



### Biophysics

DOI: 10.1002/ange.201713167 Deutsche Ausgabe: Internationale Ausgabe: DOI: 10.1002/anie.201713167

# **Cytochrome-P450-Induced Ordering of Microsomal Membranes Modulates Affinity for Drugs**

Zuschriften

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Abstract: Although membrane environment is known to boost drug metabolism by mammalian cytochrome P450s, the factors that stabilize the structural folding and enhance protein function are unclear. In this study, we use peptide-based lipid nanodiscs to "trap" the lipid boundaries of microsomal cytochrome P450 2B4. We report the first evidence that CYP2B4 is able to induce the formation of raft domains in a biomimetic compound of the endoplasmic reticulum. NMR experiments were used to identify and quantitatively determine the lipids present in nanodiscs. A combination of biophysical experiments and molecular dynamics simulations revealed a sphingomyelin binding region in CYP2B4. The proteininduced lipid raft formation increased the thermal stability of P450 and dramatically altered ligand binding kinetics of the hydrophilic ligand BHT. These results unveil membrane/ protein dynamics that contribute to the delicate mechanism of redox catalysis in lipid membrane.

Structure, function and regulation of membrane proteins are elicited through interaction with cell membrane.<sup>[1]</sup> A systematic "mapping" of lipid-protein interactions is one of the current challenges of molecular and cell biology, [1] and could represent a determinant advancement for understanding biological processes at membrane level. Indeed, more than 60% of drug targets are located at the cell surface and other membrane subcellular interfaces.<sup>[2]</sup> Drugs and xenobiotics detox functions are also regulated by membrane-anchored cytochrome P450s<sup>[3]</sup> located in the cytosolic side of the endoplasmic reticulum (ER); they have a broad catalytic activity which includes monooxygenation and dealkylation of a variety of ligands, including > 70% of commercially available drugs, as well as xenobiotics; [4] several P450 isoforms are also involved in hormone synthesis and the arachidonic acid cascade.[3] The biophysical challenges posed by the lipid membrane, as well as its molecular intricacies, have impeded a molecular characterization of P450-lipid dynamics. There is consensus about the significance of the lipid membrane in the catalytic activity of P450:<sup>[5]</sup> it provides the landscape for electron transfer to occur, through physical interaction with the electron transferase CPR. [6] Membrane anchoring is one of the speculated mechanism that prevents P450 trafficking towards other subcellular compartments.<sup>[7]</sup> Also, lipid membrane is the main access pathway for hydrophobic substrates, playing a critical role in P450 drug metabolism and pharmacokinetics.[8-12] There is mounting evidence that protein-protein interactions are also driven by cross-talking between transmembrane domains (TMD), [13-15] which further amplifies the role of lipids on the structural stability and function of P450. Thus, there is a need for new biophysical tools that can enlighten such presently precluded dynamics, and provide critical knowledge on the function of P450.

For membrane proteins, the information regarding subcellular organization in different membrane compartments is obtained through several methods, including indirect detergent-based assay (DRM), single-molecule imaging and spectroscopy, [16] and mass spectrometry. [17] Notwithstanding the critical role of membrane in P450 catalysis, only a few groups have addressed the fundamental questions regarding the organization of P450 in ER. DRM experiments reported that P450 isoforms and CPR likely to co-exist in ER microdomains.<sup>[18]</sup> Nonetheless, given the controversies associated with DRM, [19] as well as the more recent definition of lipid rafts as small (<10 nm) and transient SM-rich domains, [20] new strategies are needed, particularly to reveal the underestimated role played by proteins in driving lipid segregation. Since P450 strongly interacts with lipid membrane through its TMD and FG-loop, [15,21] we hypothesized that any proteindriven change in lipid composition could be trapped at the nanometric scale of a nanodisc.

The 4F peptide was able to form nanodiscs (see Figure S1 in the Supporting Information) that retained the ER membrane composition (Table S1) after SEC purification (Figure S1). Peptide-based nanodiscs have several advantages over membrane scaffolding protein nanodiscs (MSP) for studying membrane proteins. First, they allow detergent-free protein incorporation, which avoids post-reconstitution purification steps, [22,23] as shown by both size-exclusion chromatography (SEC, Figure S3a) and dynamic light scattering

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https://doi.org/10.1002/anie.201713167.

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(DLS, Figure S1b and S3b-d). Second, the surrounding nanodisc belt consists of 4F peptide units (ca. 20 per nanodisc);[22,23] compared to MSP nanodiscs, the size of peptide nanodisc is determined by the lipid:peptide ratio (Figure S1b). At 1:0.75 w/w lipid:peptide ratio, the diameter of the nanodisc was  $8 \pm 1$  nm, which is suitable for P450 monomerization.<sup>[23,24]</sup> When incorporated in 4F-nanodiscs, CYP2B4 showed a predominantly low-spin heme absorption spectrum (Figure S1d), which is indicative of negligible interactions between the heme prosthetic group and the lipid components. Contrary to MSP nanodiscs, peptide-based nanodiscs allow size-rearrangement and lipid exchange between nanodiscs.<sup>[22]</sup> Our approach based on the hypothesis that the presence of a membrane protein in a fraction of nanodiscs would perturb the exchange equilibrium in a way that reflects the ability of the protein itself to recruit or exclude specific membrane components in its proximate boundaries.

CYP2B4 was reconstituted in excess 4F-nanodiscs (Figure 1 a), the fraction containing the protein was subsequently purified by SEC (Figure 1 b), and lipids and cholesterol were quantified (Figure 2 a,b). The lipid composition uniquely reflects the perturbation induced by CYP2B4. The first observation is that, compared to the initial ER composition



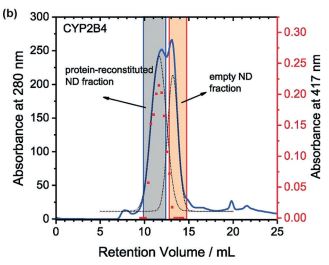


Figure 1. Peptide-based nanodiscs as a tool to unveil lipid-boundary regions of microsomal CYP2B4. a) Schematic of the lipid-exchange experiment. CYP2B4 was reconstituted with a 2.5-fold excess of empty nanodiscs and incubated overnight to allow lipid exchange. b) SEC profiles showing two partially overlapping peaks, representing the reconstituted and empty fractions, respectively; the protein-containing fraction was also characterized by the absorbance at the Soret maximum (417 nm).

("ER" in the rest of the manuscript), the lipid profiles were altered (Figure 2b). Nanodiscs after lipid exchange, ("ER<sub>ex</sub>") were enriched of SM (ca. 2.5-fold increase, p < 0.05) and cholesterol (ca. 1-fold increase, p < 0.001), and with a concomitant decrease of zwitterionic PE (ca. 1-fold, p < 0.01). Cytb<sub>5</sub>—whose cytosolic domain poorly interact with the membrane<sup>[5]</sup>—was used as a control, and as expected it did not perturb the lipid composition of the nanodiscs (Figure 2b). Lateral accumulation of saturated SM and cholesterol is indicative of raft-like liquid-ordered microdomain  $(l_0)$ formation. [25] Indeed, protein-induced lipid domains in nanodiscs were corroborated by fluorescence correlation spectroscopy. For this, we evaluated the partition of the fluorescent lipid probe DiI C<sub>12</sub> into nanodiscs as altered by the presence/ absence of P450 (Figure 2c and Figure S4). When SM-rich domains were formed, a significant change in the G(0) autocorrelation function amplitude was observed (Figure 2c and Table S2), indicating a different distribution of the fluorescent dye in presence of CYP2B4. Several control experiments performed on empty nanodiscs, as well as P450reconstituted 4F-POPC nanodiscs (Figure S4c) indicated that Dil C<sub>12</sub> was not interacting with protein backbone, or entering the active site of P450 (Figure S4d), making the hemeinduced fluorescence quenching unlikely.

Next, we investigated if an ordered region surrounding the protein was important to stability, structure and ligandaffinity of P450. A first striking evidence of a change in the membrane organization was the reduction in the nanodiscs size (ca. 2.2 nm, p < 0.05), likely indicating that the enrichment of SM and cholesterol in CYP2B4 caused the ERex bilayer to be more tightly "packed" (Figure S5a). CD results indicated, compared to the protein in solution, no appreciable changes in the folding (Figure S6a,b). Protein stability to thermal denaturation is an important parameter for assessing the effect of membrane on P450. [26] Compared to solution, CYP2B4 in 4F-ER nanodiscs resulted in an improved thermal stability with increased  $T_{\rm m}$  (ca. +4°C, Figure 2d). The formation of the  $l_{\rm o}$  domain (ER<sub>ex</sub>) further increased  $T_{\rm m}$  (ca. +4 °C), with  $\Delta H$  increasing from 118 to 163 kcal mol<sup>-1</sup> (p <0.05). Although a broader profile was observed for ER<sub>ex</sub>, and could be interpreted in terms of opening of the P450 binding pocket, additional structural data are needed to confirm this interpretation; further studies are under progress which will be published in the future.

In P450, conformational shifts in the protein backbone are disclosed in the active site environment, by subtle but observable changes in the heme absorbance spectrum. In P450, the predominance of low-spin state is due to water as the sixth ligand, [27,28] and limiting the access of water to the active site can lead to high-spin equilibrium shift (Figure S1d and 2e). These structural and dynamical fluctuations at the protein/lipid interface can potentially control the opening of access channels, playing a role in the recruitment of lipophilic substrates. [29-30] The affinity of P450 for drugs is a critical step in determining their pharmacokinetics, since the intrinsic clearance of a pharmacophore is determined by its metabolic rate and its affinity for the metabolizer. [12,31] We measured the spectral binding equilibrium constants ( $K_s$ ) by observing the ligand-induced spin shift in CYP2B4, when in solution or





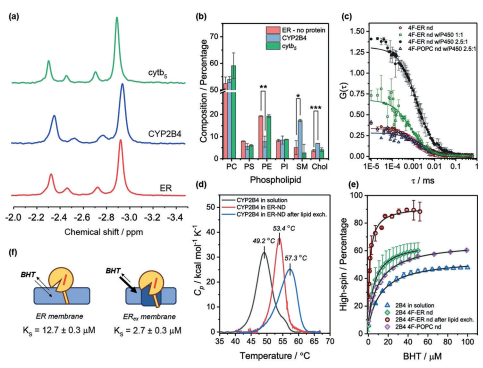


Figure 2. CYP2B4-induced liquid-ordered domain modulates drug affinity. a)  $^{31}$ P NMR on detergent-treated nanodiscs was used to assess the composition of the protein-containing fraction after overnight lipid exchange. b) Lipids and cholesterol contents in empty, CYP2B4, and cytb<sub>5</sub> 4F-ER nanodiscs after lipid exchange, as measured by  $^{31}$ P NMR and GC-MS. Data are expressed as average  $\pm$  standard deviation (n=3). \*p<0.05, \*\*p<0.05, \*\*\*p<0.01, \*\*\*\*p<0.001. c) Fluorescence autocorrelation functions for nanodiscs. Scatter plot represent average  $\pm$  standard deviation of two independent experiments; solid lines represent numerical fitting using a 3D correlation diffusion model. The lipid probe Dil C<sub>12</sub> differently partitioned in P450-containing nanodiscs, causing changes in the autocorrelation function. d) DSC curves. e) Protein-induced changes in lipid boundaries alter the affinity for BHT. f) Schematic of lipid-induced modulation of ligand affinity for BHT.

nanodiscs. We compared the effect of ligands, BHT and 4-CPI, which have different partitioning between membrane and bulk solution (their logP values are 5.54 and 1.99, respectively). UV/Vis titration with BHT led to several observations. First, in  $ER_{ex}$  nanodiscs, the initial high-spin population ([BHT] = 0) was higher than in solution (in ER + 15 %, p <0.05). Second, the equilibrium affinity  $K_s$  increased by 4-fold from solution (12  $\mu$ M) to ER<sub>ex</sub> nanodiscs (3  $\mu$ M), whereas in ER it was midway (8 μm). Third, the maximum high-spin fraction extrapolated by the sigmoidal binding curve showed a significant increase from solution (40%) to ER (60%); however, in  $ER_{ex}$  the  $l_o$  domain surrounding the protein increased the population to 100% high-spin. Similar experiments were performed with the more hydrophilic 4-CPI (Figure S7), a tight type II binder, which coordinates to the heme iron through its N-1 atom in the imidazole ring. When in solution, CYP2B4 showed lower affinity for 4-CPI ( $K_s$  =  $0.2 \,\mu\text{M}, \, p < 0.05$ ), with a sigmoidal binding curve indicative of multiple binding events, as reported. [32] The reconstitution of CYP2B4 into nanodiscs attenuated the sigmoidal shape of the binding curve and increased the affinity by 5-fold. No significant differences were observed in the affinities among the lipid nature, with the  $K_s$  values in the 0.04–0.07  $\mu$ m range (Figure S7). Hydrophobic ligands, such as BHT, are thought to partition into the membrane and reach the prosthetic group towards the 2a channel, situated between the FG loop which dips into the lipid membrane. On the other hand, hydrophilic ligands—such as 4-CPI—poorly interact with the membrane layer and are thought to enter the 2c channel located between the B' helix/BC loop on the protein cytosolic domain.

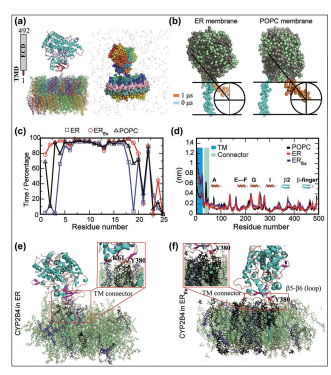
Structural characterization membrane proteins in a closely native lipid environment is not facile.[33] Alternatively, MD simulation on P450s embedded in homogeneous membrane mimetic, [9,21,34] has supplied insights into protein/ lipid interfaces, despite not fully representing the physiological lipids. We performed both all-atom and coarsegrained (CG) simulations on CYP2B4 embedded in POPC, ER and ER<sub>ex</sub> membranes. The initial TMD of CYP2B4 was oriented perpendicular to the bilayer surface for dynamics interpretation (Figure 3 a).<sup>[9]</sup> Under a 100 ns all-atom simulation, CYP2B4 showed no substantial alteration of protein folding around the catalytic

site as revealed from its backbone RSMD. The CYP2B4 soluble domain depicted a constrained conformation with an average RMSD of 2.3 (ER), 2.0 (ER $_{\rm ex}$ ) and 2.9 Å (POPC) in comparison to its full-length conformation. In contrast, the RMSD of C $_{\alpha}$  atoms of the full-length CYP2B4 were higher than 2.5 Å and the average RMSD were 3.1 (ER), 4.2 (ER $_{\rm ex}$ ) and 3.6 Å (POPC). This suggested that TMD and lipid composition greatly modulate the overall CYP2B4 dynamics, and the lipid composition modulates the dynamics of both TMD and soluble domain.

To gain further insight, we performed CG-MD at microsecond time scale. As anticipated from the all-atom MD, a comparatively high TMD dynamics was observed in POPC (Figure 3b). The perpendicularly oriented TMD-to-bilayer surface ended with a tilt of ca. 45° during 10  $\mu$ s MD simulations for both ER and POPC membrane nanodiscs. However, the translational motion of TMD (from center) was much larger in POPC and was restricted by the 4F-peptide belt. In contrast, the translational motion was well-restrained in ER membrane nanodiscs during multi-microsecond MD calculation and was not found in close proximity to the belt (Figure 3b). Transient interaction between CYP2B4's soluble domain and lipid membrane was consistently observed in all simulations. Simulations showed that in SM-enriched ER membrane (ER<sub>ex</sub>), the M1-F20 is an  $\alpha$ -helix along the entire







**Figure 3.** Atomistic insights into the full-length CYP2B4 interaction with ER membrane. a) All-atom (left) and coarse-grained (right) models of CYP2B4. Protein and lipids are shown as cartoon and stick, respectively, in the all-atom model, whereas as spheres and dotted spheres in the CG-MD models. b) CG-MD snapshots showing the TMD orientation in POPC and ER nanodiscs at 0 (cyan) and 10 (orange) μs. c) Percentage of helicity rise per residue in CYP2B4-TMD with simulation time. d) Root mean square fluctuation of CYP2B4. TMD and linker connecting TMD and soluble domain are highlighted in blue and cyan, respectively. e) MD snapshot illustrating the interaction between CYP2B4 and ER, and f) SM and cholesterol-rich ER<sub>Ex</sub>. SM and POPS are shown in black and purple, respectively.

course of simulation, whereas in both POPC and ER the M1-L5 residues were mostly disordered (Figure 3c). Monitoring the RMS fluctuation of backbone atoms of CYP2B4 in POPC, ER and ER<sub>ex</sub> membranes (Figure 3d), it was found that RMS > 2.0 Å were observed for TMD, whereas the mobility of the proline-rich (P36-P49) loop was significant only in POPC. Loops connecting helix motifs in the soluble domain showed similar fluctuations over time. In the ER membrane, the FG-loop (bearing the F' and G' helices), as well as  $\beta$ 2,  $\beta$ 5 and  $\beta$ 6 sheets (also known as  $\beta$ -finger) close to the C-terminus presented higher rigidity, due to an increased interaction with the membrane hydrophobic core (Figure 3 d).

All-atom MD simulations unveiled interactions between specific residues and lipids (Figure 3). In ER, the soluble domain was found to interact with lipid head groups, in particular the FG-loops and the  $\beta$ -finger loop (Figure 3e). Results also showed SM clustering around the TMD-connecting loop and the  $\beta$ -finger, corresponding to the region spanning the T375 to I382 chain. Further structure analysis showed H-bonding interaction between SM and Y380 (Figure 3e, insert). Simulations on ER<sub>ex</sub> showed a dramatic increase of SM clustering, with additional specific interactions with the Pro-rich TM connecting loop (G28-L40). The TM  $\alpha$ -

helical character of the domain was calculated over the 100-ns simulation time (Figure S8). Compared to ER, the enrichment of SM and cholesterol in ERex caused the helix to have an ideal angle (100°) along the entire simulation time (Figure S8a). Finally, we calculated the number of H-bonds over simulations in both ER systems, as a tool to evaluate the interaction with polar groups of the lipids (Figure S8b). As expected, the number of H-bonds was found to be substantially higher in SM-enriched ER<sub>ex</sub>, indicating more interaction with the membrane surface. Sequence alignment shows that both TMD connecting region and β-finger are largely conserved among microsomal P450s (Figure S9). Particularly, the β-finger has several conserved hydrophobic residues, including the 382 Ile-Pro-Lys384 sequence, as well as the prolinerich connector domain. This leads us to hypothesize that other microsomal P450s can possess similar ability of specific interaction with lipid components.

In conclusion, this study reveals that P450s is highly dynamic, and able to interact and modify its surrounding lipid environment to enhance both function and stability. Proteininduced formation of SM-rich domain could facilitate P450 monomerization and stability. Recent experimental and computational evidences on the lateral organization of microsomal cytochrome P450 pointed to a strong interaction of both TMD and soluble domain with the lipid bilayer. [13,15,21,29,33] The results presented here also provide significant molecular insights into the factors governing P450 ligand affinity for hydrophobic molecules. Further, the innovative combination of peptide-based nanodiscs and <sup>31</sup>P NMR experiments to quantitatively determine the different type of lipids constituting nanodiscs will be useful to study a variety of membrane proteins (like GPCR and poreforming proteins<sup>[35]</sup>) and membrane-assisted amyloid aggregation.

#### **Acknowledgements**

This study was supported by NIH (GM084018 to A.R.). We thank Sarah Cox for help with TEM, Matthew Schweiss for assistance with GC-MS, the Biophysics SMART Center and Damon Hoff for assistance with Alba confocal microscope.

### Conflict of interest

The authors declare no conflict of interest.

**Keywords:** biophysics  $\cdot$  hemeproteins  $\cdot$  lipids  $\cdot$  membranes  $\cdot$  nanodiscs

**How to cite:** *Angew. Chem. Int. Ed.* **2018**, *57*, 3391–3395 *Angew. Chem.* **2018**, *130*, 3449–3453

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Manuscript received: December 27, 2017 Revised manuscript received: January 16, 2018 Accepted manuscript online: January 31, 2018 Version of record online: February 22, 2018