

Imaging the dynamics of individual electropores

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Edited by Michael L. Klein, Temple University, Philadelphia, PA, and approved March 25, 2016 (received for review September 1, 2015)

Electroporation is a widely used technique to permeabilize cell membranes. Despite its prevalence, our understanding of the mechanism of voltage-mediated pore formation is incomplete; methods capable of visualizing the time-dependent behavior of individual electropores would help improve our understanding of this process. Here, using optical single-channel recording, we track multiple isolated electropores in real time in planar droplet interface bilayers. We observe individual, mobile defects that fluctuate in size, exhibiting a range of dynamic behaviors. We observe fast (25 s^{-1}) and slow (2 s^{-1}) components in the gating of small electropores, with no apparent dependence on the applied potential. Furthermore, we find that electropores form preferentially in the liquid disordered phase. Our observations are in general supportive of the hydrophilic toroidal pore model of electroporation, but also reveal additional complexity in the interactions, dynamics, and energetics of electropores.

droplet interface bilayer | electroporation | toroidal pores | optical single-channel recording | lipid bilayers

Maintenance of an intact cell membrane is vital for cell viability; it provides the barrier that prevents cell lysis and controls permeability to the external environment. However, intentional transient permeabilization of the membrane is also exploited as a means to introduce genes or drugs into an organism, and targeted permanent disruption of plasma membranes is an effective means to eliminate specific cells (1–3).

Electroporation (or electroporemeabilization)—the physical breakdown of a bilayer membrane under an external electric field—is a long-standing, popular method used to control the integrity of a cell membrane. Since its discovery and the first investigations in the 1960s and 1970s (4, 5), electroporation has been used in a wide range of applications, including gene transfection (6), wound and water sterilization (7, 8), tumor ablation (9, 10), electrochemotherapy (11, 12), and transdermal drug delivery (13). Furthermore, links to defibrillation damage have been highlighted (14).

Toroidal Pore Model

The transient aqueous pore hypothesis provides the basis of our current understanding of electroporation (Fig. 1). In this model, the kinetics of pore formation is governed by the transition over an energy barrier E^* created by the intersection of potentials corresponding to two distinct pore configurations: (i) a hydrophobic pore, where the lipids are simply parted with respect to an intact membrane, and (ii) a hydrophilic, toroidal pore (15). In this work, when describing a toroidal pore, we refer to a conductive pore, with the lumen lined and stabilized by lipid head groups (Fig. 1A). At small radii, hydrophilic pores reside within a local energy minimum (16); at large radii, there exists a local maximum (with associated critical radius r_c), beyond which a pore may grow indefinitely. The application of a transmembrane potential modifies the free energy of the hydrophilic pore such that its free energy, along with this barrier, is reduced (Fig. 1B). At a critical potential V_c , this barrier to unbounded pore expansion is lost, and the defect grows until the bilayer is destroyed.

Previous Imaging of Electropores

Theory and simulation have provided a rich source of predictions regarding the properties of electropores. For example, molecular

dynamics simulations have predicted the evolution of hydrophilic pores takes place from an initial membrane-spanning water file (17–19). However, there is a conspicuous mismatch between this level of predictive power and the information yielded by experiment.

Studies visualizing the presence of electropores are extremely limited. Techniques such as rapid freezing of electroporated erythrocytes (20) have provided snapshots suggestive of pore formation, but these have been disputed (21). The transient nature and small size of electropores place significant limitations on the measurement of electropore dynamics, and more fundamentally on their direct detection. Electrical single-channel recording (SCR) can resolve these dynamics at high time resolution (22, 23); however, SCR is limited in that it can only detect the total current across the membrane, and is thus unable to resolve whether conductance events are due to single or multiple permeabilization events. Indeed, the most common characteristic for electroporation in SCR would be a “noisy” trace that precedes bilayer rupture (see Fig. S5A, Lower).

A better insight into electropore formation would be provided by methods capable of observing the dynamics of electropore formation in real time. Few dynamic experimental imaging studies of these defects exist. Pioneering work in giant unilamellar vesicles (GUVs) has led to methods for assessing line tension, studying pore closure and formation times, and the dynamics of solute flow between the internal and external volumes (24, 25). For example, GUVs have been used to show transport across a membrane can be achieved by an electroporation pulse only 10 ns in duration (26). However, many of these systems typically examine small numbers of large, microscopic pores, much larger than the nanoscopic pores observed in cell systems (20–120 nm) (20), predicted by molecular dynamics simulations (17, 19) and implied by cellular uptake assays (27).

Here, we seek to improve our experimental understanding of electroporation by exploiting optical single-channel recording (oSCR) (28, 29) to image individual voltage-induced defects in a lipid membrane by detecting a fluorescent signal proportional to the flux of Ca^{2+} flowing through a pore. Very recently, we detected electroporation events using fluorescence signals generated by K^+ ionic flux (30). Although this work showed electropores formed

Significance

Transient permeabilization of the cell membrane is needed to introduce genes and drugs into cells, and electroporation (the physical breakdown of a bilayer membrane under an external electric field) is the most common method to achieve this. By applying an electric field to an artificial lipid bilayer, we have been able to visualize the presence of individual electropores in the bilayer. We exploit droplet interface bilayers (formed from the contact between an aqueous droplet and a hydrogel surface immersed in a phospholipid/oil solution) to provide simultaneous single-channel electrical recording and fluorescence imaging of the bilayer.

Author contributions: J.T.S. and M.I.W. designed research; J.T.S. performed research; J.T.S. and M.I.W. analyzed data; and J.T.S. and M.I.W. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1517437113/-DCSupplemental.

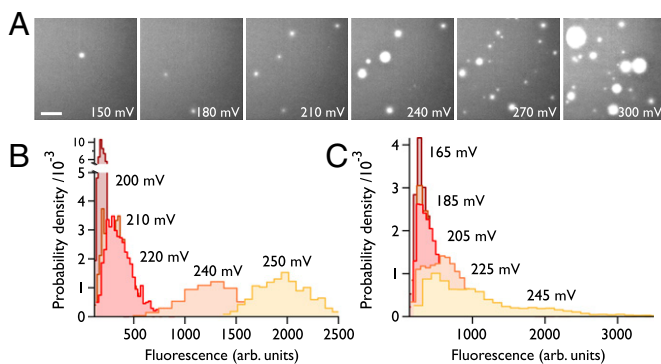


Fig. 3. Ensemble response. (A) Number and size of pores increases with voltage: maximum intensity images from >1,000-frame recordings on the same bilayer. (Scale bar: 25 μm .) (B) Histograms of oSCR intensities for a single pore within a bilayer experiencing an increasing applied potential. Pore amplitudes were obtained by 2D Gaussian fitting to the oSCR spot throughout the 1,000-frame movie. (C) Data from an ensemble of spots ($n = 2,424, 3,137, 5,641, 8,411,$ and $9,428$, respectively) as the potential is varied.

electrical recording (5, 22, 23); however, here, by isolating individual oSCR signals, we are able to extend observation of these phenomena to higher potentials, where the individual signals would be obscured in a purely electrical measurement. We have observed that all of these modes can occur at elevated potentials, with no apparent favor of one mode over another.

Visualization of pores during the application of potential across the bilayer allows us to confirm when there is a single pore present. This often occurs at low (80–110 mV) potentials, and we may determine the conductance of these defects and thus estimate our sensitivity. Our oSCR observations of isolated electropores indicate that conductances as small as 400 pS may be (optically) detected, a sensitivity over five times better than obtained using potassium-sensitive dyes (30). This current is similar in magnitude to the current measured directly at the onset of electroporation where presumably only a single electropore is present (Fig. S5A, Upper).

Approximating an electropore as a cylindrical defect (SI Experimental Methods), we obtain an approximate pore radius of 0.22 nm for the smallest pores (those detected at the onset of electroporation). This value will be an underestimate as the field experienced in the region of the pore will be reduced upon its formation (32, 33); however, it supports previous suggestions that the smallest electrically conductive pores are those able to accommodate at least a single file of ions. This value is also on the order of the smallest pores found in experimental and theoretical studies (19, 27, 34–36).

Electropore Diffusion. Individual, diffusing electropores can be tracked within the membrane (Fig. 4A). Mobile pores were generally small and exhibited a broad distribution of lateral diffusion coefficients (D_{lat}) (Fig. 4B) with a mean value of $0.67 \pm 0.58 \mu\text{m}^2\text{s}^{-1}$ (max: $2.7 \mu\text{m}^2\text{s}^{-1}$; min: $0.037 \mu\text{m}^2\text{s}^{-1}$). The lateral diffusion coefficient of these pores showed no obvious correlation with applied potential.

Phase Dependence of Electroporation. Molecular dynamics simulations predict that electroporation is favored in lipid regions of greater disorder (37). We electroporated phase-separated DIBs using two ternary mixtures, DPhPC/dipalmitoyl phosphoglycerol (DPPG)/cholesterol and DPhPC/brain sphingomyelin (bSM)/cholesterol (both molar ratio, 1:1:1), which exhibited liquid ordered (L_o) and liquid disordered (L_d) phase coexistence. Domains were visualized using 1 mol% of the lipophilic dye DiI, which partitions into the liquid disordered phase. Fig. 4C shows a median-averaged image from such an experiment, overlaid with trajectories of electropores diffusing in the membrane (see also Fig. S6 and Movies S2

and S3). Pores are seen to move within the L_d phase, moving between but not into L_o domains.

Pore Closure and Bilayer Rupture. Experiments carried out in cells have indicated that some pores can remain open for hundreds of seconds (27, 38, 39), whereas closure is typically in the nanosecond-to-microsecond range in simulations (19, 33). Within the time resolution of our experiments (16 ms), we observe pores that close immediately upon removal of the applied potential.

Uncertainty exists as to whether bilayer rupture is a result of a single expanding pore, or a collection of smaller defects (25, 40). Fig. 5A shows consecutive frames during DIB breakdown: we observe bilayer rupture to only take place via a single electropore. During the growth of this critical defect, other pores shrink and seal, likely as the sudden increase in conductance relaxes the potential across the membrane. Once started, the rupture process cannot be arrested. These observations support the transient aqueous pore model, where once a defect above the critical radius r_c is formed, there is rapid and uncontrolled expansion of the pore (15, 16). We note that the rupture patterns we observe (Fig. 5A, lower row) are similar to the floral instability patterns seen in rupturing multilamellar vesicles (41).

In phase-separated bilayers, the L_o boundary delimiting the disordered regions in which electropores can form (Fig. S6B) does not appear to restrict the expansion of the pore at elevated potentials; rupture occurs from one disordered region and proceeds to destroy the whole bilayer.

Interactions Between Electropores. A further question that has been raised is whether electropores are able to coalesce (25, 42). We only observe isolated pores and do not observe electropore coalescence in our experiments. Additionally, we have observed anticorrelation in pore currents (Fig. 5B), either when small pores are in close proximity (several tens of micrometers), or when a critical pore exists within the membrane (Fig. 5A). Such behavior would be consistent with modulation of either the local electric field or membrane surface tension by the presence of other pores.

Gating Kinetics. Large pores were observed to persist throughout the duration of our experiments; however, at the onset of electroporation, we observe discrete fluctuations in pore radius. This can be maintained for periods of up to several hundred seconds, and can be observed in both electrical and fluorescence recordings (uppermost traces, Fig. S5A and B). Given that this behavior arises at the onset of defect formation, we attribute this type of conductance to the opening and closing of the smallest possible pore. This

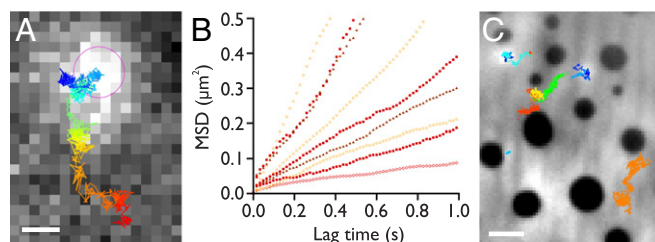


Fig. 4. Electropore diffusion. (A) A single frame from an oSCR stack (61.7 Hz) of a diffusing electropore, recorded at 200 mV, overlaid with the tracking trajectory (blue, 0 s, to red, 16.2 s). The pink circle indicates the detected location of the spot in this frame. (Scale bar: 2 μm .) (B) Representative mean-squared displacement versus time plots. Open circles: 180 mV; filled squares: 200 mV; open lozenges: 260 mV; filled triangles: 330 mV. Mean $D_{\text{lat}} = 0.67 \mu\text{m}^2\text{s}^{-1}$ ($n = 46$). We observe no obvious correlation of diffusivity with the applied potential. (C) Electroporation in a DPhPC/DPPG/cholesterol (1:1:1) phase-separated DIB. Median image of 500 frames recorded at 99.4 Hz. Electropores form in the (bright) L_d phase, diffusing around the (dark) L_o regions. Colored overlays show the trajectories of different tracked electropores. (Scale bar: 5 μm .)

the open time represents the depth of the energy well corresponding to a hydrophilic pore; the closed time corresponds to the height of the barrier for hydrophilic pore formation. The independence of open times on applied potential (Fig. 6C, *Upper*) implies that, in our experiments, there is no significant deepening of the hydrophilic pore well with applied potential. A rise in potential does broaden this local minimum, granting access to a greater range of pore radii (Fig. 3B and C), but we do not observe that pores at these higher potentials are more stable. The suggestion that the applied potential does not influence the kinetics of pore opening is consistent with previous work by Wilhelm et al. (40).

The closed time similarly does not vary with increasing potential, implying that the height of the energy barrier to hydrophilic pore formation is also relatively constant, i.e., the point where hydrophobic and hydrophilic potentials cross, E^* , is insensitive to applied potential. (This is further supported by our observation of small electropores at high potentials.) This is consistent with the proposed mechanism for the transition between a hydrophobic and a hydrophilic pore: the lipid rearrangement is predominantly driven by the minimization of the unfavorable interaction between water and the tail groups. Similarly, collapse of a conducting defect will always require the coming together of the water-lined toroidal pore walls, and the physical forces associated with this, along with those governing the transition of lipids back to a parallel, unperturbed bilayer arrangement, are unlikely to be greatly modulated by the potential. The idea that the crossing point between these potentials exists at a fixed location in terms of energy and radius has been hypothesized previously (36).

The reason for the open and closed times exhibiting two components is less clear. The histograms used to determine the pore lifetimes in Fig. 6 contain data from more than 1,900 electropores; however, analysis of individual pores shows that single pores also exhibit double-exponential behavior (Fig. S7). Each defect appears able to exist in either a short- or long-lived open or closed state. We tentatively attribute the longer timescales, $\tau_{2,open} \sim 400$ ms and $\tau_{2,closed} \sim 600$ ms, to the process of hydrophilic pore opening and closure, respectively. The observation that $\tau_{2,closed} > \tau_{2,open}$ is consistent with the barrier to conductive pore formation being larger than the barrier to pore collapse. The shorter times ($\tau_{1,open}, \tau_{1,closed} \sim 30\text{--}40$ ms) have a less obvious origin, and are not supported by the simple toroidal pore model. Bimodal rates for both opening and closing would be consistent with a number of different kinetic models, for example, a single closed state in equilibrium with two distinct open states. Molecular dynamics simulations have suggested that brief pore openings may be the result of conductive hydrophobic pores (43). However, this would require such a conductive hydrophobic pores to be long-lasting, with no favorable transition to the hydrophilic toroidal state. Alternatively, these lifetimes may be the result of brief (hydrophilic) openings as the potential only just overcomes E^* , in which case the pore opens, reduces the local electric field, and rapidly closes. Future work will help move beyond this speculation as to the nature of these kinetics.

Our observations point toward established defects influencing the nature of new ones: anticorrelation in pore fluctuations and the relaxation of the electrical stress on the membrane when a very large pore exists (Fig. 5 and Movies S4 and S5) are evidence of this. These factors are very likely the cause of the range of fluctuation regimes we observe (Fig. S5), and may contribute to the maintenance of small pores at higher potentials (Fig. 3C). The way in which biological membranes dissipate or augment these local tensile or electrical stresses during electroporation is therefore of great interest if we are to further understand this phenomenon in vivo.

Limitations. The events we detect do not provide a direct measurement of pore size: the fluorescent events corresponding to the “cloud” of ion flux flowing through each pore is larger than the pore itself. Although we can resolve the locations of individual pores to within a few tens of nanometers, we cannot image their

structure directly. Furthermore, optical detection of electropores is unavoidably constrained by the diffraction limit, and, although unlikely (42), if multiple pores exist within the diffraction-limited point spread function (FWHM = 0.59 μm), they will be essentially unresolved.

The time resolution of our current experiments (16 ms) also limits our ability to investigate electroporation kinetics. Such a restriction means that we are unable to observe electropore dynamics under the (typically) nanosecond-to-microsecond pulsed AC protocols that are typically used to electroporate cells. These rapid, transient applications of potential are designed primarily to reduce thermal damage that would otherwise kill the cell. However, in our experiments, with a small number of isolated pores over the bilayer area, we observe no measureable temperature changes associated with current flow.

Last, the presence of a supporting agarose substrate might potentially perturb electropore kinetics; however, our observations of Brownian diffusion of electropores and bimodal kinetics independent of diffusion (Fig. S7) imply that any effect of the substrate is minor.

Summary. Our results are in support of the hydrophilic toroidal pore model; however, the kinetics we resolve are not completely explained by this simple scheme. Under an electric field, electropores appear and disappear within the membrane in a stochastic manner, and exhibit constantly fluctuating radii. Pores become more numerous and fluctuate around greater radii as the applied potential is increased, as is consistent with our current understanding. Electroporation in this system appears not to be characterized by a single behavior, but by a range of fluctuation regimes. These observations point toward more complex interactions between electropores, indicating that local variations in the electric field or surface tension owing to pores already present limit the size of newly formed defects.

The ability to decompose the total current across the bilayer due to electroporation into the component contribution of individual electropores has enabled us to shed further light on the physical mechanism that controls this important phenomenon. The next steps must be to bridge the gap between the insights afforded by these in vitro models, and the realities of electroporation in vivo.

Experimental Methods

Materials. Stocks of 1,2 diphytanoyl-*sn*-glycero-3-phosphocholine (DPPC), 1,2-dipalmitoyl-*sn*-glycero-3-phosphoglycerol (DPPG), b5M, and cholesterol (all Avanti Polar Lipids) were stored in chloroform at -20 °C. The 8.7 mg·mL⁻¹ solutions of lipid in hexadecane were produced from these stocks before each experiment. All aqueous solutions were prepared using doubly deionized 18.2 M Ω ·cm MilliQ water. Low-melt agarose (Sigma-Aldrich) solutions were freshly prepared each day and kept at 90 °C to ensure homogeneity. Potassium chloride solutions were buffered with 10 mM Hepes, adjusted with potassium hydroxide, treated with Chelex resin (200–400 mesh; Bio-Rad) to remove divalent cations, and filtered using a 0.22- μm Steriflip disposable filter (Millipore). All experiments were carried out within purpose-machined poly(methyl methacrylate) devices with 16 distinct wells (44), enabling multiple bilayers to be imaged using a single device. Calcium-sensitive Fluo-8 fluorescent dye (AAT Bioquest) was prepared as a 1 mg·mL⁻¹ stock solution in distilled water and stored at -20 °C. All other chemicals were purchased from Sigma-Aldrich. Conductivity of solutions were measured at 21.1 °C using a calibrated (12.88 mS·cm⁻¹) Mettler Toledo FG3 m with LE703 probe. The conductivities were determined to be as follows: 1.5 M KCl, 10 mM Hepes: 144 mS·cm⁻¹; 750 mM CaCl₂, 10 mM Hepes: 99.5 mS·cm⁻¹.

Device Preparation and Experimental Setup. DIBs were prepared as described previously (44). Briefly, plasma-cleaned coverslips were spin-coated with 0.75% (wt/vol) aqueous agarose, and then affixed to the device. This was subsequently filled with 2% (wt/vol) hydrating agarose solution containing buffered KCl or CaCl₂. This hydrates (but does not cover) the substrate agarose by surrounding each well. Lipid-in-oil (8.7 mg·mL⁻¹ in hexadecane) was applied to the wells. After an incubation period to allow a monolayer to form ($\sim 15\text{--}30$ min), aqueous droplets in the same lipid-in-oil solution were added to the wells, forming a bilayer with the substrate under gravity. Prepared devices were placed within a Faraday cage on an inverted microscope (Eclipse TiE; Nikon).

Ag/AgCl electrodes were inserted into the hydrating agarose and the droplet, and connected to an Axopatch 200B patch-clamp amplifier (Molecular Devices) in voltage-clamp mode. DIBs were observed using a 60× TIRF oil objective (Nikon). Fluo-8 was excited by fiber-launched 473-nm CW laser radiation (~4 mW at fiber output) and imaged on an electron-multiplying CCD (iXon3 897; Andor). All experiments were conducted at room temperature. (Further detail on the experimental setup may be found in *SI Experimental Methods*.)

Phase-Separated DIBs. Bilayers were prepared as described, using mixtures of either DPhPC/DPPG/cholesterol or DPhPC/bSM/cholesterol (1:1:1 molar; total lipid concentration, 8.7 mg·mL⁻¹). The lipid-in-oil mixture also contained 1 mol% of the lipophilic dye Dil. After the droplets were added to the device wells, the device was incubated at 45 °C for 20 min to ensure lipid mixing.

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