Introduction

Chemotaxis, or directed movement, of a eukaryotic cell toward the source of chemoattractant (hereafter shorted as CA), i.e. cAMP for Dictyostelium or fMLP for neutrophils, play an essential role in many pathophysiological processes (Parent 2004, Bagorda and Parent 2008). During embryogenesis, axons are steered by nerve growth factors to form the nervous system (Ruthel and Banker 2015). In immune responses, neutrophils are directed to the correct locations, and they kill the debris (Oliveira et al 2016). Chemotaxis is also central to wound healing and is implicated in tumor metastasis (Zimmermann 2017). For efficient movement, cells must sense the difference in CA concentration between its two ends, initiate the appropriate intracellular signaling cascade to mediate cytoskeletal remodeling and finally, form the well-defined front and rear regions (Devreotes and Janetopoulos 2003, Dalous et al 2008). At the front, the assembly of cytoskeletal actin propels the protrusion (Cai and Devreotes 2011). At the opposite side, actomyosin contracts and pulls up the rear (Parent 2004). Clearly, cells should be able to reverse their polarity for efficient targeting to the new source of CA (Gerisch and Keller 1981). That is, when a polarized cell is exposed to a sufficiently high CA gradient from the opposite direction, its initial front region could be reorganized transiently as the rear region and vice versa. Thus, understanding the dynamics of eukaryotic chemotaxis remains crucial.

Dynamic seesaw model for rapid signaling responses in eukaryotic chemotaxis

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Abstract

Directed movement of eukaryotic cells toward spatiotemporally varied chemotactic stimuli enables rapid intracellular signaling responses. While macroscopic cellular manifestation is shaped by balancing external stimuli strength with finite internal delays, the organizing principles of the underlying molecular mechanisms remain to be clarified. Here, we developed a novel modeling framework based on a simple seesaw mechanism to elucidate how cells repeatedly reverse polarity. As a key feature of the modeling, the bottom module of bidirectional molecular transport is successively controlled by three upstream modules of signal reception, initial signal processing, and Rho GTPase regulation. Our simulations indicated that an isotropic cell is polarized in response to a graded input signal. By applying a reversal gradient to a chemoattractant signal, lamellipod-specific molecules (i.e. PIP3 and PI3K) disappear, first from the cell front, and then they redistribute at the opposite side, whereas functional molecules at the rear of the cell (i.e. PIP2 and PTEN) act oppositely. In particular, the model cell exhibits a seesaw-like spatiotemporal pattern for the establishment of front and rear and interconversion, consistent with those related experimental observations. Increasing the switching frequency of the chemotactic gradient causes the cell to stay in a trapped state, further supporting the proposed dynamics of eukaryotic chemotaxis with the underlying cytoskeletal remodeling.

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chemotaxis is integral to elucidate biochemically induced cell movement.

In the past two decades, much progress, particularly with Dictyostelium and neutrophils, has been made by identifying additional components of the chemotactic pathways related to cytoskeletal remodeling (Ridley et al 2003, Parent 2004). The dynamic features of eukaryotic chemotaxis are a consequence of redistribution of intracellular signaling molecules; that is, the regions rich in Rac, Cdc42, PIP3 are associated with actin branching and growth, while those rich in RhoA, PIP2, and PTEN are correlated with myosin II-induced contraction (Gardiner et al 2002, Kolisch et al 2008, Pramanik et al 2009). As the time scale of intracellular signaling responses is likely to be much shorter than those of gene expression and protein synthesis, spatiotemporal regulation of signaling pathways have become more interesting, especially when testing how cells effectively respond to spatiotemporally varied biochemical stimuli (Gardiner et al 2002).

Experimentally, state-of-the-art techniques, such as microfabrication and in vivo imaging, have greatly facilitated quantitative analyses of intracellular signaling responses. For example, a microfluidic device can generate the precisely controlled gradient of extracellular CA (Ebrahimzadeh et al 2000, Meier et al 2011). Single-molecule imaging using fluorescent proteins enables highly sensitive quantification of cellular responses and cytoskeletal remodeling by simultaneously monitoring the spatiotemporal distribution of typical intracellular molecules (Ueda et al 2001, Sako et al 2012). Despite the different eukaryotic cell types used in those experiments (Weiner 2002, Dalous et al 2008), the dynamics of cellular chemotaxis are similar and can be summarized as follows. (1) Adaptive: the cells can sense a small difference in CA concentration between the front and rear regions over a wide range of mean concentrations (Parent 1999, Chen et al 2003). (2) Rapid: the cells generate the initial polarization in approximately 1–2 min (Weiner 2002). (3) All-or-none: the cells usually have one major pseudopod at a time (Dalous et al 2008). (4) Tunable: the cells continually adjust their polarized direction in response to new stimuli (Gerisch and Keller 1981, Dalous et al 2008). These four features are a prerequisite for a cell to undergo dynamic chemotaxis and should be included in a universal framework with the known signaling pathways.

Various mathematical models have been proposed to explore different mechanisms involved in intracellular signaling responses of chemotactic cells (Devreotes and Janetopoulos 2003, Iglesias and Devreotes 2008, Vanderlei et al 2011). The earliest turning-type model argues that the stable pattern of eukaryotic chemotaxis may arise if there is autocatalytic local production of a slow diffusive activator that also causes the production of a fast diffusive inhibitor (Meinhardt 1999). This model suggests symmetry breaks at the cellular scale with multiple activated regions; hence, the activation is not restricted to the front region. In addition, the model cell cannot change the direction of its polarity in response to a new stimulus, such that the entire system exhibits a ‘lock-on’ state. Several subsequent models have been developed based on a local excitation-global inhibition (LEGI) principle (Parent 1999, Ma et al 2004). In a basic LEGI model (Parent 1999), receptor occupancy triggers a fast, local excitatory signal and a slow, global inhibitory signal. At the cell front, the concentration of the activator is greater than the inhibitor and vice versa at the rear. Likewise, a balanced-inactivation model was also developed by considering two second messengers produced at equal rates (Levine et al 2006). The diffusion of one messenger, coupled with an inactivation scheme, ensures a switch-like response to external gradients over a wide range of mean concentrations. Lacking in these two models, however, are mechanisms of the role of actin-dependent pathways in regulating eukaryotic chemotaxis. To address these issues, an additional model was established based on a LEGI-biased excitable network hypothesis, where the LEGI scheme serves as an input to the downstream biochemical network that controls cytoskeletal activity (Xiong et al 2010). Unfortunately, these models do not account for stable polarity due to the difficulties in assigning polarity-specific components. Additionally, an antagonizing pathway model proposes that external stimuli activate two divergent pathways that mutually inhibit one another via interaction with the actin cytoskeleton (Onsum and Rao 2005), which can correctly localize and stabilize specific proteins to the respective front and rear regions of the cell. It also predicts that the cell slowly rotates its original pseudopods in response to a secondary reversal signal and simultaneously forms two pseudopods in response to a pair of stimuli, which does not agree with the experimental observations. Our previous work on the bidirectional molecular transport mechanism (Feng and Zhu 2014), in which the Rho GTPase-PIs feedback loops (PIs denote to PIP3 and PIP2 hereafter) are employed and the cell is treated as an intracellular and intra-membrane signal feedback system, has ignored the F-actin-independent pathways in regulating eukaryotic chemotaxis. Thus, several models can be used to unravel how signaling molecules form a polarized distribution to initiate the dynamic chemotaxis of a cell and then respond to a reversal CA stimulus.
Here we present a multi-modular modeling framework for eukaryotic chemotaxis. Numerical simulations were designed to interpret a group of distinct but interrelated experimental observations, including initial polarization, reversal polarization, cell trapping, and cell mutation. Our modeling indicated that intracellular signaling responses with chemotactic pathways act as a bidirectional molecular transportation system controlled by a spatial gradient of active Rho GTPases that are transduced from the external signaling field. Physically, this prediction supports a surprisingly simple seesaw-like mechanism between the cell front and rear.

**Biological backgrounds**

Our modeling began by understanding that the primary components of chemotactic pathways among different kinds of eukaryotic cells are highly conserved, and signaling cascades are dominated by distinct mechanisms at different signaling layers (modules). As mentioned in our earlier work (Feng and Zhu 2014), four modules were applied, including signal reception (I), initial signal processing (II), small Rho GTPase regulation (III), and bidirectional molecular transport (IV). We propose a brief modeling strategy (figure 1 (a)) and describe detailed descriptions of the molecular mechanisms involved in each module.

**Signal reception**

Cells sense the CA using transmembrane receptors (i.e. G protein-coupled receptors (GPCR)). The GPCRs are uniformly distributed along the plasma membrane (Parent 1999, Ridley et al 2003), which serves as an important regulator for cells to accurately sense the change in extracellular gradients.

**Initial signal processing**

The cascades between heterotrimeric G proteins (α, β, and γ) and F-actin independent signaling provide the initial amplification of sensed extracellular signals and adaptation mechanisms (Parent 2004, Stephens et al 2008). Upon CA binding to GPCRs, GTP can be exchanged for GDP on Gα, which triggers a fast activation response and slower inhibition response. (III) Rho GTPase regulation module: spatial regulation of Rho GTPase members (Rac, Cdc42, and RhoA) is achieved by the localization of GEFs and the antagonistic effects between them. (IV) Bidirectional molecular transport module: spatial effects generated from Rho GTPase-PIs feedback loops control the bidirectional cytoskeletal remodeling. Curved arrows in the insert illustrate the roles of upstreaming modules for controlling the positive direction of molecular transport. For initial polarization, the required directional cue is relative weak (depicted by fine arrows); for reversal polarization, the required directional cue from the opposite direction is strong (depicted by bold arrows).

**Small Rho GTPase regulation**

Rho-family small GTPases serve as central hubs in transducing signals from extracellular CA to the actin cytoskeleton (Raftopoulou and Hall 2004). The Rho GTPases are switch-like proteins that are cycled between active membrane-bound (GTP) forms and inactive cytosolic (GDP) forms (Raftopoulou and Hall 2004, Jilkine et al 2007). Activation of Rho GTPase is mediated by guanine exchange factors (GEFs) and inactivated by GTPase activating proteins (GAPs), while translocation of Rho GTPase from the membrane to cytosol is regulated by GDP dissociation inhibitors.
(GDIs) (Raftopoulou and Hall 2004, Holmes et al 2012). Besides the self-regulation of each Rho-family member, there is considerable cross-talk between different members of the Rho-family (Jilkine et al 2007). Cdc42 is considered to serve as the upstream signal of Rac, while RhoA has mutual inhibitory interactions with Cdc42 and Rac (Gambardella and Vermeren 2013). The CA-induced spatial regulation of Rho GTPase could be initiated by Gαi and Gβγ. For example, Gαi activates specific Rho GEFs via PDZ-GEF/Lsc1 (Fenteany and Glogauer 2004), and Gβγ binds to p21-activated kinase 1 (PAK1) and then activates Cdc42 via PAK-associated GEF (PIXα) (Gambardella and Vermeren 2013).

### Bidirectional molecular transport
Once the gradient of active Rho-family GTPase is established, the spatial distributions generated by Rho GTPase and PIs tend to mediate positive/negative feedback loops (Weiner et al 2002, Charest and Firtel 2006), which drive bidirectional molecular transport. Intracellular transport of signaling molecules may orchestrate the localization of actin regulators and myosin activators, both of which are directly related to force generation mechanisms (Charest and Firtel 2006, Rottner and Stradal 2011).

**Prominent front region with highly active Cdc42/Rac**
A role for small Rho GTPases in a self-organizing feedback loop of actin-PI3K-PIP3 was proposed based on observations that latrunculin treatment and pharmacological inhibition of PI3K or Rho GTPases significantly blocks PIP3 accumulation (Charest and Firtel 2006). Concretely, active Cdc42/Rac molecules interact with various actin-binding proteins (ABPs),

| Parameter | Description | Value | Sources |
|-----------|-------------|-------|---------|
| K         | Diminution rate of CA | 1 s⁻¹ | Postma et al (2001) |
| GR        | Total number of GPCR | 80,000 | Ueda et al (2001) and Rappel and Levine (2008) |
| k⁻        | Dissociation rate | 1 s⁻¹ | Janssens and Van Haastert (1987) and Rappel and Levine (2008) |
| k⁺        | Association rate | 30 nM⁻¹ s⁻¹ | Janssens and Van Haastert (1987) and Rappel and Levine (2008) |
| δα, δτ    | Inactivation rates of activator, inhibitor | 0.2, 0.2 s⁻¹ | Levine et al (2006) |
| k_{AI}    | Inactivation rate between activator and inhibitor | 100 µm (s · molecule)⁻¹ | Levine et al (2006) |
| k_{S}     | Production rate of activator and inhibitor | 1, 1 s⁻¹ | Levine et al (2006) |
| k_{I}     | Association rate of inhibitor | 3 nM⁻¹ s⁻¹ | Levine et al (2006) |
| R_{tot}, C_{tot}, ρ_{tot} | Total levels of Rac, Cdc42, RhoA | 7.5, 3, 3 µM | Michaelson et al (2001) and Jilkine et al (2007) |
| R_{th}, C_{th}, ρ_{th} | Typical basal levels of active Rac, Cdc42, RhoA | 3, 2.4, 1.25 µM | Maree et al (2006) and Maree et al (2012) |
| δα, δC, δρ | Basal decay rates of active Rac, Cdc42, RhoA | 1, 1, 1 s⁻¹ | Zhang and Zheng (1998), Jilkine et al (2007) and Maree et al (2012) |
| I_{b}, I_{c}, I_{p} | Baseline activation rates of Rac, Cdc42, RhoA | 0.4, 0.8, 0.4 s⁻¹ | Maree et al (2006) |
| β, τ | Efficent GEF-dependent Cdc42, RhoA activation rates | 3, 3 s⁻¹ | Sakumura et al (2005) |
| α | Cdc42-dependent Rac activation rate | 1 µm (s · molecule)⁻¹ | Otsuji et al (2007) |
| γ | RhoA-dependent Rac inactivation rate | 1 µm (s · molecule)⁻¹ | Otsuji et al (2007) |
| P_{th}, P_{th} | Basal levels of PIP2, PIP3 | 10, 0.45 µM | Ma et al (2004) and Maree et al (2012) |
| K_{PI3K}^{PI3K}, K_{PTEN}^{PTEN} | PIP3 and PIP2 level for half-max PI3K, PTEN feedback | 8, 8 µM | Estimated |
| K_{PI3K}^{PI3K}, K_{PTEN}^{PTEN} | Enzymatic rates per active PI3K, PTEN | 4, 4 s⁻¹ | Estimated |
| R_{max}, ρ_{max} | Activity constants of Rac, RhoA | 4, 1 µM | Estimated |
such as SCAR/WASP and Arp2/3 complex, which initiates F-actin polymerization (Parent 2004). During this process, PI3K is recruited from the cytosol to the membrane and catalyzes the phosphorylation of PIP2 to produce PIP3 (Stephens et al. 2002, Van Haastert and Devreotes 2004). PIP3 molecules then act as binding sites for various ABPs, which in turn promote F-actin polymerization and lead to the further membrane recruitment of PI3K and the repetition of the cycle (Sasaki et al. 2007). The barbed ends of actin filaments proliferate by Arp2/3-mediated branching, extending until they reach the membrane and then exerting protrusion forces against the membrane (Rottner and Stradal 2011).

**Rear region with highly active RhoA**

Active RhoA molecules interact with phosphatidylinositol phosphatase (PTEN) (Li et al. 2005), which is recruited from the cytosol to the membrane by PIP2 (Sulis 2003). In its active form, PTEN antagonizes the action of PI3K by converting PIP3 to PIP2 (Weiner et al. 2002, Sulis 2003, Billadeau 2008). Myosin-II is colocalized with PTEN, and its activity is regulated by myosin light-chain (MLC) phosphorylation, which is positively regulated by MLC kinase (MLCK) or Rho- associated kinase (ROCK) (Van Haastert and Devreotes 2004, Besser and Schwarz 2007). Consequently, actomyosin contracts and pulls up the rear.

**Theoretical modeling**

Consider that a cell is placed at a shallow CA filed, and only the contour membrane of the cell is assumed to sense the biochemical stimuli (figure 1(b)). By focusing on the spatiotemporal regulations of intracellular signal cascades, the structural elements inside the cell body are ignored, and the lamellipodium is treated as a cylindrical disk, with the cytosol as its interior and the cell membrane/cortex as its perimeter. There are different diffusive coefficients for molecular diffusion at extracellular regimes, the cytoplasm, and membranes, given as $D_l = 50$, $D_c = 10$, and $D_m = 1 \, \mu m^2 \cdot s^{-1}$, respectively (table 1). Basic equations, derived from the above biochemical scheme (figure 1(a)), consist of a set of coupled partial differential equations (PDEs) that describe the reaction kinetics, cross-talk, diffusivity, and communication between the intermediate products. The translocation behaviors of those signaling molecules between the membrane and the cytosol are modeled in distinct ways. Proteins such as PI3K, PTEN, PAK1, and Lsc1, with limited amounts, are treated by a stochastic kinetics approach. In view of fast diffusion and a sufficient amount, the inactive cytosolic or GDP forms of Rho GTPase are assumed to be uniformly distributed in the cytosol and then calculated by the conservation law.

**Basic equations**

In this subsection, the equations and parameters are summarized for our modeling. Implementation decisions and simplifying assumptions are also discussed.

**Signal reception**

Diffusion and diminution of CA molecules were modeled by the reaction-diffusion equation, with corresponding initial and boundary conditions,

$$\frac{\partial S}{\partial t} = D_S \nabla^2 S - KS, \quad (1a)$$

$$S(x, y, t) = S_0, \quad \text{for } x \in [x_1, x_2], \ y \in [y_1, y_2],$$

$$S(x, y, 0) = 0, \quad \text{for } x \notin [x_1, x_2], \ y \notin [x_1, y_2] \quad (1b)$$

$$S(X, Y, t) = 0,$$

where $S, D_S$, and $K$ are the concentration, diffusive coefficient, and diminution rate of CA molecules, respectively. $x \in [x_1, x_2]$ and $y \in [y_1, y_2]$ define the domain of the point source, where the concentration of CA equals $S_0$. $X$ and $Y$ represent the boundary of the CA field. The binding kinetics of the CA molecules (as ligands) to the GPCRs (as receptors) are described by the following reaction equation,

$$R + L_{\alpha} = RL,$$

where $R, L, R_{\alpha}, k_+, and k_-$ represent receptor, ligand, complex, association rate, and dissociation rate, respectively.

**Initial signal processing**

The input CA signals need to be further translated into an intracellular asymmetry of signaling molecules. Here, we used a balance-inactivation mechanism to mimic this process (Levine et al. 2006). The original mechanism proposed previously involves three interacting steps. First, the local level of receptor occupancy ([RL]) drives the production of a membrane-bound species $A$ and a cytosolic species $I$ at equal rates, $k_0$. Second, the cytosolic species diffuses inside the cell and attaches to the membrane at a rate $k_i$, where the membrane-anchored species $I$ is denoted as $I_m$ for our modeling. Finally, both species $A$ and $I_m$ inactivate each other with a rate $k_i$, and also $A$ and $I_m$ spontaneously degrade at the rates $\delta_A$ and $\delta_I$ respectively, which was used to mimic their biologic antagonism. The evolution dynamics of $A, I$ and $I_m$ are described as follows:

$$\frac{\partial A}{\partial t} = D_m \nabla^2 A + k_0 [RL] - \delta_A A - k_i A_m, \quad (3a)$$

$$\frac{\partial I_m}{\partial t} = D_m \nabla^2 I_m + k_i I - \delta_I I_m - k_i A_m, \quad (3b)$$

$$\frac{\partial I}{\partial t} = D_c \nabla^2 I, \quad (3c)$$

with a boundary condition,

$$D_c \frac{\partial I}{\partial r} = k_0 [RL] - k_i I. \quad (3d)$$

Upon the original balance-inactivation model, the abstract component activator $A$ and inhibitor $I$ can be identified as $G_{\alpha \beta}$ and $G_{\alpha \alpha}$, respectively, even though
the mechanistic details responsible for their cross-talk are not explicitly explored. \( A \) and \( I_m \) may initiate the downstream Rho GTPase regulation module by interacting and/or incorporating with corresponding GEF of Rho GTPase via the intermediate messengers. In this model, we simply assumed that PAK1 and Lsc1 bind to the membrane at a rate proportional to the local density of \( A \) and \( I_m \), respectively. The translocating behaviors of PAK1 and Lsc1 from the cytosol to the membrane are described by the binding reaction equation (equation \( \text{(2)} \)), where PAK1 and Lsc1 act as ligands, while \( A \) and \( I_m \) act as receptors. Because the numbers of PAK1 and Lsc1 (similar to PI3K and PTEN) are limited and their impacts on system behavior are significant, we identified them as effectors. Spatiotemporal evolution of cytosolic effectors obeys the following standard diffusion equation,

\[
\frac{\partial E_c}{\partial t} = D_c \nabla^2 E_c, \tag{4a}
\]

with a boundary condition for the outward pointing normal derivative of the cytosolic effectors,

\[
D_c \frac{\partial E_c}{\partial n} = -k_{c} S_m E_c + k_m M_c, \tag{4b}
\]

where \( E_c = E_{\text{PAK1}}, E_{\text{Lsc1}}, E_{\text{PI3K}}, \) and \( E_{\text{PTEN}} \) represent the cytosolic concentrations of PI3X3, Lsc1, PI3K, and PTEN, respectively, \( M_c = M_{\text{PAK1}}, M_{\text{Lsc1}}, M_{\text{PI3K}}, \) and \( M_{\text{PTEN}} \) define the membrane-bound concentrations of PAK1, Lsc1, PI3K, and PTEN, respectively, and \( S_m \) denotes the corresponding receptor concentration. Converting the concentration (in \( \mu M \)) into the site density of molecules (in \( \mu m^{-2} \)) is achieved by applying a factor, \( \eta = 100 \text{mM}^{-1} \text{mm}^{-2} \) (Gerisch and Keller 1981).

**Small Rho GTPase regulation**

We considered small Rho GTPases (Rac, Cdc42, RhoA) dynamics using a simplified model described previously, where mass conservation of various signaling components and diffusion-driven instability generates their polarized distribution (Otsuji et al 2007, Maree et al 2012). We assumed that each member of Rho GTPases was cycled between active membrane-bound forms and inactive cytosolic forms. The interactions among each member of Rho GTPases formed double negative feedback loops, as presented in the schematic of the Rho GTPases interactions in figure 1(a). The spatiotemporal evolution of active Rho GTPases is described as follows,

\[
\frac{\partial G}{\partial t} = D_m \nabla^2 G + P_G \left( \frac{G}{G_{\text{tot}}} \right) - \omega G, \tag{5a}
\]

where \( G = R, C, \) and \( \rho \) represent the active (membrane-bound) forms of Rac, Cdc42, and RhoA, respectively. \( G_{\text{tot}} = G_{\text{tot}, R} + G_{\text{tot}, C} + G_{\text{tot}, \rho} \) are the total concentrations of Rac, Cdc42, and RhoA, respectively. \( G_i = G_{\text{tot}, i} \) and \( \rho_i \) are the total concentrations of the respective inactive (cytosol) forms of Rac, Cdc42, and RhoA, calculated by the mass conservation law. \( P_G \) is the activation term and expressed as:

\[
P_R = I_R + \alpha_C, \quad P_C = I_C + \beta C, \quad \text{and} \quad P_\rho = I_\rho + \tau \rho. \tag{5b}
\]

Here \( I_R, I_C, \) and \( I_\rho \) are the baseline activation rates. \( \alpha \) sets the rate of Cdc42-enhanced Rac activation, \( \beta \) is the rate of GEF-mediated activation of Cdc42, and \( \tau \) is the rate of GEF-mediated activation of RhoA. \( \omega_G \) is the inactivation term and expressed as:

\[
\omega_G = \delta_R + \gamma_R \rho, \quad \omega_C = \delta_C + \varepsilon \rho, \quad \text{and} \quad \omega_\rho = \delta_\rho + \varepsilon C. \tag{5c}
\]

Here \( \delta_R, \delta_C, \) and \( \delta_\rho \) are the GAP-mediated baseline inactivation rates. \( \gamma \) sets the rate of RhoA-mediated Rac inactivation, and \( \varepsilon \) is the mutual inactivation rate of Cdc42 and RhoA.

**Bidirectional molecular transport**

PIP3 and PIP2 are membrane lipids that play well-known regulatory roles in the actin-myosin cytoskeleton (Stephens et al 2008). PIP3 provides binding sites for various ABPs to promote F-actin assembly and, thus, serves as the marker of the front of a polarized cell (Kolsch et al 2008). PIP2 offers binding sites for myosin-II and, thus, serves as the marker of cell rear (Cai and Devreotes 2011). The spatiotemporal regulation of PIP3 \( (P_3) \) and PIP2 \( (P_2) \) forms the core of the bidirectional molecular transport mechanism, which is described by the following equations.

\[
\frac{\partial P_3}{\partial t} = D_m \nabla^2 P_3 + k_{\text{PI3K}} \left( \frac{M_{\text{PI3K}} P_3 \tilde{R}}{K_{\text{M}^{\text{PI3K}}} + P_3} \right) - k_{\text{PTEN}} \left( \frac{M_{\text{PTEN}} P_3 \tilde{\rho}}{K_{\text{M}^{\text{PTEN}}} + P_3} \right), \tag{6a}
\]

\[
\frac{\partial P_2}{\partial t} = D_m \nabla^2 P_2 + k_{\text{PI3K}} \left( \frac{M_{\text{PI3K}} P_2 \tilde{R}}{K_{\text{M}^{\text{PI3K}}} + P_2} \right) - k_{\text{PTEN}} \left( \frac{M_{\text{PTEN}} P_2 \tilde{\rho}}{K_{\text{M}^{\text{PTEN}}} + P_2} \right). \tag{6b}
\]

Here

\[
\tilde{R} = \min \left( \frac{R}{R_{\text{max}}}, 1 \right), \quad \tilde{\rho} = \min \left( \frac{\rho}{\rho_{\text{max}}}, 1 \right).
\]

In equation \( \text{(6a)} \), the first term on the right-hand side accounts for PIP3 diffusion, the second represents PIP3 production due to membrane-bound PI3K \( (M_{\text{PI3K}}) \) acting on PIP3, and the third accounts for PIP3 diminution due to membrane-bound PTEN \( (M_{\text{PTEN}}) \) acting on PIP3. The value of \( k_{\text{PI3K}} \left( k_{\text{PTEN}} \right) \) is the maximum enzymatic rate per active PI3K (PTEN) molecule. \( K_{\text{M}^{\text{PI3K}}} \) is the typical level of PIP3 \( (P_3) \) that gives half-maximum feedback via membrane-bound PI3K (PTEN). \( R(\tilde{\rho}) \) is a normalized factor that defines the effect of Rac (Rho) activity on the activation of PI3K (PTEN). \( R_{\text{max}}(\rho_{\text{max}}) \) acts as a constant for Rac (RhoA) activity. If \( R(\tilde{\rho}) \) is greater than \( R_{\text{max}}(\rho_{\text{max}}) \), the activity of Rac (RhoA) is no
longer a limiting factor for PI3K (PTEN) activation, and then \( \tilde{R}(\rho) \) equals unity.

Equation (6b) describes PIP2 dynamics. Similarly, the first term on the right accounts for PIP2 diffusion, the second represents the production of PIP2 from PIP3 via membrane-bound PTEN, and the third accounts for the reduction of PIP2 into PIP3 via membrane-bound PI3K. The total amounts of PIP3 and PIP2 remain constant over time. PI3K and PTEN act as key regulatory proteins of PIP3/PIP2 conversation, and both could translocate between the membrane and the cytosol by interacting with PIP2 and PIP3, respectively. While the active form feeds back onto its own production through cooperative binding, such a membrane-cytosol exchange cycle introduces the strongest feedback loop into the system.

**Numerical methods**

Numerical simulations were conducted with our previous work (Feng and Zhu 2014). Briefly, equations (1)–(6) were solved using the Lattice–Boltzmann method (LBM) within a static cell perimeter, in which a simple Monte–Carlo method was embedded to treat those stochastic source terms. The LBM approach is an auxiliary construct that possesses well-known computational advantages by using the fixed Cartesian meshes and straightforward implementation of boundary numerical schemes, allowing a smooth interpolation between the interior and exterior of the cell. The cell is placed in a regular grid containing 100 × 250 grid points and a grid spacing of 0.2 \( \mu \)m. Cell diameter is chosen to be 10 \( \mu \)m. A time step of 0.0025 s was used to solve the diffusion-reaction equation and implement the Monte–Carlo method. Initially, the concentrations of all signaling molecules were uniform in the interior of a circular domain, representing an unstimulated resting cell. All diffusible molecules satisfied the no-flux boundary conditions at the cell edge.

**Results**

**Basic features of cell polarization**

To assess whether the model represents basic experimental observations, we ran the full model (equations (1)–(6)) with biologically based parameter sets (table 1). The spatial positions of various molecules on the membrane were presented using a single variable \( \theta \), an angle varied between 0 and 2\( \pi \). To eliminate the effect of the initial condition, the model cell underwent self-evolution in the absence of external stimuli to achieve a resting steady state (data not shown). To replicate point source-induced chemotaxis, the model cell was first simulated with a graded field formed by a CA point source 10 \( \mu \)m away from the cell for 200 s, followed by instantaneously switching the same point source to the opposite side at the location 5 \( \mu \)m away from the cell at \( t = 200 \) s for an additional 300 s thereafter. Except for the graded external stimulus, no front or rear regions of the cell were defined prior to the simulations. The spatiotemporal distributions of the integral molecule are shown in figure 2.

**Initial graded stimulus period**

After the initial introduction of the CA point source, the diffusion of CA molecules creates a concentration gradient around the cell perimeter (figure 2(a)). This steady-state CA field is established relatively fast, in less than 5 s, with an average concentration of approximately 1 nM and a gradient of 14.3 \( \times 10^{-2} \) nM \( \mu \)m\(^{-1}\). As a result of the high and fast dissociation rate of this binding event, the extracellular gradient of CA ligands is spatially mirrored by the graded occupancy of GPCRs (figure 2(b), blue line). The asymmetry in GPCR occupancy is spatially amplified by the fast signal processing process. The nascent, uniform front and rear regions are now marked by the accumulated \( G_{\beta\gamma} \) and \( G_{\alpha} \), respectively (figure S1(a) and (b) (stacks.iop.org/PhysBio/15/056004/mmedia)), with no significant functional difference. The accumulation of \( G_{\beta\gamma} \) and \( G_{\alpha} \) introduces a local perturbation in Rho GTPase regulation by downregulating PAK1 and Lsc1 (figure S1(c) and (d)), respectively. Together with the mutual inhibition effects between Rhoa and Cdc42, the stimuli-biased distributions of Cdc42 and Rhoa are established (figures 2(c) and (d)). The reciprocal localization of Cdc42 and RhoA, along with their complementary regulation effects on Rac, ensures that Rac becomes positively regulated at the front and negatively regulated at the rear. Thus, the Rac activity displays a steeper gradient (figure 2(e) and (e')). Spatial separation of Rho GTPase members then act as a compass to direct cytoskeletal remodeling. In the regions of high Rac/Cdc42 activity, elevated Rac enhances PI3K activation and, in turn, promotes a local increase in PIP2 concentration. Local recruitment of PI3K from the cytosol to the membrane through PIP3 induces a local reduction of PIP3 which increases the dissociation of membrane-bound PTEN. Therefore, a short range positive feedback loop (Rac \( \rightarrow \) PI3K \( \Rightarrow \) PIP3) is formed, enabling PI3K and PIP2 to be localized at the front (figures 2(f) and (h)). In regions of high RhoA activity, the previously dissociated PTEN from the front rebinds with PIP2. After being activated by RhoA, these PTEN molecules cause further diminution of PIP2, which in turn forms a long-range negative feedback loop (Rhoa \( \rightarrow \) PTEN \( \Rightarrow \) PIP2) that enables PTEN and PIP2 to be localized at the rear (figures 2(g) and (i)). Such diffusion-driven bidirectional molecular transport is guided by the established internal Rho GTPase gradient and achieves a steady state when the pool of cytosol PI3K is significantly depleted. As a result, the final distribution of PIP2 achieves an all-or-none pattern, ensuring that only a single lamellipodia is maintained.

**Reversal graded stimulus period**

Following the introduction of reversal stimuli at \( t = 200 \) s, the redistributed patterns of different
Figure 2. Basic features of polarization produced by WT model cell. (a) Spatial distribution of CA outside the cell. A micropipette containing constant CA concentration (considered as a point source) was initially placed at the middle right side of the cell. The diameter of the cell was 10 µm. (b) GPCR occupancy profiles at an initial stimulating time point (t = 100 s, blue line) and a reversal stimulating time point (t = 300 s, red line). (c)–(g) Simulations of spatiotemporal evolution of active Rho GTPases, that is, Cdc42 (c), Rac (d), and RhoA (e), and PIs, that is, PIP3 (f) and PIP2 (g), along the periphery of the model cell. The ‘0’ angle is chosen to the time point toward the micropipette during the initial stimulus period. The spatial profiles of Rho GTPase (c′)–(e′) display shallow gradients (blue lines), in response to an initial stimulus, or steep gradients (red lines), in response to a stronger reversal gradient. PIP3 and PIP2 act as the markers of cell front and rear, respectively. Both display an all-or-none distribution pattern at the end of initial and reversal stimulus period. (h) and (i) Spatial distribution of PI3K (h) and PTEN (i) at t = 100 s, when the cell has already reached a well-polarized state. Initially, both PI3K and PTEN, as effectors, are set to be uniform in the cytosol with concentrations of 0.07 and 0.11 µM, respectively. Cytosolic distributions of the effectors generate fluctuations near the membrane region due to stochastic translocation behaviors. Local concentrations of membrane PI3K and PTEN are related to the local concentrations of cytosolic PIP3 and PIP2, respectively. (j) Concentration of PIP3 as a function of time at the front (blue line) and the rear of the cell (red line). There is a clear delay period generated at the beginning of the reversal stimulus period (200–240 s), whereby the cell returns to an almost non-polarity state rather than generating two pseudopods at the same time.
regulatory proteins induced the cell to correctly repolarize according to the new stimuli (figures 2(c)–(g), t = 500 s). Specifically, the change in CA gradient was immediately sensed by the cell through the biased occupancy of GPCRs (figure 2(b), red line). Due to the introduction of balance-inactivation mechanism (equation (3)), the switch-like redistribution of $G_{βγ}$ and $G_{α}$, constantly ensured the precise spatial control of PAK1 and Lsc1, such that the internal asymmetry in Rho GTPase was re-established along the new direction (figures 2(c)–(e)). The response was rapid: it reached a new steady state within 10 s. The spatial bipolar redistribution pattern of Rho GTPase was more significant. That is, active Rac and Cdc42 responses exhibited symmetric peaks facing the new point source (figures 2(c') and (d')), whereas active RhoA accumulated away from the source (figure 2(e')). Such a redistribution of Rho GTPase greatly altered the local activation rate of PI3K and PTEN and, further, the direction of molecular transport. After approximately 120 s, PIs distribution achieved an all-or-none pattern, according to the new stimuli direction, as expected. There was also a significant delay in PIP3 response to the reversal stimulus (figure 2(j), t = 200–240 s). In other words, the cell was predicted to return to an almost non-polarity state, but it was not predicted to spontaneously generate two major pseudopod. The specific molecular labels (i.e. PI3K, PIP3) possibly disappeared first from the initial front and then reappeared at the opposite side because the inversion of cell polarity was initiated by a global inhibitor that turned down actin polymerization in the entire cell, followed by a fast-acting positive feedback loop raised to establish a new front (Dalous et al 2008). This global inhibitor was not necessarily required based on the current modeling. Rather, the spatiotemporal effect generated from Rho GTPase-Pis mediated feedback loops could well explain this phenomenon. Despite the fast redistribution of active Rho GTPase (figures 2(c)–(e)), the accumulation of PIP3 at the initial front may still exist for a while (figures 2(f) and (j)), resulting in competition for cytosolic PI3K at two ends. Since the cytosolic PI3K could not effectively arrive at the new front side, the spatial effect of Rho GTPase-Pis mediated feedback loops was limited and no PIP3 accumulation occurred there. After abolishing PIP3 accumulation at the initial front, the competition for cytosolic PI3K between the two ends was won by the new front side. Henceforth, the spatial difference in Rho GTPase activity was continually amplified, which finally led to a full PIP3 response at the new front side.

Effect of CA gradient steepness on cellular polarization

We tested the role of CA gradient steepness in eukaryotic chemotaxis. First, we quantified the effect of gradient steepness on the initial polarization by adjusting the distance between the point source and the cell. All other parameters remained the same. As shown in figure S2, four positions of the source point $(a, b, c$, and $d$, from far to near) were chosen, on which the formed steepest gradients were 2.8, 4.7, 7.9, and $14.3 × 10^{-2}$ nM $\mu$m$^{-1}$, respectively. Time courses of PIP3 accumulation at the original cell front were compared, corresponding to each stimulus. As the applied stimulus gradient become steeper, the time required for the cell to achieve steady polarization became shorter and varied from 180 to 80 s. Meanwhile, the peak value of PIP3 also increased from 6 to 9 µM, suggesting that the cell developed larger pseudopod in response to a steeper CA gradient. In addition, due to the balanced-inactivation mechanism introduced in the initial signal processing module, the modeled cell can respond to a 2% difference in CA concentration between the front and the rear of the cell (data not shown). Without a doubt, the required shallow CA gradient could be derived only if the cell was a distance of $3\lambda \sqrt{D_f K}$ from the source.

By further adding a specific uniform CA field, the percentage variation of CA concentration across the cell could be simulated.

Next, we tested eukaryotic chemotaxis with different gradients of reversal stimulus. Similarly, three positions of CA point sources (c)–(g) were chosen (figure S2). From near to far, the formed reversal stimulus gradients were 43.7, 23.7 and $14.3 × 10^{-2}$ nM $\mu$m$^{-1}$, respectively. To reduce computational costs, those simulations were stated from the same planar polarized state. For each reversal stimulus condition, five successive snapshots of PIP3 redistribution are presented in figure 3((b)). In general, the steepness of the reversal gradient determined the dynamics of PIP3 redistribution and further defined the cell reorientation. Particularly, in the case of the steepest reversal gradient (point e), PIP3 first disappeared from the original front and then reappeared at the opposite side (figure 3((b), top row), suggesting that the cell reoriented itself in response to the new stimulus by generating an entirely new front. At mild steepness (point f), PIP3 redistributed with slow rotation (figure 3((b), middle), implying that the cell preferred to make a U-turn to align to the new gradient. However, in the case of shallow steepness (point g), the cell was locked on the original distribution pattern of PIP3 to a great extent (figure 3((b), bottom row), so that the cell could not reorient itself to the new stimulus.

Collectively, these simulations further illustrated the cooperativity of Rho GTPase-Pis mediated feedback loops. The appearance of PIP3 accumulation resulted from the local strength of Rho GTPase-Pis mediated feedback loops, which indeed depended on both local Rac activity and PIP3 concentration. In the initial cell polarization, Rac activity and PIP3 concentration reached maximal values at the original front. Naturally, a steeper initial CA gradient induced a steeper intracellular Rho GTPase gradient, and accordingly, it evoked the spatial effect of Rho GTPase-
PI3K feedback loops in a rapid manner. In reversal polarization, the maximal value of PIP3 concentration appeared at the original front, whereas the maximum local Rac activity occurred at the opposite side due to the fast dynamics. This mismatch in the distributions of active Rac and PIP3 made the location of maximum feedback strength appear at the rear, near the original front, or at the original front, corresponding to the steepest, mildest, and shallowest stimulus, respectively.

Simulations with time-varied CA signals
To further validate this model, we also used a microfluidic function generator with double T-junction channels to alter the gradient direction of CA in a precise and time-dependent manner (Meier et al 2011). The most interesting finding was that a very fast switch frequency in the gradient direction led to a resting state for trapping the cells, compared to those at a stepwise reduction of gradient switching periods (from 200 s to 15 s). Our simulations indicated that, at a long period of $t = 200 \text{s}$, the PIP3 response followed the varied gradients of the stimulus, indicating that the cell fully adjusts the polarity according to the respective gradient direction (figure 4(a)). At $t = 60 \text{s}$, the induced PIP3 patches were less pronounced (figure 4(b)). Notably, the PIP3 accumulation completely disappeared at the very rapid switching period of $t = 15 \text{s}$ (figure 4(c)), suggesting that cytoskeletal remodeling ceased and the cell was chemotactically trapped in rapid switching. As an ideal case, a cell exposed to spatially uniform CA field was also simulated. Here, spontaneously appearing patches of PIP3 could still be achieved (figure 4(d)), even though more time was required, compared with the gradient-induced case (figure 2(f)).

The underlying mechanism of this high frequency-dependent cell trap observation can also be well interpreted using the rational of our modeling. Basically, cell trapping is shaped by the balance between the efficient transport of intracellular molecules and the switch frequency of the external stimuli. If the switch frequency matches the rate of intracellular biochemistry reorganization, the strength of the Rho GTPase-PI3-mediated positive feedback loop becomes sufficiently strong to drive the bidirectional molecular transport. Here, the intracellular molecules display a successful shuttle run feature that is guided by temporally varied stimulus. However, if the switch frequency is much faster than the rate of intracellular molecular transport, the effect of external stimuli is limited to interrupt the spontaneous establishment of intracellular molecule patches, and no net molecular accumulation is achieved.

Chemotaxis dynamics of PTEN-mutated cells
Next, we analyzed how specific signaling molecules regulate the dynamics of eukaryotic chemotaxis. For example, the cytosolic concentration of PTEN molecules is either downregulated or upregulated for a cell exposed to the same initial or reversal stimulus (Sulis 2003, Subramanian et al 2007). Time courses of PIP3 accumulation at the original front or rear regions are calculated in figures 5(a) and (b), respectively. For PTEN downregulation, a significant decrease in PTEN level ($\langle \text{PTEN} \rangle = 0.03 \mu \text{M}$) induces a global high level of PIP3 concentration (blue lines), implying...
that the cell could develop multiple, rather than single, pseudopod(s). A slight decrease \((\text{PTEN}) = 0.07 \, \mu\text{M}\) reduced the delay in establishing new PIP3 at the original rear side \((\text{red lines})\), compared with the ‘wild-type’ or WT cell at \((\text{PTEN}) = 0.11 \, \mu\text{M} \,(\text{brown lines})\). For PTEN upregulation, a slight increase \((\text{PTEN}) = 0.13 \, \mu\text{M}\). Mresisted PIP3 accumulation, as observed in the reduced PIP3 concentration \((\text{black lines})\). Further increases in \((\text{PTEN}) = 0.15 \, \mu\text{M}\) completely abolished the establishment of reversal PIP3 accumulation \((\text{green lines})\), suggesting a threshold value of PTEN concentration, above which the cell fails to develop reversal polarity.

This response pattern for a PTEN-varied cell upon initial CA stimulus can be deduced intuitively. Since the PTEN acts as an inhibitor toward the Rho GTPase-PIs mediated feedback loops, a PTEN-downregulated cell becomes easily excited, whereas a PTEN-upregulated cell has high inertia to prevent the excitation. Thus, those cellular behaviors, upon a reversal stimulus, could be better understood by analyzing typical time courses of the amount of cytosolic PI3K and PTEN (figure S2, derived from a WT cell). During the reversal stimulus period \((t = 200–500 \, s)\), the strength of the positive feedback loop \((\text{Rac} \rightarrow \text{PI3K} \leftrightarrow \text{PIP3})\) decreases at the original front and increases at the opposite side, while the negative feedback loop \((\text{RhoA} \rightarrow \text{PTEN} \leftrightarrow \text{PIP2})\) acts oppositely. As a result, more membrane-bound PI3K molecules at the original front enter into the cytosol with a significant increase in cytosolic PI3K profile. In contrast, the PTEN profile demonstrates a significant reduction when more cytosolic PTEN molecules are recruited to the membrane. A critical time point at which the cell starts to rebuild its polarity was determined by the relative levels of cytosolic PI3K and PTEN. Naturally, it was more difficult for the PTEN-upregulated cell to achieve this critical point, showing a more significant delay phenomenon. Considering that the PTEN-downregulated cell failed to maintain only one major pseudopod upon initial and reversal polarization process, whereas the PTEN-upregulated cell had difficulty establishing initial polarity, it is

![Figure 4](image-url)
important for eukaryotic cells to maintain a proper level of PTEN. In addition, since the roles of PI3K and PTEN in regulating cell polarity are contrary (Parent 2002), decreasing PTEN or increasing PI3K has similar effect, and vice versa (data not shown).

Discussion

Considering theoretical modeling, our model can quantify the dynamics of rapid intracellular signaling responses in eukaryotic chemotaxis based on two aspects. First, in contrast to previous models treating intracellular signal cascades as a pure signal processing system relying on abstract inhibitory and activating components (Meinhardt 1999, Ma et al 2004), our model is centered on a molecular transport system and based on a realistic signaling network. Second, in contrast to most earlier models assuming that the interior of the cell is homogeneous (Onsum and Rao 2005), our model precisely accounts for cytosolic diffusion and the stochastic translocation behaviors of effector molecules, for example, the PIP3 profiles. Although the simplification of cytosolic diffusion generates little effect on initial polarization of the cell, it brings up significant differences during reversal polarization. Here, a dramatically high value of the diffusion coefficient (i.e. $D_c = 30 \ \mu m^2 \cdot s^{-1}$) leads to a reduction of the delay and corresponds to the generation of pseudopods at two ends (figure S3). This is not surprised biophysically, because low diffusivity enables an effector molecule that has detached from the membrane to be trapped locally via reversible binding and unbinding events when the concentration of substrate molecules is sufficiently high. By contrast, an extremely high or even infinite diffusivity constant abolishes the trapping of those detached effector molecules. Thus, the introduction of realistic cytosolic diffusion of effector molecules is essential for a model to reflect biologically competitive effects between two signaling poles, which was not explicitly described in earlier theoretical studies.

Our modeling consists of four layered modules, and each module is dominated by a specific mechanism. This strategy is supported by a body of experimental evidence. For example, latrunculin-treated Dictyostelium cells adopt a spatial sensing mechanism that does not depend on the intact cytoskeleton (Janetopoulos et al 2004). Direct activation of endogenous Rac in HeLa cells enables the bypass of several upstream signaling molecules, and the graded activation triggers cellular polarization similar to that observed in Dictyostelium cells (Holmes et al 2012). Moreover, various intermediate interactions exist among different modules (for example, PIP3 provides binding sites for various GEFs, and in turn, it regulates Rho GTPase (Bagorda and Parent 2008). Accordingly, there should be an additional PIP3 regulation module upstream of Rho GTPase. However, since this module only allows the cell to exceed the remodeling threshold without generating a shortboard effect on the entire system, this effect can be also included into the initial signaling processing module. Thus, we assumed a four layered module as a plausible scheme to embody the specific signal transduction cascades during eukaryotic chemotaxis (Parent 2004). By comparing this model with existing experimental observations, we also gained several additional insights.
into a system-level understanding of eukaryotic chemotaxis. One of the most important points is the conceptual role of an inhibitor. Inhibitors are required when modeling the signaling transduction of chemotaxis was initially postulated with the Turing-type model, in which the inhibitor is a rapidly diffusing protein, and its role is to induce diffusion-driven instability to achieve internal signal amplification (Meinhardt 1999). Following the popularity of the LEGI mechanism (Parent 1999, Ma et al 2004), it was widely accepted that a global inhibitor is mapped directly onto PTEN (Stephens et al 1999, Rottner and Stradal 2011); the latter asks for, at least, the CA-mediated PLC/PI3K signal pathway. Adding these functional components to models could further the understanding of eukaryotic chemotaxis. Second, only biochemical factors are considered in the current work. In reality, mechanical cues such as matrix stiffness (Kuo et al 2012, Schaefer and Hordijk 2015) and shear flow (Dalous et al 2008, Wang et al 2014) are also crucial. When cells sense the mechanical factors using distinct mechanisms with stretch-activated ion channels or mechanosensitive cellular adhesive molecules, they are transduced into biochemical signals with common downstream signaling pathways (Vogel and Sheetz 2009). Thus, a systematic study of chemotaxis under mechanical stimuli would unveil new mechanisms of mechanotaxis by incorporating mechanical sensing modules. Finally, while our modeling attempted to capture the dynamic features of intracellular signaling responses, the model cell is still static. Qualitatively, if a cell moves toward the source, its internal signaling dynamics would be accelerated, since a steeper CA gradient should be met. Quantitatively, however, one may observe from the CA gradient profile (figure S2) that the influence of cellular motility on the intracellular signaling responses is negligible, as long as a cell is located farther than 3λ from the source. Nevertheless, our ongoing work is to incorporate a motility module into the current model, and this integrated model would be especially useful for unraveling the spatiotemporal regulation effects of internal signaling cascades upon the tunable cellular responses.

Conclusions

While our simulations provide cellular responses that mirror most of the observed signaling responses in eukaryotic chemotaxis, several issues remain to be tested in the future. First, we simply used specific GPCR-mediated PI3K signaling pathways as the proxies for cytoskeletal remodeling, rather than modeling the entire signaling network. Actually, the coordinated polymerization and depolymerization of F-actin-based cytoskeleton are regulated by many other molecular mechanisms. For example, the former requires various ABPs, such as Arp2/3 and VASP (Stephens et al 2002, Rottner and Stradal 2011); the latter asks for, at least, the CA-mediated PLC/PI3K/GSK3 signaling pathway. Adding these functional components to models could further the understanding of eukaryotic chemotaxis. Second, only biochemical factors are considered in the current work. In reality, mechanical cues such as matrix stiffness (Kuo et al 2012, Schaefer and Hordijk 2015) and shear flow (Dalous et al 2008, Wang et al 2014) are also crucial. When cells sense the mechanical factors using distinct mechanisms with stretch-activated ion channels or mechanosensitive cellular adhesive molecules, they are transduced into biochemical signals with common downstream signaling pathways (Vogel and Sheetz 2009). Thus, a systematic study of chemotaxis under mechanical stimuli would unveil new mechanisms of mechanotaxis by incorporating mechanical sensing modules. Finally, while our modeling attempted to capture the dynamic features of intracellular signaling responses, the model cell is still static. Qualitatively, if a cell moves toward the source, its internal signaling dynamics would be accelerated, since a steeper CA gradient should be met. Quantitatively, however, one may observe from the CA gradient profile (figure S2) that the influence of cellular motility on the intracellular signaling responses is negligible, as long as a cell is located farther than 3λ from the source. Nevertheless, our ongoing work is to incorporate a motility module into the current model, and this integrated model would be especially useful for unraveling the spatiotemporal regulation effects of internal signaling cascades upon the tunable cellular responses.

Conclusions

Rapid signaling responses are indispensable in implementing functions of a eukaryotic cell. In this work, we developed a 2D mathematical model for eukaryotic chemotaxis at an intermediate level of the molecular network. At least four key related features were validated from our modeling. First, active Rac/Cdc42, PIP3, and PI3K were enriched at the front of the cell, whereas active RhoA, PIP2, and PTEN were concentrated in the opposite rear side. Cell polarization was dynamically regulated by these core intracellular molecules and reached a steady state within 1–2 min. Second, there was a significant delay in generating new pseudopods when the cell responded...
to a reversal stimulus, mainly due to competition for effector molecules (i.e. PI3K and PTEN) between two ends but not through an undiscovered global inhibitor. Third, the cell became trapped upon high-switching frequency of the direction of gradient stimulus. Finally, it was crucial to maintain the balance between the relative amounts of PI3K and PTEN molecules for proper initial and reversal polarization.

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