2D-ELDOR Study of Heterogeneity and Domain Structure Changes in Plasma Membrane Vesicles upon Cross-Linking of Receptors

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ABSTRACT: 2D electron–electron double resonance (2D-ELDOR) with the “full S” method of analysis is applied to the study of plasma membrane vesicles. Membrane structural changes upon antigen cross-linking of IgE receptors (IgE-FcεRI) in plasma membrane vesicles (PMVs) isolated from RBL-2H3 mast cells are investigated, for the first time, by means of these 2D-ELDOR techniques. Spectra of 1-palmitoyl-2-(16-doxyl stearoyl) phosphatidylcholine (16-PC) from PMVs before and after this stimulation at several temperatures are reported. The results demonstrate a coexistence of liquid-ordered (L_o) and liquid-disordered (L_d) components. We find that upon cross-linking, the membrane environment is remodeled to become more disordered, as shown by a moderate increase in the population of the L_d component. This change in the relative amount of the L_o versus L_d components upon cross-linking is consistent with a model wherein the IgE receptors, which when clustered by antigen to cause cell stimulation, lead to more disordered lipids, and their dynamic and structural properties are slightly altered. This study demonstrates that 2D-ELDOR, analyzed by the full S method, is a powerful approach for capturing the molecular dynamics in biological membranes. This is a particular case showing how 2D-ELDOR can be applied to study physical processes in complex systems that yield subtle changes.

INTRODUCTION

Over 16 000 missense mutations, which lead to single amino acid changes in protein sequences, have been identified and linked to human disease. Most of them occur in integral membrane proteins that have dynamic interactions with lipid molecules to form various microdomains on the surface of a cell. In recent years, the participation of plasma membrane lipids in regulating protein interactions and targeting early signaling events has become increasingly evident. The concept of “lipid rafts” emerged to explain a variety of experimental data that pointed to the possibility that the phase-like behavior of membrane lipids provides a means for selective compartmentalization in cellular processes. A simplistic view of this concept is that the fluid plasma membrane has distinctive regions characterized by disordered lipids with mostly unsaturated acyl chains (liquid-disordered, L_d-like) versus ordered lipids with mostly saturated acyl chains, and enriched in sphingomyelin, cholesterol (liquid-ordered, L_o-like) regions. Plasma membrane rafts correspond to the L_o-like regions that coexist with L_d-like regions, and available evidence supports the view that these domains are small (10–200 nm), heterogeneous, and highly dynamic. In a functionally relevant manner, small L_o-like domains are sometimes stabilized as larger compartments through stimulated protein–protein–protein–lipid interactions.

There is abundant evidence that domains exist/coexist in membranes and that small changes in lipid/protein composition can have dramatic effects on several signal transduction pathways.

A prominent example of a cell surface receptor that interacts differentially with L_o-like domains to initiate a transmembrane signal is FcεRI, the high-affinity receptor for IgE on mast cells, which plays a central role in the allergic immune response. Antigen cross-linking of IgE-FcεRI causes its coupling with Lyn kinase and resulting phosphorylation within a stabilized L_o-like membrane environment to initiate assembly of the signaling complexes. Although considerable biochemical data support this view and these are further corroborated with fluorescence measurements, knowledge of molecular dynamic properties of the domains before and after cell stimulation remains inadequate. Overall, the basic concept of coexisting domains in plasma membranes remains murky. Moreover, the nature and composition of these membrane structures very likely change with initiating and progressing events that accompany cell activation. Thus, although there is broad recognition of the importance of L_o-like membrane compartments in cell function, progress is currently limited by lack of adequate tools for their dynamic characterization.

Physical methods that have been found useful in investigating domains in membranes include fluorescence techniques (e.g., fluorescence resonance energy transfer (FRET), fluorescence anisotropy, and fluorescence microscopy) and

Received: February 18, 2011
Revised: July 22, 2011
Published: July 22, 2011
electron spin resonance (ESR). FRET and fluorescence anisotropy provide information about membrane heterogeneity and lipid order but much less about the molecular dynamics of lipids than can be obtained ESR with spectral simulation. Confocal, multiphoton, and super-resolution microscopy and single-particle tracking techniques have provided compelling, but still limited, visualization of heterogeneity of cellular membranes.

ESR offers a unique view of lipid dynamics in a complex membrane environment, and considerable effort has been devoted toward a comprehensive evaluation of membrane—phase coexistence. In model membranes, continuous-wave (cw) ESR can be used to map out the dynamic structure of the dipalmitoyl-sn-glycero-phosphatidylcholine/dialauryl-sn-glycero-phosphatidylcholine/cholesterol (DPPC/DLPC/chol) ternary phase diagram, wherein Ld, Lo, and gel phases exist or coexist. Also, a thermodynamic tie-line in the two-phase coexistence region of the ternary lipid mixtures was determined using cw ESR, and this approach has been extended to provide the complete tie-line field. Studies of cellular membranes have also been carried out. Ge et al. demonstrated the coexistence of Lo-like and Ld-like components in plasma membrane vesicles (PMVs) isolated from rat basophilic leukemia (RBL-2H3) mast cells using cw ESR with several different acyl chain spin-labeled phospholipids and a spin-labeled cholestane (CSL). More recently, Swamy et al. provided direct evidence for the

Figure 1. Experimental 2D-ELDOR spectra for 16PC in cross-linked and un-cross-linked PMV and their respective difference spectra at temperatures of 15, 20, and 30 °C, shown in the standard magnitude mode for convenience of display. (The real and imaginary parts of the full Sc − spectra are shown in the Supporting Information.) Here, Tm = 50 ns. (The spectra were first normalized to a common double integral value before taking their difference.)
The coexistence of two different types of lipid populations (i.e., Lo-like versus Ld-like) in the plasma membranes of live cells from four different cell lines by cw ESR.

The advanced ESR technique of 2D-ELDOR (two-dimensional electron–electron double resonance) has been used to study the phase structure of membranes and lipid–protein interactions in several studies.26–29 Earlier work showed that 2D-ELDOR provides unusually clear spectral distinctions between the Lo and Ld phases in model membranes,27 and more recently, it has been used to explore the phase properties of a model membrane and even deconvolute the spectra in terms of coexisting Lo and Ld phases using the “full Sc−” format30 of spectral analysis (Chiang, Costa-Filho, and Freed,28 hereafter referred to as CCF). In the present work, we extend this approach to plasma membranes, which are inherently more complex systems than model membranes. In particular, the CCF-based deconvolution of the 2D-ELDOR spectra in the two-phase region into the Lo and Ld components in equilibrium was greatly aided by the pre-knowledge of the pure Lo and Ld spectra at the phase boundaries, such that only their linear combination in the two-phase region needed to be determined. For PMV, no such prior knowledge is available. In cw ESR studies on PMV and live cells, two components were successfully resolved,16,25 but fitting with the many parameters that are needed is typically accompanied by substantial ambiguity. Thus, the challenge addressed in the present study was to show how 2D-ELDOR with full Sc− analysis could (1) clearly and unambiguously resolve the two spectral components in biomembranes and (2) provide accurate dynamics and ordering parameters to characterize these phases. Our driving interest in the present work was to show how 2D-ELDOR can be successfully utilized to study the phase structure of membranes and lipid/C0 di/sional electron interactions in several studies.26

The spectra in this study (including Figure 1), were obtained using the end-chain label 16PC, which was found to be the most favorable acyl chain label for high-quality 2D-ELDOR spectra.26–30,33,37 This is because it is more motionaly dynamic, resulting in much sharper spectral lines (especially compared to labels closer to the head group), which translates into much better signal strength, and it provides excellent distinguishability of the different phases. In addition, 16PC partitions nearly equally between the Lo and Ld phases, whereas this is not the case for lipids labeled closer to the head group, (e.g., SPC partitions with a 3:1 ratio favoring the Ld phase).

Three-pulse 2D-ELDOR experiments at 17 GHz were performed (cf. Figure 2). One collected the FID signals during the time interval t2 (after the spectrometer dead time, t0) for fixed values of the preparation time, t1, and the mixing time, Tm (cf. Figure 2A). The FID signal was Fourier-transformed in t2 and then in t1 to construct a frequency domain spectrum as a function of frequencies f1 and f2 (cf. Figure 2). The FID signal was in a “hypercomplex” format, containing two ordinary complex signals that we call Sc+ and Sc−. We only used the Sc− signal26 because the Sc+ signal typically decays rapidly within the finite spectrometer dead time. Rigorous analyses of these two signals in different motional regimes are given elsewhere.34–37

Although we show the 2D-ELDOR spectra in Figures 1 and 3 in the magnitude mode for convenience in display, we employed the full Sc− method30 to analyze them. In this method, the real and imaginary parts of the Sc− spectrum were simultaneously fit, instead of fitting the absolute magnitude of the complex
Figure 3. 2D-ELDOR spectra from 16PC for (A) un-cross-linked and (B) cross-linked PMV at 30 °C as a function of mixing time, $T_{m}$. The experimental spectra are shown in the left column, the best nonlinear least-squares fits are in the middle column, and their difference is in the right column. Spectra are shown in the magnitude mode for convenience. The $T_{m}$ are noted in the figure.
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The spectra of the PMV from RBL-2H3 mast cells in the present study include those from the un-cross-linked (U) versus cross-linked (C) samples at three temperatures, 15, 20, and 30 °C. Figure 1 shows a comparison between the magnitude 2D-ELDOR spectra for the U and C cases at these temperatures for a common mixing time, \( T_m = 50 \text{ ns} \). The U and C spectra have been normalized to a common value of the double integral (over \( f_1 \) and \( f_2 \)). Their difference is also shown on the same scale. This comparison demonstrates small but significant changes in the spectrum as a result of cross-linking IgE-FcεRI with antigen on the PMV. Figure 3 shows magnitude spectra at 30 °C as a function of \( T_m \) for both U (Figure 3A) and C (Figure 3B) cases. The spectra for all of the \( T_m \) fit simultaneously, so that the spectral changes, which provide additional information, are included. Figure 3A and B also shows the best-fit spectra, as well as the residual between experiment and fit. The fits are clearly very good. Also, the residuals are much smaller and of different character than the differences displayed between U and C spectra in Figure 1. The quality of fits that we obtain provides substantial justification for interpreting the observed differences due to cross-linking in Figure 1 as significant. As stated above, the detailed fitting was performed using the full Sc− method, and extensive examples of this fitting are given in the Supporting Information.

RESULTS

The spectra of the PMV obtained in experiments is best with phase distortions, which had previously hindered their use in fitting to theory, whereas the magnitude mode (cf. Figure 1) is not; therefore, it had previously been used at the expense of poorer resolution. The problem of phase distortions due to imperfect pulses, finite spectrometer dead times, and other instrumental effects is solved in the full Sc− method by simultaneously extracting the dynamic parameters and the phase corrections from the spectral analysis. The full Sc− results and their comparison with the theoretical fits are rather complex to display. (In fact, it was shown that the rigorous analysis requires that the theoretical spectra be phase-adjusted to correspond to the experiments.) For this reason, the complex full Sc− results are shown in the Supporting Information. As we have previously found, the accuracy in fitting the dynamic parameters is significantly enhanced by employing the full Sc− approach. It is important to note that once the phase corrections are obtained, they may be used in an approximate (although nonrigorous) fashion to convert the experimental Sc− spectra into an absorption spectrum displayed in the spin−echo correlation spectroscopy (SECSY) format (cf. Figure 2b and below). It provides much higher resolution than spectra in the magnitude mode. True pure absorption spectra, of course, obtainable from the (best-fit) theoretical spectra.

A virtue of displaying the absorption spectra in the SECSY format is that it provides a clear separation of inhomogeneous versus homogeneous broadenings, as discussed by Chiang et al. Spectral simulations were performed using the modified nonlinear least-squares fitting program called NLSPMC-fullscm, which is modified from Budil et al. to include the full Sc− fitting. The definitions for the dynamic parameters and the homogeneous line widths are given in ref 28. The spectral simulations for membrane vesicles used in the NLSPMC-fullscm program were obtained with the microscopic order with macroscopic disorder (MOMD) model.

Figure 4. The population of the \( L_0 \) component, coexisting with the \( L_d \) in the un-cross-linked versus cross-linked samples with respect to temperature.

(Figures S1−S6). Figure 3 represents an easily visualized display in the magnitude mode.

Fluorescence techniques can measure structural changes among membrane lipids upon cross-linking membrane proteins. Our ESR spectroscopic results, such as in Figure 1, not only demonstrate lipid structural changes in the PMV upon cross-linking IgE-FcεRI, but they can provide a more detailed quantitative assessment. That is, the full Sc− analysis of the 2D-ELDOR data unequivocally demonstrates that the spectra are the superposition of two distinctly different components, which can be characterized as \( L_0 \)-like and \( L_d \)-like regions in the PMV. The relative amounts of the two and how they change with cross-linking is shown in Figure 4, and the respective dynamic and ordering parameters for these components are given in Figure 5. The primary parameters, which are shown in Figure 5, are the rotational diffusion constant, \( R_I \) and the order parameter, \( S_0 \). These values for U-PMV are comparable to those previously obtained by cw ESR. The \( L_d \)-like phase is characterized by a small \( S_0 \) for the 16PC label and the \( L_0 \)-like phase by a larger ordering (e.g., 0.10 versus 0.27 at 30 °C). It is clear from Figure 4 that the \( L_0 \)-like phase is the predominant component for both U and C cases. The most important change upon cross-linking IgE-FcεRI on the PMV is the increase of the relative amount of the \( L_d \) component. Overall, the ordering and dynamic properties of both the \( L_0 \) and \( L_d \) components do not change very substantially upon cross-linking (cf. Figure 5); \( S_0 \) increases slightly for both components, but the value of \( R_I \) does decrease substantially for the \( L_d \) component and less so for the \( L_0 \) component.

The fitting and spectral analysis provide the spectral phase corrections, and it is then possible to obtain (approximate) absorption spectra in the SECSY mode. In Figure 6, examples are shown for the U and C cases at the three temperatures for \( T_m = 50 \text{ ns} \). The 1D-ESR spectrum is along the \( f_2 \) axis, and the homogeneous line width is along the \( f_1 \) axis. Their respective differences are shown as, in Figure 1, and these again reveal that significant spectral changes occur upon antigen cross-linking of IgE-FcεRI in the PMV. From the best-fit theoretical spectra, it is also possible to obtain the component \( L_0 \) and \( L_d \) spectra. We show an example in the pure absorption SECSY mode in Figure 7 for the U case at 30 °C and \( T_m = 50 \text{ ns} \). It is clear that the \( L_0 \) and \( L_d \) spectra are distinctly different, as expected; the \( L_0 \) spectrum with its low ordering is relatively small...
much sharper with narrower line widths than the \( L_d \) spectrum with its greater ordering. These results underscore the capacity of 2D-ELDOR full Sc analysis to discriminate these two lipid-phase components and to assess quantitatively their respective properties.\(^{27,28}\)

**DISCUSSION**

A biologically significant observation in the present work is that, as a result of antigen cross-linking of the receptor IgE-FcεR1 in PMV derived from RBL-2H3 mast cells, the majority \( L_o \) component is reduced relative to the \( L_d \) component for the spin

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**Figure 5.** The rotational diffusion rate, \( R_o \), and the order parameter, \( S_o \), versus temperature for un-cross-linked and cross-linked PMV.

**Figure 6.** Experimental 2D-ELDOR spectra for 16PC in un-cross-linked (left column), cross-linked (center column), and their respective difference spectra (right column) after converting to the (approximate) absorption in the SECSY format. Results are shown for 15, 20, and 30 °C and \( T_m = 50 \text{ ns} \).
label 16PC. Our ESR analysis is the first demonstration of such behavior of the lipid domains in response to receptor activation in plasma membrane preparations. We previously showed for U-PMV that the Lo component is the primary component by utilizing cw ESR, but this technique was not sensitive enough to distinguish any changes upon receptor cross-linking.16 The use of the more powerful 2D-ELDOR technique with the improved method of data processing and analysis has enabled us to clearly distinguish the subtle changes occurring. The previous cw ESR work16 showed that when the lipids were extracted from the PMV, only the Lo component remained. These results suggested that the Ld component is associated with lipids in the vicinity of the proteins. This interpretation is supported by the more recent study on the plasma membranes of live cells.25 Substantially more Ld component was observed for the live cells than for the PMV (although the Lo component remained the major one). This is consistent with the fact that the live cell plasma membranes have additional cytoskeleton-linked proteins and thereby a higher concentration of proteins; consequently, their disordering effect on the lipid structure is enhanced. Thus, our new results imply that antigen cross-linking of IgE receptors on the plasma membrane leads to increased disordering of the lipid structure. We suggest that the stimulating event of receptor cross-linking may mediate changes in membrane lipid organization that contribute to transmembrane signal initiation.

A previous study that is consistent with the implications of our 2D-ELDOR results is that of the phospholipid composition of detergent-resistant membrane (DRM) vesicles isolated from RBL cells that had been antigen cross-linked (C) or not (U) using mass spectroscopy. Fridriksson et al.43 found a large increase (12–22%) in the abundance of polyunsaturated phospholipids for the DRM from C cells versus U cells. They proposed that this large increase in the abundance of polyunsaturated phospholipids in DRMs corresponds to substantial lipid remodeling due to cross-linking of IgE-FcεRI, possibly including recruitment of intracellular membrane material into the plasma membrane as a result of stimulation. Furthermore, the increase in polyunsaturated lipids should lead to greater lipid disordering, corresponding to a Lo-like phase. These observations regarding the increased isolation of polyunsaturated phospholipids with DRMs after IgE-FcεRI cross-linking and cell activation were recently extended in a systematic study carried out by Han et al.44

Our results on the PMV indicating increased lipid disordering due to receptor cross-linking may be related to the results of these mass spectrometry studies.

In our previous cw ESR studies, we have discussed the fact that the actual relative amounts of Lo-like and Ld-like domains in biological membranes depend also on the partition coefficient of the probe, Kp. We found a Kp of approximately 1 for 16PC in a model three-component membrane system at 22 °C.25 Thus, we estimate that the fractions of the spectral components are approximately those of the Lo-like and Ld-like domains in the PMV. The increase in the Ld component with temperature in the U PMV samples (cf. Figure 4) was not observed with the other acyl chain labels used in previous work.16 (In that work, the cw ESR spectra from 16PC did not permit a separation of the Lo and Ld components, but labels further up the acyl chain did.) It is possible that the Kp of 16PC increases with temperature over the range of 15–30 °C, which would imply improved solubility in the Lo-like phase as its ordering decreases with increasing temperature. Alternatively, there may be restructuring of the lipids with temperature in the U-PMVs such that fewer disordered lipids are required to support the individual IgE receptors, and this temperature effect is significantly reduced upon cross-linking, as observed for the C-PMVs.

In conclusion, we emphasize the power of 2D-ELDOR spectroscopy in association with the full Sc—method of analysis to uncover subtle details of lipid structure and dynamics in membranes derived from cells. Our examination of effects on membrane lipids of cross-linking IgE-FcεRI in PMV demonstrates how 2D-ELDOR can be applied to study physical processes in complex systems that yield subtle changes.

**ASSOCIATED CONTENT**

§ Supporting Information. Extensive examples of detailed fittings, performed using the full Sc—method (Figures S1–S6). This material is available free of charge via the Internet at http://pubs.acs.org.

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**ACKNOWLEDGMENT**

We wish to thank Arun Gidwani for preparing the PMV samples for these experiments and David Holowka for his encouragement, advice, and critical reading of the manuscript. Work was supported by grants from the National Institutes of Health/National Center for Research Resources (NIH/NCRR) Grant P41RR016292 and National Institute of Biomedical Imaging and Bioengineering (NIH/NIBIB) Grant 2R01EB003-150 to J.H.F. and NIH Grant AI018306 to B.B. Computations.
were implemented at the Cornell Center for Materials Research Computing Facility.

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