Lipid packing defects and membrane charge control RAB GTPase recruitment

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1 | INTRODUCTION

RAB proteins are small GTPases of the RAS superfamily that are involved in many steps of transport inside the cell. There are over 60 RAB proteins in humans and they all localize to distinct membrane compartments. RAB proteins which oscillate between an active form (GTP-bound) and an inactive form (GDP-bound) can bind to membranes with the help of their prenyl group (geranylgeranyl group), a posttranslational lipid modification at their C-terminal extremity. The RAB escort protein (REP), known to be involved in RAB prenylation, and the GDP dissociation inhibitor (GDI), known to bind to soluble RABs, are known to play key roles in both the delivery and the recycling of RAB proteins to and from membranes1,2 but cannot account for their specific intracellular localization. Until now, multiple studies have suggested that RAB-specific membrane targeting could be mediated by protein factors such as guanine nucleotide exchange factors (GEF), originally known to activate RABs by nucleotide exchange,3 and GDI displacement factors, thought to influence the release of prenylated RAB proteins from GDI.4,5 Extensive sequence analysis, domain swapping and mutagenesis studies of different RAB proteins have shown that specific domains are involved in RAB targeting to membranes. Pereira-Leal and Seabra6 identified five RAB family regions that distinguish RAB proteins from the other members of the RAS superfamily and four subfamily regions that stand to differentiate each RAB subfamily. Different combinations of mutations of these domains led to mislocalization of the RAB proteins, suggesting that membrane specificity is also determined by specific RAB sequences.7 The hypervariable region of RAB35 has also been shown to be determinant for proper membrane targeting.8

While protein–protein interaction has been widely studied to explain RAB-specific membrane targeting, very little is known about the influence of the membrane itself. Diverging from the initial fluid mosaic model,9 it is now known that membranes are crowded and heterogeneous environments with lipids and proteins diffusing...
laterally allowing the formation of regions which vary in thickness and composition.20 Because of specific lipid metabolism and selective transport, cellular membranes have heterogeneous lipid compositions and can also be characterized by asymmetrical lipid compositions between the two leaflets.11 Similarly to RAB proteins, some lipids localize to specific compartments and thereby also define organelle identity. Because of their diversity in lipid composition, intracellular membranes exhibit different physicochemical properties such as charge, lipid packing defects and membrane curvature.12–14 The negatively charged phosphatidylserine lipid is mostly found at the inner leaflet of the plasma membrane,15 and each membrane of the endocytic pathway but also the trans-Golgi network membrane and the plasma membrane are characterized by specific phosphoinositide content.16,17 The plasma membrane, which contains saturated cylinder-shaped lipid species and high cholesterol levels, is characterized by tight lipid packing. On the other hand, the endoplasmic reticulum (ER) that exhibits high levels of unsaturated cone-shaped lipids and low levels of cholesterol is characterized by loose lipid packing.12 Two distinct membrane territories have therefore been described: the late secretory pathway and the endocytic pathway (plasma membrane, endosomal membranes and trans-Golgi network) characterized by tight lipid packing and a highly charged cytosolic leaflet, and the early secretory pathway (ER and cis-Golgi) characterized by loose lipid packing and a weakly charged cytosolic leaflet. Of note, these varying lipid compositions are also relevant in the context of membrane order. The plasma membrane, which is enriched in saturated lipids and cholesterol, was shown to exist in both liquid-disordered (Ld) and liquid-ordered (Lo) states (so-called raft phase), while intracellular membranes composed of unsaturated lipids and low cholesterol levels rather behave as Ld phases.18,19

Membrane curvature is also a key feature of intracellular membranes as most cellular organelles display regions of both low and high curvatures. For example, the ER is formed of a complex network of interconnected flat sheets and highly curved tubules,20,21 and endosomes display globular (low curvature) and tubular regions.22 External constraints applied, for instance, by the cytoskeleton, protein coats or insertion of amphipathic protein domains can force a lipid bilayer to bend23–25 and induce positive membrane curvature (convex surface detectable by cytosolic proteins), which gives rise to large lipid packing defects.26 Lipid packing defects as well as membrane charge were described to be essential parameters regulating the specific membrane binding of some peripheral proteins.16,26–30

In this work, we investigate the role of membrane physicochemical properties in the binding of RAB proteins using in vitro assays consisting of purified RAB proteins and giant unilamellar vesicles (GUVs) as model membranes of controlled lipid composition.21

2 | RESULTS AND DISCUSSION

Four RAB proteins that localize to distinct membranes in cells were chosen for our study: RAB1 and RAB6 which associate with pre-Golgi and Golgi/trans-Golgi network membranes, respectively, RAB5 which is present on early endosomes and RAB35 which mainly localizes to the plasma membrane22 (see Supporting information).

2.1 | RAB6 specifically localizes to the Ld phase independently of its prenylation state

To test whether RAB proteins show specific recruitment to a given lipid phase, we investigated the recruitment of purified RAB proteins to GUVs exhibiting phase separation between Lo and Ld domains.33 GUVs were formed using a lipid mixture consisting of brain sphingomyelin (BSM), cholesterol and 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC) (3:1:3 molar ratio).34 No binding of unprenylated RAB proteins was observed on these membranes (Figure S1) which is in good agreement with the commonly accepted view that RAB proteins are incorporated into biological membranes through their C-terminal geranylgeranyl groups.35 Most RAB GTPases are diprenylated in the cell with the addition of two geranylgeranyl moieties to the 2 C-terminal cysteines.36 The use of diprenylated proteins (Figure S2A) is technically challenging because of the high affinity of the GDP-bound RAB for the REP,37 which prevents binding to GUV membranes (Figure S3A-(1)). Recruitment of diprenylated RAB6 to membranes exhibiting phase separation was however achieved through the addition of the RAB binding domain of LidA (LidA201–583) and GGTase I. LidA, a RAB6 superrepressor from Legionella pneumophila, outcompetes the REP, while GGTase I shields the prenyl group from the solvent (detailed in Figure S3A). Under these conditions, diprenylated RAB6 was found to clearly segregate to the Ld phase (Figure S3A-(5)). Monoprenylated GDP-bound RAB6 (Figure S2B) also specifically binds to Ld membranes in the presence (Figure S3B) and also in the absence (Figure 1A) of these additional protein factors. These results demonstrate that LidA201–583 and GGTase I do not influence protein membrane binding specificity and that both mono- and diprenylated RAB proteins preferentially bind to Ld membranes. To confirm these results, experiments were performed with GUVs composed of pure Ld phase (DOPC and cholesterol in a 1:1 molar ratio) or with GUVs composed of pure Lo phase (BSM and cholesterol in a 1:1 molar ratio).34 GDP-bound RAB6, independently of its monoor diprenylation status, was only recruited to Ld GUVs but not to Lo GUVs (Figures S3 and 1B). Thus, RAB6 displays similar membrane binding preferences toward Ld domains independently of its mono- or diprenylation status.

Although membrane binding specificity seemed to be unchanged, the kinetics of membrane dissociation were previously described to differ drastically between proteins harboring one or two lipid groups. In vitro studies using synthetic vesicles have indeed suggested that proteins that possess two geranylgeranyl modifications have a half-life of several hours38 whereas monoprenylated proteins usually exhibit a half-life of 1 s or less.39 In the same direction, more recent in vitro studies demonstrated that nonphysiological addition of a second farnesyl group to N-RAS proteins leads to reduced membrane dissociation rate.40 To investigate the kinetics of diffusion and association with membranes of mono- and diprenylated RAB proteins, we have performed fluorescence recovery after photobleaching experiments of GDP-bound RAB6 in complex with LidA201–583 on DOPC-containing GUV membranes. RAB6 recovery by lateral diffusion was assessed by bleaching a small circular region of the GUV on the membrane close to the coverslip. Recovery curves yielded diffusion coefficients $D = 1.2 \pm 0.5 \, \mu m^2/s$ and $D = 1.3 \pm 0.2 \, \mu m^2/s$ for mono- and
diprenylated RAB6, respectively, comparable to the diffusion coefficient of a TexasRed-labeled 1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine (DHPE) lipid under the same conditions \((D = 1.2 \pm 0.3 \, \mu m^2/s)\) (Figure S4A). RAB6 recovery from the bulk was assessed by bleaching the full GUV and yielded recovery half-lives \(t_{1/2} = 62 \pm 18\) seconds and \(t_{1/2} = 414 \pm 205\) seconds for mono- and diprenylated RAB6, respectively (Figure S4B). These results indicate that monoprenylated RAB6 exhibits a faster recovery rate, which is in good agreement with previously published data.38,39

Although the recovery dynamics of mono- and diprenylated RAB6 were found to diverge, we have demonstrated that both were preferentially recruited to the same disordered membrane domains. This suggests that membrane binding kinetics have no influence on membrane binding preferences, making it possible to use monoprenylated RABs for this study.

In our experimental in vitro approach, activation (via nucleotide exchange) of diprenylated RABs leads to the dissociation of the REP–RAB complex and to the subsequent exposure of the hydrophobic prenyl groups to the solvent, thereby causing protein precipitation. In order to overcome this issue, diprenylated RABs can be activated in the presence of membranes which allows the stabilization of the prenyl groups, but consequently prevents any measurement of the amount of active GTP-bound RABs in our system. RAB proteins are however known to localize to the cytosol in their GDP-bound inactive form and to get activated (GTP-bound) by GEFs upon membrane incorporation.41 As membrane-bound RAB proteins are therefore mostly active, we decided for our study to investigate the binding of RAB proteins in their GTP-bound form. Because the efficiency of nucleotide exchange to GTP could not be measured when using diprenylated RAB proteins, this approach was only feasible using monoprenylated RAB proteins (Figure S2B). Therefore, we decided to focus our study on monoprenylated RAB proteins.

Some RAB proteins can undergo additional C-terminal modifications following geranylgeranylation, such as proteolysis and/or carboxyl methylation, depending on their prenylation motif.42,43 RAB carboxyl methylation, which consists in the addition of a carboxyl group to the exposed prenylated cysteine, was shown to enhance the hydrophobicity of the C-terminus and subsequently to increase membrane affinity.43 However, because the absence of methylation was shown to only affect the cycle of RAB membrane/cytosol partitioning, but not their specific membrane localization,43 we did not investigate the potential effects of RAB carboxyl methylation in our in vitro experiments.

2.2 | RAB proteins specifically localize to the Ld phase through their geranylgeranyl group

Monoprenylated GTP-bound RAB6 was, like its GDP-bound counterpart, only recruited to Ld domains on GUVs displaying phase separation (Figure 1A) and recruitment was only observed on Ld vesicles but not on Lo vesicles (Figure 1B), indicating that the binding specificity of RAB6 is also independent of its activation state. Similarly to RAB6, monoprenylated and activated RAB1 and RAB5 segregated specifically to Ld domains on GUVs displaying phase separation (Figure 1A) and recruitment was only observed on Ld vesicles but not on Lo vesicles (Figure 1B).
We next investigated whether the prenyl group plays a direct role in the specific recruitment of RAB proteins to the Ld phase. For that purpose, we looked at the recruitment of glutathione S-transferase (GST) to which a CAAX prenylation motif (CVIL) was added at its C-terminus. The purified and fluorescently labeled protein was enzymatically monoprenylated using the same protocol that for the RAB proteins. As shown in Figure 1, monoprenylated GST also specifically segregated to Ld domains. On the other hand, no recruitment of unprenylated GST could either be detected on Lo or Ld domains (Figure S1), confirming that the prenyl group is required and sufficient for GST membrane insertion.

Altogether, the above results suggest that the recruitment of RAB proteins to Ld membranes is mediated by the geranylgeranyl moiety. Because of its bulky C20 isoprenoid highly unsaturated chain structure, the geranylgeranyl moiety (Figure S2C) might only be able to incorporate itself into Ld membranes. In agreement with this, previous studies have demonstrated that lipidated peptides containing isoprenyl groups or unsaturated acyl chains preferentially insert themselves into Ld membranes and show negligible affinity toward Lo membranes. This preference for disordered domains was also shown for RAS proteins containing a C-terminus unsaturated C15 isoprenoid farnesyl group. In contrast, peptides incorporating saturated acyl chains such as palmitoyl were found to be significantly recruited to Lo domains. Similarly, the addition of a saturated C16 palmitoyl group to transmembrane proteins was shown to mediate their dynamic targeting to raft-like Lo phases. Thus, a likely hypothesis is that the isoprenoid unsaturated structure of prenyl groups favors their insertion into Ld membranes.

2.3 RAB35 membrane recruitment is driven by both the charged hypervariable region and the prenyl group

The great majority of RAB GTPases, including the previously tested RAB1, RAB5, and RAB6, are found associated with intracellular membranes. RAB35, on the other hand, was shown to localize to intracellular endocytic compartments and also to the plasma membrane. Thus, we wondered whether RAB35 membrane binding was governed by a similar mechanism. We first tested the recruitment of monoprenylated RAB35 to Lo and Ld GUVs. Unexpectedly, RAB35 was not recruited to either of these membranes (Figure 2A), indicating that the prenyl group is not sufficient to drive RAB35 membrane insertion.

Endosomal and plasma membranes are known to be negatively charged because of the large amount of phosphoinositides and phosphatidylserine, anionic lipids known to play major roles in signaling processes and membrane dynamics. RAB35 contains stretches of positively charged residues at its C-terminal region, the last 20 amino acid region being the most charged as compared with that of the other RABs (Table S1). In cellulo studies have shown that this polybasic region is essential for targeting RAB35 to the plasma membrane, indicating that RAB35 localization depends on electrostatic interactions between the negative charge of the inner leaflet of the plasma membrane and the positive charges of the RAB35 C-terminal region. To address the role of electrostatic interactions, we monitored the recruitment of RAB35 to negatively charged GUVs containing 30 mol% 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho-L-serine (POPS). RAB35 was efficiently recruited to POPS-containing vesicles, while no detectable interaction occurred when POPS was replaced by neutral 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC) (Figure 2A). These results demonstrate that RAB35 membrane recruitment is mediated by electrostatic interactions. We then investigated whether the prenyl group is required for RAB35 membrane binding to negatively charged POPS-containing vesicles. No binding of unprenylated RAB35 to charged vesicles was observed (Figure 2A), indicating that both the electrostatic interactions and the prenyl group are necessary for RAB35 recruitment.

In order to assess whether, as for the previously tested RABs, the prenyl group might also mediate specific membrane domain targeting, we monitored the recruitment of RAB35 to anionic Lo and Ld GUVs by replacing cholesterol with negatively charged sulfate cholesterol (see Supporting information). RAB35 membrane binding was now observed not only on Ld vesicles but also on Lo vesicles (Figure 2B). These results in combination with the absence of RAB35 recruitment to neutral Lo and Ld vesicles confirm that RAB35 membrane incorporation requires negatively charged lipids and clearly demonstrate that in some cases the prenyl group is able to interact with Lo membranes. Additionally, we quantified the area density of prenylated RAB35 (Φp) using Equation (1) and observed a 3-fold increase in recruitment to Ld vesicles as compared with Lo vesicles (Figure 2B). This is consistent with the previous observations that prenylated proteins preferentially bind to Ld domains. Taken together, our results suggest that the membrane recruitment of C-terminally charged and prenylated RAB proteins is primarily dependent on the presence of anionic lipids. This specificity for negatively charged membranes gives the ability to RAB35 to overcome the exclusive binding of the prenyl group to Ld domains. This charge dependency is crucial for RAB35 interaction with negatively charged endosomal and plasma membranes. Interestingly, when comparing the charge of the last 20 amino acids of all human RAB proteins, we found that RAB23 and RAB35 display the highest positive charge (Table S1). RAB23 has also been shown to localize to the plasma membrane suggesting that its specific recruitment to the plasma membrane might also be mediated by electrostatic interactions.

Pointing in the same direction as our results, pioneer studies of McLaughlin et al. demonstrated that myristoylated alanine-rich C-kinase substrate interacts with the negatively charged plasma membrane through a myristoyl-electrostatic switch involving the N-myristoyl group and a positively charged protein domain. Additionally, more recent studies showed that the recruitment of proteins of the RAS family (K-RAS4B and RND3) to the plasma membrane can be modulated by electrostatic interactions between the positively charged C-terminus and anionic phospholipid headgroups.

2.4 RAB proteins can sense membrane curvature through their prenyl group

Most RAB proteins, for instance RAB1 or RAB6, are present on transport vesicles which typically have a diameter of 40-60 nm and can thus be regarded as curved membranes. Therefore, we investigated
As a model, we used an optical tweezer setup to pull membrane tubes from L-α-phosphatidylcholine (Egg, Chicken) (EggPC) GUVs (with additional 0.1 mol% of TexasRed-DHPE lipids) (Figure S5A).\textsuperscript{56,57} RAB protein relative enrichment (or sorting) between the highly curved tube and the flat GUV membrane was imaged by confocal microscopy. Tuning membrane tension through micropipette aspiration of the GUV allows us to modulate the tube radius and to measure protein sorting for increasing curvature (up to 1/15 nm\(^{-1}\)). Biological membranes are two-dimensional surfaces with two principal curvatures \(C_1 = 1/R_1\) and \(C_2 = 1/R_2\) (with \(R_1\) and \(R_2\) referred to as the principal radii of curvature) along two perpendicular directions.\textsuperscript{58} The total curvature of the membrane is \(C = C_1 + C_2\). In the case of a spherical vesicle of radius \(R\), the membrane deforms equally in both directions leading to \(C_1 = C_2 = 1/R\) and a total curvature \(C_v = 2/R\). In the case of a cylindrical tube of radius \(R\), which is curved only in one direction and flat in the other, \(C_1 > 0\) and \(C_2 = 0\) yielding a total curvature \(C_v = 1/R\).\textsuperscript{58} A 15 nm radius tube will thus have the same curvature as a 30 nm radius intracellular transport vesicle,\textsuperscript{59} indicating that the typical curvatures in our experiments are biologically relevant.

Curvature sensing was assessed by calculating the sorting ratio (\(S\)) defined as the protein/lipid signal ratio on the tube divided by that of the Epsin N-Terminal Homology (ENTH) domain which is curved only in one direction and flat in the other, \(S = 1 + 1/(R_1C_p\phi_v)\), where \(S\) is the sorting ratio, \(R_1\) is the tube radius, \(C_p\) is the effective spontaneous curvature of the protein and \(\phi_v\) is the protein area fraction which is related to \(\Phi_v\) by \(\Phi_v = \rho \phi_v\) (\(\rho\) is the inverse of the area per protein). The intrinsic curvature radius of the protein \(C_p^{-1}\) can be determined by plotting \((S - 1)\phi_v\) as a function of curvature (1/\(R_1\)) and taking the resulting slope of the linear fit (Figure 3B). \(\Phi_v\) was assessed using Equation (1) and \(\rho\) was estimated by assuming that RAB proteins are spherical proteins of around 25 kDa with a corresponding average radius of 2 nm (\(\rho = 1/12.6\) nm\(^{-2}\)).\textsuperscript{59} \(C_p^{-1}\) values were, respectively, 2.1 ± 0.6, 2.6 ± 0.7 and 1.5 ± 0.5 nm for RAB1, RAB5 and RAB6. RAB proteins interact with membranes through the hydrophobic insertion of their prenyl group into the bilayer while a few amino acid residues close to the prenylation site will be in proximity to the lipid headgroups. Thus, the geometry of the inserted domain may be comparable to that of the Epsin N-Terminal Homology (ENTH) domain which senses membrane curvature through an amphipathic helix, but also to that of lipids with inverted conical shapes, such as lysophosphatidic acids (LPAs). The first was described to exhibit a spontaneous curvature radius of 1.6 nm,\textsuperscript{60} while the latter was shown to generate local positive curvature and to display a spontaneous curvature radius of 2 nm.\textsuperscript{58} The spontaneous curvature values of ENTH domains and LPAs are therefore both very similar to those we obtained for RAB proteins. Our values are also very similar to that of Amphiphysin (1.9 ± 0.4 nm) which was calculated using the same low-density

The influence of membrane curvature on RAB membrane recruitment.

Figure 2 RAB35 membrane binding is both charge and prenyl group dependent. GUVs were incubated with 2 \(\mu\)M GFP-tagged RAB35. A, Neutral Lo and Ld vesicles were formed using 1:1 (molar ratios) of BSM:cholesterol and DOPC:cholesterol, respectively. Negatively charged POPS-containing vesicles were formed using a 30% POPS:30% cholesterol:40% EggPC (mol/mol) mix while neutral POPS-containing vesicles were formed by replacing POPS with POPC. Monoprenylated RAB35 was only recruited to negatively charged POPS-containing GUVs but not to neutral vesicles. Unprenylated RAB35 was not recruited to negatively charged POPS-containing vesicles. B, Negatively charged Lo and Ld vesicles were formed by replacing cholesterol with cholesterol sulfate. Monoprenylated RAB35 was recruited to both negatively charged Lo and Ld vesicles. Quantifications of GFP-RAB35 protein densities (\(\Phi_v\)) show a 3-fold increase in RAB35 recruitment to disordered membranes. (scale bar: 10 \(\mu\)m; *** = t-test, P-value < 0.0001)
FIGURE 3  Prenylated proteins can sense membrane curvature. A highly curved membrane tube was pulled with optical tweezers from an EggPC GUV containing the fluorescent lipid marker TexasRed-DHPE (red) in the presence of 100-300 nM Alexa488 labeled monoprenylated proteins (RAB1, RAB5, RAB6 and GST). A. The plots show the protein sorting ratios (calculated using Equation (2)) as a function of tube curvature (1/Rtube) and each dot represents one sorting measurement at a given tube radius. Data were obtained from 10 (RAB1 and GST) or 7 (RAB5 and RAB6) independent experiments. Each plot was fitted with a linear regression (black line). For all prenylated proteins, sorting increases when the curvature is increased (i.e., when the tube radius is decreased). B. The protein effective spontaneous curvatures were calculated using the theoretical model from Sorre et al.57 in which the sorting ratio S is given by $S = 1 + 1/(RC_p\phi_v)$, where $R_t$ is the tube radius, $C_p$ is the effective spontaneous curvature of the protein and $\phi_v$ is the protein area fraction (related to the protein density $\Phi_v$ by $\Phi_v = \rho\phi_v$, $\rho$ = inverse of the area per protein = 1/12.6 nm$^{-2}$). (S - 1)$\phi_v$ is thus predicted to scale linearly with the tube curvature (1/R) with a slope $C_p^{-1}$. By using the same sorting ratio values as obtained in (A) and measuring the protein density on the GUV, we plotted (S - 1)$\phi_v$ as a function of curvature (1/R). The spontaneous curvature radius value was determined by fitting the plot with a linear regression and extracting the value of the slope. Error bars correspond to the experimental errors of the measurements.

2.5  Prenylated proteins show preferences for lipid packing defects

The Ld phase is characterized by the assembly of unsaturated lipids which are known to promote lipid packing defects.12 Membrane curvature was also shown to lead to the appearance of defects in the arrangement of lipids.12 To explain the preferential binding of RAB proteins to Ld membranes and their sensitivity to curvature, we hypothesized that RAB membrane recruitment is dependent on the presence of lipid packing defects in the bilayer.

To test this hypothesis, we performed recruitment experiments with GUVs containing 15 mol% 1-2-dioleoyl-snglycerol (DOG), a conical-shaped lipid that was shown to induce the formation of packing defects similar to those found on positively curved membranes.63 Control GUVs containing lower amounts of lipid packing defects were composed of 15 mol% DOPC cylindrical lipids (see Supporting information). The membrane recruitment of all monoprenylated proteins was significantly increased in the presence of DOG (Figure 4), that is, in the presence of higher amounts of lipid packing defects.

Unlike DOG, polyunsaturated fatty acids (PUFAs), such as 1-stearoyl-2-docosahexaenoyl-sn-glycero-3-phosphoethanolamine (PUFA PE), were shown to decrease the amount of lipid packing defects, especially in curved membranes.64 We measured RAB and GST binding on GUVs composed of 30 mol% PUFA PE and used as a control GUVs containing higher amounts of packing defects and composed of 30 mol% 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine.
This lipid-driven membrane binding mechanism sheds new light on how RAB GTPases could bind to membranes. Intracellular membranes are mainly composed of Ld phases,18 and many RAB proteins associate with highly curved transport vesicles.54,55 Our hypothesis is that the addition of one or two geranylgeranyl moieties on all RAB proteins serves as a core mechanism to bind them to specific membrane domains displaying lipid packing defects, the specificity for a given compartment (ER, Golgi and endosomes) relying then on other mechanisms such as the presence of specific GEFs.3

An interesting variation to this theme is given by RAB35 which has a positively charged C-terminus and is mainly found at steady state associated with the plasma membrane and the endocytic compartments.47 We showed that RAB35 membrane recruitment is primarily dependent on the presence of negatively charged lipids which are also predominantly found on endosomal and plasma membranes.11 Even though lipid packing defects enhance RAB35 membrane affinity, they are not essential for membrane binding.

In conclusion, our work illustrates that the physicochemical properties of membranes, such as charge distribution and lipid packing defects, could be prime determinants of the localization of RAB proteins to cellular membranes.

4 | MATERIAL AND METHODS

4.1 | In vitro monoprenylation and diprenylation

The prenylation reaction consists in the addition of one or two C20 geranylgeranyl moieties (geranylgeranyl pyrophosphate, GGpp; Sigma) at the C-terminal extremity of the proteins. Prenylation was achieved either through monoprenylation (addition of one geranylgeranyl group) using purified geranylgeranyl transferase type I (GGTase I) (Figure S2B) or diprenylation (addition of two geranylgeranyl groups) using the native prenylation machinery consisting of purified RAB geranylgeranyl transferase (RABGGTase or GGTase II) and REP (Figure S2A).

Monoprenylation reactions were performed at 25 °C for 1.5 hours with a molar ratio of 0.5:1:5 GGTase I, RAB and GGpp. Molar ratios for the diprenylation reaction were 1:5:0.5:0.75 RABGGTase:GGpp: RAB:REP and the reaction was performed at 25 °C for 4 hours. To control efficient protein prenylation, nitrobenzoxadiazole (NBD)-farnesyl pyrophosphate (Jena Bioscience), a C15 fluorescent analog of geranylgeranyl pyrophosphate, was used as described previously.65

4.2 | Giant unilamellar vesicles

GUVs were grown on indium tin oxide (ITO)-coated glass slides using the electroformation technique.66 Fifteen microliters of a 0.5 mg/mL lipid mix was dried on ITO-coated slides for a few minutes at 50 °C and subsequently under vacuum for at least

FIGURE 4 Increasing amounts of lipid packing defects enhance RAB membrane binding. RAB1, RAB5, RAB6 and GST were labeled using Alexa488, monoprenylated and incubated with GUVs at 2 µM final concentration. DOG-containing GUVs with a high density of lipid packing defects were formed using an 85% EggPC:15% DOG (mol/mol) mix. In control GUVs, DOPC replaced DOG. The right panel shows the quantification of protein density on the membrane (ϕ) in both DOPC and DOG containing vesicles. We observed a significant increase in protein recruitment on GUVs with higher levels of lipid packing defects. (scale bar: 10 µm; *** = t-test, P-value < 0.0001; ** = t-test, P-value = 0.0006)

(POPE) (see Supporting information). We found that the membrane recruitment of geranylgeranylated proteins significantly decreases in the presence of PUFA PE (Figure 5), that is, when the amount of packing defects is decreased.

Altogether, the above results suggest that lipid packing defects are drivers of RAB membrane recruitment and that this lipid packing defect sensing is mediated by the C-terminal prenyl group.

3 | CONCLUSION

It has been known for a long time that prenyl groups act as nonspecific membrane anchors but our results, together with recently published data,28,29 highlight a role for prenyl groups (farnesyl and geranylgeranyl) in specific membrane domain targeting. Similarly to what we found with geranylgeranylated RAB proteins, farnesylated N-RAS preferentially binds to Ld domains on flat membranes and its differential membrane recruitment was shown to rely on the presence of lipid packing defects induced by curvature and specific lipid geometrical shapes.28,29 A likely explanation is that prenyl groups are largely unsaturated and have a kinked structure allowing them to get preferentially inserted into membranes containing packing defects such as Ld or curved membranes.
0.1 mg/mL β-Casein to prevent adhesion of the GUV to the glass. Membrane binding was studied using 100-300 nM final concentration of protein.

### 4.4 Measurement of protein density on the membrane

Protein density was assessed as previously described. Briefly, fluorescence was calibrated using GUVs made of EggPC lipids and BodipyFL-C5-1-hexadecanoyl-sn-glycero-3-phosphocholine (HPC), a green fluorescent lipid, at various concentrations. The HPC area density on the GUV (φ<sub>HPC</sub>) can be calculated by assuming that the average area occupied by a single phosphatidylcholine (PC) molecule is 0.7 nm<sup>2</sup>. The fluorescent intensity of this lipid on the GUV membrane was measured (I<sub>HPC</sub> at a given confocal photomultiplier tube detector gain) for each area density. A linear fit of the fluorescence vs area density plot gave the conversion constant (A<sub>gain</sub>) (φ<sub>HPC</sub> = A<sub>gain</sub> × I<sub>HPC</sub>). Proteins were labeled with the Alexa488 fluorophore and lipids with BodipyFL-C5-HPC, two fluorophores exhibiting different spectral properties. Thus, we measured the correction factor F = I<sub>gain</sub>/I<sub>HPC</sub>, that is, the ratio of fluorescence intensities of Alexa488 and HPC, respectively, at a given concentration in solution. Both fluorescent signals in bulk scaled linearly with their concentration and F is defined as the ratio between the slopes of the Alexa488 linear fit and that of HPC. The protein labeling efficiency was taken into account by calculating the degree of labeling (n<sup>+</sup>) of the protein using Equation S1. Protein density on the GUV membrane (φ<sub>prot</sub>) was thus given by

\[
\phi_{prot} = \frac{A_{gain} \times I_{prot}(gain)}{F \times n^+ (gain)}
\]  

### 4.5 Measurement of sorting ratio

In order to quantify protein sorting to the tube, the fluorescence intensity of the Alexa488 labeled protein (I<sub>protein</sub>) was normalized by the intensity of the fluorescent lipid (TexasRed-DHPE, h<sub>lipid</sub>) at each tension step increase. The sorting ratio S corresponds to the ratio between the normalized protein intensity on the tube and the same normalized intensity on the GUV (Figure S5B):

\[
S = \frac{I_{protein}/h_{lipid}}{I_{protein}/h_{lipid}_{tube}}
\]

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Conflict of interest
The authors declare no potential conflict of interests.

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