Three Separable Domains Regulate GTP-Dependent Association of H-ras with the Plasma Membrane

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The microlocalization of Ras proteins to different microdomains of the plasma membrane is critical for signaling specificity. Here we examine the complex membrane interactions of H-ras with a combination of FRAP on living cells to measure membrane affinity and electron microscopy of intact plasma membrane sheets to spatially map microdomains. We show that three separable forces operate on H-ras at the plasma membrane. The lipid anchor, comprising a processed CAAX motif and two palmitic acid residues, generates one attractive force that provides a high-affinity interaction with lipid rafts. The adjacent hypervariable linker domain provides a second attractive force but for nonraft plasma membrane microdomains. Operating against the attractive interaction of the lipid anchor for lipid rafts is a repulsive force generated by the N-terminal catalytic domain that increases when H-ras is GTP loaded. These observations lead directly to a novel mechanism that explains how H-ras lateral segregation is regulated by activation state: GTP loading decreases H-ras affinity for lipid rafts and allows the hypervariable linker domain to target to nonraft microdomains, the primary site of H-ras signaling.

The spatial organization of signaling proteins on the plasma membrane contributes to the regulation and coordination of signaling networks. The mechanisms whereby proteins and lipids are sequestered into specific microdomains are incompletely understood. The proposal that cholesterol-rich lipid rafts, which have been clearly demonstrated in artificial membranes, also exist and operate in biological membranes has gained widespread but not general acceptance (12, 25). There is good evidence from single-molecule tracking, fluorescence resonance energy transfer (FRET), and photonic force microscopy for cholesterol- and actin-dependent microcompartmentalization of the outer leaflet of the plasma membrane (3, 6, 16, 22, 28), although the biophysical basis of this process continues to be debated (1, 3, 12, 23, 25).

Studying the inner leaflet of the plasma membrane is even more problematic. Classical fractionation protocols that rely on detergent insolubility and flotation with light membranes on density gradients suggest that a subset of inner plasma membrane proteins associate with lipid rafts but cannot offer any insight into the spatial organization of inner leaflet proteins in intact cells. FRET studies with monomeric yellow and cyan fluorescent proteins tethered by a Fyn membrane anchor strongly suggest that lipid rafts exist on the inner plasma membrane, since FRET is abolished when cell surface cholesterol is acutely depleted (29). The putative size of lipid rafts (<100 nm) is below the resolution of the light microscope, preventing direct visualization of discrete microdomains by confocal microscopy unless rafts are first cross-linked into larger structures (8).

To address the problem of directly visualizing inner plasma membrane microdomains we used electron microscopy (EM) to study the spatial organization of Ras signaling domains. Ras proteins are small GTPases that operate as molecular switches on the inner surface of the plasma membrane. Three isoforms of Ras, H-, N-, and K-ras, are ubiquitously expressed in mammalian cells. They interact with a common set of effector and regulatory proteins, and yet each isoform generates a distinct signal output (7). These signaling differences may reflect the ability of the different Ras membrane anchors to target to distinct membrane microdomains. The membrane anchors of H-, N-, and K-ras comprise a common C-terminal farnesyl cysteine carboxy methylster operating in concert with one or two adjacent S-palmitoyl cysteine residues in N- and H-ras or a polybasic domain of six lysines in K-ras (7).

Sucrose gradient fractionation and functional assays in cholesterol-depleted cells show that inactive H-ras is partially localized to lipid rafts but that constitutively activated H- and K-ras fractionate away from raft markers (17). Quantitative EM analysis of intact plasma membrane sheets shows that constitutively activated H- and K-ras occupy spatially distinct nonraft microdomains and that only inactive H-ras interacts with lipid rafts (18). An independent study (14), using fluorescence recovery after photobleaching (FRAP) on living cells, reached similar conclusions: activated H- and K-ras interact with distinct nonraft sites on the plasma membrane and only inactive H-ras has significant affinity for lipid rafts.

From our studies it has become clear that the guanine nucleotide-bound state and C-terminal sequences adjacent to the membrane anchor influence whether H-ras associates with
lipid rafts or nonraft microdomains (17). It remains unknown, however, how H-ras plasma membrane interactions are regulated and how different elements control its overall membrane affinity and preference for different microdomains. To tackle these questions we examined a series of H-ras C-terminal mutants, which have been well characterized in terms of biological and biochemical activity, using a combination of high-resolution EM spatial mapping and FRAP analysis of living cells. Our data show that separable, attractive forces provided by the membrane anchor and linker domain of the hypervariable region (hvr) govern the interaction of H-ras with the plasma membrane. These attractive forces however operate against a repulsive force contributed by the N-terminal catalytic domain that increases on GTP loading. The results suggest a novel mechanism for regulation of H-ras lateral segregation that may be applicable to other lipidated signaling proteins. The study also shows how data generated from EM (yielding high-resolution spatial information) and FRAP (providing dynamic information from living cells), two quantitative methods of studying membrane interactions, are mutually informative.

MATERIALS AND METHODS

Reagents. Affinity-purified polyclonal anti-green fluorescent protein (GFP), monomolecular anti-Ras Y13-238 antibodies, the expression constructs GFP-H-rasG12V, GFP-CTH, GFP-H-ras(wt), and GFP-H-rasG12V, and hypervariable linker region mutants of GFP-H-rasG12V (GFP-H-rasG12V-Δhvr, Δ1, Δ2, Δ1ala, and Δ2ala) have been described previously (11, 14, 17, 18). GFP-H-ras proteins were fixed 48 h posttransfection with 3% paraformaldehyde (30 min), permeabilized with 0.2% Triton X-100, and incubated (40 min) with 0.1 μg of phalloidin-tetramethyl rhodamine isocyanate (TRITC)/ml. The cells were washed and mounted in Prolong Antifade (Molecular Probes). Transfected cells were identified by GFP fluorescence, and the TRITC-stained F-actin in these cells was visualized using a Zeiss LSM 510 confocal microscope fitted with rhodamine filters.

RESULTS

Biological activities of GFP-H-ras proteins in COS-7 cells. Jaumot et al. previously reported the biological and biochemical activity of a series of constitutively active H-rasG12V proteins with deletions or alanine substitutions in the hvr linker domain adjacent to the membrane anchor (11) (for schematic representations and nomenclature, see Fig. 1A). The plasma membrane interactions of these mutants were not examined in detail, because we had not then developed EM techniques to assign proteins to morphologically featureless plasma membrane microdomains. BHK cells and PC12 cells were used for this earlier work. The FRAP analysis reported here employs COS-7 cells; therefore, we first confirmed that the biological effects of the mutant GFP-H-rasG12V proteins in these cells are similar. As a simple readout for Ras activity we examined the disassembly of actin stress fibers (27). F-actin in COS-7 cells transiently expressing GFP-H-rasG12V proteins was stained with phalloidin-TRITC and imaged by confocal microscopy (Fig. 1B). The number of transfected cells exhibiting clear stress fibers was determined (Fig. 1C). Control cells transfected with GFP vector exhibited strong stress fibers, whereas expression of GFP-H-rasG12V resulted in a nearly complete loss of stress fibers. GFP-H-rasG12V-Δhvr, GFP-H-rasG12V-Δ1, GFP-H-rasG12V-Δ2, and H-rasG12V-Δ1ala were scored biologically inactive since their expression did not cause any significant loss of stress fibers, whereas the alanine substitution mutant H-rasG12V-Δ2ala exhibited reduced activity (Fig. 1C). These results obtained with COS-7 cells agree with the previously characterized biological activities of the GFP-Ras proteins in PC-12 and BHK cells (11).

Localization of hvr to nonraft microdomains requires hvr linker domain sequences. To directly visualize and quantify the microlocalization of H-ras proteins on intact plasma membranes we used a combination of immunogold EM and spatial statistics. Intact plasma membrane sheets were prepared from BHK cells (left untreated or incubated with MβCD to deplete cholesterol) expressing GFP-H-rasG12V proteins and labeled after fixation with 4-nm-particle-size gold directly coupled to affinity-purified anti-GFP antibodies (Fig. 2). The gold patterns on multiple plasma membrane sheets were analyzed using Ripley’s K-function to determine whether they were random, clustered, or dispersed. As reported previously (18) and shown here (Fig. 3A), full-length H-rasG12V is clustered on the plasma membrane: the L(r)-r curve shows a significant Gaussian bleach profile and the mobile fraction were derived by nonlinear regression analysis, fitting to a lateral diffusion process with a single r value (15). Statistical comparisons were carried out using Student’s t test.

Cell fractionation. BHK or COS-7 cells expressing GFP-tagged proteins were subjected to hypotonic lysis, and P100 and S100 fractions were prepared from postnuclear supernatants as described previously (21). A total of 20 μg of each P100 fraction and an equal proportion of the S100 fraction were immunoblotted for GFP. Blots developed using enhanced chemiluminescence were visualized and quantified by phosphorimaging (21).
positive deviation out of the 99% confidence interval for complete spatial randomness (CSR) at a radius of 20 nm (Fig. 3A). The clustering of H-rasG12V is unaffected by cholesterol depletion, indicating association with cholesterol-independent microdomains (Fig. 3) (18). Figure 3B shows that deletion of the complete hvr linker domain (amino acids 166 to 180: H-rasG12V-hvr) has a fundamental effect on the microlocalization of H-ras: H-rasG12V-hvr is clustered, as indicated by the substantial positive deflection of the L(r)-r curve outside the 99% CSR confidence interval, but clustering is completely abolished in MβCD-treated cells. We conclude that H-rasG12V-hvr is localized to lipid rafts. Raft localization is also evident for H-rasG12V-D1, H-rasG12V-D1ala, and H-rasG12V-D2 (Fig. 3C to E; for structure, see Fig. 1A), which are significantly clustered, but their clustering is substantially reduced upon cholesterol depletion. Unlike H-rasG12V-hvr, however, H-rasG12V-D1, H-rasG12V-D1ala, and H-rasG12V-D2 must retain some affinity for nonraft microdomains, because the L(r)-r
curves do not fall completely into the CSR envelope. This indicates residual clustering after lipid raft disassembly with MβCD (Fig. 3C to E). The microlocalization of H-rasG12V-Δ2ala closely resembles that of full-length H-rasG12V and exhibits the same degree of clustering irrespective of whether cell surface cholesterol is depleted (Fig. 3F).

To confirm the assignment of H-rasG12V-Δ1, H-rasG12V-Δ1ala, and H-rasG12V-Δ2 but not H-rasG12V-Δ2ala to lipid rafts, they were coexpressed in BHK cells with the lipid raft marker GFP-tH (schematically shown in Fig. 1A). Plasma membrane sheets from transfected cells were colabeled with 4-nm-particle-size gold conjugated to anti-GFP and 2-nm-particle-size gold conjugated to anti-Ras (Y13-238). The extent of colocalization of Ras mutants with GFP-tH was then quantified using bivariate K-functions to analyze the gold patterns (Fig. 4). The null hypothesis for bivariate analysis is that the gold particle populations are randomly arrayed around each other; thus, positive deflections of an $L_{xy}(r)$-r curve above the 95% confidence interval indicate significant colocalization of the two sets of gold particles. By these criteria, H-rasG12V-Δ1, H-rasG12V-Δ1ala, and H-rasG12V-Δ2 colocalize with the raft marker GFP-tH whereas H-rasG12V-Δ2ala does not (Fig. 4).

These data suggest that hvr linker sequences are required for the interaction of H-ras with nonraft microdomains. To confirm this, we investigated whether these sequences are sufficient to alter the targeting of GFP-tH to lipid rafts. To this end we studied the microlocalization of GFP-CTH, which contains the entire H-ras hvr (amino acids 166 to 189) fused to GFP rather than just the minimal membrane anchor present in GFP-tH (Fig. 1A). In contrast to GFP-tH, which is exclusively clustered in cholesterol-dependent lipid rafts (18) (Fig. 5), GFP-CTH also clusters outside lipid rafts, as evidenced by its persistent clustering in cholesterol-depleted cells (Fig. 5). Finally, to complete the EM analysis, we show that GFP-H-ras(wt)-Δhvr clusters in lipid rafts, as demonstrated by the loss of its clustering upon cholesterol depletion (Fig. 5).

Distinct roles for H-ras anchor and hvr linker domains in regulating membrane association. EM provides high-resolution spatial information but is restricted to fixed membrane sheets. To probe the dynamics of Ras-membrane interactions in living cells we used FRAP. We first compared the fluorescence recovery of GFP-H-rasG12V and GFP-H-rasG12V-Δhvr. Typical FRAP curves are shown in Fig. 6, and the averaged results are depicted in Fig. 7. GFP-H-rasG12V had a characteristic fluorescence recovery time $\tau$ of $\sim0.4$ s with a laser beam with a Gaussian radius of 0.85 μm, in keeping with our previously reported lateral diffusion coefficient ($D$) of $\sim5 \times 10^{-9}$ cm$^2$/s (Fig. 6A) (14). Interestingly, the fluorescence recovery of GFP-H-rasG12V-Δhvr was much faster ($\tau \sim 0.1$ s; Fig. 6B), although it was still more than an order of magnitude slower than that of free GFP in the cytosol, which recovered at a rate faster than the experimental time scale ($\tau < 0.005$ s; Fig. 6C). This indicates that the fluorescence recovery of GFP-H-rasG12V is retarded relative to that of GFP-H-rasG12V-Δhvr.

To further investigate this difference we performed FRAP with laser beams of different sizes (5, 10, 13). If FRAP occurs by lateral diffusion, $\tau$ is essentially the characteristic diffusion time $\tau_D$, and is proportional to the area illuminated by the beam ($\tau_D = \omega^2/4D$, where $\omega$ is the Gaussian radius of the laser beam). When FRAP takes place by dynamic exchange between membrane-bound and cytosolic pools (as may occur for weaker, transient interactions with the membrane), $\tau$ reflects the chemical relaxation time due to exchange, which is equal on all surface regions regardless of whether they are illuminated by the beam, and therefore does not depend on the beam size (5, 10, 13). The expected ratio of $\tau(40\times)/\tau(63\times)$ for the two beam sizes generated using the $40\times$ and $63\times$ objectives, respectively, is 2.56 for pure lateral diffusion or 1 (no dependence on beam size) for pure exchange (see legend to Fig. 7) (10). In agreement with earlier results (13), GFP-H-rasG12V exhibited pure lateral diffusion, as evident by a

FIG. 2. Immunogold labeling of Ras proteins on plasma membrane sheets. Examples of plasma membrane sheets from BHK cells expressing GFP-H-rasG12V and labeled with 4-nm-particle-size gold conjugated directly to affinity-purified polyclonal anti-GFP antibody are shown. (A) Untreated cells. (B) Cells treated for 60 min with 1% MβCD before the sheets were prepared. Bar, 100 nm.
The N-terminal catalytic domain of H-ras weakens plasma membrane interactions. The conclusion that the hvr linker domain is essential for stable membrane association of GFP-H-rasG12V suggested that the lipid-modified anchor by itself might not be sufficient for stable membrane binding. To examine this possibility we investigated the membrane interactions of GFP targeted to the plasma membrane by the minimal C-terminal anchor of H-ras (GFP-tH). GFP-tH exhibited stable membrane association and pure lateral diffusion, as evident by its \( \tau(40\times)/\tau(63\times) \) values and ratio, which were similar to those of GFP-H-rasG12V (\( P > 0.1 \) in all cases; Fig. 7A). Thus, out of the context of full-length H-rasG12V, the H-ras minimal anchor is sufficient for stable membrane association. However, when this anchor is connected to the N-terminal domains of H-ras, omission of the hvr linker (GFP-H-
rasG12V-hvr) results in exchange (Fig. 7A). Taken together, these data strongly suggest that the catalytic N-terminal domains of GFP-H-rasG12V generate a repulsive force that weakens the association of H-ras with the membrane.

The N-terminal catalytic domain of Ras proteins undergoes marked conformational changes between the GDP- and GTP-bound states. We therefore examined whether GTP loading might regulate the repulsive force exerted by the N-terminal region on H-ras membrane binding. We reasoned that any differences would be more marked in H-ras proteins that lack the hvr linker domain and are solely dependent on the anchor for membrane association. In contrast to GTP-bound GFP-H-rasG12V-hvr that recovers solely by exchange, GDP-bound GFP-H-ras(wt)-hvr exhibits a mixed mode of fluorescence recovery (i.e., a contribution from both lateral diffusion and exchange), as indicated by the $\tau_{(40\times)}/\tau_{(63\times)}$ ratio (1.6), which is significantly different from both 1 and 2.56 ($P < 0.001$ in both cases) (Fig. 7A). This suggests that GTP-loaded GFP-H-rasG12V-hvr has a weaker membrane interaction than GDP-loaded GFP-H-ras(wt)-hvr, a conclusion also supported by the higher $\tau$ values measured for GFP-H-ras(wt)-hvr with both beam sizes. Together, the data show that of the GDP- and GTP-bound states of the hvr mutants, the former has the stronger and more stable interaction with the membrane, although this interaction is not as strong as that of the stably interacting (i.e., laterally diffusing) full-length GFP-H-ras(wt) (Fig. 7A). We conclude that the conformation of the catalytic N-terminal domain regulates H-ras membrane interactions, with GTP binding to H-ras increasing the repulsive force. Thus, H-ras exhibits GTP-dependent regulation of membrane affinity.

A testable prediction of this conclusion is that there may be a greater cytosolic pool of GFP-H-rasG12V-hvr than GFP-H-ras(wt)-hvr. BHK cells expressing GFP-H-rasG12V-hvr, GFP-H-ras(wt)-hvr, and a range of control proteins were therefore fractionated into S100 and P100 fractions and quantitatively immunoblotted for GFP (Fig. 8). The fraction of GFP-H-rasG12V-hvr and GFP-H-ras(wt)-hvr stably associated with cell membrane was significantly reduced ($P < 0.001$ in both cases) compared to that full-length GFP-H-rasG12V. Significantly ($P < 0.001$) more GFP-H-ras(wt)-hvr was membrane associated than GFP-H-rasG12V-hvr, in full consistency with the FRAP data. Similar results (data not shown) were obtained with COS-7 cells. Interestingly, a significant ($P < 0.05$) difference was also measurable between GFP-tH and GFP-CTH, a result consistent with GFP-CTH having a greater membrane affinity than GFP-tH (Fig. 8).

The hvr linker domain is required for stable binding to nonraft microdomains. To examine whether the anchor and linker hvr domains of H-ras interact predominantly with raft or nonraft microdomains, we studied by FRAP the effects of cholesterol depletion on the membrane affinity of GFP-H-ras. In accord with former studies (14), both GFP-H-rasG12V and GFP-H-ras(wt) exhibited fluorescence recovery by pure lateral

FIG. 4. Spatial mapping of H-ras proteins to lipid rafts using bivariate K-functions. Plasma membrane sheets were prepared from BHK cells coexpressing GFP-tH and (untagged) H-rasG12V-Δ1 (A), H-rasG12V-Δ1ala (B), H-rasG12V-Δ2 (C), or H-rasG12V-Δ2ala (D). Sheets were colabeled with monoclonal antisera to Ras (Y13-238) coupled directly to 4-nm-particle-size gold and anti-GFP antisera coupled directly to 2-nm-particle-size gold. Coincidence of the 2- and 4-nm-particle-size gold in study areas 725 by 725 nm in size was analyzed by spatial statistics. The graphs show mean bivariate K-functions expressed as $L_{biv}(r)-r$ on the 95% confidence interval for CSR (●). Each curve represents data pooled from multiple (n = 6 to 9) plasma membrane sheets. The average labeling density was 26 (4 nm) and 145 (2 nm) gold particles/μm².
diffusion \[\tau(40\times)/\tau(63\times)\] not significantly different from 2.56; \(P > 0.2\) after cholesterol depletion (Fig. 7B). However, the effects of cholesterol depletion on the lateral diffusion rates were much more pronounced for GFP-H-ras(wt), in accord with its partial localization to lipid rafts (17, 18). Thus, neither clustering of GFP-H-rasG12V (Fig. 2) nor its continuous membrane association (Fig. 7) requires lipid rafts. In contrast, cholesterol depletion had a strong impact on the normally raft-resident GFP-H-rasG12V-hvr, which became too fast to be measured at our time scale (\(\tau < 0.005\) s, in similarity to GFP results; Fig. 7B).

This indicates that the normally raft-resident GFP-H-rasG12V-hvr is very loosely associated with the membrane in the absence of rafts, in keeping with its random membrane distribution following cholesterol depletion (Fig. 3). This could be a result of strong manifestation of the repulsive forces from the N-terminal domain and/or a loss of positive attractive forces from the lipid-modified anchor.

To distinguish between these two possibilities, we performed FRAP studies on cholesterol-depleted cells expressing GFP-tH and GFP-CTH. The \(\tau\) values of GFP-tH, which has only the lipid anchor, decreased significantly (1.3-fold; \(P < 0.001\)) following cholesterol depletion (Fig. 7), indicating a reduction in interaction with the membrane. However, this reduction was
The hvr linker domain contains two regions that contribute differently to membrane affinity. The H-ras hvr linker domain has been tentatively divided into two regions: region 1 and region 2 (Fig. 1 and 9) (11). Given our data showing that the hvr linker contributes to H-ras membrane interactions (Fig. 3, 4, and 7), we evaluated by FRAP the specific contribution of regions 1 and 2 by the use of H-ras mutants in which the regions were lacking or replaced with alanines (Fig. 9). A comparison of GFP-H-rasG12V-Δ1 (missing region 1) and GFP-H-rasG12V-Δ1ala (region 1 sequence replaced by alanines; Fig. 1A and 9) showed that both proteins exhibited fluorescence recovery by pure lateral diffusion [τ(40×)/τ(63×) ratio, 2.0 to 2.3; Fig. 9A]. These values are very close to that of full-length H-rasG12V and suggest stable associations with the membrane. Cholesterol depletion significantly weakened these interactions but did not eliminate them altogether (Fig. 9B), as illustrated by the smaller τ values and the shift of the fluorescence recovery to exchange [τ(40×)/τ(63×) ratio of 0.9, not significantly different from 1; P > 0.2]. We conclude that region 2 of the hvr linker, present in both mutants, contributes to the interactions of H-rasG12V with the membrane. In addition, the interactions of H-rasG12V-Δ1ala and H-rasG12V-Δ1 appear to be with both raft and nonraft sites, in accord with the EM clustering analysis (Fig. 3C and D).

In H-ras mutants lacking region 1, the correct position of region 2 is retained immediately adjacent to the C-terminal lipid-modified anchor. However, in the mutants missing region 2 (GFP-H-rasG12V-Δ2 and GFP-H-rasG12V-Δ2ala; Fig. 1A), region 1 is retained in its original position only in the alanine substitution mutant. GFP-H-rasG12V-Δ2ala displays stable membrane interactions not significantly different from those of full-length H-rasG12V [τ(40×)/τ(63×) = 2.3; P > 0.1], and this strong interaction is insensitive to cholesterol depletion [τ(40×)/τ(63×) = 2.5; P > 0.1] (Fig. 9). On the other hand,
GFP-H-rasG12V-Δ2 (in which region 1 is out of the original context and is moved up to be adjacent to the lipid anchors) displays impaired membrane interactions: in untreated cells, it exhibits exchange $[\tau(40\times)/\tau(63\times) = 0.8]$, and cholesterol depletion further reduces its membrane interactions, resulting in a fluorescence recovery rate faster than the experimental time scale (Fig. 9B). We conclude that region 1 of the hvr linker makes a major contribution to H-rasG12V membrane interactions but that this contribution requires region 1 to be correctly positioned in the context of the protein relative to the lipid-modified C-terminal anchor. The interaction of region 1 with the membrane appears to be with cholesterol-insensitive, non-raft sites. These results are in excellent agreement with the EM data (Fig. 3E to F and 4).

**DISCUSSION**

The interaction of lipidated proteins with cell membranes is dependent on the lipid anchor. We show here, however, that for the H-ras small GTPase, plasma membrane anchoring is the product of three separable forces, of which only one is generated by the lipid anchor. The forces operating on H-ras to modulate plasma membrane attachment were revealed using a combination of FRAP to differentiate between stable and transient membrane attachment and quantitative EM to identify the spatial interactions. The minimal H-ras anchor, comprising the processed CAAX motif plus the adjacent palmitoylation sites, is sufficient to target GFP to the plasma membrane. The FRAP analysis reported here (Fig. 7) shows that GFP-tH is stably associated with the plasma membrane such that fluorescence recovery occurs exclusively by lateral diffusion. EM analysis (Fig. 4 and 5) (18) illustrates that GFP-tH is clustered in cholesterol-dependent lipid raft microdomains. Together these data show that the minimal membrane anchor of H-ras provides a high-affinity membrane interaction primarily with plasma membrane lipid rafts (see proposed model in Fig. 10). The affinity of the minimal anchor for the plasma membrane is reduced when lipid rafts are disassembled by cholesterol depletion (reflected by faster $\tau$ values) but remains sufficiently strong to prevent a shift of the FRAP to exchange (Fig. 7). In consequence, as shown by EM, there is no loss of GFP-tH from the plasma membrane in cholesterol-depleted cells but GFP-tH is no longer clustered and is randomly distributed (18).

In contrast, GFP-H-rasG12V-Δhvr, which has exactly the same membrane anchor as GFP-tH, does not interact stably with the plasma membrane. FRAP of GFP-H-rasG12V-Δhvr occurs by exchange and in cholesterol-depleted cells becomes too fast to be measured experimentally, showing severely reduced membrane affinity (Fig. 7). EM analysis (Fig. 3) shows however that GFP-H-rasG12V-Δhvr is still attached to the...
plasma membrane, clustering in lipid rafts, or distributed randomly over the membrane when cells are cholesterol depleted. The simplest interpretation of the combined data is that the N-terminal catalytic domain of H-ras generates a repulsive force that opposes the affinity of the anchor for lipid rafts (Fig. 10). Interestingly, the repulsive force is reduced when H-ras is GDP loaded: FRAP of GFP-H-ras(wt)-hvr occurs by a combination of exchange and lateral diffusion in normal and cholesterol-depleted cells. Thus, the anchor generates a more stable membrane interaction for GDP-bound H-ras(wt)-hvr than for GTP-loaded GFP-H-rasG12V-hvr, a conclusion supported by cell fractionation results (Fig. 8). In both cases, EM shows that the anchor still targets to lipid rafts.

The membrane interactions of the lipid anchor are further modified when the hvr linker domain is present. The linker domain provides the C terminus of H-ras with a second source of membrane affinity but one that is predominantly for nonraft sites on the plasma membrane (see model in Fig. 10). Several lines of evidence support this conclusion. First, the fluorescence recovery time for GFP-CTH, which has the H-ras hvr linker domain and the lipid anchor, is unaffected by cholesterol depletion (Fig. 7). This demonstrates that there is no reduction in its membrane affinity when lipid rafts are lost. Secondly, EM shows that under the same conditions GFP-CTH remains clustered on the plasma membrane (Fig. 5). Thirdly, in contrast to GFP-H-ras(wt)-hvr and GFP-H-rasG12V-hvr, full-length GFP-H-ras(wt) and GFP-H-rasG12V, which have the hvr linker sequence, exhibit high-level affinity and stable interactions with the plasma membrane, as measured by FRAP, in the presence or absence of lipid rafts. EM identifies the site of interaction of the hvr linker as a nonraft microdomain (Fig. 3 and 5). The more extensive loss of affinity of the anchor for lipid rafts when H-ras-hvr is GTP loaded strongly suggests that the membrane attachment of H-ras-GTP is more dependent on the hvr than that of H-ras-GDP; given that the affinity of the hvr linker is predominantly for nonraft microdomains, these observations offer a basic mechanism for the GTP-dependent lateral segregation of H-ras out of lipid rafts that has been reported previously (Fig. 10) (17, 18).

Earlier work has suggested that the hvr linker domain can be functionally divided into two separate regions designated region 1 and region 2 (Fig. 1A), where region 1 is sufficient to support the biological activity of H-rasG12V (11) (Fig. 1C). We can now account for these observations. The FRAP analysis (Fig. 9) showed that GFP-H-rasG12V-Δ2ala, which retains region 1 with the correct spacing from the membrane anchor, displays strong and stable (pure lateral diffusion) membrane association. The FRAP parameters for GFP-H-rasG12V-Δ2ala were very similar to full-length H-rasG12V and were unaffected by cholesterol depletion. Spatial mapping by EM showed that GFP-H-rasG12V-Δ2ala clustered in nonraft microdomains. We conclude that region 1 of the hvr linker provides a major source of affinity for nonraft microdomains. However, this region has to be spaced correctly relative to the membrane anchor to operate efficiently. This is illustrated by GFP-H-rasG12V-Δ2, in which region 1 is not spaced appropriately and functions poorly, resulting in membrane interaction characteristics approaching those of GFP-H-rasG12V-hvr: EM and FRAP show that GFP-H-rasG12V-Δ2 is localized to lipid rafts but binds to the membrane with very low affinity (fluorescence recovery is by exchange).

The FRAP studies (Fig. 9) show that the two mutants missing hvr linker region 1 (GFP-H-rasG12V-Δ1 and -Δ1ala) interact stably with the membrane (fluorescence recovery by pure lateral diffusion; Fig. 9A). Deletion of both regions 1 and 2 (GFP-H-rasG12V-Δhvr) results in much weaker membrane association (fluorescence recovery by exchange; Fig. 7A), strongly suggesting that region 2 of the hvr linker (present in H-rasG12V-Δ1 and -Δ1ala) contributes to the association of H-rasG12V with the plasma membrane. The interaction of H-rasG12V-Δ1 and -Δ1ala appears to be with both raft and...
nonraft sites, as shown by the EM analysis and FRAP studies following cholesterol depletion. The EM analysis (Fig. 3 and 4) shows that both GFP-H-rasG12V-D\(\Delta\)1 and GFP-H-rasG12V-D\(\Delta\)1ala reside in part in cholesterol-sensitive clusters, colocalized with the raft-resident GFP-I\(\alpha\). Accordingly, the FRAP studies demonstrate that cholesterol depletion weakens but does not abolish the membrane interactions of GFP-H-rasG12V-D\(\Delta\)1 and GFP-H-rasG12V-D\(\Delta\)1ala, resulting in a shift of their fluorescence recovery to exchange (Fig. 9B).

All of the data presented here can be synthesized into a relatively simple model in which three separable forces operate on H-ras at the plasma membrane (Fig. 10). The model illustrates that a repulsive force of the N-terminal domain of H-ras operating primarily against an attractive force from the lipid anchor for lipid rafts is essential for allowing H-ras to exit from lipid rafts. A second attractive force provided by the hvr linker anchor for lipid rafts is essential for allowing H-ras to exit from operating primarily against an attractive force from the lipid rafts. A prediction of this model, that the membrane affinity of GFP-C\(\alpha\)-H should be greater than that of GFP-I\(\alpha\), is supported by cell fractionation and nonraft mutants on the basis of the ability of cholesterol depletion to elevate the lateral diffusion rate (14, 24). For all such proteins [H-ras(wt), H-rasG12V, H-rasG12V-D\(\Delta\)ala, GFP-I\(\alpha\), and GFP-C\(\alpha\)], the FRAP results are in complete agreement with the EM mapping data, providing a live-cell correlate to the EM studies.

Together, our studies demonstrate that the interactions that regulate H-ras membrane affinity also determine the preference for raft or nonraft microdomains. The combined approach has allowed us to characterize the roles of specific regions of H-ras in its interactions with the membrane. Notably, our findings demonstrate for the first time that three separate regions of H-ras have specific contributions to its affinity to the plasma membrane. The concept that the GDP/GTP loading state not only affects H-ras microlocalization in raft and nonraft domains but also modulates its affinity to the plasma membrane has important implications for H-ras signaling, since Ras association with the plasma membrane is necessary for its biological function (7). In addition, the finding that cholesterol depletion has simultaneous effects on both the lateral segregation and the membrane attachment of H-ras mutants raises the intriguing possibility that the free energy of H-ras lateral segregation is comparable to that of the hydrophobic interaction mediated by the H-ras membrane anchor. Finally, it is tempting to speculate that the findings presented here have broad implications for other signaling proteins that are lipid anchored to the inner surface of the plasma membrane. For example, it is generally assumed that lipid modifications alone are sufficient for lipid raft association but, as shown here, the microlocalization of a lipid modified protein can also depend on additional structural features of the protein. Furthermore it is possible that other signaling proteins, including the Src kinases, could dynamically regulate their interactions with lipid rafts by a similar mechanism to that described here for H-ras; namely, by generating a repulsive force, modulated by conformational changes in the catalytic domain, to act against the lipid anchor.

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