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Probing the coupled adhesion and deformation characteristics of suspension cells

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By combining optical trapping with fluorescence imaging, the adhesion and deformation characteristics of suspension cells were probed on single cell level. We found that, after 24 h of co-culturing, stable attachment between non-adherent K562 cells and polystyrene beads coated with fibronectin, collagen I, or G-actin can all be formed with an adhesion energy density in the range of 1–3 × 10^{-2} mJ/m², which is about one order of magnitude lower than the reported values for several adherent cells. In addition, it was observed that the formation of a stronger adhesion is accompanied with the appearance of a denser actin cell cortex, especially in the region close to the cell-bead interface, resulting in a significant increase in the apparent modulus of the cell. Findings here could be important for our understanding of why the aggregation of circulating cells, like that in leukostasis, takes place in vivo as well as how such clusters of non-adherent cells behave. The method proposed can also be useful in investigating adhesion and related phenomena for other cell types in the future. © 2014 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4893734]

Adhesion plays important roles in processes such as cell migration, differentiation, proliferation, and cell death. For this reason, intense effort has been spent to characterize the interactions between living cells and their microenvironment. In particular, techniques like atomic force microscopy (AFM), interference microscopy, micropipette manipulation, laser spallation, and shear flow assay have all been developed to probe cell-substrate or cell-cell adhesion. A common theme of these approaches is to induce cell detachment from the substrate (or other cells) with imposed disruptive “forces” (e.g., laser shock, shear flow, and direct pulling) from which the adhesion characteristics can be quantitatively extracted. However, due to the non-trivial geometry and deformability of cells, proper theoretical models are often needed to interpret data from such experiments. One popular choice to serve this purpose is the Johnson-Kendall-Roberts (JKR) theory, where the effect of adhesion is represented by a single parameter (adhesion energy density) and relationships describing the spontaneous attachment as well as enforced separation of two elastic spheres have all been obtained in simple forms. Actually, in addition to being used to analyze cellular attachment, the JKR approach has also been adopted in examining biological adhesion of vesicles, nanoparticles, and objects with wavy surfaces.

Despite these aforementioned efforts, several important issues remain to be further explored. For one thing, most existing experiments focus on adherent cells but very little has been reported regarding the adhesion response of suspension cells. Of course, by their nature, non-adherent cells usually cannot establish strong attachment with the outside. Nevertheless, under pathological conditions such as leukostasis circulating leukocytes can form aggregates, obstruct small vessels in organs, and eventually lead to serious problems like convulsion, stroke, visual, and auditory impairment. Evidently, a precise knowledge of the adhesion characteristics of these non-adherent cells will be important for our understanding of how such phenomenon takes place as well as for finding possible prevention/treatment strategies. In addition, although it is widely known that the formation of tight contact regions (often referred to as focal adhesions) between cells and the extracellular matrices (ECMs) requires the assembly of numerous proteins in the intracellular side, very few studies have explored the question of whether and how the appearance of adhesion alters the mechanical response of cells.

Aiming to address these issues, we developed a method to quantitatively measure the adhesion response of individual suspension cells by combined optical trap manipulation and fluorescence. Specifically, precisely controlled optical pulling was applied to ligand-coated polystyrene beads that are in adhesive contact with K562 (human chronic myelogenous leukemia) cells. Evolutions of the contact area and bead movement during the detaching process were recorded which, in conjunction with the JKR description, allow us to extract the adhesion energy density of the cell-bead interface as well as the cell modulus. Fluorescence imaging was also utilized to monitor possible adhesion-induced cytoskeletal changes.

Fibronectin and collagen I, two common adhesion proteins, are known to be important for the immobilization of living cells within tissues and organs, as well as facilitate cell-substrate adhesion in vitro. In addition, it is conceivable that attachment of cells to the surface deposited with actin, a molecule heavily involved in the formation of focal adhesions and capable of binding to various membrane-associated proteins and receptors, can take place. Based on these observations, we focus our attention on the adhesion between suspension K562 cells and beads coated with the three aforementioned ligands, respectively. Specifically, cells were co-cultured with 5 μm-poly styrene beads (Polysciences), deposited with fibronectin (Sigma), actin (Molecular Probes) or collagen I (Sigma), on a fibronectin-coated confocal dish.
for 24 h, see supplementary material for details. After that, optical trap (MMI) was used to grab individual microspheres, in adhesive contact with K562 cells, and then pull them away from the cells immobilized on the cover-slip as illustrated in Fig. 1(a). Videos were taken during the entire separation process to record the bead displacement in time, \( u(t) \), as well as changes in the contact radius \( a \). Since the motion of the laser, i.e., \( \delta(t) \) as shown in Fig. 1(a), is controlled in our experiment, the quantity \( \delta - u \) represents the relative displacement between the bead and the laser beam which, in return, leads to a trapping potential on the bead tending to minimize \( |\delta - u| \).

It has been shown that optical trap works approximately like a linear spring, so that a force

\[
F(t) = -k[\delta(t) - u(t)]
\]

will be exerted on the microsphere confined within the laser beam with the negative sign representing that this force is “tensile,” i.e., trying to separate the bead from the cell. Here, \( k \) is the effective stiffness of the trap whose value depends on factors like the laser power and bead size and hence must be calibrated beforehand, see supplementary material for the calibration protocol. In this study, a constant trapping stiffness of 39.7 pN/\( \mu \)m was used. With a method of estimating the pulling force at hand, we can then construct the \( F \) vs. \( a \) curve for the detaching process, as illustrated in Fig. 1(b), which is of key interest in the study of adhesive contact. To make sure changes in the contact area can be accurately monitored, special attention has been paid to identify microspheres adhering to the side (not to the upper or lower portion) of the cell, based on which all our tests were conducted.

In addition to optical manipulation, immunofluorescent imaging was also employed to monitor possible adhesion-induced cytoskeletal changes in K562 cells. In particular, cells were fixed with 3.7% paraformaldehyde, permeabilized with 0.1% Triton X-100, and then stained against F-actin with AF488-phalloidin (Invitrogen) prior to experiment. After co-culturing with ligand-coated microspheres, a fluorescence microscope (Nikon) was used to obtain images of cells in adhesion.

A representative sequence of micrographs of a K562 cell subjected to optical pulling is shown in Fig. 1(b). As the magnitude of \( F \) increases, deformation of the cell becomes more apparent and the contact area keeps shrinking gradually. Finally, detachment between the cell and the bead takes place abruptly after \( F \) reaches a critical level. Data from multiple independent tests for each case, i.e., beads coated with fibronectin (\( n = 10 \)), collagen I (\( n = 12 \)), actin (\( n = 13 \)), or without coating (\( n = 8 \)), are gathered in Fig. 2(a). Compared to the case of non-coating beads, the presence of fibronectin or collagen I noticeably elevates the maximum pulling force (from \( \sim 90 \) pN to \( \sim 130 \) pN) that the adhesion can sustain. Interestingly, coating the bead surface with actin further increases the pull-off force to the range of \( \sim 250 \)–\( 300 \) pN. Recall that, according to the well-known JKR theory, the enforced separation between two elastic spheres in adhesive contact can be described by

\[
F = \frac{4E_r a^3}{3R} + \frac{8\pi\gamma a^3}{R^2},
\]

where \( a \) and \( F \), as defined before, are the contact radius and applied force; \( \gamma \) is the adhesion energy density representing the energy reduction per unit area when two surfaces are brought together; and \( 1/R = 1/R_1 + 1/R_2 \) where \( R_1 \) (\( \approx 10 \) \( \mu \)m) and \( R_2 \) (\( = 2.5 \) \( \mu \)m) are the radii of the cell and the bead (Fig. 1(a)). Since the microsphere is much stiffer than the cell, the so-called reduced cell modulus \( E_r \) takes the form \( E_r = \frac{E_r}{1-v_1} \), with \( E_1 \) and \( v_1 (\approx 0.5) \) being the Young’s modulus and Poisson’s ratio of the cell, respectively.

The generic shape of the \( F \) vs. \( a \) curve, according to Eq. (2), is schematically shown in Fig. 1(b) by the solid line where the critical pulling force for triggering sudden detachment is predicted to be \( F_c = -\frac{2\gamma n R}{3} \), with a corresponding contact radius of \( a_c = \left[ \frac{3\gamma n R}{16\pi\gamma} \right]^{1/3} \). As illustrated in Fig. 2(a), our data can be well explained by Eq. (2) if one chooses \( \gamma = 0.03 \) mJ/m\(^2\) and \( E_r = 100 \) Pa for actin coating.
Adopting the same $\gamma$ and $E_r$ values as those in Fig. 2(a), predictions from Eq. (3) are shown by the solid lines in Fig. 2(b) which, evidently, are in good agreement with experimental observations. We realize that the best way to visualize the comparison between our results and predictions from the JKR model is to plot $F/F_c$ against $a/a_c$, with $F_c$ and $a_c$ calculated from the corresponding $\gamma$ and $E_r$ values for each coating condition. Note that, based on the definitions of $F_c$ and $a_c$, Eq. (2) can be rewritten in the normalized form as

$$u = \sqrt{\frac{\pi \gamma}{2E_r}} \left(2\sqrt{a} + \sqrt{a_0}\right) - \frac{a^2}{R}. \quad (3)$$

Indeed, all data collapse into a single master curve in the $F/F_c$ vs. $a/a_c$ plot (Fig. 2(c)), as predicted by Eq. (4), further demonstrating that, irrespective of the types of ligands being used, the adhesion response of these suspension cells can always be well described by the JKR theory.

One thing must be pointed out is that the results obtained from our experiments are rather consistent with each other. For example, by fitting Eq. (2) to data from each pulling test, the bar graphs of extracted $\gamma$ and $E_r$ are shown in Figs. 3(a) and 3(b) which demonstrate that variation in the values of these two parameters is rather small. Interestingly, results here also suggest that a stronger adhesion is accompanied with a larger cell modulus. In particular, according to Fig. 3(b), K562 cells in contact with actin-coated beads appear to possess a modulus that is $\sim$60% higher than that exhibited by cells adhering to microspheres without coating. To confirm this finding, we have conducted additional AFM rate-jump indentations28,29 on K562 cells (20 independent tests for each coating), as detailed in the supplementary material, where similar trend was also observed (Fig. 3(b)).

It is conceivable that the formation of adhesion can trigger protein assembly in the cytoskeleton, strengthen the cell body, and eventually lead to an elevated apparent modulus. To test this hypothesis, the density of F-actin, a major cytoskeletal component, in K562 cells was monitored by fluorescence imaging. As expected, compared to the control group (i.e., cells in contact with microspheres without coating), a much denser actin cell cortex, especially in the region close to the cell-bead interface, was observed in cells attaching to actin-coated beads (Fig. 3(c) and Fig. S2 in the supplementary material).24 We further quantify these results by calculating the average fluorescent intensity in the 500 nm-thick layer underneath the membrane-bead interface as well as plotting the intensity distribution along the path starting at the center of the interface and moving into the cell, as indicated by the red dashed loop and white dashed arrow in Fig. 3(c). Clearly, results here show that coating the bead with fibronectin or collagen I leads to a $\sim$50% increase in the F-actin intensity while the deposition of G-actin roughly doubles that. These evidences support the notation that the formation of adhesion is coupled with the local reinforcement of F-actin in the cytoskeleton, resulting in a higher resistance of the cell against deformation.
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In this study, we have developed a method to quantitatively measure the adhesion and deformation of suspension cells on single cell level. Specifically, we found that adhesion between K562 cells and polystyrene beads coated with (or without) common cell-binding ligands can be formed with an energy reduction in the range of $1 - 3 \times 10^{-2} \text{mJ/m}^2$, a value that is comparable to those found for mutants of Dictyostelium or some pure bilayer systems. In comparison, this energy for several adherent cells, including T-lymphocytes and murine sarcoma S180 cells, has been reported to be of the order of $10^{-1} \text{mJ/m}^2$. The relatively weak adhesion capability of K562 cells is not that surprising given their non-adherent nature, that is, these cells normally do not adhere to each other or to the outside. Nevertheless, it is well-known that conditions such as leukostasis can trigger the aggregation and clumping of circulating blood cells and lead to serious medical issues. As such, data obtained here should help us understand how phenomena like this take place as well as identify possible prevention/treatment strategies. Our results also demonstrated that, compared to fibronectin or collagen I, coating the surface with G-actin will lead to stronger cell attachment, a finding that surprisingly has hardly been reported. Identification of the molecular mechanism behind is beyond the scope of current study and is certainly an issue that warrants further investigation.

Most existing studies treated the adhesion energy density and cell modulus as independent quantities. However, observations here clearly suggest that these two parameters are coupled for the investigated system. In particular, cells adhering to actin-coated beads appear to have a modulus that is significantly ($\sim 60\%$) higher than those bound to microspheres without coating (Fig. 3(b)). This finding could be important in the modeling of adhesion mediated processes, as well as interpreting experimental data, in the future. For example, according to the present study, cell aggregates can become much stiffer than individual cells, a feature that may have great implications in analyzing the deformation or migration behavior of cell clusters. The approach proposed here may also be of immediate use in examining issues like how membrane fluctuations or stochastic transitions in protein binding/unbinding affect weak cell-cell or cell-substrate interactions, given that forces in the range of piconewton can be precisely applied in the current setup.

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24 See supplementary material at http://dx.doi.org/10.1063/1.4893734 for additional information on bead coating and cell culture; optical tweezers calibration; rate-jump indentation; and fluorescent images of K562 cells in adhesive contact.