Intracellular membranes: Conserved mechanisms of formation and regulation throughout evolution

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Summary (one sentence max 40 words)
This review highlights how physical-chemical laws, conserved upon evolution, influence cell membrane morphology and phospholipid biosynthesis.

Abstract
Membrane remodeling and phospholipid biosynthesis are normally tightly regulated to maintain the shape and function of cells. Indeed, different physiological mechanisms ensure a precise coordination between de novo phospholipid biosynthesis and modulation of membrane morphology. Interestingly, the overproduction of certain membrane proteins hijack these regulation networks, leading to the formation of impressive intracellular membrane structures in both prokaryotic and eukaryotic cells. The proteins triggering membrane proliferation share two major common features: 1) they promote the formation of highly curved membrane domains and 2) they lead to an enrichment in anionic, cone-shaped phospholipids (cardiolipin or phosphatidic acid) in the newly formed membranes. Taking into account the available examples of membrane proliferation upon protein overproduction, together with the latest biochemical, biophysical and structural data, we explore the relationship between protein synthesis and membrane biogenesis. We propose a mechanism for the formation of these non-physiological intracellular membranes that shares similarities with natural inner membrane structures found in α-proteobacteria, mitochondria and some viruses-infected cells, pointing towards a conserved feature through evolution. We hope that the information discussed in this review will...
give a better grasp of the biophysical mechanisms behind physiological and induced intracellular membrane proliferation, inspiring new biotechnological applications.

**Keywords:** Membrane remodeling; membrane biosynthesis, membrane curvature; phospholipids; inner membrane; evolution
1. Introduction

Membranes are lipidic films that define the boundaries of cells and organelles. They constitute both a permeability barrier for aqueous compounds and a major site of exchange between the interior and exterior of these cells and compartments. Therefore, they are essential for compartmentalizing the biochemical reactions that sustain life. Membranes are composed of lipids arranged as bilayers, together with proteins that can be either inserted in the lipid layer or peripherally associated to it. Membrane organization, as well as lipid and protein constituents, vary between organisms (eukaryotic cells, bacteria, virus), but also among species of certain organism (bacteria). Furthermore, membrane composition changes in response to various signals or environmental conditions resulting in three-dimensional rearrangements, or membrane remodeling events. These spatial rearrangements occur in all life forms and are essential for important physiologic processes, such as cell division and differentiation, organelle formation and intracellular trafficking. The exact mechanisms underlying membrane remodeling have mainly been deciphered in eukaryotic cells and are starting to be understood in prokaryotes. Importantly, the knowledge gained from studying membrane remodeling in physiological context has allowed the establishment of a few basic physico-chemical principles that appear to also apply to non-physiological membrane rearrangements, such as membrane proliferation occurring upon protein overproduction.

1.1. Influence of membrane curvature in membrane remodeling

Eukaryotic cells possess various essential intracellular organelles (endoplasmic reticulum (ER), Golgi apparatus, endosome, mitochondria...), which differ in morphology, function and structure but are all bounded by bilayers with increased curvature compared to the cell membrane. In addition, multiple membrane remodeling events have to be coordinated to carry out physiological processes such as vesicular trafficking, endocytosis or exocytosis. These processes, which have been studied for decades, depend on a large array of proteins, either cytoskeletal (actin, tubulin) or directly implicated in membrane curvature and remodeling, such as clathrin, dynamins or BAR or ENTH domain-containing proteins. For a long time, these
complex membrane remodeling processes were thought to be an exclusive feature of eukaryotic cells. Recently, it has been demonstrated that prokaryotic cells also undergo multiple membrane remodeling processes which are similarly controlled by specific proteins, analogous to the ones found in eukaryotic cells (Vega-Cabrera and Pardo-López 2017). Regardless of the different protein complexes involved for each organism, from a biophysical perspective, four basic molecular mechanisms have been described to remodel biological membrane, modifying their curvature (Figure 1) (Zimmerberg and Kozlov 2005):

a) Pushing or pulling the membrane using molecular motors,

b) Bending along a rigid supramolecular protein scaffold,

c) Asymmetric interaction of proteins with only one leaflet of the lipid bilayer,

d) Insertion of wedge-shaped proteins into the membrane.

![Figure 1](image.png)

Figure 1. Different mechanisms of membrane deformation a) push force by molecular motors, b) protein supramolecular scaffolding, c) asymmetric interaction with one leaflet of the membrane and d) insertion of wedged-shaped proteins.

Those biophysical mechanisms are ubiquitous and involved in multiple physiological processes. For example, proteins with ENTH (Epsin NH2-Terminal Homology) domains in eukaryotes insert a wedged-shaped N-terminal amphipathic helix into the membrane, leading to curvature and deformation of the membrane (Figure 1a). ENTH-containing proteins are also involved in
clathrin mediated budding by regulating and promoting the scaffolding of clathrin on membrane (Figure 1b). However, for the moment proteins with ENTH domains have only been observed in eukaryotes.

Another example are BAR (Bin-Amphiphysin-Rvs) domains, which form banana-shaped dimers that can bind membrane with curvature but also induce and stabilize membrane curvature via helical oligomerization and scaffolding (Nishimura, Morone and Suetsugu 2018). BAR domain interaction also leads to phosphoinositides lipid clustering and formation of stable microdomains (Zhao et al. 2013). BAR-domain proteins functions are multiple: they regulate actin polymerization by interacting and recruiting actin assembly factors at the membrane, they cooperate with dynamin and clathrin to mediate membrane fission during endocytosis and are involved in the formation of filopodia and lamellipodia (Nishimura, Morone and Suetsugu 2018). Only 2 BAR-containing proteins have been described so far in bacteria, BdpA from *Shewanella oneidensis* is involved in the biogenesis of outer membrane extensions and trigger the formation of such extensions when expressed in a heterologous host (*E. coli*) (Phillips et al. 2020). MamY is a BAR-domain protein implicated in magnetosome formation (Tanaka, Arakaki and Matsunaga 2010).

The Dynamin superfamily of proteins includes some of the best studied membrane remodeling proteins; in particular, those involved in endocytosis, organelle formation and maintenance, and cytokinesis in eukaryotic cells. Dynamins are molecular motors that modify membrane curvature (Figure 1a). They are characterized by a GTPase domain, an elongated α-helical bundle that drives self-assembly, and the capacity to interact with lipids and membrane. They promote membrane tubules formation by self-assembling as an helical scaffold. GTP hydrolysis induces a conformational change in this scaffold that leads to further membrane constriction and possibly membrane fission or fusion (Jimah and Hinshaw 2019). In bacteria, multiple dynamin-like proteins (BDLP: Bacterial Dynamin Like Proteins) capable of modifying membrane curvature have been identified but their physiological roles are not entirely known. Some have been localized to the division septum, suggesting a role in cell division (Bramkamp 2012). The two BDLP DynA and DynB for instance, are necessary for the cytokinesis event
preceding sporulation in *Streptomyces venezuelae* (Schlimpert et al. 2017). Also, *Bacillus subtilis* DynA, a BDLP localized at the septa, mediates immunity against phage infection and membrane stress and it is also thought to involve membrane remodeling, since this protein is able to perform lipid mixing and membrane fusion *in vitro*, in a GTP-independent process (Bürmann et al. 2011; Sawant et al. 2016; Guo and Bramkamp 2019). A similar function in membrane repair and maintenance has been assigned to the *M. tuberculosis* BDLP IniA, which is capable to modify the membrane curvature of cardiolipin (CL)-containing liposomes and induce GTP-dependent membrane fission *in vitro* (Wang, Zhang and Xiao 2019). Finally, *Escherichia coli* LeoA, B and C are periplasmic BDLP implicated in outer membrane vesicles formation, although a direct interaction with lipids or a capacity to remodel membranes has not been demonstrated (Michie et al. 2014).

Another interesting illustration of physiological event requiring major membrane remodeling and membrane curvature modifications is bacterial cell division. Bacterial cells can be bounded by one or two membranes. Monodermic bacteria only possess one membrane, the cytoplasmic membrane, which is usually surrounded by a thick layer of peptidoglycan. In contrast, didermic bacteria have two membranes: the inner membrane (which corresponds to the cytoplasmic membrane of monodermic bacteria) and the outer membrane, separated by the so-called periplasm, which contains a peptidoglycan layer. During division, all those membrane layers must be remodeled to yield two independent bacteria. Membrane invagination required for bacterial cell division appears to be driven by forced membrane bending. Septum formation first requires the spatially regulated, hierarchical assembly of a multiprotein complex called the divisome. The correct positioning of the divisome at midcell is ensured by the Min and the Nucleoid Occlusion (NOc) systems, which are themselves dependent on and regulated by membrane lipids, especially anionic lipids microdomains found at the poles and septum (Mileykovskaya and Dowhan 2005; Barák and Muchová 2013). Amongst the more than 20 proteins that constitute the divisome, the tubulin-like GTPase FtsZ plays a central role in membrane invagination by oligomerizing into a dynamic, cytoplasmic Z-ring attached to the membrane via ZipA and FtsA. The complex further serves as a docking site for other divisome components. The role of FtsZ in membrane invagination has been debated. Its ability to display
intrinsic curvature in its polymeric state and deform various artificial membranes in vitro (Kretschmer et al. 2019), led to a model in which the cytoplasmic membrane is pulled inward by Z-ring constriction during cytokinesis. However, it has also been suggested that FtsZ only serves as a scaffold onto which the peptidoglycan remodeling machinery assembles. In the latter case, it is the growth of the septal cell wall that pushes the membrane toward the center of the cell (den Blaauwen, Hamoen and Levin 2017). In addition, the actin-like ATPase FtsA, which interacts with phospholipids via its C-terminus and bridges FtsZ to the membrane, was shown to induce membrane rearrangement in vitro and vesicle formation upon overexpression in E. coli (Conti, Viola and Camberg 2018). It was thus proposed to facilitate membrane invagination by deforming the membrane at the septum site (Krupka et al. 2014; Conti, Viola and Camberg 2018). Whatever its exact mechanism of formation, the membrane curvature generated upon membrane invagination in turn participates in the recruitment of negative curvature-specific proteins such as DivIVA, which further binds other players of cell division and localizes them at the septum site. The final steps of bacterial cell division (fusion and fission of the membrane(s) leading to the separation of the two daughter cells) are not characterized yet and it is still unclear whether specific fusion/fission proteins complexes are necessary or if membrane fission occurs spontaneously as a consequence of membrane curvature and/or protein crowding (Snead et al. 2017; Steinkühler et al. 2020). A role for FtsA in this process has been proposed based on the occasional scission observed when FtsA was added to FtsZ-liposome in vitro (Osawa and Erickson 2013). However, this is inconsistent with the fact that FtsA, FtsZ and ZipA leave the septum before cell separation (den Blaauwen, Hamoen and Levin 2017).

Sporulation is another event occurring in bacteria that involves extensive membrane remodeling. Under unfavorable conditions, the vegetative cells of some species of Gram-positive bacteria, such as Bacillus or Clostridium, produce resistant and metabolically dormant structures called spores. Sporulation starts with an asymmetrical cell division which generates a small cell (forespore) connected to the larger mother cell by two membranes separated by a thin phosphatidylglycerol (PG) layer. This step requires the same divisome complex described for vegetative growth. However, whereas septum formation during vegetative growth avoids
the nucleoid, during sporulation the septum closes over the chromosome, which is then translocated to the forespore. In *B. subtilis*, DNA translocation is performed by SpoIIIE, a homologue of *E. coli* FtsK which localizes at septal midpoint, possibly by its ability to sense regions of increased membrane curvature (Fleming et al. 2010). SpoIIIE is also required for membrane severing of the cytoplasmic bridge remaining between the mother cell and the forespore after DNA translocation. Although the exact mechanism by which SpoIIIE mediates this event is unknown, it was proposed that the protein forms multimeric channels in the mother cell and forespore membranes. Those channels finally assemble into an intramembrane trans-channel whose disassembly triggers membrane separation (Liu, Dutton and Pogliano 2006).

After severing of the membranes, the mother cell progressively engulfs the forespore, in a phagocytosis-like process, until the forespore is liberated in the cytoplasm of the mother cell, where it will further mature. During engulfment, the mother cell membrane generated at the division site expands and migrates on each side of the forespore until it totally surrounds it and the two leading edges reconnect next to the cell pole. The process of engulfment depends on two complementary ratchet-like mechanisms: the mother cell SpoIIDMP protein complex localized at the leading edge of the engulfing membrane, processively degrades the peptidoglycan synthesized ahead by the forespore and by doing so pulls the membrane forward (Meyer et al. 2010; Khanna et al. 2019). In addition, the forespore protein SpoIIQ interacts with the mother cell SpoIIAH in a zipper-like mechanism, which renders the membrane movement irreversible (Broder and Pogliano 2006). Once the two leading edges have meet, they fuse to liberate the forespore in the mother cell cytoplasm. In *Bacillus subtilis*, this final fusion event depends on SpoIIIE and FisB (Fission protein B) (Liu, Dutton and Pogliano 2006; Doan et al. 2013). FisB is a bitopic membrane protein with a large periplasmic region and a small cytoplasmic domain. Although the exact mechanism by which FisB triggers membrane fusion is still unknown, the protein is able to induce lipid mixing *in vitro*, in a process depending on the specific interaction between its periplasmic domain and CL. It is thus assumed that FisB, via this interaction, brings the two leading engulfment membranes in close contact before fusion (Doan et al. 2013). As expected, sporulation is dependent on a reactivation of *de novo* membrane lipids synthesis (Pedrido et al. 2013). CL for instance, is strongly enriched in the septum, the forespore and the mother cell engulfment membranes during sporulation (Kawai
et al. 2004) and accumulates in the mature spore (Kawai et al. 2006). Mutant strains producing only trace amounts of CL show delay in spore formation and produce reduced amounts of spores that are unable to germinate when placed back in favorable conditions (Kawai et al. 2006). CL enrichment might thus be important for the function of membrane proteins required for sporulation (e.g. FisB) or for their recruitment to specific regions of curvature. Membrane curvature-dependent localization has indeed been shown for B. subtilis SpoVM, which is necessary for spore maturation and localizes at the forespore surface by detecting positively curved membranes and inserting in them by an atypical amphipathic α-helix (Ramamurthi et al. 2009; Gill et al. 2015).

1.2. Evolutive origin of intracellular organelles

Although prokaryotic cells have been historically claimed as organelle-free organisms, several examples of intracellular membrane-restricted compartments have now been identified. For instance, intracellular membrane structures are naturally present in α-proteobacteria, an evolutive ancestor of γ-proteobacteria (Gupta 2000), where they either increase the efficiency of the cell bioenergetic metabolism (anaerobic anoxygenic photosynthesis, nitrifying and/or methanotrophic bacteria, etc.) or provide an evolutive advantage (magnetosome) (Muñoz-Gómez et al. 2017).

Those membrane structures have been proposed as potential ancestors of mitochondria inner membrane cristae after the discovery of a common membrane remodeling protein: alphaMic60 and Mic60 in α-proteobacteria and mitochondrion, respectively (Muñoz-Gómez et al. 2015). The growth of intracellular membrane structures in both α-proteobacteria and mitochondria requires the assembly of the photosynthetic or respiratory protein complexes, which are known to induce strong membrane curvature (Woronowicz et al. 2013; Horvath et al. 2015; Niederman 2016). Mic60 and its analogue alphaMic60 are part of the protein complex that presumably bends the membrane and stabilizes the cristae junctions in mitochondria (or inner membrane invagination points in α-proteobacteria). Although modern γ-proteobacteria lack the gene encoding alphaMic60 (Huynen et al. 2016) and have lost the ability to physiologically produce inner
membrane structures, heterologous overproduction of eukaryotic Mic60 restores the capacity of E. coli to produce those inner membrane structures (Tarasenko et al. 2017). This result implies that the ancestral mechanism for inner membrane proliferation of α-proteobacteria is not completely lost in γ-proteobacteria, but only dormant, and that it can be restored when certain conditions are met. The presence of CL in both prokaryotes and mitochondria is another argument often used to defend the endosymbiotic origin of mitochondria (Tian, Feng and Wen 2012). CL fulfills many biological roles in mitochondria and bacteria (Mileykovskaya and Dowhan 2009; Paradies et al. 2019). Membrane curvature and CL seem to be linked, as CL depleted mutants show altered internal ultrastructure and function of mitochondria (Matsumura et al. 2018). Furthermore, a recent study illustrates the importance of the formation of highly curved cristae for the correct accumulation of CL inside the mitochondria (Kojima et al. 2019). Notably, CL seems to be also related to physiological membrane remodeling processes in bacteria such as sporulation (section 1.1). Still, how CL, membrane curvature and membrane biosynthesis are exactly related remains a mystery.

Magnetosomes are ~30-120nm spherical, membrane-bound compartments that contain iron-rich magnetic particles. They organize as chains along the cell and allow magnetotactic bacteria to sense and orient in the geomagnetic field. They derive from the cytoplasmic membrane to which they may remain attached or not (Barber-Zucker and Zarivach 2017). The mechanism of membrane invagination and vesicle formation, which seems to precede biomineralization, has not been completely deciphered but depends on the product of multiple mam (magnetosome membrane-associated) genes. Individual deletions of mam genes have identified four proteins involved in the biogenesis of the membrane of magnetosomes (Maml, MamL, MamQ and MamB) (Murat et al. 2010; Raschdorf et al. 2016). However, the structure of those proteins has not been resolved and only speculations are available on the mechanism they use to bend the membrane and create the invaginations necessary for magnetosome formation (Komeili 2012; Barber-Zucker, Keren-Khadmy and Zarivach 2016). Besides, the overexpression of each one of those four genes alone is not sufficient to trigger membrane proliferation. More recent studies however suggest that at least some Mam protein might directly induce membrane curvature. In particular, MamY, a BAR domain-containing protein, interacts with liposome and induces
liposome tubulation *in vitro*. Because a ∆mamY *Magnetospirillum magneticum* mutant showed altered magnetosome size-distribution, MamY was first proposed to be involved in membrane constriction (Tanaka, Arakaki and Matsunaga 2010). However, MamY *in vitro* tubulation activity is specifically increased upon CL interaction, suggesting that MamY might recruit CL to the site of magnetosome formation to induce the formation of highly curved membranes (Tanaka *et al.* 2018). Still, overexpression of MamY in *E. coli* or *M. magneticum* did not alter cell membrane morphology, confirming that *in vivo*, other factors are certainly needed to trigger membrane curvature and vesicle formation (Tanaka, Arakaki and Matsunaga 2010). More recently, another study proposed that MamY represents a membrane positive curvature-sensing element and serves as a scaffold to properly align the chain of magnetosome parallel to the axis of the cell (Toro-Nahuelpan *et al.*, 2019). The role of CL in this function was however not tested.

In some photosynthetic bacteria, intracytoplasmic vesicles called chromatophores contain pigments and light-harvesting proteins used to perform photosynthesis. Chromatophores function depend on the light-harvesting complexes 1 (LH1) and 2 (LH2) together with the reaction center (RC). These complexes, which are also directly implicated in chromatophore formation and shape determination, are thought to induce membrane curvature through a combination of wedging and scaffolding mechanism (Figure 1). Indeed, the ability of these integral membrane proteins to bend and deform membranes depends on their capacity to oligomerize. The RC-LH1 complex, when monomeric, cannot bend membrane. However, RC-LH1 in complex with the small protein PufX forms dimers with the two monomers bent by a 146° angle (Qian, Bullough and Hunter 2008; Hsin *et al.* 2009; Tucker *et al.* 2010). In the absence of LH2, these dimers form tubular chromatophores *in vivo* (Chandler *et al.* 2008). LH2 is also sufficient to induce membrane curvature in *R. sphaeroides*. The protein forms hexagonally packed complexes, which are localized at high membrane curvature regions and, according to molecular dynamic simulation, could also induce membrane curvature (Chandler *et al.* 2009; Hsin *et al.* 2009; Scheuring *et al.* 2014). The combined action of LH2 and RC-LH1-PufX would thus allow for the formation of spherical shaped chromatophore.
1.3. Hijacking membrane remodeling: lessons learned from viral infection

In addition to the aforementioned membrane-remodeling physiological events, intracellular membranes can also be reshaped during infection by peculiar viruses able to usurp host lipid metabolism to create new compartments dedicated to their replication or replication organelles (Figure 2). Viruses infecting a large variety of hosts, ranging from bacteria and unicellular eukaryotes to vegetal and animal cells, have been described that trigger this phenomenon (Romero-Brey and Bartenschlager 2016). Among those, positive-sense single-stranded RNA viruses (+RNA) infecting eukaryotic cells are the most studied. Because the +RNA of those viruses has the same sense as the cellular messenger RNA it is immediately translated when it reaches the cellular cytosol. Thus, viral proteins capable of modifying membrane curvature, which will be discussed in more detail in the following sections, are readily produced in the early infection stage (Harak and Lohmann 2015; Shulla and Randall 2016).

Figure 2. Electronic tomography reconstruction of the replication organelles of some +RNA viruses. a) Left: Interconnected reticular network induced by dengue virus infection. The cytosolic face of the intracellular membranes is shown in brown and the ER lumen in black. Right: Viral particles (red) found in continuous ER cisternae. ER membranes are colored in light brown and inner vesicle membranes in dark brown (Welsch et al. 2009). b) Left: Surface model of Kunjin virus replication organelles; ER membranes are colored in red, ribosomes in white and viral RNA in yellow. Right: Vesicles (white) connected to each other and to the ER membranes (red) (Gillespie et al. 2010).
In addition, a correlation between membrane curvature and lipid biosynthesis during +RNA viral infection has been proposed (Miller and Krijnse-Locker 2008). Indeed, viral proteins modulating membrane curvature were also shown to promote the formation of membrane contact sites and the recruitment of host factors involved in lipid metabolism (van der Schaar et al. 2016), in particular phosphatidylinositol-4-phosphate (PI4P) or phosphatidylethanolamine (PE) synthesis (Altan-Bonnet 2017). Interestingly, the accumulation of PIP4 or PE is often accompanied with an enrichment in sterol that might contribute to the stabilization of membrane curvature and is important for the replication of the virus (Zhang et al. 2019).

In addition to +RNA viruses, other viruses containing double-stranded RNA (Reoviruses) as well as DNA viruses (Poxvirus, Vaccina virus, African swine fever virus, Frog Virus 3 and Paramecium Bursaria Chlorella Virus, giant Mimivirus Acanthamoeba polyphaga) also induce massive host membrane rearrangements. Although less studied, those viruses also rely on the production of proteins modifying the curvature of the host membrane (Weisberg et al. 2017; Moss 2018, Mutsafi et al. 2013)). The membrane-enveloped double stranded RNA bacteriophages from the cystoviridiae family, such as phage φ6, are the only known enveloped phages and are evolutionarily related to the +RNA eukaryotic virus picornavirus (Koonin, Dolja and Krupovic 2015). They also produce proteins capable of bending the inner membrane of their hosts (Gram negative bacteria) which are necessary for virus replication (McGraw, Mindich and Frangione 1986).

In summary, certain viruses hijack the lipid metabolism of their hosts using specific proteins that modify membrane curvature and host co-factors to alter the lipid composition of the membrane to favor their own replication. However, how those factors are related to de novo membrane biosynthesis and viral replication organelles assembly remains elusive and should be further investigated.

1.4. Inner membrane proliferation upon overproduction of some membrane proteins.
Overproduction of recombinant membrane proteins is usually difficult due to various limitations, including a shortage of membrane space needed to accommodate the produced proteins. In a few peculiar cases however, overproduction of membrane proteins, either in prokaryotic or eukaryotic cells, has revealed an unexpected and intriguing ability of cells to synthesize an excess of internal membranes. In fact, these newly synthesized inner membranes often contain large amounts of well-folded recombinant proteins, holding great promises for biotechnological applications. Since the pioneer observation of Weiner et al. (Weiner et al. 1984), only a few dozen membrane-associated proteins from prokaryotic (Table 1) and eukaryotic (Table 2) origin have been described that trigger non-physiological lipid membrane proliferation or “inducible intracellular neo-membranes” (Figure 3).

Figure 3. Negative-staining TEM pictures of some examples of inner membrane proliferation upon membrane protein overproduction. a) Longitudinal (left) and transversal (right) sections of *E. coli* inner membrane tubules after fumarate reductase overproduction (Weiner et al. 1984).
b) Onion-like vesicles formed upon overproduction of protein 3A of Foot and Mouth Disease Virus (FMDV) (Weber et al. 1996).

c) S. cerevisiae cell with the cytosol (cyt), nucleus (n) and the staked membranes “Karmellae” (k) around the nucleus (n) (left) and detail of those membranous structures (right) after 3-Hydroxy-3-methylglutaryl-CoA reductase (Hmg-CoA) overproduction (Wright et al. 1988).

d) Vesicles formed in S. cerevisiae upon overproduction of poliovirus protein 2BC (Barco and Carrasco 1995).

From a morphological point of view, those inducible intracellular neo-membranes can be related to the bioenergetic compartments of α-proteobacteria and mitochondria (Arechaga 2013), and/or replication organelles of +RNA viruses (Miller and Krijnse-Locker 2008). Furthermore, similarly to those “natural” intracellular compartments most of the proteins triggering inducible intracellular neo-membranes (Table 1 and Table 2) also create zones with high membrane curvature (Jamin et al. 2018). For this reason, “natural” and “induced” intracellular membrane proliferation might share a more profound relation that goes beyond simple morphological resemblance.

2. Mechanisms of protein-induced membrane curvature

Modulation of membrane curvature is often at the midst of both physiological and induced membrane remodeling processes. The induced membrane proliferation upon protein overproduction has, however, the advantage of being decoupled from the cell physiological regulations. For this reason, we will examine the current knowledge on the mechanism of inner membrane proliferation, focusing on the influence of membrane curvature not only on membrane morphology but also on phospholipid biosynthesis. In particular, four important questions about inner membrane proliferation upon protein overproduction remain: 1) how can overproduced proteins induce the deformation of the inner membrane creating different morphologies; 2) what are the characteristics of the proteins triggering lipid biosynthesis and, thus, inner membrane proliferation; 3) how are protein overproduction and de novo phospholipids biosynthesis coordinated; and 4) can we find regulatory mechanisms conserved
across evolution explaining internal membrane proliferation in both prokaryotic and eukaryotic cells.

In order to yield the observed morphologies (vesicles, tubules, stacks of flat membranes, etc.) listed in Table 1 and Table 2 and illustrated in Figure 3, proteins inducing inner membrane proliferation must modify membrane curvature by means of one (or the combination of several) of the general mechanisms previously proposed (molecular motors, supramolecular scaffolding, asymmetric membrane interaction, wedging, see Figure 1). However, the production of a pulling or pushing force will not be discussed in this section since none of the reviewed proteins is a molecular motor, nor a scaffolding protein interacting with any molecular motor.

2.1 Protein-protein supramolecular interactions

The construction of a 3D supramolecular scaffold via supramolecular interactions is an efficient way of controlling cell membrane curvature and is used by many proteins involved in membrane remodeling processes (e.g. endocytosis, fission, motility, membrane trafficking, etc.) (Simunovic et al. 2016). Most of the proteins inducing intracellular membrane proliferation (Table 1 and Table 2) have been described to form supramolecular assemblies around the lipid bilayer. The membrane curvature and, consequently, the inner membrane morphology observed in electronic microscopy will depend on the shape, nature and concentration of monomers constituting the supramolecular scaffold.

Heterologously expressed caveolin-1 in E. coli cells is perhaps the best-characterized example of how a single membrane protein can shape the morphology of the newly synthesized lipid bilayer. Caveolin-1 is a scaffolding protein involved on the formation of vesicles (caveolae) arising from the plasma membrane of eukaryotic cells (Parton 2018). The formation of heterologous-caveolae (h-caveolae) derived from E.coli inner membrane is linked to the assembly of caveolin-1 into a supramolecular cage (Walser et al. 2012). This cage contains around 160 caveolin-1 monomers and is similar in structure and size to eukaryotic caveolae. The three membrane-interacting domains and the oligomerization domain of caveolin-1 are
required for inner membrane proliferation (Ariotti et al. 2015). The formation of a regular and well-defined caveolin-1 scaffold seems to impose a strong local curvature on the cell membrane, causing the budding of vesicles coated with caveolin-1, and triggering the biosynthesis of phospholipids. As a consequence, monodisperse vesicles of the same size as those found in eukaryotic cells, accumulate in the *E.coli* cytosol.

The overproduction of phage PM2 protein P6 represents another instance of membrane proliferation presumably induced by a supramolecular cage (Männistö et al. 1999). Although the structure of P6 has not been studied when it is overproduced in *E. coli* membrane, the structure of the entire PM2 phage has been determined by crystallography. This phage, which infects Gram negative bacteria from the *Pseudoalteromonas* genus (Abrescia et al. 2008), is composed of an icosahedral protein capsid containing a lipid membrane that encloses a double stranded DNA (dsDNA). P6 interacts with the viral lipid membrane but it is not a capsid forming protein. It associates with the P3 protein to form a well-ordered supramolecular structure that confers icosahedral symmetry to the lipid bilayer. P6 is inserted at the edges, whereas two P3 dimers stabilize the facets of the icosahedra. In other words, P6 is in charge of “welding” the lipid bilayer to create the icosahedral vertices. In the absence of P3, P6 is still able to impose a curvature to the inner *E.coli* cell membrane, producing vesicles (Armour and Brewer 1990). Foot-and-mouth disease virus (FMDV) protein 3A is another example of a viral protein modulating membrane curvature. During the early stages of infection, FMDV, like other +RNA virus, remodels the ER membrane of its hosts (mammalian cells) to form the viral replication organelle, which provides a platform for viral RNA replication (Breese and Graves 1966). The interaction between the non-structural viral proteins (such as FMDV protein 3A) and the host phospholipids seems to be the trigger for host membrane remodeling (Deng et al. 2010). When overexpressed in *E.coli*, FMDV protein 3A alone is able to deform the inner membrane, producing onion-like vesicles instead of its characteristic replication organelles (Weber et al. 1996). The lack of other viral proteins or the differences in the nature and composition of phospholipids between eukaryotes and prokaryotes might explain this change in morphology. The precise mechanism of membrane deformation by FMDV protein 3A is unknown, but it requires the central amphipathic helix of the protein, together with the two cytosolic N- and C-
terminal domains (González-Magaldi et al. 2014), which might interact with other viral proteins or induce oligomerization with other copies of FMDV 3A. Similarly, phage φ6 is a dsRNA bacteriophage, evolutionarily related to eukaryotic +RNA viruses (section 1.4), which infects Gram negative bacteria from the *Pseudomonas* genus. The overproduction of P9 and P12 phage φ6 proteins induce the proliferation of intracellular vesicles in *E. coli* and *P. syringae* (Johnson and Mindich 1994; Sarin *et al.* 2012; Myhrvold, Polka and Silver 2016; Lyytinen, Starkova and Poranen 2019). Both proteins (P9 and P12) are required to create the lipidic envelope of the phage (Mindich, Sinclair and Cohen 1976). P9 is a transmembrane protein, which is produced in large quantities in the early stages of infection and inserted in the inner membrane (McGraw, Mindich and Frangione 1986; Johnson and Mindich 1994; Jung, Jung and Lim 2015). P12 is a non-structural protein and its role in the creation of the viral envelope is unknown, although it seems to somehow inhibit the degradation of P9 by the host proteolytic enzymes (Johnson and Mindich 1994; Lyytinen, Starkova and Poranen 2019).

Overexpression of *E. coli* fumarate reductase results in the formation of an array of densely packed lipid tubules in *E. coli* cytosol, that are severed from the inner membrane (Elmes, Scraba and Weiner 1986). These lipid tubules are stabilized by a scaffold of fumarate reductase packed in a regular helical configuration containing 10 proteins per helix turn (Weiner *et al.* 1984). Similar tubules are also observed when succinate dehydrogenase is overexpressed in *E. coli* (Maklashina, Berthold and Cecchini 1998). Probably, the mechanism of tubule stabilization is similar to that observed with fumarate reductase due the structural and functional similarities between these two enzymes (Ruprecht *et al.* 2009; Starbird *et al.* 2018). Although the supramolecular packing of succinate dehydrogenase has not been studied in depth, the authors observed different morphologies (tubules or vesicles) depending on the expression level of the protein (Maklashina, Berthold and Cecchini 1998). The supramolecular array of succinate dehydrogenase necessary to stabilize the tubule morphology might only be formed if the protein is produced at sufficient level. Thus, below a critical concentration, succinate dehydrogenase is still able to deform the membrane and yield vesicles but it is not capable of maintaining the tubule structure. The *sn*-Glycerol-3-P acyltransferase is another example of protein inducing membrane tubules formation upon overproduction (Wilkinson *et al.* 1986). The
individual molecules of sn-Glycerol-3-P acyltransferase are arranged in dumbbell-shaped dimers, which are packed in a left-handed helix along the tubule axis (Wilkison et al. 1992). The association of six sn-Glycerol-3-P acyltransferase dimers completes a helix turn.

The whole F₀F₁ ATP synthase (von Meyenburg, Jogersen and van Deurs 1984), and more efficiently, its b subunit alone (Arechaga et al. 2000), also produce vesicles and tubules detached from the inner membrane when overexpressed in E. coli. There is no data about the supramolecular packing of the b subunit of F₀F₁ ATP synthase in the lipid bilayer. Nevertheless, interactions between adjacent proteins seem to be important, since the removal of the cytosolic dimerization domain of the b subunit of F₀F₁ ATP synthase (residues from 53 to 122 (Revington, Dunn and Shaw 1999)) inhibits tubules formation (Arechaga et al. 2000).

The overproduction of the serine chemotaxis receptor (Tsr) from E. coli also triggers inner membrane proliferation (Lefman et al. 2004). Tsr is a transmembrane protein with a periplasmic domain that binds small molecules (Tsr is specific to serine) and a cytoplasmic domain associated with the adaptor protein CheW and the kinase CheW (Grebe and Stock 2004). In normal physiological conditions, cytoplasmic domains of adjacent Tsr form trimers of Tsr dimers, and self-assemble in two-dimensional clusters concentrated at the bacterial cell poles (Kim, Wang and Kim 2002). When overproduced, Tsr is also organized as trimeric assemblies of dimers (Lefman et al. 2004). However, because Tsr amounts are significantly increased, the two-dimensional clusters of Tsr can interact with each other creating a three-dimensional pseudo-hexagonal crystalline array that folds the inner membrane (Lefman et al. 2004). If this crystalline array is destroyed, e.g. by overproducing Tsr partners (CheW and CheW) at the same levels as Tsr, membrane proliferation is inhibited, even at high Tsr concentration in the membrane (Zhang et al. 2007). This result suggests that the high membrane curvature imposed by the crystalline array of Tsr is necessary to trigger phospholipid biosynthesis.

Membrane curvature induction by protein overproduction is not restricted to prokaryotic hosts and has also been observed in eukaryotic cells (Table 2). Unfortunately, structural data on the arrangement of the recombinant proteins in the newly synthesized inner membranes are lacking.
Still, there are some hints pointing to the presence of supramolecular scaffolds. For example, the minimal protein fragment of cytochrome P450 NADPH reductase and of canine ribosome receptor (RRp), which both induce membrane proliferation, include the charged residues flanking the hydrophobic transmembrane domain, as described for prokaryotic proteins (Kärgel et al. 1996; Becker et al. 1999). Moreover, it has been reported that, in some cases, the morphology of the newly produced membranes depends on the amount of overproduced membrane protein (Albertini et al. 2001). Disturbance of the 3D structure of the protein by introducing proline mutations (Vergères et al. 1993), GFP fusions in critical positions (Rubino, Di Franco and Russo 2000), partially misfolding the proteins (Zimmer et al. 1997) or by deleting oligomerization domains (Profant et al. 1999), results in an altered morphology of the inner membrane structures. In addition, protein-protein supramolecular interactions are also required for inner membrane proliferation in mammalian cells (Snapp et al. 2003). The production of chimeras with cytochrome b5 transmembrane domains and a dimerization-prone GFP changes the morphology of proliferating membranes in the ER from stacks of membranes to a bi-continuous phase with cubic symmetry. Furthermore, membrane proliferation is also induced by fusing dimerization prone GFP to ER resident proteins; whereas each proteins overproduced separately, do not trigger membrane proliferation. Taken together, these features are in line with the observations made in prokaryotes, and suggest that protein-protein supramolecular interactions are an ubiquitous mechanism to control membrane curvature cells.

2.2 Asymmetric interaction of proteins with only one leaflet of the lipid bilayer.

Besides the formation of a supramolecular scaffold, the asymmetric insertion of a membrane protein into one leaflet of the lipidic bilayer can lead to a modification of the membrane curvature. Such a mechanism have been hypothesized for monotopic proteins inducing membrane proliferation such as MurG (Van Den Brink-van Der Laan et al. 2003), LpxB (Metzger IV and Raetz 2009) and PtmA (Linna et al. 2017) and confirmed by sequence comparison and in silico studies of aMGS (Eriksson et al. 2009; Ge et al. 2014). In addition, PtmA and aMGS have been purified and their ability to remodel synthetic liposomes into lipid tubules has also been demonstrated in vitro (Ariöz et al. 2014; Ge et al. 2014; Linna et al. 2017).
2.3 Insertion of wedge-shaped proteins into the membrane.

The insertion of wedge-shaped protein into the lipid bilayer can also modulate membrane curvature (Lee 2004). Most transmembrane domains of membrane proteins introduce a packing mismatch in the lipid bilayer, which alters the membrane curvature. A well-known example of this mechanism is the wedge-shaped bacteriorhodopsin, a light-driven proton pump expressed in archaeabacteria under anaerobic conditions. Bacteriorhodopsin is found in highly curved, specialized membrane microdomains (purple membranes), in which it forms trimeric, hexagonal units packed in a 2D crystalline lattice together with archaeal lipids (Oesterhelt and Stoeckenius 1971; Henderson 1977). These lipids also contribute to the 2D crystalline packing as bacteriorhodopsin mutants with a constitutive wedge-shaped structure still need a specific lipidic environment to induce membrane curvature (Krebs and Isenbarger 2000; Rhinow and Hampp 2010; Yokoyama et al. 2014). Therefore, both phospholipid composition and protein tertiary 3D structure work together to modulate the membrane curvature in purple membranes.

It is remarkable how newly produced inner membranes of prokaryotes are enriched in cone-shaped non-bilayer forming lipids (cardiolipin (CL) or lyso-phospholipids) (Table 1). Similarly to wedge-shaped proteins, those cone-shaped phospholipids can also modulate membrane curvature. Their effect is expected to be less important than those induced by proteins, but not be negligible. Indeed, eukaryotic caveolae are enriched in phosphatidylinositol (PI) and cholesterol, which are important for the organization of lipid rafts. Similarly, E.coli adapts the phospholipid composition of h-caveolae replacing anionic PI by PG and non-bilayer forming cholesterol for lyso-phospholipids, only found in trace concentration in normal physiological conditions (Walser et al. 2012). Symmetrically, in silico simulations show that membrane bending is facilitated by the incorporation in the lipid bilayer, of CL at levels similar to the ones observed in cases of membrane proliferation upon protein overproduction (Boyd, Alder and May 2017). Furthermore, two examples have recently shown the importance of CL enrichment in the morphology of overproduced inner membranes. Firstly, E. coli mutants with reduced amounts of CL (2% vs 24% mol in wild type) changed the organization of the produced inner
membrane from organized tubules to onion like vesicles upon overproduction of the b subunit of F₀F₁ ATP synthase (Carranza et al. 2017). E. coli mutants completely depleted of CL confirmed this change of morphology and, in addition, were less efficient in triggering the proliferation of inner membranes (Carranza et al. 2017). Secondly, CL is necessary for the formation of inner membrane vesicles by PmtA, as it has been demonstrated both in vitro and in vivo (Linna et al. 2017). In vitro, PmtA is unable to deform synthetic liposomes lacking CL and no vesicle-like structures were observed in vivo after PmtA overproduction in CL-deficient bacterial mutants. Together, both examples illustrate the central role of CL in the proliferation of inner membranes.

The phospholipid composition of the newly produced membranes in eukaryotes is largely unknown. To our knowledge, only a few indirect observations are available. For instance, “Karmellae” (or stacks of membranes surrounding the nucleus) produced by overproduction of 3-Hydroxy-3-methylglutaryl-CoA reductase (Hmg-CoA) seem to be enriched in neutral lipids, e.g. sterols (Lum and Wright 1995). However, no lipid quantification or identification was performed and this observation is only derived from a preferential staining with Nile Red, which has low specificity and indiscriminately stains all cell membranes. As previously discussed, the expression of some viral proteins, (2BC from poliovirus, 36K from Carnation Italian ringspot virus or protein A from Flock House virus) also induce different types of membrane in yeast (Table 2) (Barco and Carrasco 1995; Rubino, Di Franco and Russo 2000; Miller et al. 2003). In these cases, the creation of sterol-rich microdomains are required to assist membrane-remodeling proteins to effectively modulate membrane curvature (Ivankin, Kuzmenko and Gidalevitz 2012; Krishna and Sengupta 2019). It is worth mentioning here the bacteriophage φ6, which induces a 9-fold increase in CL levels in infected bacteria (Mindich, Sinclair and Cohen 1976), reinforcing the importance of CL in intracellular membrane proliferation in prokaryotes.

2.4 Summary and final considerations about protein-induced membrane curvature
Most of the proteins triggering membrane proliferation are able to locally modulate the membrane curvature. Independently of the precise mechanism involved in the creation of highly curved membrane microdomains, some common features are found in many of the reviewed proteins:

- The production of the membrane-interacting domains alone is not sufficient to modulate the membrane curvature. Additional protein domains are often required, suggesting that the induction of highly curved membrane microdomains is not a general consequence of membrane protein overproduction and insertion into the intracellular membrane.

- In some cases, the membrane curvature depends on the concentration of the overproduced proteins. A certain threshold of overproduced protein is necessary to trigger membrane proliferation. Different levels of protein concentration can lead to different membrane morphologies in vivo.

- The membrane curvature induced by proteins is often accompanied by a production of non-bilayer forming phospholipids (e.g. CL). These phospholipids can assist in the creation and stabilization of curved membrane domains and favor some membrane morphologies.

The membrane morphologies observed in vivo (vesicles, tubules, flat stacks of membranes, etc.) depend on the balance between protein 3D structure of the recombinant protein, its concentration, and the phospholipid membrane composition. Recently, Bonazzi et al. attempted for the first time to theoretically model the membrane morphology produced by arc-shaped proteins (or protein assemblies) modifying membrane curvature (Bonazzi and Weikl 2019). This theoretical model not only predicts the dependence of membrane morphology on protein concentration but also the existence of a vesicle-to-tubule transition, observed for some proteins listed in Table 1 and Table 2. Furthermore, tubule-shaped membranes are formed independently of the shape of the protein when the concentration of proteins is high enough to cover more than 40% of the membrane surface area, which explains the prevalence of tubular membrane structures upon protein overexpression. This study strongly suggests that the mechanisms involved in membrane curvature induction and, as a consequence, the
morphology observed upon protein overproduction, are predictable and determined by physical laws.

3 Influence of membrane curvature on phospholipid biosynthesis in prokaryotic cells

The viability of bacteria directly depends on their ability to maintain membrane homeostasis and the electrochemical gradient in response to different environmental conditions. Lipid biosynthesis and modification are the most energy-intensive processes of membrane homeostasis, therefore it is not surprising that lipid metabolism is tightly regulated both transcriptionally and enzymatically (Cronan, Jr. and Rock 2008). Impressively, proteins discussed in this review have found a way to hijack this vital regulation, forcing bacteria to produce increased amounts of inner membranes.

Because fatty acid and phospholipid biosynthesis are coupled in *E.coli*, we will focus exclusively on the regulation of the biosynthesis of phospholipid polar heads, overlooking fatty acid metabolism (Cronan, Jr. and Rock 2008). Phospholipid metabolism and its regulation in prokaryotes (Figure 4) have been extensively studied (Parsons and Rock 2013; Sohlenkamp and Geiger 2015; Lin and Weibel 2016). Still, even in the archetypical *E.coli*, many pieces are lacking to construct a complete vision of lipid metabolism. One of these missing pieces is how (membrane) protein expression is synchronized with lipid biosynthesis. Protein synthesis stopped after inhibition of lipid synthesis and re-started when lipid metabolism was restored (Cronan, Jr. and Bell 1974; Bell and Cronan, Jr. 1975; McIntyre et al. 1977). Consequently, it appears that protein and lipid metabolisms are intimately connected and cross-regulated. However, the nature or mechanism of this regulation is unknown, even though some clues point to a regulation via multiple stress pathways (Seyfzadeh, Keener and Nomura 2006; Bury-Moné et al. 2009; Gopalkrishnan, Nicoloff and Ades 2014).

3.1 The importance of phospholipid homeostasis
*E. coli* maintains a constant ratio between zwitterionic phosphatidylethanolamine (PE), which accounts for about 75% wt. of total phospholipids, and anionic PG and CL, whose relative amounts depend on the physiological state (log- or stationary-phase) of the cells (Cronan, Jr. and Rock 2008). A feedback mechanism between the cross-regulated enzymes controlling the synthesis of PE and PG/CL (PssA and PgsA, respectively) maintains the homeostasis in phospholipid headgroup diversity (Figure 4). PssA is a monotopic membrane protein that acts as a sensor, detecting changes in relative phospholipid composition (PE vs PG/CL) in the lipid bilayer (Louie, Chen and Dowhan 1986; Satomi *et al.* 1996). It is active when associated with anionic phospholipids (PG and CL) and catalyzes the synthesis of PE. On the contrary, when anionic phospholipids become less available, PssA is deactivated, causing PgsA metabolic route to accelerate and to increase the synthesis of PG and CL.

Besides the aforementioned enzymatic regulation, phospholipid homeostasis is also subject to genetic control. Alterations in phospholipid composition stimulate the activation of several stress response pathways that ensure the maintenance of the bacterial envelope integrity (σE, Cpx, Bae and Rcs) (Figure 4) (Rowlett *et al.* 2017). Even though all these regulation pathways are entangled, the Cpx system is of particular interest for phospholipid homeostasis in *E. coli* (Bury-Moné *et al.* 2009). It activates the transcription of more than 100 genes, especially genes coding for inner membrane proteins and phospholipid metabolism, including Psd (an enzyme involved in the synthesis of PE, Figure 4) (Price and Raivio 2009; Raivio, Leblanc and Price 2013; Raivio 2014). Furthermore, the Cpx regulation pathway is controlled by CpxA, a transmembrane protein kinase located in the *E. coli* inner membrane, sensitive to modifications of the relative concentrations of anionic phospholipids in the lipid bilayer (Stenberg-Bruzell *et al.* 2015).

It seems clear that any alteration in the inner membrane phospholipid balance is compensated by the biosynthesis of the complementary type of phospholipids, either via enzymatic and/or genetic regulation. The enrichment in anionic phospholipids (especially CL) observed in most of the protein-triggered inner membrane proliferation (Table 1), most likely represent such an alteration. The constant biosynthesis of membrane proteins continuously alters phospholipid
homeostasis and pushes *E. coli* to produce new phospholipids. The enrichment for specific phospholipids could be explained either by a selective interaction of the overproduced protein with the phospholipid type (PE or PG/CL), or by the creation of membrane microdomains with impaired accessibility to the homeostasis membrane sensors.

3.2 Electrostatic interactions between anionic phospholipids and positively charged proteins

Overproduction of a membrane protein able to selectively bind anionic lipids, such as CL, could create a phospholipid imbalance in the inner cell membrane. Evidences supporting this
hypothesis have been obtained in the case of the monotopic glucosyltransferase alMGS (Eriksson et al. 2009). The alMGS domains that interact with phospholipids are enriched in positively charged amino acids (Ge et al. 2014), suggesting that they may selectively bind anionic phospholipid through electrostatic interactions. Indeed, the capacity of alMGS to alter phospholipid metabolism through its capacity to bind anionic phospholipids was later confirmed (Ariöz et al. 2014). In addition, when overproduced, alMGS displaces the cyclopropane fatty acid synthase (CFA synthase) from its membrane binding site, inhibiting the synthesis of cyclopropanated fatty acids. Finally, phospholipid biosynthesis and alMGS overproduction are linked, as the supplementation of the culture media with lipid metabolism precursors leads to increased amounts of inner membranes with a concomitant increase in alMGS production (Ariöz et al. 2014). Finally, alMGS overproduction also activates the σE and Cpx envelope stress responses, which can activate phospholipid metabolism as discussed in section 3.1. Besides alMGS, all of the monotopic membrane proteins listed in Table 1 (MurG, LpxB, caveolin-1 and PmtA) are known to bind to the lipid bilayer via electrostatic interactions with anionic phospholipids (Van Den Brink-van Der Laan et al. 2003; Metzger IV and Raetz 2009; Walser et al. 2012; Linna et al. 2017). Electrostatic interactions with transmembrane proteins have been less studied. Nevertheless, the activity of many respiratory complexes often depends on CL concentration (Arias-Cartín et al. 2012). Thus, it is not surprising that CL has been found to selectively interact with succinate dehydrogenase and F_{0}F_{1} ATP-synthase (Yankovskaya et al. 2003; Laage, Tao and McDermott 2015). Indeed, it was recently demonstrated that the mitochondrial F_{0}F_{1} ATP-synthase possesses a CL interaction site, conserved between yeast and cattle, which is enriched in positively charged amino acids (Mehdipour and Hummer 2016; Duncan et al. 2018).

The presence of positively charged residues flanking transmembrane domains in the cytosolic leaflet of E.coli inner membrane is not, however, an exclusive feature of proteins triggering inner membrane proliferation. In fact, the “positive charges inside” rule (von Heijne 1989), is a highly conserved mechanism to control the orientation of membrane proteins in the inner membrane (Elazar et al. 2016). This “positive charges inside” rule is conserved in most membrane proteins, not only in E. coli, but also in other prokaryotes and eukaryotes (Baker et
al. 2017), while intracellular membrane proliferation has only been observed with a dozen of specific membrane proteins. To determine whether proteins able to induce membrane proliferation display specific charges distribution, we calculated the surface electrostatic potential of the proteins listed in Table 1 for which a structure is available in the PDB. Those proteins present a positive electrostatic lobe located nearby the phospholipid polar heads in the cytosolic leaflet of the inner membrane (Figure 5a). However, this feature is far from being limited to the proteins inducing membrane proliferation. Indeed, MsbA and G3P transporter, two proteins that do not trigger intracellular membrane proliferation (Gubellini et al. 2011), also exhibit a marked positive electrostatic lobe near the cytosolic leaflet of the inner membrane (Figure 5b). In conclusion, this positive electrostatic lobe is probably useful for membrane association and/or insertion in the correct orientation, but it is not sufficient per se to trigger inner membrane proliferation.

Figure 5. Isopotential electrostatic surfaces calculated for proteins available in the PDB. Charges were calculated at pH 7.5 with PDB2PQR server http://nbcr-222.ucsd.edu/pdb2pqr_2.0.0/ using Amber force field and naming schemes. In blue, positive isosurfaces at +10 kT/e - in red, negative isosurfaces at -10 kT/e - a) Proteins triggering membrane proliferation when overproduced (monotopic MurG, PDB code 1F0K),(Ha et al. 2009)
and transmembrane Fumarate reductase, PDB code 6AWF (Starbird et al. 2018); b) proteins whose overproduction does not trigger membrane proliferation (MsbA, PDB code 6BPL (Mi et al. 2017; Oh et al. 2018) and G3P transporter, PDB code 5XJ9 (Huang et al. 2003)).

3.3 Formation of CL microdomains

An alternative explanation for the perturbation of phospholipid homeostasis in a reduced accessibility of certain type of phospholipids to the membrane homeostasis sensors, that could be a consequence of lipid microdomains formation. Lipid rafts enriched in sterols, sphingolipid and some proteins are a well-established membrane organization element in eukaryotic membranes (Sezgin et al. 2017). The existence of membrane lipid microdomains, analogous to the lipid rafts of eukaryotes, has also been reported in prokaryotes (Bramkamp and Lopez 2015). These membrane microdomains associate with specific membrane protein complexes and are enriched in particular lipids (notably CL) (Mileykovskaya and Dowhan 2009; Lin and Weibel 2016). Moreover, the spontaneous breaking of membrane symmetry with subsequent enrichment in CL in areas of high negative curvature has been theoretically modeled and also experimentally observed in E.coli membranes (Renner and Weibel 2011; Boyd, Alder and May 2017, 2018). Very recently, this CL clustering and enrichment of highly curved membrane areas has been experimentally quantified (Beltrán-Heredia et al. 2019). It should be noted that in both theoretical and experimental designs an external force is necessary to impose high membrane curvature, as a mere accumulation of CL alone is not sufficient to curve the membrane enough to induce further CL clustering.

In this regard, the insertion of a membrane protein (or supramolecular complex of proteins) inducing a high local curvature would be needed to provide the necessary force to bend the membrane and induce CL clustering. This could in turn lead to an anionic phospholipids depletion in the non-curved zones of the membrane, which will subsequently be detected by the phospholipid homeostasis sensors as a signal to start lipid biosynthesis. Then, the newly synthesized lipid membranes would allow for more membrane protein insertion, thus, closing the cycle (Figure 6). As previously discussed in section 2, all the proteins triggering inner
membrane proliferation have the capacity to modify the membrane curvature and, hence, to induce CL clustering. Of note, this hypothesis can explain not only the production of inner membranes but also the enrichment in CL (instead of anionic PG that would be expected in the case of the electrostatic interactions hypothesis) observed in most examples of intracellular membrane proliferation upon protein overproduction. Furthermore, it could be the missing link to the ancestral mechanism of inner membrane compartments observed in α-proteobacteria and mitochondria, where both the insertion of membrane curvature-inducing protein complexes and CL clustering are required.

Figure 6. Proposed mechanism for phospholipid biosynthesis triggered by membrane protein overproduction.
Table 1. Proteins triggering membrane proliferation in prokaryotes

<table>
<thead>
<tr>
<th>Protein</th>
<th>Organism</th>
<th>Type</th>
<th>Origin</th>
<th>Expressed</th>
<th>Morphology</th>
<th>Protein arrangement</th>
<th>Lipid composition</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fumarate Reductase</td>
<td>E. coli</td>
<td>TM</td>
<td>E. coli</td>
<td>Tubules</td>
<td>Helical (10 units per turn)</td>
<td>CL enrichment (~15% mol)</td>
<td>(Weiner et al. 1984; Elmes, Scraba and Weiner 1986)</td>
<td></td>
</tr>
<tr>
<td>Succinate Dehydrogenase</td>
<td>E. coli</td>
<td>TM</td>
<td>E. coli</td>
<td>Tubules and Vesicles</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>(Maklashina, Berthold and Cecchini 1998)</td>
</tr>
<tr>
<td>P6 (sp6.6 gene product)</td>
<td>Phage PM2</td>
<td>TM</td>
<td>E. coli</td>
<td>Vesicles</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>(Johnson and Mindich 1994; Sarin et al. 2012; Myhrvold, Polka and Silver 2016; Lyytinen, Starkova and Poranen 2019)</td>
</tr>
<tr>
<td>P9 and P12</td>
<td>Phage ϕ6</td>
<td>TM</td>
<td>P. syringae E. coli</td>
<td>Vesicles</td>
<td>-</td>
<td>CL enrichment (~10% mol in P.pseudoalcaligenes infected by phage ϕ6)</td>
<td>-</td>
<td>(Armour and Brewer 1990)</td>
</tr>
<tr>
<td>AlkB (alkane oxidation system)</td>
<td>P. oleovorans</td>
<td>TM</td>
<td>E. coli</td>
<td>Vesicles</td>
<td>-</td>
<td>CL enrichment (not quantified)</td>
<td>-</td>
<td>(Nieboer, Vis and Witholt 1996)</td>
</tr>
<tr>
<td>Tsr chemotaxis receptor</td>
<td>E. coli</td>
<td>TM</td>
<td>E. coli</td>
<td>Inner membrane invaginations</td>
<td>Crystalline pseudohexagonal (3 dimers of Tsr per repeating unit).</td>
<td>-</td>
<td>(Lefman et al. 2004; Zhang et al. 2007)</td>
<td></td>
</tr>
<tr>
<td>(b subunit of) F0F1 ATP synthase</td>
<td>E. coli</td>
<td>TM</td>
<td>E. coli</td>
<td>Tubules and vesicles</td>
<td>Unknown. Dimer formation.</td>
<td>CL enrichment (~24% mol)</td>
<td>-</td>
<td>(von Meyenburg, Jogersen and van Deurs 1984; Miroux and Walker 1996; Arechaga et al. 2000; Carranza et al. 2017)</td>
</tr>
<tr>
<td>P3A</td>
<td>FMDV</td>
<td>M</td>
<td>E. coli</td>
<td>Onion-like vesicles</td>
<td>Unknown. Possible oligomerization?</td>
<td>-</td>
<td>-</td>
<td>(Weber et al. 1996)</td>
</tr>
<tr>
<td>Caveolin-1</td>
<td>H. sapiens</td>
<td>M</td>
<td>E. coli</td>
<td>Vesicles</td>
<td>Well-defined supramolecular cage (160 monomers per cage)</td>
<td>PG and lysophospholipids enrichment (not quantified)</td>
<td>(Walser et al. 2012; Ariotti et al. 2015)</td>
<td></td>
</tr>
<tr>
<td>MurG</td>
<td>E. coli</td>
<td>M</td>
<td>E. coli</td>
<td>Vesicles</td>
<td>-</td>
<td>CL enrichment (~22% mol)</td>
<td>-</td>
<td>(Van Den Brink-van Der Laan et al. 2003)</td>
</tr>
<tr>
<td>LpxB</td>
<td>E. coli and H. influenzae</td>
<td>M</td>
<td>E. coli</td>
<td>Tubules and vesicles</td>
<td>-</td>
<td>PG and CL enrichment (not quantified)</td>
<td>-</td>
<td>(Metzger IV and Raetz 2009)</td>
</tr>
<tr>
<td>PmtA</td>
<td>A. tumefacensis</td>
<td>M</td>
<td>A. Tumefacensis</td>
<td>Vesicles</td>
<td>-</td>
<td>CL needed for membrane proliferation (not quantified)</td>
<td>-</td>
<td>(Linna et al. 2017)</td>
</tr>
<tr>
<td>alMGS</td>
<td>A. laidlewii</td>
<td>M</td>
<td>E. coli</td>
<td>Vesicles</td>
<td>-</td>
<td>PG and CL enrichment (not quantified)</td>
<td>-</td>
<td>(Eriksson et al. 2009; Anibz et al. 2013, 2014; Ge et al. 2014)</td>
</tr>
</tbody>
</table>

TM: Transmembrane protein; M: monotypic protein; FMDV: Foot and Mouth Disease Virus
4. Influence of membrane curvature on phospholipid biosynthesis in eukaryotic cells

Unicellular eukaryotes (especially yeasts) are, like prokaryotes, well-known hosts used for protein production and biotechnological applications. As in prokaryotes, intracellular membrane proliferation can also be induced by some membrane proteins overexpression in unicellular eukaryotes (mostly documented in *S. cerevisiae* and *P. pastoris*). To date, only transmembrane (not monotopic) proteins have been reported to trigger membrane proliferation in yeasts (Table 2). The first reports of membrane proliferation in eukaryotes described nuclear membrane structures (*Karmellae*), but since then, some have been shown to also originate from other cell organelles such as the endoplasmatic Reticulum (ER), mitochondria, peroxisome or vacuole (Table 2).

Membrane proliferation can now be specifically targeted to particular cellular organelles by engineering chimera proteins containing *ad hoc* signal peptides (Miller *et al.* 2003).

4.1 The Unfolded Protein Response

The relationship between expression levels of recombinant membrane proteins and phospholipid biosynthesis has been studied more in depth in eukaryotes than in prokaryotes. Early reports on the activation of the inositol response pathway and the Unfolded Protein Response (UPR) in *S. cerevisiae* overproducing Hmg1 were the first attempts to link membrane proliferation to protein overproduction (Cox, Chapman and Walter 1997). In yeast, UPR is controlled by Ire1, an integral ER membrane protein with an ER luminal domain sensitive to the presence of misfolded proteins (*via* the release of the chaperone Kar2, also known as BiP) and a transmembrane and proximal domain sensing alterations of lipid composition in the ER membrane (Volmer and Ron 2015; Ho, Xu and Thibault 2018). Either an accumulation of unfolded proteins or an imbalance of lipid homeostasis constitute life-threatening events, which are tightly controlled by Ire1 (Figure 7) (Fun and Thibault 2019). Activation of Ire1 triggers the production of Hac1, a transcription factor that controls ca. 381 genes, including those of phospholipid biosynthesis, ER-associated protein degradation, protein translocation across ER membrane, vesicular trafficking, cell wall biogenesis, and vacuolar protein sorting (Travers *et al.* 2000). It was then postulated that the overproduction of a membrane protein is a perturbation that could activate UPR response and, indirectly phospholipid biosynthesis through Ire1 sensor. Ire1 knock-out yeasts (*S. cerevisiae Δire1*) were constructed to test this hypothesis. However, membrane proliferation upon overproduction of P450 cytochrome or Pex15p was not impeded in *Δire1* yeasts (Menzel *et al.* 1997; Stroobants *et al.* 1999). Interestingly, *Δire1* yeasts were still able to produce Kar2/BiP, which is a part of the UPR response. Thus, the authors postulated an alternative (and unknown) Ire1-independent mechanism for UPR activation. Later, these
observations were expanded to HMG-CoA reductase isozyme (Hmg1) overproduced in \( \Delta \text{ire1} \) yeasts. In this case, membrane proliferation was achieved in complete independence from Ire1 and secretion of Kar2/Bip chaperone. Consequently, the authors concluded that, at least for Hmg1, membrane proliferation phenomena should be unrelated to UPR. However, the activation of UPR pathway seems somewhat advantageous for inner membrane proliferation, as the overproduction of the transcription factor Hac1 (the main product of UPR), using an external expression plasmid improved the production yield of membrane proteins and intracellular membranes (Guerfal \textit{et al.} 2010; Vogl \textit{et al.} 2014). Interestingly, overproduction of Hac1 alone changes the morphology of the ER membrane to a cubic phase and increases the Kar2/BiP chaperon levels (Guerfal \textit{et al.} 2010). As previously discussed, bi-continuous cubic phase ER is often observed during the infection +RNA virus (van der Schaar \textit{et al.} 2016), which are known to induce membrane proliferation in both prokaryotes and eukaryotes. Additionally, stacks of \textit{Karmellae}-like membranes surrounding the yeast nucleus are produced upon coexpression of Hac1 with a membrane protein, which alone would not trigger membrane proliferation (Vogl \textit{et al.} 2014).

Figure 7. UPR phospholipid sensor. Ire1 is able to detect the presence of misfolded proteins or alterations of phospholipid composition in the membrane of the ER and it is activated by a dimerization and autophosphorylation. This activation leads to an unconventional splicing of \textit{HAC1} mRNA and the translation of Hac1, which activates gene expression and phospholipid biosynthesis.

4.2 Inositol regulation pathway and the importance of phosphatidic acid
How to reconcile the fact that membrane proliferation in yeasts is at the same time influenced by and independent of UPR? A plausible explanation is the existence of redundant sensors capable of activating, independently of UPR, the biosynthesis of phospholipids. PI and its precursor, inositol, are implicated in the inositol regulation pathway (INO1 genes), which is activated during membrane proliferation after canine RRb production in yeast cells (Block-Alper et al. 2002). INO1 promotes the synthesis of phospholipids (and many other inositol-sensitive genes) and is regulated independently of UPR by the couple of activators Ino2/Ino4 and repressed by Opi1 (Figure 8) (Henry, Gaspar and Jesch 2014). This regulation system is clearly related to membrane proliferation, as mutants deficient of Ino2 failed to produce intracellular membranes upon RRb expression, whereas deletion of the repressor Opi1 favors the proliferation of inner membranes (Block-Alper et al. 2002). All the genes under inositol regulation (including INO1) are, in fact, controlled by phosphatidic acid (PA) levels, which is a central regulator of both the synthesis of phospholipids and reserve lipids (triacylglycerides) in S. cerevisiae (Carman and Henry 2007). Opi1 is a soluble protein associated to a transmembrane protein (Scs2), which acts as a membrane sensor for PA microdomains (Loewen et al. 2004; Young et al. 2010). When PA concentration is sufficient, Opi1 remains inactive and anchored to Scs2 in the ER membrane. On the contrary, when PA concentration drops down, Opi1 is released and imported to the nucleus where it represses the phospholipid synthesis.

![Figure 8](image_url)

Figure 8. Inositol dependent phospholipid sensor. When PA levels are high (i.e. low inositol concentration), Opi1 interacts with Scs2 and is sequestrated in the ER membrane (left) and INO1 and other inositol-sensitive genes are expressed. When the PA levels drops down, Opi1 detaches from Scs2 and migrates to the nucleus where it represses the expression of inositol-sensitive genes (right).

4.3 Other regulatory mechanisms

In addition, the regulation of phospholipid homeostasis in S. cerevisiae is controlled by multiple biochemical and genetic factors and goes well beyond UPR and inositol regulation pathways (Carman and Han 2011).
Therefore, the existence of additional regulatory pathways to control phospholipid biosynthesis upon protein overproduction cannot be ruled out. In fact, intracellular membrane proliferation of ER in *S. cerevisiae* is associated with altered membrane trafficking. For example, overproduction of Sec12p blocks the ER-to-Golgi intracellular trafficking of *S. cerevisiae* and induces the formation of clusters of the chaperone Kar2/BiP, like in UPR pathway (Nishikawa, Hirata and Nakano 1994). Similar blockage of intracellular trafficking was also observed after overexpression of the poliovirus 2BC protein and the peroxisomal Pex15p protein, which both accumulate newly formed ER membranes (Barco and Carrasco 1995; Elgersma *et al.* 1997). Conversely, the overproduction of the canine RRp enhances the secretory pathways in *S. cerevisiae* (Becker *et al.* 1999). Of note, this altered intracellular trafficking is not a general scenario for all the protein-induced intracellular membrane proliferation in eukaryotes, as exemplified with Hmg1 (Nishikawa, Hirata and Nakano 1994). It is not clear whether these alterations of cellular trafficking are only a consequence of membrane proliferation or, on the contrary, contribute to inner membrane proliferation in eukaryotes. In any case, regardless of the precise mechanism, the biosynthesis of phospholipids also seems to be controlled by the membrane composition in eukaryotes.

4.4 Cardiolipin and phosphatidic acid membrane microdomains: a universal regulator mechanism for phospholipid biosynthesis conserved through evolution?

PA seems to be a central actor in phospholipid biosynthesis regulation in eukaryotes (Carman and Han 2011). For instance, increased PA cellular levels, either by a lack of PA degradation due to lower PA phosphatase activity or by an increase in PA concentration due to an overproduction of diacylglycerol kinases, leads to an expansion of the nuclear membrane (Santos-Rosa *et al.* 2005; Han, Siniossoglou and Carman 2007; Han *et al.* 2008). It should be noted that this membrane expansion in the absence of any membrane protein modulating membrane curvature leads to yeasts with an aberrantly large nucleus, without any organized membrane morphology (*e.g.* stacks of membranes, tubules or vesicles).

From a physicochemical point of view, the phospholipids found at the core of membrane proliferation in eukaryotes and prokaryotes (PA and CL), are strikingly similar (Figure 9) (De Kroon, Rijken and De Smet 2013). Both PA and CL are cone-shaped anionic phospholipids that accumulate in zones of negative curvature in biological membranes. Furthermore, both of them have two ionizable positions (Figure 9), determined by two distinct pKₐ values (Kooijman and Burger 2009; Sathappa and Alder 2016). Their exact values and hence, the ionization state of PA and CL, depend on the membrane local environment. Indeed, different ionization
states of PA controlled by lipid-protein interactions have been proposed to explain the regulatory role of PA in eukaryotes (Kooijman and Burger 2009). Similarly, CL undergoes changes in lipid packing as a function of its environment, external pH and divalent cations (Khalifat *et al.* 2011; Kensbock, Ahrens and Helm 2019).

As discussed before, proteins inducing membrane proliferation in both prokaryotic and eukaryotic cells are characterized by their ability to bend the membrane. At the same time, cone-shaped anionic PA and CL share a marked preference for negatively curved regions of the membrane. Therefore, the disturbance of phospholipid homeostasis via the creation of membrane microdomains enriched in PA or CL might be a central regulator of phospholipid metabolism conserved through evolution.

**5. Membrane overproduction as a platform for biotechnological applications**

Beyond the exotic observation of inner membrane proliferation in prokaryotes and eukaryotes, the phenomena of inducible membranes have interesting implications for biotechnology. The increased phospholipid amount due to membrane expansion could be useful in the field of biofuel production by fermentation. Modification of metabolic pathways aiming at diverting carbon fluxes towards the desired target compound has been tried and
is far from being straightforward (Hollinshead, He and Tang 2014; Zhou et al. 2016). In this context, the overexpression of a protein triggering membrane proliferation could represent a simple, alternative strategy to redirect lipid metabolism and enhance biofuel production yield. In the same line, production of the P9 and P12 viral proteins have been proposed to increase the yield of useful hydrophobic active principles (Myhrvold, Polka and Silver 2016).

For structural biologists, membrane protein production still represents a major biochemical challenge (Zoonens and Miroux 2010). Despite the emergence of eukaryotic expression systems, microbial expression systems are the most popular vehicle for membrane protein production (Dilworth et al. 2018). Historically, genetically modified strains have been developed to enhance membrane protein production (Miroux and Walker 1996; Baumgarten et al. 2017; Angius et al. 2018). Recently, a novel strategy has arisen for membrane protein production both in prokaryotes and eukaryotes, relying on tuning the cell membrane phospholipid composition to accommodate higher amounts of recombinant membrane proteins (Pichler and Emmerstorfer-Augustin 2018; Kanonenberg et al. 2019). Some of the proteins triggering internal membrane proliferation, like the P9 protein of phage φ6, have been proposed as a fusion partner to membrane proteins to trigger membrane expansion and increase the yield of membrane protein production (Jung, Jung and Lim 2015). The next generation of membrane protein production platforms may combine those three strategies: genetic regulation of protein expression, modulation of phospholipid composition, and membrane expansion triggered by protein overproduction.

In addition, these production platforms could find applications in nanotechnology. Almost all cell types secrete nano- and micro-sized vesicles used for intercellular communication (Raposo and Stahl 2019). Granting control over the production and composition of those vesicles hold great promises in nanotechnology and nanomedicine (Busatto et al. 2020). In this regard, the protein-induced intracellular membrane proliferation could increase the vesicle production yield. For example, membrane proliferation upon overproduction of the b subunit of F₈F₁-ATP synthase has been recently used to prepare proteoliposomes (Royes et al. 2019). This method represents an attractive alternative to in vitro proteoliposomes reconstitution, alleviating several steps of protein extraction, purification and reconstitution in liposomes. In the same line, the preparation of bacteria-derived lipid vesicles presenting antigenic proteins from pathogens on their surface have been used for vaccine preparation (Farjadion et al. 2018). The production of chimera proteins containing a membrane-proliferation domain and an adequate antigen could dramatically improve vaccine safety and mass production. In any case, the mechanistic considerations revealed in this review can help to better understand membrane proliferation.
upon membrane protein overproduction and harness this phenomenon to inspire and design new biotechnological applications.

6. Conclusions and perspectives

Despite the variety of inducible membranes and host cells reported in the literature, the proteins triggering membrane proliferation share some common properties. They are able to modify membrane curvature and hijack the regulation of phospholipid synthesis, where anionic, non-bilayer-forming phospholipids (CL and PA) seem to play a central role. We reviewed the mechanisms involved in phospholipid homeostasis, identifying their possible coordination with membrane protein biosynthesis. The hypothesis that phospholipid production might be induced by electrostatic interactions between anionic phospholipids and a positively charged lobe of the protein has been ruled out. Taking into account all the reviewed information, we propose a general mechanism of intracellular membrane proliferation in which the overproduced recombinant proteins can induce high curvature local areas, creating clusters of anionic, cone-shaped phospholipids (CL in prokaryotes and PA in eukaryotes). The accumulation of these phospholipids in the curved microdomains causes a phospholipid imbalance in the non-curved areas of the cell membrane. This local imbalance is likely to be detected by the phosphate homeostasis sensors, which in turn, will stimulate phospholipid biosynthesis. Upon the continuous accumulation of the recombinant protein (or their supramolecular assemblies), new curved microdomains are produced, and the cell is forced to constantly synthesize new phospholipids to maintain phospholipid homeostasis. The final consequence is an expansion of the intracellular membranes displaying the different morphologies observed in cellulo. This general mechanism of intracellular membrane proliferation shares common features with the formation of energetic compartments in α-proteobacteria (and mitochondria), and for some viruses, which remodel the inner membrane structures of the host to create their replication organelles. In both cases, the membrane expansion is triggered by the insertion of proteins that modulate membrane curvature, causing the accumulation of non-bilayer forming lipids (CL or PA). In summary, the modification of membrane curvature can act in cellulo as an inducer of phospholipid biosynthesis and membrane expansion and it is probably a conserved feature through evolution.
7. Acknowledgements

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Table 2. Proteins triggering membrane proliferation in eukaryotes

<table>
<thead>
<tr>
<th>Organism</th>
<th>Protein</th>
<th>Origin</th>
<th>Expressed</th>
<th>Morphology</th>
<th>Target organelle</th>
<th>Observations</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>HMG-CoA</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>“karmellae” stacked membranes around nucleus</td>
<td>ER</td>
<td>Soluble domain not required. Transmembrane helix alone not sufficient.</td>
<td>(Wright et al. 1988; Lum and Wright 1995; Profant et al. 1999)</td>
</tr>
<tr>
<td>Cytochrome b5</td>
<td>R. norvegicus</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>“karmellae” stacked membranes around nucleus</td>
<td>ER</td>
<td>Transmembrane domain disturbed by proline hinders membrane proliferation</td>
<td>(Vergères et al. 1993)</td>
</tr>
<tr>
<td>PMA2 (H+ ATPase)</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td></td>
<td>Tubules</td>
<td>ER</td>
<td>-</td>
<td>(Supply et al. 1993)</td>
</tr>
<tr>
<td>RRp - 180kDa</td>
<td>C. lupus</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>“karmellae” stacked membranes around nucleus</td>
<td>ER</td>
<td>RBS not required for membrane proliferation. Increase of secretory pathway.</td>
<td>(Wanker et al. 1995; Becker et al. 1999)</td>
</tr>
<tr>
<td>Dβ2 receptor</td>
<td>H. sapiens</td>
<td>P. pastoris</td>
<td></td>
<td>“karmellae” stacked membranes around nucleus</td>
<td>ER</td>
<td>-</td>
<td>(Grünewald et al. 2004)</td>
</tr>
<tr>
<td>sk2 Channel</td>
<td>H. sapiens</td>
<td>P. pastoris</td>
<td></td>
<td>“karmellae” stacked membranes around nucleus and ER</td>
<td>ER</td>
<td>-</td>
<td>(Licata et al. 2006)</td>
</tr>
<tr>
<td>B2 receptor</td>
<td>H. sapiens</td>
<td>P. pastoris</td>
<td></td>
<td>“karmellae” stacked membranes around nucleus</td>
<td>ER</td>
<td>-</td>
<td>(Song et al. 2015)</td>
</tr>
<tr>
<td>LaminB receptor</td>
<td>G. domesticus</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>“karmellae” stacked membranes around nucleus</td>
<td>ER</td>
<td>-</td>
<td>(Smith and Blobel 1994)</td>
</tr>
<tr>
<td>Pex12p</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td></td>
<td>Multilayered membranes</td>
<td>Peroxisome</td>
<td>Observed morphology is dependent on expression level</td>
<td>(Albertini et al. 2001)</td>
</tr>
<tr>
<td>Pex15p</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td></td>
<td>Multilayered membranes</td>
<td>Peroxisome, ER</td>
<td>ER to peroxisome transport blocked</td>
<td>(Egersma et al. 1997)</td>
</tr>
<tr>
<td>2BC</td>
<td>Poliovirus</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>Vesicles</td>
<td>Vacuole</td>
<td>ER transport blocked</td>
<td>(Barco and Carrasco 1995)</td>
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<tr>
<td>36k protein IRV</td>
<td>Carnation Italian Ringspot Virus</td>
<td>S. cerevisiae</td>
<td></td>
<td>Vesicles</td>
<td>Mitochondria</td>
<td>-</td>
<td>(Rubino, Di Franco and Russo 2000)</td>
</tr>
<tr>
<td>Protein A</td>
<td>Flock House Virus</td>
<td>S. cerevisiae</td>
<td>S. cerevisiae</td>
<td>Vesicles</td>
<td>Mitochondria</td>
<td>Retargeting of protein A to ER possible with ER specific sequence</td>
<td>(Miller et al. 2003)</td>
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</table>

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