How cholesterol interacts with membrane proteins: an exploration of cholesterol-binding sites including CRAC, CARC, and tilted domains

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INTRODUCTION

Transmembrane domains of proteins cross the lipid bilayer of biological membranes to ensure the insertion of a subset of amino acid residues within the membrane. These proteins are usually referred to as integral membrane proteins. Deciphering the molecular mechanisms allowing the interaction of these proteins with membrane lipids has been and still is the subject of intense research efforts at the crossroad of several scientific areas including biochemistry, biophysics, cell biology, and bioinformatics. Basically, a transmembrane domain is an α-helical segment of ca. 20–25 apolar amino acid residues flanked at each end by more polar residues allowing them to “float” at the lipid-water interface region of the membrane, thereby stabilizing the helix within the lipid bilayer (Lee, 2003). These “interfacial” amino acid residues have to manage the transition between an apolar and a polar environment. For this reason, Lys, Arg, Tyr, and Trp residues are most commonly found at these flanking positions. Lys and Arg have a long apolar side chain buried in the apolar section of the membrane, and a positively charged basic group that “breathes” at the surface of the membrane. This unique topology and its associated functional counterpart are metaphorically referred to as the “snorkeling” effect (Strandberg and Killian, 2003). By the same token, Trp, and Tyr have an aromatic structure compatible with the apolar region of the membrane, but contain an –OH group (Tyr) or an N atom (Trp) capable of forming hydrogen bonds with polar groups (Lee, 2003). Nonhelical transmembrane β structures have also been characterized, for instance in bacterial cytolysins (Harris and Palmer, 2010), but are less represented than the widespread α-helical domains in resident plasma membrane proteins. In this review, we will focus on TM domains with an α-helical structure and their different modes of interaction with membrane cholesterol.

CHOLESTEROL STRUCTURE, DYNAMICS, AND MEMBRANE TOPOLOGY

Cholesterol is a polycyclic amphipathic molecule derived from the sterane backbone (Fantini and Barrantes, 2009). Its polar section is restricted to a single hydroxyl (OH) group which can form two distinct types of hydrogen bond (acceptor and donor) with a polar group belonging to either a membrane lipid or a protein. The apolar section of cholesterol has an asymmetric structure with two distinct faces, referred to as α and β according to the system numeration of ring compounds proposed by Rose et al. (1980) (Figures 1A and B). The α face displays a planar surface,
in contrast with the β face which has a significantly rougher surface owing to the presence of several aliphatic groups (two methyl groups and a terminal iso-octyl chain that are linked to the sterane backbone) (Fantini and Barrantes, 2009) (Figure 1C). The side chains of branched amino acids such as Ile, Val, or Leu can interpenetrate these aliphatic “spikes” and are thus particularly suited for an association with the β face of cholesterol through van der Waals interactions. This is the case for the cholesterol binding domain of α-synuclein (Fantini et al., 2011). Moreover, aromatic side chains can stack onto the α face of cholesterol through CH−π interactions (Nishio et al., 1995). However, this should not be taken as an absolute rule since the aliphatic side chains of an α-helical segment could also form a groove with a planar surface fitting the α face of cholesterol (Di Scala et al., 2013). Conversely, an aromatic ring oriented normally with respect to the main axis of an α-helical region could perfectly well accommodate the rough β face of cholesterol by intercalating the aromatic structure between the aliphatic spikes of the lipid.

Another key parameter which determines how cholesterol interacts with a TM domain of a protein is the membrane phase to which it belongs. It should be kept in mind that although cholesterol is concentrated in sphingolipid-enriched membrane microdomains such as lipid “rafts” (Simons and Ikonen, 1997; Anderson and Jacobson, 2002) it is also present outside these microdomains, i.e., in the liquid disordered (Ld) phase of the plasma membrane that contains high amounts of glycerophospholipids such as phosphatidylcholine (Fantini et al., 2002). Studies with model membranes indicate that cholesterol interacts more favorably with sphingomyelin than with phosphatidylcholine (Mattjus and Slotte, 1996) (Figure 2), a behavior which has been attributed to the presence of a saturated acyl chain in sphingomyelin compared to a cis-unsaturated chain in phosphatidylcholine (Fantini et al., 2002). The saturated chain in sphingomyelin, together with the trans-unsaturated sphingosine backbone, would allow maximal van der Waals interactions with cholesterol. Moreover, phosphatidylcholine (Figure 2A) has carbonyl groups which could act as hydrogen bond acceptors, but is devoid of any hydrogen bond donor group such as the amino group of sphingomyelin (Figure 2B). Therefore, the association between cholesterol and phosphatidylcholine relies on weakly discriminative van der Waals forces plus limited hydrogen bond capabilities. When cholesterol is associated with phosphatidylcholine, both its α and β faces (Rose et al., 1980) are potentially available for an interaction with a TM domain (Figure 2A). Moreover, the −OH group of cholesterol is not buried in the phosphatidylcholine/cholesterol complex and thus remains accessible for establishing a hydrogen bond with a TM domain. In contrast, cholesterol forms condensed lipid complexes with sphingolipids (either sphingomyelin or glycosphingolipids) as shown in Figure 2B (Radhakrishnan et al., 2000). In these molecular assemblies, the −OH group of cholesterol is available for the formation of a stabilizing hydrogen bond with the polar head...
group of the sphingolipid. Thus, this −OH group is not initially available for a hydrogen bond with a TM domain of a protein. Sphingolipids generally interact with the α face of cholesterol, leaving the β face available for the TM domain (Fantini and Barrantes, 2009). Finally, it is interesting to note that in model membranes, cholesterol can form two distinct types of dimers that are stabilized through van der Waals interactions: (1) transbilayer tail-to-tail dimers (Harris et al., 1995; Mukherjee and Chattopadhyay, 1996; Rukmini et al., 2001) (Figure 3A); and (2) dimers formed by the association of two cholesterol molecules interacting with their respective smooth α faces, leaving the opposite β faces available for protein binding (Figure 3B). In this latter case, the cholesterol dimer can recruit for instance two G-protein coupled receptors, thereby inducing their functional dimerization (Figure 3C) (Hanson et al., 2008). That such cholesterol dimers actually exist in natural plasma membranes has not been formally demonstrated. Although the intracellular dynamics of cholesterol has been the subject of intense research efforts during the last decade (Mesmin and Maxfield, 2009), the transbilayer distribution of cholesterol has remained uncertain (Ikonen, 2008). Indeed, despite the fact that cholesterol preferentially interacts with sphingolipids in the exofacial leaflet, fluorescence quenching studies suggested that it is in fact more abundant in the cytoplasmic leaflet (Mondal et al., 2009). The molecular basis for such an asymmetric transbilayer distribution of plasma membrane cholesterol is unknown. However, the limited accessibility of cholesterol in the exofacial leaflet, due to its tight interactions with sphingolipids, could lead to an underestimation of its content in this leaflet. Nevertheless, whichever membrane layer has the highest cholesterol content, cholesterol is present in both leaflets of the plasma membrane, providing various interaction possibilities with TM domains. In all cases, unraveling the molecular mechanisms involved in the interaction of TM domains with cholesterol may require (1) the description of the cholesterol/α-helix complex at the molecular/atomic level and (2) consideration of the lipid/protein environment in which this interaction occurs. With this in mind we will now describe the different types of protein structures capable of interacting with membrane cholesterol.

THE CRAC DOMAIN

There is little doubt that the most popular cholesterol-binding domain in the scientific literature is the **Cholesterol Recognition/interaction Amino acid Consensus** sequence, generally referred as the CRAC domain (Li and Papadopoulos, 1998). CRAC is a short linear motif which fulfills a very simple algorithm, that is, in the N-terminus to C-terminus direction: a branched apolar Leu or Val residue, followed by a segment containing 1–5 of any residues, then an aromatic residue that is mandatory Y, then again a segment containing 1–5 of any residues, and finally a basic Lys or Arg, i.e., in the one letter amino acid code, \((L/V)\)-\(X_{1-5}\)-(Y)-\(X_{1-5}\)-(K/R). The looseness of the CRAC definition for a motif that mediates binding to a unique lipid that does not display any structural variation is rather unexpected, raising some skepticism about its predictive value (Palmer, 2004; Epand, 2006). However, the motif has been found in various proteins known to bind cholesterol and in many cases the interaction between cholesterol and CRAC has

![Figure 3](https://www.frontiersin.org)
been confirmed by physicochemical approaches. Moreover, single mutations in the CRAC domain have been found to markedly decrease or abolish the interaction, as is the case for the central aromatic residue which is necessarily Tyr and cannot be replaced by any other aromatic residue (Jamin et al., 2005; Epand, 2006; Epand et al., 2006).

Molecular modeling studies have shown that the CRAC motif belonging to TM domains can have a good fit for cholesterol, as illustrated for the 5th TM domain of the human type 3 somatostatin receptor (Figure 4A). This CRAC domain lies between amino acid residues 221–231 and has the following sequence: VICLCYLLIVVK. It can therefore be either interpreted as LCYLLIVVK or as VICLCYLLIVVK (amino acid residues that fulfill the CRAC algorithm in bold and underlined). Detailed analysis of the energy of interaction between cholesterol and this CRAC domain showed that the complex involves essentially five residues, four of which belong to the CRAC motif (V-221, C-225, L-228, and I-229) and the fifth remaining outside (K-232), for a total energy of interaction of $-43\text{kJ.mol}^{-1}$ (Baier et al., 2011). Surprisingly, the central Tyr residue does not interact with cholesterol, although it is mandatory in the definition of CRAC. Moreover, the energetic pattern of the cholesterol-binding site, i.e., 221-VICLCYLLIVVK-232 (residues involved in cholesterol binding in bold and underlined) does not overlap with any of the two CRAC motifs defined above (LCYLLIVVK or VICLCYLLIVVK). Overall this may indicate that although the CRAC algorithm has some predictive value for identifying cholesterol-binding motifs in the TM domain, cholesterol can choose a slightly different fit around the CRAC domain to adjust its shape to the three-dimensional structure of the TM domain.

There is another caveat in the predictive value of the CRAC algorithm for the specific case of TM domains: the CRAC domain is an oriented motif, with an apolar amino acid residue at the N-terminus and a highly polar, positively charged basic residue at the C-terminus. This means that if a CRAC motif belongs to a TM domain and allows this domain to interact with cholesterol, several parameters other than the classical CRAC algorithm have to be fulfilled. First, the basic residue at the C-terminus should be located at the lipid-water interface to ensure an optimal interaction with the membrane structure. Therefore, if the TM domain containing the CRAC motif crosses the membrane in the N-term to C-term direction (i.e., with the N-term region extracellular and the C-term region cytoplasmic), then it will interact with cholesterol in the cytoplasmic leaflet of the plasma membrane (Figure 4A). Conversely, if the TM domain crosses the plasma membrane in the opposite direction (i.e., the N-term region cytoplasmic and the C-term region extracellular), then it will interact with cholesterol in the extracellular leaflet. In both cases, the amino acid residues of the variable segments separating L/V from Y and Y from K/R can still vary, but they must be apolar because they are embedded in the apolar part of the membrane. Since the original definition of the CRAC domain did not specifically take the membrane insertion of the motif into consideration, the X amino acid residues could be any residue.

For CRAC motifs belonging to TM domains, the definition should therefore be restricted to (L/V)-X$_{-1}$-(Y)-X$_{-1}$-(K/R) with apolar X residues compatible with the hydrophobic membrane environment, otherwise the CRAC algorithm could incorrectly predict the presence of a potential cholesterol-binding domain that in fact lies outside the membrane. This particular case is illustrated for the human delta-type opioid receptor, which contains a CRAC motif the juxtamembrane domain just upstream of the 2nd TM domain (Figure 4B). The sequence of this CRAC domain is 74-IVRYTKMK-81, whereas the 2nd TM domain encompasses residues 85–102. As a matter of fact, the very high polarity of this CRAC motif restricts its location outside the membrane. Nevertheless, as a bona fide CRAC domain, the IVRYTKMK sequence has a high affinity for cholesterol, with a total energy of interaction of $-49\text{kJ.mol}^{-1}$, as calculated from docking studies (Baier et al., 2011). The amino acid residues involved in cholesterol binding are I-74, V-75, R-76, Y-77, and T-78, which can be summarized as a contiguous IVRYTKMK motif containing two of the three residues that define the CRAC domain, including the central and mandatory Tyr residue. This aromatic residue (Y-77) contributes $-26\text{kJ.mol}^{-1}$, which represents about 50% of the total energy of interaction of the CRAC/cholesterol complex. Indeed, the side chain of Y-77 binds to cholesterol through a CH-$\pi$ stacking interaction with the B ring of sterane, whereas its OH group can contribute to a network of electrostatic interactions that also includes the polar OH group of cholesterol (Figure 5). Nevertheless, the predicted membrane topology of the human delta-type opioid receptor suggests that this CRAC domain lies outside the membrane so that its interaction with cholesterol, although theoretically possible, is unlikely (Figure 4B). Despite these caveats there are numerous cases of CRAC motifs within TM domains, illustrating the overall robustness of the CRAC algorithm for predicting...
cholesterol-binding sites in TM segments of integral membrane proteins (Epand, 2006; Epand et al., 2010; Gimpl, 2010; Paila and Chattopadhyay, 2010), including G-protein coupled receptors of several distinct neurotransmitters (Jafurulla et al., 2011; Oddi et al., 2011; Sengupta and Chattopadhyay, 2012). However, in some instances, bioinformatics analysis of the amino acid sequence of some membrane proteins failed to identify any CRAC motifs encased within TM domains. This is the case for the human nicotinic acetylcholine receptor (AChR) and this particular situation led to the development of a new algorithm for predicting cholesterol-binding domains in TM segments of membrane proteins.

THE CARC DOMAIN

The new cholesterol-binding domain is similar to the CRAC sequence, but exhibits the opposite orientation along the polypeptide chain (i.e., constitutes an “inverted CRAC” domain), i.e., (K/R)-X1–5-(Y/F)-X1–5-(L/V) from the N-term ending to the C-term ending (Baier et al., 2011), for which reason the sequence was coined “CARC.” Besides the reverse orientation, CARC is distinct from CRAC in that the central aromatic amino acid can be either Tyr or Phe. Basically, the CARC motif is based on a sequence of amino acids which includes, from N-term to C-term endings, a basic (Lys or Arg), an aromatic (Tyr or Phe) and a branched aliphatic residue (Leu or Val). The presence of the basic residue ensures that the CARC motif is correctly positioned at the polar-apolar interface of a TM domain, exactly where cholesterol is supposed to be. This is due to the “snorkeling” effect discussed above (Strandberg and Killian, 2003) which can be attributed to the fact that the long and flexible side chain of lysine (or arginine) is buried in the hydrophobic part of the membrane, whereas the cationic group emerges at the membrane surface (Segrest et al., 1990; Baier et al., 2011). Interestingly, Arg and Lys residues are more frequently found in the N-terminal than in the C-terminal section of a TM domain, indicating that snorkeling is an asymmetric phenomenon (Strandberg and Killian, 2003). The specific topology of the CARC sequence (K/R…Y/F…L/V, from the N-terminus to the C-terminus) implies that the key residues following K/R in the motif (i.e., Y/F and L/V) actually belong to a TM domain, and this is consistent with the fact that both aromatic and aliphatic residues have an apolar side chain.

Moreover, even if there is no direct interaction between cholesterol and K/R, the basic amino acid in the first position is critical for identifying a functional CARC motif belonging to a TM domain. Incidentally, this also explains why the previously characterized CRAC motif, which has an inverse topology (L/V…Y…K/R, from the N-terminus to the C-terminus), does not always belong to a TM domain (Figure 4B). Furthermore, the tyrosine residue (an absolute requirement for CRAC) can be functionally replaced by Phe in the CARC motif. In the CRAC motif, the phenol group of tyrosine is often required to form a H-bond with the OH group of cholesterol (Epand et al., 2010), and this would not be possible with Phe. In the case of CARC, the interaction between the aromatic amino acid and cholesterol occurs in the apolar region of the membrane, far from the OH group of cholesterol, and the interaction with cholesterol is mediated almost exclusively by the CH-π stacking arrangement between the aromatic ring of the amino acid (either Tyr or Phe) and one of the sterane rings of cholesterol (Figure 6). Finally, the requirement for Leu or Val is justified by the need to accommodate the crevices and asperities of the cholesterol molecule (Fantini and Barrantes, 2009) through numerous van der Waals contacts between these residues and cholesterol. The human type 3 somatostatin receptor illustrates these biochemical principles: in this case, the CARC domain corresponds to the 203-[RAGFIYTAAL]-213, which overlaps the extracellular leaflet of the 5th domain of the receptor (this TM domain encompasses the segment 206–231). Thus, though as discussed above the Arg residue does not belong to the TM domain, it is important...
for defining the correct orientation of the TM domain with the CARC motif in the exofacial leaflet of the plasma membrane. All the key amino acid residues that define the CARC motif (R-203, F-206, Y-209, and L-213) do interact with cholesterol, rendering a total energy of interaction of $-54 \text{kJ.mol}^{-1}$ (Figure 6). What is particularly interesting in this CARC motif is that it contains not just one but two aromatic residues that both stack onto cholesterol through near-perfect CH-π interactions. Correspondingly, the energetic contribution of these aromatic residues for cholesterol binding is high: $-18.2 \text{kJ.mol}^{-1}$ and $-14.7 \text{kJ.mol}^{-1}$ for F-206 and Y-209, respectively. The interaction of L-213 with the isooctyl chain of cholesterol accounts for $-10.0 \text{kJ.mol}^{-1}$. This illustrates the efficiency of van der Waals interactions between the branched side chain residues and the rough β face of cholesterol. Overall, these unique biochemical features explain why the CARC domains detected in the AChR and its homologous ion channels cover a wide evolutionary span from bacteria to humans (Baier et al., 2011). The CARC motif is also present in the TM domains of the important group of G-protein coupled receptors, which, together with the AChR and other neurotransmitter receptors, are thought to play a role in the pathogenesis of Alzheimer’s disease, in connection with α-synuclein, the neural protein associated with Parkinson disease, which can form oligomeric ion channels in the plasma membrane of neurons (Fantini and Yahi, 2010, 2011). This protein contains a CRAC domain which has been shown to bind cholesterol with low affinity. However, the domain is located outside the membrane-spanning regions of the proteins that supposedly interact with cholesterol and display neither CRAC nor CARC motifs (Fantini et al., 2011). Nevertheless, a second cholesterol-binding domain has been successfully characterized in α-synuclein, corresponding to a tilted peptide known to be toxic for cultured neurons, namely the segment 67-GGAVVTGVTAVA-78 (Fantini et al., 2011). What is intriguing about this peptide is that it does not contain the basic and aromatic residues that are mandatory in both the CRAC and CARC algorithms. Indeed, the definition of tilted peptides is functional, not sequence-based. Tilted peptides are short helical protein fragments that are able to disturb the organization of the molecular system into which they insert. They are characterized by an asymmetric distribution of their hydrophobic residues, which induces a tilted orientation (around 45°) toward the membrane plane (Lins et al., 2008). Because they induce a significant distortion of the membrane structure, tilted peptides are involved in the fusion process triggered by viral glycoproteins.

TILTED PEPTIDES

There are several instances in which a cholesterol-binding site is functionally characterized in a TM segment without the help of the CRAC or CARC algorithms. This has recently been shown to be the case for α-synuclein, the neural protein associated with Parkinson disease, which can form oligomeric ion channels in the plasma membrane of neurons (Fantini and Yahi, 2010, 2011). This protein contains a CRAC domain which has been shown to bind cholesterol with low affinity. However, the domain is located outside the membrane-spanning regions of the proteins that supposedly interact with cholesterol and display neither CRAC nor CARC motifs (Fantini et al., 2011). Nevertheless, a second cholesterol-binding domain has been successfully characterized in α-synuclein, corresponding to a tilted peptide known to be toxic for cultured neurons, namely the segment 67-GGAVVTGVTAVA-78 (Fantini et al., 2011). What is intriguing about this peptide is that it does not contain the basic and aromatic residues that are mandatory in both the CRAC and CARC algorithms. Indeed, the definition of tilted peptides is functional, not sequence-based. Tilted peptides are short helical protein fragments that are able to disturb the organization of the molecular system into which they insert. They are characterized by an asymmetric distribution of their hydrophobic residues, which induces a tilted orientation (around 45°) toward the membrane plane (Lins et al., 2008). Because they induce a significant distortion of the membrane structure, tilted peptides are involved in the fusion process triggered by viral glycoproteins.
This is the case with the tilted peptide of α-synuclein, which binds cholesterol with a tilt angle of 46° (Crowet et al., 2007). Despite the lack of aromatic residues, the contribution of apolar aliphatic residues accounts for a total energy of interaction of ~53 kJ.mol\(^{-1}\), which is comparable to the values obtained for both CRAC and CARC domains. This further illustrates how efficient van der Waals interactions between branched apolar residues and the β face of cholesterol can be. Similarly, a linear motif (22-EDVGSNKGAIIGLM-35) including a part of the tilted domain of Alzheimer’s β-amyloid peptide has been identified as a high affinity binding site for cholesterol (Di Scala et al., 2013).

An example of the interaction between a fusogenic tilted peptide and cholesterol is shown in Figure 8. This is the tilted peptide of the transmembrane glycoprotein gp41 of HIV-1. The tilted orientation of the peptide with respect to cholesterol is clearly visible in the model, and the tilt angle of 41° is close to the experimental value (Charloteaux et al., 2006). The association of the tilted peptide with cholesterol is mediated by a series of van der Waals interactions for a total energy of interaction of ~48.5 kJ.mol\(^{-1}\). Tilted peptides have been detected in various proteins otherwise known to require cholesterol for their membrane insertion process, including amyloidogenic and viral fusion proteins (Fantini et al., 2011). This suggests that tilted peptides have evolved in such a way as to acquire the cholesterol-binding properties that facilitate their biological functions.

**OTHER CHOLESTEROL-BINDING MOTIFS**

An important issue to resolve is whether cholesterol can interact with TM domains of protein lacking a CRAC, a CARC or a tilted domain. If we consider any possible interaction between a membrane protein and its surrounding lipids, including cholesterol, then the answer might be yes. In this respect, two distinct types of lipid-protein interactions in the membrane environment should be considered. Lipids in the first boundary shell, or annulus, are referred to as annular or belt lipids (see review in Marsh and Pali, 2012). Such annular lipids interact with the membrane-embedded surface of the protein (Lee, 2011). In addition, there is a second category of lipids that are buried to a greater or lesser extent in the protein. Such lipids are found bound between TM domains and are referred to as non-annular lipids (Lee, 2011; Marsh and Pali, 2012). Because they are bound to the protein surface, annular lipids can be exchanged for other membrane lipids. This is usually not the case for non-annular lipids that are less or not accessible at all to surrounding membrane lipids. Indeed, non-annular binding sites may involve several TM domains that form a 3D binding site for the lipid. For instance, several potential cholesterol-binding residues located in distinct TM domains of the 5-HT1A serotonin receptor were identified by comparison with the conserved cholesterol-consensus motifs in the β2-adrenergic receptor—as predicted from its crystal structure (Paila et al., 2009). Interestingly, the evolutionary conserved residues of this non-annular binding site are typically those found in CRAC and CARC domains, i.e., a combination of aromatic (Y-73 in TM2, W-161 in TM4), basic (R-151 in TM4) and aliphatic (I-157 in TM4) residues. Thus, the molecular mechanisms of cholesterol binding described for the CRAC and the CARC domains should probably also apply for this 3D binding site.

Overall, the interaction between membrane proteins (or peptides) and annular/non-annular lipids has been studied by three main approaches: \(^2\text{H}\) nuclear magnetic resonance (NMR), spin-label electron paramagnetic resonance (ESR) and X-ray crystallography (see reviews in Marsh, 2008, 2010). This has enabled determination of the stoichiometry of the lipid–protein interaction, the selectivity of the protein for different lipids and the amino acid residues physically involved in lipid binding. X-ray crystallography is of particular value since it can give a precise description of lipid-protein contacts at the atomic scale.

Unfortunately, very few 3D structures of protein-cholesterol complexes are available in the literature. In 2008, Hanson et al. published the 2.8Å resolution crystal structure of a thermally-stabilized human β2-adrenergic receptor bound to cholesterol. The cholesterol binding site of this receptor was defined as a 3D motif involving the 1st, 2nd, 3rd, and 4th TM domains (referred to as TM1, TM2, TM3, and TM4). Among these domains, TM2 and TM4 appeared particularly critical for cholesterol binding. Interestingly, TM4 contains an aromatic tryptophan residue (W-158) that is highly conserved among human G-protein coupled receptors. The aromatic side chain of W-158 stacks onto cholesterol through a CH–Pi interaction with ring D of the sterol. The binding site also involves two other amino acid residues of TM4 (R-151 and L-163) and a second aromatic residue located in TM2 (Tyr-70).

As noted by Hanson et al. (2008), the amino acid sequence of TM4 contains the combination of basic (R), aromatic (W) and aliphatic (L/V) residues found in CRAC domains (151-RVIIIMVIVSGILTSLPIQMHWY-174). However, this sequence does not fulfill the criteria of the CRAC domain. Moreover, it is even closer to a CARC domain (with the typical R—W—L motif), except that there are 6 residues between R-151 and W-158, which exceeds by one unit the maximal number of 5 residues allowed by the CARC algorithm. Finally, the...
CONCLUSIONS AND PERSPECTIVES

At first glance, this review of cholesterol-binding sites encased in TM domains of proteins may appear rather complex. Two consensus domains, CRAC and CARC, have been characterized. As a cholesterol-binding domain present in a TM domain, CARC appears more consistent than CRAC in predicting cholesterol-recognition motifs in integral membrane proteins, especially for predicting cholesterol-binding sites located in the exofacial leaflet of the plasma membrane. The CRAC algorithm is generally relevant (Sengupta and Chattopadhyay, 2012), but it can occasionally mis-predict unrealistic cholesterol-binding domains outside TM domains (Figure 4B). Fusogenic tilted peptides lacking a CARC or a CRAC domain can interact with cholesterol, although they do not contain the basic and aromatic amino acid residues that are mandatory for both the CARC and CRAC algorithms. Finally non-annular cholesterol-binding sites can be formed by the cooperation of several TM domains of G-protein coupled receptors, rendering the prediction of such binding domains particularly difficult.

Linear and 3D binding sites for cholesterol might have distinct biological functions. In bringing together several TM domains, cholesterol could exert a condensing effect on the whole protein, which, e.g., in the case of the 5-HT1A receptor could help the protein to acquire some of its functional characteristics, including the delineation of the ligand-binding pocket (Paila et al., 2011). In contrast, cholesterol interacting with only one TM domain at the periphery of a receptor protein would be more suited to bringing together two distinct receptor macromolecules and triggering their dimerization, a key step in signal transduction cascades (Figure 3C).

This may explain why various types of cholesterol-binding motifs can be found in TM domains. These motifs may vary greatly in length, amino acid sequence, and can either be linear such as the CRAC (Li and Papadopoulos, 1998), CARC (Baier et al., 2011), and tilted peptides (Fantini et al., 2011) or three-dimensional, e.g., cholesterol consensus motifs (Hanson et al., 2008) and sterol sensing domains (Garmy et al., 2005). Nevertheless, common molecular mechanisms seem to control the interaction of transmembrane proteins with cholesterol. These mechanisms include the delineation of a polar area to accommodate the OH group of cholesterol at the membrane-water interface, and the establishment of numerous van der Waals interactions in the apolar zone of the membrane (Epa nd et al., 2006; Baier et al., 2011). These van der Waals interactions comprise London forces and CH–π interactions. London forces require an optimal geometric fit between cholesterol and the apolar side chains of the amino acid residues that belong to the motif. This explains why branched amino acid residues such as Ile, Val, and Leu are overrepresented in cholesterol-binding sites. Aromatic residues can stack onto the smooth α face of cholesterol or intercalate between the aliphatic spikes emerging from the rough β face of cholesterol. These simple biochemical rules probably apply to most if not all cholesterol-binding sites that are part of TM domains of proteins.

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