Cotranslational recruitment of ribosomes in protocells recreates a translocon-independent mechanism of proteorhodopsin biogenesis

Highlights

- Generated a simple artificial cell model for membrane protein insertion
- We identified protein-inherent control of translational targeting without chaperones
- Ribosomes, artificially tethered to GUVs increased membrane protein insertion
Cotranslational recruitment of ribosomes in protocells recreates a translocon-independent mechanism of proteorhodopsin biogenesis

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SUMMARY
The emergence of lipid membranes and embedded proteins was essential for the evolution of cells. Translocon complexes mediate cotranslational recruitment and membrane insertion of nascent proteins, but they already contain membrane-integral proteins. Therefore, a simpler mechanism must exist, enabling spontaneous membrane integration while preventing aggregation of unchaperoned protein in the aqueous phase. Here, we used giant unilamellar vesicles encapsulating minimal translation components to systematically interrogate the requirements for insertion of the model protein proteorhodopsin (PR) – a structurally ubiquitous membrane protein. We show that the N-terminal hydrophobic domain of PR is both necessary and sufficient for cotranslational recruitment of ribosomes to the membrane and subsequent membrane insertion of PR. Insertion of N-terminally truncated PR was restored by artificially attaching ribosomes to the membrane. Our findings offer a self-sufficient protein-inherent mechanism as a possible explanation for effective membrane protein biogenesis in a “pre-translocon” era, and they offer new opportunities for generating artificial cells.

INTRODUCTION
How cellular life first emerged on Earth remains one of the most fundamental questions in biology. The evolution of lipid-based membranes with embedded protein components is considered a crucial step in the emergence of primordial cells (Lane and Martin, 2012). Alpha-helical membrane proteins are ubiquitous and perform a startling number of essential tasks. Owing to their predominantly hydrophobic nature, these proteins readily integrate into the amphiphilic environment of a cell membrane (White and Wimley, 1999). A four-step thermodynamic model for spontaneous insertion of α-helical membrane proteins has been proposed and rigorously tested (Cymer et al., 2015; MacKenzie, 2006; White and Wimley, 1999; Wimley and White, 1996). Briefly, an unstructured hydrophobic peptide chain experiences a large free-energy-barrier-preventing membrane insertion (Wimley and White, 1996). The adoption of hydrogen-bonded α-helical structure lowers this free-energy-cost-making membrane partitioning favorable (Almeida et al., 2012; BenTal et al., 1997; Ladokhin and White, 1999). The bilayer interface region, a complex chemical environment dominated by lipid headgroups, facilitates the adoption of α-helical structure and thus aids in the spontaneous integration process (Ladokhin and White, 1999; Ulmschneider et al., 2011). However, in the aqueous environment, away from the bilayer interface region, aggregation of hydrophobic peptides predominates and inevitably leads to permanently aggregated states unable to adopt native structure and partition into the bilayer (Cymer et al., 2015).

Modern cells have evolved various chaperoning pathways to prevent aggregation in the cytoplasm and facilitate membrane insertion (Steinberg et al., 2018). The signal recognition particle (SRP) chaperoning pathway is the most common and facilitates the cotranslational targeting and insertion of the vast majority of α-helical membrane proteins. Via interactions with its membrane bound receptor, SRP positions translating ribosomes in close proximity to a membrane-embedded insertion apparatus such as the Sec translocon (Atopian et al., 2013; Rapoport, 2007). Given the complexity of this machinery, it is likely that membrane proteins were originally inserted into the membrane spontaneously and that mechanisms such as SRP targeting and the Sec translocon evolved later, becoming essential as compartmentalization and functional diversity of cells increased (Bohnsack and Schleiff, 2010; Pohlschroder et al., 2005).

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Given the propensity for detrimental aggregation in aqueous environments, the vast majority of \( \alpha \)-helical membrane proteins are inserted into the membrane cotranslationally (Cymer and von Heijne, 2013; Luirink et al., 2012). The efficient recruitment of translating ribosomes to the membrane becomes crucial as cell size and complexity increase (Soga et al., 2014). This raises the question whether \( \alpha \)-helical membrane proteins that emerged before the evolution of chaperoning/translocon systems have inherent features that facilitate membrane recruitment and insertion. Previous work has demonstrated that many membrane proteins can spontaneously integrate and fold into simple lipid membranes (Findlay et al., 2016; Harris et al., 2017; Pellowe and Booth, 2019; Soga et al., 2014). However, whether protein-inherent features of the emerging nascent chain affect recruitment of ribosomes to the membrane and thus tip the balance from aggregation toward integration remains to be investigated. Such a mechanism would represent an intriguing theory for how early cells could have overcome the lack of a dedicated chaperoning and translocon system. A bottom-up approach using artificial, cell-mimicking systems offers excellent opportunities to probe these hypotheses surrounding membrane protein biogenesis in the absence of a translocon (Xu et al., 2016).

To examine the possibility of protein-inherent membrane recruitment, we turned to the recombinant “cell-free” protein synthesis system known as the Protein synthesis Using Recombinant Elements (PURE). This system allows the investigation of protein folding in a minimal environment devoid of any chaperoning proteins or other insertion mediating factors (Kuruma and Ueda, 2015; Shimizu et al., 2001). When coupled with liposomes and lipid nanodiscs, this system has proven valuable for enhancing our understanding of the spontaneous membrane protein insertion process (Berhanu et al., 2019; Findlay et al., 2016; Harris et al., 2017; Matsubayashi et al., 2014). The PURE system can also be readily encapsulated within cell-size mimicking giant unilamellar vesicles (GUVs) and has been used in this way to highlight the importance of the vesicle surface area/volume ratio for membrane protein insertion/aggregation (Soga et al., 2014). To date, the focus of such studies has been on the insertion process itself, while the question of how these highly hydrophobic proteins avoid aggregation in a cell-mimicking context has not been thoroughly addressed.

Here, we have developed methods to interrogate the cotranslational recruitment of ribosomes to the membrane in a minimal context. We encapsulated PURE reactions within GUVs and used confocal imaging to investigate the minimal requirements for membrane localization and insertion of the model \( \alpha \)-helical membrane protein proteorhodopsin (PR) (Beja et al., 2000). We could show that the cotranslational recruitment of ribosomes to the membrane is the major determinant of the fate of proteorhodopsin in GUVs even in the absence of specific chaperoning and targeting pathways. We found that cotranslational ribosome recruitment is an inherent effect driven by the physical characteristics of the N-terminal hydrophobic domain of PR. The importance of this mechanism was further proven by artificially tethering the ribosomes to the membrane, which rescued the insertion of mislocalized PR lacking the N-terminal domain. The fundamental insights and techniques developed here can now be used to investigate bottom-up membrane protein assembly in a more biologically relevant context and will allow the expansion of the current tool set for the generation of simple artificial cell models.

RESULTS

The N-terminal domain of de novo synthesized proteorhodopsin enhances membrane insertion in cell-size-mimicking giant vesicles

As a minimal, cell-size-mimicking chassis, we used GUVs generated by the droplet transfer approach (Altmura et al., 2017; Noireaux and Libchaber, 2004), encapsulating PURE system components within the vesicle lumen. Plasmid DNA constructs containing the truncated or full-length forms of PR (PR\(_{\text{AN}}\) and PR, respectively) (Figures 1A, 1B, and S1) were then supplemented into inner solutions allowing the investigation of protein localization, insertion, and function using a range of confocal fluorescence microscopy assays. Enhanced green fluorescent protein (EGFP) fusion tags were first added to the C-terminus (Figure 1C) and the ratio of fluorescence obtained from the lumen and membrane of individual vesicles (compared with a soluble EGFP control Figure S2) enabled us to assess protein localization.

PR\(_{\text{AN}}\) lacking the N-terminal domain exhibited weaker membrane localization when compared with full-length PR, which showed a remarkably high association to the membrane (Figures 1D and 1E). We analyzed the effects of GUV size on membrane localization by performing a correlation analysis. A moderate positive correlation was observed for both PR and PR\(_{\text{AN}}\) suggesting that increasing vesicle size slightly reduced PR localization. Heterogeneity between individual GUVs was to be expected given that the efficiency of the
Figure 1. The N-terminal domain of proteorhodopsin (PR) is required for membrane localization and insertion
(A) Schematic indicating the two potential fates (aggregation and insertion) of membrane proteins synthesized in cell-size-mimicking biomimetic systems.
(B) Schematic representation of the topology PR with the N-terminal hydrophobic domain removed (PRΔN) and full-length PR.
(C) Experimental system for analysis of EGFP localization.
(D) Confocal microscopy images of GUVs after internal de novo protein synthesis of soluble EGFP (left), PRΔN-EGFP (center) and PR-EGFP (right). From top to bottom: brightfield image, fluorescence emission and 3-dimensional representations of fluorescence. Scale bars are 10 μm.
(E) Lumen/membrane fluorescence intensity ratio derived from radial profiles of EGFP fluorescence emission of individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are shown for 20 individual GUVs (filled circles). Different letters represent statistically significant differences (p < 0.001; one-way ANOVA using Tukey [HSD] post-hoc analysis).
(F) Experimental system for analysis of fluorescent HA antibody binding.
(G) Confocal images of GUVs after protein synthesis and incubation with Alexa-Fluor-647-conjugated HA antibody. Images are organized as in (D). Scale bars are 10 μm.
(H) Extravesicular/membrane fluorescence intensity ratio derived from radial profiles of Alexa-Fluor-647-conjugated HA antibody fluorescence of individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are based on 45 individual GUVs (filled circles). Asterisks represent statistical significance (P < 0.001) using a two-sample t test assuming equal variance. Data were normalized against a control ratio taken from GUVs expressing PR-EGFP with no HA epitope.
(I) Fluorescence intensity of DiBAC4(3) in the membrane of GUVs containing a non-functional PR mutant (D97N), PR without the N-terminal domain (PRΔN) and full-length PR. Individual vesicles were analyzed every 5 s under constant excitation with a 488-nm laser to excite both DiBAC4(3) and PR. 12 GUVs from 3 individually prepared experiments were analyzed for each protein construct with results showing the mean value from each experiment. Data are fitted with second order polynomials and 99% confidence intervals. All data were normalized to PRD97N data to account for photobleaching of DiBAC4(3).
See also Figures S1–S6.
encapsulation process is not saturated (Figure S3). Importantly, the observed membrane localization was indicative of protein insertion into the membrane as determined through the use of an antibody-binding assay (Figure 1F). Externally supplied fluorescently labeled hemagglutinin (HA) antibody showed preferential membrane localization when an HA epitope was introduced into the first periplasmic/extravesicular loop of PR (Figure S4), when compared with PR containing an HA epitope insertion in the third intracellular loop and to all PR D Nc constructs (Figures 1G, 1H, and S5).

A functional analysis was carried out to further correlate the observations made regarding localization and insertion. A fluorescent indicator of membrane potential, DiBAC4(3), was introduced into GUV membranes and dye emission was tracked upon the excitation of PR. The nonfunctional PR mutant D97N, which is incapable of proton transport (Pfleger et al., 2009), was used as a negative control. Full-length PR containing the N-terminal domain exhibited enhanced function in GUVs compared with PR D N. However, PR D N still showed low levels of functionality indicating that at least a small proportion of the protein was inserted and folded correctly in GUV membranes (Figures 1I and S6).

These combined results show that full-length PR is recruited and inserted into the membrane of GUVs in the absence of any soluble chaperones or translocon proteins, highlighting a crucial role for the N-terminal domain, which may form an eighth membrane-embedded α-helix.

The N-terminal hydrophobic domain of PR acts as a membrane anchor and facilitates downstream helix insertion
To further study the function of the N-terminal domain of PR, constructs were generated allowing PURE-mediated synthesis of the N-terminal domain in isolation (N), the first transmembrane helix (TM1), and the N-terminal domain together with the first transmembrane helix (N + TM1), all with a C-terminal EGFP fusion (Figure 2A). The N-terminal domain alone (N) was able to localize to the membrane despite the absence of downstream transmembrane helices, while TM1 exhibited very little membrane localization. The membrane localization of TM1 was recovered when the N-terminal domain was present (N + TM1) (Figures 2B and 2C).

The depth of peptide insertion into the hydrophobic core of the bilayer was then probed using the lipophilic environment-sensitive fluorescent probe nile red which reports on the fluidity of lipid tail groups (Mukherjee et al., 2007). Nile red undergoes a blue-shift in fluorescence emission in response to increasing membrane rigidity. We confirmed the sensitivity of nile red in GUVs with a range of POPC:cholesterol ratios (Figure S7) and then measured the red/green fluorescence emission ratio of nile red in the membranes of GUVs without (control) or with the PURE system enclosed. The N-terminal domain of PR alone (N) did not alter the lipid environment of the GUVs when synthesized in isolation, despite the high levels of membrane localization previously observed. By contrast, N + TM1 caused a significant decrease of red/green ratio indicating that it was able to constrain the lipid tails at the outer leaflet and increase membrane rigidity (Figures 2C and 2D). These data suggest that the N-terminal hydrophobic domain alone does not penetrate deeply into the core of the bilayer but can facilitate the insertion of TM1 which is otherwise not inserted into the membrane despite its high relative hydrophobicity.

Localization and insertion of de novo synthesized PR in GUVs requires cotranslational targeting
Our results indicate that the mechanisms involved are cotranslational and require ribosome recruitment to the membrane which in turn relies on protein-inherent features of the N-terminal hydrophobic domain. PR synthesized in the absence of a bilayer was unable to localize to posttranslationally supplied GUV membranes (Figure S8). Using a stalling peptide (SecM from E. coli), which causes translational arrest at a terminal proline residue, we generated constructs that result in the accumulation of stalled ribosome nascent chain complexes (RNCs). Figure 3 shows that stalled RNCs (fluorescently labeled) were only recruited to GUV membranes when the N-terminal hydrophobic domain was present. This behavior is consistent with previous results, both in vivo and in vitro, showing that most α-helical membrane proteins are inserted into the membrane cotranslationally (Cymer and von Heijne, 2013; Harris et al., 2017; Rapoport, 2007). We also confirmed that the mechanism is not linked to the membrane localization of mRNA (Jagannathan et al., 2014; Nevo-Dinur et al., 2011), regardless of the presence or absence of the coding sequence for the N-terminal hydrophobic domain (Figure S9). Consequently, the N-terminal domain could be used to guide
translating ribosomes to the membrane, thus increasing the efficiency of membrane insertion over aggregation in the lumen.

**The artificial tethering of ribosomes enables the localization and insertion of PR lacking the N-terminal domain**

In association with lipids, PR is known to adopt its native structure both in vivo and in vitro regardless of the presence of the N-terminal hydrophobic domain (Gourdon et al., 2008; Soto-Rodriguez and Baneyx, 2019). However, cotranslational ribosome membrane recruitment could become critical as vesicle size increases.
If this is the case, one would predict that artificial tethering of ribosomes to the membrane would restore insertion of a PR lacking the N-terminal domain. Using vesicles doped with the synthetic lipid 1,2-dioleoyl-sn-glycero-3-[(N-(5-amino-1-carboxypentyl) iminodiacetic acid) succinyl] (nickel salt) (DGS-NTA(Ni)) (Peters et al., 2015), we attached modified 70S ribosome complexes from the *E. coli* strain JE28 containing hexahistidine tags on the four L7/L12 proteins of the large ribosomal subunit (Figure 4A) (Ederth et al., 2009), visualized using propidium iodide fluorescence staining of rRNA (Figures 4B, 4C, and S10). Functionality of the purified His-tagged ribosomes was confirmed by replacing commercially supplied ribosomes in bulk PURE reactions, resulting in the successful synthesis of α-hemolysin (Figure S11).

Enhanced membrane localization of PRΔN-EGFP was observed when ribosomes were tethered to the membrane (Figures 4D and 4E). Recovery of membrane insertion of PRΔSP was also confirmed by the HA epitope assay used previously (Figures 4F and 4G). A control experiment was performed to ensure that the antibody did not bind directly to the NTA-containing lipid (Figure S5). These experiments showed that the requirement
Figure 4. Pretranslational membrane tethering of ribosomes rescues the localization and insertion of PR lacking the N-terminal domain

(A) Schematic representation of the His-tagged bacterial 70S ribosome complex (PDB entry 4V4P). Yellow and gray surfaces represent the 50S and 30S subunits, respectively; green and red surfaces represent the histidine-tagged L7/12 proteins and exit tunnel proteins, respectively.

(B) Confocal images of GUVs encapsulating 1 μM of 70S ribosomes stained with 100 μg/mL propidium iodide (PI). GUVs were generated with and without 2.7 mol% DGS-NTA(Ni) in the membrane. Scale bars are 10 μm. DGS-NTA(Ni) tethers His-tagged ribosomes to the membrane. A control experiment was performed to ensure that the antibody did not bind directly to the NTA containing lipid (Figure S6).

(C) Lumen/membrane ribosomal fluorescence intensity ratio derived from radial profiles of PI emission in individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are based on 20 individual GUVs (filled circles). Asterisks represent statistical significance (P < 0.001) using a two-sample t test assuming equal variance.

(D) Confocal images of GUVs after synthesis of PRΔN-EGFP by His-tagged ribosomes tethered to the membrane (2.7) or not (0). Scale bars are 10 μm.

(E) Lumen/membrane fluorescence intensity ratio derived from radial profiles of EGFP fluorescence emission of individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are shown for 20 individual GUVs (filled circles). Asterisks represent statistical significance (P < 0.001) using a two-sample t test assuming equal variance.

(F) Confocal images of GUVs after synthesis of PRΔN-EL1HA by His-tagged ribosomes tethered to the membrane (2.7) or not (0) and incubation with Alexa-Fluor-647-conjugated HA antibody. Scale bars are 10 μm.

(G) Extravesicular/membrane fluorescence intensity ratio derived from radial profiles of Alexa-Fluor-647-conjugated HA antibody fluorescence of individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are based on ≥45 individual GUVs (filled circles). Asterisks represent statistical significance (P < 0.001) using a two-sample t test assuming equal variance. Data were normalized against a control ratio taken from GUVs expressing PR-EGFP with no HA epitope.

See also Figures S10 and S11.
of the N-terminal domain of PR for membrane localization and insertion can be replaced by artificially attaching the ribosomes to the membrane. The data indicate that the proximity of ribosomes to the membrane is a crucial limiting factor for the spontaneous, cotranslational insertion of PR into the membrane of cell-size-mimicking vesicles, at least when no receptor-mediated targeting mechanisms are present.

**DISCUSSION**

The use of bottom-up approaches to investigate the membrane protein biogenesis process have led to remarkable discoveries which begin to address some of the fundamental questions concerning the evolution of cellular membranes. It has long been argued that the final adopted structure of an α-helical membrane protein represents a thermodynamically favorable state (Cymer et al., 2015; Popot and Engelman, 1990). It is therefore logical to assume that many membrane proteins already contain within their amino acid sequences the necessary tools to overcome energetic barriers arising during membrane insertion. Thus, they may not require the assistance of sophisticated chaperoning and insertion complexes as long as the cellular context is simple, for example, lacking functionally diverse cellular compartments. Recent studies have indeed provided some evidence that translocons are not an absolute requirement for the insertion and folding of many membrane proteins, at least in simplified biomimetic systems, and are only needed for the insertion of proteins with large extracellular soluble domains (Baars et al., 2008; Berhanu et al., 2019; Matsubayashi et al., 2014). We have shown here that proteorhodopsin (PR), a model α-helical membrane protein is able to spontaneously integrate into a simple lipid membrane of giant vesicles in the absence of a translocon or chaperoning proteins.

If a purely thermodynamically driven pretranslocon mechanism of insertion prevailed early in evolution (Pohlschroder et al., 2005), membrane recruitment of translating ribosomes was likely crucial to avoid misfolding of aggregation-prone hydrophobic proteins before they came into contact with a membrane. Some modern cells still contain clues that hint at the importance of such a mechanism. For example, yeast mitochondrial ribosomes are permanently attached to the inner mitochondrial membrane which lacks a Sec translocon (Glick and VonHeijne, 1996; Jia et al., 2003). This attachment facilitates the insertion of membrane proteins encoded by the mitochondrial DNA (Szyrach et al., 2003).

Given the structural ubiquity of rhodopsins throughout all domains of life and thus the likely ancient nature of such proteins, PR seems to be a good model to investigate this hypothesis. We have shown here that the N-terminal hydrophobic domain of PR is critical for the localization and insertion of the membrane-spanning hydrophobic α-helices into cell-size-mimicking GUVs. The function of PR in GUVs is also drastically reduced when the N-terminal domain is absent. It seems likely that this drop in function is owing to a reduction in protein insertion and subsequent increase in aggregation. The large volume/surface area ratio of GUVs has previously been shown to affect the spontaneous insertion and function of another α-helical membrane protein, EmrE, when synthesized de novo using the PURE system, but the protein-inherent mechanisms underpinning localization remained unexplored (Soga et al., 2014). It should be noted that a recent in vivo study identified the N-terminal domain of PR as important for efficient biogenesis even in cells containing chaperoning and insertion pathways (Soto-Rodriguez and Baneyx, 2019). This hints at the intriguing possibility that thermodynamically driven membrane affinity mechanisms aid membrane protein biogenesis in vivo to this day, at least for microbial rhodopsins, although this idea has yet to be explored.

Using a well-defined minimal system, we could show that the localization effect mediated by the N-terminal domain of PR does not require additional cellular factors such as SRP, FtsY, or other chaperones and efficiently recruits translating ribosomes to the membrane. It should be noted that the levels of translation were markedly lower for PRΔN. This was likely owing to protein misfolding and aggregation when the N-terminal domain was absent and translating ribosomes were not efficiently attached to the membrane. Because the N-terminal domain alone did not seem to penetrate deeply into the membrane, it is likely that it recruits the ribosomes through interaction with the lipid interface region. We cannot entirely rule out the possibility that the N-terminal domain fully penetrates the bilayer as the resolution of our nile red assay may be limiting.

Our experiments with TM1 suggest that this N-terminal interaction also guides the insertion of the downstream helices of PR, which would otherwise tend to aggregate in the aqueous interior of GUVs. Previous studies have implicated the bilayer interface as an important chemical environment that facilitates the adoption of α-helical structure and subsequent partitioning of hydrophobic peptides into the bilayer interior (Ullschneider et al., 2011, 2018; White et al., 2001). It is an intriguing possibility that the N-terminal hydrophobic regions of other membrane proteins beyond the rhodopsins also have a high affinity for
this region of the bilayer. Such affinity may even act in parallel to SRP-mediated targeting to enhance the efficiency of membrane protein biogenesis. Recent work has shown that SRP in both prokaryotes and eukaryotes recognizes hydrophobic helices along the entire length of the nascent chain and is not specific, as previously thought, for the N-terminal helix (Chartron et al., 2016; Costa et al., 2018; Schibich et al., 2016). Further research is clearly required to address these exciting questions.

Crucially, the N-terminal-dependent localization and insertion of PR can be mimicked by direct, pretranslational binding of ribosomes to the lipid membrane. Given that PR shows moderate functionality in the absence of its N-terminal domain, it seems reasonable to assume that this reduced function is due to reduced protein insertion and increased aggregation as seen in Figure 1. While the translation of PRΔN is markedly lower than full-length PR, there is likely a modest increase in the efficiency of translation following membrane tethering of ribosomes, possibly owing to a reduction in protein aggregation at the exit tunnel of the ribosome, although we cannot be sure of the precise reason for this observation. A recent study has shown that tethering an emerging nascent chain to the membrane increases insertion (Ando et al., 2018). However, these techniques are limited owing to the retention of the N-terminus on the extravascular surface. Our study builds on this work by attaching ribosomes to the membrane and by using a more realistic, cell-size-mimicking model (GUVs). As mentioned previously, membrane tethering of ribosomes is a well-known strategy used by mitochondria to support effective biogenesis of inner membrane α-helical proteins encoded by the mitochondrial genome (Bonnefoy et al., 2009; Szyrach et al., 2003). Our work presents proof that this strategy can be replicated for synthetic membrane protein assembly in a cell-size-mimicking vesicle system. The technique can now be used to generate simple artificial cell models requiring the insertion of aggregation-prone membrane proteins into vesicle membranes.

In conclusion, we have identified an inherent mechanism of the N-terminal domain of PR that is both necessary and sufficient for the cotranslational recruitment of ribosomes to the membrane of minimal cells. This recruitment then drives the subsequent insertion of downstream α-helices. Our findings using a single model protein hint at a possible explanation for effective membrane protein biogenesis during early evolution in a “pretranslocon” era, and they offer new opportunities for generating artificial cells from the bottom-up. It will be interesting in the future to investigate further examples of this phenomenon and whether such fundamental processes still play a role in membrane protein biogenesis that is masked by the existence of chaperoning systems in modern cells.

Limitations of the study

Our study used the model membrane PR to delimit requirements for membrane insertion in a simple protocell model. To broaden the impact of this research further testing should be carried out to determine whether and which of the reported observations also apply to other membrane proteins and how they are integrated with more complex integration machineries in living cells. Furthermore, in GUVs, we were not able to quantify the kinetics or the efficiency of the insertion process. Kinetics of membrane protein folding in a minimal context has been previously investigated (Harris et al., 2017) and was beyond the scope of our study. Determining insertion efficiencies within GUVs was not possible owing to the lack of robust methodologies for achieving a relevant result. It is likely that the simple protein-inherent process presented here has much lower insertion efficiency than translocon-mediated mechanisms.

Resource availability

Lead contact

Further information and requests should be directed to and will be fulfilled by the Lead Contact, Ross Eaglesfield (ross.eaglesfield@glasgow.ac.uk).

Materials availability

All plasmids used in this study are available upon reasonable requests from the lead contact. Requests relating to E. coli strain JE28 should be directed to S.S.

Data and code availability

All raw images and data generated and analyzed during this study are available from the lead contact upon request.
METHODS
All methods can be found in the accompanying transparent methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102429.

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AUTHOR CONTRIBUTIONS
R.E., J.R. and A.A. designed the study. R.E. performed experimental work and data analysis. M.A.M. supported experimental work. S.S. provided the JE28 strain and relevant expertise. R.E. and A.A. wrote the manuscript with input from all authors. All authors were involved in manuscript editing and revision.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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Supplemental information

Cotranslational recruitment of ribosomes in protocells recreates a translocon-independent mechanism of proteorhodopsin biogenesis

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Figure S1. SDS-PAGE analysis of batch PURE reactions. Related to Figure 1. Proteins EGFP, PR-EGFP and PRΔN-EGFP were synthesized in bulk in the presence of large unilamellar vesicles (LUVs) and BODIPY-labelled lysine amino acids. Following protein synthesis reactions were run on an SDS-PAGE gel and imaged using the fluorescence of incorporated BODIPY-labelled lysine and coomassie staining. Translation products are marked with white arrows.
Figure S2. Examples of radial profile fluorescence analysis. Related to Figure 1. (A) Confocal microscopy image of a GUV with membrane localized fluorescence. The red circle indicates the region of interest used for radial profiling. Scale bar is 10 µm. The corresponding line plot shows the radial fluorescence profile for the image shown. Membrane fluorescence was determined by calculating an average of three pixels with the highest signal being the middle pixel. Lumen fluorescence was determined by taking an average of the subsequent 15 pixels moving into the vesicle lumen. Values were normalized to background fluorescence measured as an average of three arbitrary points on the image without GUVs present. (B) Confocal microscopy image of a GUV with lumen localized fluorescence. The red circle indicates the region of interest used for radial profiling. Scale bar is 10 µm. The corresponding line plot shows the radial fluorescence profile for the image shown. When no membrane fluorescence peak was observed the average of three pixels was taken at the point where fluorescence stopped increasing initially. Lumenal fluorescence was determined as before by averaging the fluorescence intensity of 15 pixels further into the vesicle lumen. Background adjustments were carried out as before. All images were captured at a resolution of 2048 x 2048 in order to maintain a consistent pixel size.
Figure S3. Correlation analysis of membrane localized EGFP and GUV size. Related to Figure 1. Fluorescence ratios from Fig. 1E were plotted against GUV radius and a pearson correlation analysis was performed. The resulting model is shown with a 95% confidence interval. (A) EGFP (B) PR-EGFP (C) PRΔN-EGFP.
Figure S4. Extra- and intracellular loop modification of PR with hemagglutinin epitope. Related to Figure 1. The topological structure of PR is shown with the N-terminal hydrophobic domain shown on the left followed by the 7 transmembrane helices. HA epitope insertion by Gibson assembly yielded the extracellular (green) modification in PR-EL1HA and the intracellular (purple) modification in PR-IL3HA. Negatively charged residues are blue and positively charged residues are red.
Figure S5. Data for Alexa Fluor-647 conjugated HA antibody-binding assays. Related to Figure 1. (A) Confocal images of GUVs for each protein construct and condition. Images shown are brightfield, fluorescence emission > 650nm and 3-dimensional representations of fluorescence. Scale bars are 10 µm. DGS-NTA(Ni) only sample does not contain a PURE reaction but was used as a control to show that antibody could not bind directly to the NTA containing lipid. (B) Extravesicular/membrane fluorescence intensity ratio derived from radial profiles of Alexa Fluor-647 conjugated HA antibody fluorescence of individual GUVs (n = 16 – 48). Mean values (solid lines) and standard deviations (dashed lines) are based on individual GUVs (filled circles). Different letters represent statistically significant differences (p < 0.001; one-way ANOVA using Tukey (HSD) post-hoc analysis). The lowest ratio detected in control vesicles lacking an HA epitope is marked in red and was used to normalize all subsequent data.
Figure S6. Functionality of PR synthesized inside GUVs. Related to Figure 1. Fluorescence intensity of DiBAC₄(3) in the membrane of GUVs containing a non-functional PR mutant (D97N), PR without the N-terminal domain (PRΔN) and full-length PR. Individual vesicles were analyzed every 5 seconds under constant excitation with a 488nm laser to excite both DiBAC₄(3) and PR. 12 GUVs from 3 individually prepared experiments were analyzed for each protein construct with results shown as the mean value for each experiment. Panel 1 shows all data points with fitted 2nd order polynomials and 99% confidence intervals. Red represents PR D97N, green PRΔN and blue PR. The remaining panels show individual data points for all protein constructs with individual protein constructs highlighted according to color as in panel 1. The top row shows raw data with the second row showing data normalised to the displayed polynomial function for PR D97N to account for photobleaching of DiBAC₄(3). Residuals for each fitted polynomial function are shown under the respective panel.
Figure S7. Nile red calibration in POPC:cholesterol GUVs. Related to Figure 2. (A) Schematic showing the nile red fluorescence emission response to increased cholesterol and thus lipid order. Loosely packed lipids without cholesterol lead to probe emission being red-shifted. Increasing concentrations of cholesterol leads to a blue-shift in the emission spectrum due to increased lipid tail order. (B) Quantification of nile red fluorescence ratio (red/green) for GUVs containing increasing concentrations of cholesterol. Mean values (solid lines) and standard deviations (dashed lines) are shown for ≥11 individual GUVs (filled circles). Different letters represent statistically significant differences (p < 0.001; one-way ANOVA using Tukey (HSD) post-hoc analysis).
Figure S8. Membrane recruitment and insertion of PR does not occur post-translationally. Related to Figure 3. (A) Schematic representation of the experimental design used to investigate post-translational protein localization. Bulk reactions were performed without lipids. Following completion of reactions solutions were used as the inner solution to generate GUVs by droplet transfer. Following a short incubation, GUVs were visualized using confocal microscopy. (B) Confocal image showing GUVs encapsulating PR-EGFP and PRΔSP synthesized prior to vesicle formation without an amphiphile present. Scale bars are 10 µm. (C) Lumen/membrane fluorescence intensity ratio derived from radial profiles of EGFP fluorescence emission of individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are shown for 20 individual GUVs (filled circles). n.s. represents no statistical significance (P = 0.182) using a two-sample t-test assuming equal variance.
Figure S9. Spinach2-tagged mRNA is localized diffusely throughout the lumen of GUVs following transcription. Related to Figure 3. (A) Sequence and secondary structure of the RNA aptamer spinach2. (B) Chemical structure of the conditionally fluorescent dye 3,5-difluoro-4-hydroxybenzylidene imidazolinone (DHFBI). (C) Spinach2 aptamer attached to the 3’ end of mRNA transcripts. (D) Fluorescence time-series showing fluorescence of PR-spinach2 (open circles), PRΔN-spinach2 (open circles) and a negative control containing no template DNA (filled circles). (E) Confocal image showing GUVs after mRNA synthesis and treatment with DHFBI. Scale bars are 10 µm. and (F) Lumen/membrane fluorescence intensity ratio derived from radial profiles of DHFBI fluorescence emission of individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are shown for 20 individual GUVs (filled circles). n.s. represents no statistical significance (P = 0.756) using a two-sample t-test assuming equal variance.
Figure S10. Synthetic attachment of histidine-tagged 70S ribosome complexes to GUV membranes using nitrilotriacetic acid (NTA) affinity. Related to Figure 4. (A) Confocal microscopy images of GUVs encapsulating 1 µM of 70S ribosomes stained with 100 µg/ml propidium iodide (PI). GUVs were generated with and without 2.7 mol% DGS-NTA(Ni) in the membrane. Scale bars are 10 µm. (B) Lumen/membrane ribosomal fluorescence intensity ratio derived from radial profiles of PI emission in individual GUVs. Mean values (solid lines) and standard deviations (dashed lines) are based on 30 individual GUVs (filled circles). Different letters represent statistically significant differences (p < 0.001; one-way ANOVA using Tukey (HSD) post-hoc analysis).
Figure S11. Purified, histidine tagged ribosomes are functional in the PURE system. Related to Figure 4. SDS-PAGE analysis of PURE reactions containing no ribosomes (lane 2), 1 µM kit supplied ribosomes (lane 3) or 1 µM self-purified, histidine-tagged ribosomes (lane 4). Reactions were supplemented with pDNA encoding α-hemolysin and fluorescently-tagged lysine amino acids. The resulting gel was probed using coomassie (left hand image) and BODIPY fluorescence (right hand image).
**Transparent Methods**

**Chemicals**

All lipid products were purchased from Avanti Polar Lipids. Alexa Fluor-conjugated antibodies were purchased from Thermo Fisher Scientific. All other chemicals were purchased from Sigma-Aldrich unless otherwise stated.

**Cloning procedures**

Plasmid DNA containing the gene encoding green-light absorbing proteorhodopsin from the SAR86 group of marine γ-proteobacteria was kindly provided by Prof. Edward DeLong (Beja et al., 2000). The vector backbone used for the construction of all cell-free expression constructs was the control plasmid supplied with the PURExpress kit (New England Biolabs). This vector contains a T7 promoter, ribosome binding site and T7 terminator sequence with an ampicillin resistance gene for selection. The PR encoding gene was amplified by PCR to remove the endogenous stop codon and was ligated into the cell-free expression vector between KpnI and BamHI (for fusion constructs) or XhoI (for PR only constructs) restriction sites. Meanwhile, PR truncations were introduced by PCR with a start codon replacing each removed residue. These sequences were then ligated into the expression vector using the same restriction sites as the full-length construct. The gene encoding EGFP was then amplified by PCR and inserted downstream of both full-length PR and PR truncations in between BamHI and XhoI restriction sites. For EGFP only plasmids, PR was removed by PCR before re-ligation of the linearised vector. The 84-residue encoding linker sequence of tolA was amplified by PCR directly from E. coli colonies and was ligated downstream of PR and PRΔN between BamHI and SphI restriction sites. The 17-residue encoding sequence of secM was also amplified directly from E. coli and ligated downstream of tolA between SphI and XhoI restriction sites.

The sequence for the mRNA aptamer Spinach2 was 5’ – GATGTAACTGAATGAAATGGTGAAGGACGGGTCCAGTAGGCTGCTTCGGCAGCCTACTTGTTGAGTGAGTGTGAGCTCCGTAACTAGTTACATC- 3’ and was amplified directly from DNA oligonucleotides (Integrated DNA Technologies) by PCR and inserted at the 3’ end of PR and PRΔN constructs between BamHI and XhoI restriction sites.

The sequence encoding the HA epitope was 5’ -TATCCGTATGAGCCGGATTATGCGC- 3’ and was used directly from DNA oligonucleotides (Integrated DNA Technologies) with 20 base pair complementary overhangs for Gibson Assembly reactions to generate PR and PRΔN constructs with HA epitopes in the second periplasmic/extracellular loop and the third intracellular loop of PR. All constructs were confirmed by sequencing prior to experimental use.

**Bulk cell-free protein synthesis using the PURE system**

Cell-free reactions were carried out using the PUREfex1.0 (GeneFrontier) cell-free transcription-translation system. Reaction volumes used for bulk synthesis were 10 µl and were set up according to the manufacturer’s protocol. Each reaction was supplemented with 0.3 µl of FluoroTect™ GreenLys in vitro translation labelling system (Promega), 0.5 µl of Murine RNase inhibitor (New England Biolabs), 20 ng µl-1 of plasmid DNA and 1 mg/ml of POPC (Avanti Polar Lipids) LUVs. LUV stocks (10 mg/ml in 50 mM HEPES-KOH (pH 7.6), 100 mM potassium glutamate and 15 mM magnesium acetate) were generated by manual extrusion through 0.1 µm polycarbonate membranes using a mini-extruder (Avanti Polar Lipids). Reactions for PR constructs were supplemented with 100µM all-trans-retinal. Reactions were incubated at 37°C for three hours in the dark. SDS-PAGE loading buffer (4x) was then added directly to reaction mixtures and samples were incubated for 15 minutes at 65°C to avoid the formation of precipitates. Samples were then used directly for SDS-PAGE. In-gel fluorescence from labelled amino acids was analysed using a Typhoon FLA 9000 (GE Healthcare) with an excitation wavelength of 473 nm and a long-pass emission filter for detection of wavelengths > 510 nm. Gels were then recovered and stained with coomassie to confirm the presence of the protein components of the PURE system.

**Preparation of giant unilamellar vesicles (GUVs) and cell-free reaction encapsulation**
GUVs were generated using the droplet transfer method as previously described (Altamura et al., 2017; Noireaux and Libchaber, 2004). An aqueous/lipid interface was generated by gently layering 300 µl of 0.5 mM POPC (Avanti Polar Lipids) solubilized in light mineral oil on top of an outer aqueous solution supplemented with 200 mM glucose (see below for solution make-up). The resulting two-phase solution was left at room temperature for 2 hours to allow saturation of the interface with POPC molecules. Meanwhile, 20 µl of an inner solution supplemented with 200 mM sucrose was prepared and transferred into 600 µl of fresh 0.5 mM POPC in light mineral oil. This solution was then emulsified by pipetting for 30 seconds and gently layered on top of the previously prepared, now lipid saturated, aqueous/lipid interface. The entire mixture was then centrifuged at 2,500 x g for 10 minutes and pelleted GUVs were collected using a pipette following the careful removal of the oil phase.

For cell-free reactions, the inner solution was composed of the PURE cell-free reaction mixture supplemented with 200 mM sucrose, 1 µl murine RNase inhibitor (New England Biolabs) and 20 ng µl-1 of the relevant plasmid DNA. Additionally, for cell-free reactions with His-tagged ribosomes, the supplied ribosome solution was omitted and 1 µM of self-purified His-tagged ribosomes was added (see below for purification and labelling methods). The outer solution was composed of the small molecular weight components of the PURE system (0.3 mM of each of the 20 amino acids, 1.5 mM spermidine, 3.75 mM ATP, 2.5 mM GTP, 1.25 mM of both CTP and UTP, 25 mM creatine phosphate, 1.5 mM 1,4-dithiothreitol (DTT), 18 mM magnesium acetate, 280 mM potassium glutamate, 50 mM HEPES-KOH pH7.6 and 0.02 mg ml-1 folinic acid), supplemented with 200 mM glucose and osmotically matched to the inner solution with NaCl (typically 75 mM) using a Vapro 5520 vapor pressure osmometer (Wescor). Following collection, GUVs were incubated for three hours at 37°C in the dark. All reactions involving the synthesis of PR or PRΔN were supplemented with 100µM all-trans-retinal following GUV collection and prior to incubation.

For ribosome binding experiments the inner solution was composed of 100 mM potassium glutamate, 18 mM magnesium acetate, 50 mM HEPES-KOH pH7.6, 1 µM 70S ribosomes and 200 mM sucrose. The outer solution was composed of 100 mM potassium glutamate, 18 mM magnesium acetate, 50 mM HEPES-KOH pH7.6 and 200 mM glucose. GUVs were collected and visualised immediately.

For mRNA localisation experiments, ribosomes were omitted from the reaction mixture to avoid translation of mRNAs and 3,5-difluoro-4-hydroxybenzylidene imidazolinone (DHFBI) was added to a final concentration of 20 µM in the inner and outer aqueous solutions. Following collection GUVs were incubated at 37°C for one hour before visualization.

**DiBAC4(3) fluorescence quenching assay**

PURE encapsulating GUVs were prepared as previously described and incubated for 20 minutes in the dark at 37°C. GUV suspensions were then transferred to 4°C and were either used immediately or stored for no more than 2 hours. DiBAC4(3) was added to a final concentration of 1 µM and vesicle suspensions were allowed to settle in microscopy chambers for 20 minutes in the dark prior to imaging. GUVs containing PR D97N, PRΔN and PR were then excited with a 488 nm argon laser for excitation of both DiBAC4(3) and PR. Vesicles were constantly excited for 290 seconds with quantification of membrane bound DiBAC4(3) fluorescence performed every 5 seconds.

**Hemagglutinin (HA) antibody binding assay**

GUVs collected following synthesis of the relevant protein were re-suspended in fresh outer solution supplemented with 1 mg ml-1 of Alexa Fluor 647-conjugated HA antibody (Thermo-Fisher Scientific). The solution was mixed gently by pipetting and was incubated at 37°C for 30 minutes. GUVs were then pelleted by centrifugation at 2,500 x g for 10 minutes at room temperature, antibody-containing solution was carefully removed, and GUVs were re-suspended in fresh outer solution.

**Affinity purification of histidine-tagged ribosomes and fluorescent labelling of ribosomal proteins**
E. coli strain JE28 was used for isolation of tetra-histidine-tagged 70S ribosomes using a previously established protocol (Ederth et al., 2009). Briefly, JE28 cultures were grown shaking at 37°C in LB media supplemented with 50 µg ml-1 kanamycin. At OD600 ≈ 1 cultures were slowly cooled to 4°C. Cells were then pelleted by centrifugation at 4,000 rpm for 30 minutes at 4°C and re-suspended in buffer A (20 mM HEPES-KOH pH 7.6, 10 mM magnesium chloride, 150 mM potassium chloride and 30 mM ammonium chloride) supplemented with 0.5 mg ml-1 lysozyme and 10 µg ml-1 RNase-free DNase I (Promega). Cell suspensions were further lysed by sonication. Cell lysates were subjected to two rounds of centrifugation at 20,000 x g for 30 minutes at 4°C to remove insoluble material. Cleared lysates were then applied to a pre-packed Ni-NTA column equilibrated with un-supplemented buffer A. The column was washed with buffer A containing 5 mM imidazole before ribosomes were eluted with buffer A containing 150 mM imidazole. Pooled elution fractions were dialyzed twice for 2 hours against 5 L of buffer A at 4°C to remove imidazole. At this stage ribosomes were either pelleted by ultracentrifugation through a 30 % (w/v) sucrose cushion at 130,000 x g for 16 hours at 4°C and re-suspended in polymix buffer (5 mM magnesium acetate, 5 mM ammonium chloride, 95 mM potassium chloride, 0.5 mM calcium chloride, 8 mM putrescine, 1 mM spermidine, 5 mM potassium phosphate (pH 7.6) and 1 mM DTT), or were fluorescently labelled (see below). Ribosome concentration was determined by measuring absorbance at 260 nm and using an extinction coefficient of 3.91 × 107 M-1 cm-1 (Becker et al., 2012). Ribosomes were further concentrated to 20 µM using 10,000 Da MWCO concentration columns (BioVision).

For fluorescent labelling, Alexa Fluor 488 5-SDP ester (Thermo-Fisher Scientific) was added to 5 ml of imidazole-free ribosome solution at a concentration of 50 µg ml-1. The resulting solution was gently mixed at room temperature for one hour and then dialyzed against 5 L of dye-free solution for 2 hours at 4°C. This solution was then ultracentrifuged through a 30 % (w/v) sucrose cushion at 130,000 x g for 16 hours at 4°C. The resulting pellet containing fluorescently labelled ribosomes was re-suspended in polymix buffer and concentrated to 20 µM as before.

Nile red lipid order assay

GUVs collected following synthesis of the relevant protein using the PURE system were re-suspended in fresh PURE outer solution supplemented with 200 mM glucose and 0.1 µM nile red before being visualised using confocal microscopy. A 488 nm argon laser was used to excite nile red and emission was collected using band-pass filters for 510 - 590 nm and for 650 - 750 nm.

Microscopy

Confocal microscopy images were taken using a Leica TCS SP8 laser scanning confocal microscope equipped with a 63x oil immersion objective. Samples were placed in 8-well, uncoated polymer µ-slides (Ibidi) and allowed to settle to the bottom of the chambers for 10 minutes prior to imaging. Image analysis was performed using Fiji software equipped with a plugin to measure radial fluorescence profiles. Three-dimensional surface plots using a fire LUT were also generated using Fiji in order to enhance the visualization of samples with low fluorescence. Background adjustments were performed by taking three background readings from arbitrary points on each image and calculating the mean intensity.

Statistical analysis

Tests performed for statistical analysis were one-way ANOVA with Tukey (HSD) post-hoc analysis and two-tailed, two sample Student’s t-test. All statistical tests were performed using R version 3.4.3 and statistical significance was assumed with a p-value of < 0.001 unless otherwise stated.

Supplemental References

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