Mechanochemical modeling of neutrophil migration based on four signaling layers, integrin dynamics, and substrate stiffness

Shiliang Feng¹,² · Lüwen Zhou¹,² · Yan Zhang¹,² · Shouqin Lü¹,² · Mian Long¹,²

Abstract
Directional neutrophil migration during human immune responses is a highly coordinated process regulated by both biochemical and biomechanical environments. In this paper, we developed an integrative mathematical model of neutrophil migration using a lattice Boltzmann-particle method built in-house to solve the moving boundary problem with spatiotemporal regulation of biochemical components. The mechanical features of the cell cortex are modeled by a series of spring-connected nodes representing discrete cell–substrate adhesive sites. The intracellular signaling cascades responsible for cytoskeletal remodeling [e.g., small GTPases, phosphoinositide-3-kinase (PI3K), and phosphatase and tensin homolog] are built based on our previous four-layered signaling model centered on the bidirectional molecular transport mechanism and implemented as reaction–diffusion equations. Focal adhesion dynamics are determined by force-dependent integrin–ligand binding kinetics and integrin recycling and are thus integrated with cell motion. Using numerical simulations, the model reproduces the major features of cell migration in response to uniform and gradient biochemical stimuli based on the quantitative spatiotemporal regulation of signaling molecules, which agree with experimental observations. The existence of multiple types of integrins with different binding kinetics could act as an adaptation mechanism for substrate stiffness. Moreover, cells can perform reversal, U-turn, or lock-on behaviors depending on the steepness of the reversal biochemical signals received. Finally, this model is also applied to predict the responses of mutants in which PTEN is overexpressed or disrupted.

Keywords Chemotaxis · Cytoskeletal remodeling · Mathematical model · Biochemical · Biomechanical

1 Introduction
Understanding the migration mechanism of neutrophils is a crucial issue in immune responses (Kolaczkowska and Kubes 2013). Generally, as a fast moving amoeboid-type cell, a neutrophil migrates by invoking a complexly controlled and integrated sequence of events, protrusion, adhesion, and contraction (Bagorda and Parent 2008; Fenteany and Glogauer 2004). Initially, the assembly of cross-linked actin filaments drives the protrusion of a thin sheet-like structure, called the lamellipod, at the leading edge (Fenteany and Glogauer 2004; Ridley et al. 2003). Next, the cell strengthens its adhesion to the extracellular matrix (ECM) at the leading edge and weakens its adhesion at the rear edge (Nagano et al. 2012; Yang et al. 2017). Finally, myosin molecular motors bind the bundled actin filaments and exert contractile stress to pull the cell rear forward (Rottner and Stradal 2011; Shin et al. 2010). Under in vivo conditions, the direction of neutrophil migration is largely determined by the gradient of pathogen-associated biochemical signals (Jin 2013), while the cell morphology and migration velocity are dramatically affected by the mechanical properties of the substrate (Zaman et al. 2006).

Interestingly, even in the absence of biochemical guidance, an adhesive neutrophil is not static, as it constantly extends and retracts protrusions, suggesting that its cytoskele-
tal remodeling process could be excited by internal signaling noise (Swaney et al. 2010). Recently, measurements on endogenous Rac activation (Lin et al. 2012) and exogenous PIP3 delivery (Weiner et al. 2002) via bypassing upstream cascades confirmed that the polarity of the cytoskeleton is largely built upon positive/negative feedback loops mediated by small Rho GTPases (including Cdc42, Rac, and RhoA) and membrane lipids (including PIP2 and PIP3, hereafter defined as PIs) (Weiner 2002). If a neutrophil is stimulated by a graded biochemical signaling field, the GPCR-controlled, cytoskeleton-independent pathway (involving the feedback cycle of PI3K-Ras) is initiated and serves as a compass mechanism for controlling the directional regulation of Rho GTPases (Damoulakis et al. 2014), and thus, cytoskeletal remodeling (Sasaki et al. 2007).

When a neutrophil moves, complex mecano-sensing signaling pathways are activated due to adhesive bond stretching, thereby generating mechanical–chemical feedback from the substrate to the cell (Giannone and Sheetz 2006). Among various molecular bases (i.e., paxillin, FAK, vinculin, talin, and integrin) (Giannone and Sheetz 2006), integrins are the most important molecules underlying the adhesion mecano-sensing phenomenon (Shibata et al. 2013). Integrins serve as both the molecular hub for localizing adhesive molecular signaling (Scales and Parsons 2011) and as the cell–substrate physical connection for detecting mechanical properties of the substrate (Schaefer and Hordijk 2015). Using multiple techniques, such as atomic force microscopy, magnetic tweezers, and optical tweezers, the binding kinetics and forced dissociation of distinct integrin–ligand bonds have been isolated and identified (Sako et al. 2012; Elosegui-Artola et al. 2014; Liang et al. 2008). For example, the binding of β2-integrin member LFA-1 to the intercellular adhesion molecule 1 (ICAM-1) ligand yields a smaller dissociation rate and a larger association rate than that of another member, Mac-1, to ICAM-1 (Li et al. 2013).

Numerous mathematical/computational models of eukaryotic cell mobility have previously been developed (Iglesias and Devreotes 2008; Karsenti 2008; Danuser et al. 2013). While these models provide qualitative and/or quantitative insights into the distinct features of cell motility based on specific simplifications, the coupling between biochemical and biomechanical environments to regulate cytoskeletal remodeling and focal adhesion dynamics remains oversimplified. For example, previous models typically treated the cytoskeleton as a mechanical framework consisting of a mesh of discrete nodes connected by elastic springs and viscous dashpots (Sarvestani and Jabbari 2009; Dokukina and Gracheva 2010; Bottino et al. 2002). While solving the force-balance equations at each node reasonably agrees with the experimentally measured cell moving velocity, these models require that the cell be polarized (i.e., the cell has a well-defined front and rear). Meanwhile, the cell boundary alteration induced by a regulatory signaling network inside the cell is described by a conceptual polarization mechanism (Wolgemuth et al. 2011). Generally, biomarkers for the cell protrusive front are elaborately defined, but those for the contractive rear are unelucidated. A more recent model on integrative cell migration (Kim et al. 2012) incorporates focal adhesion dynamics and cytoskeletal remodeling but does not include the biophysical properties of ECM and therefore fails to capture the spatiotemporally coordinated events of intracellular signaling dynamics.

Here, we developed an integrative neutrophil migration model that couples the cellular focal adhesion dynamics and intracellular signaling responsible for cytoskeletal remodeling and morphological change. We extended our previously proposed signaling models (Feng and Zhu 2014; Feng et al. 2018) in two aspects. First, a focal adhesion dynamics module dominated by force-dependent integrin–ligand binding kinetics and integrin recycling was incorporated to describe the mecano-sensing machinery. Second, a viscoelastic mechanical model was introduced to simulate cytoskeleton-mediated cell deformation and movement. An in-house-developed lattice Boltzmann-particle (LBP) method was utilized to treat the moving boundary problem (MBP) efficiently, wherein the particle method was adopted to reassess the mechanical boundary condition and the lattice Boltzmann (LB) method for solving the reaction–diffusion system on the newly born cell domain. Additionally, a simple Monte Carlo (MC) method was embedded to treat the stochastic reaction source terms. This model can account for the spontaneous, directional, and turning dynamics of cell migration in response to uniform, graded, and reversal biochemical stimuli, respectively. It generates mutant migration via the modification of effector molecular concentrations. Moreover, varying integrin–ligand binding kinetics can serve as an adaptation mechanism to varied substrate stiffness.

### 2 Materials and methods

In this section, we summarize the theoretical modeling with simplified assumptions (for more detailed model assumptions, see the Supporting Information). This model consists of three modules. The first two modules describe the signaling cascades responsible for cytoskeletal remodeling and focal adhesion dynamics, respectively, and are governed by coupled reaction–diffusion equations. The third module denotes cell mechanics and motility and is governed by force-balance equations.

#### 2.1 Module for four signaling layers

This module involves four layers (Fig. 1a), (I) signal reception, (II) initial signaling processing, (III) small Rho GTPase
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Fig. 1  A working computational model for neutrophil chemotaxis. a Four-layered signaling cascades responsible for cytoskeletal remodeling. (i) Signal reception layer: A cell detects soluble factor fMLP by G protein-coupled receptors (GPCRs). (ii) Initial signal processing layer: G protein dissociates into $G_\alpha$ and $G_\beta\gamma$ subunits, which triggers a fast activation and a slow inhibition response. (iii) Small Rho GTPase regulation layer: Spatial regulation of Rho GTPase members (Rac, Cdc42, and RhoA) is achieved by the localization of GEFs (i.e., PAK1 and Lsc1) and antagonistic effects among the members (*denotes their active forms). (iv) Bidirectional molecular transport layer: Spatial effects generated from Rho GTPase-PI feedback loops manipulate bidirectional cytoskeletal remodeling. The curved blue and red arrows indicate the feedback loops for generating protrusive and contractive forces, respectively. The inserts show the molecular details that are treated implicitly in our model. The arrows colored blue and red in the inserts represent the pathways for providing membrane binding sites, indicating the applicability of the mass conservation law. b Schematic of the focal adhesion dynamics module. Inactive, unbound active, and bound integrins are indicated by blue, green, and red springs, respectively. The dense springs represent the stiff matrix. Four steps in integrin regulation are depicted: (i) integrin binding; (ii) once a sufficient stretch force is exerted on the bond, the bond can serve as a source point for initiating mechano-sensing signaling cascades; (iii) the local strength of intracellular integrin activation is mediated by the active Rac concentration; and (iv) ultimate amplification. c LBP method implementation of the model. Mechanically, the cell is modeled by a series of spring-connected nodes (top layer). The entire computational domain is discretized by the D2Q9 lattice model. Lattices colored red, green, and blue are specified as the membrane, cytosol, and extracellular environment, respectively. Molecular translocation described by reaction equations occurs at the shared lattices. d Simulation procedures. (i) Solving reaction–diffusion equations. (ii) Specifying nodal forces based upon the lattice concentration. (iii) Movement. (iv) Lattice–particle remapping (see Fig. S1 for more details)
regulation, and (IV) bidirectional molecular transport. Details on the overall modeling strategy and justifying the assumed cross talk among these modules have been previously described (Herant et al. 2005) and are briefly discussed below.

(I) Signal reception Diffusion and diminution of a chemoattractant, i.e., fMLP, are modeled by the reaction–diffusion equation with corresponding initial and boundary conditions

\[
\frac{\partial S}{\partial t} = D_f \nabla^2 S - KS, \quad (1)
\]

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

\[
S(x, y, 0) = 0, \quad \text{for} \quad x \notin [x_1, x_2], y \notin [y_1, y_2],
\]

\[
S(X, Y, t) = 0. \quad (2)
\]

Here, S, D_f, and K are the concentration, diffusion coefficient, and delay rate of fMLP, respectively. \(x \in [x_1, x_2]\) and \(y \in [y_1, y_2]\) define the domain of the source point wherein the concentration of fMLP equals \(S_0\). X and Y represent the boundary of the fMLP field. The binding kinetics of fMLP molecules (as ligands) and \(R, L, RL, k_+, k_-\) respectively. \(A\) is the receptor, ligand, bond, association rate, and dissociation rate, respectively.

(II) Initial signal processing A balance-inactivation mechanism is used to mimic the initial signaling process (Levine et al. 2006). This mechanism involves three interacting steps: (i) The local receptor occupancy level ([RL]) drives the production of membrane-bound species A and cytosolic species I at equal rates, \(k_+\); (ii) the cytosolic species diffuses inside the cell and attaches itself to the membrane at a rate \(k_1\) and becomes the membrane-anchored species \(I_m\); and (iii) both species A and \(I_m\) inactive each other with a rate \(k_-\), and A and \(I_m\) spontaneously degrade at rates \(\delta_A\) and \(\delta_I\), respectively. The system equations can be written as

\[
\frac{\partial A}{\partial t} = D_m \nabla^2 A + k_+[RL] - \delta_A A - k_1 I_m, \quad (4)
\]

\[
\frac{\partial I_m}{\partial t} = D_m \nabla^2 I_m + k_1 I - \delta_I I_m - k_1 A I_m, \quad (5)
\]

\[
\frac{\partial I}{\partial t} = D_c \nabla^2 I, \quad (6)
\]

with the boundary condition

\[
D_c \frac{\partial I}{\partial n} = k_3[R_L] - k_1 I. \quad (7)
\]

The translocating behaviors of PAK1 and Lsc1 from the cytosol to the membrane (like those of PI3K and PTEN) are described by the binding reaction equation [Eq. (3)], where PAK1 and Lsc1 act as ligands, while A and \(I_m\) function as receptors. The spatiotemporal evolution of cytosolic components obeys the following standard diffusion equation:

\[
\frac{\partial E_c}{\partial t} = D_c \nabla^2 E_c, \quad (8)
\]

with the boundary condition

\[
D_c \frac{\partial E_c}{\partial n} = -k_+ S_m E_m + k_- E_m, \quad (9)
\]

where \(E_c = E_{PAK1}, E_{Lsc1}, E_{PI3K}, \) and \(E_{PTEN}\) represents the cytosolic concentrations of PAK1, Lsc1, PI3K, and PTEN, respectively, \(S_m\) denotes the corresponding receptor concentration, and \(E_m\) represents the corresponding membrane-bound form.

(III) Small Rho GTPase regulation Each Rho GTPase member may cycle between active membrane-bound and inactive cytosolic forms (Lin et al. 2012). The spatiotemporal evolutions of active Rho GTPases are described as follows:

\[
\frac{\partial G}{\partial t} = D_m \nabla^2 G + P_G(R, C, \rho) \left( \frac{G_1}{G_{tot}} \right) - \omega_G G, \quad (10)
\]

where \(G = R, C, \) and \(\rho\) represent the active (membrane-bound) forms of Rac, Cdc42, and RhoA, respectively. \(G_{tot} = C_{tot}, \) and \(\rho_{tot}\) are the total amounts of Rac, Cdc42, and RhoA, respectively, \(G_1 = R_1, C_1, \) and \(\rho_1\) are the total amounts of the respective inactive (cytosol) forms of Rac, Cdc42, and RhoA that are calculated by the conservation law. \(P_G\) is the activation term and is expressed as

\[
P_R = I_R + \alpha C, \quad P_C = I_C + \beta E_C, \quad P_\rho = I_\rho + \tau E_\rho. \quad (11)
\]

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

\[
\omega_R = \delta_R + \gamma \rho, \quad \omega_C = \delta_C + \varepsilon \rho, \quad \omega_\rho = \delta_\rho + \varepsilon C. \quad (12)
\]

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

**IV) Bidirectional molecular transport** 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:

$$\frac{\partial P_3}{\partial t} = D_m \nabla^2 P_3 + \frac{H_m P_2 R}{k_{PTEN}^{cat}} \left( \frac{P_3}{k_{PTEN}^{cat}} + P_2 \right)$$

$$- \frac{\partial P_2}{\partial t} = D_m \nabla^2 P_2 - \frac{H_m P_2 R}{k_{PTEN}^{cat}} \left( \frac{P_3}{k_{PTEN}^{cat}} + P_2 \right) + \frac{M P_T R}{k_{PTEN}^{cat}} \left( \frac{P_3}{k_{PTEN}^{cat}} + P_2 \right)$$

where $\bar{R} = \min\{R/R_{max}, 1\}$ and $\bar{\rho} = \min\{\rho/\rho_{max}, 1\}$. In Eq. (13), the first term on the right-hand side accounts for PIP3 diffusion, the second term accounts for PIP3 production due to membrane-bound PI3K ($H_m$) acting on PIP2, and the third term accounts for PIP3 diminution due to membrane-bound PTEN acting on PIP3. The parameters $k_{PTEN}^{cat}$ ($k_{PTEN}^{cat} M$) and $k_{PI3K}^{cat}$ ($k_{PI3K}^{cat} M$) are based on the steady state levels of PI3. $\bar{R}$ ($\bar{\rho}$) is the normalized factor reflecting the effect of Rac (Rho) activity on PI3K (PTEN) activation, and $R_{max}$ ($\rho_{max}$) acts as a constant of Rac (RhoA) activity. If $\bar{R}$ ($\bar{\rho}$) is greater than $R_{max}$ ($\rho_{max}$), the activity of Rac (RhoA) is no longer a limiting factor for PI3K (PTEN) activation, and $\bar{R}$ ($\bar{\rho}$) then equals unity. Equation (14) describes PIP2 dynamics. Similarly, the first term on the right accounts for PIP2 diffusion, the second term accounts for PIP2 production from PIP3 via membrane-bound PTEN ($T_m$), and the third term accounts for the reduction of PIP2 into PIP3 via membrane-bound PI3K.

**2.2 Module for focal adhesion dynamics**

Formation of the focal adhesion complex is a stochastic process due to the intrinsic features of binding and unbinding between integrins on the cell membrane and their ligands on the substrate surface. A general one-step multivalent reversible reaction can be written as

$$v^i m_i + v^j m_j \xrightleftharpoons[k_{off}]{k_{on}} v^b m_b,$$

where $m_i$, $m_j$, and $m_b$ denote integrin, ligand, and integrin–ligand bonds, respectively, and $v^i$, $v^j$ and $v^b$ denote the corresponding stoichiometric coefficients ($v^i = v^j = v^b = 1$ in the current work). $k_{off}$ and $k_{on}$ represent the off and on rates, respectively, which are updated after each simulation time interval ($\Delta t_i$). $k_{off}$ is commonly described by the Bell model (Bell 1978), and $k_{on}$ is determined by the site density of active integrins. Because the timescale of integrin recycling, i.e., endocytosis or self-delay, is much longer than that of neutrophil polarization, i.e., more than 30 versus 2–3 min (Böttcher et al. 2012), the total numbers of membrane-bound active and inactive integrins are assumed to be conserved. The total numbers of active and inactive integrins are assumed to be conserved. The transformation of integrins from the inactive state to the active state is determined by the stretch force ($f_{ij}$) applied (see Supporting Information). The evolution of inactive ($w$) and active integrins ($w^*$) is schematically shown in Fig.1b and is described by the following reaction–diffusion equations:

$$\frac{\partial w}{\partial t} = D_I \nabla^2 w + \zeta w^* - \rho_{Rac} \frac{v w}{w_0 + w},$$

$$\frac{\partial w^*}{\partial t} = D_A \nabla^2 w^* - \zeta w^* + \rho_{Rac} \frac{v w}{w_0 + w},$$

where $D_I$ and $D_A$ are the diffusion coefficients for the inactive and active integrins, respectively, $\zeta$ is the self-delay rate of active integrins, and $w_0$ is the concentration of inactive integrins that gives the half maximal conversion rate. $v$ is the source term defined as follows:

$$v = \begin{cases} v_0 + \delta \Psi & \text{if } f_{ij} \geq F_r \\ v_0 & \text{if } f_{ij} < F_r \end{cases}$$

where $v_0$ is the basal rate of integrin activation, $\delta \Psi$ is the rate of force-mediated integrin activation, and $F_r$ is the threshold force. The dynamics of focal adhesion are regulated by chemoattractant concentrations based on the active level of Rac ($\rho_{Rac}$).

**2.3 Module for cell mechanics**

We use the membrane to model two biological entities, the plasma membrane and the underlying cytoskeleton or cortex. The neutrophil membrane is represented by a discrete spring-connected particle circle (with $N$ elastic springs connected at $N$ nodes). The motion of each node $i$ is described by the following force-balance equation:

$$m_i \frac{dv_i}{dt} = F_{i}^{elas} + F_{i}^{vis} + F_{i}^{pro} + F_{i}^{con} + F_{i}^{drag},$$

where $v_i$ is the velocity vector, $m_i$ is the nodal mass, and $F_{i}^{elas}$, $F_{i}^{vis}$, $F_{i}^{pro}$, $F_{i}^{con}$, and $F_{i}^{drag}$ are the elastic energy, viscous energy, protrusive, contractive, and drag forces, respectively. A detailed explanation of each term is given as follows:
Passive viscoelastic force The elastic energy stored in the springs due to stretching or compression is given by

$$E_1 = \frac{1}{2} \sum_{i=1}^{n} K_1 \left( \frac{l_i - l_0}{l} \right)^2,$$

where \(l_i\) is the length of the \(i\)th spring that is iterated at every step, \(l_0\) is the relaxed (zero-force) length, and \(K_1\) is the effective stiffness constant of the spring. The elastic energy stored in the spring due to bending is given by

$$E_b = \frac{1}{2} \sum_{i=1}^{n} K_b \tan^2 \left( \frac{\theta_i - \theta_0}{2} \right),$$

where \(\theta_i\) is the angle between a pair of consecutive springs \((i, i+1)\) that is iterated at every step, \(\theta_0\) is the relaxed (zero-force) angle, and \(K_b\) is the spring constant for bending. In addition, an area constraint is also applied to ensure that the cell area is conserved within 1% during the simulation. This is implemented via an energetic penalty as follows:

$$E_s = \frac{1}{2} K_s \left( \frac{s - s_0}{s} \right)^2$$

where \(s\) and \(s_0\) are the instantaneous and equivalent areas of the cell, respectively, and \(K_s\) is the penalty coefficient. The total elastic energy \((E_m)\) of the cell membrane is the sum of all three types of energy:

$$E_m = E_1 + E_b + E_s.$$ 

The elastic force acting on the membrane particles is then calculated using the principle of the virtual work as follows:

$$F_i^{\text{elas}} = -\frac{\partial E_m}{\partial P_i},$$

where \(P_i\) is the position vector of the \(i\)th node. The viscous force is given by

$$F_i^{\text{vis}} = -\gamma v_{ij},$$

where \(\gamma\) is the viscosity coefficient of the cellular cytoskeleton, and \(v_{ij}\) is the relative displacement rate of the neighboring nodes \(i\) and \(j\).

Active protrusive and contractive forces Physically, the generation of protrusive and contractive forces can be described by the Brownian ratchet model (Danuser et al. 2013) and the force–velocity relation of the molecular motor (Kim et al. 2012), respectively. Here, we simply assume that the nodal protrusive force \((F_i^{\text{pro}})\) is related to the local concentration of PIP3 \((P_3)\), and the nodal contractive force \((F_i^{\text{con}})\) is correlated with that of PIP2 \((P_2)\). The general forms of \(F_i^{\text{pro}}\) and \(F_i^{\text{con}}\) are given by

$$F_i^{\text{pro}} = \frac{v_{\text{pro}} \eta_i}{2P_3} \left( l_i \hat{n}_i + l_{i+1} \hat{n}_{i+1} \right),$$

$$F_i^{\text{con}} = \frac{-v_{\text{con}} \eta_i}{2P_2} \left( l_i \hat{n}_i + l_{i+1} \hat{n}_{i+1} \right),$$

where \(v_{\text{pro}}\) (\(v_{\text{con}}\)) is the protrusive (contractive) force–concentration transfer factor, which is derived from the maximum active force generated by a cell, \(P_{3s}\) (\(P_{2s}\)) is the saturation concentration of PIP3 (PIP2), and \(\hat{n}_i\) (\(\hat{n}_{i+1}\)) is the outward unit normal vector. \(\eta_i\) is a normalized factor that reflects the magnitude dependence of \(F_i^{\text{pro}}\) (\(F_i^{\text{con}}\)) on the cell–substrate adhesion strength and calculated by the following Langmuir–Hill equation:

$$\eta_i = \frac{(N_i^{b})^2}{(N_{\text{opt}})^2 + (N_i^{b})^2}. \tag{28}$$

Here, \(N_{\text{opt}}\) is the typical number of closed integrin–ligand bonds (formed only from active integrins) above which the adhesion strength is saturated, and \(N_i^{b}\) is the number of closed bonds on the \(i\)th node.

Substrate frictional drag The drag force \((F_i^{\text{drag}})\) comes from two sources, the viscous drag force and the energy dissipated by the rupture of bonds under the stretch force, which is proportional to the cell velocity \((v_i)\). \(F_i^{\text{drag}}\) can be written as

$$F_i^{\text{drag}} = (\eta_0 + \eta) v_i, \tag{29}$$

where \(\eta_0\) is the minimum value of viscosity. Based on the approach proposed in Dokukina and Gracheva (2010), \(\eta\) is assumed to be a linearly increasing function of substrate stiffness \((k_{\text{sub}})\), i.e., \(\eta = c_{\text{vis}} k_{\text{sub}}\), where \(c_{\text{vis}}\) is its slope.

2.4 Numerical simulations

To investigate mechanical–chemical coupling in cell shape and cell motion, suitable models and computational algorithms are required. Here, an LBP method developed in-house is applied to solve the MBP (Figs. 1c, d, S1). The solution of the reaction–diffusion system is then coupled to the mechanical model since it appears in the force terms \((F_i^{\text{pro}}\) and \(F_i^{\text{con}}\)), and the solution of the force-balance equations thus determines the location of the domain edge. Two time steps, \(\Delta t_L\) and \(\Delta t_M\), are defined for numerically solving the coupled system, representing time discretization in the diffusion–reaction and force-balance equations, respectively. During one simulation cycle \((T)\), assuming there are
concentration fields of PIP 3 and PIP 2 are correlated with kPa and an ICAM-1 site density of 30 A neutrophil placed on a flat substrate with a stiffness of 5 kPa and an ICAM-1 site density of 30 µm⁻² is considered to represent a typical physiological condition. As shown in Fig. 2a, the trajectories of the cell centroid are totally different in each run, indicating spontaneous cell migration, and the spatiotemporal responses of PI3s and integrins in a typical case are presented. Apparently, small Rho GTPase members undergo only self-evolution and are not spatially regulated due to the lack of directional biochemical stimuli. However, their baseline activities appear to provide a limited “driving force” to evoke cytoskeletal remodeling. In fact, acting as effectors, PI3K and PTEN, can translocate from the cytosol to the membrane. After being activated by active Rac and RhoA, respectively, the catalytic effect of PI3K leads to the conversion of PIP2 to PIP3 and vice versa for PTEN. During this process, the stochastic noise generated from effector translocation is amplified locally due to the positive feedback loop of Rac → PI3K → PIP3, leading to the appearance of PIP3 patches (Fig. 2b, top). Note that one of the PIP3 patches grows at t ~ 160 s and becomes gradually dominant. Meanwhile, as PIP2 has the same diffusivity with PIP3 and their total amounts are constant, the pattern of PIP2 regulation mirrors that of PIP3 (Fig. 2b, bottom). Considering that the concentration fields of PIP3 and PIP2 are correlated with protrusive and contractive forces, respectively, the resulting anisotropy force fields (Fig. 2c) drive cell migration, and the integrin-ligand bonds provided by the basic active integrins are stretched. The local mechanical-chemical feedback loop could be initiated if the tensile force bypasses a threshold. Thus, more active integrins are produced from the pool of inactive integrins, which, in turn, accelerates cell migration by strengthening cell-substrate adhesion. Successively, the spatiotemporal regulation of both inactive and active integrins reaches a steady state (Fig. 2d). Here, active integrins present biased distributions due to the increased local amplification from the mechanical-chemical feedback loop and limited diffusivity (Fig. 2d, bottom), while inactive integrins are distributed uniformly (Fig. 2d, top). By contrast, no biased distribution is found for bound integrins (Fig. 2e), suggesting that the accumulation of active integrins is not strong enough to introduce anisotropic cell-substrate adhesion in the absence of directional Rac activity. Time courses of cell velocity in this case (red line) and in two other cases (brown and blue lines) are also presented in Fig. 2f. In these typical cases, the cell undergoes the transition of an initial increase (0–40 s), a linear increase (40–400 s), and a stable plateau (400–700 s) along with the spatiotemporal regulation of PI3s and active integrins. On the other hand, since the cell merely develops its motility by amplifying stochastic noise, these time courses of cell velocity naturally manifest substantial differences upon each simulation (Fig. 2f).

### 3 Results

#### 3.1 Modeling spontaneous migration

A neutrophil placed on a flat substrate with a stiffness of 5 kPa and an ICAM-1 site density of 30 µm⁻² is considered to represent a typical physiological condition. As shown in Fig. 2a, the trajectories of the cell centroid are totally different in each run, indicating spontaneous cell migration, and the spatiotemporal responses of PI3s and integrins in a typical case are presented. Apparently, small Rho GTPase members undergo only self-evolution and are not spatially regulated due to the lack of directional biochemical stimuli. However, their baseline activities appear to provide a limited “driving force” to evoke cytoskeletal remodeling. In fact, acting as effectors, PI3K and PTEN, can translocate from the cytosol to the membrane. After being activated by active Rac and RhoA, respectively, the catalytic effect of PI3K leads to the conversion of PIP2 to PIP3 and vice versa for PTEN. During this process, the stochastic noise generated from effector translocation is amplified locally due to the positive feedback loop of Rac → PI3K → PIP3, leading to the appearance of PIP3 patches (Fig. 2b, top). Note that one of the PIP3 patches grows at t ~ 160 s and becomes gradually dominant. Meanwhile, as PIP2 has the same diffusivity with PIP3 and their total amounts are constant, the pattern of PIP2 regulation mirrors that of PIP3 (Fig. 2b, bottom). Considering that the concentration fields of PIP3 and PIP2 are correlated with protrusive and contractive forces, respectively, the resulting anisotropy force fields (Fig. 2c) drive cell migration, and the integrin-ligand bonds provided by the basic active integrins are stretched. The local mechanical-chemical feedback loop could be initiated if the tensile force bypasses a threshold. Thus, more active integrins are produced from the pool of inactive integrins, which, in turn, accelerates cell migration by strengthening cell-substrate adhesion. Successively, the spatiotemporal regulation of both inactive and active integrins reaches a steady state (Fig. 2d). Here, active integrins present biased distributions due to the increased local amplification from the mechanical-chemical feedback loop and limited diffusivity (Fig. 2d, bottom), while inactive integrins are distributed uniformly (Fig. 2d, top). By contrast, no biased distribution is found for bound integrins (Fig. 2e), suggesting that the accumulation of active integrins is not strong enough to introduce anisotropic cell-substrate adhesion in the absence of directional Rac activity. Time courses of cell velocity in this case (red line) and in two other cases (brown and blue lines) are also presented in Fig. 2f. In these typical cases, the cell undergoes the transition of an initial increase (0–40 s), a linear increase (40–400 s), and a stable plateau (400–700 s) along with the spatiotemporal regulation of PI3s and active integrins. On the other hand, since the cell merely develops its motility by amplifying stochastic noise, these time courses of cell velocity naturally manifest substantial differences upon each simulation (Fig. 2f).

#### 3.2 Modeling chemotactic migration

To further simulate the chemotactic behaviors of neutrophils, the substrate stiffness is set as described above, and a micropipette loaded with an fMLP solution is added close to the cell, a source point with a constant fMLP concentration [Eq. (2)]. The system runs for 350 s, and the cellular responses and spatiotemporal evolutions of key signaling components are recorded. Five seconds after introducing the source point, a steady fMLP field with a moderate gradient (7.65–8 nM) is formed surrounding the cell (Fig. 3a). The uniform distribution of GPCRs along the membrane allows the cell to sense the spatial difference in the fMLP concentration. However, the profile of occupied GPCRs at any time point (i.e., t = 50 s in Fig. 3b) inevitably displays strong stochastic noise due to the reversible binding/unbinding events. The stochastic signal received is quickly amplified by cytoskeleton-independent signaling cascades that are modeled by the balanced-inactivation mechanism [Eq. (3)]. Consequently, the spatial distributions of membrane components, represented by G_{R_f} (activator) and G_a (inhibitor), mark visible front and rear zones (Fig. S2a and b), which further transfer a clear spatial message to the downstream signaling layer of small Rho GTPase regulation by providing the binding sites for PAK1 and Lsc1, respectively (Fig. S2c and d).

The spatial profiles of the three members in the steady state are illustrated in Fig. 3c for the sake of clarity. These bipolar distributions of active Rho GTPase members sufficiently evoke the spatial effects of Rho GTPase-PI-mediated feedback loops to control cytoskeletal remodeling. Particularly, higher Rac activity at the cell front provides a higher activating rate of membrane-bound PI3K, whereas higher RhoA activity at the cell rear offers a higher activation rate of membrane-bound PTEN. As the PIP3 level is evaluated by active PI3K at the front and decreased by PTEN...
Fig. 2 Spontaneous neutrophil migration in the absence of a graded fMLP stimulus. a Cell centroid trajectories from 20 simulation runs for 700 s each. The cross denotes the starting point. b Spatiotemporal responses of PIP$_3$ (upper) and PIP$_2$ (lower) from a typical simulation. c Protrusive (outward) and contractive (inward) force fields calculated at $t = 200$ s. Local protrusive ($F_{\text{pro}}$) and contractive forces ($F_{\text{con}}$) are assumed to be proportional to the concentrations of PIP$_3$ and PIP$_2$, respectively (see Model assumptions in Supporting Information). d Spatiotemporal responses of inactive (upper) and active (lower) integrins. As the cell moves, the stretching of integrin–ligand bonds initiates mechano-sensing pathways, thus transferring more active integrins than inactive integrins. e Profile of bound integrins along the cell periphery at the end of a typical simulation. Note that no polarized distribution is observable, suggesting that the extent of active integrin accumulation is not sufficient to mediate anisotropic adhesion. f Time courses of cell velocities for three randomly selected simulations presented in blue, red, and brown. In the absence of directional cues, the establishment of cell polarity is triggered by stochastic fluctuations such that the procedures for developing cell motility differ from each other.
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Fig. 3 Directional neutrophil migration upon application of a gradient fMLP stimulus. a Spatial distribution of the fMLP field in the steady state. The source point, mimicking a micropipette containing fMLP, is applied in a spatiotemporally varied manner. b GPCR occupancy profile along the periphery of a cell at \( t = 100 \) s. The cell reads extracellular directional signals by evaluating local GPCR occupancy with a displayed strong noise. c Concentration profiles of active Rho GTPase members Rac (blue), Cdc42 (red), and RhoA (brown) in the steady state. Due to the antagonistic effects among the members and the spatial regulation effect of PAK1 and Lsc1, both active Cdc42 and Rac have higher concentrations at the cell front facing the fMLP source point, while active RhoA is centered on the opposite side. d Spatiotemporal responses of PIP3 (left) and PIP2 (right). Bipolar distribution patterns of Rho GTPase members evoke the spatial effects of Rho GTPase-PI feedback loops that drive bidirectional molecular transport, thus resulting in the all-or-none distributions of PIP3 and PIP2. e Spatial distributions of membrane-bound and cytosolic PI3K (left) and PTEN (right). Cytosolic PI3K and PTEN diffuse evenly throughout the inner region of the cell, while the distributions of their membrane-bound forms are in accordance with those of PIP3 and PIP2, respectively. f Spatiotemporal responses of inactive (left) and active integrins (right). The mechanical–chemical feedback loop initiated by stretching the integrin–ligand bond is spatially amplified upon activation of the Rac gradient (Block et al. 2016), and active integrins consequently become shapely localized at the cell front. g Bound integrin profile along the periphery of a cell at \( t = 200 \) s. The strong accumulation of active integrins at the cell front leads to a polarized distribution of bound integrins, suggesting that the cell may establish firmer adhesion at the front. h Time courses of cell velocities for three simulations in which the initially applied fMLP gradient varies from 0.0055 (blue), to 0.0035 (red) to 0.0015 nM/μm (brown). Increasing the fMLP gradient accelerates the process of achieving stable movement.
at the rear, a baseline intracellular PIP3 gradient is formed in accordance with that of extracellular fMLP. By providing the binding sites for cytosolic PI3K on the cell membrane, the gradually sharper PIP3 gradient drives the forward transport of PI3K in the cytosol. Correspondingly, since PIP2 has been replaced by PIP3 at the cell front, membrane-bound PTEN tends to enter the cytosol. These PTEN molecules reach the rear via backward cytosolic diffusion, wherein they rebind with PIP2. This bi-directional molecular transport process ceases when cytosolic PI3K is significantly depleted. By then, all-or-nothing PI distributions are formed; membrane-bound PI3K and PIP3 accumulate at the front, while membrane-bound PTEN and PIP2 accumulate in the rear (Fig. 3d and e). Accordingly, the cell generates active protrusive/contractive forces, becomes elliptical, and moves along the fMLP field gradient. On the other hand, the biased distribution of active Rac evokes the spatial effects of the mechanical–chemical feedback loop responsible for integrin activation. The local difference in Rac activity can be transferred into that of integrin activity. As a result, active integrins constantly accumulate at the front of the cell (Fig. 3f, right). This process ceases until the pool of inactive integrins becomes depleted, and a spatially biased distribution of bound integrins is formed (Fig. 3f and g), suggesting that the cell tends to establish firmer cell–substrate adhesion at the front. Time courses of cell velocity in this case (blue line) and in two other cases (brown and red lines) wherein the input fMLP gradient is either decreased or increased are also presented (Fig. 3h). The average migration rate of neutrophil is reported to be 0.059 ± 0.005 μm/s (around 30 μm/700 s) (Weckmann et al. 2017), which is comparable to the rate provided by our simulations and our verification experiments (∼ 24.5 μm/700 s) (Fig. S3, Video S1 and S2). Increasing or decreasing the input fMLP gradient tends to accelerate or decelerate the process, respectively.

3.3 Impacts of substrate stiffness and integrin–ligand binding kinetics

In addition to cytoskeletal remodeling, focal adhesion dynamics are another key factor that constrain neutrophil motility, and both processes typically occur in a coordinated manner. Despite the complexities of the aforementioned molecular mechanisms, the strength of focal adhesion is proportional to the local concentration of active integrins, which is controlled not only by intracellular biochemical factors, i.e., Rac activity (Block et al. 2016), but also by extracellular mechanical factors, i.e., substrate stiffness and integrin–ligand binding affinity itself (Peng et al. 2012).

In this section, cells expressing different types of integrins are set to undergo directional movement upon the application of substrates with varying stiffness. In each simulation, the same graded chemotactic stimuli are applied, and the displacement and shape of the cells are monitored at given time intervals. Cell velocity at the steady state is plotted against substrate stiffness and integrin–ligand binding kinetics (Fig. 4a). Generally, the estimated cell velocity is a biphasic function of substrate stiffness, which can be interpreted as follows: Cell motility results from balance among cellular adhesion strength, contractive force, and protrusive force, which are associated with the force built upon integrin–ligand bonds. This molecular-level force should either exceed a threshold that is required for successful integrin activation when more integrins become activated or reduce the dissociation rate of integrin–ligand bonds, based on Bell’s model (Bell 1978). Upon application of a certain substrate stiffness (i.e., $E_{\text{sub}} > 10\ \text{kPa}$), the buildup of the force is efficient [as explained by a serial two-spring model (Schwarz et al. 2006)], yielding a high probability of a bond beyond the threshold (Fig. S4a). Since a higher level of integrin activation corresponds to firmer cell–substrate adhesion, the magnitude of active forces is saturated. Further increasing the substrate stiffness evaluates the drag force and eventually results in diminished cell motility. In contrast, since the bond force buildup is not sufficient on soft substrates (i.e., $E_{\text{sub}} < 2\ \text{kPa}$), the relatively weak cell–substrate adhesion restricts the magnitude of the active force, and the cell velocity is thus decreased. Unsurprisingly, cells generate maximal migration velocity only at intermediate substrate stiffness.

Additionally, the dependence of cell velocity on substrate stiffness differs based on specific integrin–ligand binding kinetics, suggesting that the expression of integrin variants acts as an adaptation mechanism to varied substrate stiffness. sMac-1 (red line), where the highest velocity occurs at ∼3 kPa, serves as an example comparison. Here, LFA-1 yields higher dissociation kinetics with ICAM-1, and the estimated highest velocity shifts left upon the addition of a softer substrate (at ∼2 kPa, brown line) because the mechanical–chemical feedback loop is efficiently evoked upon the inclusion of softer substrates for integrins that possess lower dissociation kinetics. By contrast, Mac-1 processes lower dissociation kinetics with ICAM-1, and the highest velocity shifts right upon the addition of a stiffer substrate (at ∼5 kPa, blue line), as expected. Since the dissociation kinetics of specific integrins govern the level of active integrins at distinct substrate stiffness values, the association kinetics further determine the amount of bound integrins, and thus, the cell velocity.

The cell morphologies and active integrin distributions at the steady state are exemplified in Fig. 4b. For three pairs of two integrins (Mac-1, sMac-1, and LFA-1) with distinct binding kinetics, increasing substrate stiffness enhances integrin activation; the global level of active integrins is low (∼3 μM) on a soft substrate (1 kPa) and reaches saturation on intermediate (5 kPa) and stiff substrates (20 kPa). The shape of
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Substrate stiffness (kPa)

Velocity (μm/min)

Mac-1
sMac-1
LFA-1

Fig. 4 Impacts of substrate stiffness and integrin–ligand binding kinetics on neutrophil migration. The substrate stiffness varies from 1 to 20 kPa. Three sets of binding kinetics (k^off_0(s), A_kon(10^{-6} μm^3 s^{-1})), including Mac-1-sICAM-1 (1.12, 0.90) (blue in a), sMac-1-ICAM-1 (0.43, 0.39) (red in a), and LFA-1-ICAM-1 (0.5, 0.5) (brown in a), are given in the simulations. a Average migration velocity as a function of substrate stiffness and binding kinetics. In general, the stiffness dependence of cell velocity is biphasic. Altering the binding kinetics shifts the location of the peak velocity value, suggesting that the expression of distinct integrins serves as an adaption strategy upon the addition of different substrates. b Morphologies and integrin responses at varying substrate stiffness values and reaction kinetics. As the stiffness increases, the activation of integrins becomes saturated due to the increased efficiency of triggering mechanical–chemical feedback. The cell is more asymmetric on intermediate substrates (5 kPa) and less so on soft (1 kPa) and stiff (20 kPa) substrates.

3.4 Turnability of cell motility

This model can be further extended to elucidate cell behaviors in response to reversal stimuli with varying steepness values. The simulations are started from the non-polarized cell state. The same initial stimulus is applied at t = 0 s, and a specific reversal stimulus is applied at t = 350 s for an additional 550 s. The cell centroid trajectories (colored lines) and snapshots of cell shape (colored by PIP3 concentration) are presented in Fig. 5. In response to a steep reversal gradient (i.e., 0.095 nM/μm), the cell approaches the new stimulus by reversing its original motility in a nearly straight trajectory (Fig. 5a and Video S3). Interestingly, since the cell first loses its original polarity and then repolarizes correctly in accordance with the new stimulus direction, a remarkable time lag exists (e.g., from 350 to 405 s) during which the cell returns to a non-polarized, round state, and its net motility is nearly abolished. At a moderate reversal gradient (i.e., 0.055 nM/μm), the cell similarly maintains its original movement direction during the time lag period, but the polarity of the cell yields a slow rotation rather than a direct reversal (Fig. 5b and Video S4). Consequently, the cell makes a U-turn in response to the change in the gradient direction. When a shallow reversal gradient is applied (i.e., 0.035 nM/μm), the cell makes a U-turn in response to the change in the gradient direction. When a shallow reversal gradient is applied (i.e., 0.035 nM/μm), the cell maintains its migration along its original direction (Fig. 5c and Video S5).

To elucidate how cell motility can be fundamentally tuned by the steepness of the reversal stimulus, the spatiotemporal patterns of PIP3 and the time courses of cytosolic PI3K/PTEN amounts corresponding to various gradients are presented in Fig. S5. When a cell receives stimuli from the opposite direction, Rho GTPase members effectively redistribute their activities; the activated Rac and Cdc42 are detected at the incipient front and reduced at the incipient rear, while RhoA acts oppositely. Since Rho GTPase members (in their active forms) yield faster redistributing dynamics than PI3Ks and steeper reversal stimuli result in sharper Rho GTPase gradients (data not shown), the location where the highest Rac → PI3K ⇌ PIP3 positive-feedback loop strength occurs could be varied by specific reversal stimulus gradients. Accordingly, different spatiotemporal patterns of PIP3 are produced (Fig. S5a–c). By monitoring the time course of cytosolic PI3K amounts (Fig. S5d), a quantitative understanding of cell turnability can be achieved. Once the stimulus is reversed, the evaluation of RhoA activity at the incipient front reduces local PIP3 via active PTEN, which induces the dissociation of...
Fig. 5 Influence of reversal gradient steepness on neutrophil turnability. The cell originally set in a fMLP gradient and moving to the right receives the reversal fMLP stimulus at $t = 350$ s at different reversal gradient steepness values. Time-lapsed snapshots of the cell trajectories and shapes are illustrated. The colored bar on the right denotes PIP3 concentrations. 

(a) Responses of a cell to a steep reversal stimulus (0.095 nM/µm); the cell reorients itself by first losing its original polarity and then reestablishing its PIP3 polarity toward the new stimulus. Note that a time lag period exists (from 350 to 405 s) during which PIP3 accumulation nearly disappears (also see Video S3).

(b) Responses of a cell to a moderate reversal signal (0.055 nM/µm), where PIP3 redistribution is achieved in a slow rotation manner such that the cell prefers to make a U-turn to the new stimulus (also see Video S4).

(c) Response of a cell to a shallow reversal signal (0.035 nM/µm), where PIP3 accumulation is frozen such that the cell becomes stuck in its original direction (also see Video S5).
membrane-bound PI3K and thus increases the cytosolic PI3K profiles (Fig. S5d). Specifically, in response to steeper reversal stimuli, the strong rejection effect of membrane-bound PI3K, elevated from the increased RhoA activity at the original cell front, results in the fast increase in cytosolic PI3K amounts (blue line, 350–410 s). Accordingly, the original rear efficiently accesses cytosolic PI3K and forms the newly born pseudopod corresponding to the decrease in the PI3K profile (blue line, 410–470 s). When the reversal stimulus gradient decreases, the increase in cytosolic PI3K becomes slow (red line, 350–490 s) or completely ceases (brown line). The previously formed pseudopod still runs such that the cell displays a “U-turn” or “lock-on” behavior.

3.5 Responses of mutated cells

This model is also applied to test a typical biological scenario for the responses of cell mutants with distinct PTEN expressions. Using the same time-dependent initial and reversal fMLP stimuli (cf. Fig. 5a), the corresponding cell trajectories and the cell shapes are presented at a series of successive time points (Fig. 6). PTEN-overexpressed cells (i.e., [PTEN] = 0.14 µM) require more time to accumulate PIP3 and become fully polarized (Fig. 6a) because the inhibiting effect of cytoskeletal remodeling in mutated cells is stronger than that in wild-type cells. In addition, these mutants have lower migration velocities, exhibiting reductions of ~70% compared with those of wild-type cells. By contrast, cells become easily excited when PTEN is deficient (i.e., [PTEN] = 0.035 µM). The shortage of the inhibitor allows the cell to generate a pseudopod along its entire periphery. Here, the mutated cell is pushed at both the front and the rear, restricting its migration velocity. Meanwhile, the cell has a defective contractive rear (Fig. 6b).

Once the reversal stimulus is received, the mutant cells respond differently. For example, a PTEN-overexpressed cell tends to return to a non-polarized state (Fig. 6a, 350–900 s). Although a high PTEN expression level results in a remarkable inhibiting effect of membrane-bound PI3K at the original cell front, the relative ratio of [PTEN] to [PI3K] in the cytosol is sufficiently high enough that the cell fails to be repolarized. On the other hand, relatively excessive PI3K expression promotes the cell to establish its new front directly opposite from where PTEN is deficient (Fig. 6b). Collectively, the above simulations imply that maintaining proper levels of core signaling molecules in the cell is substantially important. Since the role of PI3K in mediating PI signaling is opposite to that of PTEN, cells overexpressing PI3K may exhibit behaviors similar to those of PTEN-deficient cells and vice versa.

4 Discussion

To elucidate the migration behaviors of neutrophils in different physiological environments, we herein proposed a mechanochemical model with an intermediate level of molecular details. The model is multi-modularized, containing a four-layered signaling module, a focal adhesion regulation module for mechano-sensing, and a motility module. By exploring the integrative model, we gained the following insights that help to understand the mechanisms of neutrophil chemotaxis: First, the four-layered signaling structure with the strongest feedback at the bottom layer can properly capture the spatiotemporal regulation of intracellular molecules during neutrophil migration (Figs. 2 and 3). Second, the expression of integrin variants is considered an adaptation strategy for efficient neutrophil migration upon the application of substrates of varying stiffness (Figs. 4 and S4). Third, rather than treating cytosolic effectors as homogeneous species, as assumed in earlier modeling studies (Postma and Haastert 2001; Onsum and Rao 2007), this model demonstrates that proper cytosolic diffusion generates dual effects, effective molecular transport and moderate molecular trapping, both of which are required for appropriate cell reorientation (Fig. S6). Finally, the generation of different neutrophil turning behaviors (i.e., reversal, U-turn, and lock-on) is attributed to the synergistic effects of instantaneous intracellular molecular transport and the original cellular movement (Figs. 5 and 6). To further explain the above insights in more detail, three key issues, (i) core principals for constructing our model, (ii) comparisons to experimental data and early modeling studies, and (iii) limitations of our model, are discussed below.

4.1 Core principles for constructing our model

One of the most theoretical difficulties in simulating cell motility is how to adequately describe the complicated signaling cascades responsible for cytoskeletal remodeling. Generally, these cascades are organized into a network that describes the relationship among different molecules. Cells treated with actin polymerization inhibitors become rounded and lose polarity, yet they continue to sense the direction of chemotactic gradients (Van Haastert and Devreotes 2004). By contrast, the exogenous activation (Lin et al. 2012) or delivery of signaling molecules (Weiner et al. 2002) while bypassing upstream cascades may induce chemotactic behaviors without biochemical stimuli. These experimental data imply that the signaling cascades responsible for cytoskeletal remodeling could be multilayered. From this viewpoint, the signaling process can be deciphered as follows: First, a neutrophil utilizes the GPCR occupancy to depict the local concentration of the surrounding biochemical field (signal reception, Fig. 1a-i). Second, G protein dis-
Fig. 6  Responses of a PTEN-mutated cell to the same initial and reversal stimuli. Initial and reversal stimuli gradients are set to 0.035 and 0.095 nM/µm, respectively. Cellular responses vary dramatically upon the alteration of PTEN expression. a Cell centroid trajectory and PIP₃ response to a 200% increase in the basal PTEN basal. The cell accumulates a small amount of PIP₃, corresponding to the development of a small pseudopod, and fails to repolarize in response to the secondary stimulus. b Trajectory of a cell centroid and PIP₃ response to a 50% decrease in the basal PTEN level. The cell establishes a wide range of PIP₃ accumulations and fails to develop a contractive rear.

associates into $G_\alpha$ and $G_{\beta\gamma}$ subunits, which are responsible for regulating fast activation and slow inhibition responses, respectively (Levine et al. 2006). Meanwhile, the diffusion of signaling molecules on the membrane effectively eliminates the noise from the received signal (initial signal processing, Fig. 1a-ii). Third, GEF localization along with the antagonistic effects of Rho GTPase members promotes Rho GTPase bipolar distribution patterns (Rho GTPase regulation, Fig. 1a-iii) (Raftopoulou and Hall 2004). Fourth, short-range positive and long-range negative feedback loops are generated from the depletion of effector molecules in the cytosol and the interconversion of substrate molecules (i.e., PIP₃ and PIP₂) at the membrane (Kölsch et al. 2008), resulting in molecular transport in one direction enhancing the transport in the opposite direction (bidirectional molecular transport, Fig. 1a-iv). Based on our previous studies (Feng and Zhu 2014; Feng et al. 2018) as well as those from other laboratories (Marée et al. 2012; Otsuji et al. 2007), the mechanisms that enable the spatial separation of Rho GTPase members and PIs (i.e., PIP₃ and PIP₂) are different. The former is caused by
mutually exclusive effects between Rho GTPase members, as modeled by double-negative feedback loops. The latter is induced by the spatial effects of Rho GTPase-PI-mediated feedback loops. Physically, it also relies on the translocating features of the effectors (i.e., PI3K and PTEN) along with fast molecular diffusivity in the cytosol. Our model results verified that once the clear all-or-nothing PI distribution pattern is established, the actin network is pushed at the front and pulled at the rear such that the cell achieves the most efficient movement upon the guidance of shallow, graded biochemical stimuli. In the absence of directional stimuli, however, the generation and growth of such PIP$_3$/PIP$_2$ patches rely on only the amplification of stochastic noise, and achieving significant cell motility thus requires more time.

This work also proposes a novel numerical method, LBP, which is helpful for solving MBPs of inherently deformable, non-stationary domains with redistributed biochemical signaling components. In the past two decades, several computational frameworks for MBPs have been proposed. Although these frameworks differ in their computational methodologies, their procedures are similar (Holmes and Edelstein-Keshet 2012). In principle, two separate aspects are required. First, a method of choice, such as the level-set (LS) method (Neilson et al. 2011; Shi et al. 2013), phase-field (PF) method (Camley et al. 2013), or cellular Pott (CP) method (Marée et al. 2012, 2006), is implanted to track the moving boundary. Second, any conventional numerical method, such as the finite difference (FD) method, finite element (FE) method, or finite volume (FV) method, is invoked to solve the interior reaction–diffusion problem. Comparably, in the current LBP method, the particle method is applied to specify the moving boundary conditions via coupled, nonlinear reaction–diffusion equations, wherein the partial differential equations in the irregular domains are solved by the LBM. In a sense, the greatest advantage of the LBP method derives from implementation of the LB method (Ayodele et al. 2011). On the one hand, since the computational domain mesh is achieved using small, regular lattices, the specifications of spatiotemporally changed domains can be reached by means of an elementary Boolean operation. These features provide a simpler description of complicated geometry compared with those of conventional numerical methods. On the other hand, arithmetic operations in conventional computing are performed in only the collision process (see Supporting Information), which is a localized operation, such that the LB method-based MBP framework is substantially efficient and suitable for parallel processing.

4.2 Comparison to experimental data and early modeling works

Our model can quantitatively interpret a group of distinct but interrelated experimental neutrophil chemotaxis observations. The first set of our cell chemotaxis simulations (Figs. 2 and 3) aims to compare the integrated model against the following basic aspects of experimental observations. The outputs of neutrophil migration experiments come from both cellular and molecular levels. At the cellular level, the neutrophil undergoes directional migration upon the application of graded stimuli or spontaneous migration upon the application of uniform stimuli (Nelson et al. 1975). At the molecular level, the spatial patterns of signaling molecules at different time points are presented (Wong et al. 2006; Gardiner et al. 2002). Although cellular observations have been captured by early modeling works, most rely on a single signaling mechanism achieved by abstract signaling components (Shi et al. 2013) or simple GTPase models (Wolgemuth et al. 2011; Vanderlei et al. 2011). Here, by adapting a four-layered approach to the complicated signaling cascades, we propose that when perceiving external stimuli in real time, isolation of the directional signal from strong noise, having two poles defined for actin self-organization, and effective bidirectional molecular transport are prerequisites for cytoskeletal remodeling in neutrophil chemotaxis. To this end, our model links experimental observations at the two levels. Models belonging to such multilayered categories do exist (Marée et al. 2012; Dawes and Edelstein-Keshet 2007), but they have differing dominant mechanisms for the different layers derived from distinct model cells.

The second set of simulations (Fig. 4) aims to compare with experimental data showing that the chemotaxis behavior of neutrophils is substrate dependent. Recent measurements using neutrophils (Stroka and Aranda-Espinoza 2009) as well as other eukaryotic motion cells, i.e., fibroblasts (Pelham and Wang 1997), smooth muscle cells (Peyton and Putnam 2005), and endothelial cells (Yeung et al. 2005), have demonstrated that the optimal cellular migration efficiency is achieved with the substrate of intermediate stiffness (5–10 kPa) (Janson and Putnam 2015). With stiff substrates (> 20 kPa), the cell develops mature focal adhesions at both the front and the rear, which is too firm to break and therefore cannot effectively move. Conversely, with soft substrates (< 1 kPa), the cell forms a round morphological shape and presents unstable adhesions, which are unable to support the appreciable motility. Theoretically, our model is not the first to address this issue, as experimentally observed biphasic dependences of cell migration speeds on substrate stiffness have been previously explained merely by the force balance (Dokukina and Gracheva 2010; Gracheva and Othmer 2004). By contrast, our model and the related simulations are more biologically relevant, as they link cell migration speed and receptor–ligand binding kinetics. Unlike chemoattractant stimuli, alteration of the mechanical environment is fast within a short spatial range, and biological tissue stiffness reflects significant changes over large temporal and spatial scales in accordance with healthy or abnormal tissue (Janmey...
and Miller 2011). Correspondingly, as a neutrophil adapts to chemoattractant stimuli via spatiotemporal regulation with quick activation and slow inhibition, it also adapts to substrate stiffness via expressing various types of integrins.

The third set of simulations (Fig. 5) focuses on unraveling the specific features of cell turning. The turning behaviors of neutrophils in response to a secondary reversal stimulus have been investigated since 1981 (Gerisch and Keller 1981). Accordingly, two distinct turning forms are defined with similar timescales of 1–2 min, (i) U-turn (neutrophil follows the existing lamellipodium at the leading edge to make a stepwise movement toward the source) and (ii) reversal, (cell loses its previous lamellipodium, after which a new front is induced at the opposite end and points to the new attractant source). Monitoring the time courses of F-actin and myosin II fluorescence intensities detected a significant time delay (approximately 30 s) in F-actin redistribution after reversing external stimuli (Dalous et al. 2008), providing signal processing evidence at the molecular level and the “sequence of reversal” at the cellular level. On the other hand, theoretically interpreting cellular turning behaviors is still diverse and not fully understood. For example, in the divergent pathway model (Postma and Haastert 2001) specific to neutrophils, the U-turn feature is explained by only the delayed response of intracellular signaling molecules, while the effect of the strength of the secondary stimulus is not considered. In a cellular Potts model (Marée et al. 2012) of steady keratinocyte motility, cell shape changes may feed back to intracellular signaling dynamics regulation, resulting in a faster U-turn rate. Compared with these early modeling works, our model replicates both “U-turn” and “reversal” turning features and reports the latter with the correct sequence. Such improvements are achieved by two factors. First, we successfully captured the intracellular signaling responses on a proper spatiotemporal scale (i.e., the PIs achieved the all-or-none distribution pattern in 2 min) via the definition of signaling layers specific to neutrophils. Second, such specific chemotactic features are attributed to the molecular trapping effect induced by cytosolic diffusion and the stochastic translocation of effector molecules (Fig. S6). Overall, by combining with the motility module, the actual cell fate in chemotaxis is governed by competition between the mechanical strengths of retaining the existing lamellipodium and forming a new lamellipodium. Adopting the competitive strategy enables a neutrophil to choose its target most efficiently.

The final set of simulations (Fig. 6) aims to predict how a mutated cell migrates along the chemotactic gradient. Motivated by the prioritized role of PTEN in neutrophil migration, PTEN is selectively depleted in neutrophils with a loxP-Cre system wherein Cre recombinase is expressed under the control of the granulocyte-elastase promoter (Ela2CrePtenfl) (Heit et al. 2008). PTEN-deficient neutrophils produce two or more lamellipodia and an unidentifiable uropod. Correspondingly, Ela2CrePtenfl mice also show less cell recruitment during infection, which is manifested as a lack of motility in PTEN-deficient neutrophils. On the other hand, the behaviors of cells overexpressing PTEN are equivalent to those with reduced PI3K since the roles PI3K and PTEN are reciprocal. Blocking PI3K signaling using a wortmannin inhibitor shows that the cell exhibits a poorly developed lamellipodium, crawls much more slowly, and fails to persist in forward movement, which unlikely enables its migration all the way to the stimulus source point (Wang et al. 2002). These observations are consistent with our model predication in Fig. 6b and a, respectively.

4.3 Limitations of our model

While our predictions are, in principle, consistent with those of experimental studies, and the proposed LBP method is suitable for our numerical calculations in MBPs, several issues remain. First, at least two aspects of the four-layered signaling module could be improved, one of which is in the initial signaling processing layer. Although introduction of the balanced-inactivation mechanism provides our model great simplicity, such integral treatment relying on abstract components could also be combined with other mechanisms, such as the local modulation of chemoattractant (Mackenzie et al. 2016) or a phase separation mechanism (Gamba et al. 2005). The other improvement involves the small Rho GTPase regulation layer, as only one Rho GTPase cross-talk scheme was selected from many other candidates for simplicity (Holmes et al. 2012; Jilkine et al. 2007), and this selection needs to be extended to other related signaling pathways (i.e., FilGAP) (Houk et al. 2012).

Second, while the focal adhesion regulation module is minimized to contain the two essential aspects of integrin regulation, i.e., the inside-out and outside signaling pathways (Yap and Kamm 2005), more detailed mechano-sensing mechanisms are required to achieve a broad range of neutrophil motility. For example, numerical simulation has shown that membrane tension could regulate the growth of nascent focal adhesion (He and Ji 2016). In an integrin cluster model, the force-dependent integrin clustering is considered by local chemical potential reduction (Kong et al. 2010). From a theoretical modeling viewpoint, the local activation of an existing integrin has the same effect as integrin cluster. However, this microscopic model may provide a physical interpretation of the threshold force ($F_t$) adopted here. Besides, mechanical factors are well known to also affect the regulation of Rho GTPases, especially RhoA (Schaefer and Hordijk 2015). Myosin II-dependent contraction is activated by RhoA and Rho-dependent kinase (ROCK) signaling, while the increased contraction consequently increases Rho/ROCK signaling.
Considering that a neutrophil moves relatively fast on soft substrates and no correlation between traction force and focal adhesion size is observed for mature adhesion (Stricker et al. 2011), the effect of a force-mediated RhoA signaling feedback loop is not a limiting factor for neutrophil motility in free space. However, such an effect may increase or even become dominant when neutrophils are required to enter a confined space [e.g., via a hepatic sinusoid or the ECM network (Cao et al. 2016)] whereupon the cell needs to enhance its contractility to override the resistance raised from a stiffer nucleus.

Additionally, the cytoskeletal remodeling module is quite simplified in our model. The shortage of earlier models in combination with previous observations in signaling cascade disruption experiments led to our proposal of a mechanochemical model of neutrophil chemotaxis based on signaling polarity. To a large extent, cytoskeletal remodeling is likely governed by the polarities of signaling molecule distributions; however, their intercorrelation is far more sophisticated. A variety of mechanisms, including G-actin treadmilling (Danuser et al. 2013), dendritic nucleation at the leading edge (Mogilner and Edelstein-Keshet 2002), and hydrodynamic flow (Niwayama et al. 2011), may function together to regulate the dynamic patterns of actin–myosin flow. Future models may involve the related molecular components in cytoskeletal remodeling based on the experimental results of disrupting actin-binding proteins.

Finally, as an old saying goes, every coin has two sides; every numerical simulation platform has its advantages and limitations. In the current framework of the LBP method, combining the auto-parallelization feature of the LB method, treating the cell edge as a collection of particles, and introducing a constriction area allow us to treat the signaling cascades very efficiently and with high numerical stability. Accordingly, in the future, the LBP-based system may be further extended to a multi-cellular system, wherein the chemical interactions between cells and intracellular dynamics are of great interest. However, application of the LBP method comes at the price of losing mechanical details. Currently, problems such as the stochastic features of cell edge protrusions–contractions, the effects of cell shape feedback on cell motility, and the interactions of cells with obstacles have not been included. To these regards, the cellular Potts method (Marée et al. 2012) is more ideal.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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