A mesoscale mechanical model of cellular interactions

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ABSTRACT Computational models of cell mechanics allow the precise interrogation of cell shape change. These morphological changes are required for cells to survive in diverse tissue environments. Here, we present a mesoscale mechanical model of cell-substrate interactions using the level set method based on experimentally measured parameters. By implementing a viscoelastic mechanical equivalent circuit, we accurately model whole-cell deformations that are important for a variety of cellular processes. To effectively model shape changes as a cell interacts with a substrate, we have included receptor-mediated adhesion, which is governed by catch-slip bond behavior. The effect of adhesion was explored by subjecting cells to a variety of different substrates including flat, curved, and deformable surfaces. Finally, we increased the accuracy of our simulations by including a deformable nucleus in our cells. This model sets the foundation for further exploration into computational analyses of multicellular interactions.

INTRODUCTION

A cell’s ability to change shape and interact with its environment is essential for growth, proliferation, tracking and consuming nutrients, and survival. In single-cell and multicellular systems, a variety of morphological changes are required for cells to interact with other cells and their local environment. Cell consumption events such as phagocytosis, entosis, endocytosis, and micropinocytosis all are characterized by a cell deformation in which external materials, including nutrients, media, particles, or other cells, are engulfed. These events represent a diverse array of biological processes that are present across a wide range of cell types. The mechanical properties of these cell engulfment processes are well characterized and therefore provide a good basis to begin interrogating multicellular interactions through the reduced computational requirements of the two-cell scenarios.

Simulating cell engulfment events requires considering cellular interactions and understanding how cells adhere to these surfaces, which include the substrate or neighboring cells. Cell-cell interactions can be modeled at various length scales, from subcellular simulations considering interactions of individual receptors that enable cell adhesion (1,2) to tissue-level simulations representing hundreds of cells (3,4). To accomplish modeling across these ranges of complexity requires a variety of different computational techniques. Here, we consider a mesoscale-scale model involving simulations of one to two cells with a focus on recreating the corresponding cell shape changes. Many techniques are available for modeling deformable cells with high geometric resolution (5). Most of these have been used to simulate migration (6–9), although some have been used in other biological processes such as cell division.
(10–12) and phagocytosis (13). We implement our approach using the level set method (LSM), which was originally developed to represent interactions across phases and can track moving boundaries accurately.

Though the LSM has been successfully used to simulate various cellular processes that involve individual deformable cells (10,14,15), to our knowledge it has not been previously used to interrogate cell-substrate contacts or cell-cell interactions such as engulfment. To incorporate these phenomena, which involve receptor-mediated adhesions, requires that we consider a model for these adhesions. Our implementation assumes that adhesions arise from catch-slip bonds. Catch-slip bonds are characterized by having biphasic behavior depending on the force applied. During the catch phase, the dissociation decreases transiently as the strength of the tensile force increases (16,17). In contrast, during the slip phase, after sufficient tensile force has been applied to the bond, the rate of dissociation increases. These can be represented mathematically as in the Bell model (18). Many catch-slip proteins such as cadherins are necessary for healthy cell development, migration, and physiology (19,20).

In this study, we begin by briefly describing the implementation of the LSM for deformable cells, followed by a description of how active stresses and adhesion are implemented in this framework with both solid and deformable substrates. The model is first tested with cells interacting with flat solid substrates to verify the implementation of adhesion. We then explore the landscape of more biologically relevant cell engulfment events through modulation of mechanical parameters and stress production. We further increase the biological accuracy of these simulations through the introduction of a mechanically deformable nucleus. Ultimately, this model lays the foundation to understand better the mechanistic details of cellular interactions and cell consumption events and to predict cell behavior across different scenarios.

MATERIALS AND METHODS

Model overview

A cell’s shape is affected by a variety of different mechanical parameters that are specific to the cell type as well as its environment. We considered a range of physical processes, which include experimentally measured physical parameters such as cortical tension and viscoelasticity as well as various active and passive stresses (Fig. 1 A; Table 1). Cortical tension is the energy cost for adding a unit of surface area to the cell (21). Cortical tension, along with the local curvature of the cell, creates a Laplace-like pressure that drives the cell to minimize the surface area/volume ratio. Additionally, the viscoelasticity of the cell is determined by the components of the cytoskeletal network, which provide a time-dependent response to stresses imposed (22). We represented the viscoelastic cell with a mechanical equivalent circuit representing the cell cortex and membrane (Kelvin-Voigt) in series with a purely viscous cytoplasm (Fig. 1 B). The resulting cell shape is determined by the balance of stresses. These stresses range from the small stochastic membrane fluctuations (Fig. 1 C) that contribute to cell adhesion to active stresses driven by actin polymerization and myosin contractility to several passive stresses, such as the propensity to conserve cellular volume and the Laplace-like pressure described above (Fig. 1 D). We include adhesion between the cell and its environment, which can be either an extracellular matrix or a neighboring cell.

Implementing the LSM

The LSM is a numerical technique that tracks the evolution of an interface with high geometric accuracy while improving computational efficiency by defining boundaries on a fixed Cartesian grid and eliminating the need for parameterization (24). Originally developed to simulate propagating fronts with curvature-dependent speeds (31,32), the LSM has been used in various biological settings (5). Here, we highlight some of the basics; a detailed description of the LSM implementation in modeling cells is provided in (24).

The LSM uses an implicit scheme to define arbitrary surfaces. The cellular boundary, $\Gamma(t)$, is defined as the zero-level isocountour of potential function $\phi(z, t)$, defined as one dimension higher than $\Gamma(t)$. In our simulations, $z = (x, y) \in \mathbb{R}^2$; this renders a two-dimensional (2D) simulation representing a cross section through the cell. Though several possibilities exist for choosing $\Gamma(t)$, there are numerical advantages to choosing a signed distance function to describe the potential function $\phi(z, t)$ (33). In particular, the distance from point $z$ to the boundary $\Gamma(t)$ is described by

$$\phi(z, t) = \pm \min_{z' \in \Gamma(t)} \|z - z'\|,$$

where the sign is +1 for points outside of the boundary and −1 for points inside. Throughout the evolution of the simulation, if the boundary deviates from the signed distance function (the normal of the gradient of $\phi$ is greater than a defined parameter threshold), it can be reinitialized using the function

$$\frac{\partial \phi}{\partial t} + \text{sign}(\phi)(|\nabla \phi| - 1) = 0.$$

In simulations in which cells undergo significant deformations or experience significant stresses, reinitialization was required more frequently.

Inclusion of stresses in LSM

The evolution of $\phi$ is described by the equation

$$\frac{\partial \phi}{\partial t} + v \cdot \nabla \phi = 0,$$

where $v$ describes the local velocity of the potential function. To calculate the velocity, we first compute the various stresses acting on the cell and use a viscoelastic mechanical equivalent circuit of the cell to determine their effect on the local velocity. The velocity of the cell is then given by $v(z, t) = \dot{\delta}_{\text{bound}}/\partial t$, where $\delta_{\text{bound}}$ takes into account the deformation of the cytoplasm and cortex, as described below (Eq. 5). As per the level set formalism, the total stresses acting on the cell ($\sigma_{\text{tot}}$) are applied outward or inward normal to the cell boundary and include passive and active stresses.

We consider two passive stresses:

$$\sigma_{\text{passive}} = \sigma_{\text{ten}} + \sigma_{\text{area}}.$$

Surface tension ($\sigma_{\text{ten}}$) captures the Laplace-like pressure found at the interphase between the cell and its environment; it is given by

$$\sigma_{\text{ten}} = \gamma_{\text{ten}}\kappa(z, t),$$
where $\gamma_{\text{ten}}$ is the local cortical tension, $\kappa = \nabla \cdot \mathbf{n}$ is the local curvature, and $\mathbf{n}$ is the unit normal vector. We assume that the cytosol is incompressible, which implies that the cell volume remains constant. In this two-dimensional model of the cell, we enforce this by constraining the area enclosed by the cell boundary. To this end, we implement a stress given by

$$s_{\text{area}} = \gamma_A (A(t) - A_0),$$

where $A(t)$ and $A_0$ are the areas enclosed by the cell boundary at times $t$ and $t = 0$, and $\gamma_A$ is a conservation term used to ensure that the cell’s area does not change over the course of the simulation (Fig. S1). Note that we do not assume that the perimeter is conserved because *Dictyostelium* is known to shuttle membrane so that it is not limiting (34). When considering other cells, the perimeter could also be constrained in a similar manner.

The addition of active stresses,

$$\sigma_{\text{active}} = \sigma_{\text{Brown}} + \sigma_{\text{pro}} + \sigma_{\text{ret}},$$

increases local velocity, thereby displacing the cell membrane. Active stresses are a combination of Brownian motion ($\sigma_{\text{Brown}}$) due, in part, to thermal fluctuations at the membrane; protrusive stress ($\sigma_{\text{pro}}$), representing actin polymerization at the front end; and retractive stress ($\sigma_{\text{ret}}$), representing myosin contractility at the rear end. All stresses are applied as a vector normal to the boundary of the cell. Protrusive and retractive stresses are modeled deterministically; an example of their spatial distribution can be seen in Fig. S2. In our model, we do not explicitly model subcellular actomyosin concentrations or activity, nor do we consider how they may change over time in response to a mechanical stimulus (35,36). Instead, we apply active stresses based on the experimentally measured actomyosin force production at regions experiencing active protrusive or retractive stresses on predetermined regions of the cell.
Brownian stress SD deviation given by catch bond $f$ E-cadherin binding parameters slip bond $s$ attachment threshold $G_s$ on rate $G_s$ Protrusive stress $g$ Cytoplasmic viscosity $g$ (brane, and their magnitude at each point is recomputed at each iteration 50 nm (41,42). To determine the stress required to produce these fluctuations (25,26,22). These values have been well quantified experimentally (25,26,22).

Initial cell-substrate contact can be initiated by stochastic contact through Brownian motion (Fig. 1 C). Observable membrane fluctuations can be caused by thermal noise and by underlying cellular activity driven by the cytoskeleton (37–40). When computing the magnitude of Brownian motion, we do not model temperature explicitly. Instead, we use experimental measurements of supported intermembrane junctions which suggest that a membrane fluctuates with a root mean-square displacement ($\Delta r^2$) of ~50 nm (41,42). To determine the stress required to produce these fluctuations, we worked backward. As shown in the supplemental file, the magnitude of the stress required to produce these fluctuations has a standard deviation given by

$$\sigma_{\text{Brown}} = \left(\frac{2k_c \gamma_c}{dt} \Delta r^2\right)^{1/2}.$$  

(1)

We assume that these fluctuations primarily affect the membrane of the cell, which in our model is represented as the compound cortex (consisting of the membrane and underlying actomyosin cytoskeleton); therefore, only the Kelvin–Voigt element of the viscoelastic circuit was included in the calculation. The frequency with which the fluctuations occur varies as a function of the time step ($dt$) assigned for the simulation. The Brownian fluctuations ($\sigma_{\text{Brown}}$) are uniformly distributed across the entire cell membrane, and their magnitude at each point is recomputed at each iteration of the simulation to capture the stochasticity of the process. This, combined with the protractive and retractive stresses ($\sigma_{\text{area}}, \sigma_{\text{cell}}$), acts as the active stress ($\sigma_{\text{active}}$). The total stress (expressed in nanonewtons per square micron) is then given by

$$\sigma_{\text{tot}} = \sigma_{\text{passive}} + \sigma_{\text{active}}.$$  

(2)

When only passive stresses are considered, they are sufficient to instigate contact between a cell and a substrate. When active stresses are added, these passive stresses become nearly negligible. At points of contact, the cell shape is also restricted to ensure that it remains excluded from the volume of the interacting surface. This is done by constructing a mask potential function representing the shape of the surface with which the cell is interacting (43). At points at which the cell’s potential function intersects the mask’s potential function, the cell’s shape is clipped by

$$\phi(t + \Delta t) = \min(\phi(t + \Delta t), \psi),$$

where $\psi$ is the potential function of the mask. When considering two deformable cells, each exerting active $\sigma$, each cell applies a stress to the other proportional to the local velocity. The total stress for each cell is computed independently before applying a stress to the opposite. Specifically, if two cells, denoted 1 and 2, exert total stresses $\sigma_{\text{tot}1}$ and $\sigma_{\text{tot}2}$, respectively, at a common contact point, then the net stress affecting cell 1 is given by

$$\sigma_{\text{1- net}} = \sigma_{\text{tot}1} - |\sigma_{\text{tot}2}|,$$

with a similar formula for the stress on cell 2.

**Modeling a cell with a deformable nucleus**

To model a cell with a deformable nucleus, a second level set function was used and was chosen so that initially, the nucleus was near the cell centroid. The nucleus was assumed to occupy 40–50% of the area of the cell, with elasticity ranging from 0.25 times to four times that of a wild-type cell. The nucleus is modeled as a viscoelastic material based on the same mechanical equivalence circuit shown in Fig. 1 B. In this way, we are not considering the more complex nuclear dynamics that occur after mechanical deformation (44,45). The outer cell membrane acts as a mask for the inner nucleus and is described as

$$\phi_1(t + \Delta t) = \max(\phi_1(t + \Delta t), \phi_2(t + \Delta t)), $$

where $\phi_1$ and $\phi_2$ are the potential functions for the nucleus and the cell, respectively. Masking prevents the nucleus from leaving the boundary of the cell membrane. As the outer cell deforms according to the velocity resulting from the total stress as defined by Eq. 2, the nucleus will move tracking the cell by modeling molecular linkages of the nucleus to the cytoskeleton as elastic springs following Hooke’s law:

$$\sigma_{\text{link}} = k_{\text{Hooke}} \left\| z_1(t) - z_2(t) \right\|,$$

where $z_1(t) = (x_1(t), y_1(t))$ and $z_2(t) = (x_2(t), y_2(t))$ are the centroids of the nucleus and cell at time $t$, respectively, and $k_{\text{Hooke}}$ is the Hooke constant. This term resulted in a biologically relevant stress regime of fewer than 2 nN/μm² (46,47). This stress was applied as a Gaussian on the perimeter of the nucleus and is described as

$$\phi_1(t + \Delta t) = \min(\phi_1(t + \Delta t), \psi),$$

where $\psi$ is the potential function of the mask. When considering two deformable cells, each exerting active $\sigma$, each cell applies a stress to the other proportional to the local velocity. The total stress for each cell is computed independently before applying a stress to the opposite. Specifically, if two cells, denoted 1 and 2, exert total stresses $\sigma_{\text{tot}1}$ and $\sigma_{\text{tot}2}$, respectively, at a common contact point, then the net stress affecting cell 1 is given by

$$\sigma_{\text{1- net}} = \sigma_{\text{tot}1} - |\sigma_{\text{tot}2}|,$$

with a similar formula for the stress on cell 2.

**Inclusion of cell adhesion**

We implement adhesion in the form of catch-slip bonds based on measured values for different cell-substrate and cell-cell adhesion proteins. From a
biomechanical perspective, these types of adhesion bonds may be incorporated somewhat generically based on a biophysically relevant stress landscape representing many adhesion proteins. Catch-slip bonds are force dependent, where at lower force, the bond lifetime decreases until reaching a transition point, and at forces higher than this transition point, the bonds release, or “slip” (Fig. 1 E) (48). We have modeled catch-slip bonds by assuming that attachment leads to a stress threshold that must be overcome before the attachment can be broken. The attachment of cells that undergo a catch-to-slip transition can be described by the following equation (23):

$$\frac{dN}{dt} = -N(e_{p}N + e_{c}N) + \gamma(N_{t} - N). \quad (3)$$

Here, $N_{t}$ is the total number of adhesion proteins in a cluster that can bond, and $N$ is the number of these bonds that form attachments. $F$ is the total force acting on this cluster, and $\gamma$ is a protein-specific attachment rate. The two variables, $\phi$, and $\phi_{s}$ describe the catch- and slip-bond terms, respectively. We consider several experimentally measured bonds, including E-cadherins, P-selectins, integrins, and fibronectins (28,29,49–51). In our model, we assume that this equation is at equilibrium, leading to a uniform number of bonds per unit length, and that each unit length of bond immediately reaches steady state upon contact, as experimentally measured bond kinetics show that E-cadherins and P-selectins reach steady state in less than 2 s (27,52,53), which is considerably faster than the time taken for the cellular deformations that we consider. Eq. 3 has two equilibria (Fig. 2 A), given by

$$F(N) = N\phi_{max} \pm N\cosh^{-1}\left(\frac{1}{2N}\sqrt{e_{s}N_1 - N_2}\right), \quad (4)$$

where $\phi_{max} = \frac{1}{2}(\phi_{c} + \phi_{s})$. The stable (− sign) and unstable (+ sign) solutions meet at a point at which the number of attached bonds reaches a maximum, $N_{max}$, which is achieved for applied external force

$$F_{max} = N_{max}\phi_{max} - N_{max}\frac{\gamma e^{1/2}(\phi_{c} - \phi_{s})}{\gamma e^{1/2}(\phi_{c} - \phi_{s}) + 2}. \quad (5)$$

In our model, we take this to be a threshold determining whether the membrane is attached. To implement this threshold, we use this calculation to compute an adhesion stress threshold $\sigma_{tot}$ based on substituting an adhesion protein density for $N_{t}$. For example, for E-cadherin, which has $\phi_{max} = 27$ proteins/μm, $\gamma = 5.3 \times 10^{-2}$, and $N_{t} = 64$ proteins/μm², the stable and unstable solutions of Eq. 3 converge at an applied stress of $1.7$ nN/μm². We assume that the adhesion proteins are uniformly distributed along the boundary of the cell and the substrate. Then, the two-dimensional level set function is discretized onto a Euclidean grid (dx) to determine points of contact between the cell and the substrate. We then determine the magnitude and direction of the total stresses acting on the cell at each point of contact. We use the result from Eq. 5 as a threshold $\sigma_{tot}$ that must be overcome to pull the cell away from the surface. For example,

$$\sigma_{tot-net} = \min(\sigma_{tot} + \sigma_{max}, 0). \quad (6)$$

In this way, we are not modeling individual bonds but use a threshold that is based on experimentally measured, protein-specific bond density and kinetics as well as bond breakage forces. If the total stress at that point is less than the threshold of adhesion, the point remains adhered. Alternatively, if the total stress is larger, then the point detaches. Using the bond numbers described above, we found threshold values in the range of 1–5 nN/μm², which are consistent with experimental measurements of cell attachment (54,55).

Implementation of mechanics

With these parameters implemented into the viscoelastic model of cells, we can begin to dissect the machinery and stresses needed for cell-cell interactions such as those we see in entosis and other cell consumption events. Through experimental measurements using micropipette aspiration and atomic force microscopy, we know that cell mechanics can be modeled using the Kelvin-Voigt-Dashpot viscoelastic model (24,56). In particular, our viscoelastic model assumes that the cell is a compound material consisting of a mostly elastic membrane and cortex and a mostly viscous cytoplasm (Fig. 1 B). The evolution of the local deformation along the normal direction of the boundary is given by

$$\dot{x}_{cyto} = \frac{1}{\gamma_{c}}\sigma_{tot} \quad (7a)$$

and

$$\dot{x}_{cort} = -\left(k_{c}/\gamma_{c}\right)x_{cort} + \left(1/\gamma_{c}\right)\sigma_{tot}, \quad (7b)$$

where $\sigma_{tot}$ is the total stress applied on the cell; $x_{cyto}$ and $x_{cort}$ are the local displacements of the cortex and cytoplasm, respectively; and $k_{c}$ (0.098 nN/μm²), $\gamma_{c}$ (6.1 nN/μm³), and $\gamma_{c}$ (0.064 nN/μm³) are viscoelastic components of the cell describing the elasticity ($k_{c}$) and viscosity ($\gamma_{c}$) of the cortex and the viscosity ($\gamma_{c}$) of the cytoplasm. Thus, the normal velocity of the boundary is the sum of $\dot{x}_{cyto}$ and $\dot{x}_{cort}$.

Computational implementation

A summary of the steps in the model is given in Fig. S3. The model was implemented and simulated using MATLAB (The MathWorks, Natick, MA) and the LSM Toolbox (43). Using the LSM Toolbox derivative function called Upwind First First (first-order upwind approximation of the first derivative), the integrator function called ode CFL1 (first-order Euler scheme with a Courant-Friedrichs-Lewy constrained time step of ~0.01), and a grid size (dx) of 0.1 μm, we can model ~1.5 biological hours per hour of computational time. On a grid size of 40 × 50 μm, run on a MacBook Pro 2.4 GHz 8-Core Intel Core i9, 1 h of simulation time resulted in a little less than an hour of biological time.

RESULTS

Modeling cell-to-substrate adhesion

We began by simulating the interaction between a spherical cell and a solid surface. To move away from its equilibrium state, a cell must experience force, which may be internally generated or externally applied. One of the smallest forces seen in a biological system arises from Brownian fluctuations. The magnitude of thermal fluctuations of the membrane has been measured experimentally using a variety of techniques, including fluorescence interference contrast microscopy (41,42). The resulting measurements indicate that a detached membrane fluctuates with a root mean-square displacement of ~50 nm. Using the Kelvin-Voigt portion of the mechanical equivalent circuit, we calculated the stress required to displace the membrane to the experimentally measured parameters (~55 pN/μm², Materials and methods). The resultant simulated fluctuations were smaller than the experimentally measured values (which only consider membranes), as our system is modeled as a compound material,
consisting of the membrane, cortex, and underlying cytoplasm, all of which limit the magnitude of deformation resulting from the applied stress (Fig. 2A).

If the Brownian fluctuations were sufficiently large to instigate contact between a point on the cell perimeter and the substrate, that point would adhere. In our model, adhesion is implemented as a catch-slip bond. The lifetime of these bonds is a function of the applied mechanical load. The threshold that transitions a bond between the catch

regime and the slip regime was modeled based on an adaptation of the original Bell model (Fig. 2B) (18,23). This formulation allows the strength, density, and kinetics to be modulated to represent diverse types of adhesion molecules increasing the flexibility of our model. We computed the adhesion landscape across a range of catch-slip adhesion molecules, which depends both on the adhesion protein density and the characteristic forces necessary for the catch-slip transition for different adhesion molecules. We found that

FIGURE 2 Implementation of cell-substrate adhesion. To model multicellular interactions effectively, a biologically accurate model of adhesion must be incorporated. (A) The stress required to form thermal fluctuations was used as an input to the mechanical circuit and resulted in small deformations uniformly around the cell perimeter. The kymograph illustrates the magnitude of the resulting cellular deformations across the cell boundary over time. (B) The threshold between catch and slip was determined by the intersection of the stable (solid line) and unstable (dotted line) equilibria for E-cadherin (red) and P-selectin (blue). This point represents the maximal stress on the bonds that transitions the adhesion from the catch regime to the slip regime. (C) The point of catch-slip transition varies as a function of adhesion protein density and the necessary stress to mediate the transition between catch and slip, creating an adhesion landscape. Here, the density of adhesion molecules and the detachment force of the molecules are varied in a biologically relevant regime. The color gradient indicates the maximal stress the bonds can withstand before transitioning into the slip regime. (D) The adhesion landscape illustrated above is used to determine the cell-specific cell mechanics and the strength of the specific adhesion molecule (shown here are E-cadherins and P-selectins). For all tested molecules, as cortical tension of the cell decreases, we see increased cell settling, which is maintained for longer periods. Shading indicates the 95% confidence interval. (E) Cell compression simulations validate the strength of the adhesion between the cell and the substrate. When the adhesion between the bottom substrate and cell is higher (top), the cell remains attached to the bottom. In contrast, when the adhesion of the top moving beam and cell is higher, the cell will remain attached to the top beam as it moves upward with a known velocity. The surface with higher adhesion is illustrated in yellow. WT, wild-type. To see this figure in color, go online.
this implementation of integrin adhesion is linearly dependent on the density of the adhesion proteins (Fig. 2 C).

When we simulated the effect of thermal fluctuations along with catch-slip bonds, we observed moderate adhesion to the surface. With a catch-slip threshold approximating E-cadherin adhesion (1.7 nN/μm²) and wild-type cortical tension (1 nN/μm) (24), the bottom 1 μm of the cell perimeter adhered for ~30 min before the passive stresses of the cell (volume, curvature, and cortical tension) overwhelmed the bonds and detached (Fig. 2 D). We found that the amount of adhesion to the substrate was a function of the cortical tension of the cell. As the cortical tension was decreased, the fraction of the adhered cell perimeter increased, and the length of time that it remained adhered increased as well. Increasing the threshold from catch to slip of the molecular linkage, as seen in P-selectin (catch-slip threshold, 5.2 nN/μm²), raised the extent of cell perimeter adhesion for longer periods compared to E-cadherin (Figs. 2 D and S4).

To validate the implementation of adhesion further, we simulated a cell compression experiment (55). To mimic the experimental design, the simulated cell was compressed between two solid substrates before the top surface was moved away with a known velocity (Fig. 2 E). In simulations in which the adhesion between the cell and the bottom surface was 5× stronger than adhesion to the top surface, the cell remained attached to the bottom, whereas the bonds broke between the top surface and the cell. Conversely, when the adhesion was stronger on the top, the cell remained adhered to the top surface after detaching from the bottom (Fig. 2 E).

Modeling active stress production

Brownian fluctuations of the cell membrane allow for stochastic contact between cells and substrates that are within proximity. These fluctuations, along with the implementation of adhesion as described above, led to some deformation but were insufficient to overcome passive cortex retraction to displace the cell significantly on flat or curved surfaces (data not shown). We next considered the role of active stresses in ensuring cell adhesion. To this end, we added a protrusive stress of magnitude 2 nN/μm² to the bottom 2 μm directly adjacent to areas of adhesion (Fig. S2). With the added protrusive stress, the cell deformed considerably more compared with the simulations with Brownian fluctuations exclusively (Fig. 3 A). As before, the cells began with a single point of contact between the cell boundary and the substrate. For a cell with a cortical tension of 1 nN/μm, the bottom ~3 μm of the cell contacted the bottom

![Figure 3](https://example.com/figure3.png)

**FIGURE 3** Effects of internal and external stresses on cells. The addition of active stress modeling actin polymerization results in cellular deformations. (A) A three-dimensional visualization of a two-dimensional simulation rotated about the axis of symmetry showing a cell settling with an active normal stress of 2 nN/μm² on the bottom 2 μm of nonadhered cell perimeter. (B) Cell perimeter contact over time for cells experiencing 2 nN/μm² active normal stress on the bottom 2 μm of nonadhered cell perimeter. For both E-cadherins and P-selectins, as the cortical tension of the cell decreases, the amount of cell perimeter that forms a contact with the bottom substrate increases. (C) Active normal stress can be applied to the cell in different directions to simulate varied experimental designs. Here, the stress is applied outward normal to the right. The outward normal stress was sufficient to navigate around obstacles in its path. To see this figure in color, go online.
substrate. As before, the fraction of the cell perimeter able to make and sustain contact with the surface increased as a function of decreasing cortical tension. We observed similar behaviors in the simulations of adhesion proteins with characteristics of either E-cadherin or P-selectin adhesion proteins. In fact, the deformation between these two simulations was quite similar (Fig. 3 B). Despite the difference in the strength of adhesion between these two different proteins, the magnitude of the added active stress was sufficient to overcome the passive retractive stresses that are inherent to the cell.

To simulate more diverse biological environments, we expanded the region of the cell perimeter over which these stresses are applied. To simulate a shear stress in a microfluidics-type environment, the stress can be applied outward normal to the right portion of the cell perimeter. In these simulations, the cell first was allowed to settle (as described above) and made initial contact with the bottom surface before the cell moved to the right. The environment could be made more complex by adding obstacles around which the cell had to deform (Fig. 3 C).

**Interactions with solid curved surfaces**

To establish the foundation of cell-cell interactions such as cell consumption events, we simulated an active cell as described above interacting with a solid passive cell. In cell engulfment events such as entosis and phagocytosis, a cell deforms around a target cell, ultimately consuming it. By testing a variety of different parameter sets, we determined a landscape of cell-cell interactions.

We first simulated active cells interacting with passive cells (more specifically, passive circular objects) of varying sizes (Fig. 4 A). In our simulations, an outer engulfing cell with a diameter of 10 μm interacted with passive cells with diameters ranging from 2 to 14 μm. We considered an outer cell generating outward normal stresses of either 3 or 6 nN/μm² over either 1 or 2 μm of unadhered cell perimeter (25,26). Cells that experienced higher stress over a larger region of the cell perimeter (6 nN/μm² over 2 μm) completed engulfment events faster than cells experiencing less stress over a smaller area (3 nN/μm² over 1 μm). In some circumstances (e.g., outer cells with active stresses only over the bottom 1 μm of the cell), the stress was not sufficient to complete the engulfment. Additionally, we saw that cells could consume passive cells up to an equivalent size to themselves. However, if the engulfing cell was smaller than the passive target cell, then it was unable to complete an engulfment event even at the highest stresses tested. At points of contact, the cells have adhesion with a catch-slip threshold of 1.7 nN/μm². For a 6 μm cell (active normal stress of 5 nN/μm² over the bottom 3 μm of the cell perimeter), as the size of the passive cell increased from 1 to 6 μm, the time to engulf the surface also increased from 30 to 144 min, respectively (Fig. 4 B).

We next considered the role of cortical tension during engulfment. Simulations showed that as the cortical tension of cells was reduced locally, the cells were able to complete cell consumption events more efficiently at all sizes of passive cells tested. Additionally, through this local cortical tension reduction, cells that were previously unable to complete an engulfment event were now able to with the same stress load (Fig. 4 C). For example, when the local cortical tension of the active 10 μm cell was reduced by 50%, the cell could engulf a cell of an equivalent size, whereas previously, with wild-type cortical tension, it was unable to accomplish this.

This cortical tension consequence was seen most prominently in simulations containing larger passive cells. In many cases, cortical tension is proportional to viscoelasticity (57). To parse apart the role of cortical tension versus viscoelasticity, we reduced the viscoelasticity of cells independent of cortical tension (Fig. 4 D). In this scenario, the active cells engulfed the passive cells with less efficiency compared to the cells for which local cortical tension was reduced. This indicates that cortical tension resists efficient cell engulfment events more significantly than viscoelasticity.

Not all obstacles with which a cell encounters are perfectly spherical. Bacteria, for example, come in a diverse array of shapes. We therefore simulated an active cell interacting with a passive cylindrical-shaped cell to model an event that more accurately represents phagocytosis (Fig. 4 E). This simulation sets the foundation to expand the infrastructure to more cell types.

**Interactions with deformable bodies**

Although we successfully modeled how a cell encounters and engulfs solid objects of various sizes and shapes, we next moved to expand the biological range of our modeling framework. We incorporated deformability into the objects with which the active viscoelastic cells interacted. To this end, LSM potential functions were used for each cell. This allowed for independent modulation of the mechanics and stress production of each of the cells independently. To represent the interactions between deformable bodies accurately, we applied stresses to each cell that were proportional to the velocity of the interacting cell (Fig. 4 F). In this simulation, the engulfing cell is twofold more mechanically deformable than the cell being engulfed and is experiencing an active stress of 5 nN/μm² over the bottom 2 μm of non-adhered cell perimeter. This mechanical heterogeneity is often seen in successful cell-cannibalism events and is seen frequently in diseased states such as cancer (58).

With both cells having viscoelastic characteristics, we can investigate more biologically diverse and accurate cell-cell interactions such as entosis (59,60). We found that when both cells were deformable, both cells did deform in response to the stresses. With this ability to deform, less stress was needed to consume a viscoelastic cell compared with a solid cell of the same size.
Cellular deformations with a nucleus

In previous simulations, we assumed that the cytoplasm of the cell is a spatially homogeneous viscoelastic material. In practice, the internal compartments will greatly affect the ability to change cell shape (61). Thus, to increase the biological accuracy of the simulations, we included a deformable nucleus. To understand the effect of a deformable nucleus on cell shape, we modeled a cell settling including adhesion (threshold 1.7 nN/m$^2$). We see that with a stress of 8 nN/m$^2$ on the bottom third of the cell perimeter, the cell flattens, adhering to the surface leaving room for the nucleus in the center of the cell body (Fig. 5 A). Owing to its size, the nucleus is often the limiting factor to cellular deformation especially in confined environments (61). We found that when an active cell (radius 5 μm) includes a nucleus (radius 2 μm) and then engulfs a passive cell with a radius less than that of the active cell, the nucleus did not impact the time required to complete engulfment (cf. Fig. 4 A vs. 5 B).

We also considered how the presence of a stiff nucleus affected cell deformability. To this end, we simulated the movement of a cell (radius 5 μm) into a confined space
(2 \mu m in width) with a stress of 4 nN/\mu m^2 on the bottom over 3 \mu m of cell perimeter. In these simulations, the nucleus was required to deform within the cell to fit through the space (Fig. 5 C). When the width of the confinement was reduced to 1.5 \mu m, the nucleus was unable to deform sufficiently to fit through the confined space (Fig. 5 D). Depending on the cell type, the elasticity of the nucleus can vary anywhere from 0.5 to 6 times that of the cell (49,62,63). We simulated this situation by varying the nuclear mechanical stiffness, ranging from 0.25 \times to 4 \times wild-type elasticity (\sim 70 pN/\mu m^2). At the higher stiffness, the nucleus restricted the cell’s ability to move into the confinement (Fig. 5 D). This impact was readily observed by comparing the deformation of the nucleus at the 117 min time point. Compared to the nucleus with 25% wild-type elasticity, the cell with the 4 \times nucleus stiffness deformed considerably less. Finally, we altered the cortical tension of the cell body with a deformable nucleus (elasticity 4 \times wild-type). These cells experienced an outward normal stress of 8 nN/\mu m^2 over the bottom third of the cell perimeter (Fig. 5 E). As the cortical tension of the cell increased from 0.25 \times to 4 \times wild-type, we saw that the cell’s ability to deform is a function of cortical tension. When the cortical tension was low, the cell easily deformed into a confined environment. When we increased the cortical tension to 4 \times that of a wild-type cell, the cell was unable to deform into the same confined environment.

DISCUSSION

This work provides a foundation for testing a variety of cellular interactions using LSM. The results from these simulations are consistent with what has been seen experimentally. For example, experimentally we saw that engulfment events of small cells and particles completed on the order of minutes (64,65), whereas large cells completed on the order of hours (59). Additionally, we have explored the cell engulfment parameters giving insight into the robustness of cell engulfment. Testing the sensitivity of the system allowed us to understand better which mechanical parameters are most responsible for successful cell engulfment events. Using cell engulfment as a testbed, we expanded our modeling framework by including a deformable cell nucleus and exploring cells in confinement and can now begin to have predictive power over diverse biological systems.

One limitation to achieving this goal using the LSM is the computational load that would be required. As previously mentioned, the computational load scales proportionally to the number of cells being simulated. In its current implementation, it would be difficult to scale these simulations to model the number of cells that are seen in other modeling techniques such as vertex models or cellular Potts models. These techniques can model systems on the order of hundreds of cells, although they lose the geometric accuracy that is achieved with the LSM (66–69). One candidate
approach to address this limitation is to use a hybrid modeling strategy with increasing degrees of course grain-
ing as one moves away from the primary cells of interest.

Our simulations demonstrate one way in which adhesion between cells or cell and substrate can be implemented in the LSM framework. Using the phase-field method, which has many similarities to the LSM (see the review (5)), there have been several implementations of both multicellular interactions (6,7) as well as cells that incorporate a nucleus (11). For example, Shao et al. incorporate adhesions through an elastic force term that acts as a drag term on cell motion (6). These bonds stretch but can break with a probability that depends on their extension. A similar approach was used by Löber et al. (7,8), who tracked a local density of adhesive sites and make the protrusive strength depend on the number of active bonds. These authors also consider the interactions between two cells by introducing two terms into the phase-field equations: one that penalizes overlap between the phase-field functions of interacting cells and another that represents adhesion by advecting the phase-field function of one cell along the normal vector to the interface of another cell. To account for the presence of a nucleus inside a cell during divi-
sion, Zhao and Wang (11) incorporate two different auxiliary functions (similar to the way that we do here) and apply a steric interaction term to penalize the overlap between the cytoplasm and the nucleus. One apparent advantage of these approaches is the relative ease by which the interacting bodies can be considered. On the other hand, this is done by avoiding a sharp interface, which is one of the hallmarks of the LSM method.

The continuum models, such as the one presented here, provide an excellent platform for whole-cell simulations, although they omit details about the subcellular components. In this way, they represent a minimal model of cell shape change based on a deformable boundary. Using a mechanical equivalence circuit, all net forces and stresses are applied at this boundary, and the effect of the intracellular mechanics is incorporated through the viscoelastic model. Agent-based methods (70,71) provide an alternative approach whereby inter-
actions between specific components are considered. Owing to their computational cost, these modeling tech-
niques are presently limited for modeling at the cellular level except for the simplest cell types (72). Another simplification in our modeling approach is that for computational effi-
cy, the simulations have been done in two dimensions. One attractive feature of continuum interphase methods, such as the LSM or the phase-field approach, is that the computational steps are readily extended to deal with three-dimensional systems. This has been done in several situ-
tions when simulating other behaviors, such as cell divi-
sion (10–12) or migration (73,74), in which the degree of cellular deformation is similar to those considered here. In these cases, the differences between the corresponding two- and three-dimensional models are small.

Having a solid infrastructure in place to model cellular interactions is important as we consider the role of important multicellular interactions in the body. The body is a complex system of cells interacting in predictable ways. In tissue envi-
ronments, cells sense and respond to their surroundings and often change shape in response to these queues. This mechanosensation (the ability to sense and respond to me-
chanical stimuli) is an important part of a cell’s ability to survive and maintain homeostasis in a tissue environment. Our models have focused on the mechanical modulation and stress production of a primary cell driving the engulf-
ment of a second cell. Current literature shows that the en-
gulfed cell can play a significant active role in cell engulfment events, as seen in processes such as entosis (59). Moving forward, we plan to interrogate the role of the engulfed cell as a driver for cell-cell interactions.

Disease states often arise when a cell loses this ability to appropriately sense and adapt to its surroundings (75). For example, in cancer, changes in cellular deformability and altered mechanoresponsive protein expression lead to aberrant cellular morphologies (62). This results in mechanical heterogeneity between cells, often causing unconventional cellular interactions. The foundation of designing an in silico tissue environment starts with a single cell-cell interaction. With the ability to model diseased tissue sys-
tems using theory and in silico methods, we could have pre-
dictive power over the system, which would help guide our future experimental design.

SUPPORTING MATERIAL
Supporting material can be found online at https://doi.org/10.1016/j.bpj.2021.10.021.

AUTHOR CONTRIBUTIONS

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