

Intercellular signaling through secreted proteins induces free-energy gradient-directed cell movement

Nataly Kravchenko-Balasha^{a,1,2}, Young Shik Shin^{b,1,3,4}, Alex Sutherland^a, R. D. Levine^{b,c,d,4}, and James R. Heath^{a,4}

^aNanoSystems Biology Cancer Center, Division of Chemistry, California Institute of Technology, Pasadena, CA 91125; ^bDepartment of Molecular and Medical Pharmacology, David Geffen School of Medicine, University of California, Los Angeles, CA 90095; ^cDepartment of Chemistry and Biochemistry, University of California, Los Angeles, CA 90095; and ^dThe Institute of Chemistry, The Hebrew University of Jerusalem, Jerusalem 91904, Israel

Edited by Michael L. Klein, Temple University, Philadelphia, PA, and approved March 31, 2016 (received for review February 8, 2016)

Controlling cell migration is important in tissue engineering and medicine. Cell motility depends on factors such as nutrient concentration gradients and soluble factor signaling. In particular, cell–cell signaling can depend on cell–cell separation distance and can influence cellular arrangements in bulk cultures. Here, we seek a physical-based approach, which identifies a potential governed by cell–cell signaling that induces a directed cell–cell motion. A single-cell barcode chip (SCBC) was used to experimentally interrogate secreted proteins in hundreds of isolated glioblastoma brain cancer cell pairs and to monitor their relative motions over time. We used these trajectories to identify a range of cell–cell separation distances where the signaling was most stable. We then used a thermodynamics-motivated analysis of secreted protein levels to characterize free-energy changes for different cell–cell distances. We show that glioblastoma cell–cell movement can be described as Brownian motion biased by cell–cell potential. To demonstrate that the free-energy potential as determined by the signaling is the driver of motion, we inhibited two proteins most involved in maintaining the free-energy gradient. Following inhibition, cell pairs showed an essentially random Brownian motion, similar to the case for untreated, isolated single cells.

cell–cell force | cell motility | surprisal analysis | Langevin equation | Brownian dynamics

Changes in free energy define the direction for spontaneous changes in chemistry and physics. As examples, chemical gradients and electrical potential changes across membranes (1) may be viewed as chemical potentials that define a direction (2, 3). Other examples include chemotaxis (2–4) and active transport (1), both of which show that overcoming a concentration gradient requires work in the sense of expenditure of free energy. In this study, we aim to show that the thermodynamic analog of the free energy of an intercellular signaling system, mediated by secreted proteins, acts to determine the direction of change in cell–cell movement. Secreted proteins are a vehicle for cell–cell communication and signaling (5) and, once received by a cell, can initiate intracellular signaling cascades, resulting in changes in gene transcription, protein expression, and the activation of cellular functions. Such functions might include cell division, the secretion of a new group of proteins, or, as investigated here, cell motility (1).

Our experiment is a system of two interacting but otherwise isolated cells for which we measure both cell motion trajectories over a period of several hours and, at the terminal time point, the expression levels of a panel of secreted proteins. The experimental platform is the single-cell barcode chip (SCBC), which permits measurements of statistically significant numbers of cells (6–9).

Our information-theoretic analysis (10) of the experimental data regards the signaling proteins as species mediating the exchange of information between cells. This analysis is used to determine the changes with distance of the free energy of the cell–cell signaling and to show that the cell–cell relative motion can be described as a constrained Brownian motion. We show that the direction of change in cell movement is toward a more stable cellular arrangement where cell–cell signaling is balanced. Inhibiting that signaling results in a loss of the directed movement;

the cells move in a purely Brownian-type random walk, similar to the case of single isolated cells. In other words, we determine a cell–cell potential that characterizes the cell–cell motion in a way that is similar to other two-body interacting systems in physics and chemistry. We show that this potential is established through the exchange of secreted protein. By identifying those key signaling proteins, we can experimentally control the cell–cell motion.

Results

Our experimental design is guided by previous observations (10) that a pair of glioblastoma (GBM) cancer cells will exhibit a stable steady state that is characterized by a narrow range of intercellular separation distances. A hypothesis tested in this work is that one or more secreted proteins mediates that stable separation and directs cell migration toward that stable state. Thus, we sought to capture both relative motion trajectories for pairs of cells, as well as the levels of secreted proteins that might influence those trajectories. To determine an optimal panel of secreted proteins for monitoring single cells and cell pairs, we initially assayed for 12 secreted cytokines and growth factors from single and small populations of U87EGFR cells (Fig. S1). U87EGFR cells are model GBM cancer cells characterized by overexpression of the epidermal growth factor receptor (EGFR) (11). One protein served as a negative control, whereas the others were known to participate in cell–cell signaling [see

Significance

We demonstrate the role of free energy in determining the direction of motion in a biological process. A thermodynamics-motivated approach is used to show that free-energy changes in cell–cell signaling determine the force gradient for directed cell motion. Using isolated cell pairs and assays of secreted protein levels, we compute the free energy of the cell–cell signaling network as a function of cell–cell separation. The changes in free energy as a function of separation correspond to a potential-energy gradient that can influence cell–cell motion. Recordings of cell motion trajectories were compared with the direction of the gradient. Neutralizing the secreted proteins most involved in establishing the free-energy gradient cancels the directed motion.

Author contributions: N.K.-B., Y.S.S., R.D.L., and J.R.H. designed research; N.K.-B. performed theoretical research; Y.S.S. performed experimental research; N.K.-B., Y.S.S., A.S., R.D.L., and J.R.H. contributed new reagents/analytic tools; N.K.-B., Y.S.S., R.D.L., and J.R.H. analyzed data; and N.K.-B., Y.S.S., R.D.L., and J.R.H. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

¹N.K.-B. and Y.S.S. contributed equally to this work.

²Present address: Bio-Medical Sciences Department, The Faculty of Dental Medicine, The Hebrew University of Jerusalem, Jerusalem 9112001, Israel.

³Present address: NanoIVD, Inc., Redondo Beach, CA 90278.

⁴To whom correspondence may be addressed. Email: youngshik@gmail.com, rafi@fh.huji.ac.il, or heath@caltech.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1602171113/-DCSupplemental.

$\alpha = 1$, HGF and IL-6 dominate. Accordingly, we measured the cell motion trajectories for two cell microchambers under the influence of neutralizing antibodies to inhibit IL-6 and HGF signaling. Fig. 7 shows that, even after a delay of 6 h, there is hardly any directed motion, with zero-centered Gaussian distributions of slowly increasing widths with time, as expected for Brownian motion. The data clearly demonstrate that inhibiting unbalanced signaling hinders any directed motion but hardly affects the Brownian part. More generally, the data also illustrate that the information theory analysis of the proteomic data correctly identifies those signaling proteins that influence relative cell motion the most. It also shows that experimentally influencing the activity of those proteins provides an experimental handle for influencing cell organization and migration.

Discussion

Cell motion is an active biological process (20) that cannot be fully described by random Brownian motion (20, 21). The U87EGFR cancer cells investigated here are representative of the highly invasive disease of GBM, and so understanding the motility of such cells has both fundamental and practical value. We report on a search for a pairwise cell interaction potential that is governed by cell–cell signaling. The minimal point of this potential is the steady state of the cell–cell signaling. The potential induces cells to move toward that steady-state separation. We applied a thermodynamics-based theory to a dataset from hundreds of isolated cell pairs that integrated measurements of secreted protein levels with 6 h of cellular motion trajectories. Surprisal analysis of that dataset revealed the existence of a free-energy gradient, with an energy minimum at a cell separation of 200 μm . Analysis of the relative motion of the cell pairs revealed directed cell movement toward this steady-state separation, implying that, at this separation, cell–cell signaling is balanced and corresponds to the most stable state of the cell–cell potential. Away from this separation distance, the signaling is unbalanced, and cell motion is described by two tendencies: isotropic motility and directed movement due to the cell–cell potential gradient. Simulations of the directed motion using the high friction limit, the Langevin equation, and the potential derived from the signaling closely reproduced the experimental observations. We showed that the unbalanced cell–cell signaling is the cause of the directed motion by inhibiting those two proteins most responsible for the free-energy gradient. In summary, we show that soluble factor signaling between two cells can define a free-energy gradient, which, in turn, directs the relative cell motions. Experimental control over the levels of those soluble factors provides a handle for controlling cellular motion in a predictive fashion.

Materials and Methods

Surprisal Analysis. The analysis was carried out as described in some detail before (19, 25) and in *Supporting Information*. The particular application to pairs of cells has also been presented (10) for the purpose of determining the most stable steady-state separation in U87EGFRV8 and U87PTEN cell types. It was shown therein that this separation as determined for isolated cell pairs accounts for the most probable cell–cell distance in bulk cell cultures. This is similar to molecular liquids or crystals where the two-body force provides a realistic first approximation for the many-body energy.

The Langevin Equation. The equation is solved numerically by drawing random values between 1 and -1 for the random force, making a new drawing at each time step because the force is taken to be uncorrelated. To derive the interpretation in terms of relaxation time, we expand the cell–cell potential about the steady-state separation $r_{ss} = 200 \mu\text{m}$, namely $U(r) \cong U(r_{ss}) + k(r - r_{ss})^2/2$. Near r_{ss} the Langevin equation reads $d(r - r_{ss})/dt = -(k/\gamma)(r - r_{ss}) + \sqrt{2DR}(t)$, showing that for consistency of units γ/k has the dimension of “time” and that it has the interpretation of the relaxation time to the steady separation. The value for the relaxation time of 8 h is determined by the fit to the data for cells that are near r_{ss} .

The experimental results for the distribution of cell–cell separations at the earliest time after acclimation (Fig. S7A), and after an additional delays of 2 and 6 h (Fig. S7B), show that cells that were initially located at short cell–cell

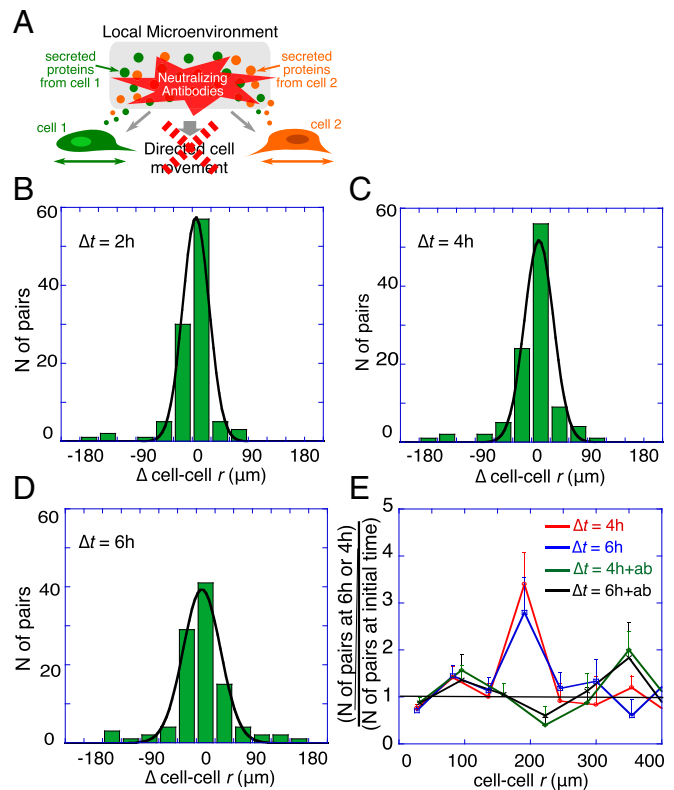


Fig. 7. Cell–cell motion after treatment with neutralizing antibodies against IL-6 and HGF. (A) Schematic illustration of the local microenvironmental condition. (B–E) Following the inhibition of signaling, the changes in the distribution of the cell–cell displacements ($\Delta\text{cell-cell } r$) after $\Delta t = 2 \text{ h}$ (B), $\Delta t = 4 \text{ h}$ (C), and $\Delta t = 6 \text{ h}$ (D). The results shown are for about 150 U87EGFR cell pairs that were initially separated by less than 200 μm . The measured displacements ($\Delta\text{cell-cell } r$) were binned into histograms. The histograms were fitted to Gaussian distribution ($R^2 > 0.95$). Even after a delay of 6 h, the histogram of the cell–cell distances could be well fitted by a Gaussian distribution ($R^2 = 0.97$). (E) The results shown in Fig. 3A and the corresponding results for a similar number of cell pairs treated with the neutralizing antibodies. For the antibody-treated cells (+ab), the probability of finding a cell pair at any distance range after 4 and 6 h was similar to the value at the initial time (green and black curves fluctuating about 1).

separation distances are not strongly attracted toward the steady-state separation of $r_{ss} = 200 \mu\text{m}$. However, the harmonic approximation for $U(r)$ introduced above, implies a very strong restoring force at short distances, which is inconsistent with the experimental results (Fig. S7) at small separations $r \ll r_{ss}$. This is also inconsistent with the results of surprisal analysis of the signaling as shown in Fig. 2B and Fig. S6. At short distances, the signaling is almost stable (Fig. 2B and Fig. S6). Therefore, for separations below 50 μm , we reduce the force constant k down to 10% of its value for $r = 200 \mu\text{m}$.

Cell Culture and Reagents. U87EGFR cells were kindly provided by Paul Michel’s Laboratory (University of California, San Diego, La Jolla, CA). Cells were routinely maintained in DMEM (American Type Culture Collection) containing 10% (vol/vol) FBS in a humidified 5% (vol/vol) CO_2 incubator at 37 $^\circ\text{C}$. See Tables S1 and S2 for lists of DNAs and antibodies used in the study. For neutralization assay, anti-IL-6 and anti-HGF antibodies were added to single-cell suspension at a final concentration of 0.5 $\mu\text{g}/\text{mL}$, respectively, before loading into the device.

SCBC Design and Fabrication. The SCBC is composed of a two-layer elastomer microfluidics layer bonded on top of a barcode-patterned glass slide. Details of microchip design and fabrication can be found in *Supporting Information*. In brief, molds for polydimethylsiloxane (PDMS) microfluidic device were fabricated by photolithography. These molds were used to mold PDMS elastomer for flow and control layers that form 2,640 microchambers after thermal bonding. For experiments, cells are randomly loaded to microchambers and allowed to acclimate for 2 h. The cell numbers and cell positions are recorded

