Dual Modes of Cdc42 Recycling Fine-Tune Polarized Morphogenesis

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SUMMARY

In budding yeast, the highly conserved small GTPase Cdc42 localizes to the cortex at a cell pole and orchestrates the trafficking and deposition of cell surface materials required for building a bud or mating projection (shmoo). Using a combination of quantitative imaging and mathematical modeling, we elucidate mechanisms of dynamic recycling of Cdc42 that balance diffusion. Rd1, a guanine nucleotide dissociation inhibitor (GDI), mediates a fast recycling pathway, while actin patch-mediated endocytosis accounts for a slower one. These recycling mechanisms are restricted to the same region of the nascent bud, as both are coupled to the Cdc42 GTPase cycle. We find that a single dynamic parameter, the rate of internalization inside the window of polarized delivery, is tuned to give rise to distinct shapes of Cdc42 distributions that correlate with distinct morphogenetic fates, such as the formation of a round bud or a pointed shmoo.

INTRODUCTION

Polarized morphogenesis refers to the processes that give rise to distinct asymmetric cell shapes such as those of a neuron, epithelial cell, and filamentous fungus, etc, which are critical for specialized functions and physiology of these cells and organisms. The establishment of cell polarity, which results in the localization of signaling and cytoskeletal components that subsequently organize the growth of distinct cell structures, is a key step in polarized morphogenesis. While recent studies have led to considerable insights into the initial symmetry breaking processes that establish cell polarity (Li and Gunderson, 2008; Onsum and Rao, 2009), it remains poorly understood how polarized distributions of key regulatory molecules are stably maintained and are fine-tuned to give rise to different morphogenetic outcomes in response to diverse physiological or developmental signals.

A key regulator of cell polarity in many eukaryotes is the Rho family GTPase Cdc42 (Etienne-Manneville, 2004), first discovered in the yeast Saccharomyces cerevisiae (Johnson, 1999).

G1 phase yeast cells polarize to initiate bud formation or in response to pheromone to form the mating projection (shmoo). During both these processes, Cdc42 localizes to a small cortical domain, which becomes the presumptive bud or shmoo site (Richman et al., 2002; Ziman et al., 1993). The localized Cdc42 orchestrates the morphological development of a bud or shmoo by controlling the formation of oriented actin cables that direct transport of membrane vesicles and organelles, the assembly of a ring of septins that define the bud neck, and the specification of the site of exocytic vesicle fusion (Park and Bi, 2007; Wedlich-Soldner and Li, 2004). Cdc42 cycles between two guanine nucleotide (GTP and GDP) -bound states. This cycle is controlled by Cdc42’s guanine nucleotide exchange factor (GEF), Cdc24, and several GTPase-activating proteins (GAPs) (Park and Bi, 2007). In its GTP-bound form, Cdc42 interacts with a wide range of effectors that control downstream functions (Park and Bi, 2007). Effector interactions and the assembly of a polarized actin cytoskeleton also constitute feedback loops that enhance the accumulation of active Cdc42 during the initial symmetry breaking process (Wedlich-Soldner and Li, 2004).

After symmetry breaking at the initiation of budding, a polarized distribution of Cdc42 is stably maintained to ensure rapid growth of a unique bud for cell division. Although the Cdc42 polar cap appears to be stable, individual molecules of Cdc42, as well as the GEF Cdc24 and the adaptor protein Bem1, are highly dynamic, as shown by fluorescence recovery after photobleaching (FRAP) experiments (Wedlich-Soldner et al., 2004). Cdc42 exhibits a considerable membrane diffusion rate of 0.036 μm²/s, measured with a mutant, Cdc42Q61L, which is stably associated with the membrane and unable to enter the cytosol (Marco et al., 2007). For exploration of how a stable cap of Cdc42Q61L can be dynamically maintained, a mathematical model was built to describe balance of flux due to Cdc42Q61L membrane diffusion, internalization, and polarized delivery (Marco et al., 2007). This analysis concluded that the rate of internalization of Cdc42Q61L was optimized to achieve maximum polarity.

While the above work provides a useful framework for understanding how the polarized distribution of a membrane protein can be maintained at a cortical domain in a dynamic manner, it did not explain how cells maintain the dynamic distribution of Cdc42 in its native form. The fluorescence recovery rate of wild-type (WT) Cdc42 in FRAP measurements is an order of magnitude higher than that of Cdc42Q61L, in part because of the rapid exchange of Cdc42, but not Cdc42Q61L, molecules...
between the membrane and cytosolic pools (Wedlich-Soldner et al., 2004). This result implicates a possible role for Rdi1, the lone guanine-nucleotide dissociation inhibitor (GDI) for Rho family GTPases in yeast, in rapid Cdc42 recycling. GDI proteins are known to play a role in maintaining soluble GTPases in the cytosol through their ability to bind the GTPase’s COOH terminal prenyl group (Bustelo et al., 2007; DerMardirossian and Bokoch, 2005; Johnson et al., 2009). In yeast, Rdi1 has been shown to interact with Cdc42 at the membrane, extract Cdc42, and lead to an increase in its abundance in cytosolic fractions.

In this study, through both experimental exploration and mathematical modeling, we show that Rdi1 indeed plays an active role in fast Cdc42 recycling. A combination of the Rdi1-mediated recycling pathway with the actin-mediated endocytic recycling, which exhibit very different dynamic constants, determines the steady-state Cdc42 distribution. The dual-pathway model predicts that windows of the two recycling pathways must overlap spatially and be of similar sizes. The internalization rates for WT Cdc42 are not necessarily optimized for polarity but are instead tuned to give rise to different Cdc42 distributions that correlate with distinct morphogenetic fates.

RESULTS
Cdc42 in Budding Cells Is Recycled at the Polarized Site via Two Mechanisms
We began the study by testing the possibility that the yeast GDI, Rdi1, known to play a role in maintaining cytosolic Cdc42, is involved in fast Cdc42 recycling. GFP-Cdc42 was introduced under its native promoter into \( \text{\textit{Jrdi1}} \) mutant yeast cells, and FRAP data were obtained. For simplicity, “GFP” is omitted hereafter when we refer to fluorescently tagged Cdc42 or Cdc42 mutant proteins, as GFP was the only fluorescent tag for Cdc42 used in this study. GFP-Cdc42 is functional as it can fully rescue growth in \( \text{\textit{Jcdc42}} \) background (Roland Wedlich-Soldner, personal communication).

As in the previous work (Wedlich-Soldner et al., 2004), we focused on the polar cap of Cdc42 prior to bud emergence in a dividing cell population (Figure 1A) to avoid data heterogeneity.
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Figure 2. Modeling Dynamic Distribution of Cdc42 on the Cortex
Basic depiction of the model, where $D_i$ is the lateral diffusion in the membrane and $m$ and $n$ are the internalization rate inside and outside the delivery window, respectively. $\chi$ marks the delivery window, and $h$ is the rate of delivery of molecules to the membrane (for more details, see the Experimental Procedures and Supplemental Data). A shows the simplest scenario considering only one recycling pathway. B and C show the model with overlapping Rdi1 and actin-based delivery windows of the same size or of different sizes, respectively (see the Supplemental Data).

due to different cell-cycle stages. To determine the quantitative relationship between recycling parameters and the shape of Cdc42 steady-state distribution, we focused on cells that had already formed a stable Cdc42 polar cap. Cdc42 in $\text{Jrd1}^+$ cells exhibited a drastically reduced FRAP rate ($r = 1/t_{1/2}$) compared to Cdc42 in WT cells (Figures 1B and 1C) ($p < 10^{-10}$). The recycling defect of Cdc42 in $\text{Jrd1}^+$ can be mimicked by the Cdc42R66E mutant, which does not bind GDI (Gibson and Wilson-Delfosse, 2001) ($p = 0.3$) (Figure 1D). Treatment of WT cells with latrunculin A (LatA) to remove the actin-dependent recycling pathway slowed recovery rate, as previously described (Wedlich-Soldner et al., 2004), while treatment of $\text{Jrd1}^+$ cells with LatA to depolymerize actin further reduced $r$. Membrane fluorescence did eventually recover under these conditions (see below); however, these measurements are not easily comparable to WT or $\text{Jrd1}^+$ cells as the Cdc42 polar cap rapidly dissipated in $\text{Jrd1}^+$ cells treated with LatA, yielding nonpolarized cells (Figure 1E). This residual recovery persisted in the presence of cyclohexamide and thus is not a result of new protein synthesis (data not shown). The finding that Cdc42 remained polarized in either $\text{Jrd1}^+$ cells, or WT cells treated with LatA, but not in $\text{Jrd1}^+$ cells treated with LatA (Figures 1E and 1F; see Figure S1A available online), suggests that Rdi1 and actin play redundant but essential roles in maintaining Cdc42 polarization.

As LatA disrupts both actin cables and actin patches, the latter being endocytic structures, we asked whether endocytosis is important for Cdc42 recycling by using the temperature sensitive $\text{arp3-2}$ mutant (Winter et al., 1997). This mutant disrupts the Arp2/3 complex, the actin nucleation factor required for assembly of actin patches, at the nonpermissive temperature 35°C. Similar to the effect of LatA alone, Cdc42 recycling was slowed in $\text{arp3-2}$ at 35°C. Recycling of Cdc42R66E in $\text{arp3-2}$ was also impaired at 35°C, but not at the permissive temperature (Figure 1D). The recycling rate of Cdc42 in WT cells was roughly the sum of the rate in LatA-treated or $\text{arp3-2}$-treated cells with that of Cdc42 in $\text{Jrd1}^+$ or Cdc42R66E (Figure S1B). These data suggest that endocytosis and the GDI work in parallel to control Cdc42 recycling at the polar cap in WT cells. The GDI-based mechanism accounts for fast recycling of Cdc42, while endocytosis represents a slow recycling pathway.

Overview of the Steady-State Model of Cdc42 Polar Cap
In order to understand at a quantitative level how each and both recycling pathways contribute to the dynamic maintenance of a Cdc42 polar cap, we used an approach that combined quantitative imaging with mathematical modeling. The model shares the same framework as the previous one (Marco et al., 2007) but has several differences. As in the previous model, the flux of Cdc42 is governed by membrane diffusion ($D_i$), a window in the plasma membrane where Cdc42 is delivered to the cortex from the internal pool with rate ($h$), and the rate of internalization of Cdc42 from the cortex inside ($m$) and outside ($n$) this window (Figure 2A, Equation 1, and Supplemental Data). The assumption of a discrete window for polarized delivery is a simplification of the reality but is justified based on the sharp polarized distribution of structures involved in Cdc42 recycling (Figure S2A) and excellent fit between model output and experimental measurements (see below).

We used a combination of FRAP and imaging experiments to extract model parameters (Supplemental Data and Figure S2B). We then applied these model parameters to a steady-state version of Equation 1 to examine the effect of the parameters on the steady-state distribution of Cdc42, after the initial establishment of polarity. In the case of WT Cdc42, the model had to be expanded from the previous version (Marco et al., 2007) to describe two recycling pathways (Figures 2B and 2C). Each of the above parameters, with the exception of $D_b$, has in theory two values, each corresponding to one of the pathways (Figures 2B and 2C). $D_b$ was held constant between the two pathways, as previous analysis suggested that this was largely governed by the prenyl group that anchors Cdc42 into the membrane (Marco et al., 2007). As opposed to the previous model, $m$ and $n$ were assessed separately in this study. $m$ was also independently validated by inverse FRAP (iFRAP) (see below). Before considering the model of dual pathways, we first assessed each pathway individually: $\text{Jrd1}^+$ cells enabled isolated analysis of Cdc42 polar...
cap maintained by actin-based recycling, while LatA treatment to remove actin allowed specific assessment of the Rdi-dependent mechanism.

Modeling the Steady-State Cdc42 Cap through Actin or Rdi1-Mediated Recycling

To assess the situation in which there is only actin-based recycling (Cdc42 in \( \Delta r d i 1 \)), we determined the delivery window by measuring the distribution of the formin protein Bni1 on the plasma membrane (Figure 3A). Bni1 nucleates the formation of actin cables and stays at actin barbed ends as the filaments elongate (Evangelista et al., 2003). Therefore, the Bni1 distribution on the cortex is likely to correlate with the ends of actin cables and thus the window of cable-based delivery. The average full width half maximum (FWHM) of a Gaussian fit of the perimeter line-scan of cells expressing Bni1-GFP was 13% of the cell perimeter in WT cells; thus, the total width of the actin-based delivery window was roughly 26% of the perimeter (Figure 3A). The Bni1 distribution was not dramatically changed in \( \Delta r d i 1 \) cells or upon expression of Cdc42\(^{266L} \) (Figure 3A). The Bni1 distribution was slightly narrower than polarized Cdc42 (Figures 3A and S3), consistent with spreading due to membrane diffusion of Cdc42 once delivered to the cortex.

Using this delivery window size, we applied the partial differential equation-based model to our time-dependent FRAP data and the relative integrated membrane-bound Cdc42 intensity inside and outside this window to arrive at a unique solution of \( m \), \( n \), and \( h \) for each cell (Figure 3D) (see the Supplemental Data). An output of the model is the amount of internal Cdc42 relative to the total Cdc42 (\( G_c \)), (see the Supplemental Data). Comparison of the computed \( G_c \) to the experimentally measured value provides an independent validation of the model and was remarkably accurate on a cell-to-cell basis (Figure 3B).

As a second validation, values of internalization rate inside the delivery window (\( m \)) obtained from the modeled FRAP data were compared to fluorescence loss rates in independent iFRAP experiments. In iFRAP, the internal (cytosolic + internal membrane-bound) Cdc42 was photobleached, and the rate of fluorescence decrease at the polar cap was monitored as molecules left this region (for example, see Figure 3C). As opposed to FRAP, which is defined by a combination of events described by Equation 1, we expect that the rate of fluorescence loss in iFRAP is an approximation for \( m \), because only 10% to 20% of Cdc42 molecules are present in the polar cap, while a vast abundance of Cdc42 molecules take part in the equilibrium between cytosol and the cap. Indeed, the rate of Cdc42 fluorescence loss measured by iFRAP matched well with \( m \) values obtained by modeling of FRAP data (Figure 3D).

The model predicted the ratio of Cdc42 internalization rates by the actin-based pathway inside and outside the delivery window (\( m/n \)) to be 2.8 ± 0.3. Because of a low amount of Cdc42 outside the delivery window and thus a reduced signal to noise in fluorescence measurements, it was not possible to validate internalization rate outside the window (\( n \)) by iFRAP. To assess the ratio \( m/n \) experimentally, we measured the density of actin patches inside and outside the delivery window with two-color confocal imaging of cells expressing Bni1-GFP and mCherry-tagged Arc40, a subunit of the Arp2/3 complex concentrated in actin patches. The ratio of actin patches inside to outside the Bni1-defined window was 3.1 ± 0.2 (\( n = 18 \)) (Figure 3E), in close agreement with the ratio of \( m/n \) calculated by the model (\( p = 0.35 \)). Along with the FRAP data in the arp3-2 strain, this supports the idea that Cdc42 is internalized by endocytosis through actin patches at rates proportional to the local density of actin patches. The strong polarization of Bni1 (Figure 3A) and actin patches (Figure S2) justifies the use of a discrete, window-based model for describing steady-state polarity.

As opposed to the actin-dependent recycling system, where the window of delivery is marked by the formin Bni1, the basis of the delivery window size for the Rdi1 pathway was unknown. To explore the simplest possibility, we analyzed Cdc42 FRAP data in cells treated with LatA with the model using the same window size as the actin-based pathway. Recycling parameters were calculated by application of the model to the FRAP data and are shown in Figure 3D. \( h \) for the Rdi1 pathway was higher than that for the actin-based pathway, as \( w \) were similar between the two pathways. Values of \( G_c \) as an output of the model matched the experimentally measured values well (Figure 3B). The model-calculated values of \( m \) were slightly higher than, though in a same range as, rates from iFRAP in cells treated with LatA (Figure 3D). We also examined the possibility that the Rdi1 recycling window was larger or smaller in the dual recycling model (see below).

A Model of the Steady-State Cdc42 Polar Cap Maintained by Dual Recycling

To explore how the two recycling mechanisms work together to dynamically regulate Cdc42 distribution at the cortex of WT cells, we again first applied the simplest possible scenario: that the two recycling mechanisms are spatially overlapping and employ concentric delivery windows of the same size (depicted in Figure 2B). When two overlapping windows of the same size are considered mathematically, their rates become additive (see the Supplemental Data); i.e., the overall recycling parameters of \( m \), \( n \), and \( h \) for WT Cdc42 are expected to be the sum of parameters from the two individual pathways (Figure 2B).

With the above consideration, we applied the single-pathway model to FRAP data of Cdc42 in WT yeast cells, using a window size of 26% of the perimeter. Values of \( G_c \) predicted from the model agreed well with experimental \( G_c \) measurements (Figure 3B). The average value of \( G_c \) for Cdc42 in WT cells (61%) was qualitatively consistent with the high percentage of cytosolic Cdc42 found in a fractionation experiment (Wedlich-Soldner et al., 2004). Interestingly, we indeed observed that the values of \( n \), \( h \), and \( m \) calculated from Cdc42 FRAP and imaging data in WT yeast cells using a single-pathway model were statistically equivalent to the distributions obtained by simply adding all possible combination of parameters measured for \( \Delta r d i 1 \) cells and cells treated with LatA (Figure 3D). In addition, the values of \( m \) predicted from the model for treatment of WT Cdc42 with a single pathway or \( m \) resulting from the sum of \( m_1 \) and \( m_2 \) from the two individual pathways both closely matched internalization rate of Cdc42 in WT cells measured independently by iFRAP (Figure 3D). This analysis suggests that dynamics of Cdc42 in WT cells can be adequately described experimentally and mathematically as a sum of the Rdi1-dependent and actin-dependent pathways.
Figure 3. Determination of Cdc42 Recycling Parameters

(A) The window of the actin-based recycling mechanism was estimated by the location of the formin Bni1. Example images of Bni1-GFP are shown, along with an example fit of a perimeter trace to a Gaussian model. The window width is approximated as two times the full width half maximum (FWHM) of the Gaussian distribution. The average and standard error of the mean (SEM) of fits of n between 19 to 21 cells is shown. Bni1-GFP was used in WT and Drd1 cells, while Bni1-mCherry was used in cells expressing GFP-Cdc42Q61L.

(B) Relationship of $G_c$, or percentage of internal Cdc42 relative to total, as an output of the model relative to the experimental measurement.

(C) Example iFRAP curves, along with a montage of an example iFRAP measurement for a cell expressing GFP-Cdc42 in Drd1. The scale bar represents 2.0 μm. The bleached region is circled in red. Fluorescence loss is measured in the cap.

(D) Output of model parameters. Model calculated values of internalization rate inside the window ($\bar{m}$) (in black) were compared to values measured experimentally by inverse FRAP (iFRAP) (in red). The values shown for the sum of the individual pathways represent all possible sums of all combinations. Box width represents the SEM, and whiskers represent the standard deviation (SD).

(E) Ratio of internalization rate inside the window to rate outside the window ($\bar{m}/\bar{n}$) from modeled parameters for the actin recycling pathway compared to the measured ratio of density of actin patches (using Arc40-mCherry) inside and outside the window defined by Bni1. Box width represents the SEM, and whiskers represent the SD. An example of a summed time series for the actin patch marker Arc40-mCherry is shown, with the window marked (same cell as the left cell in A). The scale bar represents 2.0 μm.
While the choice of identical windows for the two pathways has so far enabled excellent agreement between model predictions and experimental measurements, we explored the possibility that the Rdi1-based exchange window may be smaller or larger than the actin-based delivery window (Figure 2C). We reworked the model to allow for combination of two pathways of different sizes (Figure 2C), with the window size of the actin-dependent delivery held constant (see Equations S18–S22 in the Supplemental Data). Because the rate of internalization measured for Cdc42 in WT cells by iFRAP and the overall recycling rate as measured by FRAP were roughly the sum of the rates for the individual pathways (Figures 1, 3D, and S1) and the Cdc42 polar caps are generally symmetric, we know to good approximation that the actin-dependent and Rdi1-dependent recycling windows are concentric. Furthermore, if the individual recycling pathways are described correctly, their combination should effectively recapitulate the observed Cdc42 distribution in WT cells.

FRAP data of LatA-treated cells were modeled with the use of a smaller or larger window than previously employed. We then combined the new parameters of the Rdi1 recycling pathway with the actin pathway parameters in a dual recycling pathway scenario, and calculated steady-state distributions (Figure 4). Combination of parameters from the two pathways with a smaller Rdi1 pathway window than that of the actin pathway resulted in a predicted Cdc42 cortical distribution with a valley in Cdc42 intensity at the center of the cap (Figures 4D and 4E). This pattern of Cdc42 distribution was not observed in WT cells. If the Rdi1-based delivery window was assumed to be larger than that of the actin-based recycling, the resultant WT Cdc42 distribution upon combination of the pathways was very wide, covering nearly half of the membrane (Figures 4D and 4E). This wide distribution also was not observed experimentally (Figure S3). In addition, the average $G_c$ values calculated from the resultant steady-state distribution after combination of the smaller or larger Rdi1-mediated recycling window with the actin recycling window were 0.34 and 0.42, respectively, both far outside the range of experimentally measured $G_c$ for Cdc42 in WT cells (average 61%) (Figure 3B). In contrast, direct combination of recycling parameters from individual actin and Rdi1-dependent processes using identical window sizes (26% of perimeter) produced a distribution identical to that obtained by modeling FRAP data of WT Cdc42 with a single-window model (as depicted in Figure 2A; Figure 4E). We note that the steady-state distributions calculated from the model parameters fit the experimental distributions well, despite the noise in the experimental distributions (Figure 4F). On the basis of these analyses, the only way to accurately represent Cdc42 distributions in WT cells through two independent recycling pathways is to assume concentric windows of similar sizes.

One point of interest is the observed fluorescence recovery, albeit slow, even in the absence of both Rdi1 and actin (Figures 1C and 1D). To assess the potential impact of a third recycling pathway on our system, we revised the model to include a large window encompassing the entire cell surface, as recovery by this pathway was uniform and did not maintain polarity (see the Supplemental Data, section 2.5). We applied the revised model to $d_{rdi1}$ cells treated with LatA, and the resultant value of $h$ was very small: $0.00023 \pm 0.00005 \, [1/\text{um}^2\text{sec}]$ (Figure S4). The order of magnitude lower value of the delivery rate observed here compared to those for the actin or Rdi1-based pathway allowed us to neglect this mechanism in the model.

Figure 4. Effect of Varying Window Size for Rdi1-Dependent Recycling on Cdc42 Distribution
A–C) Example images depicting the orientations used for examination of the Cdc42 distribution. The red line in (B) represents the perimeter used as the x axis in D and E. “V” marks a vacuole. (D and E) Normalized output obtained by combining parameters for actin-based recycling with parameters for the Rdi1-pathway and using the dual-pathway model (Figure 2C). The actin pathway window size was held constant (26% perimeter), and Rdi pathway window sizes were varied as labeled. For comparison, the distribution obtained by modeling of Cdc42 in WT to a single pathway (window size: 26% perimeter) is shown. The y axis represents the protein abundance in arbitrary units, while the x axis represents the perimeter in scaled arbitrary units. Intensity map in (E) follows the scale used in (C). (F) Overlay of steady-state Cdc42 distributions observed experimentally in individual WT cells (dots) and those as calculated using parameters extracted from imaging and FRAP data of the same cells (smooth line).
The Cdc42 GTPase Cycle Regulates the Two Recycling Pathways

The result that two windows of Cdc42 recycling must colocalize on the cortex and be of similar size is a useful and unexpected finding of the model. A potential mechanism underlying their colocalization may be that both pathways rely upon the Cdc42 GTPase cycle, since Cdc42 mutants locked in either nucleotide-bound state (Cdc42Q61L or Cdc42D57Y) exhibited much reduced FRAP rates compared to WT Cdc42 (Wedlich-Soldner et al., 2003, 2004). The mutant FRAP rates were not further reduced in the Δrdi1 mutant (Figure 5A), suggesting that Rdi1 no longer participates in the slow recycling of these mutants.

We tested whether Cdc42Q61L and Cdc42D57Y mutant proteins might be deficient in GDI-complex formation in the cytosol by using live-cell fluorescence cross-correlation spectroscopy (FCCS) (Bacia et al., 2006) (for methods, see the Supplemental Data). FCCS assesses protein complex formation by measuring codiffusion, or cross-correlation, of red- and green-labeled species. We showed previously that cross-correlation between mCherry and GFP-tagged proteins expressed at their endogenous levels in live yeast cells is an effective method to quantitatively examine cytosolic interactions (Slaughter et al., 2007, 2008). Strong cross-correlation (20%) was indeed observed between GFP-Cdc42 and Rdi1-mCherry in the cytosol of polarized yeast cells, compared to that of the negative control of unlinked GFP and mCherry (0%). The positive control of linked GFP-mCherry (45%) was less than 100%, probably because of incomplete folding or dark states of autofluorescent proteins (also see Slaughter et al., 2007) (Figures 5B and 5C). We note that Rdi1-mCherry is an active protein, at least in its ability to...
rescue rapid recycling of Cdc42 in the \textit{Rdi1} background (data not shown). The observed cross-correlation of Rd1-mCherry with GFP-Cdc42 was abolished by the R66E or C188S mutation of Cdc42, both expected to disrupt the interaction of Cdc42 with GDI (Gibson and Wilson-Delfosse, 2001; Koch et al., 1997; Masuda et al., 1994). Surprisingly, both GFP-Cdc42^{D57Y} and GFP-Cdc42^{Q61L} exhibited drastically reduced levels of cross-correlation with Rd1-mCherry, suggesting that formation of the cytosolic Cdc42-Rd1 complex in vivo is dependent on an active GTPase cycle.

To determine whether defects in the GTPase cycle also affect the actin-based recycling of Cdc42, we used the model to analyze FRAP results of Cdc42^{Q61L} in the \textit{Rdi1} background. Values of \( G_c \) predicted with the model for Cdc42^{Q61L} agreed well with experimentally measured values and were lower than \( G_c \) values for WT Cdc42 (Figure 3B). The internalization rates inside the window \((m)\) predicted for Cdc42^{Q61L} in \textit{Rdi1} were much lower than those for WT Cdc42 in \textit{Rdi1} cells and agreed well with experimentally measured internalization rates from iFRAP, while the rate of delivery \((h)\) for Cdc42^{Q61L} was only slightly lower compared to that of WT Cdc42 in \textit{Rdi1} (Figure 3D). Interestingly, while \( m \) was reduced for Cdc42^{Q61L}, the rate of internalization outside the window \((n)\) predicted for Cdc42^{Q61L} in \textit{Rdi1} was statistically indistinguishable from the \( n \) value for WT Cdc42 in \textit{Rdi1} \((p = 0.61)\) (Figure 3D). These results suggest that the defect in Cdc42 GTPase activity impaired Cdc42^{Q61L} recycling via endocytosis within the delivery window, but had little effect outside the delivery window.

One potential mechanism by which the Q61L mutation could reduce the Cdc42 recycling rate is stabilization of the interaction of Cdc42-GTP with its effectors, should the effector-bound Cdc42 molecules be protected from endocytosis. To test this hypothesis, we introduced the mutation T35A to Cdc42^{Q61L}, which was shown to disrupt binding of CRIB-domain-containing effectors (Gladfelter et al., 2001), though T35 may also cause a structural change of the entire effector loop in the GTP-bound state of Cdc42 or Ras (Adams and Oswald, 2007; Spoerner et al., 2001). Supporting our hypothesis, Cdc42^{Q61L,T35A} was recycled at a much more rapid rate than Cdc42^{Q61L}, whereas the T35A mutant did not improve the recycling of the GDP-bound Cdc42^{Q61L}, which does not bind effectors (Figure 5A). Moreover, recycling of Cdc42^{Q61L,T35A} was not changed further in the \textit{Rdi1} mutant cells, suggesting that recycling of this mutant was through endocytosis in an Rd1-independent manner (Figure 5A). Taken together, the results described above suggest that the GTPase cycle of Cdc42 controls both pathways of recycling, by allowing the formation of the complex with Rd1 and by triggering the release of effectors to allow internalization through endocytosis. The latter mechanism was further supported by the effect of overexpression of the effector Gic2 on recycling (see below).

If the Cdc42 GTPase cycle is required for both recycling pathways, the recycling windows may colocalize with regulators of the Cdc42 GTPase cycle. While the GEF Cdc24 promotes the exchange of GDP for GTP on Cdc42, the GTPase activity of Cdc42 is controlled by several partially redundant GAP proteins (Knaus et al., 2007; Sopko et al., 2007; Zheng et al., 2007). We examined two of these GAPs, Bem2 and Bem3, because they localize to the bud tip and have been shown to regulate Cdc42 during bud formation (Knaus et al., 2007). Indeed, in the \textit{Δbem2Δbem3} double mutant, Cdc42 recycling rate was significantly reduced (Figure 5A). Furthermore, measurement of GFP-tagged GAP or GEF proteins showed that these proteins were concentrated as polar caps in polarized cells that displayed similar widths as that of Bni1 (Figure 5D). Strains expressing both Bni1-GFP and Bem2-mCherry or GFP-Cdc42 and Bem2-mCherry further demonstrated nearly perfect colocalization (Figure 5E). This colocalization was unaffected in \textit{Rdi1} cells (data not shown). In addition, Bem2 and Rga1 (but not Bem3 [Knaus et al., 2007]) remained strongly polarized upon LatA treatment (data not shown), consistent with the idea of the Rd1 window remaining in place independently of actin. The above data support the idea that the Cdc42 GTPase cycle is required for both recycling pathways and therefore may underlie the colocalization of the two recycling pathways.

**Relationship between the Rate of Internalization and Strength of Polarity**

The earlier model of Cdc42^{Q61L} maintenance in G1-arrested cells predicted a nonmonotonic relationship between the rate of internalization and polarity, measured as the ratio of the Cdc42 distribution peak height over the peak width (Marco et al., 2007). It was found that the observed rate of internalization inside the window \((m)\) falls on the value that correlated with maximal polarity. To investigate whether polarity is also optimized by the rate of internalization of WT Cdc42 during the physiological process of budding, we calculated a steady-state distribution (Equation S5, in nondimensional units) from average parameters of \( m, n, \) and \( h \) for each condition, from which polarity was calculated as previously defined (peak height divided by window width). To compare our results directly to the previous model, we fixed \( m/n \) ratios to the average of the data obtained for each specific condition and plotted the calculated theoretical polarity as a function of \( m \). This analysis showed that for Cdc42^{Q61L} in \textit{Rdi1}, the observed internalization rate inside the window \((m)\) was indeed found to be optimized for maximum polarity, consistent with the previous finding (Marco et al., 2007). However, this optimization was not found for WT Cdc42 recycled by either individual pathway or by dual pathways. In each of these cases, polarity maximum fell to the left of the measured values of \( m \). In other words, internalization rates were faster than those that would result in polarity maximum.

Since similar values of internalization rate outside the window \((n)\) for the actin-dependent recycling pathway in \textit{Rdi1} were obtained for Cdc42 and Cdc42^{Q61L}, it was more reasonable to compare the two forms of Cdc42 with \( n \) held constant as opposed to a constant \( m/n \) ratio as previously assumed. Figure 6E plots polarity as a function of \( m \) for fixed \( n \) and \( h \) (note that the measured values of \( h \) were similar but not identical for the two conditions shown here, and this plot used their average). The resulting plot demonstrates that for these values of \( n \) and \( h \), polarity will increase monotonically with reduced \( m \). This comparison can be extended to include also the recycling of Cdc42 by the Rd1 pathway (i.e., in the presence of LatA). In this case, the internalization rate outside the window \((n)\) was similar between all three conditions, but the rate of delivery \((h)\) was different, making possible a three-dimensional plot.
examining polarity as a function of $m$ and $h$ for fixed $n$ (Figure S5A). The 3D plot confirms the generality that a reduced rate of internalization inside the window relative to other parameters increases polarity. To explore the relationship of all model parameters to polarity in general, we searched parameter space for combinations of $m$, $n$, and $h$ that would satisfy specified requirements for a polarized system at three values of $D$: 0.36, 0.036, and 0.0036 $\mu$m$^2$/sec. The three-dimensional parameter-space plot and its projections are shown in Figures S5B and S5C, and the result is discussed in detail in the Supplemental Data.

Recycling Parameters Affect the Shape of the Cdc42 Polar Cap Distribution and Subsequent Morphogenesis

As the observed rates of internalization of WT Cdc42 do not result in maximal polarity, we tested whether these rates might instead be related to adaptation to specific morphogenetic functions. We first compared the steady-state distribution of Cdc42 or Cdc42Q61L on the cortex, modeled with average values of $m$, $n$, and $h$ obtained for each of the conditions indicated (Equation S5) (Figure 7A). The height of the distribution was increased for Cdc42Q61L relative to WT Cdc42. At the opposite end, Cdc42 in WT cells treated with LatA (thus recycled by the Rdi1 pathway) had a reduction in the amount of Cdc42 within the window relative to the total Cdc42. The qualitative trend of the above comparison of the modeled Cdc42 distributions was also confirmed with imaging of Cdc42 intensity within the window relative to total Cdc42 (Figure 7B).

Interestingly, the predicted WT Cdc42 distribution had a distinct shape from that of Cdc42Q61L; the WT Cdc42 membrane distribution was broader inside the window than the distribution of Cdc42Q61L, giving the former a more box-like shape. To quantify this difference through a single parameter, we chose the slope value at half the distance from the cap center to the window edge in the average WT Cdc42 distribution and termed this the critical slope. We then calculated the distance from the center of the polar cap to where this slope was achieved for each modeled cell. The farther from the center of the cap that the membrane distribution reaches this critical slope, the more “box-like” the distribution is inside the cap. This method of quantification allows for examination of curves without the need to force their fit to a function. The pointed nature of the distribution for Cdc42Q61L is revealed in this analysis compared to other conditions examined (Figure 7C).

As Cdc42 establishes the site of polarized secretion in yeast through multiple downstream mechanisms (see the Introduction), the shape and strength of its distribution are likely to be the main determinant of the shape of polarized growth. The more box-like polar cap observed with WT Cdc42 compared to that of Cdc42Q61L may ensure that growth is better spread out within the polar cap to form a round rather than pointed bud. To examine this possibility, we labeled live cell surfaces with fluorescent mannoprotein marker concanavalin-A (Tkacz et al., 1971; Tkacz and Lampen, 1972). Using confocal imaging, we indeed observed more pointed buds for cells expressing Cdc42Q61L than for cells expressing WT Cdc42 (Figure 7D). This effect was not due to a difference in the expression levels of Cdc42 and Cdc42Q61L (Figure S6A). The difference in bud shape was quantified by examination of the relationship between the length and width of buds. Larger length relative to width was observed for cells expressing Cdc42Q61L than those expressing WT Cdc42 in buds of all sizes (Figures 7D–7F). While in large buds this could be explained by a lack of depolarization in cells expressing Cdc42Q61L at a later cell-cycle stage, the difference was statistically significant in small buds with widths less than 3 μm (Figure 7F). This result suggests that a higher internalization rate for WT Cdc42 within the delivery window is important for achieving a round bud morphology.

To test further whether tuning $m$ can modulate bud morphology, we took FRAP data and modeled Cdc42 recycling in the presence of an increased level of the CRIB domain-containing effector Gic2 (Brown et al., 1997; Chen et al., 1997) expressed under the GAL1 promoter, since effector binding may negatively regulate Cdc42 internalization (Figure 5). Gic2 overexpression indeed slowed recycling of Cdc42 (Figure S6B). Moreover, the internalization rate $m$ was reduced while the internalization
Figure 7. Effects of Cdc42 Recycling on Polarized Morphogenesis

(A) The steady-state Cdc42 distribution calculated from average m, n, and h values is shown for each condition. The y axis represents the relative protein abundance in arbitrary units, while the x axis represents the perimeter in scaled arbitrary units. Window size is scaled to 1.0 (see the Supplemental Data).

(B) Relative intensity (arbitrary scale) of Cdc42 inside the window relative to total Cdc42 (membrane and internal), both from experimental measurements and the model-calculated steady-state distributions. Average and SEM are shown.

(C) The distance from the cap center at which a critical slope is reached (see the main text) is shown. Average and SEM are shown.

(D) Cells labeled with FITC-concanavalin-A to mark the cell wall. Representative images are shown. The scale bar represents 2.0 μm.

(E) Plots of y versus x in cells expressing Cdc42 or Cdc42Q61L stained as described in D. Each point represents a measured cell.

(F) Quantification of y/x ratio for buds with width smaller than 3 μm. Box width represents the SEM, and whiskers represent the SD.

(G) Cdc42 localization and characteristic pointed shape of WT cells after 1.5 hr of treatment with 75 μM α-factor, compared to cycling cells. The scale bar represents 2.0 μm.
rate outside the window \( n \) was unchanged (Figure S6B). Again, we find that reduction in \( m \) relative to \( n \) resulted in a pointed steady-state Cdc42 distribution, and subsequent pointed morphology (Figures S6C and S6D). The value of \( h \) was also reduced, though reduction in \( h \) serves only to reduce the overall amplitude of the steady-state distribution but does not affect distribution shape (for given \( m \), \( n \), and cell size) (Figure S6E).

The pointed cell morphology due to experimentally reducing the internalization rate \( m \) by either expressing Cdc42\textsuperscript{Q61L} or overexpressing the effector Gic2 led us to ask whether tuning of the internalization rate occurs naturally as a means to achieve a pointed morphology in yeast. For example, in mating, a sharp shmoo tip is desired for efficient cell fusion (Madden and Snyder, 1998). We applied the model to FRAP data of cells arrested with the mating pheromone \( \alpha \) factor (Figure 7G). A slightly smaller window size was used (17%), as measured from the Bni1-GFP distribution in these cells (Figure S6F). The delivery rate \( h \) and internalization rate outside the window \( n \) for Cdc42 were found to be similar between pheromone-arrested cells and cycling cells, whereas the internalization rate inside the window \( m \) was significantly reduced (Figure 7H). This reduction in \( m \) was verified by independent iFRAP measurements of Cdc42 (red points in Figure 7H).

We next determined whether the reduction in internalization rate, \( m \), observed for WT Cdc42 in pheromone-arrested cells compared to that in cycling cells was due to a change in actin or Rdi1-based recycling. Interestingly, the iFRAP rate in \( Jrd1 \) cells showed no difference between cycling cells and shmooing cells (Figure 7I), while a reduced rate of internalization was observed in shmooing versus cycling cells upon treatment with LatA (Figure 7I), recapitulating the difference in Cdc42 internalization rates between WT shmooing and cycling cells. These data suggest that in the case of pointed shmoo, it is the Rdi1 pathway that is modulated to achieve a slower Cdc42 internalization rate, leading to a pointed Cdc42 distribution relative to cycling cells (Figures 7C, 7J, and S6H) and subsequent characteristic shmoo morphology.

**DISCUSSION**

The results presented above demonstrate that Cdc42 is recycled through two mechanistically distinct pathways, one involving membrane trafficking, and the other requiring the conserved Cdc42 regulator, GDI. While GDI is well known to bind and extract Cdc42 from the membrane into the cytosol (Cole et al., 2001; Li et al., 1995), the presence of two parallel pathways for Cdc42 recycling underscores the importance of this process and provides robustness for dynamically maintained cell polarity. In addition, the parameters of these recycling pathways can be tuned to control the shape of the Cdc42 distribution. Only wild-type Cdc42 recycled through dual pathways was able to maintain relatively high polarity as well as a “box-like” shape that ensures type of the Rdi1 deletion mutant strain in cycling cells may be explained by the presence of a redundant pathway for Cdc42 recycling (endocytosis). However, inhibition of both pathways led to rapid loss of Cdc42 polarity.

Our data suggest that the mechanisms by which the GTPase cycle of Cdc42 facilitates internalization of Cdc42 through the dual pathways are complex. With FCCS analysis, a strong interaction of Rdi1 with Cdc42 was observed in the cytosol of live yeast cells, consistent with its proposed mechanism of action. However, formation of the Rdi1-Cdc42 complex was drastically reduced by mutations that lock Cdc42 in the GTP or GDP bound forms. This result suggests that in vivo the GTPase cycle of Cdc42, but not a particular nucleotide-bound form, is important for the formation of Rdi1-Cdc42 complex, and that not all Cdc42 molecules in the mobile pool are free to bind Rdi1. Recent in vitro experiments in lipid bilayers found that the rate of dissociation of Cdc42-GDI complex from the membrane is unaffected by the nucleotide-bound state of Cdc42, even though the affinity of GDI for GDP-bound Cdc42 is nearly 10-fold higher than it is for GTP-bound Cdc42 (Johnson et al., 2009). Our results suggest that GTPase hydrolysis presents another layer of regulation in vivo where bound regulators or effectors may prevent the formation of the Cdc42-GDI complex and corresponding Cdc42 dissociation from the membrane.

The GTPase cycle also regulates Cdc42 recycling via endocytosis. The observation that the effector-binding mutation T35A increased the recycling rate of Cdc42\textsuperscript{Q61L} to near that of WT Cdc42 in \( Jrd1 \) cells suggests that simply abrogating effector binding, without restoring GTPase activity, was sufficient to enable Cdc42 to be recycled more efficiently through endocytosis. Although T35A might also lead to a gross structural change in Cdc42 that can affect its interaction with other regulatory proteins, the effect of overexpression of Gic2 supports the possibility that binding to specific or a wide range of effectors could protect Cdc42 from endocytic internalization. However, this mechanism does not explain the slow recycling of Cdc42\textsuperscript{D57Y}, which is not expected to bind effectors and consequently did not exhibit enhanced recycling in the presence of the T35A mutation. Slow recycling of Cdc42\textsuperscript{D57Y} relative to Cdc42\textsuperscript{Q61L,T35A} suggests that Cdc42 must be in the GTP-bound active form in order to efficiently enter or stimulate the formation of endocytic structures, consistent with the known role of active Cdc42 in stimulating the formation of cortical actin patches (Lechler et al., 2001; Li et al., 1995).

The presence of two parallel pathways for Cdc42 recycling underscores the importance of this process and provides robustness for dynamically maintained cell polarity. In addition, the parameters of these recycling pathways can be tuned to control the shape of the Cdc42 distribution. Only wild-type Cdc42 recycled through dual pathways was able to maintain relatively high polarity as well as a “box-like” shape that ensures
growth of a round bud that is characteristic of the organism and may be optimized for receiving segregating organelles and nuclear materials during mitosis. In contrast, the reduced internalization rate inside the delivery window in Cdc42-261L-expressing cells results in a sharper Cdc42 peak that leads to a pointed bud shape. More interestingly, a reduction in the internalization rate inside the delivery window (m) was observed in cells undergoing pheromone response and predicts a sharper Cdc42 distribution than that in cycling cells. While other factors could also be involved, it is likely that the sharper Cdc42 distribution contributes to the pointed shmoo shape, which is required for efficient mating.

Taken together, our results suggest that parameters of Cdc42 recycling in yeast, especially the internalization rate within the polarized region, are adapted not to achieve maximum polarity, but to fulfill specific morphogenetic outcomes that may be advantageous to either vegetative growth or mating (Figure 7K). The observed difference between Cdc42-261L and wild-type Cdc42 may help explain why Rho GTPases are not simply membrane proteins but instead are evolutionarily conserved, nimble regulators of cell polarity. By involving the GTPase cycle in their rapid recycling, the dynamic parameters of Cdc42 recycling can potentially be tuned through GAPs, GEFs, and effectors. These in turn are targets of further upstream signals, thus allowing the GTPase protein distribution to adapt to different downstream morphogenetic requirements. As Cdc42 and regulators of its recycling are all highly conserved, the principles observed in this study may be extended to other systems of polarized morphogenesis.

**EXPERIMENTAL PROCEDURES**

**Yeast Culture**

Yeast strains used in this study are listed in Table S1. Yeast cells were grown in synthetic complete media to mid-log phase prior to analysis. For experiments with Lat-A (BioMol), 100 μM LatA was added for 10 to 15 min prior to data acquisition. Yeast cells were immobilized on glass for analysis. For experiments using the GAL1 promoter for Cdc42, cells were grown to mid-log phase in synthetic media with raffinose and then induced with 2% galactose for 1.5 to 2 hr. For overexpression of Gic2 with pGAL1, 2.5 hr of induction were used. For studies in pheromone-arrested cells, 75 μM α factor (Pi Proteomics) was added for 1.5 hr. For examination of bud shape, 200 μl cells were treated with 15 μl of 1 mg/ml FITC-concanavalin-A for 30 min. The cells were washed twice with phosphate-buffered saline and then imaged.

In most experiments, GFP-Cdc42 was controlled under the CDC42 promoter; however, because Cdc42-261L is dominantly lethal, the inducible GAL1 promoter was used in experiments involving comparisons with this mutant. Induction of Cdc42 expression with the GAL1 promoter for 90 to 120 min led to expression of Cdc42 proteins at a level similar to the expression level using the CDC42 promoter (Figure S6A).

**Imaging Experiments and Data Analysis**

Confocal images were acquired with an inverted Zeiss 200 m outfitted with a spinning-disc confocal system (Yokogawa) and a EM-CCD (Hamamatsu C9100). FRAP and iFRAP data were acquired with this system using an attached MicroPoint Mosaic bleaching system (Photonic Instruments) integrated with Metamorph acquisition software (Molecular Devices). Extraction of data from the images was performed with ImageJ software, and least-squares fitting was performed with OriginLab Pro. For determination of size of the membrane distribution to cell perimeter for Bni1, Bem2, Bem3, Rga1, and Cdc24, a line-scan was drawn around the cell perimeter of a confocal slice (Figure 3C) and fit to a Gaussian distribution. Total distribution width was approximated as two times the FWHM of the Gaussian distribution.

**FCCS**

Fluorescence cross-correlation spectroscopy (Bacia et al., 2006) data were obtained in live yeast as previously described (Slaughter et al., 2007, 2008). In brief, GFP-Cdc42 and Rd1-mCherry were excited with the 488 nm and 561 nm laser lines, respectively, of a Zeiss ConfoCor 3. Emission filters were BP 505–540 and LP580 for the red and green channels, respectively. An emission dichroic of HFT590 was used, compared to the HFT565 used previously, to minimize crosstalk from the green to red channel. Correlation curves were fit and amplitudes of the correlation curves were used to calculate a percent bound fraction of the two species (Rigler et al., 1988).

**General Model**

We consider a model of Cdc42 protein dynamics on the surface of a polarized yeast cell (Marco et al., 2007). The dynamics of the distribution of Cdc42 along the membrane can be represented by Equation 1, where \( f(r, \phi, t) \) denotes the surface (membrane) density of Cdc42, \( D_v \frac{df}{dt} \) describes the diffusion along the membrane, and \( m \) represents the internalization rate inside the delivery window. The parameter \( n \) is the internalization rate outside the window, \( h \) is the rate of delivery of molecules to the polar cap, and \( F_c \) is the amount of cytosolic (or internal) Cdc42 (see Figure 2). \( \gamma \) equals 1 inside the confines of the window and equals 0 outside.

\[
\frac{df}{dt} = -mf + n(1 - \gamma)\frac{f}{h + F_c}.
\] (1)

As the FRAP process is essentially nonstationary, we use Equation 1 as a starting point and use time dependent FRAP data along with imaging to determine model parameters (Supplemental Data). We then choose to use model parameters from FRAP data to examine a steady-state version of Equation 1 for description of a polarized protein undergoing dynamic equilibrium at steady state, and not during initial stages of polarity establishment. All parameters except the diffusion coefficient are found from the combination of FRAP and steady-state imaging experiments as described in the Supplemental Data. The value of \( D_v \) of 0.036 μm²/s was used as published (Marco et al., 2007).

For details of imaging experiments, data analyses, and FCCS, see the Supplemental Data.

**SUPPLEMENTAL DATA**

Supplemental Data include Supplemental Experimental Procedures, seven figures, and one table and can be found with this article online at http://www.cell.com/developmental-cell/supplemental/5153-5807/0900445-6.

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