Flippase-mediated phospholipid asymmetry promotes fast Cdc42 recycling in dynamic maintenance of cell polarity

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Lipid asymmetry at the plasma membrane is essential for such processes as cell polarity, cytokinesis and phagocytosis¹⁻³. Here we find that a lipid flippase complex, composed of Lem3, Dnf1 or Dnf2 (ref. 4), has a role in the dynamic recycling of the Cdc42 GTPase, a key regulator of cell polarity⁵, in yeast. By using quantitative microscopy methods, we show that the flippase complex is required for fast dissociation of Cdc42 from the polar cortex by the guanine nucleotide dissociation inhibitor. A loss of flippase activity, or pharmacological blockage of the inward flipping of phosphatidylethanolamine, a phospholipid with a neutral head group, disrupts Cdc42 polarity maintained by guanine nucleotide dissociation inhibitor-mediated recycling. Phosphatidylethanolamine flipping may reduce the charge interaction between a Cdc42 carboxy-terminal cationic region with the plasma membrane inner leaflet, enriched for the negatively charged lipid phosphatidylserine. Using a reconstituted system with supported lipid bilayers, we show that the relative composition of phosphatidylethanolamine versus phosphatidylserine directly modulates Cdc42 extraction from the membrane by guanine nucleotide dissociation inhibitor.

Establishing stable cell polarity is crucial for cellular morphogenesis and differentiation. The Cdc42 small GTPase localizes to the site of polarized growth in yeast and orchestrates structural and signalling events required for budding and mating^{5,6}. Previous work showed that the polarity of Cdc42 distribution is maintained dynamically through two co-localized recycling pathways to counter Cdc42 diffusion^{7–9}. Whereas a slow pathway works through actin-based membrane trafficking, a fast pathway is mediated through Rdi1, the yeast guanine nucleotide dissociation inhibitor (GDI) for Rho family GTPases¹⁰. GDI is known to extract Rho proteins from the membrane by forming a complex involving both protein–protein contacts and binding of GDI to the prenyl lipid anchor at the COOH terminus of these proteins^{11,12}. When actin polymerization is inhibited, Rdi1-mediated Cdc42 recycling is required for the polarization of Cdc42 at the presumptive bud site⁹.

To gain insights into the molecular mechanisms regulating the location and rate of Rdi1-mediated Cdc42 recycling, we carried out a suppressor screen taking advantage of the fact that overexpression of Rdi1 under the Gal1 promoter represses yeast growth, probably owing to overextraction of Cdc42 and other essential Rho family GTPases from membrane compartments¹³ (also see Fig. 1). We reasoned that deletion of a gene encoding a protein that facilitates Cdc42 extraction by Rdi1 might rescue the Gal-Rdi1-induced growth defect. As there are many possible mechanisms for the suppression of the growth defect, the candidates would be further narrowed down to those encoding proteins co-localizing with Cdc42 at the site of polarization. A centromeric plasmid expressing Rdi1 from the Gal1 promoter was transformed into each strain of the yeast haploid non-essential deletion library14. Growth of each strain overexpressing Rdi1 was quantified and normalized relative to its growth without Rdi1 overexpression. This screen revealed ~277 initial candidates rescuing the overexpression growth defect, among which 15 encode plasma-membrane-localized proteins (Supplementary Fig. S1b). Among these 15 genes, LEM3 encodes a protein localized at the site of polarization (Fig. 1a and Supplementary Figs S1a,S2a). Lem3 is a Cdc50 family protein forming a heterodimeric complex with P-type ATPases Dnf1 or Dnf2, which flips aminophospholipid from the outer to inner plasma-membrane leaflet^{4,15,16}. Dnf2 was a weak rescuer from the screen (Supplementary Fig. S1a). Both Dnf1 and Dnf2 also localize to the site of polarization, similarly to Lem3 (Supplementary Fig. S2a-c). This observation and their reported involvement in cell polarity¹⁷ led us to carry out an in-depth study on the possible role for aminophospholipid flipping in

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Figure 1 $\Delta lem3$ suppresses the growth and Cdc42 localization defects due to *RDI1* overexpression. (a) Images of wild-type and the $\triangle lem3$ mutant (four replicate spots) grown on solid media with or without Rdi1 overexpression (pGal-RDI1) from the genome-wide screen. (b) Example autocorrelation curve of GFP-Cdc42 in wild-type cells under its endogenous promoter. FCS experiments were carried out in live yeast cells (wild type or △rdi1) expressing GFP-tagged Cdc42, Cdc42^{R66E} or Cdc42 $^{\mbox{c1885}}.$ FCS data were averaged over 13 cells and fitted with a one-component or two-component diffusion model. Inset: Weighted residuals obtained from each fitting. (c) Autocorrelation curves were fitted to a two-component model to extract percentages of slow- and fast-diffusing molecules. Error bars represent the estimated s.e.m. in the parameters determined from fitting average autocorrelation curves obtained from 12, 10, 13 and 22 cells from left to right corresponding to each of the Cdc42 mutants (Methods). Wt, wild type. (d) Molar concentration (nM) of soluble Cdc42 calculated from the FCS data as

regulating Cdc42 recycling at the polar cortex. To avoid heterogeneity in polarization stages, all the microscopy analyses in this work were carried out with unbudded polarized cells with a Cdc42 polar cap at the incipient bud site. described in c (Methods). (e) The average autocorrelation curve obtained from FCS experiments in live cells (n = 12) expressing Rdi1-mCherry (a red fluorescent protein) was also fitted with one- and two-component models in comparison with the Cdc42 curve. Inset: Distribution of weighted residuals obtained from each fitting. (f) Mobile Cdc42 concentrations (nM) of different pools (as indicated) determined at different times of pGal1-Rdi1 induction in wild-type cells. Shown are means and estimated s.e.m. from the analysis of average autocorrelation curves obtained from 21, 26 and 27 cells (from left to right). (g) Representative images of GFP-Cdc42 at the polar cap in unbudded wild-type and $\Delta lem3$ cells at different times of Rdi1 induction (as indicated above each column). Scale bar, 2 µm. (h) Fluorescence intensity profiles of GFP-Cdc42 along the perimeter of wild-type or △lem3 cells at different times of Gal-RDI1 induction. Fluorescence traces from 10-12 cells were first peak-aligned and then averaged to yield each of the profiles shown. (i) Same analysis and show as in **f** for $\triangle lem3$ cells (14, 21 and 23 cells from left to right).

To test if Δlem_3 reduces the ability of Rdi1 to extract Cdc42, we first used fluorescence correlation spectroscopy (FCS; ref. 18) to measure the mobile cytosolic concentration of green fluorescent protein (GFP)–Cdc42 expressed under the *CDC42* promoter^{7,9} in



Figure 2 Lem3 regulates Rdi1-mediated Cdc42 internalization and polar-cap morphology. (a) Schematic representation of the mathematical model as previously described⁹. The grey horizontal bar represents the overlapping Rdi1 and actin-based delivery window (χ), approximately 25% of the perimeter (Supplementary Fig. S2e–g; ref. 9); *d*_t, lateral diffusion rate of Cdc42 in the membrane; *m* and *n*, internalization rates inside and outside the delivery window, respectively; *h*, delivery rate inside the window. (b) Example normalized FRAP curves of GFP–Cdc42 from wild-type and Δ *lem3* cells. Solid lines represent the exponential fit of the FRAP data.

wild-type and $\Delta lem3$ cells with or without Rdi1 overexpression. FCS records fluorescence intensity fluctuations using a confocal microscope equipped with a photon counter. The autocorrelation curve of these data can be fitted to extract the molar concentration of the diffusing species from the amplitude of the curve (G_0) and the diffusion time (τ_D) from the shape of the curve¹⁸. Autocorrelation curves of GFP-Cdc42 fitted poorly with a single-component diffusion model but significantly better ($P < 1 \times 10^{-10}$) with a two-component diffusion model (Fig. 1b). This suggests the existence of at least two pools of mobile Cdc42 population, 64% showing fast diffusion $(\tau_D = 2.9 \pm 0.3 \text{ ms})$ and 36% showing slow diffusion $(\tau_D = 70 \pm 8 \text{ ms};$ Fig. 1c,d). The fast pool was significantly reduced in the $\Delta r di1$ background and for the Cdc42^{R66E} mutant, defective in binding Rdi1 (ref. 19; Fig. 1c,d). The vast majority (>95%) of Rdi1 exists in a fast-diffusing population ($\tau_D = 2.2 \pm 0.1$ ms; Fig. 1e). The slow pool of Cdc42 was diminished by the *cdc42*^{C188S} mutation (Fig. 1c,d), which prevents prenylation. This analysis, along with the observed in vivo cross-correlation of Cdc42 and Rdi1 (ref. 9), suggests that the fast pool of Cdc42 consists largely of the Cdc42-Rdi1 complex, whereas the slow pool is dependent on prenylation and likely to be membrane vesicle associated.

(**c**–**e**) Extracted model parameters from FRAP data. Each point represents a value from a single cell. The small square is the mean; the box range is s.e.m.; whiskers represent s.d.; the line is the median. (**f**) Examples of normalized iFRAP curves from cells expressing GFP–Cdc42. (**g**) iFRAP rates from cells expressing GFP–Cdc42. Box plots are as described in **c**–**e**. (**h**) Example kymographs obtained from bleach-corrected movies (300–385 s long) along the perimeter of Δ *lem3* cells expressing GFP–Cdc42 in the presence of solvent control (dimethylsulphoxide) or 100 µM LatA (three examples shown). Scale bar, 2 µm.

Induction of Rdi1 overexpression resulted in a time-dependent increase in total mobile Cdc42 concentration in the cytosol (Fig. 1f). This increase is accompanied by a similar increase in Cdc42 concentration in the fast but not the slow pool, consistent with extraction of Cdc42 by Rdi1 from the plasma membrane into the soluble pool (Fig. 1f). Consistently, depletion of Cdc42 from the plasma membrane, especially from the polar cortex, can be observed (Fig. 1g,h). In $\Delta lem3$ cells, the Cdc42 polar cap was more prominent than in wild-type cells before Gal-Rdi1 induction (Fig. 1g,h), accompanied by a smaller mobile Cdc42 pool (Fig. 1i). Whereas Rdi1 overexpression reduced the concentration of Cdc42 in the polar cap of wild-type cells, in $\Delta lem3$ cells a nearly wild-type pre-extraction level remained (Fig. 1g,h) even after 150 min of Rdi1 induction. The cytosol Cdc42 pool also remained constant (Fig. 1i). As the mean fluorescence intensity of GFP-Cdc42 over the entire cell was slightly reduced in $\Delta lem3$ when compared with that in wild type (by \sim 15%, Supplementary Fig. S2d), the reduced Cdc42 soluble pool after 150 min Gal induction in $\Delta lem3$ (by ~40%, Fig. 1f,i) can be explained as a combined effect of a slightly reduced expression and the reduced extraction from the polar cortex.

We next carried out fluorescence recovery after photobleaching (FRAP) experiments to examine the recycling of GFP-Cdc42 at the



Figure 3 The Lem3–Dnf1/Dnf2 complex regulates the polar Cdc42 dynamics and phosphatidylserine distribution. (a) iFRAP rates of GFP–Cdc42 in polarized cells with single or double deletion of *DNF1* and *DNF2* in box plots as described in Fig. 2 caption. (b) iFRAP rates shown as in **a** for GFP–Cdc42 in wild-type and $\Delta lem3$ cells after treatment with 50 µM Ro peptide for 30 min at room temperature. Solvent control was with the same volume of water:acetonitrile (1:1). (c) Representative confocal microscopy images of wild-type and $\Delta lem3$ cells expressing GFP–Lact-C2, a phosphatidylserine biosensor. The white arrows point to the polar caps. Scale bar, 2 µm. (d) Representative GFP–Lact-C2 fluorescence intensity traces along the cell

polar cortex9 and analysed these data using a previously established mathematical model for Cdc42 recycling that enables extrapolation of the rates for Cdc42 dissociation and association from FRAP data9 (Fig. 2a,b and Supplementary Fig. S2e-g). Model computation found that $\Delta lem3$ most significantly reduced *m*, the rate of Cdc42 dissociation from the polar-cap region, but only slightly reduced h, the rate at which Cdc42 is targeted to the polar cap, whereas n, the internalization rate outside the cap, remained similar (Fig. 2c-e). Under such a scenario the model also predicts a slightly more pointed polar cap9, which could be observed in $\Delta lem3$ cells (Supplementary Fig. S2h,i). The reduced *m* in $\Delta lem3$ was confirmed with inverse FRAP (iFRAP), which directly measures the rate of internalization of GFP-Cdc42 at the polar cap9 (Fig. 2f,g). For comparison, the Cdc42 internalization was not significantly affected by the $\Delta pil1$ mutation (iFRAP rate of $0.19 \pm 0.028 \text{ s}^{-1}$, compared with 0.21 ± 0.01 s⁻¹ in wild type), which disrupts the eisosomes and normal ergosterol distribution in the plasma membrane²⁰.

perimeter. The white area represents the total fluorescent intensity in the half of the plasma membrane containing the polar cortex; the opposite half is shown in grey. (e) Confocal microscopy images of wild-type or Δ *lem3* cells expressing GFP–2xPH-PLCô (a PtdIns(4,5)P₂ biosensor). Scale bar, 2 µm. (f) Representative GFP–2xPH-PLCô fluorescence intensity traces along the cell perimeter as shown in **d**. (g) GFP–Lact-C2 fluorescence intensity ratios of the half of the plasma membrane containing the polar cap (white area in **d**) over the rest. The small square is the mean; the box range shows the s.e.m.; the whiskers are the s.d.; the line is the median. (h) The GFP-2xPH-PLCô fluorescence intensity ratio shown as in g.

To determine whether $\Delta lem3$ specifically slows down Rdi1-mediated Cdc42 internalization, iFRAP was carried out in the presence of latrunculin-A (LatA), which disrupts endocytosis²¹. LatA further slowed down Cdc42 dissociation in $\Delta lem3$ to an extent comparable to LatA's effect on wild-type cells (Fig. 2f,g). By contrast, the combined effect of $\Delta rdi1$ and $\Delta lem3$ on Cdc42 dissociation rate was non-additive when compared with $\Delta rdi1$ alone (Fig. 2g). These results suggest that Lem3 selectively promotes Rdi1-mediated Cdc42 dissociation from the polar cortex. LatA treatment also led to marked reduction of polarity in $\Delta lem3$ cells (Fig. 2h), consistent with a role for Lem3 as an important regulator of Rdi1-mediated Cdc42 recycling in the maintenance of cell polarity.

To test if the flippase complex is involved in fast Cdc42 dissociation from the polar cortex, we carried out iFRAP on $\Delta dnf1$ and $\Delta dnf2$ mutant cells. Cdc42 dissociation was significantly slowed in these mutant cells (Fig. 3a). We further tested if acute inhibition of



Figure 4 The influence of the charge property of a Cdc42 carboxy-terminal region on Cdc42 dynamics. (a) Internalization rates of Cdc42, Cdc42^{S185K} and Cdc42^{S185A} at the polar cortex measured with iFRAP in wild-type and Δ *lem3* cells; box plots as described in

lipid flipping had the same effect by using the tetracyclic peptide Ro09-0198 (Ro, also called cinnamycin), which specifically binds to phosphatidylethanolamine exposed on the outer plasma-membrane leaflet when added to the culture media^{1,22}. Ro peptide at 50 μ M, which did not cause significant cell death when compared with solvent control (5.6±0.29%, compared with 4.3±0.7%), slowed down Cdc42 dissociation from polar cortex in wild-type cells but not in Δ *lem3* cells (Fig. 3b). These results show that phosphatidylethanolamine flipping is required for fast Cdc42 dissociation from the polar cortex.

Consistent with a previous report²³, a defect in phosphatidylethanolamine flipping in $\Delta lem3$ correlated with an increase in the negatively charged phospholipid phosphatidylserine at the inner membrane leaflet of the polar cortex, as shown by using the phosphatidylserine biosensor GFP-Lact-C2 (ref. 2; Fig. 3c,d,g and Supplementary Fig. S3a,b). The distribution of phosphatidylinositol 4,5-bisphosphate (PtdIns $(4, 5)P_2$), monitored with the biosensor GFP-2xPH - PLCô (PH - PLCô, pleckstrin homology domain of phospholipase C δ) (ref. 24), was not significantly different between wild-type and $\Delta lem3$ cells (Fig. 3e,f,h). As Cdc42 contains a highly conserved cationic tail immediately adjacent to the prenylation site, the slowed Cdc42 dissociation in $\Delta lem3$ may be due to an enhanced charge interaction between Cdc42 and membrane lipids due to phosphatidylserine enrichment. To test this, we increased the Cdc42 polycationic charge by mutating Ser-185 to lysine (KKS185KK to KKK¹⁸⁵KK) and carried out iFRAP on wild-type and $\Delta lem3$ cells expressing Cdc42^{S185K}. In wild-type cells, the Cdc42^{S185K} dissociation rate from the polar cortex was significantly lower than that of Cdc42, approaching the same slow rate as Cdc42 dissociation in $\Delta lem3$ (Fig. 4a). In $\Delta lem3$, Cdc42^{S185K} also exhibited a moderate but significant decrease in dissociation rate when compared with that of Cdc42 (Fig. 4a).

Fig. 2 caption. (**b,c**) Representative confocal microscopy images of wild-type and Δ *lem3* cells expressing GFP-tagged Cdc42^{S185K} (**b**) or Cdc42^{S185D} (**c**). The white arrows point to the polar caps. Scale bar, 2 µm.

As Ser-185 might possibly act as an electrostatic switch through phosphorylation²⁵, we also mutated this residue to alanine, but the Cdc42^{S185A} did not exhibit a reduced iFRAP rate when compared with Cdc42 in wild-type cells and even showed a slightly increased iFRAP rate when compared with Cdc42 in $\Delta lem3$ (Fig. 4a). We also mutated the Ser-185 residue to the negatively charged aspartic acid, and the resulting mutant protein (Cdc42^{S185D}), unlike Cdc42^{S185K}, was poorly localized to the polar cap in both wild-type and $\Delta lem3$ cells (Fig. 4b,c). These observations support the notion that charge interaction between the Cdc42 polycationic tail and the phospholipids at the polar cortex regulates Cdc42 dissociation from the plasma membrane.

To test if the above regulation is direct, we employed an in vitro system involving supported lipid bilayers (SLBs). SLBs were assembled from liposomes with two different lipid compositions, one enriched for phosphatidylserine (referred to as 40% phosphatidylserine, 15% phosphatidylethanolamine, Methods), and the other for phosphatidylethanolamine (referred to as 40% phosphatidylethanolamine, 15% phosphatidylserine). Prenylated Cdc42 with an amino-terminal tetracysteine motif (TC-Cdc42) was purified from a yeast plasmamembrane fraction and labelled with the biarsenical dye FlAsH (ref. 26 and Supplementary Fig. S4). The labelled TC-Cdc42 was added to the SLB and appeared as a fluorescent layer with some bright foci when observed with total internal reflection fluorescence microscopy (TIRF), and its binding to the SLB was dependent on the prenyl group of Cdc42 (Supplementary Fig. S4c-e). In the SLB with 40% phosphatidylserine, 15% phosphatidylethanolamine, the fluorescence intensity of the SLB-bound TC-Cdc42, observed with TIRF, decreased spontaneously and slowly in the presence of the BSA control, and this decay was not enhanced by added purified, bacterially expressed Rdi1 (Fig. 5a,c). In the SLB with 40% phosphatidylethanolamine, 15% phosphatidylserine,



Figure 5 Lipid composition directly regulates Rdi1-mediated Cdc42 extraction. (a) Example traces of fluorescence intensity decay due to dissociation of FIAsH-labelled TC-Cdc42 from the SLB containing 40% phosphatidylserine and 15% phosphatidylethanolamine as measured by TIRF microscopy. Fluorescence intensity values (dots) were normalized to the maximum intensity of each trace and fitted to an exponential decay model (lines). (b) The same experiment and data presentation as in **a** but from an SLB containing a higher concentration of phosphatidylethanolamine (40%) relative to phosphatidylserine (15%). (c) Rates of TC-Cdc42 dissociation from the SLB determined from the TIRF movies, for 40% phosphatidylethanolamine (PE), 15% phosphatidylserine (PS) (n = 10, 10, 18 and 18 SLB areas from left to right) and for 40% phosphatidylserine, 15% phosphatidylethanolamine (n = 10, 10, 15 and

addition of Rdi1 stimulated the dissociation of Cdc42 from the membrane in a dose-dependent manner (Fig. 5b,c), confirming that phosphatidylethanolamine strongly facilitates Rdi1-mediated Cdc42 dissociation from the membrane.

On the basis of the data presented above, we propose a model whereby the Lem3–Dnf1/2 flippase complex acts as an electrostatic switch to promote fast Cdc42 dissociation from the plasma membrane by Rdi1 (Fig. 5d). At the steady state, the polar cortex region is enriched for phosphatidylethanolamine in the outer leaflet and phosphatidylserine in the inner leaflet^{2,17}. Such lipid asymmetry facilitates the localization of Cdc42 to the polar cortex²⁷, possibly by enhancing the charge interaction between the polycationic tail of Cdc42 and the negatively charged membrane surface. A transient increase in neutral lipids such as phosphatidylethanolamine as a result of flippase activity destabilizes the charge interaction of Cdc42 with the inner membrane leaflet, increasing the chance of exposure of the prenyl moiety on Cdc42, thus promoting its capture by the cytosolic Rdi1 protein and fast internalization.

A previous study implicated the Lem3 flippase activity in regulating the activity of the GTPase-activating proteins (GAPs) for Cdc42 and proposed this regulation to be important for the apical to isotropic switch of cell growth at G2/M (ref. 17). Two lines of consideration 10 SLB areas from left to right). Bar graphs show mean and s.e.m. (d) A schematic model depicting the flippase complex regulating Rdi1-mediated dissociation of Cdc42 from the plasma membrane by changing the charge property of the inner membrane leaflet. On the left, the Cdc42 prenyl group (red) is inserted in the inner leaflet of the plasma membrane at the polar cortex, with electrostatic interaction between the Cdc42 cationic C-terminal region (+) and the negatively charged phospholipid head groups (blue circles) providing stability for this association. The flippase complex mobilizes phospholipids with neutral head groups such as phosphatidylethanolamine from the outer to the inner leaflet, reducing the electrostatic interaction between Cdc42 and the membrane and enhancing the access of Rdi1 to the Cdc42–Rdi1 complex.

suggest that the regulation of Cdc42 dynamics observed in our study by the flippase complex is not a result of GAP regulation. First, the conclusion that phosphatidylethanolamine flipping regulates GAP activity in the previous work was based on the in vitro demonstration that phosphatidylethanolamine and phosphatidylserine stimulated the GAP activity of Rga1 toward Cdc42, contrasting the inhibitory effect of PtdIns(4, 5)P2. However, as shown in this work, the phosphatidylethanolamine-flipping defect in $\Delta lem3$ is accompanied by an increased enrichment of phosphatidylserine, not PtdIns(4,5)P₂, at the inner membrane leaflet of the polar cortex. As phosphatidylserine and phosphatidylethanolamine are equally effective in stimulating Cdc42 GAP in vitro¹⁷, the reciprocal enrichment of phosphatidylserine and phosphatidylethanolamine at the inner and outer membrane leaflets in the mutant, respectively, would predict little net effect on the GAP activity. Second, in the previous study the key in vivo result supporting the authors' conclusion was that overexpression of the GAP Rga1 rescued the elongated bud morphology in $\Delta lem3$, leading to the idea that lipid flipping regulates GAP activity in vivo. However, in budded cells beyond the nascent-bud stage, Rga1 is localized at the bud neck²⁸, whereas Lem3 is found at the bud tip¹⁶. Thus, it is unlikely that a direct regulation of the GAP by bud-tip-localized

flippase activity underlies the observed genetic interaction. Consistently, Rga1 overexpression does not rescue the defect in Cdc42 dynamics in $\Delta lem3$ but shows synthetic growth defect with $\Delta lem3$ (Supplementary Fig. S5a,b). Our model, on the other hand, is based on the change in the charge property of the inner membrane leaflet as a result of phospholipid flipping, and the *in vitro* data support a direct effect of the lipid composition on Cdc42 dissociation by Rdi1. Thus, our study reveals a mechanism by which dynamic lipid asymmetry regulates a highly conserved signalling process during cellular morphogenesis.

METHODS

Methods and any associated references are available in the online version of the paper at http://www.nature.com/naturecellbiology

Note: Supplementary Information is available on the Nature Cell Biology website

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AUTHOR CONTRIBUTIONS

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A.D. and R.L. designed the experiments; A.D. carried out all experiments and prepared the manuscript, with help from B.D.S. and J.R.U.; R.A. and W.D.B. assisted in the whole genome screening; B.R. assisted in data analysis; R.L. conceived and supervised the project and revised the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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METHODS

Yeast strains and genome-wide screening for suppressors of Gal-Rdi1. A detailed list of yeast strains used in this study is provided in Supplementary Table S1. The yeast haploid non-essential deletion library¹⁴ was transformed with a centromeric plasmid containing Rdi1 under a galactose-inducible promoter (CENpGAL-RDI1:His). Cells were grown for 2 days at 30 °C so that all the strains reached maximum growth with approximately the same optical density. At this point, strains were diluted (1:100) and spotted in quadruplicates on Sc + 2% galactose + 2% raffinose (non-transformed control) or Sc - His + 2% galactose + 2% raffinose agar (transformed experimental strains). A His+ wild-type control strain was included on every plate. A wild-type strain (RLY2530; Wt-control) on non-selective raffinose plates and a wild-type strain with Gal-Rdi1 (RLY3811; Wt-experimental) were included in several plates throughout the library to obtain the wild-type reference value. The plates were incubated at 23 °C for 98 h before saturation. Growth of each spot from the scanned images was quantified as the area and averaged over four spots to obtain growth for each strain. Area values were normalized with respect to the His+ wild-type control strain on the same plate. A growth ratio for each deletion strain was then obtained by dividing the normalized growth on the experimental plate by that on the control plate. The growth ratio for the wild-type control obtained by averaging from several plates was 0.26. On the basis of the distribution shown in Supplementary Fig. S1a, strains with a growth ratio equal to or above 0.56 were considered as rescuer candidates. On the basis of gene annotation in the Saccharomyces Genome Database, genes encoding proteins involved in the galactose-regulated transcription pathway and gene expression were excluded.

Live-cell imaging and data analysis. FRAP and iFRAP experiments and data analysis were carried out as described⁹. High signal-to-noise ratios for the still images were obtained by summing 20 middle confocal slices of a short movie of 20 frames. Image analysis and data extraction were carried out using ImageJ software; statistical analysis and data plotting were carried out using OriginLab Pro software.

Fluorescence correlation spectroscopy. FCS experiments were carried out as described in ref. 29. In brief, live yeast cells were mounted on coverslips and FCS measurements were taken in the cytosol. GFP-Cdc42 was excited with the 488 nm laser line of a Zeiss Confocor 3 through an HFT 488/561 dichroic. HFT 565 was used as an emission dichroic and BP 505-540 as an emission filter to collect fluorescence data. Autocorrelation curves generated from the data were fitted to a two-component diffusion model³⁰ with a triplet blinking component of 250 µs (ref. 31). Fitting was carried out using weighted nonlinear least squares³². The autocorrelation curves from many short (4s) data acquisitions were averaged for analysis. Curves that demonstrated bleaching or non-diffusive dynamics as judged by visual inspection of the binned trajectories and individual correlation traces were eliminated from the analysis. Weights were estimated from the s.e.m. at each time lag point. Errors in the fitted parameters were estimated by the Monte Carlo method as the s.d. in fits of 500 simulated correlation curves with Gaussian random errors corresponding to the estimated weights and the observed χ^2 parameter³². These errors represent the estimated standard error in the fit parameters. An f-test was carried out to compare the one-component and two-component diffusion models³². The two-component model represented a strong improvement in the quality of the fit as judged by the f-test $(P < 1 \times 10^{-10})$ and the visual inspection of quality of the fit and residuals (Fig. 1b). The FCS data obtained from Rdi1-mCherry were also fitted with one- and two-component models. In either case, the autocorrelation curve for Rdi1 is dominated by a rapidly diffusing component (more than 95% of population). An f-test between the models preferred the two-component fit with a P-value of less than 1×10^{-10} . However, given the small amplitude of the slow component, we conclude that it represents a relatively insignificant portion of the Rdi1 population.

For measurements of fast- and slow-pool concentrations in the presence of Gal–Rdi1, it is reasonable to assume that fast and slow diffusion times will remain unchanged. Therefore, the diffusion times were fixed to the values obtained for the wild-type strain at zero time for Gal induction. Indeed, the curves were reasonably well fitted by this model in every case. This allows for unbiased estimation of fast- and slow-pool concentrations as a function of Gal induction time.

From the fitting we obtained G_0 , the amplitude of the correlation function at zero time lag, and the average diffusion time τ_D required for the molecules to traverse the focal volume. The average number of molecules (*N*) present in the focal volume was calculated from

$$N = \gamma / G_0 \tag{1}$$

where γ is the shape factor of the focal volume. For one-photon excitation, the γ -factor is 0.36 (ref. 33). The focal volume was calculated using fluorescein in 0.1 M NaOH with the standard method³³⁻³⁶.

Modelling. For each cell, the mathematical model was applied for the analysis of the FRAP movie. Model parameters and assumptions are as described previously⁹. Considering the overlapping window size between the actin and Rdi1-based recycling mechanisms, we measured the window size from the formin Bni1 distribution in $\Delta lem3$ cells (Supplementary Fig. S2e–g) and found it similar to the distribution of Bni1 in wild-type cells⁹.

Recombinant protein purifications. We constructed a plasmid with a 'pGAL1-GST-HA-TC-Cdc42' sequence (referred to as TC-Cdc42; GST, glutathione Stransferase; HA, haemagglutinin). We used a genetically encoded tetracysteine motif (HRWCCPGCCKTF) (TC; ref. 37) that binds efficiently to the biarsenical dye FlAsH²⁶, and absorbs at 488 nm and emits at 525 nm. The tagged Cdc42 was overexpressed in yeast to maximize the amount of protein at the plasma membrane and purified from a plasma membrane fraction. Cell fractionation to prepare plasma membrane and protein purification was carried out as previously described^{12,38} with modifications. After homogenization of spheroplasts, unlysed cells were pelleted at 400g for 10 min. The pellet (P1) obtained at this step was rewashed with 5 ml lysis buffer and the supernatants were pooled (S1). The plasma-membrane-containing S1 pool was centrifuged twice at 158,420g for 45 min to pellet plasma-membrane fractions (P2). P2 was resuspended in solubilization buffer (50 mM Tris at pH 7.5, 300 mM NaCl, 5 mM MgCl₂, 1% Triton X-100 and 0.8 M sorbitol) and briefly homogenized 15 times on ice, and the resultant solution was incubated under shaking for 45 min at 4 °C. The solution was further centrifuged at 9,000g for 20 min and the supernatant (S3) was loaded on glutathione-agarose beads (Sigma, G4510) pre-equilibrated with solubilization buffer. The beads were washed with several column volumes of wash buffer (50 mM Tris at pH 7.5, 300 mM Nacl, 5 mM MgCl₂ and 0.1% 3-[(3-cholamidopropyl)-dimethylammonio]-1-propanesulphonate (CHAPS)) supplemented with protease inhibitors, phenylmethylsulphonyl fluoride and dithiothreitol, followed by the elution buffer (Tris-HCl at pH 8.0 and 0.1% CHAPS). TC-Cdc42 was eluted with elution buffer containing 10 mM reduced glutathione (Sigma-Aldrich, G4251). Recombinant non-prenylated Cdc42 was purified from bacteria using a GST tag. For Rdi1 purification, yeast Rdi1 was subcloned into the pET-28a vector with a C-terminal hexahistidine tag. Protein purification was done essentially as previously described¹² with minor modifications.

Liposome preparation. To prepare liposomes we purchased chloroform solutions of porcine brain phosphatidylethanolamine (catalogue no 840022C), porcine brain phosphatidylserine (catalogue no 840032C) and porcine brain phosphatidylcholine (catalogue no 840053C) from Avanti Polar Lipids. Cholesterol was purchased from Nu-Chek Prep, and dissolved in chloroform. Lipids were mixed in a clean glass tube, dried under a gentle stream of nitrogen gas and further dried under vacuum for 1 h. The dried lipid mixture was hydrated and resuspended with TBSM buffer (50 mM Tris, 150 mM NaCl, 5 mM MgCl₂ at pH 7.5) to a final concentration of 2 mM (ref. 39) and bath sonicateds for 30 min at 45 °C to produce the final liposome solution⁴⁰.

Cdc42 extraction assay using TIRF microscopy. Supported lipid bilayers were prepared on no 1.5 glass coverslips. Coverslips were made hydrophilic with a rinse with a concentrated H₂SO₄ and H₂O₂ (3:1) mixture, thoroughly washed with ultrapure water and dried with a stream of nitrogen gas. Liposomes with the composition 40% phosphatidylserine, 15% phosphatidylethanolamine, 5% phosphatidylcholine and 40% cholesterol, or 40% phosphatidylethanolamine, 15% phosphatidylserine, 5% phosphatidylcholine and 40% cholesterol, were added to this coverslip, incubated for 20 min and washed thoroughly with TBSM buffer. (50 mM Tris, 150 mM NaCl, 5 mM MgCl₂, pH 7.5) Prenylated Cdc42 with the tetracysteine FlAsH binding motif was labelled with 166.6 µM FlAsH dye (Invitrogen, catalogue no T34561) for 1 h at room temperature and added to the lipid bilayer at 63-126 nM final concentration, followed by 10 min incubation. Unbound and loosely bound protein was thoroughly washed from the lipid bilayer with TBSM buffer. TIRF imaging was carried out using a Carl Zeiss Axiovert 200 M inverted microscope equipped with the Laser TIRF accessory and a Plan-Apochromat ×100, 1.46 NA objective, and a C9100-13 EM-CCD digital camera from Hamamatsu Photonics. To quantify Cdc42 extraction, rectangular areas excluding the bright foci of Cdc42 aggregates were sampled from each experiment. The fluorescence intensity (I) decay for each area was fitted for to an exponential decay model, $I = A_0 + A_1 \exp(-\alpha t)$, where α is the exponential decay rate, using OriginLabPro software.

Statistical analysis. Statistical differences between two sets of data other than FCS data (see above) were analysed with a two-tailed unpaired Student *t*-test.

METHODS

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b Rescuer candidates that localize to the plasma membrane.

LEM3	Membrane protein of the plasma membrane and ER, interacts specifically in vivo with the phospholipid translocase (flippase) Dnf1p; involved in translocation of phospholipids and alkylphosphocholine drugs across the plasma membrane.
THI73	Putative plasma membrane permease proposed to be involved in carboxylic acid uptake and repressed by thiamine; substrate of Dbf2p/Mob1p kinase; transcription is altered if mitochondrial dysfunction occurs
PUN1	Plasma membrane protein with a role in cell wall integrity; co-localizes with Sur7p in punctate membrane patches; null mutant displays decreased thermotolerance; transcription induced upon cell wall damage and metal ion stress
FUS1	Membrane protein localized to the shmoo tip, required for cell fusion; expression regulated by mating pheromone; proposed to coordinate signaling, fusion, and polarization events required for fusion; potential Cdc28p substrate
ROD1	Membrane protein that binds the ubiquitin ligase Rsp5p via its 2 PY motifs; overexpression confers resistance to the GST substrate o-dinitrobenzene,zinc, and calcium; proposed to regulate the endocytosis of plasma membrane proteins
SSO1	Plasma membrane t-SNARE involved in fusion of secretory vesicles at the plasma membrane and in vesicle fusion during sporulation; forms a complex with Sec9p that binds v-SNARE Snc2p; syntaxin homolog; functionally redundant with Ssc2p
SSO2	Plasma membrane t-SNARE involved in fusion of secretory vesicles at the plasma membrane; syntaxin homolog that is functionally redundant with Sso1p
YPS1	Aspartic protease, member of the yapsin family of proteases involved in cell wall growth and maintenance; attached to the plasma membrane via a glycosylphosphatidylinositol (GPI) anchor
PSR2	Functionally redundant Psr1p homolog, a plasma membrane phosphatase involved in the general stress response; required with Psr1p and Whi2p for full activation of STRE-mediated gene expression, possibly through dephosphorylation of Msn2p
РМР3	Small plasma membrane protein related to a family of plant polypeptides that are overexpressed under high salt concentration or low temperature, not essential for viability, deletion causes hyperpolarization of the plasma membrane potential
YPS3	Aspartic protease, member of the yapsin family of proteases involved in cell wall growth and maintenance; attached to the plasma membrane via a glycosylphosphatidylinositol (GPI) anchor
SNC2	Vesicle membrane receptor protein (v-SNARE) involved in the fusion between Golgi-derived secretory vesicles with the plasma membrane; member of the synaptobrevin/VAMP family of R-type v-SNARE proteins
LDB19	Protein involved in regulating the endocytosis of plasma membrane proteins by recruiting the ubiquitin ligase Rsp5p to its target; localization changes in response to nutrient levels; null mutant has reduced affinity for alcian blue dye
SUR7	Plasma membrane protein that localizes to furrow-like invaginations (MCC patches); component of eisosomes; associated with endocytosis, along with Pli1p and Lsp1p; sporulation and plasma membrane sphingolipid content are altered in mutants
TPN1	Plasma membrane pyridoxine (vitamin B6) transporter; member of the purine-cytosine permease subfamily within the major facilitator superfamily; proton symporter with similarity to Fcy21p, Fcy2p, and Fcy22p

Figure S1 Genome-wide screen for rescuers of Rdi1 over-expression growth defect. (a) Distribution of In(growth-ratio) over the entire yeast haploid nonessential deletion library after 98 hrs of growth at 23C (see Supplementary Methods). Growth ratio is defined as the ratio of growth of a given strain with to without Rdi1 overexpression. Green line represents the Wt value and the cyan box represents the cutoff range of the In(growth-ratio) for rescuer candidates, including Δ Iem3 (the pink line). Δ dnf2 (blue line) had a slightly lower In(growth-ratio) than the cutoff and was therefore not in the initial rescuer list. (**b**) A list of the rescuer candidates that localize to the plasma membrane, with SGD description (www.yeastgenome.org).



Figure S2 The Lem3 flippase complex is polarized to the polar cortex and the effect of Δ *lem3* on the shape of Cdc42 distribution. Shown are confocal images of unbudded polarized cells expressing Lem3-GFP (**a**), Dnf1-GFP (**b**) and Dnf2-GFP (**c**), and box plots for quantification of the fluorescence intensity ratios of the half of the plasma membrane containing the polar cap over the other half as described in Figure 3g legend. P-value is measured using one-tailed one sample t-test to compare whether the intensity ratios are greater than 1.0 which corresponds to uniform distribution. Small Square is the mean; box range shows SEM; whiskers are standard deviation (SD); line is median. (**d**) Quantification of the mean fluorescence intensity over the entire cell for Wt (n = 10, 10, 11, from left to right) and Δ lem3 (n = 11, 9, 11,

from left to right) cells at different Rdi1 induction time points as indicated. Error bars represent standard error of mean (SEM). (e) The delivery window size in Δ lem3 was measured by the distribution of GFP tagged formin Bni1 as described previously¹. Confocal images represent unbudded polarized cells expressing Bni1-GFP. (f) The fluorescence distribution along the perimeter was fitted to a Gaussian model. (g) The window width is approximated as 2 times the full width half max (FWHM) of the Gaussian distribution. Shown is the mean and SEM (n=16). (h) Representative images of GFP-Cdc42 at the polar cap in unbudded Wt and Δ lem3 cells. (i) Fluorescence intensity traces along the perimeter are normalized and aligned with respect to peak intensity and shown adjacent to each cell type. Scale bar: 2 μ m.

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GFP-Lact-C2

mCherry-Cdc42



Colocalization

b



GFP-Lact-C2



∆lem3

Figure S3 PS polarization overlaps with Cdc42 and PS distribution in budded cells. (a) Confocal images of unbudded polarized cells in a cycling population expressing Cdc42 tagged with mCherry at the N-terminus and

GFP-Lact-C2. Scale bar: 2 μ m. (b) Images of budded Wt or Δ Iem3 cells expressing GFP-Lact-C2. Arrow points to enriched GFP signal in the small buds of Δ Iem3 cells compared to Wt cells. Scale bar: 4 μ m.

















Figure S4 SDS-PAGE followed by Coomassie staining of the purified proteins. Prenyl group is required for TC-Cdc42 association with the SLB. (a) Prenylated TC-Cdc42 (GST-HA-TC-Cdc42) protein was expressed and purified from the yeast plasma membrane (See Supplementary method). (b) C-terminally hexahistadine-tagged Rdi1 protein was expressed and purified from bacteria cell lysate. Asterisks mark the respective protein band of interest. (c) TIRF image of the SLB with 40% PS, 15% PE loaded with 126 nM FIAsH labeled prenylated Cdc42

(purified from yeast plasma membrane). (d) SLB with 40% PS, 15% PE similarly labeled as above with 126 nM FIAsSH labeled non-prenylated Cdc42 (purified from bacteria), with the rest of the conditions and image acquisition identical to c. Images were presented with identical contrast. Scale bar: $10 \ \mu$ m. (e) Mean fluorescence intensity sampled over 13-15 different optical fields on SLBs loaded with FIAsH-labeled prenylated or nonprenylated Cdc42 as explained above. Box plots are shown as described for Figure S2c.



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Figure S5 Over-expression of Rga1 (Cdc42 GAP) does not rescue Cdc42 dissociation rate at $\Delta lem3$ polar cortex. (a) Serial tenfold dilutions of cultures for the indicated strains were spotted on SGal-Leu and SD-Leu plates. Plates were scanned after 4 days incubation at 23⁰C. Notice

that Rga1 overexpression causes a growth defect, which is exacerbated by $\Delta lem 3$. (b) iFRAP rates (1/s) of GFP-Cdc42 in Δ lem3 cells bearing vector control or pGAL1-RGA1. Box plots are shown as described for Figure S2c.

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