Fluid surface coatings for solid-state nanopores: comparison of phospholipid bilayers and archaea-inspired lipid monolayers

Olivia M Eggenberger1, Geoffray Leriche2, Takaoki Koyanagi2, Cuifeng Ying1, Jared Houghtaling1,3, Thomas B H Schroeder1,4, Jerry Yang2, Jiali Li5, Adam Hall6 and Michael Mayer1,3

1 Adolphe Merkle Institute, University of Fribourg, Fribourg, Switzerland
2 Department of Chemistry and Biochemistry, University of California, San Diego, La Jolla, CA, United States of America
3 Department of Biomedical Engineering, University of Michigan, Ann Arbor, MI, United States of America
4 Department of Chemical Engineering, University of Michigan, Ann Arbor, MI, United States of America
5 Department of Physics, University of Arkansas, Fayetteville, AR, United States of America
6 Virginia Tech-Wake Forest University School of Biomedical Engineering and Sciences, Wake Forest School of Medicine, Winston-Salem, NC, United States of America

E-mail: michael.mayer@unifr.ch

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Abstract
In the context of sensing and characterizing single proteins with synthetic nanopores, lipid bilayer coatings provide at least four benefits: first, they minimize unwanted protein adhesion to the pore walls by exposing a zwitterionic, fluid surface. Second, they can slow down protein translocation and rotation by the opportunity to tether proteins with a lipid anchor to the fluid bilayer coating. Third, they provide the possibility to impart analyte specificity by including lipid anchors with a specific receptor or ligand in the coating. Fourth, they offer a method for tuning nanopore diameters by choice of the length of the lipid’s acyl chains. The work presented here compares four properties of various lipid compositions with regard to their suitability as nanopore coatings for protein sensing experiments: (1) electrical noise during current recordings through solid-state nanopores before and after lipid coating, (2) long-term stability of the recorded current baseline and, by inference, of the coating, (3) viscosity of the coating as quantified by the lateral diffusion coefficient of lipids in the coating, and (4) the success rate of generating a suitable coating for quantitative nanopore-based resistive pulse recordings. We surveyed lipid coatings prepared from bolaamphiphilic, monolayer-forming lipids inspired by extremophile archaea and compared them to typical bilayer-forming phosphatidylcholine lipids containing various fractions of curvature-inducing lipids or cholesterol. We found that coatings from archaea-inspired lipids provide several advantages compared to conventional phospholipids; the stable, low noise baseline qualities and high viscosity make these membranes especially suitable for analysis that estimates physical protein parameters such as the net charge.
of proteins as they enable translocation events with sufficiently long duration to time-resolve dwell time distributions completely. The work presented here reveals that the ease or difficulty of coating a nanopore with lipid membranes did not depend significantly on the composition of the lipid mixture, but rather on the geometry and surface chemistry of the nanopore in the solid state substrate. In particular, annealing substrates containing the nanopore increased the success rate of generating stable lipid coatings.

Supplementary material for this article is available online

Keywords: nanopore, single protein, lipid coating, archaea-inspired lipids, stability, noise

(Some figures may appear in colour only in the online journal)

1. Introduction

1.1. Background

Nanopore-based resistive pulse recordings are emerging as a powerful single-molecule characterization approach with applications in nucleic acid sequencing [1–7], biomolecule detection [8–11], characterization of single proteins [12–17], and biomarker-based diagnostics [18]. For instance, we showed recently that solid-state nanopores whose walls are coated with a fluid lipid membrane make it possible to fingerprint proteins with regard to their volume, approximate shape, charge, dipole moment and rotational diffusion coefficient [15].

In nanopore-based single molecule characterization, a high-gain current amplifier applies a constant potential difference between two electrolyte compartments separated by a thin insulating membrane. The amplifier records the ionic current through a single, electrolyte-filled nanopore in the membrane that connects the two compartments. As non-conducting macromolecules transit the pore, they typically cause transient reductions in the ionic current that contain information about the volume, charge, shape, and dipole moment of the translocating species [15, 19–22].

The earliest and much of the recent work in nanopore sensing employed biological protein pores spanning a planar lipid bilayer [23, 24]. The dimensions of these pores are well defined and typically constant from one protein nanopore to the next, as long as the protein’s amino acid sequence is not altered; in addition, protein pores often display minimal non-specific interactions with most analytes of interest [3, 25]. The diameters of the narrowest constriction of wildtype protein pores are, however, limited to a range between 0.4 and 3.3 nm, precluding the translocation of midsize and large proteins as well as other large analytes [2]. Recently, nanopores fabricated in synthetic substrates have enabled the analysis of larger particles [11, 26, 27], but surface interactions between these solid-state nanopores and biomolecules can result in undesired non-specific adsorption and clogging [16, 28, 29].

To reduce non-specific adsorption to the substrates of solid-state nanopores, various groups explored surface treatments such as silanization [30–32], self-assembled monolayers of thiols on gold films [33–35], atomic layer deposition of metal oxides [36, 37], and surfactant coatings [38, 39]. Some of these coatings also functionalized the pore surface [14, 40, 41] to improve detection efficiency [13, 42, 43]. In 2011, our group introduced a fluid lipid membrane coating [16], which eliminated or minimized non-specific interactions with the pore walls and enabled conjugation of proteins to mobile lipid anchors in the membrane [16]. This lipid anchor strategy significantly slows the translocation speeds and rotational dynamics of tethered proteins while circumventing undesired non-specific interactions with nanopore walls [15]. Slow translocation speeds enable current recordings that resolve most translocation events fully in magnitude and time and yield a large amount of data from individual translocation events. Together, these benefits of long-lived translocation events bolster the information content and statistics from individual resistive pulses. The strategy of tethering proteins to lipid anchors therefore enables the characterization of individual proteins beyond the determination of their volume and translocation time [15]. In particular, by eliminating or minimizing non-specific interactions with the pore walls, lipid coatings make it possible to quantify and interpret translational and rotational dynamics of lipid-anchored proteins as they move through the nanopore [15]. This approach makes it possible to extract the protein’s dipole moment, and rotational diffusion coefficient in addition to its shape and volume from an individual translocation event and is a first step towards identifying and counting individual proteins in mixtures [15]. While alternative nanopore coatings can also minimize clogging of nanopores and have the advantage that they are more straightforward to apply to the pore walls and chemically more robust than lipid coatings, their disadvantage is that they have not yet been able to minimize non-specific interactions to pore walls sufficiently to allow for the quantification of single molecule protein characteristics that rely on translocation and rotation, such as protein charge or dipole [35, 44]. Notably, the ability to anchor a protein to the bilayer allows for the determination of charge based on the distribution of translocation times for that protein [15, 16].
Here, we sought an ideal lipid-based surface coating that coats synthetic nanopores in silicon-nitride substrates in a straightforward and reproducible manner, while maintaining a stable and low-noise electrical baseline and slowing down the translational and rotational speed of non-spherical, lipid-anchored proteins such that orientation-dependent modulations in current during a translocation event could be resolved and quantified in time. In order to evaluate the quality and integrity of these bilayer coatings, we took advantage of measurements of protein charge. To do so, we recorded translocation time distributions of three lipid-anchored proteins and fit these distributions to the equation for biased first-passage time [45] with protein charge as a fitting parameter. Close agreement of the protein charge determined from the fit with the theoretically expected charge indicated that proteins translocated ideally and as theoretically predicted by the first-passage time equation; if non-specific interactions within the pore would have slowed down the proteins, then the translocation time could not be fitted well by the biased first-passage time equation and the determined protein charge would be significantly lower than expected.

1.2. Coating parameters

Applying a fluid lipid coating to a solid-state nanopore can increase the low frequency noise in current recordings, likely due to undulations of the coating [46]. As the ability to identify translocation events in a current recording is limited by the signal-to-noise ratio [47], an ideal nanopore coating would add minimal additional noise to the system or would even reduce the noise [16]. A coated nanopore must also yield a baseline current that varies minimally over various time scales, as fluctuations can lead to artifacts in the analysis.

The lateral diffusion coefficient \( (D_L) \) of lipids in a membrane is an important parameter in resistive pulse experiments that employ lipid-tethered analytes [16]. Yusko et al and Plesa et al demonstrated that only a small fraction of untethered proteins translocating through nanopores take sufficiently long to be time-resolved due to limitations of the recording bandwidth and the signal-to-noise ratio of the equipment [16, 48]. Proteins with lipid tethers that are anchored in a viscous lipid membrane coating, however, have more than 100-fold prolonged dwell times \( (\tau_d) \) in the pore compared to untethered proteins because the lateral diffusion coefficient of a tethered protein is dominated by the diffusion of its lipid anchor [16]. Slow protein movement is favorable since it increases the fraction of translocation events with dwell times longer than the time-resolution threshold of the recordings [16].

The work presented here, together with earlier work [15, 16, 49, 50], shows that nanopore fabrication methods and post fabrication processing steps affect the ease of coating a pore and the stability of the coating during nanopore experiments. In this paper, we coated pores that were made by three different fabrication techniques: ion-beam sculpting [7, 51], ion-beam drilling [52–54], and dielectric breakdown [50, 55]. A final goal of the work presented here was to identify the attributes of lipid coatings and pore fabrication strategies that facilitate reproducible formation of membrane coatings.

1.3. Experimental design: lipids tested

Figure 1 shows the chemical structure of the different classes of lipids tested in this study. In an attempt to decrease noise levels and increase the overall stability of the coating, we added various proportions of cholesterol into bilayers of 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC). Cholesterol increases stability in lipid membranes [56] because of its effect on membrane structure and packing [57–59]. Additionally, membranes with increasing proportions of cholesterol are more viscous, with correspondingly reduced \( D_L \) values [16, 60]. Here, we hypothesized that increasing the cholesterol concentration in the coating would prolong protein dwell times and lead to coatings with good stability.

We also tested fluid lipid coatings inspired by extremophile organisms of the kingdom archaea, which often have cell membranes composed of bolaamphiphilic lipids [61, 62]. These lipid structures have two terminal polar head groups connected covalently by a hydrophobic alkyl chain and form lipid membranes composed of fluid monolayers instead of bilayers. These lipids are thought to stabilize cell membranes, allowing extremophiles to survive in environments with extreme temperature, pH and salinity [61, 62]. To test if coatings from these lipids would be beneficial, we coated nanopores with a class of chemically synthesized archaea-inspired bolaamphiphilic lipids containing two glycerophosphocholine heads, which were attached to glycerol by an ether-bond to one pendant phytanyl chain and to one end of a membrane-spanning hydrocarbon chain (figures 1(A)–(C)). We have recently examined the ion permeation kinetics of membranes composed of these lipids [63–67] and since we had a selection of these lipids available, we tested eight of them.

For comparison, we additionally tested coatings composed of two commercially available and widely used non-bolaform archaea-inspired lipids: 1,2-diphytanoyl-sn-glycero-3-phosphocholine (DiPhyPC) and 1,2-di-O-phytanyl-sn-glycero-3-phosphocholine (Di-O-PhyPC).

Finally, we hypothesized that the discrepancy in ease of coating between different pores may be due to heterogeneities in the ‘sharpness’ of the radii of curvature of the corners at the brims of the pores. As we have reported previously, the success rate of forming a lipid coating on a nanopore chip can vary from 10% to 50% depending on the chip and we found that nanopores with diameters smaller than 17 nm [50] and pores with sharp edges of extreme curvature could, so far, not be coated with lipid membranes. Similar limitations of forming lipid membranes with extreme curvature have been reported for phospholipid vesicles [68] and for lipid coatings of nanoparticles [69–71]. We therefore tested if lipid membrane compositions that form more flexible coatings with regard to membrane curvature, when compared to those composed of cylindrically shaped lipids, would be able to conform more readily to sharp corners and thereby coat
nanopores with extremely small radii of curvature. In order to enhance the coating’s ability to conform to highly curved surfaces, we chose a ‘curvature-tolerant’ lipid composition containing lipids with a cone-shaped geometry (1-oleoyl-2-hydroxy-sn-glycerol-3-phosphocholine or LysoPC), with a complementary inverted cone shape (1,2-dioleoyl-sn-glycerol-3-phosphoethanolamine or DOPE), and with 1,2-dioleoyl-sn-glycerol-3-phosphocholine (DOPC), a cylindrically shaped lipid [72].

2. Results and discussion

2.1. Baseline quality: noise and stability

Figure 2 shows that coating a nanopore with a lipid membrane had only a relatively small effect on the noise in the most important frequency range, above 5 kHz for most lipid compositions, as reflected in the power spectra (distributions of noise power versus frequency, figures 2(B)–(D)). In contrast, lipid coating increased mostly the noise present in current recordings in the low frequency range (1/f noise) [47]. Figure 2(A) shows plots of the root mean square (rms) current noise after coating a nanopore chip divided by the rms noise before coating a chip with lipid membranes of various compositions. Bilayer coatings of the phytanyl-bearing lipids DiPhyPC and Di-O-PhyPC produced the largest increase in noise, likely due to unruptured liposomes near the pore entrance, shown in figure 3. Curvature-tolerant lipid bilayers also significantly increased the noise in current recordings. These elevated noise levels putatively stem from the diversity of lipid shapes present in the bilayer. Given that membranes containing conical lipids are prone to forming curved regions [73], membrane undulations along the pore walls are likely more pronounced when the membrane is a mixture of curvature-tolerant lipids compared to membranes from non-conical lipids.

Increasing the proportion of cholesterol in POPC membranes had no significant effect on the noise; in general lipid coatings of pure POPC with 10 to 40 mol% cholesterol showed the smallest increase in noise upon coating with a tendency towards the lowest noise values with 20 mol% cholesterol. These favorable noise properties (figures 2(A) and (C)) are likely due to the structural changes cholesterol imparts to lipid membranes [57, 74]: since the addition of cholesterol improves the packing of membranes, cholesterol-doped membranes may be more stable and fluctuate less than in the absence of cholesterol. Overall, we observed the lowest noise recordings with coatings from POPC-cholesterol mixtures and from archaea-inspired lipid membranes that contained 80 to 100 area% of archaea-inspired lipids. Comparing the various types of archaea-inspired lipids shown in figure 1(A), we observed no differences with respect to the noise and stability of nanopore coatings.

With most of the lipid compositions tested here, the current baseline measured from lipid-coated nanopores was stable for the average duration of a protein translocation experiment (around one hour). Experiment-ending instability
was rare, and when it did occur, it generally began after the addition of protein. Lipid coatings made from DiPhyPC and Di-O-PhyPC bilayers were more prone to unstable baseline currents than coatings of other lipid compositions. Epi-fluorescence microscopy revealed that during the coating procedure with these two lipids, many small unilamellar vesicles (SUVs) did not burst despite the osmotic pressure gradient. The resulting surface-attached residual SUVs likely contributed to baseline fluctuations due to the proximity of these liposomes to the nanopore entrance. Figure 3 displays a fluorescent image of a well-coated nanopore made with liposomes from POPC lipids, with the corresponding stable current trace in figure 3(B). In contrast, figure 3(C) shows a fluorescent image of liposomes from DiPhyPC lipids on the chip window with the corresponding unstable current trace in figure 3(D).

2.2. Lateral diffusion of lipids

Figure 4 shows $D_L$ values obtained via fluorescence recovery after photobleaching (FRAP) experiments that were carried out on the SiN$_x$ membrane of nanopore chips after coating with various lipid compositions. The inset shows that $D_L$ values of coatings from POPC lipids could be varied by a factor of $\sim 3$ by including 10 to 50 mol% of cholesterol in the lipid composition. Overall, the results in figure 4 show that $D_L$ values measured in nanopore coatings composed of membrane-spanning archaea-inspired lipids were approximately
10-fold smaller than for the \( D_f \) of bilayers of pure POPC or DiPhyPC lipids. DiPhyPC (and Di-O-PhyPC) bilayers formed coatings with poor baseline stability. (A) Epifluorescence image of a pore coated with POPC lipids. (B) Representative current trace for a pore coated with POPC lipids. (C) Epifluorescence image showing unruptured liposomes on a nanopore chip coated with DiPhyPC lipids. (D) Representative current trace for a pore coated with a DiPhyPC bilayer. All lipid compositions contained 0.8 mol% Rhodamine-PE lipids; unruptured liposomes appear as bright spots. The scale bars on both images represent 20 \( \mu \)m. The \( y \)-axis scale on the current traces represents 2 nA and the \( x \)-axis scale represents 2 s.

2.3. Ease of coating

During this investigation, we found that nanopores fabricated by ion beam sculpting produced stable coatings more frequently than nanopores fabricated by ion beam drilling or dielectric breakdown [50, 52]. None of the lipid compositions that we tested, including the curvature-tolerant lipid mixture, had a significant effect on the success rate or ease of coating. We found, however, that pores made by ion beam drilling or dielectric breakdown, which initially resisted coating, often could be coated more readily by all lipid compositions after ten or more cleaning cycles (see Materials and Methods). Since these cleaning cycles were frequently accompanied by an increase in the pore diameter as quantified from current measurements, we hypothesized that pores fabricated by these two methods initially contained rougher surfaces or sharper edges than those formed by ion beam sculpting and that these edges were smoothed over time by the slow surface etching induced by the cleaning solution or by the aqueous electrolyte during experiments [76].

In contrast, most of the nanopores that coated reliably well in this work were fabricated using an ion beam sculpting technique, in which He ions, which are directed towards a focused ion beam-drilled pore (~100 nm diameter), cause the pore to shrink to a desired diameter through mass flow of silicon nitride [7]. Once the sculpting process was complete, each ion beam sculpted pore was temperature treated during an annealing process in an inert atmosphere [76]. Most of the pores produced by ion beam drilling or dielectric breakdown that were difficult to coat initially did not undergo annealing during production [52, 77]. Rollings et al [76] and we [50] have shown that annealing smoothens the shape at the rim of silicon nitride nanopores and that this effect likely renders
nanopores amenable to coating with lipids. Rollings et al also demonstrated an increase in the stability towards slow etching in electrolyte solution of uncoated ion-beam sculpted pores after annealing, possibly due to a decrease in the number of dangling bonds left over from the fabrication process [76]. AFM images taken by Rollings and TEM images taken by us indeed indicated a smoother surface after annealing [50, 78]. Asghar et al showed that annealing also alleviates residual stress in silicon oxide nanopores [79]. We hypothesized that these effects together may have increased the success rates for generating lipid coatings of high quality. This hypothesis is supported by the observation that pores formed by controlled dielectric breakdown [55, 80, 81] initially were difficult to coat, while an annealing step doubled the success rate of coating and improved the noise characteristics in both uncoated and coated states [50]. Altogether, these results indicate that pore morphology is a critical parameter for the success of coating nanopore chips with lipid membranes.

2.4. Determining protein charge

One major advantage of coating nanopore walls with fluid lipid coatings is that this approach makes it possible to slow down the translocation of proteins by means of attaching them to a lipid anchor. The viscous drag of this anchor slows down the translocation of tethered proteins by two orders of magnitude, making it possible to resolve complete distributions of translocation times, $t_d$. These $t_d$ distributions reveal the net charge of the protein by fitting the distribution to the equation for a biased first-passage time process [16, 45]:

$$P(t_d) = \frac{t_p}{\sqrt{4\pi D_L t_d^3}} e^{-\frac{(t_d-v)^2}{4D_L t_d}},$$

where $v = |z|eV_pD_L/\nu k_BT$. The parameter $|z|$ (unitless) represents the net number of unit charges carried by the protein. $V_p$ (V) is the drop in electrical potential across the nanopore, $k_B$ (J K$^{-1}$) is the Boltzmann constant and $T$ (K) is the temperature of the system. This approach assumes that the lateral diffusion coefficient of the lipid anchor with the attached protein is the same as the average $D_L$ value of the lipids in the bilayer coating [16].

For charge determination in previous work [15, 16], we have typically fit the experimentally-determined probability distribution of $t_d$ values with equation (1). However, when we applied this approach to data from experiments with nanopore coatings from archaea-inspired lipids, the determined charge values were unrealistically high and the model could not fit the data (with adjusted $R^2$ values <0.5 in all cases). Close inspection of the recorded $t_d$ data revealed a large population of short dwell times (<40 μs) that dominated the distribution and were too short to originate from proteins that were lipid-anchored to a highly viscous coating from archaea-inspired lipids. We hypothesized that this first population of short translocation times primarily represented false events due to fluctuations in the baseline noise [75] and proteins that have lost their lipid tether during the course of the experiment.

Figure 4. Box plots displaying the lateral diffusion coefficients $D_L$ of all the major lipid compositions examined in this work as determined by FRAP on the SiN$_x$ windows of nanopore chips. The lower whiskers reach to the outermost data points that fall within the lower inner fence, defined as the 25th percentile − (1.5 × interquartile range), the upper whiskers reach to the outermost data points that fall within the upper inner fence, defined as the 75th percentile + (1.5 × interquartile range), the box encompasses the interquartile range, the line represents the median and the small square represents the mean. The box plot in blue represents the combined data from coatings with archaea-inspired lipids and mol% for compositions containing only POPC lipids, the red curve represents a composition containing archaea-inspired lipids and mol% for compositions containing only POPC lipids.
To account for these artifacts, we first integrated equation (1) to obtain the cumulative density function (CDF). This change allowed us to perform fitting on the data without the influence of binning. Next, we expanded the CDF to allow for the fitting of three separate populations within the cumulative distribution of measured \( t_d \) values. This expansion of the fitting equation made it possible to account for (1) the first population of \( t_d \) values shorter than 40 \( \mu \text{s} \) that we attributed to artifacts, (2) an intermediate population with the range of expected \( t_d \) values, and (3) a small, third population with dwell times longer than 1 ms, representing events of a few proteins that may have interacted non-specifically with the pore. We then determined the charge of three different proteins from the fit to the 2nd population in the middle of the distribution, resulting in mean charge values within \( \pm 30\% \) of reference values (see figure S1, available online at stacks.iop.org/NANO/30/325504/mmedia) [15]. Further details on the integration of equation (1) and the function for three-population fits can be found in the Supplementary Information.

### 3. Conclusions

Table 1 summarizes the conclusions from this investigation. By the quality criteria discussed here, we recommend either coatings from either pure POPC or with 10–20 mol% cholesterol or coatings from either pure archaea-inspired lipids or with 10–20 area% POPC for optimal nanopore-based characterization. Among these, the monolayer-forming lipids inspired by archaea led to the longest translocation times of all the lipids tested while maintaining stable, low-noise baselines. The low event frequency as a consequence of the high viscosity of these monolayer coatings can be addressed by increasing the proportion of lipid anchors in the coating and the accumulation of short-lived, false translocation events from statistical baseline fluctuations over extended recording times [75] can be minimized by restricting the analysis only to events with dwell times >40 \( \mu \text{s} \) through multiple-population fitting. We further recommend annealing of nanopore chips (regardless of fabrication method) to improve their stability in aqueous electrolytes as well as their ability to support a lipid membrane coating.

### 4. Materials and methods

We obtained all non-bolaform phospholipids from Avanti Polar Lipids. We synthesized archaea-inspired lipids as described by Koyanagi et al [63–67]. We purchased L-lactate dehydrogenase (59747) from Sigma Aldrich. We purchased polyclonal anti-biotin IgG-Fab fragments (800-101-098) from Rockland.

Unless specified otherwise, we performed protein translocation experiments using silicon chips containing 275 nm thick, 30 \( \mu \text{m} \times 30 \mu \text{m} \) silicon nitride windows, each bearing one nanopore, sculpted by a focused ion beam, as described by Li et al [7]. The narrowest constriction of these ion beam sculpted pores had a length of 10–30 nm with diameters ranging from 25–45 nm and a pore shape that can be approximated by a cylinder [15]. This nanopore opens up to a channel that is approximately 265 nm long. The channel is roughly cyclindical, expanding within a few nanometers from the pore to a diameter of about 100 nm. In addition to these chips, we used nanopores prepared by drilling with a He ion beam, as described by Yang et al [54]. These pores had a length of 20 nm with diameters ranging from 25–40 nm and a slight hourglass-like geometry [53]. As the third type of nanopore, we fabricated pores by dielectric breakdown using a modified procedure based on the one first described by Kwok et al [55]. Specifically, we enlarged the diameter of the nanopore slowly to 20 nm by applying a biopolar electric field with reduced amplitude that resulted in a cylindrically shaped pore [50].

We performed annealing as described by Rollings et al [76] by heating the chips in an inert nitrogen atmosphere over the course of 1 h from room temperature to 700 °C, holding the chips at this temperature for 30 min and letting them cool overnight to room temperature.

Before experiments, the nanopores were cleaned in freshly mixed, hot piranha solution (3:1 (v/v) concentrated sulfuric acid and 30% (v/v) aqueous hydrogen peroxide), rinsed copiously with deionized water and dried with \( \text{N}_2 \) gas.

During each experiment, we coated the freshly piranha cleaned nanopore substrates with a supported lipid membrane by depositing and fusing small unilamellar vesicles (SUVs) composed of the appropriate lipid composition onto the

### Table 1. Comparison of the lipid compositions tested with regard to their suitability for resistive pulse recordings based on four characteristics.

|lipid composition of coating| Stable baseline | Low noise | Slow translocation | Straightforward to coat |
|---------------------------|----------------|----------|------------------|-----------------------|
|100% POPC                 | +              | +        | +                | +                     |
|25, 50, 80, 90, 100% archaea lipids + 75, 50, 20, 10, 0% POPC | +              | +        | +                | +                     |
|100% DiPhyPC              | −              | −        | −                |  +                    |
|100% Di-O-PhyPC           | −              | −        | −                | +                     |
|60% DOPC + 20% DOPE + 20% LysoPC | +              | −        | −                | +                     |
|10, 20, 0, 40% Cholesterol +90, 80, 70, 60% POPC | +              | +        | +                | +                     |
|50% Cholesterol + 50% POPC | +              | +        | +                | −                     |

++ Indicates that a coating performed very well, + indicates that a coating performed well, − represents a negative influence as a result of this coating and −− is for a very negative influence. All percentage values are in area% for compositions containing archaea-inspired lipids and mol% for compositions containing bilayer-forming lipids.
Almost all lipid compositions included 0.8 mol% rhodamine NBD. Briefly, we formed lipid coatings by filling the upper compartment of the PDMS-nanopore setup with aqueous solution (150 mM KCl buffered at pH 7.4 with 10 mM HEPES) containing the SUVs (lipid concentration of 2 mM) and added the same buffer without SUVs to the bottom compartment. After an incubation period of 20 min, we rapidly imersed the entire setup in a large (∼500 ml) beaker of deionized water to favor bursting of residual liposomes by the resulting osmotic pressure gradient and to remove excess liposomes. We replaced the buffer with 2 M KCl containing either 10 mM HEPES buffer, pH 7.4 for experiments with Fab or 10 mM potassium citrate, pH 6.1 for experiments with L-lactate dehydrogenase.

In the case of membrane-spanning, monolayer-forming, archaea-inspired lipids, we quantified the proportion of lipids as area% instead of mol% because from the point of view of the lipid coating that is exposed towards the lumen of the pore, the relevant parameter is what proportion of the exposed area is composed of archaea-inspired lipids and what proportion is composed of bilayer-forming lipids. This approach assumes that the area per lipid of a membrane-spanning archaea-inspired lipid with a PC headgroup is the same as the area per lipid of a bilayer-forming lipid with a PC head group. Hence, for the membrane-spanning archaea-inspired lipids used here, a membrane with 80 area% archaea-inspired lipids and 20 area% POPC lipids corresponds to a membrane with 40 mol% archaea-inspired lipids and 60 mol% POPC lipids.

To confirm that each nanopore had successfully acquired a lipid coating, we used at least two methods: (1) comparing pre- and post-coating resistance values of the nanopore to the expected theoretical values [15, 16, 50] and (2) inspecting the fluorescence intensity by epifluorescence microscopy (as described in Yusko et al [16]). Because the addition of a lipid membrane decreases the nanopore diameter and increases the nanopore length, the current value observed under a constant voltage decreases by a predictable extent after coating and the fluorescence intensity typically increases in the area of the nanopore chip with the nanopore where a lipid membrane covers the top and bottom [16]. The most stringent quality criterion of lipid coatings, however, was provided by fits to translocation time distributions. Only high quality coatings make it possible to determine protein charge by fitting these data to the equation for a biased first-passage time process [15, 16].

We used a Nikon E600FN upright microscope equipped with an X-cite 120 Fluorescence Illumination System and a CoolSnap HQ camera (Photometrics) and a 60× water-dipping objective (NA = 1.00) to image the lipid coatings and perform FRAP [16].

We measured rms noise properties from the standard deviation of the measured electrical baseline current at an applied potential of ±100 mV after applying a digital Gaussian low pass filter with a cutoff frequency of 15 kHz to the recorded current trace.

We anchored proteins to the lipid membrane in two ways: for anti-biotin Fab fragments we employed 0.15 mol% of a lipid that presented a biotin-group as its headgroup in the membrane (1 area% for membranes composed of archaea-inspired lipids). We attached the other proteins with bis (succinimidyl) penta(ethylene glycol), an amine-reactive crosslinker (Thermo Scientific, 21581), by reacting this crosslinker with 1 mol% lipids with an ethanolamine headgroup within the membrane coating [15]. Specifically, we dissolved the crosslinker in a 2 M KCl buffer containing 100 mM KHCO3 at pH 8.4 and added the solution to the top of a lipid-coated nanopore chip at a final crosslinker concentration of 10 mg ml⁻¹. After 10 min incubation, we thoroughly rinsed away the crosslinker and added L-lactate dehydrogenase dissolved in the same carbonate buffer at the highest possible final protein concentration (between 1 and 3 μM). After at least 30 min, we rinsed away the excess protein and began recording with the specified recording buffer.

We performed protein sensing experiments as described by Yusko et al [15]. We measured current traces using Ag/AgCl pellet electrodes (Warner Instruments) with a patch-clamp amplifier (Axopatch 200B, Molecular Devices) at constant applied voltage of −100 mV. The buffer-filled compartment on the top was connected to the headstage of the amplifier and the bottom compartment was connected to ground. The analog low-pass filter of the amplifier was set to a cutoff frequency of 100 kHz. We acquired and stored data using a multifunction I/O device (PCI 6281, National Instruments) with a 500 kHz sampling frequency along with a custom LabView (National Instruments) program. All protein translocation analysis was performed using MATLAB (Mathworks). We first digitally filtered the data with a Gaussian 15 kHz low-pass filter and then applied a peak-finding algorithm developed by Pedone et al [82] to select regions of each current versus time recording where the measured current was greater than five times the standard deviation of the baseline current. The end of a resistive pulse was defined as the moment when the current returned to within one standard deviation of baseline current. We defined dwell times as the full-width at half-maximum value of the resistive pulse.

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