Cholesterol plays a decisive role in tetraspanin assemblies during bilayer deformations†

Marcelo Caparotta\textsuperscript{a} and Diego Masone\textsuperscript{b,c}\

The tetraspanin family plays key roles in many physiological processes, such as, tumour invasion, cell motility, virus infection, cell attachment and entry. Tetraspanins function as molecular scaffolds organised in microdomains with interesting downstream cellular consequences. However, despite their relevance in human physiology, the precise mechanisms of their various functions remain elusive. In particular, the full-length CD81 tetraspanin has interesting cholesterol-related properties that modulate its activity in cells. In this work, we study the opening transition of CD81 under different conditions. We propose that such conformational change is a collaborative process enhanced by simultaneous interactions between multiple identical CD81 tetraspanins. With molecular dynamics simulations we describe the crucial role of a ternary lipid bilayer with cholesterol in CD81 conformational dynamics, observing two emergent properties: first, clusters of CD81 collectively segregate one tetraspanin while favouring one opening transition, second, cumulative cholesterol sequestering by CD81 tetraspanins inhibits large membrane deformations due to local density variations.

\textsuperscript{a} Facultad de Ciencias Exactas y Naturales, Universidad Nacional de Cuyo (UNCuyo), 5500, Mendoza, Argentina.\n\textsuperscript{b} Instituto de Histología y Embriología de Mendoza (IHEM) - Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Universidad Nacional de Cuyo (UNCuyo), 5500, Mendoza, Argentina.\n\textsuperscript{c} Facultad de Ingeniería, Universidad Nacional de Cuyo (UNCuyo), 5500, Mendoza, Argentina.\n\textsuperscript{*} Corresponding author: diego.masone@ingenieria.uncuyo.edu.ar.
\textsuperscript{†} Electronic Supplementary Information (ESI) available: ESI includes PMF convergence analysis for free energy profiles; umbrella sampling technical details; distance-angle 2D scattered maps for atomistic simulation; atomistic molecular dynamics initial snapshots states for 2 and 4 CD81 tetraspanins; backconverted snapshots from the coarse-grained space to all-atom resolution and area relaxation plots for atomistic systems containing 1,2 and 4 CD81 tetraspanins. See DOI: 00.0000/00000000.
1 Introduction

Tetraspanins are a particular kind of membrane proteins expressed in animals, plants and fungi. Although they are the largest family of transmembrane proteins in mammals, little is known about their molecular mechanisms of action. Having a strong propensity to multimerise, they self-organize along the plasma membrane in high density clusters (tetraspanin-enriched microdomains or tetraspanin webs) that play a central role in various cellular functions.

Tetraspanins are known as molecular facilitators and master organizers of the membrane architecture, controlling fundamental immune cell functions and a wide range of physiological processes. They are of key importance in cell morphology, invasion, motility, fusion and signalling, in the brain, in the immune system and in tumours. Tetraspanins are involved in cell proliferation, pathogen uptake and fertilization. Therefore, there is great interest in describing tetraspanins as novel drug-targets for infectious diseases and cancer.

In particular, CD81 (cluster of differentiation 81), or Tspan28, is involved in cell-surface assembly of the human immunodeficiency virus (HIV), the influenza A virus and the hepatitis C virus (HCV). Several studies have highlighted the affinity of CD81 for highly curved membranes or its influence in membrane remodelling and deformation. Therefore, there is great interest in describing tetraspanins as novel drug-targets for infectious diseases and cancer.

Structurally, tetraspanins contain four membrane-spanning domains connected extracellularly by a large and a small loop. Tetraspanins interact directly with lipids and other membrane proteins to control the architecture of the membrane. In particular, CD81 tetraspanin (PDB ID: 5tcx) contains a cholesterol binding pocket between its transmembrane domains which has been shown (both experimentally and computationally) to influence the open/close conformational change whether cholesterol is present or not. Such opening events increase the flexibility of the external region containing γ and δ loops (res 158-174 and res 176-189 respectively), known to be important in CD81:CD81 direct interactions. In this sense, molecular dynamics has proved indispensable in both pure and applied research providing the methodology for detailed microscopic modelling on the molecular scale.

In this work, we studied CD81 assemblies as a collective phenomenon, observing their collaborative properties due to simultaneous interactions between multiple tetraspanins. First, we showed that the conformational opening transition already described for CD81 inserted in a lipid bilayer, takes place only partially in an isolated tetraspanin outside the bilayer. Second, we described this opening transition as favoured by assemblies of multiple CD81 tetraspanins solvated in explicit solvent, still outside the bilayer. Third, we observed that when inserted in a lipid bilayer containing cholesterol, CD81 tends to cluster and to segregate one tetraspanin. Moreover, we determined that the segregation of cholesterol, CD81 tend to cluster and to segregate one tetraspanin. Moreover, we determined that the segregation of cholesterol, CD81 tends to cluster and to segregate one tetraspanin. Moreover, we determined that the segregation of cholesterol, CD81 tends to cluster and to segregate one tetraspanin. Finally, we described the effects of CD81 tetraspanins in bending the bilayer, showing that cholesterol sequestering prevents large deformations induced by local density variations.

2 Results and discussion

2.1 CD81 open transition is a collaborative event

![Fig. 1 Open transition in a single CD81 tetraspanin. a) Free energy profile using Phe58-Phe126 Cα distance as reaction coordinate (θ). Molecular dynamics snapshots depict the transition steps. Angle θ is measured between Cα atoms of residues Val114, Val136 and Gin88 (orange spheres). Error bars are standard errors calculated by individually splitting the profile in independent blocks. See supplementary information for PMF convergence details. b) Distance-angle 2D scattered map for all umbrella windows used to recover the free energy profile in panel a). c) Distance-angle 2D scattered map for 1μs unbiased molecular dynamics of a single CD81 in water. d) Averaged distribution of the angle and the distance along the 1μs unbiased simulations. e) Molecular dynamics snapshots show unbiased initial (t=0μs) and final (t=1μs) states. CD81 is represented in black ribbons, with Phe58 and Phe126 highlighted in vDW representations following the Corey-Pauling-Koltun (CPK) colouring convention for distinguishing atoms. Water molecules are not shown. f) Phe58-Phe126 averaged distance over unbiased molecular dynamics in the μs time scale. To observe the conformational change of CD81 from closed to open states, we have obtained the free-energy profile describing this transition in the full-length human CD81 tetraspanin (PDB ID 5tcx) after removing cholesterol molecules (apo-protein) and solvated in TIP3 water molecules under the CHARMM36 force-field. The reaction coordinate used to explore the energy path was the 3D distance between Cα atoms in residues Phe58 and Phe126. This convenient distance definition was originally proposed by Blacklow and collaborators to characterise, with unbiased molecular dynamics, the opening transi-
tion of a single CD81 tetraspanin inserted in a 124 palmitoyl-
oleoyl-phosphatidylcholine (POPC) lipid bilayer, using TIP3 water
molecules and the CHARMM36 force-field, as well. They showed
that in 3 out of 9 trajectories the apoprotein successfully adopts an
open conformation in ≈ 500ns of simulation time. This result was
also repeated by Palor et al., using a 148 POPC lipid bilayer, TIP3
water molecules and also the CHARMM36 force-field, in 500ns un-
biased molecular dynamics.

Here, we have conducted umbrella sampling simulations for
Potential of Mean Force (PMF) calculations, spanning the
Phe58:Phe126 Cα distance (d) from 0.5nm to 2nm to induce the
opening transition of a single CD81 tetraspanin in water, without
the bilayer. Molecular dynamics simulations for the opening path
were biased with harmonic restrain of the form \(\frac{1}{2}k(d - d_0)^2\),
where d is the Phe58:Phe126 Cα distance, k is the force constant
(in kJ/mol/nm²) and d_0 is the equilibrium value for each umbrella
sampling window. See supplementary information for technical
details on umbrella sampling simulations. Panel 1a shows the
free energy profile. The free energy landscape is dominated by
an absolute free energy minimum with two wells at d≈0.5nm and
d≈0.65nm. There are two other local minima at d≈1nm and
d>2nm, where feasible although metastable conformations
can exist. We have measured the angle (θ) formed between Cα
atoms of residues Val114, Val136 and Gln88 to establish effective
disengagement of the extracellular large and a small loops during
the opening transition (see snapshots insets in panel 1a with Cα
atoms of Val114, Val136 and Gln88 residues highlighted as orange
spheres).

Although conformations with distances well into open-state val-
ues were visited during umbrella sampling (d≈2nm), θ remained
in the [70,90] degrees interval (see panel 1b), indicating rather
poor articulation of the Val114:Val136:Gln88 hinge and not a com-
plete transition into the open state. Panel 1b maps all pairs of
distance-angle points for each conformation adopted during all
umbrella windows. Although most conformations remain in the
[80,100] degrees region, some states with larger angles exist
(≈140 degrees). However, these conformations correspond to dis-
tances d≈1.6nm in the highest region of the energy profile which
belongs to transient states less likely to produce biologically repre-
sentative structures.

We also performed μs-length unbiased atomistic molecular dy-
namics of a single CD81 tetraspanin in water. Panel 1c shows all
distance-angle points for each conformation visited during this
simulation time. Averaged distributions for the distance and the
angle are indicated in panel 1d and the averaged distance \(<d>\)
over time is plotted in panel 1f. The combined observation of
distance and angle θ shows that the opening event does not
completely occur for a single CD81 tetraspanin in water with no
bilayer. Besides, after 1μs of unbiased molecular dynamics the
averaged distance \(<d>\) hardly reaches ≈0.6nm, indicating that
conformations have not yet passed the second well of the absolu-
t minimum (d≈0.65nm). Panel 1e shows molecular dynamics
snapshots of the initial (t=0μs) and final (t=1μs) states of the un-
biased simulation. Consequently, we suggest that the large confor-
mational change needed to fully open one CD81 tetraspanin might
be highly facilitated by protein-lipid interactions, in agreement to
others observations.

To evaluate whether this opening event could also driven by the
interactions with other CD81 tetraspanins, we have performed
additional unbiased atomistic simulations including more CD81
tetraspanins. We have run unbiased molecular dynamics for two
and four CD81 equidistant replicas solvated in explicit solvent and
initially disposed in line (see figure 2). Again, each structure
corresponded to the full-length human CD81 tetraspanin (PDB
ID 5tcx) in its closed conformation, after removing cholesterol
molecules (apoproteins).

Fig. 2 Unbiased atomistic molecular dynamics for multiple CD81 tetraspanins: Top: 2 CD81 tetraspanins. Bottom: 4 CD81 tetraspanins. a) and d) Phe58:Phe126 averaged distance. b) and e) Averaged distribution of the angle and the distance. c) and f) Molecular dynamics snapshots showing final (t=100ns) states. CD81 tetraspanins are represented in black, red, green and blue ribbons, with Phe58 and Phe126 highlighted in vDW representations following the Corey-Pauling-Koltun (CPK) colouring convention for distinguishing atoms. Water molecules are not shown.
Panel 2a plots the Phe58:Phe126 averaged distance of a system including 2 identical CD81 tetraspanins initially separated. Importantly, one of the two tetraspanins reaches distances $d \approx 0.75\text{nm}$ (black curve) in a relative short simulation time (more than an order of magnitude less, in comparison to figure 1f). Panel 2b maps averaged distributions for $d$ and $\theta$ of all conformations visited during these unbiased simulations, see supplementary information for 2D scattered maps. Panels 2c depicts final (t=100ns) molecular dynamics snapshots highlighting the plasticity of $\gamma$ and $\delta$ loops (res 158-174 and res 176-189 respectively) during this first-stage opening (see supplementary information for the initial configuration). Overall, the presence of another protein facilitates an initial transition of one of them with much more easiness than the single-tetraspanin system. Undoubtedly, the free energy landscape may be also modified by the presence of second tetraspanin, simplifying an initial opening transition.

For the system including 4 CD81 tetraspanins in water (yet, with no bilayer) the effect is even more clear. Right after minimization and equilibration (t=0ns) one of the four tetraspanins is already in its opening transition, with $d \approx 0.65\text{nm}$, (panel 2d green curve). Panel 2e maps averaged distributions for $d$ and $\theta$ of all conformations visited during these unbiased simulations, see supplementary information for 2D scattered maps. It can be observed that the cooperative interactions between four identical CD81 induce one CD81 to spontaneously increase its $<d>$ distance (see green curve in the distance distribution in panel 2e) accompanied by a marginal increase in $\theta$ (see green curve in the angle distribution in panel 2e). Panel 2f show a significant change of conformation in the $\gamma$ and $\delta$ loops, that simultaneously pull while increasing Phe58:Phe126 distance ($d$) (see supplementary information for the initial configuration).

These results are consistent with the molecular mechanism proposed by Lang and collaborators that the $\delta$-loop flexibility is a key element to enhance CD81:CD81 direct interactions$^{62}$ and though to large-scale protein oligomerisation$^{34,35}$. Additionally, Cunha et al. showed that open-closed conformational plasticity is pH modulated$^{16}$ and suggested that this conformational dependence is mediated by the glycoprotein E2 in the Hepatitis C virus. Therefore, we propose that outside the lipid bilayer the opening transition is enhanced by the interplay between CD81 tetraspanins, being the event a cooperative phenomenon.

### 2.2 Segregation and opening are coupled events

To study large protein-membrane systems, coarse-grained molecular dynamics have shown to be an adequate tool, in particular, to observe emergent properties difficult to anticipate at lower scales$^{46,49,63}$. Here, we have performed $\mu$s-length unbiased coarse-grained molecular dynamics separately for one, two, three and four CD81 embedded in a 1024 DPPC:DOPC:CHOL (0.58:0.12:0.3) lipid bilayer, solvated in explicit solvent. This ternary bilayer follows a suitable mixture used before to calculate the bending modulus in multi-component lipid membranes in different thermodynamic phases$^{45}$. Therefore including a saturated component with dipalmitoylphosphatidylcholine (DPPC), an unsaturated one with dioleoylphosphatidylcholine (DOPC) and (most importantly for CD81 dynamics) cholesterol (CHOL).

We have observed that CD81 tend to bind together, keeping one tetraspanin always segregated when more than 2 of them are present in a $\approx 16\times16\text{nm}$ bilayer patch. Figure 3 shows Radial Distribution Functions (RDF) of the protein–protein distances (panels 3a,3b and 3c) and the total number of direct contacts between CD81 (panels 3d,3e and 3f), for system setups containing 2, 3 and 4 CD81 inserted in the bilayer. In all cases RDFs have been calculated based on protein centres of mass (COM) and the number of CD81:CD81 direct interactions accounted only for those contacts $<0.6\text{nm}$. These measurements were performed over 5$\mu$s of unbiased simulations.

In panel 3a the RDF of the distance between the first (black protein) and second (red protein) tetraspanin is plotted, showing its peak at $\approx 2.5\text{nm}$. As expected, for $n=2$ both CD81 bind together exhibiting a large number of direct protein-protein contacts over time (panel 3d). In panel 3b the RDF of the distance between the third (green protein) and the first (black protein) tetraspanins has its maximum at $\approx 6\text{nm}$ (green curve). The same happens for the third (green protein) and the second (red protein) tetraspanin (red curve). These RDFs indicate that the third tetraspanin (green protein) is predominantly far away from tetraspanin 1 (black protein) and 2 (red protein), as confirmed by counting the number of direct contacts (panel 3e), where no green neither red curves exist. On the other hand, the black curve shows that tetraspanins 1 (black protein) and 2 (red protein) are bound together with a maximum RDF of their protein-protein COM distance $\approx 2.5\text{nm}$, just like in the system with two tetraspanins inserted in the bilayer (panel 3a). Again, this result is verified by the number of direct contacts in panel 3e, (black curve).
Analogously, in panel 3c blue, cyan and turquoise curves show the RDFs of the distances measured from the fourth tetraspanin (blue protein) to the other three ones (black, red and green proteins). In all three cases the RDFs maxima are $\approx 7-8$ nm, indicating that the fourth tetraspanin (blue protein) remains significantly separated from the other tetraspanins during the majority of the simulation time. On the contrary, the other 3 tetraspanins have distance RDF maxima at lower values, revealing that they are permanently bound together (black, red and green curves). Panel 3f confirms these RDF results by measuring $<0.6$ nm CD81:CD81 contacts for $n=4$. In the snapshot inset it is observed that 3 tetraspanins form an assembly (black, red and green proteins) leaving one tetraspanin (blue protein) out of their group. Accordingly, no contacts $<0.6$ nm between the blue tetraspanin and any other one exist (no blue, nor cyan, nor turquoise curves in panel 3f).

Remarkably, the tetraspanin that is segregated in systems with $n>2$ appears to more readily adopt an open conformation than the ones bound together. Figure 4 shows Phe58:Phe126 distances for each CD81 in the same 4 simulation setups: the lipid bilayer containing either 1, 2, 3 or 4 identical CD81. Panels 4c and 4d show one CD81 with significantly larger Phe58:Phe126 distance maintained during 5$\mu$s of unbiased simulations. Panel 4c shows Phe58:Phe126 distance (green curve) during the segregation of the third tetraspanin (green protein) and panel 4d the Phe58:Phe126 distance (blue curve) during the segregation of fourth tetraspanin (blue protein). These two tetraspanins (green and blue proteins) are the same ones that systematically locate apart from the others in previous figure 3 for $n=3$ and $n=4$.

In summary, we have observed again a cooperative phenomenon, this time for CD81 inserted in a ternary bilayer including cholesterol. Although proteins tend to cluster in all cases, when $n>2$ a single tetraspanin is regularly left apart. Notably, that segregated tetraspanin exhibits a significant increase in its Phe58:Phe126 distance showing a first-stage or partial-switch to an open conformation. Such collective protein-protein partnership is again in line with the capabilities of tetraspanins to oligomerise via the flexibility of the external loops, forming tetraspanin webs and organizing themselves in homomeric clusters.

2.3 CD81 sequesters membrane cholesterol and inhibits large deformations

![Figure 4](image)

**Figure 4** Phe58:Phe126 distances for each individual CD81 embedded in the membrane. Measured between BB beads in Phe58 and Phe126 residues (see inset in panel b) along 5$\mu$s of unbiased simulations for 4 different systems with increasing amounts of CD81: a) $n=1$, b) $n=2$, c) $n=3$ and d) $n=4$. All molecular dynamics snapshots show proteins in surface representations in black, red, green and blue. The lipid bilayer is grey with PO4 beads highlighted in yellow. Top and side views are provided. For clarity, water molecules are not shown.

![Figure 5](image)

**Figure 5** PMF calculations for membrane bending in the $\Psi$ space. Membrane-only (magenta, $n=0$) and membrane with a single CD81 (black, $n=1$). Error bars are standard errors calculated by individually splitting the profiles in independent blocks. Insets show the bilayer with (top) and without (bottom) CD81. Snapshots are averaged densities for all lipid molecules showing steady-state curvature for $\Psi=-0.49$. See supplementary information for PMF convergence details.

Like other members of the tetraspanin superfamily (i.e. CD9), it has been suggested that CD81 could influence the curvature of the cell membrane. Here, to induce bilayer curvature and to quantify its associated free-energy cost, we have used a collective variable specifically design for that scope, which we have already used in other proteins adsorbed on the membrane surface. The collective variable, defined $\Psi$, forces lipid molecules to increase their local density at the centre of the bilayer, inducing the membrane to make slight displacements along its normal axis ($\Psi$). This rearrangement of lipids bends the membrane in a saddle-like shape which emerges spontaneously. Unlike other approaches, $\Psi$ does not allow for a direct control of the bilayer curvature but is rather an indirect mechanism analogous to the protein-membrane surface interplay, as previously described for $\alpha$-
synuclein$^{13,36}$ and the N-BAR domain$^{52}$, both known for inducing curvature in lipid bilayers$^{57,74}$. Besides, the complex saddle-shape mode in which the bilayer bends follows two principal curvatures defined by two fitted circles with different radii$^{3,4,22}$ and requires significantly more energy than other modes of simpler geometry. Mathematically, $\Psi$ is a dimensionless reaction coordinate defined by eq. 1:

$$\Psi = \frac{\sum_{i=1}^{N} \tanh \left( \frac{1}{2} \sqrt{d_{x,i}^2 + d_{y,i}^2 + d_{z,i}^2} \right) - N_0}{N - N_0}$$

(1)

Here, the sum ranges over the centres of mass of every lipid molecule in the bilayer ($N = 1024$). Distances $d_{x,i}$, $d_{y,i}$ and $d_{z,i}$ are the XYZ components of the three-dimensional distance between the centre of mass of bilayer and the centre of mass of lipid molecule $i$. Constant $N_0$ is the average equilibrium value of the summation in an unbiased simulation of a flat bilayer (for this bilayer, $N_0=991.771$). For technical details on the development and implementation of $\Psi$, see Masone et al.$^{52}$.

![Mean Square Displacement (MSD) for different groups of cholesterol molecules](image)

**Fig. 6 Transmembrane region in CD81 captures cholesterol molecules.** a) Mean square displacement of all cholesterol molecules in the bilayer (black) and cholesterol molecules inside the protein pocket (red). b) One CD81 (yellow) with cholesterol molecules (cyan) bound to its transmembrane domain and the detail of cholesterol molecules sequestered inside the protein pocket.

By exploring the phase space in $\Psi$, we have found that a single CD81 tetraspanin increases the free energy cost to bend the bilayer in $\approx 200$ k$_B$T (see figure 5, black curve) with respect to the same value of $\Psi \approx 0.49$ in an identical bilayer-only system (magenta curve). Given that $\Psi$ is a function of the membrane (not the protein) any changes in the free energy are exclusively due to the interference of CD81 in the bilayer dynamics.

The reason for this free-energy increment during bending is explained by the strong interactions taking place between CD81 transmembrane region and the surrounding cholesterol molecules. One single tetraspanin transmembrane domain captures between 11 and 13 cholesterol molecules that remain strongly attached to the protein surface (see panel 6b) and are unable to move along the normal axis to the membrane, though resisting to the curvature mechanism imposed by the reaction coordinate. Moreover, at least 3 of these cholesterol molecules remain trapped inside the protein pocket (see detail in panel 6b).

To quantitatively demonstrate this effect, we have calculated the Mean Square Displacement (MSD) for different groups of cholesterol molecules (all cholesterol molecules in the bilayer and cholesterol molecules inside the protein binding pockets, see panel 6a) to estimate their degree of adherence to CD81 in unbiased 5µs-length simulations. Then, the cholesterol lateral diffusion coefficient in [nm$^2$/ms] can be extracted by fitting a straight line to the MSD at the linear region, in other words, the lateral diffusion coefficient is the slope of the linear region of MSD plot. Comparing MSDs in panel 6a, it is observed that cholesterol molecules inside the protein binding pockets exhibit much lower values (red curve), though a much lower lateral diffusion coefficient is expected when compared to the complete set of cholesterol molecules in the bilayer (black curve), quantitatively verifying the confinement phenomenon.

![Four membranes with increasing amounts of CD81 (n=1, 2, 3 and 4) under identical restrain conditions](image)

**Fig. 7 Four membranes with increasing amounts of CD81 (n=1, 2, 3 and 4) under identical restrain conditions ($\Psi = -0.49$).** Averaged cholesterol-only density maps depicting how the membrane is unable to bend in its characteristic saddle-shape, when 2 or more CD81 are inserted in the bilayer ($n \geq 2$).

For more CD81 tetraspanins in the membrane the bending-resistant effect increases drastically. As more cholesterol molecules strongly bind to more transmembrane domains, more difficult is for the membrane to bend as a response to the local density increase imposed by $\Psi$. Without increasing the force constant used by the reaction coordinate, with two, three and four CD81 the bilayer is unable to bend. Figure 7 shows the lipid bilayer with $n=1$, 2, 3 and 4 CD81 under the same harmonic restrain applied with $\Psi$. The saddle-shaped curvature emerges only for $n=1$ and there is no significant permanent curvature for $n \geq 2$. As expected, the density control strategy used by $\Psi$ to successfully induce curvature in membrane-only systems and in membranes with surface proteins$^{13,52}$ is almost nullified here by the strong protein-cholesterol binding and further confinement inside the binding pockets. This is observed for more than one CD81 inserted in this $\approx 16\times16$nm
bilayer patch containing 306 cholesterol molecules (see figure 7). To evaluate cholesterol molecules fluctuations in 2D, figure 8 shows density maps for the Root Mean Square Fluctuations (RMSF) of all cholesterol molecules in the bilayer, averaged along 5µs of unbiased molecular dynamics, for 4 setups: the bilayer with 1, 2, 3 or 4 CD81 tetraspans. In all cases, less fluctuations are observed for molecules located near the transmembrane protein domains (violet regions). This result describes the unique mechanism used by CD81 to interact with the bilayer. Not only an amount of cholesterol molecules are slowed down in the surroundings of each protein, but also each CD81 sequesters a small amount of cholesterol inside its binding pocket (see figure 6). This is consistent with other reports highlighting the strong link between cholesterol and tetraspanin biology.\textsuperscript{15,19,64}

4 Computational methods
All simulations were performed with GROMACS-2018.3\textsuperscript{1,58,70} patched with Plumed 2.4.3\textsuperscript{58}, a plug-in for free energy calculations and complex collective variable implementation. The calculation of Ψ was implemented in Plumed, which by using Lepton C++ Mathematical Expression Parser is able to evaluate symbolic expressions, such as the hyperbolic tangent used by Ψ as switching function. Radial distribution functions were calculated with GROMACS built-in gmx rdf while inter-residue distances were measured with gmx mdist. RMSF calculations were performed using gmx rmsf. Coarse-grained bilayers and proteins were prepared in part using CHARMM-GUI web server.\textsuperscript{48} Molecular dynamics snapshots were visualized using Visual Molecular Dynamics (VMD).\textsuperscript{38} Data graphics were plotted with Grace (GRaphing, Advanced Computation and Exploration of data). Figure panels were organized using Inkscape. Density maps from molecular dynamics simulations were generated using Gromaps.\textsuperscript{49}

4.1 Atomistic molecular dynamics
All atomistic molecular dynamics were performed in the isotropic NPT ensemble at T = 303.15K\textsuperscript{42,44,61} using the Nosé-Hoover thermostat\textsuperscript{53} with a relaxation time of 1ps. The compressibility was set to 4.5x10\textsuperscript{−3} bar\textsuperscript{−1} using a Parrinello-Rahman barostat\textsuperscript{56} with a time constant of 5ps and a reference pressure of 1 bar. Hydrogen bond lengths were constrained using the LINCS algorithm.\textsuperscript{60} Lennard-Jones and real-space electrostatic interactions radii were set to 1.2nm. Hydrogen bond lengths were constrained using the LINCS algorithm\textsuperscript{52}. The Verlet cutoff scheme\textsuperscript{54} was used updating the neighbour lists every 20 steps. Particle Mesh Ewald (PME) method\textsuperscript{20} was used for long-range electrostatic interactions. Periodic boundary conditions were applied in all three dimensions. The Transferable Intermolecular Potential with 3 points (TIP3) model was used for water molecules. The chosen force field in all cases was CHARMM36.\textsuperscript{10} For simulations including 1 CD81 tetraspanin solvated in 45000 water molecules the simulation box length resulted in L=11nm (cubic shape). For simulations including 2 CD81 tetraspanin solvated in 510000 water molecules the simulation box length resulted in L=12nm. For simulations including 4 CD81 tetraspanin solvated in 3300000 water molecules the simulation box length resulted in L=22nm. Values were set by the CHARMM-GUI protein solvation algorithm\textsuperscript{41}. Unbiased atomistic simulations including 1 CD81 tetraspanin solvated in explicit TIP3 water run for 1us. Umbrella windows for the free energy profile in figure 1 run for 100ns each, adding a total of 1.5us. For 2 and 4 CD81 tetraspanin solvated in explicit TIP3 water, unbiased simulations run for 100ns. All atomistic production runs used a time step of 2fs and collected data every 50000 steps.

4.2 Coarse-grained molecular dynamics
The MARTINI coarse-grained model\textsuperscript{50} is widely used for protein-lipid molecular modelling.\textsuperscript{14,26,31,40,71} In this particular reduced resolution space, beads are classified according to their mutual interactions as: polar, non-polar, apolar and charged. Cholesterol molecules are represented by 8 beads (6 modelling the sterol and 2 the tail) while keeping a mainly planar geometry. Phosphatidylcholine (PC) headgroups include two hydrophilic groups: choline...
charged positively and phosphate charged negatively. All production molecular dynamics simulations were run in the semi-isotropic NPT ensemble at T = 303.15K with the V-rescale thermostat and a relaxation time of 1ps. The compressibility was set to 3x10⁻⁴ bar⁻¹, using a Parrinello-Rahman barostat with a time constant of 12ps. Long-range reaction field electrostatic interactions were used with a Coulomb cut-off set to 1.1nm. Before production runs, all systems were minimized and equilibrated for at least 100ns to generate properly relaxed initial configurations. Periodic boundary conditions were applied in all three dimensions. For equilibration runs the pressure was set at 1.0 bar using the Berendsen barostat with a 5ps coupling constant. In all cases we used the polarisable water model for MARTINI. All bi-layers containing 1024 molecules were solvated in ~18x10⁶ water molecules, satisfying the ample water condition for MARTINI. In the coarse-grained space the simulation box dimensions resulted in Lx=16nm, Ly=16nm and Lz=12nm. All unbiased simulations in the coarse-grained space run for 5us each. Umbrella windows for the free energy profile in figure 5 run for 100ns, adding a total of 5.1us. As suggested for MARTINI, all coarse-grained production runs used a time step of 20fs and collected data every 5000 steps.

Due to particular artifacts pointed out in the MARTINI model regarding the overestimation of protein-protein and protein-membrane interactions, we have back-converted final configurations (t=5μs) to atomic resolution using the CHARMM-GUI all-atom converter for MARTINI. We have applied this conversion to systems containing 1, 2, and 4 CD81 tetraspans inserted in the DPPC:DOPC:CHOL bilayer fully solvated in water, following the same arrangement used along the atomic simulations to verify their physical feasibility. Once in the atomic space, all configurations were relaxed with a short minimization using the steepest descent method and subsequently equilibrated under NVT and NPT conditions with time steps of 1fs and 2fs. Plots of the area dimensions. For equilibration runs the pressure was set at 1.0 bar using the Berendsen barostat with a 5ps coupling constant. In all cases we used the polarisable water model for MARTINI. All bi-layers containing 1024 molecules were solvated in ~18x10⁶ water molecules, satisfying the ample water condition for MARTINI. In the coarse-grained space the simulation box dimensions resulted in Lx=16nm, Ly=16nm and Lz=12nm. All unbiased simulations in the coarse-grained space run for 5us each. Umbrella windows for the free energy profile in figure 5 run for 100ns, adding a total of 5.1us. As suggested for MARTINI, all coarse-grained production runs used a time step of 20fs and collected data every 5000 steps.

4.3 PMF calculations

PMF profiles were computed with umbrella sampling and were recovered using the Weighted Histogram Analysis Method (WHAM) in GROMACS as gmx wham and also with the WHAM implementation by Prof. Grossfield version 2.0.9.1. Convergence was assessed by repeating these calculations on consecutive trajectory blocks. For technical details on umbrella sampling windows distribution and force constants values see supplementary information.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Supercomputing time for this work was provided by CCAD (Centro de Computación de Alto Desempeño de la Universidad Nacional de Córdoba) and by SNCCAD-MinCyT Iniciativa de Proyectos Acelerados de Cálculo (IPAC), grant numbers: 2017-75-APN-SECACT#MCT, 2018-6-APN-SECACT#MECCYT and 2019-72-APN-SECACT#MECCYT. Also, grants from CONICET (PIP-13CC001), ANPCyT (PICT2017-1002) and GPU hardware by the NVIDIA Corporation are gratefully acknowledged.

Notes and references

1. Mark James Abraham, Teemu Murtola, Roland Schulz, Szilárd Páll, Jeremy C. Smith, Berk Hess, and Erik Lindahl. Gromacs: High performance molecular simulations through multi-level parallelism from laptops to supercomputers. SoftwareX, 1-2:19–25, 2015.
2. Riccardo Alessandri, Paulo C. T. Souza, Sebastian Thallmair, Manuel N. Melo, Alex H. de Vries, and Siewert J. Marrink. Pitfalls of the martini model. J. Chem. Theory Comput., 15(10):5448–5460, October 2019.
3. Haleh Alimohamadi and Padmini Ranganami. Modeling membrane curvature generation due to membrane-protein interactions. Biomolecules, 8(4), 2018.
4. P. Bassereau, Benoît Sore, and Aurore Lévy. Bending lipid membranes: experiments after w. helfrich’s model. Advances in colloid and interface science, 208:47–57, 2014.
5. H J C Berendsen, J P M Postma, W F van Gunsteren, A DiNola, and J R Haak. Molecular dynamics with coupling to an external bath. J. Chem. Phys., 81(8):3684, 1984.
6. Hugues Bersini. Emergent phenomena belong only to biology. Synthese, 185(2):257–272, March 2012.
7. C. Boucheux and E. Rubinstein. Tetraspans. Cellular and Molecular Life Sciences CMLS, 58(9):1189–1205, August 2001.
8. Claude Boucheux, Guy Huynh Thienn Duc, Claude Jasmin, and Eric Rubinstein. Tetraspans and malignancy. Expert Reviews in Molecular Medicine, 3(4):1–17, 2001.
9. Rodolfo Briones, Christian Blau, Carsten Kutzner, Bert L. de Groot, and Camilo Aponte-Santaramaría. Gromaps: A gromacs-based toolset to analyze density maps derived from molecular dynamics simulations. Biophysical Journal, 116(1):4–11, January 2019.
10. BR Brooks et al. Charmm: The biomolecular simulation program. Journal of computational chemistry, 30(10):1545–1614, July 2009.
11. Giovanni Busi, Davide Donadio, and Michele Parrinello. Canonical sampling through velocity rescaling. The Journal of Chemical Physics, 126:014101, 2007.
12. Carlos Cabanas, Maria Yanez-Mo, and Annemiek B. van Spriel. Editorial: Functional relevance of tetraspans in the immune system. Frontiers in Immunology, 10:1714, 2019.
13. Marcelo Caparotta, Diego M. Bustos, and Diego Masone. Order-disorder skewness in alpha-synuclein: a key mechanism to recognize membrane curvature. Phys. Chem. Chem. Phys., 22:5255–5263, 2020.
14. Marcelo Caparotta, Claudia N. Tomes, Luis S. Mayorga, and Diego Masone. The synaptogamin-1 c2b domain is a key regulator in the stabilization of the fusion pore. J. Chem. Theory Comput., 16(12):7840–7851, December 2020.
15. Stéphanie Charrin, Serge Manié, Christoph Thiele, Martine Billard, Denis Gerlier, Claude Boucheux, and Eric Rubinstein. A physical and functional link between cholesterol and tetraspans. Eur. J. Immunol., 33(9):2479–2489, September 2003.
16. Eva S. Cunha, Pedro Sfriso, Adriana L. Rojas, Pietro Roveri, Adam Hospital, Modesto Orozco, and Nicola G. A. Abrescia. Mechanism of structural tuning of the hepatitis c virus human cellular receptor cd81 large extracellular loop. Structure, 25(1):53–65, January 2017.
17. Selma Dahmane, Christine Doucet, Antoine Le Gall, Célia Chamontin, Patrice Dosset, Florent Murcy, Laurent Fernan-
dez, Desiree Salas, Eric Rubinstein, Marylène Mougel, Marcelo Nollmann, and Pierre-Emmanuel Milhiet. Nanoscale organization of tetraspanins during HIV-1 budding by correlative dstorm/afm. *Nanoscale*, 11(13):6036–6044, 2019.

18 de Jesus, J. Armando, Noah Kastelowitz, and Hang Yin. Changes in lipid density induce membrane curvature. *RSC advances*, 3(33):13622–13625, September 2013.

19 Alix Delaguillaumie, Julie Harriague, Sylvie Kohanna, Georges Bismuth, Eric Rubinstein, Michel Seigneuret, and Hélène Conjeaud. Tetraspanin cD22 controls the association of cholesterol-dependent microdomains with the actin cytoskeleton in T lymphocytes: relevance to co-stimulation. *J. Cell Sci.*, 117(22):5269, October 2004.

20 U Essmann, L. Perera, M. L. Berkowitz, T. Darden, H. Lee, and L. G. Pedersen. A smooth particle mesh ewald method. *J. Chem. Phys.*, 103:8577–8593, 1995.

21 Luise Florin and Thorsten Lang. Tetraspanin assemblies in virus infection. *Frontiers in immunology*, 9:1140–1140, May 2018.

22 Felix Frey and Timon Idema. More than just a barrier: using physical models to couple membrane shape to cell function. *Soft matter*, January 2021.

23 Michaela Frolikova, Pavla Manaskova-Postlerova, Jiri Cerny, Jana Jankovicova, Ondrej Simonik, Alzbeta Pohlva, Petra Secova, Jana Antalikova, and Katerina Dvorakova-Hortova. Cd9 and cd81 interactions and their structural modelling in sperm prior to fertilization. *International journal of molecular sciences*, 19(4):1236, April 2018.

24 Alan Gross. Wham: the weighted histogram analysis method. [http://membrane.urmc.rochester.edu/wordpress/](http://membrane.urmc.rochester.edu/wordpress/).

25 Yehya Homsi, Eric Rubinstein, Michel Seigneuret, and Hélène Conjeaud. Membrane remodeling and stimulation of aggregation following alpha-synuclein adsorption to phosphotidylserine vesicles. *J. Phys. Chem. B*, 125(6):1582–1594, February 2021.

26 Nathan H. Roy, Jurgen Schneider-Schaulies, Pierre-Emmanuel Milhiet, and Markus Thali. HIV-1 assembly differentially alters dynamic phases. *PLOS Pathogens*, 9(9):3866–3871, 2013.

33 Nikolas Hochheimer, Ricarda Sies, Anna C. Aschenbrenner, Dirk Schneider, and Thorsten Lang. Classes of non-conventional tetraspanins defined by alternative splicing. *Scientific Reports*, 9(1):14075, October 2019.

34 Yahya Homsi and Thorsten Lang. The specificity of homomeric clustering of cd81 is mediated by its δ-loop. *FEBS Open Bio*, 7(2):274–283, February 2017.

35 Yahya Homsi, Jan-Gero Schloetel, Konstanze D. Scheffer, Thomas H. Schmidt, Nicolas Destainville, Luise Florin, and Thorsten Lang. The extracellular delta-domain is essential for the formation of cd81 tetraspanin webs. *Biophysical journal*, 107(1):100–113, July 2014.

36 Brandon M. Hoover, Zhizhang Shen, Curran G. Gahan, David M. Lynn, Reid C. Van Lennep, and Regina M. Murphy. Membrane remodeling and stimulation of aggregation following alph-synuclein adsorption to phosphotidylserine vesicles. *J. Chem. Theory Comput.*, 6(12):3713–3720, December 2010.

37 William Humphrey, Andrew Dalke, and Klaus Schulten. VMD: a visualization tool for exploring molecular dynamics. *Journal of Molecular Graphics*, 14:33–38, 1996.

38 Sunhwan Jo, Tae Hoon Kim, and Wonpil Im. Automated builder of database of protein/membrane complexes for molecular dynamics simulations. *PLoS ONE*, 2(9):e880, August 2007.

39 Jochen S. Hub, Bert L. de Groot, and David van der Spoel. Wham—a free weighted histogram analysis implementation including robust error and autocorrelation estimates. *J. Chem. Theory Comput.*, 6(12):3713–3720, December 2010.

40 Sunhwan Jo and et al. Charmm-gui: A web-based graphical user interface for charm. *J. Chem. Theory Comput.*, 6(12):3713–3720, December 2010.

41 Sunhwan Jo and et al. Charmm-gui: A web-based graphical user interface for charmm. *J. Chem. Theory Comput.*, 6(12):3713–3720, December 2010.

42 Sunhwan Jo, Tae Hoon Kim, and Wonpil Im. Charming alpha-synuclein adsorption to phosphotidylserine vesicles. *Biophysical Journal*, 100(8):1940–1948, 2011.

43 Sunhwan Jo, Tae Hoon Kim, and Wonpil Im. Automated builder and database of protein/membrane complexes for molecular dynamics simulations. *PLoS ONE*, 2(9):e880, August 2007.

44 Sunhwan Jo, Tae Hoon Kim, Vidyashankara G. Iyer, and Wonpil Im. Charming-gui: A web-based graphical user interface for charmm. *Journal of Computational Chemistry*, 29(11):1859–1865, 2008.

45 Sunhwan Jo, Joseph B Lim, Jeffery B Klauda, and Wonpil Im. Charming-gui membrane builder for mixed bilayers and its application to yeast membranes. *Biophysical Journal*, 97(1):50–58, April 2009.

46 George Khelashvili, Benjamin Kollmitzer, Peter Heftberger, Georg Pabst, and Daniel Harries. Calculating the bending modulus of multicomponent lipid membranes in different thermodynamic phases. *Journal of chemical theory and computation*, 9(9):3866–3871, 2013.

47 Heidi Koldso, David Shorthouse, Jean Helie, and Mark S. P. Sansom. Lipid clustering correlates with membrane curvature as revealed by molecular simulations of complex lipid bilayers. *PLOS Computational Biology*, 10(10):1–11, 10 2014.

48 Sunhwan Jo, Joseph B Lim, Jeffery B Klauda, and Wonpil Im. Charming-gui membrane builder for mixed bilayers and its application to yeast membranes. *Biophysical Journal*, 97(1):50–58, April 2009.
P. A. Kollman. The weighted histogram analysis method for free-energy calculations on biomolecules. i. the method. J. Comput. Chem., 13:1011–1021, 1992.

Siewert J. Marrink, Valentina Corradi, Paulo C. T. Souza, Helgi I. Ingolfsson, D. Peter Tieleman, and Mark S. P. Sansom. Computational modeling of realistic cell membranes. Chemical reviews, 119(9):6184–6226, May 2019.

Siewert J. Marrink, H. Jelger Risselada, Sergi Yefimov, D. Peter Tieleman, and Alex H. De Vries. The martini force field: Coarse grained model for biomolecular simulations. J. Phys. Chem. B, 111(27):7812–7824, 2007.

Diego Masone and Diego M. Bustos. Transmembrane domain dimerization induces cholesterol rafts in curved lipid bilayers. Phys. Chem. Chem. Phys., 21:268–274, 2019.

Diego Masone, Marina Uhart, and Diego M. Bustos. Bending lipid bilayers: A closed-form collective variable for effective free-energy landscapes in quantitative biology. J. Chem. Theory Comput., 14(4):2240–2245, April 2018.

S. Nosé. A molecular dynamics method for simulations in the canonical ensemble. Mol. Phys., 52(2):255–268, 1984.

Szlárd Páll and Berk Hess. A flexible algorithm for calculating pair interactions on simd architectures. Computer Physics Communications, 184(12):2641–2650, December 2013.

Machaela Palor, Lenka Stejskal, Piya Mandal, Annasara Lenman, Maria Pia Alberione, Jared Kirui, Rebecca Moeller, Stefan Ebner, Felix Messner, Gisa Gerald, Adrian J. Shepherd, and Joe Grove. Cholesterol sensing by cd81 is important for hepatitis c virus entry. Journal of Biological Chemistry, 295(50):16931–16948, December 2020.

M. Parrinello and A. Rahman. Polymorphic transitions in single crystals: A new molecular dynamics method. Journal of Applied Physics, 52(12):7182–7190, December 1981.

Jason D. Perlmutter, Anthony R. Braun, and Jonathan N. Sachs. Curvature dynamics of alpha-synuclein familial parkinson disease mutants: molecular simulations of the micelle-and bilayer-bound forms. The Journal of Biological Chemistry, 284(11):7177–7189, March 2009.

Sander Pronk, Szilárd Páll, Roland Schulz, Per Larsson, Par Bjelkmar, Rossen Apostolov, Michael R. Shirts, Jeremy C. Smith, Peter M. Kasson, David van der Spoel, Berk Hess, and Erik Lindahl. Gromacs 4.5: a high-throughput and highly parallel open source molecular simulation toolkit. Bioinformatics, 29(7):845–854, 2013.

D. C. Rapaport. The Art of Molecular Dynamics Simulation. Cambridge University Press, Cambridge, 2 edition, 2004.

B Roux. The calculation of the potential of mean force using computer simulations. Comput. Phys. Commun., 91:275–282, 1995.

Huan Rui, Kyu Il Lee, Richard W. Pastor, and Wonpil Im. Molecular dynamics studies of ion permeation in vdac. Biophysical Journal, 100(3):602–610, 2011.

Thomas H. Schmidt, Yahya Homsi, and Thorsten Lang. Oligomerization of the tetraspanin cd81 via the flexibility of its δ-loop. Biophysical Journal, 110(11):2463–2474, June 2016.

David Shorthouse, George Hedger, Heidi Koldos, and Mark S. P. Sansom. Molecular simulations of glycolipids: Towards mammalian cell membrane models. SI: Lipids: from (bio)synthesis to function, 120:105–109, January 2016.

Olivier Silvie, Stéphanie Charrin, Martine Billard, Jean-Francois Franetich, Krista L. Clark, Geert-Jan van Gemert, Robert W. Sauerwein, Francois Dautry, Claude Boucheix, Dominique Mazier, and Eric Rubinstein. Cholesterol contributes to the organization of tetraspanin-enriched microdomains and to cd81-dependent infection by malaria sporozoites. J. Cell Sci., 119(10):1992, May 2006.

Jaqueline M. Tarrant, Lorraine Robb, Annemiek B. van Spriel, and Mark Dexter Wright. Tetraspanins: Molecular organisers of the leukocyte surface. Trends in Immunology, 24(11):610–617, 2003.

Christina M. Termini and Jennifer M. Gillette. Tetraspanins function as regulators of cellular signaling. Frontiers in cell and developmental biology, 5:34–34, April 2017.

G. M. Torrie and J.P. Valleau. Nonphysical sampling distributions in monte carlo free-energy estimation: Umbrella sampling. Journal of Computational Physics, 23(2):187–199, 1977.

Gareth A. Trillo, Massimiliano Bonomi, Davide Branduardi, Carlo Camilloni, and Giovanni Bussi. Plumed 2: New feathers for an old bird. Computer Physics Communications, 185(2):604–613, 2014.

Rie Umeda, Yuhkoh Satohu, Mizuki Nakada-Nakura, Kehong Liu, Takeshi Yokoyama, Mikako Shirouzu, So Iwata, Norimichi Nomura, Ken Sato, Masahito Ikawa, Tomohiro Nishizawa, and Osamu Nureki. Structural insights into tetraspanin cd9 function. Nature Communications, 11(1):1606, March 2020.

David Van Der Spoel, Erik Lindahl, Berk Hess, Gerrit Groenhof, Alan E. Mark, and Herman J. C. Berendsen. Gromacs: Fast, flexible, and free. Journal of Computational Chemistry, 26(16):1701–1718, 2005.

Floris J. van Eerden, Tom van den Berg, Pim W. J. M. Fredrix, Djuurre H. De Jong, Xavier Periole, and Siewert J. Marrink. Molecular dynamics of photosystem ii embedded in the thylakoid membrane. J. Phys. Chem. B, pages 3237–3249, September 2016.

Semen Yesylevskyy and Himanshu Khandelia. Encurv: Simple technique of maintaining global membrane curvature in molecular dynamics simulations. J. Chem. Theory Comput., 17(2):1181–1193, February 2021.

Semen O. Yesylevskyy, Lars V. Schafer, Durga Sengupta, and Siewert J. Marrink. Polarizable water model for the coarse-grained martini force field. PLOS Computational Biology, 6(6):1–17, 06 2010.

Ying Yin, Anton Arkhipov, and Klaus Schulten. Simulations of membrane tubulation by lattices of amphiphysin n-bar domains. Structure, 17(6):882–892, 2009.

Xin A. Zhang and Chao Huang. Tetraspanins and cell membrane tubular structures. Cellular and Molecular Life Sciences, 69(17):2843–2852, September 2012.

Brandon Zimmerman, Brendan Kelly, Brian J. Mcmillan, Tom C. M. Seeger, Ron O. Dror, Andrew C. Kruse, and Stephen C. Blacklow. Crystal structure of a full-length human tetraspanin reveals a cholesterol-binding pocket. Cell, 167(4):1041–1051.e11, November 2016.