Temporal and spatial regulation of Rho-type guanine-nucleotide exchange factors: the yeast perspective

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Cell polarity is involved in almost every aspect of cell and developmental biology and is functionally important for differentiation, proliferation, and morphogenesis in single-cell as well as multicellular organisms (Drubin 2000). Establishment of cell polarity is a multistep process that is regulated by internal cues or extracellular signals. It requires the selection of a specific site at the cell cortex (“landmark”) followed by the recruitment and activation of components involved in actin or microtubule polymerization at that site. These cytoskeletal rearrangements lead to various forms of polarized growth, including asymmetric cell division, phagocytosis, the formation of filopodia, lamellipodia or stress fibers, and in the most extreme cases, the formation of axons and dendrites. Sustained polar growth further requires the stabilization of the polarization machinery at the cortex to maintain cytoskeletal polarization and targeting of secretion vesicles toward that region. Finally, the polarization machinery must be disassembled and/or inactivated to end the polar growth phase.

Actin rearrangements are regulated by small GTPases of the Rho and Ras subfamilies (Hall 1998), which are activated by a variety of growth factors, cytokines, and adhesion molecules. GTPases function as molecular switches, being inactive when bound to GDP and active when bound to GTP. Exchange of GTP for GDP induces a conformational change, which allows the effector domain to interact with downstream effectors. Many targets of Rho-GTPases have been identified, which together regulate the dynamic changes of the actin cytoskeleton in response to various intra- and extracellular signals. Several recent reviews have summarized and discussed the function of these downstream effectors in regulating the various stages of actin dynamics (Van Aelst and D’Souza-Schorey 1997; Aspenstrom 1999; Johnson 1999; Kaibuchi et al. 1999). However, much less is known about the regulation of the GTPases themselves. This is a challenging task, as Rho-type GTPases need to be controlled both temporally and spatially. For example, in a migrating cell, the same GTPase that is activated at the leading edge must be turned off at the trailing edge. The aim of this review is to highlight recent advances and emerging concepts describing the spatial and temporal regulation of guanine–nucleotide exchange factors (GEFs), the key regulators of Rho-type GTPases.

Known regulators of GTPases

Biochemical and genetic studies have led to the identification of three classes of regulatory proteins that control the nucleotide state of Rho family proteins. These are the guanine–nucleotide exchange factors (GEFs), the GTPase activating proteins (GAPs), and the guanine–nucleotide dissociation inhibitors (GDIs; Fig. 1). In addition, Rho-GTPases are isoprenylated at their C termini by specific geranyl–geranyl- (GGTases) or farnesyl-transferases (FTases), and this lipid modification may target them to membrane microdomains. Rho-GTPases form an inactive cytosolic complex with GDIs, in which the GDI masks the C-terminal isoprenyl group and thereby prevents membrane association of the GTPase. This complex is in equilibrium with the membrane-bound form of the GTPase. GDIs preferentially bind to the GDP-bound form and prevent nucleotide release, thereby maintaining the GTPases in the inactive state. At the membrane, exchange of nucleotide is stimulated by GEFs, which catalyze the exchange of GTP for GDP by facilitating the release of GDP and transiently stabilizing the activated GTPase from the GEF. The level of Rho-GTP can also be regulated by altering the activity of GAPs, which increase the intrinsic GTPase activity, thus favoring the conversion from the GTP- to the GDP-bound form.

In principle, local activation of GTPases in response to intra- or extracellular signals could occur by local activation of GEFs or inhibition of GAPs and/or GDIs. These different possibilities are not mutually exclusive, and it is likely that full activation of Rho-GTPases requires activation of GEFs as well as inactivation of their GAPs and/or GDIs. For example, the ERM proteins (ezrin,
radixin, and moesin) have been shown to interact with Rho-GDI and inhibit its activity (Takahashi et al. 1997). Thus, the regulated release of compartmentalized GTPase may increase recruitment and activation at the plasma membrane, possibly as part of an amplification loop triggered by the GTPase itself. Recent studies in budding yeast strongly support a critical role for temporal and spatial regulation of Cdc24p, the GEF for Cdc42p, during cellular polarization in response to internal and extracellular cues. Below, we will summarize current knowledge of cell polarity and Cdc42p regulation in yeast, and focus in particular on the regulation of its GEF Cdc24p. On the basis of the yeast model, we propose four separable steps that may be involved in the regulation of Rho-GEFs. In the last section of this review, we discuss a few selected examples to illustrate that these basic principles may be conserved in multicellular organisms.

Cell polarity and polarized growth in yeast

The yeast *Saccharomyces cerevisiae* exhibits polarized growth at several stages of its life cycle, making it an excellent model organism to study the biochemical pathways leading to rearrangements of the cytoskeleton. During vegetative growth, a bud forms late in the G1 phase of the cell cycle (Pruyne and Bretscher 2000). The position on the cell cortex where bud growth will be initiated (incipient bud site) is not random but is regulated by several gene products in a cell-type dependent manner (Chant 1999). For example, the transmembrane protein Bud10p/Axl2p is required to guide the budding machinery to the correct position in haploid cells (Halme et al. 1996; Roemer et al. 1996), whereas Bud8p and Bud9p mark the site of polarization in diploid cells (Zahner et al. 1996). During mating, a pointed projection coined shmoo develops to allow contact and fusion with a cell of the opposite mating type. In this case, cell polarity is determined by a gradient of mating pheromones secreted by the mating partner (Arkowitz 1999; Bähler and Peter 2000). On nitrogen starvation, yeast cells elongate from one pole, forming chains of linked cells that spread across the substratum (Kron and Gow 1995; Roemer et al. 1996). All these asymmetric growth processes require polarized organization of the actin cytoskeleton.
Central to the initiation of actin polarization is the local activation of the GTPase Cdc42p (Fig. 2A), and much will be learned if we understand when, where, and how Cdc42p is activated and inactivated. Regulation of Cdc42p involves its GEF Cdc24p, several putative GAPs (Bem2p, Bem3p, Rga1p, and Rga2p), the GDI Rdi1p, and a subunit of the geranyl-geranyl transferase I Cdc43p (Johnson 1999; Pruyne and Bretscher 2000). Whereas little is known about the regulation of Rdi1p or the GAP molecules, much progress has been made in understanding the spatial and temporal regulation of Cdc24p. Cdc24p is the sole essential GEF for Cdc42p (Bi et al. 2000). As do all Rho-GEFs, it contains a domain with strong similarity to the Dbl-family of exchange factors [residues 283–452, Dbl-homology domain or DH] and a nearby pleckstrin-homology domain [residues 472–681, PH]. Cdc24p also contains a calponin-homology [CH] domain in its N terminus [residues 154–226], which in some proteins has been implicated in binding to actin (Stradal et al. 1998). Finally, Cdc24p has two calcium-binding domains [residues 649–658 and 820–831]. Whereas the PH domain is thought to serve as a membrane-targeting signal (Bottomley et al. 1998), the function of the CH- and calcium-binding motifs in regulation of Cdc24p is not clear.

Activation of Cdc24p is required during all stages of the yeast life cycle that involve polarized growth, but its role is best studied during the cell cycle and in response to mating pheromones. We will thus focus the following discussion on the regulation of Cdc24p during these two stages.

Recruitment and activation of Cdc24p at bud emergence

Cells in the early G₁ phase of the cell cycle grow isotropically until they reach a critical size, at which time activation of the cyclin-dependent kinase (CDK) Cdc28p–Clnp initiates actin polarization: Cortical actin patches congregate at the selected site, and actin cables become oriented along the mother-bud axis, resulting in polarized secretion and bud emergence (Lew and Reed 1995). The role of Cdc28p–Clnp in triggering cell polarization at bud emergence is not understood, but recent evidence suggests that it may be required for activating Cdc42p [Gulli et al. 2000]. Surprisingly, Cdc42p is sequestered in the nucleus during the G₁ phase of the hap-

Figure 2. Mode of activation of Cdc24p, the yeast GEF for Cdc42p. (A) Upstream regulatory elements involved in Cdc24p activation during budding and mating. Budding: The presence of Bud5p, the GEF for Rsr1p/Bud1p at the preexisting budding site at the plasma membrane (internal landmark) allows local activation of Rsr1p/Bud1p. Once bound to GTP, Rsr1p/Bud1p recruits and likely activates Cdc24p. Thus, Cdc42p-mediated actin polarization during budding involves a cascade of GTPases. Mating: Secreted pheromones bind to G-protein-coupled mating receptors. On receptor engagement, Gα binds to GTP and dissociates from Gβγ. Far1p targets Cdc24p to Gβγ-activated receptor (extracellular signal-mediated landmark). In addition to bridging Cdc24p interaction to Gβγ, Far1p may activate Cdc24p. (B) Proposed model for the activation of Cdc24p. In the absence of any activating signal, the N-terminal calponin homology domain [CH] may inhibit the catalytic Dbl homology domain [DH] of Cdc24p. Binding of the linker proteins Rsr1p/Bud1p-GTP, Far1p, or as-yet-unidentified additional activators to the CH-domain may open Cdc24p to its active conformation by removing the N-terminal inhibitory domain, hence relieving the catalytic DH-domain.
loid cell cycle by binding to the adapter Far1p (Toenjes et al. 1999; Nern and Arkowitz 2000b; Shimada et al. 2000). Cdc28p–Clnp phosphorylates Far1p at bud emergence, leading to ubiquitin-mediated degradation of Far1p (Henchoz et al. 1997) and subsequent release of Cdc24p from the nucleus (Fig. 3; Nern and Arkowitz 2000b; Shimada et al. 2000). However, phosphorylation of Far1p and release of Cdc24p are not the only functions of Cdc28p–Clnp at bud emergence, because cells expressing a cytoplasmic form of Cdc24p [Cdc24p-m1; Nern and Arkowitz 1998] still require Cdc28p–Clnp to induce cellular polarization. Indeed, Cdc28p–Clnp activity is necessary to recruit cytoplasmic Cdc24p to the incipient bud site (Gulli et al. 2000). This membrane recruitment requires neither an intact actin cytoskeleton nor activated Cdc42p, indicating that Cdc28p–Clnp plays an as yet unidentified role at bud emergence (Gulli et al. 2000). Cdc28p–Clnp might phosphorylate Cdc24p, thereby inducing a change in its conformation that may allow its interaction with binding partners at the plasma membrane. Alternatively, Cdc28p–Clnp might phosphorylate components at the plasma membrane, which in turn, bind to Cdc24p.

Recent evidence suggests a crucial role for the small GTPase Rsr1p/Bud1p in the local activation of Cdc24p at bud emergence. Rsr1p/Bud1p was originally isolated as a multicopy suppressor of cdc24-4 mutant cells (Bender and Pringle 1989) and subsequently shown to bind directly to Cdc24p in a GTP-dependent manner (Park et al. 1997). Rsr1p/Bud1p is localized uniformly at the plasma membrane (Michelitch and Chant 1996), suggesting that its local activation may target Cdc24p to the incipient bud site. Indeed, Cdc24p localizes to random positions in rsr1/bud1Δ cells, and conversely, overexpressed Bud1p-GTP uniformly recruits Cdc24p to the plasma membrane (Y. Shimada, P. Wiget, and M. Peter, unpubl.). Rsr1p/Bud1p is regulated by its GEF Bud5p and its GAP Bud2p, both of which localize to the incipient bud site (Park et al. 1999). Interestingly, Bud5p colocalizes with the transmembrane protein Axl2p/Bud10p, a component of the budding landmark, and directly interacts with its cytoplasmic tail (H.-O. Park, pers. comm.), suggesting

![Figure 3. Regulation of Cdc24p membrane targeting during budding and mating. In the G1 phase of the cell cycle, Cdc24p is sequestered in the nucleus by Far1p. Pathway A: At bud emergence, activation of the G1 cyclin–CDK complex Cdc28p–Clnp triggers phosphorylation and subsequent ubiquitin-dependent degradation of Far1p. As a consequence, Cdc24p redistributes to the cytoplasm. In addition, Cdc28p–Clnp promotes the recruitment of Cdc24p to the preexisting budding site by an unknown mechanism. Recruitment and activation of Cdc24p by Rsr1p/Bud1p-GTP at the predetermined budding site allows polarization of the actin cytoskeleton toward that site. Pathway B: In the presence of a gradient of secreted pheromones, the kinase complex Cdc28p–Clnp is inhibited and Far1p is stabilized. The Far1p–Cdc24p complex is exported by the exportin Msn5p, and Far1p targets Cdc24p to Gβγ at the site of activated receptor. Recruitment and activation of Cdc24p then allows polarization of the actin cytoskeleton toward the mating partner.](genesdev.cshlp.org)
that Bud5p may be recruited to the incipient bud site by binding to Axl2p/Bud10p. Polarized localization of Bud5p and Bud2p are mutually interdependent (H.-O. Park, pers. comm.), and maintenance of Bud2p at the polarization site requires Rsr1p/Bud1p (Park et al. 1999). Thus, both the GAP and GEF of Rsr1p/Bud1p colocalize at the incipient bud site and may locally modulate the rapid cycling of Rsr1p/Bud1p between its GDP- and GTP-bound states. Taken together, these results suggest that Rsr1p/Bud1p is an important regulator of Cdc24p, which may recruit Cdc24p to the incipient bud site (Figs. 2A, 3). Importantly, preliminary evidence suggests that Rsr1p/Bud1p may not only recruit but also activate Cdc24p at the plasma membrane. Rsr1p/Bud1p interacts with the conserved CH motif in the N-terminal regulatory domain of Cdc24p (Fig. 2B; Y. Shimada and M. Peter, unpubl.), which is involved in its activation in vivo. We propose that the N-terminal domain of Cdc24p inhibits its catalytic DH-domain and that binding of Rsr1p/Bud1p to the CH domain of Cdc24p relieves this autoinhibition, thereby activating the GEF (Fig. 2B, see below). Thus, local activation of Rsr1p/Bud1p may spatially and temporally activate Cdc24p, implying that a cascade of two GTPases triggers polarization toward the bud site in the G1 phase of the cell cycle (Fig. 2A). Because deletion of BUD2 is lethal in cells lacking Cln1p and Cln2p (Benton et al. 1993; Cyrtova and Nasmyth 1993), it is tempting to speculate that the Rsr1p/Bud1p module is regulated by the Cdc28p–Clnp kinase. However, in contrast to Cdc24p and Cdc28p–Clnp, the Rsr1p/Bud1p GTPase module is not essential for viability, implying that other regulators must exist that activate Cdc24p in the absence of Rsr1p/Bud1p (Fig. 2B).

Recruitment and activation of Cdc24p in a morphogenetic gradient during mating

To ensure efficient conjugation, two mating partners have to grow toward each other for subsequent fusion. The two cells are able to locate each other by interpreting a pheromone gradient and to polarize their actin cytoskeleton in the direction of the highest pheromone concentration. Pheromones bind to cell-type specific seven-transmembrane receptors (Ste2p or Ste3p), which in turn activate the associated heterotrimeric G protein (α, β, γ encoded by GPA1, STE4, and STE18 respectively), by inducing its dissociation into Gα-GTP and Gβγ subunits (Sprague and Thorner 1992; Leberer et al. 1997). It has been estimated that as little as a 1% difference in receptor occupancy is sufficient to correctly orient the cytoskeleton in the morphogenetic gradient (Segall 1993), suggesting that mechanisms must exist that amplify small differences to stabilize the axis of polarization.

Formation of an oriented mating projection requires the same basic polarization machinery as bud emergence. However, activation of Cdc42p does not occur at the presumptive bud site but, instead, Cdc42p is specifically activated at a new site determined by the morphogenetic signal. Thus, cells have to override their internal budding program and target the polarization machinery to the site of maximal receptor activation. The spatial information of the pheromone gradient is transduced intracellularly by the Gβγ heterodimer, which guides the Cdc42p-module to the correct site. The critical effector mediating this response is Far1p, which binds via two separable domains to Gβγ and Cdc24p (Butty et al. 1998).

Thus, Far1p functions as an adapter, which recruits Cdc24p to activated Gβγ during mating (Figs. 2A, 3). Recent evidence suggests that Far1p may not only recruit Cdc24p to the correct site but that, like Rsr1p/Bud1p, Far1p binds to the CH-domain of Cdc24p (Nern and Arkowitz 1998) and may be involved in its activation at the plasma membrane. Thus, Far1p and Rsr1p/Bud1p may play analogous roles under different physiological conditions: Rsr1p/Bud1p may target and activate Cdc24p specifically at an intrinsic polarity cue (bud site) during the cell cycle, whereas Far1p may target and activate Cdc24p at a site determined by an extracellular signal (pheromone gradient) during mating (Fig. 2A). How is the positional information of the bud site overridden in the presence of a mating partner? Recent evidence provides some intriguing answers to this question (Fig. 3): Whereas activation of Cdc28p–Clnp at bud emergence triggers degradation of Far1p, inhibition of Cdc28p–Clnp by pheromones stabilizes Far1p (Blondel et al. 2000). This allows the exportin Msn5p to relocate the Far1p–Cdc24p complex into the cytoplasm (Blondel et al. 1999), where it can bind to Gβγ at the plasma membrane (Nern and Arkowitz 2000b, Shimada et al. 2000). We propose that when complexed with Far1p, Cdc24p is unable to interact with Rsr1p/Bud1p at the incipient bud site, thereby favoring targeting of Cdc24p to the mating site (Fig. 3). Thus, regulated nuclear retention of Cdc24p–Far1p complexes may allow cells to choose between a preexisting landmark and an extracellular signal-mediated landmark to establish polarity (O’Shea and Her skowizt 2000).

Stabilization of Cdc24p at the polarization site

After recruitment and activation of Cdc24p at a specific site at the plasma membrane, the polarization complex needs to be stabilized to maintain production of Cdc42p-GTP and, thus, polar growth. Binding to Cdc42p inhibits the intrinsic as well as GAP-stimulated GTPase activity of Rsr1p/Bud1p (Zheng et al. 1995; Park et al. 1997). Cdc24p thus stabilizes the GTP-bound form of Rsr1p/Bud1p, and this reinforces the Rsr1p/Bud1p–Cdc24p interaction (Zheng et al. 1995). Through a C-terminal domain, Cdc24p also interacts with the adapter molecule Bem1p (Peterson et al. 1994), which contains two SH3-domains (Chenevert et al. 1992). Interestingly, Cdc24p is able to localize to the incipient bud site in bem1Δ cells, but binding of Cdc24p to Bem1p is required to stabilize active Cdc42p at the plasma membrane (Gulli et al. 2000). As a consequence, bem1Δ cells fail to grow in a polar manner, implying that the continued production of Cdc42p-GTP is responsible for polar bud growth (Gulli et al. 2000). Although Bem1p is unable to activate Cdc24p
in vitro (Zheng et al. 1995), it is possible that binding of Bem1p to Cdc24p stabilizes its active conformation. Interestingly, Bem1p also interacts with activated Cdc42p [Butty et al. 1998; Bose et al. 2001], and recruitment of Bem1p to the polarization site is dependent on Cdc42p-GTP [A.-C. Butty and M. Peter, unpubl.], providing a positive feedback loop that locally stabilizes activated Cdc24p at the plasma membrane. In addition, Bem1p has been shown to interact preferentially with GDP-bound Rsr1p/Bud1p [Park et al. 1997]. Thus, after conversion of Rsr1p/Bud1p to its GDP-bound form, Bem1p may replace Rsr1p/Bud1p and stabilize activated Cdc24p complexes at the incipient bud site, thereby promoting efficient bud emergence. Consistent with such a model, a BUD5 mutation (which is expected to reduce Rsr1p/Bud1p-GTP levels) with a mildly deleterious mutation in BEM1 results in a severe growth defect at high temperatures and accumulation of round unpolarized cells (Chant et al. 1991).

How is activation of Cdc42p restricted to a single site? It is likely that site-specific activation together with amplification loops that stabilize activated Cdc24p at the plasma membrane prevent polarization toward additional sites. In addition, GAPs or other negative regulators may prevent Cdc42p from initiating polarization at other sites on the plasma membrane [Richman and Johnson 2000]. Intriguingly, Mdg1p, which was isolated in a genetic screen for high-gene-dosage suppressors of other sites on the plasma membrane (Richman and Johnson 2000). Likewise, certain cdc42 mutants exhibit hyperpolarized bud growth (cdc42V44A and cdc42-129V36T), and at least Cdc42p-V44A fails to interact with Cdc24p, Bem1p, and the effector Gic2p at their tips [Gulli et al. 2000]. Interestingly, phosphorylation of Cdc24p leads to its dissociation from the complex (Bose et al. 2001; Gulli et al. 2000). Importantly, this feedback mechanism may help to maintain the proper temporal order of the different steps at bud emergence because inactivation of Cdc24p could not occur before

**Inactivation of Cdc24p at the site of polarized growth**

During a normal cell cycle, Cdc24p is only transiently found at bud tips and dissociates at the end of polar bud growth [Nern and Arkowitz 1999; Shimada et al. 2000]. Importantly, removing Cdc24p from bud tips may require the activity of the PAK-like kinase Cla4p [Gulli et al. 2000]. Like Cdc24p, Cla4p binds to Bem1p, and for-
Cdc42p had been activated and a stable complex with Bem1p had been formed. However, a possible role for the Cdc42p GAPs in switching off polar growth after bud emergence is not excluded, and cells deleted for BEM3 and RGA1 display misshapen cells, some exhibiting an elongated morphology [Stevenson et al. 1995]. Therefore, dissociation of Cdc24p and GTP-hydrolysis on Cdc42p by GAPs may together inactivate Cdc42p after bud emergence.

General principles for the regulation of GEFs

Although in recent years much has been learned on the regulation of cell polarity in budding yeast, it is not clear to what extent similar mechanisms exist in multicellular organisms. Many mammalian Rho-GEFs have been identified, but in most cases their in vivo specificity toward Cdc42, Rac, or Rho GTPases is not clear. Nevertheless, although the mechanistic details may be different, we draw some parallels to the mechanisms of Cdc24p regulation in yeast. We propose that temporal and spatial regulation of mammalian Rho-GEFs can similarly be divided into four basic steps [Fig. 5]: First, the appropriate GEF has to be recruited to a specific site at the cell cortex, where, second, it will be activated. In yeast, the molecules that target Cdc24p to the correct site at the plasma membrane may also be responsible for its activation. Third, the activated GEF is stabilized at the plasma membrane by binding to an adapter protein, allowing for a stable platform for the local production of GTP-bound GTPase. Finally, inactivation of the GEF and/or GTP-hydrolysis terminates polarized GTPase signaling. Below, we discuss some selected examples to illustrate that these four steps may also be useful to describe the regulation of Rho-GEFs in multicellular organisms. For a detailed discussion of the regulation of mammalian GEFs, we alert the reader to recently published reviews (Cerione and Zheng 1996; Bustelo 2000; Symons and Settleman 2000).

Membrane recruitment of mammalian GEFs by binding to activated receptors

During yeast mating, polarization is initiated by specifically targeting Cdc24p to the site of receptor activation at the plasma membrane. Similarly, there is now increasing evidence for the translocation of many Rho-GEFs to the plasma membrane on receptor engagement [Bustelo 2000]. Among the best-characterized mammalian GEFs is Vav1, which binds to activated T-cell or B-cell receptors [TCR and BCR, respectively] to induce transcriptional activation and cytoskeletal reorganization (Bustelo 2000). In addition, the broadly expressed Vav2 and Vav3 isoforms are recruited to the plasma membrane by binding via their SH2 [ Src-homology 2] domains to the EGF and PDGF receptors in HeLa cells. In some cases, binding to the receptors is not direct but may be bridged by adapter molecules [see below]. Membrane association of Vav1 and other GEFs is likely to require the cooperative function of receptor binding and the N-terminal PH domain, which is able to bind phosphatidylinositol derivatives PI[3,4]P2 [PIP2] and PI[3,4,5]P3 [PIP3]. These PIPs are produced by PI kinases, which may themselves be activated by tyrosine kinase receptors [Katada et al. 1999]. Thus, maximal activation of Rho-type GEFs may involve the cooperative action of two signals: one recruiting the GEF to the membrane, and the other locally producing PIPs. The PH domain of yeast Cdc24p is important for the function of Cdc24p in vivo [Y. Shimada and M. Peter, unpubl.], but it is not known whether it interacts with PIPs. Finally, although some mammalian GEFs have been detected in the nucleus [Clevenger et al. 1995; Bertagnolo et al. 1998; Ichiba et al. 1999; Adam et al. 2000], no functional significance for their regulation has been described.

Recent evidence suggests that at least some mammalian GTPases may themselves be targeted independently to specific sites at the cell cortex [Gingras et al. 1998] and, thus, restrict GTPase activity to these cortical domains. It is possible that the C-terminal isoprene modification of GTPases contributes to their recruitment to membrane microdomains such as caveolae [Michaely et al. 1999]. Localized GTPase activation is of particular relevance for Rac and Cdc42 that play essential roles in directed cell migration and cell polarity. Coordinated translocation of both GEFs and GTPases to a specific membrane compartment may increase the efficiency of the nucleotide exchange reaction and prevent spontaneous activation of GTPases.

Membrane recruitment of Rho-GEFs in response to integrin activation: a role for Rap1-like GTPases

In the examples above, Vav is thought to directly bind to activated receptors, whereas in yeast, the recruitment of Cdc24p to the plasma membrane requires Far1p or the small GTPase Rsr1p/Bud1p. However, a role for Rap1-GTPases in integrin signaling has recently emerged [Caron et al. 2000; Katagiri et al. 2000; Reedquist et al. 2000]. The extracellular matrix [ECM] controls cell morphology and migration and signals directly to cells through transmembrane receptors such as integrins [Fig. 5A]. Ligand binding to integrins leads to their clustering and subsequent recruitment of actin filaments to the cytoplasmic domain of integrins [Schoenwaelder and Burridge 1999]. It was shown recently that adhesion of fibroblasts to fibronectin promotes transient Rac activation [Price et al. 1998; del Pozo et al. 2000] and more prolonged activation of Rho [Ren et al. 1999], demonstrating that integrin engagement by ligand leads to activation of Rho-type GTPases, presumably through activation of a GEF. Interestingly, mammalian Rap1 (also called Krev-1 or smg21A) has been implicated in controlling cellular adhesion by modulating integrin function [Reedquist et al. 2000], and Rap1p is required for integrin-dependent phagocytosis [Caron et al. 2000]. Rap1 is similar in sequence to yeast Rsr1p/Bud1p [Bender and Pringle 1989], and indeed, mammalian
### Stepwise regulation of Rho-GEFs

**Membrane recruitment**

| Receptors: | Receptors: |
|-----------|------------|
| ?         | Receptor Tyrosine kinases (RTKs): |
|           | EGF-R, PDGF-R... (Vav1) |
|           | other receptors: |
|           | BCR, TCR... (Vav1) |
| G protein-coupled receptors (GPCRs): | G protein-coupled receptors (GPCRs): |
| Ste2p, Ste3p; linker protein Fer1p | N-Furmyl peptide receptor (?) |
| Transmembrane protein: | G13α (p115RhoGEF) |
| Axl1p/Bud2p; linker protein Bud1p-GTP | Integrins: |
| Budding yeast | CD3+VLA-4, LFA-1; via Rap1? (?) |
| Phosphoinositides: | cMβ2; via Rap1? (?) |
| ? | Phosphoinositides: |
|  | PI3 Kinase binding (Pix) |

**Activation**

| Binding of proteins: | Binding of proteins: |
|----------------------|----------------------|
| Bud1p-GTP            | Adenomatous polyposis coli (Apc) |
| Far1p                | Phosphorylation: |
| ?                    | Tyrosine kinase receptors: EGF-R, PDGF-R... (Vav1) |
| Phosphorylation:      | Syk, Zap70, Jak, Fyn... (Vav1) |
| Cdc25p-Clnp?         | Binding of second messengers: |
| ?                    | cAMP (Epac1,2) |
| Binding of second messengers: | Phosphoinositides (Vav1) |
| Ca2+                 | |
| Phosphoinositides?   | |

**Stabilization**

| Binding to adapter proteins: | Binding to adapter proteins: |
|------------------------------|-------------------------------|
| Benj1p                       | LAT (Vav1) |
|                               | Dock/Nck; via Pak (Pix) |
|                               | Paxillin; via Pkl (Pix) |

**Inactivation**

| Phosphorylation: | Phosphorylation: |
|------------------|------------------|
| Pak/Mst (?)      | Pak/Mst (?) |
| Ubiquitin-dependent degradation: | Ubiquitin-dependent degradation: |
| ?                | SOCS1-induced degradation (Vav1) |

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**Figure 5.** Stepwise regulation of Rho-GEFs. Schematic representation of molecular mechanisms controlling the membrane recruitment [1], activation [2], stabilization [3], and inactivation [4] of Cdc24p in *Saccharomyces cerevisiae* and selected examples of Rho-GEFs in multicellular organisms. When known, the identity of the GEF involved is indicated in brackets and italics.
Rap1A can suppress the growth defect of cdc24-4 temperature sensitive [ts] cells [Ruggieri et al. 1992]. Putative Rap1-homologs are also present in Caenorhabditis elegans and Drosophila. In flies, Rap1 mutations disrupt cell migration and cause abnormal cell shapes [Asha et al. 1999], suggesting that Rap1 is a critical regulator of morphogenesis in vivo, possibly by regulating integrin function. The role of Rap1 in integrin signaling and control of morphogenesis is not clear, but by analogy to yeast, we speculate that Rap1 may recruit and activate a Rho-GEF. The endogenous GEFs that control activation of GTTPases by integrins are not known. However, Vav1 has been implicated in integrin signaling, and integrin-mediated adhesion increases tyrosine phosphorylation of Vav in several cell types [Bustelo 2000]. Several Vav isoforms contain a CH motif in their N-terminal domain, which at least in Cdc24p is required for the interaction with Rs1p/Bud1p. However, it is not known at present whether Vav or other Rho-GEFs are effectors of Rap1 GTTPases.

Membrane-recruitment of Rho-GEFs by activated heterotrimeric G-proteins

As in yeast, heterotrimeric G proteins are implicated in the establishment of cell polarity in several organisms and orient cell migration and asymmetric cell division in response to extracellular chemotactants or intracellular polarity cues. Neutrophils respond to a large number of chemotactic signals, including formylated peptides produced by bacteria, the C3a component of the complement system, and several chemokines produced by endothelial cells, immunocytes, and other inflammatory cells at sites of tissue injury [Weiner et al. 2000]. All these signals activate specific G protein-coupled receptors (GPCR) coupled to the Gi class of trimeric G-proteins, leading to dissociation of Go and Gβγ, which are thought to activate Cdc42 and Rac1 required for actin rearrangements during chemotaxis [Glogauer et al. 2000]. Interestingly, recent evidence suggests that Rac-dependent actin reorganization following activation of the N-formyl-methionyl-leucyl-phenylalanine tripeptide (FMLP) receptor [FPR] is mediated through Gβγ and class Ia PIP3 kinase [Belisle and Abo 2000], but the GEF mediating this response remains to be identified. Moreover, Go13, has been shown to bind and activate the p115Rho-GEF [Hart et al. 1998]. Thus, similar to the pheromone response in yeast, Rho-GEFs may be recruited to and activated at GPCRs by directly or indirectly interacting with activated heterotrimeric G proteins.

Spatial activation of specific heterotrimeric G proteins may also be responsible for orientation of the plane of cell cleavage during asymmetric divisions in early Drosophila and C. elegans embryos. Genetic evidence suggests that Goa and Gβγ are both required for spindle orientation in early C. elegans embryos [Gotta and Ahringer 2001]. Interestingly, Gβγ is important in regulating centrosome migration around the nucleus and, hence, in orienting the mitotic spindle, whereas Goa is required for asymmetric spindle positioning. Recent results from asymmetric cell divisions in the Drosophila nervous system now suggest that heterotrimeric G proteins may be locally activated in a receptor-independent manner [Schweissguth 2000]. Pins [partner of inscuteable] is required for oriented cell division and was found to specifically bind to Goi-subunits through its Go-Loco-domains [Siderovski et al. 1999]. Pins homologs are also found in man and C. elegans, but it is not known whether they are involved in the regulation of cell polarity. In addition, database searches have identified Go-Loco domains in several proteins [K. Hoffmann, pers. commun.], some of which may be involved in activating heterotrimeric G proteins in response to a wide variety of internal signals or asymmetric determinants. Available evidence suggests that Pins may be functionally analogous to GPCRs in that it is able to bind Go subunits [Scheafer et al. 2000] and locally liberating activated Gβγ heterodimers. Indeed, expression of the Go-Loco domains of Pins is able to activate the pheromone response pathway in yeast cells lacking the pheromone receptor [Cismowski et al. 1999], suggesting that Pins may activate Gβγ in a receptor-independent fashion. Because the Pins protein is asymmetrically localized by virtue of its ability to bind to the polarity determinant Insuteable [Scheafer et al. 2000; Yu et al. 2000], it may be able to activate heterotrimeric G proteins at specific sites. The Go and Gβγ effectors that mediate the cytoskeletal response are not known, but it is likely that they will activate Cdc42 or other Rho GTTPases. Drosophila Cdc42 has been identified [Sasamura et al. 1997], but a function in asymmetric cell division has not been demonstrated. In contrast, RNAi experiments in C. elegans embryos suggest that Cdc42 is required for asymmetric spindle positioning [Jantsch-Plunger et al. 2000; M. Gotta, pers. comm.]. Activated Cdc42 has recently been shown to interact with Par6, Par3, and PKCζ [Joberty et al. 2000; Lin et al. 2000; Qiu et al. 2000], which are required for asymmetric cell divisions and spindle positioning. No proteins with obvious homology to Far1p have been found; thus, it remains to be determined how GEFs are targeted to activated Goa and Gβγ in these systems.

Activation of Rho-GEFs at the plasma membrane

Although the activation of yeast Cdc24p at the plasma membrane remains elusive, this step is comparatively well understood in several mammalian Rho-GEFs. The available evidence strongly suggests that the catalytic domain of many Rho-GEFs is inhibited intramolecularly by its N-terminal regulatory domain. Mutant forms, which have truncations of their N-terminal domain, are constitutively active and often exhibit oncogenic activity [Cerione and Zheng 1996; Whitehead et al. 1997]. Recently, structural determination confirmed that the Dbl-homology domain of Vav1 is autoinhibited by an N-terminal extension, which lies in the GTTPase-interaction site. Thus, activation of Rho-type GEFs is likely to involve dissociation of an intramolecular inhibitory do-
main from the catalytic Dbl-domain, which may be achieved by binding to a transmembrane receptor or an activator protein such as Rsrlp/Budlp or Farlp [Fig. 2B], to a second messenger, or by posttranslational modification including phosphorylation.

A strong case for the activation of Vav1 by tyrosine phosphorylation has recently emerged. Vav1 is phosphorylated on Tyr 174 by proximal tyrosine kinases on receptor stimulation and membrane recruitment, and this phosphorylation strongly increases its exchange activity [Crespo et al. 1997; Han et al. 1997]. Indeed, NMR spectroscopy data show that the N-terminal inhibitory domain encompassing Tyr 174 is released from the Dbl-homology domain on phosphorylation [Aghazadeh et al. 2000], demonstrating that tyrosine phosphorylation relieves autoinhibition by exposing the GTase interaction surface of the Dbl-homology domain. GEF activity is further promoted by PIP2 and PIP3, which bind to its PH domain [Han et al. 1998]. GEFs of the Vav or Pix family also contain an N-terminal CH domain that, in Cdc24p, appears to interact with its activators Rsrlp/Budlp and Farlp [Fig. 2B]. It is thus possible that binding of a GTPase or other activators to the CH domain of Vav cooperates with tyrosine phosphorylation for full activation, or that binding of activators may be a prerequisite for subsequent tyrosine phosphorylation.

Activation of a Rho-GEF can be triggered by binding of an activator, as illustrated by the following example. Recent evidence shows that the Rac-GEF Asef binds to the tumor suppressor adenomatous polyposis coli (APC; Kawasaki et al. 2000), which links cell adhesion and cell proliferation [Goss and Groden 2000]. The activity of Asef is negatively regulated by its N-terminal region, which contains the APC-binding site. Thus, binding of APC to Asef is predicted to relieve autoinhibition, thereby allowing the catalytic domain of Asef to interact with Rac1.

Finally, the Rap1-GEFs Epac1 and Epac2 are activated by directly binding to the second messenger cAMP (de Rooij et al. 1998, 2000). In the absence of cAMP, an interaction between the GEF domain and the cAMP-binding domain inhibits the catalytic activity of Epac1. Epac1 becomes activated by a release of this inhibition on cAMP binding. Consistent with this model, deletion of the cAMP-dependent regulatory sequences results in constitutive GEF activity (de Rooij et al. 2000).

Taken together, these activators are different in nature, they may use a common mechanism, in that they directly bind to the N-terminal domain, thereby relieving autoinhibition imposed on the catalytic domain of GEFs [Fig. 2B]. Thus, a search for factors that bind to the regulatory domain of GEFs may be fruitful to identify potential Rho-GEF activators.

Stabilization of GEFs at the plasma membrane by adapters

A second class of proteins that interact with GEFs are the adapters. In yeast, the adapter Bem1p binds to the C terminus of Cdc24p, but this interaction is not essential for Cdc24p activation in vivo [Jaquenoud et al. 1998]. Importantly, Bem1p is not required for the recruitment of Cdc24p to the plasma membrane, but it is involved in stabilizing activated Cdc24p at sites of polarization. Many mammalian adapter proteins are involved in GTPase signaling [Buday 1999], but their role in GEF activation or localization is poorly understood. For example, the palmitoylated adapter protein LAT [linker for activation of T cell] binds to Vav and thereby facilitates the targeting of Vav to glycolipid-enriched microdomains in activated T cells [Salojin et al. 2000]. However, time-lapse microscopy is required to distinguish whether LAT is involved in membrane recruitment or stabilization of Vav at the plasma membrane. An intensively studied adapter protein is Nck, which is composed exclusively of one SH2 and three SH3 domains and has been shown to interact with many proline-rich proteins including the GEFs Pix and Trio [McCarty 1998]. Nck associates via its SH2 domain with activated EGF and PDGF receptors and can be recruited into signaling complexes in response to activation of the vertebrate guidance receptors EphB1 and EphB2 [Holland et al. 1997; Stein et al. 1998], two Eph receptor tyrosine kinase family members. Consistent with these findings, the Dro sophila Nck homolog Dreadlocks (DOCK) is capable of binding ligand-activated Eph1 [Stein et al. 1998] and has recently been implicated in growth cone guidance and assembly of focal adhesion complexes by binding to Pix and Trio. Indeed, Dock/Nck expression is detected in R-cell growth cones in the target region [Garrity et al. 1996] and is also found in focal adhesions. Thus, the GEFs Trio and Pix together with the adapter Dock/Nck may regulate GTPase activity during photoreceptor axon pathfinding in Dro sophila and the formation of focal adhesions in mammalian cells. Finally, paxillin has emerged as a focal adhesion adapter involved in the integration of growth factor- and adhesion-mediated signal transduction pathways [Turner 2000]. Paxillin is a multidomain protein that binds and recruits PAK, Nck, and Pix to the plasma membrane, thereby directing GTPase-mediated reorganization of the actin-based cytoskeleton [Turner et al. 1999]. Thus, Dock/Nck and paxillin may function analogously to Bem1p in yeast and stabilize the GEFs Pix and Trio at sites of polarized growth.

Inactivation of GEFs: a role for PAK-like kinases?

In yeast, a negative feedback loop mediated by the PAK-like kinase Cla4p may inactivate Cdc24p at the plasma membrane by inducing its dissociation from the adapter Bem1p [Fig. 4; Gulli et al. 2000]. However, Cla4p has also been shown to promote actin polymerization [Eby et al. 1998; Lechler et al. 2000]. In mammalian cells, constitutive active Pak1 or overexpression of Cdc42-GTP or Rac1-GTP induces rapid loss of stress fibers and a dramatic loss of focal adhesions, resulting in retraction of the cell periphery in migrating fibroblasts [Manser et al. 1997]. These results suggest that activated Pak1 may promote turnover of these structures.

Recent studies on axonal guidance in the Dro sophila nervous system provide a beautiful example for the dual
role of Pak-like kinases in regulating dynamic remodeling of actin structures [Hing et al. 1999]. Whereas the directed extension of growth cones leads axons toward their destination, rearrangements of cytoskeletal structures on reaching the target area are required to inhibit growth cone motility and to allow the formation of synaptic connections. Pak1/Msn has been shown to play a role in the shutdown of growth cone motility when axons reach their target region [Ruan et al. 1999]; Overexpression of Pak/Msn induces pretarget growth cone termination, and conversely, many growth cones fail to stop at their target lamina in Pak/Msn mutants. Thus, target-derived stop signals may activate Pak/Msn, which in turn coordinates cytoskeletal reorganization in decelerating growth cone motility. Like Bem1p in yeast, the adapter Dock/Nck forms a complex with Pak1/Msn in growth cones and at the leading edge of embryonic epithelial cells undergoing dorsal closure, where they are found in structures resembling focal adhesions. The axonal stop signal may thus activate the function of Pak/Msn through Dock/Nck by either positioning the kinase close to its substrate or directly stimulating its activity, leading to termination of the growth cone in the target area. A possible candidate for a regulator of Pak1/Msn is the neuron-specific Rac-effector p35/cdk5, which concentrates at the leading edge of axonal growth cones and regulates neurite outgrowth in cortical neurons in culture. Active p35/cdk5 causes hyperphosphorylation of Pak1, which results in down-regulation of Pak1 kinase activity [Nikolic et al. 1998]. Thus, it is possible that inhibition of p35/cdk5 activates Pak1 at the target lamina, leading to inactivation of growth cone motility. The substrates of Pak/Msn responsible for the stop signal of growth cones or the turnover of focal adhesion complexes are not known, but by analogy with the results in yeast, they may include phosphorylation of GEFs. In this regard, it may be interesting to determine whether Trio or Pix dissociate from tips of growth cones once the neurites reach their target area.

Other than a possible role for Pak-like kinases in the negative regulation of GEFs, it has been reported recently that Vav1 is degraded via the ubiquitin–proteasome system [De Sepulveda et al. 2000]. SOCS1 is an inducible SH2-containing protein that negatively regulates cytokine and growth factor signaling required for thymic development. SOCS1 interacts with elongins B and C, which are both components of a SCF/VCP-ubiquitin ligase complex [Tyers and Jorgensen 2000], suggesting that SOCS1 may function as a substrate-specific recognition component for Vav1. Because phosphorylation of several substrates is required for their binding to the SCF complex [Deshaies 1997], phosphorylation of Vav1 may also play a negative role by initiating its subsequent degradation. Taken together, ubiquitin-dependent degradation of Vav1 may provide a novel mechanism to down-regulate its activity in response to cytokine signaling. It is not known whether a similar mechanism may also apply to Cdc24p, although its levels do not fluctuate during the cell cycle [Gülli et al. 2000].

Conclusions

Studies on Rho-GEFs in yeast and mammalian systems identified many regulators, which we postulate can be grouped in four distinct steps (Fig. 5). Impressive progress has been obtained on the activation step itself: Available results strongly indicate that Rho-GEFs are activated by releasing intracellular inhibition exerted by their N-terminal regulatory domain. Thus, a major future challenge will be to identify the various Rho-GEF activators, which are predicted to bind to their N-terminal domains. In addition, full activation of GEFs may require the cooperation of multiple activators, which may respond to different signal transduction pathways.

An important but poorly understood aspect of Rho-GTPase signaling is their local activation. Time-lapse microscopy of Cdc24p-GFP in living cells during yeast budding and mating combined with mutational analysis provided a first glimpse at the dynamic subcellular localization and the regulated assembly and disassembly of GEF complexes at the plasma membrane. However, many of the conclusions about GTPase signaling in mammalian cells are based on overexpression of either constitutively active or dominant-negative mutant forms of GEFs, Rho-type GTases, or their targets. Because these mutant forms are in most cases uniformly expressed at the cell surface, it is difficult to access local effects. In addition, the existence of feedback loops may further complicate the interpretation of the observed phenotypes. Thus, new tools and experimental approaches are needed to address the dynamic and spatial aspects of GTPase signaling. Recently, a system allowing local recruitment of activated Cdc42p to a cell surface receptor provided interesting insights into local actin polymerization and filapodia formation [Castellano et al. 1999]. In addition, an approach (FLAIR) to visualize activated Rho-GTPases based on specific binding domains and fluorescence-resonance energy transfer (FRET) microscopy has been developed, which may allow us to follow GTPase signaling in real-time in vivo [Kraynov et al. 2000].

Although we have focused this review on the regulation of Rho-GEFs, it is likely that intra- and extracellular signals also regulate the localization and/or activity of GAPs and GDIs. Thus, the coordinated control of all these regulators may be required to spatially and temporally active and inactivate Rho-GTPases in vivo.

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