserine (PS), phosphatidic acid (PA), cholesterol (chol), sphingomyelin (SM), 1-palmitoyl-2-oleyl-sn-glycero-3-phosphocholine (POPC), palmitoyl sphingomyelin (PSM), fluorescence resonance energy transfer (FRET).
Abstract

Cytochrome P450 (P450) function requires the interaction of P450 and NADPH-cytochrome P450 reductase (CPR) in membranes, and is frequently studied using reconstituted systems composed solely of phosphatidylcholine. There is increasing evidence that other endoplasmic reticulum (ER) lipids can affect P450 structure, activity, and interactions with CPR. Some of these lipid effects have been attributed to the formation of organized liquid-ordered (l_o) domains. The goal of this study was to determine if l_o domains were formed in P450 reconstituted systems mimicking the ER membrane. CYP1A2, when incorporated in “ER-like” lipid vesicles, displayed detergent insolubility after treatment with Brij 98 and centrifugation in a sucrose gradient. Lipid probes were employed to identify domain formation in both ER-like vesicles and model membranes known to form l_o domains. Changes in FRET using an established donor/acceptor FRET pair in both ER-like and model l_o-forming systems demonstrated the coexistence of l_o- and liquid-disordered (l_d) domains as a function of cholesterol and sphingomyelin content. Similarly, laurdan, a probe that reports on membrane organization, showed that cholesterol and sphingomyelin increased membrane order. Finally, brominated-phosphatidylcholine allowed for monitoring of the location of both CPR and CYP1A2 within the l_o regions of ER-like systems. Taken together, the results demonstrate that ER-like vesicles generate microdomains, and both CYP1A2 and CPR predominantly localize into l_o membrane regions. Probe fluorescent responses suggest that lipid microdomains form in these vesicles whether or not enzymes are included in the reconstituted systems. Thus, it does not appear that the proteins are critical for stabilizing l_o domains.
Introduction

Cytochromes P450 (P450s) are membrane-bound proteins predominantly found in the endoplasmic reticulum (ER) and are essential enzymes in the oxidation of endogenous and exogenous compounds (Nelson, 2003). P450s must interact with their redox partner, NADPH-cytochrome P450 reductase (CPR) to receive electrons that are required for substrate metabolism (Gigon, et al., 1969). Another obligatory component of this monooxygenase system is phospholipid. Although phospholipid has been known to be an important element in P450 function for many years (Strobel, et al., 1970), more recently, there has been increasing evidence that specific phospholipids affect this system significantly. A number of different lipids have been demonstrated to directly affect catalytic activity, stability of the enzymes, and incorporation of P450 into the membrane (Ingelman-Sundberg, et al., 1996; Reed, et al., 2006; Blanck, et al., 1984). Furthermore, a few studies have suggested that P450s can promote anionic lipid segregation and domain formation, which in turn, affects P450 function (Jang, et al., 2010; Kim, et al., 2003; Kim, et al., 2007).

Lipid microdomains in the plasma membrane also have been linked to protein function and a multitude of cellular events (Brown and London, 1998). There are two types of lipid domains that exist in the plasma membrane under physiologic conditions. Liquid-ordered (l_o) phases are enriched in saturated fatty acid chains and cholesterol, which decreases the fluidity of the lipid environment and prevents them from being solubilized by non-ionic detergents, lending to their informal name – detergent-resistant membranes (DRMs) (Brown and London, 1998). Conversely, liquid-disordered (l_d) membrane regions are mainly composed of unsaturated fatty acid chains that allow for increased membrane fluidity (Quinn and Wolf, 2009). DRMs have been extensively characterized in the plasma membrane, but only recently have been described in...
the ER (Browman, et al., 2006; Pielsticker, et al., 2005; Hayashi and Fujimoto, 2010; Brignac-Huber, et al., 2011).

Our lab has previously characterized ER-DRMs in rabbit liver microsomes that, similar to plasma membrane DRMs (Pike, 2004), are enriched in cholesterol and sphingomyelin (Brignac-Huber, et al., 2011). CYP1A2 and CPR were found to reside in these ER-DRMs, which also were determined to be cholesterol-dependent. Moreover, upon using previously published protocols (Reed, et al., 2008; Reed, 2010) to integrally incorporate CYP1A2 and CPR into purified vesicles that mimicked DRM, the apparent $K_m$ of CPR-CYP1A2 complex was decreased when compared to vesicles composed solely of bovine liver phosphatidylcholine. These findings suggested that the specific lipid composition of the ER-DRMs increased CYP1A2-dependent metabolism by altering the efficiency of CPR and CYP1A2 interaction. Although these studies showed that the function of P450 was influenced by its incorporation into vesicles comprised of lipids with a composition similar to ER-DRMs, ordered lipid domain formation in these artificial membranes was not conclusively established.

Numerous studies have analyzed lipid microdomain formation in model lipid systems that mimic the plasma membrane (Dietrich, et al., 2001; Baumgart, et al., 2007; Loura, et al., 2001; de Almeida, et al., 2005). However, very few studies have done analogous experiments in lipid systems that mimic the ER. Therefore, in this study, we provide physical evidence that lipid domains are formed in vesicles made up of physiologic lipids at the relative concentrations observed in the ER. First, we demonstrate that purified CYP1A2 and CPR, when incorporated together, localize to DRMs in these vesicular reconstituted systems in a manner that is similar to their localization in rabbit liver microsomes (Brignac-Huber, et al., 2011). In addition, the existence of lipid microdomains is demonstrated in the “ER-like” reconstituted systems by
employing several fluorescent probes that have been shown to preferentially localize to either liquid-ordered (lo) or liquid-disordered (ld) regions in plasma membrane models. Alternatively, one of the probes utilized in this study, laurdan, displays different fluorescent characteristics when isolated in lo and ld domains. Studies using these types of probes typically have been performed in plasma membrane model systems containing three well-defined lipid components and have led to the construction of phase diagrams showing the lipid proportions that define the boundaries for the lo, ld, and gel phases as well as those that delineate the boundaries of coexisting phases (de Almeida, et al., 2003).

By using a ternary-lipid model system (de Almeida, et al., 2003) as a reference and lipid vesicles containing only PC as a “negative” control, our results using the fluorescent probes provide evidence for the existence of lipid microdomains in the ER-like reconstituted systems in the absence of proteins. When proteins were integrally incorporated into the vesicular reconstituted systems, the results indicate that CYP1A2 and CPR both localize to DRMs. Furthermore, the addition of proteins to the vesicular reconstituted systems does not seem to affect lipid microdomain distribution suggesting that lipid domain formation occurs independently of protein and in turn, influences the behavior of the enzymes of the P450 system.
**Materials and Methods**

**Materials.** Lipids were purchased from Avanti Polar Lipids (Alabaster, AL). The lipids used for the reconstituted system mimicking the ER (natural membrane systems) were natural extracts from a variety of sources; phosphatidylcholine (liver, bovine), phosphatidylethanolamine (liver, bovine), phosphatidylinositol (liver, bovine), phosphatidylserine (brain, porcine), phosphatidic acid (egg, chicken), sphingomyelin (brain, porcine). The lipids used for reconstituted systems of model membranes were the synthetic lipids, 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) and palmitoyl-sphingomyelin (PSM). Cholesterol was added to reconstituted systems using water-soluble cholesterol from Sigma-Aldrich (St. Louis, MO). N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (NBD-PE), Lissamine™ rhodamine B 1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (Rh-PE), and 6-dodecanoyl-2-dimethylaminonaphthalene (laurdan) were purchased from Invitrogen (Carlsbad, CA). Protease cocktail inhibitor was purchased from Roche (Indianapolis, IN). The 5-µm syringe filters were purchased from Osmonics (Greenville, SC). YM-30 filters were purchased from Millipore (Billerica, MA), and 96-well, half area flat bottom plates were purchased from Corning (Lowell, MA). Bio-Beads SM-2 were purchased from Biorad (Hercules, CA).

**Enzyme Source.** CYP1A2 was isolated and purified from β-naphthoflavone-treated rabbit liver microsomes as described previously (Coon, et al., 1978). P450 levels were determined by measuring the absorbance of the carbon monoxy-ferrous complex at 450 nm (Omura and Sato, 1964).

**Lipid Vesicle Preparation.** Except for the details described below, vesicular lipid reconstituted systems containing 5 µM CYP1A2 were prepared at a 500:1 total lipid:P450 ratio.
as previously described (Reed, et al., 2008). Various lipid compositions were prepared in our reconstituted systems containing either naturally occurring lipids or synthetic lipid to study model membranes. Phosphatidylcholine vesicles (V-PC), mixed phospholipid vesicles (V-MixPL), vesicles mimicking the total microsomal membrane (V-ER lipids) and vesicles with the lipid composition of the DRM fractions (V-DRM) were prepared as previously described (Brignac-Huber, et al., 2011). Model membranes were prepared with the lipid compositions (mol%) listed in Table 1. All lipids except for cholesterol (added as described below) were dissolved in chloroform and slowly dried under a stream of N₂ at room temperature until chloroform was completely removed (~1 hour). Lipid was rehydrated (at 1/5th the final volume of the vesicle preparation) with 250 mM HEPES (pH 7.5) containing 5% sodium glycocholate and bath-sonicated until it clarified. When appropriate, the solubilized lipid solution was then added to the purified enzymes (to achieve final protein concentrations of 5 µM and 1 µM of P450 and CPR, respectively) in four equal aliquots of the rehydrated lipid suspension. This step limited the destruction of P450 by detergent. N₂ was layered over the P450/lipid solution between each addition before mixing the sample by inversion. When proteins were not present, the lipid solution was added to deionized water. The mixture was then incubated at 4°C for 1 hour before adding 125 mg of Biobeads SM-2 to remove the detergent. The samples were rocked for 2 hr at 4°C and the vesicles were subsequently separated from the Biobeads by drawing the sample into a syringe using a 26.5 gauge needle. The beads were then washed twice with 100ul of reaction buffer (50 mM HEPES, 15mM MgCl₂, 5mM EDTA, pH 7.5) to recover any residual vesicle volume. These washes were added to the original volume extracted from the beads and the entire sample was filtered through a 5-µm syringe filter. Vesicles that required the addition of cholesterol (indicated in Table 1) were treated as previously described using water-soluble
cholesterol (Niu and Litman, 2002; Brignac-Huber, et al., 2011). The water-soluble cholesterol was dissolved in H₂O for a final cholesterol concentration of 4µg/µl. Water-soluble cholesterol was added to the appropriate reconstituted systems and samples were rocked at room temperature for 2 hours upon which they were filtered using an YM-30 filter at 1300g for 15 minutes. This centrifugation allows for the MβC molecule to flow through the filter, leaving the vesicle system above the filter. The vesicle preparation was removed and the filter was washed twice with 75 µl of reaction buffer. Measurement of the ferrous-CO P450 complex (Omura and Sato, 1964) was used to determine P450 recovery after vesicle preparation. All lipid probes, in a chloroform stock solution, were added to lipid mixtures before drying. For FRET interactions between NBD-PE and Rh-PE, the fluorescent probes were added at concentrations of 0.1 mol% and 0.5 mol%, respectively. Laurdan was added to the lipid mixture in 1:1000 dye:lipid ratio. Brominated PC was added at 15 mol%.

Isolation of Detergent –Resistant Membranes by Brij 98 solubilization and Discontinuous Sucrose Gradient Centrifugation. A portion (0.2 ml) of the vesicular reconstituted systems, containing CYP1A2 and/or CPR (described above) was taken to 1 ml with 0.75 ml of reaction buffer and 0.05 ml of either 1% or 10% Brij 98 stock solution in water (as indicated in the results). The vesicles were solubilized, and DRMs were isolated by centrifugation (19 h at 210,000g at 4°C) on a discontinuous sucrose density gradient as described in a previous study (Brignac-Huber, et al., 2011). Eleven, 1 ml fractions were removed from the top of the gradients, and the pellets were re-homogenized in reaction buffer, and protein localization was determined by western blotting of the individual fractions. DRMs were isolated in fractions 2-5 of the discontinuous gradient.
**Fluorescence Spectroscopy.** All fluorescence spectroscopy measurements, unless otherwise stated, were carried out at room temperature. The excitation wavelength for NBD-PE is 450 nm with an emission at 535 nm, whereas Rh-PE is excited at 535 nm and emits at 605 nm. Therefore, in FRET studies between Rh-PE and NBD-PE, the excitation wavelength was 450 nm and emission was monitored at 605 nm. Values of the relative FRET efficiencies (%) were determined by the following:

\[
\frac{F}{F_{PC}} \times 100
\]

where \( F \) is the fluorescence intensity at 605 nm of each mixed lipid system and \( F_{PC} \) is the fluorescence intensity at 605 nm of either 100% bovine liver PC for the “physiologic” membrane systems or 100% 1-palmitoyl-2-oleyl-\( sn \)-glycero-3-phosphocholine (POPC) for the model membrane systems (de Almeida, et al., 2003).

The laurdan emission spectrum provides an assessment of the degree of membrane order. Laurdan is a lipophilic dye that distributes equally into \( l_o \) and \( l_d \) membrane regions, but exhibits spectral properties that are dependent on lipid packing (Dinic, et al., 2010; Parasassi, et al., 1986). The laurdan emission spectrum exhibits a blue shift in the presence of \( l_o \) domains, which is attributed to a dipolar relaxation phenomenon (Parasassi, et al., 1986). The laurdan emission spectra were measured from 370 to 620 nm with excitation at 350 nm. General polarization (GP) of laurdan is a spectroscopic property, which can be used to quantify the relative proportion of coexisting lipid phases with an increase in GP being reflective of an increase in membrane order. The GP values are measured as follows:

\[
GP = \frac{(I_B-I_R)}{(I_B+I_R)}
\]
where \( I_B \) and \( I_R \) are the peaks of emission intensity at the blue and red edges of the emission spectra.

The fluorescence of tryptophan residues in CYP1A2 and CPR was measured at 341 nm after excitation at 295 nm. The effect of brominated PC on protein tryptophan fluorescence was expressed as the percent fluorescence relative to that observed when proteins were incorporated in lipid vesicles that did not contain brominated PC:

\[
\text{Relative \% Trp fluorescence} = \left( \frac{F}{F_o} \right) \times 100
\]

Where \( F \) is the fluorescence intensity at 341 nm with Br-PC present and \( F_o \) is the fluorescence intensity at 341 nm in the absence of Br-PC.

All of the lipid components were included at identical relative concentrations in the two sets of vesicles, as bovine liver PC was substituted for brominated PC in the control vesicles.

When the tryptophan fluorescence of CPR was studied, the reconstituted system was centrifuged through an Illustra Microspin S-400 HR size-exclusion spin column (GE Healthcare, Pittsburgh, PA) according to the manufacturer’s specifications. This step was performed to remove any CPR that was not physically incorporated in the lipid bilayer of the reconstituted system (Reed, et al., 2006; Reed, et al., 2008). It was determined that brominated lipids did not affect the incorporation of CPR into the lipid vesicles (data not shown).
Results

Isolation of DRMs from purified, reconstituted P450 systems containing physiologic lipids. To elucidate the role of lipid microdomains in the function of the P450 enzyme system, it would be useful to replicate the domains that occur within a natural membrane in a reconstituted system. The advantage of this approach lies in the ability to examine the interactions of these proteins under defined and controlled conditions. In our previous study (Brignac-Huber, et al., 2011), we showed that the interaction of CYP1A2 and CPR was facilitated and in turn, the catalytic activity was stimulated when the enzymes were integrally reconstituted in vesicles that had the same lipid composition as DRMs. However, the previous study did not demonstrate that lipid microdomains were formed in the vesicular reconstituted systems or that the P450 and CPR interacted with putative domains in these systems. As a result, these possibilities are addressed in this study by using our previously published method to prepare vesicular, reconstituted P450 systems that have the enzymes integrally incorporated in the lipid bilayer (Reed, et al., 2008; Reed, 2010).

The most direct way to demonstrate that DRMs were present in the reconstituted systems and were associated with the P450 enzyme system was to solubilize the reconstituted vesicles with Brij 98 and isolate the DRMs using discontinuous sucrose density centrifugation (Brignac-Huber, et al., 2011). Due to their high lipid to protein ratio, detergent-resistant membranes coalesce around the interface between the 5% and 38% sucrose layers (i.e. fractions 2-5) (Pielsticker, et al., 2005). Figure 1 shows the results of this type of experiment and compares P450 reconstituted systems made up of only bovine liver PC (V-PC) and those made up of a mixture of natural lipids at the same relative proportions occurring in the ER (V-ER).
Figure 1 (top two profiles) shows the gradient fractions obtained after detergent treatment and centrifugation of a lipid reconstituted system containing only CYP1A2. In order to demonstrate the progressive solubilization of the vesicles, two concentrations of Brij 98 (0.05 % and 0.5%) were used in the experiments. The left panel shows the results when the vesicles were comprised of only PC, and panel 1B shows the corresponding results with V-ER reconstituted systems. It may be noted that lower Brij 98 concentrations were used to solubilize the model membranes than those used to solubilize rabbit liver microsomes (Brignac-Huber, et al., 2011). This is consistent with previous studies of model membrane vesicles showing that lower detergent concentrations are necessary for solubilization relative to those needed for natural membrane solubilization (Dietrich, et al., 2001; Sengupta, et al., 2008; Pathak and London, 2011).

The results showed complete solubilization of the CYP1A2 in PC vesicles (none of the protein was isolated in fractions 2-5). Thus, when the reconstituted system was comprised of a single phospholipid, PC, and not capable of forming segregated lipid domains, all of the CYP1A2 was solubilized by treatment with 0.5% Brij 98. Conversely, a significant proportion of the enzyme (approximately 25%) was still located in the uppermost fractions of the gradient in the V-ER system treated with 0.5% Brij 98. Thus, similar to what was observed in rabbit liver microsomes, CYP1A2 was resistant to solubilization with Brij 98 when it was incorporated in a reconstituted system that had the lipid composition of the endoplasmic reticulum.

Figure 1 also shows the corresponding sucrose density gradient profiles using reconstituted systems containing only CPR (third and fourth profiles from the top). Unlike the results with CYP1A2 reconstituted systems, CPR was solubilized from both V-PC and V-ER vesicles at 0.5% Brij 98 (Panels A and B, respectively). Although a small amount of CPR still remained in fractions 4 and 5, the great majority of the protein was no longer found in the
fractions representing the DRM. This finding suggests that CPR does not have the same inherent affinity for DRMs as CYP1A2 when reconstituted alone into the V-ER vesicles.

In addition, Figure 1 shows the sucrose gradient profiles when both CPR and CYP1A2 were reconstituted into the V-PC and V-ER systems (fifth and sixth profiles from the top). As observed in the reconstituted systems containing CYP1A2 alone, the P450 also isolated to domains when reconstituted with CPR. Interestingly, CPR also was localized to domains (approximately 28%) in the presence of CYP1A2, suggesting that the presence of CYP1A2 “draws” CPR into the domain. Because CPR was also observed to be in DRMs when examined in microsomes, it seems the presence of P450s in the domains either changes the localization of CPR or increases the tightness by which CPR binds to the DRM regions.

**Assessment of the degree of membrane order using fluorescent lipid probes.** We examined the potential of reconstituted vesicles to form lipid domains by utilizing several lipid probes in the absence and presence of either CYP1A2 or CPR. The general structures of the fluorescent lipid probes and brominated phospholipids used in this study are illustrated in Figure 2. There were two types of comparisons made with regard to different reconstituted systems. One set of comparisons used model systems comprised of different concentrations of a chemically defined phospholipid, 1-palmitoyl-2-oleyl-sn-glycero-3-phosphocholine (POPC), a chemically defined sphingomyelin, palmitoyl-sphingomyelin (PSM), and cholesterol. The second set of comparisons used reconstituted systems comprised of different proportions of naturally occurring phospholipids and sphingomyelin (with or without cholesterol).

Domain formation has been well characterized using the model system lipids, and the processes involved in microdomain formation using vesicles derived from the model lipids are
believed to be representative of those that occur in the plasma membranes (de Almeida, et al., 2003; de Almeida, et al., 2005). The ternary diagram shown in Figure 3 indicates the lipid compositions that correspond to vesicles comprised entirely of the l_d phase; a mixture of l_d and l_o phases; entirely of the l_o phase; a mixture of l_d, l_o, and gel (s_o) phases; and a mixture of l_o and s_o phases. As a basis of comparison for the behavior of fluorescent lipid probes in membranes comprised of natural lipids, we examined lipid probe fluorescence in four types of vesicles made from the model lipids. The compositions (shown as black, numbered dots in Figure 3) were representative of membranes containing varying proportions of lipid phases that would be expected to form in physiologic membranes. More specifically, we wanted to study probe fluorescence with the following types of membrane: 1) one comprised entirely of liquid-disordered lipid (POPC); 2) one comprised of a mixture of lipids that forms co-existing liquid-ordered and liquid-disordered regions but represents a composition near the phase boundary for liquid-disordered membranes (and thus, only a small proportion of the membrane would consist of the l_o phase); 3) same as 2) but the lipid mixture represents a composition in the middle of the domain for co-existing lipid phases; and 4) same as 2) but the mixture represents a composition near the liquid-ordered phase region. The proportion of lipid in the l_o phase would be expected to increase with the proportion of cholesterol (de Almeida, et al., 2003). The lipid composition of the vesicles tested is reported in Table 1.

Probe fluorescence was also examined in the following four types of lipid vesicles comprised of natural lipids (i.e. those extracted from animal tissues): 1) PC alone (V-PC), 2) a mixture of phospholipids in proportions comparable to those of the ER (V-Mix PL), 3) a mixture of phospholipids, cholesterol, and sphingomyelin in proportions equal to those of the ER (V-ER), and 4) a mixture of phospholipids, cholesterol, and sphingomyelin in proportions equal to those
of DRM isolated from the ER (V-DRM) as described previously (Brignac-Huber, et al., 2011). By observing the behavior of lipid probes in model systems with different extents of phase separation, we could compare the probe responses in “physiologic” vesicles to assess whether similar processes were occurring.

It should be noted that the lipid vesicles prepared with naturally occurring lipids contain a mixture of fatty acid chains. Studies have shown that this aspect of these systems make lipid phase transitions more subtle than those occurring in the otherwise comparable model systems. For instance, Jin et al. (Jin, et al., 2006) used a probe (di-4-ANEPPDHQ) that responds to alterations in membrane organization through shifts in its emission scan. This study found that, although the reporter probe was useful for vesicles, the emission spectrum shift was far less dramatic for vesicles comprised of natural lipids when compared to vesicles made of synthetic lipids. This effect was attributed to the more heterogeneous mixture of acyl chains found with natural lipid systems and would likely lessen the order of \( l_0 \) domains when compared to \( l_0 \) domains formed from synthetic lipids (Jin, et al., 2006). Similarly, a calorimetric study by Barenholz et al. examined the thermotropic behavior of lipid vesicles composed of natural and synthetic sphingomyelin lipid species. It was reported that membrane organization of sphingomyelin is a complex function of its specific composition and that the transition temperature of natural sphingomyelin was much broader than the synthetic forms (Barenholz, et al., 1976). Because it is less likely to obtain data with fluorescent probes that implicate lipid phase transition in vesicles made from naturally occurring lipids, more confidence can be placed in the data that do support the occurrence of these processes in these systems.

**Segregation of fluorescent probes into lipid microdomains using FRET.** There are a number of useful probes to evaluate phase separation in pure lipid systems. Two such probes
which have been extensively used and characterized are N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (NBD-PE) and Lissamine™ rhodamine B 1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine, triethylammonium salt (Rh-PE) (Figure 2). These probes are fluorescently labeled at the head group of the lipid and have different acyl chain properties. NBD-PE has two saturated acyl chains that allow this lipid to reside in more ordered domains (l_o) of the membranes. Conversely, Rh-PE has two unsaturated acyl chains, which are kinked and therefore prefers to reside in more fluid regions (l_d) of the membrane (Dietrich, et al., 2001; Crane and Tamm, 2004; de Almeida, et al., 2003). FRET between the two probes is dependent on the distance between them and can be measured by exciting NBD-PE at 450 nm (emission 535 nm) and monitoring Rh-PE emission at 605 nm. Rh-PE is excited directly through irradiation at 535 nm. Because the probes selectively partition into the ordered and disordered membrane regions, respectively, the extent of FRET varies in the presence of cholesterol and sphingomyelin when these lipid constituents lead to microdomain formation. Incorporation of the probes into a homogeneous fluid membrane would produce a baseline FRET signal between the Rh and NBD probes. As the proportions of cholesterol and sphingomyelin are increased across the l_d/l_o phase boundary (Fig. 3), the FRET interaction would be expected to decrease. The maximal decrease in FRET fluorescence would be observed where the size of the lipid domains leads to the greatest segregation of the probes; however, as the size the domains vary on either side of this point of optimal segregation, the FRET signal would be expected to increase. Thus, by comparing the FRET signal in vesicles consisting entirely of an l_d environment with mixed lipid systems that possess l_o and l_d domains, the potential for lipid microdomain formation can be assessed.
First, potential FRET interactions in the model system were monitored as described above. In the pure POPC vesicles (Model 1), there would be no phase separation and thus the two probes should be free to interact. The Model 2 lipid vesicles (Table 1) are located on the phase diagram (Figure 3) at a point in which phase separation changes from \( l_d \) to \( l_d + l_o \). The last two systems, Model 3 and Model 4 contain higher proportions of cholesterol and sphingomyelin and correspond to compositions at which the \( l_o \) regions comprise larger proportions of the entire lipid system.

FRET values were calculated relative to the FRET signal in Model 1 (100% POPC). As shown in Figure 4, the relative FRET signal at 23°C (Figure 4A) and 37°C (Figure 4B) between NBD-PE and Rh-PE was significantly decreased in the Model 2 vesicles suggesting that phase segregation of the two probes was greatest in these vesicles. As the sphingomyelin (PSM) and cholesterol concentrations were increased (Model 3), the degree of FRET between the two probes appeared to increase and was even greater in the Model 4 vesicles. This effect was likely due to re-localization of the NBD-PE probe into the \( l_d \) regions as suggested by Crane and Tamm (Crane and Tamm, 2004). Crane and Tamm performed fluorescence recovery after photobleaching (FRAP) studies in model membranes to monitor the localization and mobility of specific probes in \( l_o \) and \( l_d \) domains of model membranes. Consistent with the predicted outcome described above, the NBD-PE probe partitioned less selectively into \( l_o \) regions as the cholesterol composition was increased above 15% in the model membranes. The Crane and Tamm study used higher SM concentrations than in the present study, therefore there cannot be a direct comparison of the two results, but it is envisaged that the same effect could occur in these lipid systems with high cholesterol content. Ideally, to confirm a FRET interaction, an increase in the fluorescence signal of the acceptor molecule (in this case, Rh-PE) of the FRET pair should be
matched by a decrease in the fluorescence signal of the donor molecule (NBD-PE). However, this relationship is complicated with this FRET pair by the well-known ability of NBD-PE to self-quench its fluorescence (Brown, et al., 1994; Rodgers and Glaser, 1991). The selective partitioning of NBD-PE into relatively small l_o domains formed at lipid compositions near the l_d/l_o + l_d boundary (model lipids 2) would be expected to result in a greater degree of crowding of the probe in the domains and consequently a greater degree of self-quenching of its fluorescence. Thus, in this study, the fluorescence emission of the acceptor is used as an indicator of the physical interaction of this FRET pair and not the fluorescence emission of the donor.

The interaction between NBD-PE and Rh-PE also was monitored in lipid systems that were composed of various natural lipids at physiologically relevant compositions (Brignac-Huber, et al., 2011) (Table 1). Relative to vesicles made up of 100% PC (V-PC), the addition of other phospholipids (V-MixPL) did not significantly influence the FRET response at either 23°C (Figure 5A) or 37° (Figure 5B). However, the addition of 5 mol% cholesterol and 4 mol% sphingomyelin to the mixed phospholipids (as shown in the V-ER lipid system) produced a 15% and 20% decrease in the FRET signal at 23°C and 37°C, respectively. These results suggest effective segregation of the NBD-PE and Rh-PE probes into l_o and l_d domains, respectively. The FRET quenching in the V-ER system was most comparable to the FRET results associated with Model 2 and was consistent with a lipid composition in the l_o + l_d region of the phase diagram near the boundary for the l_d region (Figure 3), favoring the formation of relatively small ordered domains. Similar to the model system, as the cholesterol and sphingomyelin content was increased in the DRM vesicles, the FRET signal was increased. Again, this suggests that the selective partitioning of NBD-PE into the l_o domains was diminished as the size of the domains increased with increasing cholesterol and sphingomyelin content.
Detection of lipid microenvironment using the laurdan fluorescent probe. While the FRET results demonstrate segregation of the \( l_0 \) and \( l_d \) probes into different membrane regions, these experiments alone do not directly establish the phase state of the membrane (Bagatolli, 2006). In an effort to confirm microdomain formation in these membrane systems, laurdan was used to probe the effect of alterations in lipid composition on membrane characteristics. Laurdan is a lipophilic dye that distributes equally into \( l_0 \) and \( l_d \) membrane regions, but exhibits spectral properties that are specific to different types of lipid packing (Dinic, et al., 2010; Parasassi, et al., 1986). The laurdan emission spectrum exhibits a blue shift in the presence of \( l_0 \) domains (Parasassi, et al., 1986). The relative proportions of \( l_0 \) and \( l_d \) phases can be assessed by generalized polarization (GP), which is a normalized intensity ratio of the two phases (Parasassi, et al., 1991). Laurdan emission scans were first monitored with the model systems composed of different SM and cholesterol compositions. As illustrated in Figure 3, 100% POPC vesicles (Model 1) form a fluid bilayer (\( l_d \)) (de Almeida, et al., 2003). The laurdan emission spectra for these vesicles (Figure 6) produced a peak of fluorescence intensity around 461 nm. Increasing the concentration of PSM and cholesterol would be expected to produce a higher degree of membrane organization (Figure 3), which would affect the relaxation properties of the laurdan probe. The incorporation of small amounts of PSM and cholesterol (to 5% and 15%, respectively) into POPC vesicles (Model 2) led to a depression of the fluorescence peak at 461 nm with an increase in the shoulder at 434 nm (Figure 6). Further increases in PSM and cholesterol increased this response, causing a significant shift of the laurdan emission spectra, producing peak intensity at 434 nm with decreased fluorescence intensity at 461 nm. These data indicate that the presence of cholesterol and PSM influenced the micro-environment of the laurdan probe to be increased in membrane order.
Based on the results obtained from the model systems, the goal of the next experiments was to determine if membranes with lipid structure and composition similar to those found in the ER exhibit similar changes in laurdan emission. As expected, the laurdan emission spectra of 100% PC vesicles (V-PC) (Figure 7) produced a similar spectrum as the 100% POPC system (Figure 6), suggestive of a homogeneous, fluid membrane. Interestingly, vesicles comprised of a mixture of phosphoglycerides (V-Mix PL) produced a significant decrease in fluorescence intensity at 461nm, and an alteration in the 434 nm/461 nm ratio. These changes are similar to those associated with the presence of cholesterol- and sphingomyelin-containing l_o domains and may reflect the putative ability of anionic phospholipids to form ordered domains in mixed PL vesicles (Ahn and Yun, 1998; Kim, et al., 2003; Kim, et al., 2007). Vesicles having a lipid composition similar to the total ER membrane (V-ER) produced a laurdan emission spectrum with depressed fluorescence at 461 nm and a relative increase in the intensity at 434 nm, again suggesting transition to more ordered membrane regions. Lastly, the DRM lipid vesicles (V-DRM) produced a further shift in the spectra to 434 nm with a concomitant decrease of fluorescence intensity at 461 nm. The generalized polarization (GP) values were calculated from each scan and are shown in Table 2. In both the model systems and the physiologic lipid systems, the maximum GP values were seen in the vesicles containing the highest amount of cholesterol and sphingomyelin. Based on the magnitudes of the GP values, domain formation in the physiologic lipid systems did not appear to be as dramatic as in the model systems; however, it should be recognized that the alterations in cholesterol concentrations were more modest than those made with the defined model systems. Nonetheless, these results are consistent with l_o domain formation with the more complex lipid membranes.
The changes described above examined the potential for lipid domain formation in the absence of protein. Additionally, we examined the effect of the presence of the P450 protein CYP1A2 on the phase properties of these lipid vesicles. A direct comparison of the laurdan emission scans in the absence (solid lines) and presence of CYP1A2 (dashed lines) showed no significant effect on the GP values for laurdan emission in the presence of CYP1A2 (Figure 7). These results suggest that the addition of CYP1A2 to the vesicle systems did not significantly affect the lipid packing properties of the vesicles. GP values in all lipid systems with and without CYP1A2 are shown in Table 2.

Generalized polarization of laurdan is strongly affected by membrane organization (Table 2) and also by temperature (Parasassi, et al., 1991). Higher temperatures elicit a transition of the \( l_0 \) and \( l_0 + l_d \) boundary in the phase diagram causing membranes to become more fluid \( (l_d) \). Laurdan emission scans were performed with all lipid systems as a function of temperature. This change in membrane fluidity is illustrated from the laurdan emission scan of POPC vesicles containing 20% PSM and 30% cholesterol (Model 3) at temperatures ranging from 23-60°C in Figure 8A. The same scans were repeated for each model lipid system, generating similar results (Supplemental Figure 1). The results demonstrate that as the temperature was increased, the maximum fluorescence intensity shifted from 434 nm to 461 nm, demonstrating a progressive decrease in membrane order (Parasassi, et al., 1991). GP values were then calculated for each of the scans and plotted against temperature (Figure 8B) and showed a break in the curve between 40° and 45 °C, which is representative of the lipid transition from \( l_0 + l_d \) to \( l_d \) phases (Parasassi, et al., 1991). The same experiment was done in the vesicles comprised of natural lipids and results from the V-ER lipids are shown in Figure 9A. The data for all the other vesicle systems can be found in Supplemental Figure 2. Again, GP values were calculated for each scan and
plotted against temperature (Figure 9B). The data illustrate that vesicles composed of physiologically relevant lipid compositions exhibited similar changes in membrane organization and fluidity as a function of temperature as was observed with defined model systems. Therefore, \( l_0 \) and \( l_d \) phases coexist not only in the synthetic lipid systems, but also in membranes with lipid compositions that more closely mimic the ER bilayer.

**Brominated Phospholipid.** Although the FRET experiments demonstrated that \( l_0 \) domains existed in the presence of enzyme, we also wanted to determine whether CYP1A2 and CPR partitioned into \( l_d \) or \( l_0 \) regions of these vesicles. To address this question, we employed the use of 1,2-di-(9,10-dibromo)stearoyl-sn-glycero-3-phosphocholine (BrPC), which contains brominated tags on the acyl chains (Figure 2). The bulky bromine tags produce kinks in the tails causing this lipid to have properties similar to unsaturated PC chains that segregate into \( l_d \) regions of lipid vesicles (Carney, et al., 2007; East and Lee, 1982). Additionally, the bromine tags quench tryptophan fluorescence; therefore, we used this probe to assess its interaction with CYP1A2, given that this P450 contains 8 tryptophan residues throughout its sequence (Ozols, 1986), with 6 being located on the enzyme surface (Sansen, et al., 2007). Since BrPC prefers to reside in \( l_d \) membrane regions, the location of CYP1A2 can be determined based on tryptophan quenching in the different lipid systems. We performed these experiments in V-PC and V-ER lipid systems because we achieved the greatest partitioning of phase-selective probes in the V-ER systems according to FRET measurements using Rh-PE and NBD-PE. This would increase the likelihood of being able to detect significant changes in the CYP1A2 Trp fluorescence. First, various amounts of BrPC were titrated into PC vesicles (V-PC) with 5 µM CYP1A2 to identify concentrations where efficient quenching of CYP1A2 tryptophan fluorescence was observed (Supplemental Figure 3). Based on these results, 15 mol% BrPC was selected and substituted for...
PC in each of the vesicle systems. CYP1A2 tryptophan fluorescence at 341 nm was then monitored in each lipid system with and without the brominated lipid present. In V-PC, the fluorescence of tryptophan residues in CYP1A2 was quenched approximately 20% by the BrPC at 23°C (Figure 10A). The degree of fluorescence quenching was significantly less in V-ER lipids suggesting separation of the BrPC and CYP1A2 into different domains. Results were similar at 23°C and 37°C. Since, it is known that the BrPC resides in the l_d regions of the membrane (Carney, et al., 2007; East and Lee, 1982), these data support the localization of CYP1A2 into the l_o regions of the membrane and confirm our previous findings regarding the membrane localization of CYP1A2 in rabbit liver microsomes (Brignac-Huber, et al., 2011).

The similar experiment was also performed with lipid reconstituted systems containing only CPR (Figure 10B). Surprisingly, there was significantly less quenching of CPR tryptophan fluorescence by brominated lipids when the enzyme was reconstituted in the V-ER system (relative to that in V-PC). These findings suggest that CPR also localizes to a separate lipid domain from that containing the brominated lipids in the V-ER. These results are in contrast to those obtained when CPR-containing vesicles were solubilized and centrifuged on a discontinuous sucrose density gradient as the latter indicated that CPR was not localized to DRMs (Figure 1).
Discussion

The ER lipid membrane contains low levels of sphingomyelin and cholesterol and has been described as an organelle of loosely packed lipids, allowing for insertion and export of lipids and proteins (van Meer, et al., 2008). But many groups have shown that there is potential for organized microdomain formation within the ER membrane (Brignac-Huber, et al., 2011; Browman, et al., 2006; Jang, et al., 2010; Sevlever, et al., 1999). Our lab recently isolated ER detergent-resistant microdomains (ER-DRMs) from rabbit liver microsomes and found that CYP1A2 and CPR reside primarily in these resistant domains (Brignac-Huber, et al., 2011). Enzymatic studies both with microsomes and with reconstituted systems demonstrated that CYP1A2 activities were increased in the more complex systems containing SM and cholesterol – due to an increase in the apparent binding affinity of the CPR·CYP1A2 complex. The goals of the present study were to determine whether ordered lipid microdomains could be detected in reconstituted lipid vesicles having compositions similar to those found in the ER and to demonstrate whether CYP1A2 and CPR, when incorporated in the vesicles, predominantly resided within the ordered domains.

The formation of lipid domains in model membranes has been investigated for many years with the use of fluorescent probes that preferentially reside in ordered (l₀) or disordered (lₐ) lipid regions (de Almeida, et al., 2003; Baumgart, et al., 2007; Crane and Tamm, 2004). These investigations have led to the development of phase diagrams that identify the lipid compositions corresponding to different possible physical states of lipid organization within a membrane (de Almeida, et al., 2003). Although microdomain formation was established with model systems, the concentrations and specific types of lipid constituents are commonly outside the physiologic range for biological membranes. Therefore, we compared the behaviors of lipid probes in the
cholesterol:POPC:sphingomyelin model system and in reconstituted systems with compositions similar to ER membranes to assess the potential for lipid microdomain formation in the systems containing physiologic lipids.

FRET measured between NBD-PE and Rh-PE probes in the vesicles comprised of physiologic lipids was consistent with the existence of lipid microdomains and corresponded very well to the FRET measured in analogous model systems. The lowest degree of FRET was observed in the Model 2 system and in V-ER reconstituted systems which contained the lowest amount of cholesterol and sphingomyelin (Figures 4 & 5). At this lipid composition, there is a predicted phase transition within the Model 2 membrane that allows the co-existence of l_d and l_o regions (Figure 3). Because the lipid composition lies near the phase boundary, there is selective segregation and maximum separation of the two probes, leading to decreased FRET between the probes. It would be expected for FRET between NBD-PE and Rh-PE to be similar in the V-ER and Model 2 lipid reconstituted systems because it has been shown that biological membranes have lipid compositions that are very close to the l_o/l_d phase transition (Lingwood, et al., 2008). In addition, these results illustrate that the ER membrane composition is capable of phase separation and membrane heterogeneity at physiological temperatures (Figure 5B). The subsequent increase in FRET with increasing cholesterol and sphingomyelin compositions in V-DRM and Model 3 vesicles is a phenomenon that has been described elsewhere and is attributed to decreased selectivity of the NBD probe as cholesterol levels are increased (Crane and Tamm, 2004). This effect is further demonstrated by the fact that as the cholesterol concentration is additionally increased in the Model 4 system, there is a continued increase in the FRET signal.

While the selectivity of the probes discussed above can be dependent on the lipid composition, laurdan is a useful tool that indicates the proportions of l_o and l_d domains in a
manner that is independent of the domain size. This particular probe reflects changes in lipid organization through variations in its fluorescence emission. Moreover, since laurdan distributes equally within $l_0$ and $l_d$ phases, a generalized polarization (GP) value can be calculated from the emission scan of laurdan, which quantifies the relative proportion of $l_d$ and $l_0$ phases (Parasassi, et al., 1991). A higher degree of membrane organization will produce a higher GP value than disordered domains. Again, spectral shifts of laurdan were similar in the model systems and the “natural membrane” reconstituted systems. As the cholesterol and sphingomyelin compositions increased in either system, the excitation spectra blue-shifted providing support for the existence of organized domains within the vesicles (Figures 6 & 7). Furthermore, the GP values corroborated these results as the highest values were observed in V-DRM lipids.

The V-Mix PL preparation was tested because of research showing that different types of phospholipids, especially anionic phospholipids, stimulate P450 catalytic activity and promote the interaction of P450 and CPR (Ingelman-Sundberg, et al., 1981; Ingelman-Sundberg, et al., 1996; Blanck, et al., 1984; Ahn, et al., 1998; Ahn, et al., 2005). In addition, several studies have described domain formation by anionic phospholipids in the presence of P450s. For example, Kim et al. (Kim, et al., 2003) utilized various lipid probes to illustrate that the presence of phosphatidylethanolamine (PE) promoted anionic phospholipid-enriched domains in ternary systems composed of phosphatidylcholine (PC), PE, and phosphatidylserine (PS) or phosphatidic acid (PA). Moreover, this group found that the presence and clustering (domain formation) of anionic phospholipids caused CYP3A4 to be more catalytically active and more deeply inserted into the lipid vesicles. Furthermore, it was postulated that this phenomenon may be important in CYP3A4 interactions with other proteins in membranes (Kim, et al., 2003). Another study found that CYP2B1 induced the formation of anionic lipid domains in vesicles containing PC, PE, and
anionic phospholipid (PA, PS, or phosphatidylinositol were all tested) with PA having the
greatest clustering effect in the presence of CYP2B1 (Kim, et al., 2007). In addition, the presence
of PA caused a conformational change in CYP2B1 and increased CYP2B1 activity. It should be
noted that these studies used much higher anionic phospholipid concentrations than those found
in the ER membrane.

The results from the present study differ from these previous reports in that they suggest
that microdomain formation is governed by the lipids present and not created by the introduction
of protein – at least for CYP1A2 at lipid compositions encountered in the ER. Actually, our
study is consistent with that of Ahn and Yun (Ahn and Yun, 1998), which used the fluorescence
of lipid probes to show that lipid microdomains were formed in the absence of any added
protein. However, the Ahn and Yun study used anionic lipid concentrations that were much
higher than would be encountered in physiologic membranes. Our study demonstrates the
protein-independence of lipid microdomain formation by the comparison of the laurdan emission
spectra in the absence and presence of CYP1A2 (Figure 7). The presence of CYP1A2 did not
affect the laurdan spectra or the resulting GP values (Table 2). Although there are numerous
differences when comparing these results to literature reports, including the relative
compositions of the lipids used and the different types of P450s added to the reconstituted
system, our results are consistent with the localization of CYP1A2 into previously formed l_m
microdomains, rather than the microdomains being generated as a result of the presence of the
P450 enzyme. Nonetheless, the laurdan experiments in the present study support the idea of
anionic phospholipid domain formation. Interestingly, the laurdan emission scans were
suggestive of an intermediate degree of membrane organization in the presence of
physiologically relevant concentrations of anionic phospholipids (V-Mix PL) without the
inclusion of cholesterol and sphingomyelin (Figure 7). In accordance with the qualitative difference of the laurdan scan in the V-Mix PL vesicles, the GP value was 7-fold higher than that determined in the V-PC membranes (Table 2), suggesting a more ordered lipid bilayer in V-Mix PL vesicles.

Lastly, the results using brominated phospholipids established that both CYP1A2 and CPR preferentially resided in the l_o regions of the V-ER systems (Figure 10). These results are supportive of previous studies that showed CYP1A2 DRM residence in microsomal tissue (Brignac-Huber, et al., 2011; Bae, et al., 2004). However, the findings with CPR-containing systems contradicted the conclusions derived from detergent solubilization of the lipid vesicles as no CPR was observed in DRMs of the vesicles. This may indicate that CPR, like CYP1A2, preferentially associates with l_o domains but may associate less tightly to the domains than the P450 binds so that its localization is not maintained after solubilization of the V-ER with Brij 98.

The present study demonstrates that organized lipid domains do form in the presence of physiologically relevant ER lipids and that CYP1A2 preferentially resides in the ordered regions of the membrane. Furthermore, the presence of CYP1A2 does not appear to significantly influence microdomain formation. CPR appears to preferentially associate with domains loosely in the absence of CYP1A2 which allows it to be easily solubilized by detergent treatment; however, CPR may be drawn into the more ordered domains through its association with CYP1A2. Further work is needed to elucidate the nature of the interaction of CPR with DRMs. Taken together these results underscore the important roles that lipid microdomain formation play in the localization and function of the enzymes of the P450 monooxygenase system.
Authorship Contribution

Participated in Research Design: Brignac-Huber, Reed, Eyer, Backes
Conducted Experiments: Brignac-Huber, Eyer
Performed Data Analysis: Brignac-Huber, Reed, Backes
Wrote or contributed to the writing of the manuscript: Brignac-Huber, Reed, Backes


Dinic J, Biverstahl H, Maler L and Parmryd I (2010) Laurdan and di-4-ANEPPDHQ do not respond to membrane-inserted peptides and are good probes for lipid packing. *Biochim Biophys Acta*.


Sevlever D, Pickett S, Mann KJ, Sambamurti K, Medof ME and Rosenberry TL (1999) Glycosylphosphatidylinositol-anchor intermediates associate with triton-insoluble membranes in subcellular compartments that include the endoplasmic reticulum. *Biochem J* **343 Pt 3**:627-635.


Footnotes

These studies were supported by a US Public Health Services research grant from the National Institute of Environmental Health Sciences [Grant R01 ES004344]; the predoctoral research fellowship from the State of Louisiana Board of Regents [Grant LEQSF(2005-10) GF-08].

Both Lauren M. Brignac-Huber and James R. Reed contributed equally to the preparation of this manuscript and are designated as co-first authors.
Figure Legends

Figure 1. Discontinuous sucrose density gradient profiles of enzyme-lipid reconstituted systems after detergent solubilization with Brij 98. Reconstituted systems containing CYP1A2 and/or CPR (as indicated in the center of the figure) were prepared and solubilized with Brij 98 (at the concentrations indicated) as described in Materials and Methods. The resulting suspensions were layered on a discontinuous sucrose gradient, and 1 mL fractions from the top to bottom of the gradient (indicated by the lane numbering) were analyzed by western blotting for either CPR or CYP1A2. The * indicates the lane location of the 50 kDa and 70kDa molecular weight marker for CYP1A2 and CPR, respectively. Panel A shows the profiles generated from V-PC reconstituted systems, and panel B shows profiles generated from V-ER systems. When both enzymes were included in the reconstituted system, the word “Both” is used, and the enzyme detected in the western blots is indicated in parentheses. The results are representative of at least three separate experiments.

Figure 2. Lipid Probes. Chemical structure of the fluorescent lipid analogs employed and preference of lipid domain location.

Figure 3. Phase diagram of model lipid systems. Lipid phase diagram of POPC/PSM/Chol at 23°C adapted with permission from Elsevier (de Almeida, et al., 2003). Black numbered circles are experimental points used for vesicles in the model lipid systems and represent the following conditions: 1) one comprised entirely of liquid-disordered lipids; 2) one comprised of a mixture of lipids that forms a membrane with co-existing liquid-ordered and liquid-disordered regions but represent a composition near the phase boundary for liquid-disordered membranes; 3) same as 2) but the lipid mixture represents a composition in the middle of the domain for co-existing
lipid phases; 4) same as 2) but the mixture represents a composition near the liquid-ordered
phase region. Lipid compositions are described in Table 1.

**Figure 4. FRET interaction of NBD-PE and Rh-PE in Model lipid vesicles.** Degree of FRET
interaction was calculated as stated in Materials and Methods for vesicles composed of model
lipids. Measurements were done at 23°C (A) and 37 °C (B). These data represent the mean ±
S.E.M for 3 determinations, where significant differences from 100% POPC (Model 1) vesicles
are represented by *, p ≤ 0.05; **, p ≤ 0.01; ***, p ≤ 0.001.

**Figure 5. FRET interaction of NBD-PE and Rh-PE in lipid vesicles mimicking the ER
membrane.** Degree of FRET interaction was calculated as stated in Materials and Methods for
vesicles composed of natural lipids mimicking the ER membrane. Measurements were done at
23°C (A) and 37 °C (B). These data represent the mean ± S.E.M for 3 determinations, where
significant differences from V-PC vesicles are represented by **, p ≤ 0.01; ***, p ≤ 0.001.

**Figure 6. Laurdan emission spectra in Model lipid vesicles.** Laurdan emission spectra (400
nm- 620 nm) were monitored using 350 nm excitation wavelength in lipid vesicles prepared
with model lipids. Scans are representative of 4 independent experiments.

**Figure 7. Laurdan emission spectra in lipid vesicles mimicking the ER membrane.** Laurdan
emission spectra (400 nm- 620 nm) were monitored using 350 nm excitation wavelength in lipid
vesicles prepared with lipids mimicking the ER membrane in the absence of CYP1A2 (solid
lines) and presence of 5µM CYP1A2 (dashed lines) as described in Materials and Methods.
Scans are representative of 4 independent experiments.

**Figure 8. Laurdan emission spectra and GP profiles as a function of temperature in model
systems.** Laurdan emission scans (400 nm- 620 nm) were monitored using 350 nm excitation as
a function of temperature. The effect of temperature increase on the laurdan emission spectra is illustrated from the Model 3 vesicle system (A). Generalized polarization values were determined for each lipid vesicle at each temperature (B). Emission scans are representative of 4 independent experiments and GP profiles are representative of the mean ± S.E.M for 4 determinations.

Figure 9. Laurdan emission spectra and GP profiles as a function of temperature in “natural membrane” vesicles. Laurdan emission scans (400 nm- 620 nm) were monitored using 350 nm excitation as a function of temperature. The effect of temperature increase on the laurdan emission spectra is illustrated from the V-ER system (A). Generalized polarization values were determined for each lipid vesicle at each temperature (B). Emission scans are representative of 4 independent experiments and GP profiles are representative of the mean ± S.E.M for 4 determinations.

Figure 10. CYP1A2 and CPR tryptophan quenching by Brominated-PC in V-PC and V-ER vesicles. The tryptophan fluorescence of enzymes (panel A: CYP1A2; panel B: CPR) was measured (excitation 295/ emission 341) in the presence (F) and absence (Fo) of 15 mol % BrPC in V-PC and V-ER systems. Fluorescence quenching (F/Fo) was measured as stated in Materials and Methods at 23°C and 37°C. These data represent the mean ± S.E.M for 4 determinations, where significant differences are represented by *, p ≤ 0.05; **, p ≤0.01.
Table 1. Lipid Composition (mol %) of Synthetic Reconstituted Systems. Purified lipid vesicles were prepared with the lipid composition listed in order to analyze the potential of lipid domain formation. Phosphatidylcholine in the “natural membrane” systems was a bovine liver extract\(^a\), whereas the phosphatidylcholine in the model systems was 1-palmitoyl-2-oleoyl-snglycero-3-phosphocholine (POPC)\(^b\). Sphingomyelin in the “natural membrane” systems was porcine brain sphingomyelin\(^c\), whereas in the Model systems, palmitoyl-sphingomyelin (PSM)\(^d\) was used.

<table>
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<th>Vesicle</th>
<th>PC</th>
<th>PE</th>
<th>PI</th>
<th>PS</th>
<th>PA</th>
<th>SM</th>
<th>Chol</th>
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<tr>
<td>PC</td>
<td>100(^a)</td>
<td>-</td>
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<td>-</td>
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<td>-</td>
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<tr>
<td>Mix PL</td>
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<td>10</td>
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<td>1</td>
<td>4(^c)</td>
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<td>-</td>
<td>5(^d)</td>
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<td>-</td>
<td>20(^d)</td>
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<tr>
<td>Model 4</td>
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<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>30(^d)</td>
<td>40</td>
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Table 2. Generalized polarization (GP) emission values. GP values from Figures 5 and 6 were determined at 23°C for each lipid system as described in Materials and Methods. These data represent the mean ± S.E.M for 3 determinations.

<table>
<thead>
<tr>
<th>Lipid</th>
<th>Generalized Polarization Values</th>
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<td>Model 2</td>
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<tr>
<td>Model 3</td>
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<td>Model 4</td>
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<td>- CYP1A2</td>
<td>+ CYP1A2</td>
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<td>V-PC</td>
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<tr>
<td>V-Mix PL</td>
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<td>V-ER</td>
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<td>V-DRM</td>
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</tr>
</tbody>
</table>
Figure 2
Figure 5

(A) 23°C

(B) 37°C

FRET (%) = \frac{(F - F_{PC})}{F_{PC}} \times 100

---

V-PC  V-Mix PL  V-ER  V-DRM

---

V-PC  V-Mix PL  V-ER  V-DRM
Figure 6

![Fluorescence Intensity vs Wavelength](image-url)
Figure 9

(A) Fluorescence intensity as a function of wavelength (nm) at different temperatures: RT, 30°C, 35°C, 40°C, 45°C, 50°C, 55°C, and 60°C.

(B) Generalized polarization as a function of temperature (°C): V-PC, V-Mix PL, V-ER, and V-DRM.
Figure 10

(A) CYP1A2

Relative Trp Fluorescence (F/F₀) x 100

23°C  37°C

V-PC  V-ER

**  *

(B) CPR

Relative Trp Fluorescence (F/F₀) x 100

23°C  37°C

V-PC  V-ER

*  *