Effect of Packing Density on Rhodopsin Stability and Function in Polyunsaturated Membranes

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ABSTRACT  Rod outer segment disk membranes are densely packed with rhodopsin. The recent notion of raft or microdomain structures in disk membranes suggests that the local density of rhodopsin in disk membranes could be much higher than the average density corresponding to the lipid/protein ratio. Little is known about the effect of high packing density of rhodopsin on the structure and function of rhodopsin and lipid membranes. Here we examined the role of rhodopsin packing density on membrane dynamic properties, membrane acyl chain packing, and the structural stability and function of rhodopsin using a combination of biophysical and biochemical techniques. We reconstituted rhodopsin into large unilamellar vesicles consisting of polyunsaturated 18:0:22:6n3PC, which approximates the polyunsaturated nature of phospholipids in disk membranes, with rhodopsin/lipid ratios ranging from 1:422 to 1:40. Our results showed that increased rhodopsin packing density led to reduced membrane dynamics revealed by the fluorescent probe 1,6-diphenyl-1,3,5-hexatriene, increased phospholipid acyl chain packing, and reduced rhodopsin activation, yet it had minimal impact on the structural stability of rhodopsin. These observations imply that densely packed rhodopsin may impede the diffusion and conformational changes of rhodopsin, which could reduce the speed of visual transduction.

INTRODUCTION

G-protein coupled receptors (GPCRs) are the largest family of membrane surface receptors and respond to a broad spectrum of extracellular stimuli, e.g., hormones, neurotransmitters, odor and taste molecules, light, etc., and generate intracellular responses by coupling to G-proteins (1,2). A prominent member of the GPCR superfamily is the photoreceptor rhodopsin. A high-resolution structure has been obtained and its function in visual signaling is well characterized (3–5). Upon light activation, rhodopsin rapidly converts into the active conformation, metarhodopsin II (MII). Each MII binds and activates several hundred of the visual G-protein transducin in a process that depends upon lateral diffusion. The activated transducin subsequently activates the effector enzyme, cGMP phosphodiesterase, triggering the hydrolysis of cGMP. The decline of cytosolic cGMP leads to the closure of the cGMP gated Na+/Ca2+ channel and hyperpolarization of the plasma membrane, and eventually triggers the visual response.

Over 90% of the membrane proteins in rod disk membranes are rhodopsin. Rhodopsin is densely packed in disk membranes with an average packing density of ~25,000 rhodopsin/μm² (6). Recent high-resolution atomic force microscopy images of disk membranes revealed rhodopsin molecules to be unevenly distributed in disk membranes (7). The packing density of rhodopsin in dense regions is about twice as high as the overall average density. At this point, the physiological relevance of such densely packed rhodopsin molecules observed in atomic force microscopy images is somewhat controversial (8,9), and it is unknown what structural and functional role the high packing density of rhodopsin may play. High packing density of rhodopsin in disk membranes endows the rod cell with high sensitivity, but may also create a slower visual response, which is limited by the diffusion-controlled protein coupling in the visual cascade (10,11). A recent in vivo study established that a reduction of rhodopsin density in disk membranes, which reduces the molecular crowding in membranes, increased the speed of visual transduction (12).

The effect of membrane protein packing density on the dynamic and equilibrium properties of lipid membranes has been studied for many membrane proteins (13–17). Several of these studies were carried out in membranes consisting of phospholipids with disaturated acyl chains, which are not found in abundance in biological membranes. The phospholipid acyl chains in disk membranes consist largely of polyunsaturated fatty acids. About half of the phospholipid acyl chains in rod disk membranes are made of docosahexaenoic acid (22:6n3) (18). These docosahexaenoic acid chains provide a maximal fluidity or maximal free volume of acyl chain packing in disk membranes, which is responsible for optimal visual function (19–23). The effect of membrane protein packing density on membrane physical properties and protein function in membranes consisting of polyunsaturated lipids is largely unknown.

In this study we reconstituted rhodopsin into vesicles consisting of 18:0:22:6n3PC at rhodopsin/lipid ratios varying from 1:422 to 1:40. The membrane physical properties, including acyl chain packing and membrane dynamics, were
examined using time-resolved 1,6-diphenyl-1,3,5-hexatriene (DPH) fluorescence, the thermal stability of rhodopsin was measured by differential scanning calorimetry (DSC), and the activity of rhodopsin was quantified by the equilibrium between the metarhodopsin I (MI) and MII conformational states. We found that increased rhodopsin density led to increased acyl chain packing order and reduced rhodopsin activation, whereas it had minimal impact on rhodopsin thermal stability. The importance of these findings for visual transduction and their implications for the effects of membrane crowding in membrane structure and function will be discussed.

**MATERIALS AND METHODS**

**Sample preparations**

Concanavalin A sepharose was purchased from Amersham Biosciences (Piscataway, NJ). Phospholipid 18:0,22:6n3PC was obtained from Avanti Polar Lipids (Alabaster, AL). The fluorescent probe DPH was from Molecular Probes (Eugene, OR). Bovine retinas were from James and Wanda Lawson (Lincoln, NE).

Rod outer segments (ROS) were isolated from bovine retinas using the modified Shake-ate method (24). Rhodopsin was purified from the concanavalin A affinity column as described (25). Rhodopsin-containing vesicles consisting of 18:0,22:6n3PC with varied rhodopsin/lipid ratios were prepared using the rapid dilution method (26), which yielded vesicle sizes on the order of 100 nm. The phospholipid 18:0,22:6n3PC contains six double bonds at the sn-2 chain, which is highly susceptible to oxidation. Such oxidation was minimized by addition of antioxidant butylated hydroxytoluene (BHT) and removal of oxygen during preparation. BHT was added into samples at a lipid/BHT ratio of 500:1. All sample preparations were conducted in an argon-filled glove box and all buffers were degassed with argon gas before use. The final samples were suspended in Pipes-buffered PBS before loading into the sample cell. The dialysis buffer was loaded into the reference cell as a reference buffer. Samples were loaded under argon in complete darkness using infrared night vision goggles. The cells were sealed in the dark and maintained at a pressure of 2.8 atm to maintain a stable baseline. Samples were scanned at a scan rate of 0.25°C/min from 45°C to 85°C. A second heating scan was repeated after the sample was cooled and equilibrated at 20°C. DSC scans were converted into excess molar heat capacity ($C_p$) using Cp-Cal 2.1 (Calorimetry Sciences), and the $C_p$ curves were analyzed for $T_m$ and $\Delta H$ using Origin Lab (Northampton, MA) as described (31).

**Fluorescence measurements**

Vesicles were suspended in PBS buffer at a phospholipid concentration of 0.1 mM. DPH was dissolved in tetrahydrofuran and added at a phospholipid/DPH ratio of 500:1. Fluorescence lifetime and differential polarization measurements were performed with a K2 multifrequency cross-correlation phase fluorometer (ISS, Urbana, IL). Excitation at 351 nm was provided by an Innova 307 argon ion laser (Coherent, Santa Clara, CA). All measurements were conducted at 37°C in PBS buffer. Fifteen modulation frequencies, logarithmically spaced from 5 to 150 MHz were used for both lifetime and differential polarization measurements. For lifetime measurements, the emission polarizer was set at the magic angle of 54.7° relative to the vertically polarized excitation beam, and $p$-bis[2-(5-phenyloxazolyl)]-benzene in absolute ethanol was used as a lifetime reference. For each differential polarization measurement the instrumental polarization factors were measured, found to be between 1 and 1.05, and the appropriate correction factor applied. Scattered excitation light was removed from the emission beam by a 390-nm highpass filter. At each frequency, data was accumulated until the standard deviations of the phase and modulation ratio were below 0.2° and 0.004, respectively, and these values were used as the standard deviation for the measured phase and modulation ratio in all subsequent analysis. Both total intensity decay and differential polarization measurements were repeated a minimum of three times with each sample.

Total fluorescence intensity decays were analyzed with the sum of two exponential decays. The resulting time constants were averaged with intensity weighting and reported as $\langle r \rangle$. Measured polarization-dependent differential phases and modulation ratios for each sample were combined with the measured total intensity decay to yield the anisotropy decay, $r(t)$. All anisotropy decay data were analyzed using the Brownian rotational diffusion (BRD) model (28), which yields the order parameters (P2) and (P4). These two order parameters were used to construct an orientational distribution function, $f(\theta)$, of the probe molecule. In general, the orientation of a molecule with cylindrical symmetry in lipid vesicles is completely described by the angle, $\theta$, between its symmetry axis and the local membrane normal. The results of the BRD model-based analysis were interpreted in terms of an angular distribution function, which is symmetric about $\theta = \pi/2$, $f(\theta)$. The extent to which the equilibrium orientational freedom of DPH is restricted by the phospholipid acyl chains was quantified using the parameter, $f_c$ (29,30). DPH fluorescence anisotropy decays were also analyzed using an empirical sum-of-three-exponentials model. The results of this analysis are reported as the weighted average of the three rotational correlation times, $\langle \phi \rangle$ (30). All analyses of differential polarization data were performed with NONLIN, with subroutines specifying the fitting function written by the authors.

**Differential scanning calorimetry**

Differential scanning calorimetry measurements were performed with a Nano-scan II calorimeter equipped with capillary cells (Calorimetry Sciences, Provo, UT). Rhodopsin-containing vesicles were dialyzed in PBS buffer before loading into the sample cell. The dialysis buffer was loaded into the reference cell as a reference buffer. Samples were loaded under argon in complete darkness using infrared night vision goggles. The cells were sealed in the dark and maintained at a pressure of 2.8 atm to maintain a stable baseline. Samples were scanned at a scan rate of 0.25°C/min from 45°C to 85°C. A second heating scan was repeated after the sample was cooled and equilibrated at 20°C. DSC scans were converted into excess molar heat capacity ($C_p$) using Cp-Cal 2.1 (Calorimetry Sciences), and the $C_p$ curves were analyzed for $T_m$ and $\Delta H$ using Origin Lab (Northampton, MA) as described (31).

**MI-MII equilibrium measurements**

The equilibrium spectra of MI and MII were collected and deconvolved according to Straume et al (32). Briefly, rhodopsin-containing vesicles were diluted to 0.3 mg/ml in pH 7.0 PBS buffer and equilibrated at 37°C in a thermally regulated sample holder. A set of four absorption spectra were collected sequentially in an HP 8453A diode array spectrophotometer. These included the spectra acquired 1) after the sample was equilibrated at 37°C; 2) 3 s after the sample was 15–20% bleached by a 520-nm flash; 3) 10 min after addition of 30 mM hydroxyamine to remove bleached rhodopsin; and 4) after complete bleach of the sample. Individual MI and MII spectra were deconvolved from their equilibrium mixture using a nonlinear least-squares method, and the equilibrium constant, $K_{eq}$, was calculated according to $K_{eq} = [MII]/[MI]$.

**RESULTS**

**DPH fluorescence**

The fluorescence intensity decay of DPH yielded two exponential time constants (Table 1). The major contribution of DPH fluorescence decay was from the longer lifetime decay. Increasing the rhodopsin/lipid ratio, which increases
TABLE 1 Summary of DPH fluorescence lifetime in reconstituted rhodopsin/18:0,22:6n3PC membranes

<table>
<thead>
<tr>
<th>Rhodopsin/18:0,22:6n3PC</th>
<th>$a_1$</th>
<th>$\tau_1$ (ns)*</th>
<th>$a_2$</th>
<th>$\tau_2$ (ns)*</th>
<th>$\langle \tau \rangle$ (ns)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:422</td>
<td>0.87</td>
<td>7.05 ± 0.07</td>
<td>0.13</td>
<td>1.15 ± 0.10</td>
<td>6.92 ± 0.05</td>
</tr>
<tr>
<td>1:184</td>
<td>0.75</td>
<td>5.34 ± 0.16</td>
<td>0.25</td>
<td>1.11 ± 0.08</td>
<td>5.07 ± 0.15</td>
</tr>
<tr>
<td>1:94</td>
<td>0.72</td>
<td>5.07 ± 0.20</td>
<td>0.28</td>
<td>1.00 ± 0.11</td>
<td>4.78 ± 0.17</td>
</tr>
<tr>
<td>1:40</td>
<td>0.65</td>
<td>4.02 ± 0.08</td>
<td>0.35</td>
<td>0.78 ± 0.03</td>
<td>3.72 ± 0.07</td>
</tr>
</tbody>
</table>

The fluorescence intensity decay of DPH was analyzed with the sum of two exponential decays, which yielded $\tau_1$ and $\tau_2$, $a_1$ and $a_2$ are the fractions of the two exponential decays. $\langle \tau \rangle$ is the intensity-weighted average lifetime.

*DPH lifetime is expressed as ± SD obtained from three independent measures.

rhodopsin packing density in membranes, resulted in progressive quenching of the longer lifetime decay of DPH and reduced the overall average lifetime, $\langle \tau \rangle$. At the lowest rhodopsin/lipid ratio (1:422), the value of $\langle \tau \rangle$ was 6.92 ± 0.05 ns, which is close to that found in pure 16:0,22:6n3PC membranes measured under similar conditions (30). As the rhodopsin/lipid ratio increased to 1:40, $\langle \tau \rangle$ was reduced by ∼50%.

Rotational dynamics of DPH probe in membranes was obtained from the analysis of DPH differential polarization data using two different methods: the empirical sum-of-exponential model and the BRD model as described in the Materials and Methods section. The average rotational correlation time, $\langle \phi \rangle$, was ∼1.0 ns in 18:0,22:6n3PC membranes with the lowest rhodopsin density (Fig. 1 A), in agreement with the value obtained in pure 16:0,22:6n3PC membranes (30), indicating rapid rotational motion of DPH in membranes. Increasing the packing density of rhodopsin in membranes hindered the rotational motion of DPH, as demonstrated by the progressive increase of $\langle \phi \rangle$. The rotational diffusion coefficient of DPH, $D_\perp$, obtained from the BRD model also showed that an increase of rhodopsin density in membranes led to slower rotational motion of DPH (Fig. 1 B). There was an ∼2-fold reduction of $D_\perp$ between the samples with lowest and highest rhodopsin/lipid ratios (1:422 and 1:40), further demonstrating that the DPH diffusion is impeded by increased rhodopsin density in membranes.

Analysis of DPH fluorescence anisotropy decays in terms of the BRD model yields the orientational distribution function for DPH in membranes, $f(\theta)$. Comparison of the orientational probability functions, $f(\theta)\sin\theta$, for membranes with varying levels of rhodopsin showed the effect of membrane protein density on equilibrium acyl chain order (Fig. 2 A). The BRD model depicts DPH distribution in membranes as two orthogonal populations that are perpendicular and parallel to the membrane normal (30). Increased rhodopsin density in membranes reduced the DPH population perpendicular to the membrane normal and increased the DPH population parallel to the membrane normal. At a rhodopsin/lipid ratio of 1:422, ∼50% of the DPH population is in the population that is distributed perpendicular to the membrane normal (Fig. 2 B). This fraction was reduced to ∼30% at a rhodopsin/lipid ratio of 1:40, indicating the ordering of membranes by rhodopsin. The overall acyl chain packing free volume in membranes, which is summarized by the parameter $f_v$, was sensitive to the rhodopsin density in membranes. As the rhodopsin density increased, a progressive reduction of $f_v$ was observed (Fig. 3), indicating the ordering of membrane packing by rhodopsin.

Rhodopsin thermal stability

Previous studies show that the thermal stability of rhodopsin is regulated by the composition of lipids in membranes (31,33,34). Rhodopsin is stabilized in membranes consisting of more saturated phospholipids or in the presence of cholesterol. The stabilization of rhodopsin is manifested by the
increase of the denaturation temperature of rhodopsin. For all reconstituted rhodopsin samples studied, the thermal denaturation of rhodopsin exhibited a single thermal transition peak (Fig. 4). The transition temperature of rhodopsin denaturation was centered around 68.5°C for all rhodopsin/lipid ratios, indicating that the number of lipid molecules surrounding rhodopsin had no influence on the structural stability of rhodopsin. The denaturation enthalpy was slightly increased by the increase of rhodopsin density in membranes (Table 2). There is a ~14% difference of ΔH between the samples with the lowest and highest rhodopsin/lipid ratios. The ΔH at the highest rhodopsin/lipid ratio is close to that in native disk membranes (35,36).

**Rhodopsin activation**

The effect of rhodopsin packing density on rhodopsin activation was evaluated by the level of activated rhodopsin, which is quantified by the equilibrium constant, $K_{eq}$, for the MI-MII equilibrium. Rhodopsin was functional at all rhodopsin/lipid ratios, as demonstrated by the ability to form the active MII conformation (Fig. 5 A). However, the level of MII formation was influenced by the packing density of rhodopsin in membranes; rhodopsin is most active in membranes with the lowest rhodopsin packing density. Increased...
rhodopsin density in membranes led to reduced \( K_{eq} \) values. The reduction of \( K_{eq} \) was more dramatic at higher rhodopsin density. An ~2-fold increase of rhodopsin/lipid ratio from 1:422 to 1:184 resulted in a 9% reduction of \( K_{eq} \), whereas a similar increase of rhodopsin/lipid ratio from 1:94 to 1:40 resulted in a 19% reduction of \( K_{eq} \).

The decline of \( K_{eq} \) induced by the increase of rhodopsin density in membranes correlated with the decline of \( f_v \). Such correlation between \( K_{eq} \) and \( f_v \) was observed in many previous studies (34,37,38). Since the conversion of MI to MII involves large conformational change (39), membranes with lower \( f_v \) would impose higher energy barrier for such conformational change, thus a lower level of MII formation is expected. However, the correlation between \( K_{eq} \) and \( f_v \) was nonlinear, which differs from previous observations. \( K_{eq} \) values approached a plateau at higher values of \( f_v \). This suggests that changes in membrane acyl chain ordering produced by increasing protein density and by changes in acyl chain composition are not identical.

**DISCUSSION**

Recent studies suggest that disk membranes are laterally heterogeneous and contain microdomains or lipid rafts (7,40,41). In rhodopsin-dense regions, the average rhodopsin density is ~2-fold higher than the overall average rhodopsin density in the ROS disk; 48,300 rhodopsin/\( \mu \text{m}^2 \) compared to 25,000 rhodopsin/\( \mu \text{m}^2 \) (7). At present, it is not clear how such a heterogeneous structure is formed in disk membranes or whether such regions of densely packed, paracrystalline rhodopsin are physiologically relevant (8). An earlier DSC study observed a broad phase transition in ROS disks and lipid extract from ROS disks (42). This leads to the possibility that laterally segregated microdomains may be induced by a nonphysiological temperature below the broad phase transition observed by Miljanich et al. (42). In this study, we determined the structural and functional consequences of densely packed rhodopsin in polyunsaturated membranes. Our results showed that high packing density of rhodopsin slows the dynamics of the fluorescence probe DPH and increases the order of phospholipid acyl chain packing, producing reduced rhodopsin activation. This suggests that densely packed rhodopsin in the ROS disk membranes would lower visual transduction by retarding lateral diffusion and reducing the amount of rhodopsin in the active MII conformation.

**Effect on membrane properties**

A 10-fold increase of the rhodopsin/18:0, 22:6n3PC ratio resulted in an ~50% reduction of the DPH rotational diffusion coefficient and an ~50% reduction in \( f_v \). This is in general agreement with the reduced membrane dynamics and increased membrane acyl chain packing order observed in previous studies, which used DPH fluorescence anisotropy to assess ensemble acyl chain packing (13,14). There are 22–24 phospholipid molecules located in the annular lipid layer surrounding rhodopsin (43,44). Those phospholipids have hindered motion due to direct interaction with rhodopsin (44–47). At rhodopsin/lipid ratios ranging from 1:422 to 1:40, ~6% to 60% of the phospholipids are located in the

### TABLE 2 Summary of the thermal denaturation measurements of rhodopsin

<table>
<thead>
<tr>
<th>Rhodopsin/18:0,22:6n3PC</th>
<th>( T_m ) (°C)</th>
<th>( \Delta H ) (kcal/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:422</td>
<td>68.5 ± 0.1</td>
<td>137.9 ± 0.5</td>
</tr>
<tr>
<td>1:184</td>
<td>68.0 ± 0.1</td>
<td>144.1 ± 0.8</td>
</tr>
<tr>
<td>1:94</td>
<td>68.1 ± 0.1</td>
<td>148.4 ± 0.6</td>
</tr>
<tr>
<td>1:40</td>
<td>68.4 ± 0.1</td>
<td>157.9 ± 0.8</td>
</tr>
</tbody>
</table>

Values are expressed as ± SD propagated from the analysis of DSC thermograms.
annular layer. This increase in the percentage of acyl chains in contact with rhodopsin with increasing protein content would be expected to produce more ordered ensemble acyl chain packing and slower membrane dynamics detected by DPH. However, there is evidence that the effects of transmembrane protein on acyl chain order may depend upon the degree of acyl chain unsaturation. A recent $^3$H-NMR study of rhodopsin in 16:0,18:1PC recombinant membranes (rhodopsin/lipid ratio of 1:100) reported that rhodopsin slightly decreased the order of segments $C_\text{pr}-C_{15}$ of the sn-1 chain in 16:0,18:1PC (48). The observed difference between 16:0,18:1PC and 18:0,22:6n3PC suggests that a membrane containing 22:6n3 acyl chains responds differently to the presence of a transmembrane protein than a membrane composed of monounsaturated acyl chains. The difference could also be due to the different nature of the DPH fluorescent technique and the $^3$H-NMR measurement. The DPH fluorescence measurements used in this study reflect the dynamic property of DPH in membranes on a nanosecond timescale, whereas the 2H-NMR order parameter measurement reports the averaged correlation time of the C-D bond on a microsecond timescale.

The change of DPH rotational diffusion induced by increased rhodopsin density in 18:0,22:6n3PC membranes is small compared to the changes in membrane diffusional properties observed in other reconstituted membranes consisting of disaturated phospholipids (13,14,49). This suggests that polyunsaturated lipids may dampen the reduction of membrane diffusional properties induced by increased rhodopsin density.

**Effect on receptor stability**

Rhodopsin is a transmembrane protein that has direct contact with lipid membranes, thus it is not difficult to envision that lipid membranes would influence its stability. Previous studies show that membrane composition modulates rhodopsin thermal stability (33,34). Membranes consisting of more saturated phospholipids or cholesterol, which have more ordered acyl chain packing, stabilize rhodopsin with respect to thermal denaturation. Thus, the denaturation temperature of rhodopsin is higher in these membranes. In this study we observed that increased rhodopsin density in membranes caused reduction of ensemble acyl chain packing free volume, $f_v$. However, the denaturation temperature of rhodopsin remained unchanged. Raising the number of lipids per rhodopsin increases the number of nonannular or bulk lipids that are not in contact with rhodopsin, but rhodopsin is surrounded by the same lipid, 18:0,22:6n3PC. The nature of the rhodopsin-lipid interaction remains the same regardless of the variation of rhodopsin/lipid ratio. The thermal denaturation measurements demonstrate that the thermal stability of rhodopsin is independent of the acyl chain packing properties of the membrane or how many lipids are in the bulk. Rather, rhodopsin thermal stability is dependent upon the composition of the phospholipids that interact directly with rhodopsin.

**Effect on receptor function**

The initial step in rhodopsin activation is the transient formation of the MI and MII equilibrium. The conformational change from MI to MII involves a volume expansion (39), which is restricted by the surrounding lipid matrix. Previous studies demonstrate that the conformation change from MI to MII is modulated by phospholipid acyl chain composition (21–23,38,50–53), membrane cholesterol (32,34,37), and phospholipid headgroup composition (53–55). The effects of phospholipid acyl chain composition and cholesterol in a PC membrane on MII formation are related to ensemble acyl chain packing, as indicated by the linear correlation between MII formation and $f_v$ (32,37,38). This empirical relationship between ensemble acyl chain packing and MII formation in bilayers with a low level of curvature stress may represent a special case of the more generalized flexible surface model proposed by Brown and co-workers (48,53,55). By considering bilayer deformation in terms of the curvature of the individual monolayer, the flexible surface model provides an elegant description of the energetics involved in the stabilization of MII by lipids with a tendency to form nonlamellar phases, such as phosphatidylethanolamines.

**CONCLUSIONS**

In this study, we focused on the effect of rhodopsin density on rhodopsin activation in PC-containing membranes. Our results showed that the energy barrier is higher in membranes with lower $f_v$ values; thus, the formation of MII is reduced. In this study, increased rhodopsin density resulted in reduction of $f_v$ in membranes and a reduced level of MII formation, in general agreement with the previously observed correlation between MII formation and $f_v$ (21,23,34,38,56). However, these earlier studies always found a strong linear relationship between $K_{eq}$ and $f_v$ for changes induced by temperature or membrane composition. In this study, changes in rhodopsin density produced a nonlinear correlation between $K_{eq}$ and $f_v$. Fig. 5 B. This difference in the $K_{eq}$-$f_v$ relationship is likely due to the different environments where DPH is located. The linear correlation between $K_{eq}$ and $f_v$ was observed in membranes with fixed protein/lipid ratios, whereas in this study the protein/lipid ratio was varied. At higher protein/lipid ratios, the majority of DPH probes are located in the vicinity of rhodopsin, whereas at lower protein/lipid ratios, a large fraction of the DPH fluorescence signal originates from probe molecules distant from rhodopsin. Thus, in membranes with a high number of lipids per rhodopsin, DPH may detect a greater increase in $f_v$ than is sensed by the MI-MII transition, and the plot of $K_{eq}$ versus $f_v$ approaches a plateau, as seen in Fig. 5 B.
Our results show that there is a clear difference in how rhodopsin packing density affects rhodopsin stability and rhodopsin function. The structural stability of rhodopsin is determined by the nature of the lipid-protein interaction at the lipid-protein interface, dependent on lipid composition and independent of protein/lipid ratio. In contrast, the conformational change from MI to MII is strongly correlated with membrane free volume, dependent on both lipid composition and the protein/lipid ratio. This study also demonstrates that increased membrane protein density reduces membrane dynamic properties, increases membrane acyl chain packing order, and lowers rhodopsin activation. These changes were generally less than changes observed in membranes with saturated phospholipids, suggesting that the presence of 22:6n3 attenuates these changes. Calvert et al. (12) demonstrated that reduced rhodopsin density in vivo increased the speed of visual transduction, and interpreted the increase in speed as an enhancement in lateral diffusion. Our results support their interpretation and suggest that the enhanced production of MII could be an additional factor in their finding of increased speed of visual transduction.

REFERENCES


