Mini review

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The disordered boundary of the cell: emerging properties of membrane-bound intrinsically disordered proteins

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Abstract: We define the disordered boundary of the cell (DBC) as the system formed by membrane tethered intrinsically disordered protein regions, dynamically coupled to the underlying membrane.

The emerging properties of the DBC makes it a global system of study, which cannot be understood from the individual properties of their components. Similarly, the properties of lipid bilayers cannot be understood from just the sum of the properties of individual lipid molecules.

The highly anisotropic confined environment, restricting the position and orientation of interacting sites, is affecting the properties of individual disordered proteins. In fact, the collective effect caused by high concentrations of disordered proteins extend beyond the sum of individual effects.

Examples of emerging properties of the DBC include enhanced protein-protein interactions, protein-driven phase separations, Z-compartmentalization, and protein modulated electrostatics.

Keywords: Intrinsically disordered proteins; Membrane proteins; Cell Boundaries; Signalling environment; Bio nano composites; Crowding; Liquid phase separation; Z-compartmentalization; Src family kinases.

Introduction

Compartmentalization is an essential feature of cells and life that allows spatiotemporal control of processes. Lipid membranes define the boundary of the cell and cellular compartments. Within the cells, compartments may also be generated in phase-separated regions, like the lipid driven raft-like structures in membranes, or in the liquid protein or protein/RNA droplets. Information is transmitted across compartments through dynamic protein interaction networks, which acquire, amplify, transmit, and eventually respond to signals that are channelled along the signaling cascades.

Regarding the protein interaction networks, standard paradigms assume that: (i) the functionally relevant regions of proteins are folded domains, (ii) protein complexes adopt well-defined 3D structures, and (iii) interaction networks obey an isotropic and unrestricted reaction-diffusion model. Each of these paradigms has been challenged by recent findings. First, a large proportion of eukaryotic proteins, ranging from one [1] to two thirds [2], depending on the criteria used, are predicted to be intrinsically disordered proteins (IDPs). Interestingly, IDPs are far less abundant in prokaryotic organisms, suggesting that some emerging properties associated with extensive protein flexibility may have enabled the jump in complexity from bacteria and archaea to multi-cellular organisms [3]. Secondly, some complexes formed by two or more IDPs remain disordered [4] and some IDPs participate in liquid-like phases that demix from homogeneous solutions [5]. Also, a network of IDP interactions contributes to the selectivity of nuclear pores [6] or cell-to-cell channels [7]. Finally, essential signaling interactions communicating the cell with its environment take place in the highly anisotropic vicinity of lipid membranes [8].

The region extending into the cytoplasm from membrane-associated proteins is especially rich in intrinsically disordered regions (IDR). Iakoucheva et al. reported that 70% of transmembrane proteins involved
in signaling have IDRs longer than 30 residues [9]. The enrichment of IDRs in the proximity of lipid membranes might be just the consequence of the association of IDRs with the needed signaling at the cell interface. We suggest that there is a boundary layer enriched with disordered domains coupled to the lipid bilayer that provides a singular environment, with properties distinct from the bulk cytoplasm or an isolated membrane model. We name this region, which includes the lipid membrane and its boundary layer, as the disordered boundary of the cell (DBC). We refer to the IDP-rich region of the cytoplasm in direct contact with the lipid bilayer, as the ε-layer (Figure 1). In this perspective article, we want to discuss the emergent properties expected for the ε-layer and their possible implications in cell signaling. While disordered regions appear more frequently in the cytoplasmic side of the membrane, some of the concepts are equally applicable to the other side of the membrane, forming the boundary with the extracellular matrix.

Cell membrane and boundary regions

Membrane models emphasize the lipid-dominated hydrophobic environment with embedded proteins. Models have been updated regularly since the original “fluid mosaic” concept of Singer and Nicholson [10]. Engelman emphasized the crowded behaviour and heterogeneity of membrane protein composition [11]. Successive studies shed light on the structural complexity and continuous dynamics undergoing in cellular boundaries as lipid phases, lipid-protein interactions, macromolecular polymer adsorption, membrane curvature, and flip-flop lipid motions, among others nicely reviewed by Goñi et al. [12].

Membrane attached proteins, cytoplasmic regions of transmembrane proteins, or cytoplasmic proteins adsorbed on the membrane surface play important roles in coupling the membrane to the bulk cytoplasm. The interactions of the cytoplasmic regions of membrane embedded proteins and the underlying actin cytoskeleton may restrict lipid diffusion and create membrane compartments [13]. Abundant long coiled-coil proteins, called golgins, attached to the golgi have been suggested to create a tentacular matrix helping to recruit Rab-coated vesicles [14]. The emphasis of this review is, however, on the expected emergent properties when the membrane associated protein chains are disordered and present at high local concentrations.

The aqueous layer in contact with the hydrophobic membrane is a boundary layer. In purely physical terms, the potential of boundary layers to generate emergent properties is well-known. However, the ε-layer is predicted to have unique features that arise from its “composite” nature. Composites are materials combining different constituents with very distinct individual properties, that when mixed together, render a material with novel properties [15]. Well-known examples in other fields are reinforced concrete or animal bones, in which metal bars or collagen fibers modify a homogeneous matrix of cement or apatite, respectively. In contrast to these macroscopic examples, the ε-layer is not a rigid material, but has the properties of a fluid. The IDPs act as polymers anchored to the fluid lipid membrane that fill-in and modulate the properties in its immediate vicinity. The matrix would be the surrounding solution experiencing strong electrostatic and hydrodynamic fields.

In an isolated lipid membrane, the electric and hydrodynamic fields would show smooth continuous distributions, with gradients originating from the proximity of the charged bounding layer. However, the presence of abundant charged residues, a characteristic feature of the IDP sequences, introduce discontinuities in the ε-layer. The position of these charges is determined by their interaction with the electric field generated on the lipid surface or originate from neighboring charged residues. Additionally, the location of the perturbing charges is modulated by constraints on the IDP network. Considering that IDPs have nanoscopic dimensions and are dynamically coupled to their environment, the DBC can be described as a fluid nano-composite which emphasizes the essential roles of its small scale and complex dynamics.

Abundance of intrinsically disordered proteins in membranes

The abundance of IDRs is a hallmark of eukaryotes and is functionally associated to their higher regulatory complexity as compared to prokaryotic organisms. IDRs are more abundant in the subset of transmembrane human proteins than in the complete proteome (57.4% versus 35.2% defined as the percentage of proteins having consecutive regions longer than 30 residues predicted to be disordered by DISOPRED2 [16, 17]). This implies that about ~65% of the IDPs are transmembrane proteins. The cytoplasmic side of membrane proteins is particularly enriched in IDRs [18].
Non-integral cytoplasmic proteins anchored to the membrane by long intrinsically disordered regions terminated by covalently bound lipid chains, such as a myristoyl or a palmitoyl groups, represent an additional source of IDPs in the $\varepsilon$-layer. Prominent examples are MARCKS (myristoylated alanine rich C kinase substrate), the alpha subunits of G proteins [19], and the members of the Src family of kinases (SFKs) [20]. The combined abundance per cell of these IDP families is 15% of that of actin, the major component of the cytoskeleton [21].

The abundance of IDPs that interact in a permanent or reversible fashion with the membrane surface, unveils a scenario with a high density of disordered proteins confined in a narrow perimembranal layer. Such high density would result in emergent properties, i.e. beyond the simple juxtaposition of the individual components. The critical concentration at which emerging properties would appear will depend on the nature of the actual IDPs and the underlying lipid composition. For instance, simulations on the phase separation and gelation of multivalent proteins with disordered linkers emphasize as relevant determinants: the fraction of charged residues in the linkers that affects the effective solvation volume, the average spatial distance between residues, and the cooperativity of the interactions between individual interacting regions [22].

Less abundant but still highly relevant, are those proteins with exposed disordered segments in the extracellular part of lipid bilayer, specially enriched in endocytic processes [23] and cell-matrix or cell-cell interactions [24]. Although there is a small number of proteins with long extracellular disordered regions (more than 200 residues), these are usually highly glycosylated cell adhesion molecules, also called proteoglycans. The CD44 antigen is a prototypical example, containing a variable disordered stem region that expands from 80 to
almost 500 residues depending on alternative splicing [25, 26]. The cytoplasmic and extracellular loops connecting the transmembrane helices of integral polytopic proteins are often disordered and play crucial roles in substrate recognition and receptor activation/repression (Figure 2B and 2C). The extracellular loops of G-protein coupled receptors (GPCRs) are paradigmatic examples. They have an average length of 50 residues and have key functions for ligand recognition [25, 27].

Some of the general concepts related to IDP dynamics in the proximity of membranes are equally applicable to both sides of the membrane, albeit major differences exist between them. The interior of the cell presents an overall larger density in IDRs, with special highlight on the role of IDRs from non-transmembrane proteins that undergo reversible binding to the membranes (Figure 2A and 2D). Also, the cytoplasmic side accumulates negatively charged lipids. The emphasis of this manuscript is on the cytoplasmic disordered boundary of the cell.

**Tethering disordered proteins to the membrane through lipid anchors**

Lipid anchors allow the reversible association of proteins to the cytoplasmic side of the cell membrane [28]. Examples of lipid anchors include the N-terminal myristoylation that occurs in all SFKs and in the α-subunit of heterotrimneric G-proteins. Additional palmitoylation of cysteine regions is observed in most SFKs and G α-proteins (Figure 2A). Palmitoylation also occur in the cytoplasmic disordered tail of GPCRs constitutively connected to the integral membrane region. Thus, in this case, the lipid anchor probably acts as a modulator and is not the main driver of the protein-lipid interactions [29].

Small monomeric GTPases of the Ras family are modified at their C-terminus by isoprenoid moieties: farnesyl (3 isoprene units, 15 carbons) or geranylgeranyl (4 isoprene units, 20 carbons) located at the end of a Hypervariable Region. Additional palmitoylation of a second cysteine occurs in most Ras proteins [19]. The two-signal hypothesis indicates that two-lipid binding moieties are required for stable attachment of peripheral membrane proteins [28]. In the case of c-Src and K-Ras, that have only one lipid modification, N-terminal myristoylation and C-terminal farnesylation, respectively, a polybasic sequence seems to provide the required additional phospholipid-binding element (Figure 2A).

The myristoylated alanine rich c-kinase substrate (MARCKS) [30] is an abundant 35 kDa intrinsically disordered protein that contains a N-terminal myristoylation site and an effector domain in the center of the protein. This domain contains multiple positively charged lysine residues that provide an electrostatic attachment to negatively charged lipids. Additionally, it has multiple serine residues that, when phosphorylated by C-kinase or Rho kinase, can prevent the electrostatic interaction causing the protein detachment from the membrane into the cytosol. This is known as an electrostatic switch [31].

![Figure 2: Anchoring of IDRs to the membrane surface.](image-url)
A reciprocal effect of MARCKS on the lipid distribution has also been reported, affecting the distribution of PI(4,5)P₂, which is sequestered by the unphosphorylated effector domain, probably forming a neutral complex bringing together two or three PI(4,5)P₂ in a tight complex with the protein [32]. Thus, MARCKS provides an illustrative example of the possible cross-talk between the lipid and IDP systems in the DBC.

**Emerging properties of the DBC**

The emergent properties of the IDP-populated ε-boundary are a potential determinant of the function of the cell frontier. The disordered character of the anchored proteins has important implications at various levels: i) the modulation of the intrinsic properties of the individual IDR arising from the fact that they are tethered on the membrane surface; ii) the implications on the specific interactions between the membrane attached IDR and other signaling partners located in the proximity of the membrane, being themselves intrinsically disordered or globular; and iii) the modulation of the environment of the cell boundary affecting other components, which may include components not physically attached to the cell membrane but located in its proximity.

At the level of individual membrane anchored IDR, their conformational dynamics will be modulated by steric effects and by the presence of strong electrostatic fields. These fields are created by the negatively charged cytoplasmic membrane surface and by the abundant charged residues present in most IDRs.

The properties of individual IDR are distinctly manifested in their direct interactions. The interaction space available in the vicinity of the membrane is much more restricted than in an isotropic 3D scenario. To this extent, if a specific binding site is attached to a membrane anchored IDR, how would its potential interactions be modified?

**Dimensional scaling and rotational restriction**

Scaling of the dissociation constants from 3D to 2D has two well defined components, illustrated in Figure 3: First the collapse from an isotropic volume to a confined layer of width \( h \) results in an effective change in the concentration terms. The work of Bell et al. describes a model in which the dissociation constants between 3D and 2D would simply be scaled by an effective confinement length so that \( K_{d}^{2D} = h \cdot K_{d}^{3D} \) [33]. Secondly a restriction in the orientation of the interacting elements, so that the probability of productive encounters between potential partners resulting from their approaching in the proper orientation, is reduced [34].

Interestingly, as shown in Honig’s simulations [34], the conformational restriction induced by membrane anchoring may actually favor the clustering of membrane-attached proteins. The entropy loss intrinsic to any complex formation can be lower, if the potential interacting partners show a reduced conformational landscape due to their membrane attachment.

In Honig’s approach, membrane attachment plays only a restrictive role in the possible orientations. It was explicitly assumed that the energetic contributions from binding event would be identical in the isotropic 3D scenario and in the confined environment of the membrane. If we imagine that binding arises from the cooperative interaction between multiple weakly binding...
sites, not necessarily located contiguously along the IDR sequence, the anisotropic environment in the cell boundary can enforce distinct relative distributions of the individual sites with respect to the membrane surface.

This distribution can be strongly affected, for example, by the relative placement of charged residues and residues forming interaction sites along the membrane-anchored IDR sequence. Membrane anchoring can enhance lateral interactions by ensuring that multiple interacting sites are in register, as illustrated in Figure 4A.

Because of their density in the DBC, IDR may cause environmental effects that modulate the chemical potential of other chemical species, even if they are not physically attached to the membrane.

At least two effects are expected to be important: i) IDR induced changes in the effective polarizability of the boundary and ii) IDR induced crowding.

**IDR modulated electrostatics**

The electrostatic field originating on the negatively charged cytoplasmic lipid surface determines the equilibrium distribution of diffusing charged species in the boundary. While individual cations and anions can diffuse independently to reach equilibrium distributions, the positional distribution of charged residues in an IDR is defined by their protein sequence and depends on their conformational landscape. Thus, the electrostatic potential field is modulated by the presence of membrane attached IDRs. In parallel, the polarization of the IDR dipoles may actually extend the electrostatic influence of the charged membrane to longer distances than would be expected in an IDR-free membrane boundary (Figure 4B).
**IDR crowding**

IDR crowding (Figure 4C) represents a distinct situation from the well-known excluded volume crowding effects predicted for globular proteins. Specifically that, the centers of two rigid spherical objects cannot approach closer than the sum of their radii. Thus, the effective volume that can be occupied by one of the objects is reduced by the volume taken by the total number of the other objects present in the solution [35]. Two main IDP features depart from the simplest non-interacting hard-spheres model of crowding. First, their dynamic nature introduces a strong entropic component, and, secondly their highly accessible surface that facilitates non-specific interactions with other components of the cytoplasm, sometimes referred to as quinary interactions [36].

In the case of IDPs, the volume occupied by the molecule in a given instant is much smaller than the total volume sampled by the same molecule over a long time. At a same density of anchored proteins, IDRs can cover a larger surface than globular proteins [37]. When two IDPs approach, although they could occupy the same region of space, they cannot do it simultaneously. Therefore, their conformational space becomes restricted. The result is an entropic penalty that hinders the approximation of two IDR below a certain distance.

Crowding of membrane attached IDP is also modulated by the excluded volume arising from the proximity of the membrane. Convex membrane curvature around the protein attachment site increases the accessible volume that can be sampled by the IDP and therefore alleviates the entropic penalty.

Membrane bending by anchored polymers were theoretically predicted in 1995 [38]. Experimentally, the work of J. Stachowiak et al. has recently demonstrated the important role that disordered segments play in curvature formation and shape sensing. One of the most well-studies protein domains responsible for curvature sensing are BAR domains, that contain amphipathic helices and long disordered tails. At a fixed surface density, determined by the IDRs, the volume occupied by the same molecule over a long time. At a fixed surface density, determined by the interaction of the helical segments, convex curvature of the surface reduces the entropic penalty associated with the interaction of the IDR. Modulation of the local rigidity in disordered regions is a sophisticated mechanism used by BAR proteins for sensing curvature [39]. Conversely, enforcing the approximation of membrane anchored IDPs, for example by the interaction of neighbor globular domains, can induce membrane curvature [40].

The environmental effects of the IDR-rich DBC will affect all processes taking place in the ε-layer, including globular domains or other molecules not physically attached to the membrane. Thus, modifications of membrane attached IDRs, e.g. post translational modifications, may affect processes that are not, apparently, directly related to the modified IDP.

**Z-compartmentalization**

Compartmentalization is often found as a regulatory strategy restricting the possible partners for a given interaction by their common distribution in mutually excluding sites. In the cell boundary, the localization of interaction sites along the direction perpendicular to the membrane surface, the Z-direction, may generate an alternative compartmentalization mechanism (Figure 4D).

A example of Z-compartmentalization is illustrated by the ROCK2 kinase, a Rho-associated coiled coil kinase formed by constitutively active kinase dimers separated from the membrane anchoring regulatory domain by a long (107 nm) coiled-coil. The length of the coiled-coil is well conserved, although the sequence itself is variable, and consistent with its functioning as a molecular ruler. Truncation of the coiled-coil does not affect the kinase activity in vitro, but resulted in a complete loss of actin stress fibers when expressed in vivo. It has been suggested that ROCK2 activity is ultimately determined by the proximity of the substrate to the kinase domain, which would link phosphorylation to the distance to the membrane surface [41, 42]. MARCKS protein is a ROCK2 substrate [43].

While the rigid coiled coil provides a well-defined ruler, disordered linkers in the boundary of the cell may offer a modulation mechanism enabling or restricting the accessibility of the active site to specific substrates by changes in the flexibility of the connecting region, e.g. through post-translational modifications or alternative splicing changing the protein effective length [44]. A recent example shows that the length of the hypervariable domain (HVD) of Rab GTPases connecting their nucleotide binding domain (NBD) from their membrane anchoring, hydrophobic prenyl group, determine the differences in selectivity of two Rabs interacting with the same catalytic site. The Rabs are located at different distances from the membrane surface, resulting in spatial selectivity. A short HVD prevents the NBD from reaching the active site that stabilizes an otherwise reversible interaction with the membrane, but the effect can be reversed by artificially extending the length of the HVD [45].
Protein phase separation

Historically, “nucleolar accessory bodies”, later called Cajal bodies, were already described by Santiago Ramón y Cajal in 1903 in neuronal cells. These granules lack a surrounding phospholipid membrane and have remained obscure for a long time. One century later, it was demonstrated that these supramolecular assemblies have liquid properties, and are generated by phase separation of proteins and/or RNA. Recently, more examples have been described in biology and phase separation has emerged as a rediscovered field, but now involving cell biology, supramolecular chemistry, and polymer physics [5, 46]. Phase separation is a demixing transition driven by preferential interactions or selective exclusion within a homogeneous solution. Multiple weak interactions become cooperative when the interaction sites are part of a polymer but retain a highly dynamic exchange, resulting in liquid-like properties. The proteins associated with liquid-phases are mainly IDPs or low-complexity regions present in disordered segments. Phase transitions in multivalent signaling proteins have been suggested to be important in signaling [47].

Most of these spatially organized regions have liquid properties and are observed as “droplets” due to the surface tension. However, it is important to note that liquid-liquid phases generated by proteins and/or RNA deviate conceptually from the well-known van der Waals fluid, comprising spherical particles with isotropic interactions. Macromolecular polymers like proteins are highly anisotropic and protein-driven separation of protein fluid phases may be enhanced in the ε-layer.

Existing examples are the nephrin and LAT receptors. Nephrin receptor binds to Nck and the neural Wiskott-Aldrich syndrome protein (N-WASP) and form phase-separated liquid droplets in vitro [47]. In presence of lipid bilayers and in cells, at much lower concentrations, this assembly is reflect in form of membrane clusters [48]. Similarly, the interaction between the linker for activation of T cells (LAT), the growth factor receptor-bound protein 2 (GRB2), the GRB2-related adaptor protein 2 (GADS), the son of sevenless (SOS), and the SH2 domain-containing leukocyte of 76 kDa (SLP76) undergoes phase separation in vitro and generates membrane puncta in cells upon T cell receptor [49].

At least two ε-layer-specific features may enhance self-driven clustering of membrane-bound IDPs, positional matching of interaction sites and orientation bias. These two can be understood as an enhanced concentration of the interaction sites with respect to an isotropic solution. Positional matching implies a concentration of the interacting species by transferring from 3D to 2D. Orientation bias reflects the reduced entropic penalty associated to the cooperative interaction between two pairs of sites in separate molecules when these molecules adopt predominantly parallel dispositions, e.g. because both are tethered to the membrane surface.

The disordered region of Src family kinases: fuzzy complexes and membrane induced dimerization.

Recent studies of the membrane tethered, non-receptor SFKs have uncovered the importance of their intrinsically disordered domains and provide interesting examples of emerging properties associated to the DBC. The nine members of the SFK family have common domain architectures, with three folded domains and an N-terminal IDR that includes the membrane anchoring SH4 domain and the Unique domain [20]. The name “Unique” refers to the lack of homology of this region among the various SFKs that contrasts with the high conservation of the globular domains, including the kinase domain (SH1) and two classical regulatory domains (SH3 and SH2). Myristoylation of the SH4 domain and membrane anchoring are required for c-Src activity [50]. Titration with lipid bicelles monitored by NMR have identified additional lipid binding regions, in the Unique and SH3 domains [51]. The N-terminal myristoyl moiety binds to the SH3 domain in the absence of lipids, but is inserted into the lipid bilayer in the presence of lipid membrane (e.g. large unilamelar vesicles (LUV)). The SH3 domain nucleates an intramolecular fuzzy complex in which the Unique and SH4 domains are loosely tethered around the globular domain, while retaining their disordered character [52]. The intramolecular fuzzy complex is retained in the membrane-bound form bringing the SH3 domain in close proximity to the lipid surface [53]. Thus, globular domains “coated” with disordered proteins can also be an integral part of the DBC (Figure 5).

Interestingly, mutations that deactivate the lipid-binding region in the Unique domain result in a 50% decrease in the invasive capacity of Src-dependent colorectal cancer cells [52]. However, recent results show that in a LUV-anchored form of c-Src, containing the myristoylated SH4, Unique, and SH3 domains, the additional lipid binding site in the Unique domain is not interacting directly with the liposome. This lipid interacting region, contributes to stabilize the intramolecular interaction of the myristoyl group with the
SH3 domain in the non-membrane bound form [53] and modulates the orientation of the globular domains with respect to the membrane surface, via the intramolecular fuzzy complex. The change in orientation may explain the observed change in substrate specificity caused by mutations in the Unique lipid-binding region [Roche, S. private communication, manuscript in preparation], although more complex scenarios cannot be ruled out.

A recent report has suggested that the myristoyl group could also interact with the kinase domain of a second Src molecule to form a dimer [54]. The intermolecular interaction of the Src kinase domain with a N-terminal myristoyl group is expected to be promiscuous. Thus, the number of possible heterodimers is very large and interactions involving the kinase domain may affect its activity as well as the substrate preference, in addition to being modulated by the competing insertion of the myristoyl group in the lipid bilayer.

Dimerization on the surface of supported lipid bilayers had been previously observed for the N-terminal region of Src without the kinase domain [55,56]. In contrast to the transient binding observed by monomeric c-Src molecules to supported lipid bilayers, the membrane bound dimers were persistently attached and could be detected by antibodies even after extensive washing. Persistent binding is consistent with the simultaneous insertion of two myristoyl groups by c-Src dimers. Dimerization was not observed in solution or when the protein was deposited on a poly-lysine coated surface, thus the presence of phospholipids seems to be a requirement for membrane induced dimerization. Persistent binding was also observed in pure zwitterionic phosphatidyl choline supported bilayers, ruling out the possible partition of Src into phase-separated lipid regions.

The positive charges arising from the initial lysine rich segment (K5, K7, and K9) followed by a stretch of three arginine residues (R14, R15, and R16) contribute to increase the binding to negatively charged lipids but, surprisingly, also enhance the formation of dimers, in spite of the expected electrostatic repulsion.

Membrane induced oligomerization was observed in constructs containing the myristoylated SH4 domain attached to fluorescent proteins [57,58]. Extensive oligomerization was observed for short myristoylated peptides containing the entire SH4 region (15 residues) [55] or even in a myristoylated (2-9) c-Src peptide [59]. Interestingly, the first ten SH4 residues have a distinct role in the stabilization of the intramolecular fuzzy complex [52]. The interactions of the SH4 domain with the SH3 domain or the presence of the Unique domain seems to restrict the extent of oligomerization to dimers or trimers.

Surprisingly, the sequence of the initial segment of the c-Src SH4 domain is found in many snake venom analogues of phospholipase A2, in the juxtamembrane cytoplasmatic side of Adam 10, and, interestingly, in the C-terminal region of K-Ras B isoform, when the sequence is read in reverse order from the single farnesyl modified membrane anchoring element, that plays a similar role of the single myristoylated anchoring element in c-Src (Table 1).

Interestingly, membrane enabled dimerization of Adam 10 was found to depend on the presence of its intrinsically disordered cytoplasmic domain and required the disordered domain to be anchored to the membrane surface [60].
Table 1: Putative phospholipid binding sequences in disordered proteins.

| Protein, Isoform, Tail, (reverse sequence) | Sequence |
|------------------------------------------|----------|
| c-Src                                    | myr-GSNKSKPDASQRGR |
| Snake venoms analogues of phospholipase A2 | NKGKPDATDR |
| Adam 10 cytoplasmic tail                 | IQQPQRQRPRESYQMGHMRR |
| K-Ras isoform B                          | far-CKTKSKKKKKGDK |

Outlook: boundaries at different scales from lipid bilayers to the DBC

In spite of all their inherent complexity, membrane models provide a conceptual reference for a self-organizing dynamic system built from the assembly of layers with distinct properties, the hydrated layer containing the polar head groups and the hydrophobic core exhibiting a complex phase diagram. The dimensions of the two layers are related by one order of magnitude: roughly 0.5 nm and 5 nm.

The DBC can be viewed similarly as a self-organizing dynamic system in which the entire lipid membrane is one of the layers while the second layer is formed by the large concentration of disordered proteins tethered to the membrane surface in various ways. In this analogy, the cell membrane would play the role of the head-groups and the disordered proteins would form the core exhibiting collective properties, including protein driven phase transitions and imposing the peculiar environment that we have described as a fluid nanocomposite.

In contrast to the quasi-symmetrical arrangement of the lipid bilayers, the DBC can be better compared with a lipid monolayer facing the cytoskeleton, which is certainly not a rigid barrier. Recent super-resolution microscopy measurements give a first peak of maximum actin density from the cortical cytoskeleton at a distance of around 50 nm from the cell membrane [61]. The intervening region would correspond to the e-layer highly enriched 5 nm from the cell membrane [61]. The intervening region would correspond to the e-layer highly enriched 5 nm from the cell membrane [61].

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| Adam 10 cytoplasmic tail                 | IQQPQRQRPRESYQMGHMRR |
| K-Ras isoform B                          | far-CKTKSKKKKKGDK |

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