Locally controlled inhibitory mechanisms are involved in eukaryotic GPCR-mediated chemosensing

Xuehua Xu, Martin Meier-Schellersheim, Jianshe Yan, and Tian Jin

G protein–coupled receptor (GPCR) signaling mediates a balance of excitatory and inhibitory activities that regulate Dictyostelium chemosensing to cAMP. The molecular nature and kinetics of these inhibitors are unknown. We report that transient cAMP stimulations induce PIP₃ responses without a refractory period, suggesting that GPCR-mediated inhibition accumulates and decays slowly. Moreover, exposure to cAMP gradients leads to asymmetric distribution of the inhibitory components. The gradients induce a stable accumulation of the PIP₃ reporter PHₐCRAC-GFP in the front of cells near the cAMP source. Rapid withdrawal of the gradient led to the re-association of G protein subunits, and the return of the PIP₃ phosphatase PTEN and PHₐCRAC-GFP to their pre-stimulus distribution. Reapplication of cAMP stimulation produces a clear PHₐCRAC-GFP translocation to the back but not to the front, indicating that a stronger inhibition is maintained in the front of a polarized cell. Our study demonstrates a novel spatiotemporal feature of currently unknown inhibitory mechanisms acting locally on the PI3K activation pathway.

Introduction

Eukaryotic cells can detect and move up concentration gradients of chemoattractants, a process known as chemotaxis (Zigmond, 1978; Chung et al., 2001a; Iijima et al., 2002; Van Haastert and Devreotes, 2004). This behavior plays an important role in a number of processes, including metastasis, angiogenesis, immune responses, and inflammation (Murphy, 1994; Parent and Devreotes, 1999; Condeelis et al., 2005). Furthermore, chemotaxis is essential for cell aggregation in the life cycle of the social amoeba, Dictyostelium discoideum (Gerisch, 1982; Devreotes and Zigmond, 1988; Devreotes, 1994; Van Haastert and Devreotes, 2004). Chemotaxis is a coordinated phenomenon of three fundamental cell processes: gradient sensing, cell polarization, and cell motility. Chemotactic cells, such as neutrophils and D. discoideum, display polarized morphology, involving asymmetric distributions of many signaling molecules (Parent and Devreotes, 1999; Comer and Parent, 2002; Iijima et al., 2002; Devreotes and Janetopoulos, 2003), and heightened responsiveness to the attractant at their leading edge (Parent et al., 1998; Jin et al., 2000; Xu et al., 2003). These crawling cells extend their leading edges by assembling a force-generating network of actin filaments beneath the plasma membrane (Chung et al., 2001a; Pollard and Borisy, 2003). Elsewhere in the cell, actin collaborates with myosin to retract the rear of the advancing cell and to prevent errant pseudopod extension (Chung et al., 2001a). Consequently, the actin-depolymerizing agent Latrunculin can be used to eliminate polarization and motility of D. discoideum cells, and thus facilitate quantitative spatiotemporal analyses of the mechanisms underlying gradient sensing (Parent et al., 1998; Jin et al., 2000; Xu et al., 2005).

Gradient sensing is mediated by G protein–coupled receptors (GPCRs) and associated signaling components that detect the spatiotemporal changes of chemoattractants and translate shallow gradients of chemoattractants into steep intracellular gradients of signaling components (Parent and Devreotes, 1999; Chung et al., 2001b; Funamoto et al., 2002; Iijima et al., 2002). Binding of cAMP to the GPCR cAR1 induces the dissociation of heterotrimeric G proteins into Ga2 and Gβγ subunits (Jin et al., 2000; Janetopoulos et al., 2001; Xu et al., 2005). Free Gβγ activates Ras, thereby leading to the activation of PI3K, which converts P(4,5)P₂ (PIP₃) to P(3,4,5)P₃ (PIP₅₃) in the plasma membrane (Li et al., 2000; Funamoto et al., 2001; Stephens et al., 2002; Sasaki et al., 2004; Wessels et al., 2004). The phosphatase PTEN acts as an antagonist of PI3K, dephosphorylating...
studied by direct visualization and quantitative analysis of the mechanisms that leads to activation, adaptation, and amplification has been proposed that an increase in receptor occupancy activates two antagonistic signaling processes: a rapid “excitation” called “amplification” (Parent and Devreotes, 1999) and a slower “inhibition” that turns off those responses (Parent et al., 1998; Xu et al., 2005). To explain these features, it has been proposed that an increase in receptor occupancy activates two antagonistic signaling processes: a rapid “excitation” that triggers cell responses, such as the membrane accumulation of PIP_3, and a slower “inhibition” that turns off those responses (Parent and Devreotes, 1999). Although many of the molecular mechanisms of the excitatory process have been identified, those of the inhibitory process have remained elusive.

There are several key features of gradient sensing. First, cells have the ability to spontaneously terminate responses under a sustained cAMP stimulation in a process called “adaptation” (Parent et al., 1998; Xu et al., 2005). Second, if cAMP is removed from adapted cells, the cells will enter a de-adaptation phase—a refractory period lasting several minutes during which the cells progressively regain their ability to respond to another cAMP stimulation (Dinauer et al., 1980a,b). Third, cells have the capability of translating shallow cAMP gradients across the cell diameter into highly polarized intracellular responses, a process called “amplification” (Parent and Devreotes, 1999; Servant et al., 2000; Chung et al., 2001a). To explain these features, it has been proposed that an increase in receptor occupancy activates two antagonistic signaling processes: a rapid “excitation” that triggers cell responses, such as the membrane accumulation of PIP_3, and a slower “inhibition” that turns off those responses (Parent and Devreotes, 1999). Although many of the molecular mechanisms of the excitatory process have been identified, those of the inhibitory process have remained elusive.

The dynamic relationship between excitation and inhibition that leads to activation, adaptation, and amplification has been studied by direct visualization and quantitative analysis of the spatiotemporal changes in receptor occupancy, G protein dissociation, PIP_3 and PTEN distribution, and PIP_3 level along the membrane (Xu et al., 2005; Meier-Schellersheim et al., 2006). Over the years, models have been proposed to explain how the excitatory and the inhibitory processes interact in cells responding to chemotactic signals to achieve adaptation or amplification (Meinhardt, 1999; Parent and Devreotes, 1999; Postma and Van Haastert, 2001; Devreotes and Janetopoulos, 2003; Arriemarou and Meyer, 2005; Charest and Firtel, 2006; Levine et al., 2006; Meier-Schellersheim et al., 2006). Although inhibitors are essential components of all gradient sensing models, the spatial-temporal presence of inhibitors has not been examined experimentally.

In this study, we designed sequential stimulation protocols to detect temporal and spatial aspects of the inhibition process in single living cells. We found that repeated transient activations of cAR1 receptor trigger repetitive PH<sub>cra</sub>-GFP membrane translocations without detectable refractory periods, demonstrating that a short pulse of cAR1 activation elevates excitation but little long-lasting inhibition. This result provides evidence that cAR1 activation induces an immediate excitation and a delayed recruitment of long-term inhibition leading to PIP_3 accumulation. More significantly, we have revealed spatial distribution of the inhibition process induced by a cAMP gradient. Exposing a cell to a sustained cAMP gradient leads to a stable PH<sub>cra</sub>-GFP accumulation in the front of the cell. We found that a sudden withdrawal of the cAMP gradient from this biochemically polarized cell leads to a rapid return of G protein activation, PTEN, and PIP_3 distributions to basal levels around the cell membrane. However, there was a short time period during which reaction of receptors and G proteins around the membrane induced a clear PIP_3 response in the back but not the front of the cell. This inverted PIP_3 response indicates that a cAMP gradient induces a stronger inhibition of PI3K in the front of a cell.

Results

Brief cAMP stimuli activate excitation but not inhibition of PH<sub>cra</sub>-GFP membrane translocation

Previous studies suggest that activation of cAR1 triggers a fast increase of the excitation level and a slower elevation of the inhibition, allowing a cell to respond transiently and then adapt (Iijima et al., 2002; Devreotes and Janetopoulos, 2003; Janetopoulos et al., 2004; Xu et al., 2005). When a sustained cAMP stimulus is removed, cells that had adapted enter a refractory period during which the cells progressively regain their ability to respond to cAMP, suggesting that the inhibition returns to its prestimulus level more slowly than does excitation (Devreotes and Steck, 1979; Dinauer et al., 1980a,b). To test whether there is a temporal difference between the cAR1-induced excitatory and inhibitory processes controlling PIP_3 production, we measured the kinetics of PIP_3 levels around the membrane of cells that were stimulated by multiple transient cAMP stimuli (Fig. 1). We simultaneously visualized and quantitatively measured transiently applied cAMP stimulations and PIP_3 production, reported by the membrane translocation of PH<sub>cra</sub>-GFP, a PIP_3 reporter (Parent et al., 1998). Chemotaxis-competent PH<sub>cra</sub>-GFP-expressing cells (“PH cells”) were treated with Latrunculin B, which eliminates morphological polarity and motility by disrupting the actin cytoskeleton. A micropipette filled with cAMP (1 μM) was placed within 30 μm of PH cells and used in conjunction with a microinjector to deliver a series of brief cAMP stimulations. Alexa 594 was included in the micropipette as a measure of the applied cAMP concentration (Xu et al., 2005). Each cAMP stimulation induced a transient response of PH<sub>cra</sub>-GFP membrane translocation (Fig. 1). Temporal changes in cAMP concentration around the cell were determined as the average intensity change of the dye in the R1 and R2 regions (Fig. 1 B). The kinetics of PH<sub>cra</sub>-GFP membrane translocation were measured as intensity changes in cytosolic PH<sub>cra</sub>-GFP pool (Fig. 1 B), which is inversely related to the amount of membrane-associated PH<sub>cra</sub>-GFP (Xu et al., 2005). Quantitative analyses showed that PH<sub>cra</sub>-GFP translocation reached its maximal level in 4 s after the cAMP concentration reached its peak, which reflects the temporal delay that is expected for PIP_3 production upon the activation of cAR1 (Fig. 1 C). In our experimental setup, the shortest interval between two transient cAMP stimuli was ~24 s, a minimal time required for the cAMP concentration to return to its basal level between stimuli (Fig. 1 D). Sequential transient cAMP stimuli with as short as 24-s intervals generated repetitive transient responses of PH<sub>cra</sub>-GFP.
translocation, and transient responses displayed kinetics without a refractory period (Fig. 1 D). Our data indicate that a transient receptor activation quickly activates excitatory pathways leading to an increase in PIP3 and upon cAMP removal, these pathways quickly return to prestimulated levels and can be activated again by another cAMP stimulation. Transient cAR1 activations do not signal long enough to substantially elevate the slower inhibition process from its basal level. Therefore, this result supports the idea that cAR1-mediated excitation and inhibition processes increase and decrease by following a fast and a slow temporal mode, respectively.

Exposure to cAMP gradient diminishes responsiveness at the front of the cell

The spatial distribution of inhibition in a cell exposed to a cAMP gradient has never been tested. Several models propose that cAR1-induced inhibitors diffuse quickly in a cell, and thus are assumed to be evenly distributed in the inner membrane (Parent and Devreotes, 1999; Iijima et al., 2002; Janetopoulos et al., 2004). However, our dynamic analyses and computer simulations suggest that inhibition mechanisms act locally and predict that inhibition affecting PI3K activity is strongest in the front of a cell at steady state in a cAMP gradient (Xu et al., 2005; Meier-Schellersheim et al., 2006). Because the molecular nature of the inhibitors in chemosensing of D. discoideum is still unknown, we developed an approach to indirectly measure the relative extent of inhibition in the front and back of a biochemically polarized cell. We reasoned that after a removal of the cAMP gradient, the signaling network would rapidly return from the polarized to the resting steady state. During this transition, the time required to regain responsiveness to cAMP (the refractory period) may differ between the front and back depending on local
concentrations of inhibitors induced by the prior gradient. In our experiments (Fig. 2), PH cells were first exposed to a cAMP gradient until they achieved a stably polarized state, in which PH_Crac-GFP accumulated in the front. After a withdrawal of the cAMP gradient at 0 s, a uniform cAMP stimulation (100 nM of cAMP mixed with Alexa 594) induced PH_Crac-GFP translocation to the back (arrows at 35 s) of the cells. An animated version is in the supplemental materials [Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1]. Another example is shown as Fig. S1 and Video 2. (B, D, and F). Regions of interest used for assessing concentration changes of cAMP with time in the front (DF1-3) and back (DB1-3), and for quantitative measurement of PH_Crac-GFP dynamic changes for the selected front (PHF 1–3) and back (PHB 1–3) membrane regions for each of the three cells shown in A. (C, E, and G) Top panels show temporal changes of Alexa 594 fluorescence for cAMP concentration in the time course in the front (DF) and back (DB) regions. Bottom panels show temporal changes in relative levels of PH_Crac-GFP in the front (PHF) and back (PHB) for each of the three cells.

After the removal of a cAMP stimulus, G proteins quickly reassociate and can be reactivated

The observed inverted PIP_3 response is likely caused by a slow return of the intracellular components to their “resting” states because the average time period for cAMP binding to the receptor is in the range of seconds (Ueda et al., 2001). Several components in the pathways may be responsible for the differential refractory behaviors of the front and back of the cells. For example, receptors may remain asymmetrically desensitized or G proteins may not be completely reassociated upon the second cAMP stimulation. We addressed this issue by directly measuring the kinetics of G protein reassociation and reactivation in living cells using FRET analysis (Fig. 3). Cells expressing Go2-CFP and YFP-Gβ (“G cells”) were suddenly exposed to 10 μM cAMP (Fig. 3, A and B), a saturating dose for cAR1, or 1 μM cAMP (Fig. S2, available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1). Addition of cAMP induced a rapid FRET loss, which reached a steady state in <20 s, indicating G protein dissociation (Fig. 3 B; Fig. S2) (Xu et al., 2005). After the removal of cAMP, FRET returned to the prestimulus level in ~60 s, indicating that the G proteins were completely reassociated. A second sudden exposure to the same concentration of cAMP triggered an instant FRET loss that displayed kinetics very similar to those in response to the first stimulation (Fig. 3 B; Fig. S2), demonstrating that cAR1 receptor and G proteins rapidly returned to their prestimulation states and could be fully reactivated within 60 s after the removal of cAMP.

We then simultaneously monitored temporal changes in the receptor/G protein activation and PH_Crac-GFP translocation in the front and back of the cells previously exposed to a cAMP gradient (Fig. 3, C–F). We measured FRET changes in one G cell
and PHCrac-GFP translocation in another PH cell. The cells were located within 20 μm and thus both were exposed to very similar cAMP stimuli (Fig. 3 C). After a rapid withdrawal of a cAMP gradient at time 0, the G protein reassociated around the G cell membrane and PHCrac-GFP returned to the cytosol (Fig. 3 D). A uniformly applied stimulation at 134.7 s triggered a similar degree of G protein dissociation, which was measured as a CFP fluorescence increase (FRET loss), in both the front and back of the G cell (Fig. 3 E), while a distinctive accumulation of PHCrac-GFP only occurred in the back of the PH cell (Fig. 3, D and E). Because the subsequent cAMP stimulation induced G protein activation in both the front and back, we concluded that the possible asymmetrical desensitization of cAR1 receptors, such as cAR1 phosphorylation induced by the first cAMP gradient, could not explain the subsequent PHCrac-GFP translocation only to the back of the cell (Fig. 3 G). Thus, the inverted PIP3 response is likely caused by inhibitory mechanisms acting on the signaling components downstream of cAR1 and the heterotrimeric G proteins.

![Figure 3](http://www.jcb.org/cgi/content/full/jcb.200611096/DC1). Regions of interest for simultaneous measurement of G protein activation in one G cell and the inverted PIP3 response in a PH cell are shown. (D) Selected images show the inverted PHCrac-GFP translocation in the experiment shown in C. The arrows point to direction of membrane translocation: the original at 0 s and the inverted at 171 s. (E) The temporal changes in the G protein dissociation in the front (black) and back (gray) of the G cell, reflected as CFP intensity changes, in response to the uniform cAMP field. The basal level of G protein activation is 1, shown as the thin line. The thick gray bar indicates the temporal changes in cAMP concentration. (F) Temporal changes in PHCrac-GFP in the front and back of the PH cell in response to the uniform cAMP stimulation applied at 134 s. The basal levels of PHCrac-GFP in the front and back are 1, indicated as the thin dashed line. The thick gray bar indicates the temporal changes in cAMP concentration. (G) After a withdrawal of a gradient, a new uniformly applied cAMP field induced similar levels of G protein dissociation in the front and back of G cells, but triggered a clear PHCrac-GFP translocation only to the back of regions of PH cells. Normalized maximal FRET changes and PHCrac-GFP increases in the front and back of G and PH cells are shown as means ± SD (n = 5). Basal levels are 1, indicated as dashed line.

Spatiotemporal dynamics of PHCrac-GFP membrane translocation in PTEN null cells in response to cAMP stimuli

cAR1 activates an excitatory signaling branch that induces PTEN to translocate from the membrane to the cytosol and also elevates an inhibitory mechanism that allows cytosolic PTEN to return to the membrane (Funamoto et al., 2002; Iijima and Devreotes, 2002). To determine spatiotemporal changes in PIP3 in the cells lacking PTEN, we measured kinetics of PHCrac-GFP membrane translocation in pten− cells and compared the kinetics to those in wild-type (WT) cells (Xu et al., 2005; Meier-Schellersheim et al. 2006), in response to uniformly applied cAMP and a cAMP gradient (Fig. 4). WT and pten− cells expressing PHCrac-GFP were stimulated uniformly with cAMP (1 μM) at 0 s (Fig. 4 A). The cAMP-triggered PHCrac-GFP membrane translocation is fast and transient in WT cells. In contrast, the response in pten− cells was clearly slower, peaking in ~12 s and returning to prestimulus levels in more than 40 s (Fig. 4 A).
When the cells were suddenly exposed to a gradient (Fig. 4 B), membrane translocations of PH Crac-GFP occurred initially in both front and back regions in both WT and pten− cells. However, the kinetics of PH Crac-GFP translocation in the front was clearly abnormal in pten− cells. There was no clear decrease in PH Crac-GFP amount at the front for more than 150 s, which differs from the biphasic response in WT cells (Fig. 4 B). We also examined kinetics of the PIP3 in response to the removal of cAMP stimuli in pten− cells (Fig. 4 C). After cells were exposed to a gradient, PH Crac-GFP accumulates in the front regions of WT or pten− cells. Upon a removal of the gradient at 0 s, PH Crac-GFP gradually returned to the cytosol. The returning process was clearly slower in pten− cells than in the WT cells, whose t1/2 were ~22 s and 14 s, respectively (Fig. 4 C).

**PTEN quickly redistributes to the membrane after cAMP removal**

PTEN is involved in regulating spatiotemporal dynamics of PIP3 levels around the membrane of a cell in response to cAMP stimulation. Therefore, the dynamic distribution of PTEN affects the local PIP3 levels. When a cell reaches the “polarized” steady state in a stable cAMP gradient, PTEN is enriched at the back side of a cell (Funamoto et al., 2002; Iijima and Devreotes, 2002; Li et al., 2005). After withdrawal of the cAMP gradient, PTEN starts to redistribute itself from the polarized to the resting steady state. During this transition, a transient accumulation of PTEN in the front could potentially explain the inverted PIP3 response. To address this possibility, we measured the spatiotemporal dynamics of PTEN under these conditions (Fig. 5). After a rapid withdrawal of the gradient at time 0 (Fig. 5, A and B), PTEN redistributed from the back and became uniformly distributed around the membrane in ~80 s without over-accumulating in the front (Fig. 5). Furthermore, reapplying a uniform stimulus (Fig. 5, A–D) or gradient (Fig. 5, E–H) of cAMP induces PTEN translocation with kinetics (Fig. 5) similar to those observed in the cells that had not previously been stimulated (Meier-Schellersheim et al., 2006), indicating that the cAR1-controlled regulatory components of PTEN returned to their “resting” states and PTEN molecules in both the front and back were responsive to a second cAMP stimulation when the inverted PH Crac-GFP response occurred. Therefore, the excitatory and the inhibitory mechanisms that control PTEN membrane distribution are not the likely explanation for this inverted response.

**Inhibitory pathways controlled by Gα1 or Gα9 subunits are not essential for cAR1-mediated PH Crac-GFP responses**

Other mechanisms may also be involved in inhibition in the signaling network of cAMP gradient sensing. For example, Go9 and Gα1-mediated PLC pathways in *D. discoideum* have been shown to function as negative regulators in the cAR1-mediated signaling (Bominaar and Van Haastert, 1993; Brzostowski et al., 2004). To test whether either pathway is essential for the gradient sensing, we examined PH Crac-GFP responses in Go9 and Gα1 null cells (Fig. 6). We measured PH Crac-GFP membrane translocation by monitoring intensity changes of GFP fluorescence...
in the cell membrane (Xu et al., 2005). In response to a uniform stimulation, the spatiotemporal kinetics of PH<sub>Crac</sub>-GFP membrane translocation in either ga<sup>9−</sup> or ga<sup>1−</sup> cells were similar to those in WT cells (Fig. 6, A and C). When the ga<sup>9−</sup> or ga<sup>1−</sup> cells were suddenly exposed to stable cAMP gradients, PH<sub>Crac</sub>-GFP translocation, as in WT cells, consisted of two phases, an initial transient translocation around the cell membrane followed by a second phase producing a highly polarized distribution (Fig. 6, B and D). Because our observed dynamics in both mutant cells are similar to those displayed in WT cells (Xu et al., 2005; Meier-Schellersheim et al. 2006), we suggest that Ga<sub>1</sub> or Ga<sub>9</sub> controlled signaling are not essential inhibitory mechanisms for cAR1-mediated gradient sensing.

**Figure 5.** Spatiotemporal dynamics of PTEN in single cells upon a withdrawal of a cAMP gradient and then stimulated with a uniform dose (A–D) or a gradient (E–H) of cAMP. (A) A PTEN-GFP expressing cell was highly polarized in a stable cAMP gradient (red, 1 μM in the micropipette, red) at 0 s. The gradient was suddenly withdrawn from the cell at 0 s. A uniform cAMP stimulation (100 nM) was applied at 144 s. Regions of interest for the data reported in B and C are also shown. (B) Temporal changes in cAMP concentrations in the front (black) and back (gray) are shown in the time course. (C) Temporal changes in relative levels of membrane-bound PTEN in the front (black) and back (gray) regions are shown in the time course. (D) Kinetics of membrane-bound PTEN in the front and back are shown as means ± SD (n = 10) in response to a withdrawal and then to a uniformly applied stimulation. (E) A PTEN-GFP expressing cell was highly polarized in a stable cAMP gradient (red, 1 μM in the micropipette) at 0 s. The gradient was suddenly withdrawn from the cell start at 0 s, and reapplied at the 90 s. Regions of interest for the data reported in B and C are also shown. Animated version is in the supplemental materials (Video 4, available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1). (F) Temporal changes in cAMP concentrations in the front (black) and back (gray) are shown in the time course. (G) Temporal changes in relative levels of membrane-bound PTEN in the front (black) and back (gray) regions are shown in the time course. (H) Kinetics of membrane-bound PTEN in the front and back are shown as means ± SD (n = 13) in response to a withdrawal and reapplied of the cAMP gradient.

Local inhibition mechanism of PI3K revealed by a cAMP-gradient-induced inverted PH<sub>Crac</sub>-GFP translocation

We speculated that the inverted PH<sub>Crac</sub>-GFP translocation may be induced by a reapplied cAMP gradient. Fig. 7 shows this experiment. A WT cell expressing PH<sub>Crac</sub>-GFP was first equilibrated in a cAMP gradient to achieve the polarized state. After a withdrawal of the cAMP gradient at time 0 s, PH<sub>Crac</sub>-GFP returned to cytosol. At 81 s, the identical gradient was reapplied (Fig. 7, A and B). In response to this second gradient, the cells exhibited a much stronger transient accumulation of PH<sub>Crac</sub>-GFP in the back than in the front (Fig. 7, A and B). Fig. 7 C shows the comparison of normalized increase in cAMP concentration delivered by the second gradient in the front and back regions of the cells; and Fig. 7 D shows the maximal increase in PH<sub>Crac</sub>-GFP in the front and back regions during the inverted responses (n = 8). Our results show that, despite the cAMP stimulus being higher in the front than in the back, the cells responded only in the back regions. Thus, after a withdrawal of the gradient, the cells displayed an asymmetrical refractory period. During this period, PIP<sub>3</sub> initially accumulated in an intracellular gradient that had the opposite direction of the external cAMP gradient. The asymmetrical refractory period could be detected in some cells for more than four minutes after the withdrawal of the previous gradient (Fig. S3, available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1).

It was previously reported that cAMP receptor-mediated PI3K activation consists of two layers in chemotaxing cells. First, free Gβγ activates Ras that stimulates a small amount of preexisting, membrane-associated PI3K. The resulting actin
polymerization leads to recruitment of additional PI3K from cytosol to the membrane, thereby increasing the amount of active PI3K (Sasaki et al., 2004). In Latrunculin-treated cells, PI3Ks were uniformly distributed around the membrane of the cells even when they were exposed to the cAMP gradient (Sasaki et al., 2004; unpublished data). Therefore, under our experimental condition, we monitored the spatiotemporal regulations of PI3K activity without complications from the second layer of actin-dependent PI3K recruitment. In addition to the signaling pathway leading to PI3K activation, the cAMP receptor also regulates another pathway mediating the redistribution of membrane-bound PTEN, which is important for the proper directional response of PIP3. In \textit{pten} cells, PHCrac-GFP was still able to accumulate in the front when the cells were exposed to a cAMP gradient (Janetopoulos et al., 2004; Sasaki et al., 2004). We found that the crescents of PHCrac-GFP in \textit{pten} cells were broader than those formed in WT cells (Fig. 7, E and F; at time 0), as previously described (Janetopoulos et al., 2004; Sasaki et al., 2004). Furthermore, the directions of the crescents, unlike those in WT cells, did not always perfectly point to the direction of the gradient (Fig. 7 F; at time 0). These results indicate, as expected, that a cAMP gradient-induced PTEN redistribution ensures the PIP3 response in the restricted front region, and this directional response was not precise without PTEN. However, after a withdrawal of the cAMP gradient, PHCrac-GFP returned to cytosol. More importantly, the second cAMP gradient induced the inverted PHCrac-GFP membrane translocation in \textit{pten} cells (Fig. 7, E and F), indicating that a cAMP gradient-induced asymmetrical inhibition occurred in the absence of PTEN. Collectively, our results suggest that the previous gradient induced an asymmetrically distributed and locally controlled inhibition and this localized inhibition acts on the signaling pathway between free G\(\beta\gamma\) to PI3K.

**Kinetics of the asymmetrical inhibition induced by a cAMP gradient**

We examined the temporal appearance and disappearance of the gradient-induced asymmetrical inhibition (Fig. 8; Fig. S4, B and C, available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1). We found that a brief gradient stimulation of \(\sim 50\) s was not sufficient to induce an inverted PHCrac-GFP response (Fig. 8, A and B; Fig. S4 B). Thus, exposure to a stable gradient for \(\sim 2\) min is needed to establish an asymmetrical inhibition. Furthermore, cells that were removed from a gradient for 6 min and rechallenged with either uniform cAMP stimulation or a cAMP gradient displayed a noninverted PHCrac-GFP translocation response as in naive cells (Fig. 8, C and D; Fig. S4 C), indicating that asymmetrical inhibition disappears within 6 min after the gradient is removed.

**Discussion**

The existence of inhibitory components in GPCR-mediated chemosensing has been proposed for more than thirty years, but the molecular mechanisms are still unknown and, thus, cannot yet be visualized directly. Here, we report insights into the temporal and spatial aspects of the inhibition based on measurements of the spatiotemporal dynamics of known components of the gradient sensing machinery aided by our computational modeling study.
We have constructed a quantitative model for cAR1-mediated signaling network (Meier-Schellersheim et al., 2006). The model, which includes receptor-mediated and locally controlled inhibitory mechanisms that regulate PI3K and PTEN (Fig. 9 B), simulates experimentally determined dynamics of receptor activation, G protein dissociation, PTEN membrane localization, and PIP3 accumulation (Meier-Schellersheim et al., 2006). For example, in response to a uniform cAMP stimulus, the model generates a transient PIP3 response that quickly returns to the resting stage (adaptation). When exposed to a cAMP gradient, a cell generates a steeper PIP3 gradient by initially inducing a PIP3 increase followed by a PIP3 decrease around the membrane, and then producing a highly polarized distribution of PIP3 in 120 s (amplification) (Fig. S5, available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1). During the amplification process, the membrane-bound PTEN gradually translocates from the front to the back, while the amount of the membrane-associated PI3K remains the same around the membrane (Fig. 9 C). Temporal changes in PI3K activity in the front and back, which cannot be directly visualized, have been simulated by the model based on dynamics of PIP3 and membrane-bound PTEN (Meier-Schellersheim et al., 2006; Fig. S5). Previous study and our measurements indicated that when a cell reaches the “polarized” steady state, the amount of PI3K in the front is almost equal to that in the back (Sasaki et al., 2004). PIP3 is at a higher steady-state level in the front. However, this level does not continue to increase (ΔPIP3/Δt = 0) in spite of a lower level of PTEN. At the same time, in the back, a higher level of PTEN does not result in a continuous decrease in the PIP3 level (ΔPIP3/Δt = 0). Two possible models may explain different steady states of PIP3 in a polarized cell. First, PI3K activity is stronger than that of PTEN in the front, and PIP3s are continually produced. The PIP3 level remains steady in the front because it diffuses fast enough to be degraded by PTEN that is enriched in the back, which is expected from models containing only globe inhibition mechanisms (Parent and Devreotes, 1999; Iglesias and Levchenko, 2002; Iijima et al., 2002). Second, balances between the activities of PI3K and PTEN have been reached in both the front and back, and the balances are achieved by a stronger inhibition of PI3K activity in the front, which...
have been proposed in our model that includes local inhibition mechanisms (Xu et al., 2005; Meier-Schellersheim et al., 2006). Because different proposed mechanisms could lead to high chemotactic sensitivity in theory (Meinhardt, 1999; Postma and Van Haastert, 2001; Devreotes and Janetopoulos, 2003; Arrieumerlou and Meyer, 2005; Levine et al., 2006; Meier-Schellersheim et al., 2006), we designed experiments to determine which inhibitory mechanisms are likely used in GPCR-mediated chemosensing. In this study, we revealed spatiotemporal features of an inhibitory process that acts locally on the activation pathway between Gβγ and PI3K.

There is a general agreement that inhibition increases and decrease slowly in response to the changes of cAMP receptor occupancy (Parent and Devreotes, 1999; Postma and Van Haastert, 2001; Devreotes and Janetopoulos, 2003; Arrieumerlou and Meyer, 2005; Levine et al., 2006; Meier-Schellersheim et al., 2006), we designed experiments to determine which inhibitory mechanisms are likely used in GPCR-mediated chemosensing. In this study, we revealed spatiotemporal features of an inhibitory process that acts locally on the activation pathway between Gβγ and PI3K.

Figure 8. Kinetics of the asymmetrically distributed inhibition. (A and B) Asymmetrically distributed inhibition is not induced by exposure to an initial cAMP gradient for just 50 s. PH cells were suddenly exposed to a cAMP gradient (1 μM) at 0 s for 50 s. After a withdrawal of the gradient for ~70 s, a uniform cAMP stimulation (100 nM) induced PHcα-GFP translocation in both the original front and the back regions [A]; and a reapplied cAMP gradient triggered a clear PHcα-GFP translocation in the original front (B). (C and D) The asymmetrical inhibition disappears 6 min after the removal of the cAMP gradient. PH cells had been exposed to a cAMP gradient (1 μM) for more than two min, and PHcα-GFP became stably polarized in the front of the cells. The cAMP gradient was removed from the cells at 0 s. After a removal of the gradient for 6 min, a uniform cAMP stimulation (100 nM) induced a PHcα-GFP translocation in both the original front and the back regions (C), and a reapplied cAMP gradient triggered PHcα-GFP translocations to both the original front and back regions.
period of time during which another cAMP stimulation triggers G protein dissociation and PTEN translocation in both the front and back but induces PIP3 responses only in the back of the cell (Fig. 9). This suggests that inhibitors that are more abundant in the front block transmission of activating signals from Gβγ (Fig. 9). This suggests that inhibitors that are more abundant in the front block transmission of activating signals from Gβγ (Fig. 9). This suggests that inhibitors that are more abundant in the front block transmission of activating signals from Gβγ (Fig. 9). This suggests that inhibitors that are more abundant in the front block transmission of activating signals from Gβγ (Fig. 9).

Figure 9. Dynamic signaling events in a cell exposed to a gradient of cAMP leading to PIP3 polarization, followed by a withdrawal of the gradient and then reapplying the gradient inducing an inverted PIP3 response. (A) A cell is shown schematically at several time points, indicating the distributions of extracellular cAMP (red) and intracellular PHcrac-GFP (green) in the front (F) and back (B) regions. (B) A scheme of GPCR-mediated signaling network containing inhibitory mechanisms. Dynamics of signaling molecules that are filled with colors were measured in living cells. Activation of GPCR: cAMP (red), G protein dissociation, free Gβγ (gray), membrane-bound PTEN (blue), membrane-bound PI3K (light blue), PIP3 levels (green), and inhibitory components (black box). (C) Graphs represent a time course of relative signaling levels in the front and back of a cell. A naive cell at resting stage is suddenly exposed to a stable cAMP gradient at time 0. The cell reaches a polarized steady-state at 120 s. At the time T, the cAMP gradient is quickly removed, the cell starts to return to its resting stage. Before the cell completely returns to the resting stage, the cAMP gradient is re-applied, and the cell generates an inverted PIP3 response at time “In”. During this time course, the relative levels of extracellular cAMP (red), the extent of G protein dissociation (gray), membrane-associated PHcrac-GFP (green) as a measure of PIP3 levels, membrane-associated PTEN (blue), the amount of membrane-associated PI3K (light blue), the P13K activity has not been directly measured. Temporal changes in P13K activity was simulated, shown in Fig. S5), and the proposed inhibition (black dash lines) in the front and back of the cell are shown.

This inhibition is stronger in the front of the cell. The spatiotemporal features of the inhibition can shed light on unknown molecular mechanisms. Based on the fast-diffusive-inhibition models, small molecules, such as Ca2+ or cGMP, were suggested to be candidate inhibitors, which have not been verified by experiments. The “local excitation and global inhibition” model assumes the presence of a negative regulator, and suggests that it is likely to be PTEN. Based on our detailed spatiotemporal dynamics of PTEN and PIP3, our computational model showed that PTEN alone cannot fully explain the experimentally determined dynamics (Meier-Schellersheim et al., 2006). We proposed, in addition to PTEN, other inhibitory mechanisms that may involve reversible modifications of components in the pathway from free Gβγ to Ras and then to PI3K. Previous studies in mammalian GPCR signaling indicated several inhibitory components. After GPCR activation, free Gβγ dimers interact with the receptor-associated kinase GRK2, blocking Gβγ signaling (Lodowski et al., 2003). GPCR activation can also induce a translocation of a RasGAP, which binds to PIP3, to inner membrane deactivating Ras thereby inhibiting PI3K (Lockyer et al., 1999). In D. discoideum, it has been shown that a sustained cAR1 activation, which triggers a persistent G protein dissociation, induces a transient activation of RasG, which activates PI3K (Sasaki et al., 2004). The transient nature of RasG activation is consistent with the idea that the cAR1 activation also recruits inhibitors to the membrane to shut down signals from free Gβγ to Ras activation. Our computational model is able to simulate the observed spatiotemporal dynamics of known components in adaptation and in gradient sensing by including these putative inhibitor(s) (Meier-Schellersheim et al., 2006). Therefore, we propose that the inhibition process is performed by these negative regulators acting locally on the PI3K.
signaling branch and those on PTEN branch, which act in concert to control the spatiotemporal dynamic of PIP₃ around the cell membrane. Future studies are needed to identify inhibitors involved in the GPCR-mediated chemosensing network.

Materials and methods

Cell lines and live cell imaging

As previously described (Ku et al., 2005), D. discoideum cell lines expressing PHCRac-GFP (Xu et al., 2005), PTEN-GFP (Iijima and Devreotes, 2002), PI3K1-GFP (Sasaki et al., 2004), and both G2005); and pten -α-PI3K1-GFP cells expressing PHCRac-GFP were developed to the chemotactic stage. Cells were plated on a 1-well chamber for the microinjector delivered cAMP stimulation (Nalge Nunc International), allowed to adhere to the cover glass for 10 min, and then covered with additional DB buffer. Live cells were imaged using a Zeiss Laser Scanning Microscope, LSM 510 META, with a 40× NA 1.3 DIC Plan-Neofluar objective. To monitor cAMP and PHCRac-GFP, PI3K1-GFP cells were excited with two laser lines, 488 nm for GFP and 543 nm for Alexa 594, a water-soluble fluorescence dye. Images were simultaneously recorded in three channels. Channel one: fluorescence emissions from 505–530 nm for GFP (green); channel two: emissions from 580–650 nm for Alexa 594 (red).

Generation and measurement of applied cAMP stimulations

The temporal-spatial intensity changes of Alexa 594 and cells expressing PHCRac-GFP, PTEN-GFP, or PI3K1-GFP were directly imaged using a confocal microscope with 2-axis resolution of ~2 μm. Fluorescence intensities of Alexa 594 and GFP within the focal plane were simultaneously recorded in two different channels. To establish a steady gradient, we set an external supply pressure to 70 hPa (Femtotjet and micromanipulator S171; Eppendorf) to ensure the injection of a constant and small volume of cAMP and Alexa 594 into a one well chamber. Under this condition, a stable gradient was established within 100 μm around the tip of the micropipette. To suddenly expose a cell to a stable gradient, a micropipette filled with a mixture of cAMP and 0.1 g/μl Alexa 594 linked to a Femtotjet was positioned 1,000 μm away from the cells, and then was quickly moved to a position within 100 μm to the cells. During the experiments, we only changed the movement of the micropipette and the cells. The speed of the movement determines how fast a stable gradient can form around a cell (Xu et al., 2005). To withdraw a gradient, the micropipette was quickly moved away from a cell.

FRET measurement

Using a spectral confocal fluorescence microscope (LSM510 META), we measured intensity decrease of acceptor (YFP) and increase of donor (CFP) in response to stimuli. We monitored intensity changes of CFP (donor) and (YFP) acceptor following a stimulation using a time-lapse acquisition of Lambda Stacks. The cells were excited with a 454-nm laser line and the spectral emissions in each pixel of the fluorescence images were simultaneously recorded in 8 channels, each with a 10-nm width, from 464 to 544 nm. To separate multi-fluorescence signals, each of the fluorescence images was assembled as montages using Photoshop 7.0. Quantification of fluorescence intensities of Alexa 594, GFP, CFP, and YFP in the regions of interest was performed using the LSM 510 META software.

Online supplemental material

Fig. S1 shows inverted PIP responses. Fig. S2 shows FRET measurement of G protein dissociation and association and redissociation. Fig. S3 shows reapplication of a cAMP gradient induced an inverted PHCRac-GFP membrane translocation. Fig. S4 A shows Gα0 null cells detect cAMP gradient normally.

Fig. S4 B and C show kinetics of the formation the asymmetrically distributed inhibition. Fig. S5 shows PI3K activity, membrane-bound PTEN and the resulting dynamics of PIP₃ in a cell when it is exposed to a cAMP gradient in a computer simulation and a schematic representation of the signaling network that describes spatiotemporal changes. Videos 1 and 2 show uniformly applied cAMP stimulation triggered inverted PHCRac-GFP translocation. Video 3 shows simultaneously measurement of G protein activation in the front and back of a cell and the inverted PHCRac-GFP response. Video 4 shows dynamics of PTEN in a cell upon a withdrawal of a cAMP gradient and then reapplication the gradient. Videos 5 and 6 show a cAMP gradient induced the inverted PHCRac-GFP membrane translocation. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200611096/DC1.

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