High Refractive Index Silicone Gels for Simultaneous Total Internal Reflection Fluorescence and Traction Force Microscopy of Adherent Cells

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Abstract

Substrate rigidity profoundly impacts cellular behaviors such as migration, gene expression, and cell fate. Total Internal Reflection Fluorescence (TIRF) microscopy enables selective visualization of the dynamics of substrate adhesions, vesicle trafficking, and biochemical signaling at the cell-substrate interface. Here we apply high-refractive-index silicone gels to perform TIRF microscopy on substrates with a wide range of physiological elastic moduli and simultaneously measure traction forces exerted by cells on the substrate.

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Introduction

Animal tissues exhibit a broad range of stiffnesses, from <1 kPa in brain to ~10 GPa in bone. When grown on a substrate, animal cells sense its rigidity, especially in a range corresponding to soft tissues, with elastic moduli, $E$, of 0.1–100 kPa[1,2,3]. Recent studies have emphasized the importance of variations of the rigidity in development[4,5,6,7], tumorigenesis[3,8,9], and cell migration[10,11]. Substrate rigidity sensing is mediated by cellular adhesion structures that exert traction forces on the substrate, and selective visualization of these adhesion structures is key to understanding rigidity sensing. If the elastic modulus of the substrate is sufficiently low, substrate deformations caused by the traction forces can be measured under a microscope and the cell traction forces can be reconstructed. The spatial patterns and dynamics of the traction forces provide important information on cytoskeletal tensions and the mechanisms of cell spreading, migration, and polarization[1,12]. Because substrate deformation in a given area often results from traction forces applied at multiple adhesion points, the conversion of a map of substrate deformation into a cell traction force map is complicated, especially when the locations of the adhesion points are not known[13]. Adhesion points can be detected by employing molecular markers that are known to be recruited to cellular adhesion structures using wide-field or confocal fluorescence microscopy, but identification of adhesion points exerting traction forces can be challenging. In addition, the accurate assessment of the adhesion area, especially the detection of small adhesion points, can be difficult with these two types of microscopy because of their relatively high background level.

The level of fluorescence background is substantially lower in total internal reflection fluorescence (TIRF) microscopy[14], which selectively visualizes fluorescent molecules in a 100–200 nm thick layer above the substrate and is the method of choice to image the cell-to-substrate adhesion structures[15] and to study molecular trafficking events at the plasma membrane[16]. In TIRF microscopy with a popular through-the-lens illumination technique[14], the fluorescence excitation beam is directed through a marginal area of a high numerical aperture (high-NA) oil immersion objective lens. As a result, the excitation beam enters the boundary between the microscope cover glass and the medium behind it at a large angle of incidence, $\theta_1$ (as measured from the optical axis; Fig. 1A). If the refractive index of the medium, $n_2$, is lower than that of the cover glass, $n_1 \approx 1.52$, and $\theta_1$ is greater than a critical angle, $\theta_c = \arcsin(n_2/n_1)$, total internal reflection occurs at the boundary between the glass and the medium (Fig. 1A). The incident excitation beam is completely reflected off the boundary and only penetrates behind it as an evanescent wave that decays with the distance from the boundary, $z$, as $e^{-z/d}$, where the characteristic depth, $d$, is given by $d = \lambda/(4\pi n_1^2 \sin^2 \theta_1 - n_2^2)$, with $\lambda$ being the wavelength of the illumination beam in vacuum [14]. (We note that $\theta_1 = \theta_c$ corresponds to $n_1^2 \sin^2 \theta_1 - n_2^2 = 0$ and $d = \infty$.)

For a given microscope objective, there is a one-to-one correspondence between the distance of a ray from the optical axis in the plane of the objective back aperture and the resulting
angle of incidence, \( \theta_1 \) (Fig. 1). The maximal \( \theta_1 \) available with an objective lens is defined by the NA of the lens, \( \theta_{\text{max}} = \arcsin (NA/n) \). Live cells have refractive indices of up to \( n_c = 1.38 \) [14,17], and for total internal reflection at the boundary between the cover glass and a cell plated on it, as required for TIRF microscopy of the cell, the condition \( \theta_1 > \arcsin (n_c/n) \) must be met (Fig. 1A). Regular oil-immersion objectives have NA up to 1.40, making them not well suited for cell TIRF microscopy, because the requirements \( \theta_1 < \arcsin (NA/n) \) and \( \theta_1 > \arcsin (n_c/n) \) limit the range of \( \theta_1 \) to <2° and to satisfy these requirements for all its rays, the excitation beam needs to be very narrow and collimated. In addition, \( \theta_1 \) barely above \( \theta_0 \) results in extended depth of the evanescent wave, \( d \) (>170 nm for NA = 1.40 and \( \lambda = 500 \) nm) and the interaction of the evanescent wave with dense cellular organelles may lead to its conversion into scattered propagating light [14]. Therefore, all major manufacturers of biological microscopes introduced specialized objectives for TIRF with NA ranging from 1.45 to 1.49, greatly expanding the available range of \( \theta_1 \) (to 7°–13°), facilitating the introduction of the excitation beam, and generally improving the intensity and uniformity of illumination. Moreover, the capacities to vary \( \theta_1 \) and make it substantially greater than \( \theta_0 \) provided by specialized TIRF objectives enable adjusting \( d \) and making it particularly small (down to ~70 nm with NA = 1.49, \( n_2 = 1.38 \), and \( \lambda = 500 \) nm) [16].

For TIRF microscopy of cells plated on a soft substrate (Fig. 1B), the refractive index of the substrate must be greater than \( n_c \). The most commonly used cell substrates that have the rigidity of soft tissue and enable traction force measurements, polyacrylamide gels [19], have a refractive index close to that of water (\( n_w = 1.33 \)), making them unsuitable for TIRF microscopy. Silicone gels made of polydimethylsiloxane (PDMS), which have a long history of use in traction force microscopy [20,21,22,23], have a refractive index of \( \sim 1.41 \). Recently, TIRF microscopy of social amoebas (D. discoideum) on thin layers of silicone gels with a refractive index \( n_g = 1.40 \) has been reported [24,25]. In this configuration (cells on top of a gel on top of a cover glass; Fig. 1B), to reach the interface between the gel and a cell, the fluorescence excitation beam first needs to cross the interface between the cover glass and the gel, imposing a condition \( \theta_1 < \theta_c = \arcsin (n_g/n) \) [while the condition \( \theta_1 > \arcsin (n_c/n) \) remains unchanged for TIRF at the gel-cell interface]. Rays with \( \theta_1 \) greater than the critical angle \( \theta_c \) for the glass-PDMS interface will suffer a total internal reflection at this interface and not enter the gel, thus contributing to the reflection background rather than TIRF signal. Therefore, whereas the nominal NA of the objective used in the TIRF setup may be high (e.g., 1.45 as in [24,25]), its effective NA is limited to the refractive index of the gel, \( n_g = 1.40 \) [when \( \theta_1 < \theta_c = \arcsin (n_g/n) \) is more restrictive than \( \theta_1 < \theta_{\text{max}} = \arcsin (NA/n) \)], leading to problems and limitations in the TIRF microscopy of cells similar to those before the introduction of specialized high-NA TIRF objectives.

Here we used cover glasses coated with thin layers of silicone gels with a refractive index \( n_g = 1.49 \) (Fig. 1B) to perform TIRF microscopy of cells on soft substrates with a specialized TIRF objective (Nikon 60x/1.49), taking full advantage of its high NA of 1.49. We formulated these high refractive index (HRI) gels with elastic moduli, \( E \), of 0.4–130 kPa, covering nearly the entire physiological range, and tested their mechanical properties with a custom-built microfluidic device [26]. We plated human umbilical venous endothelial cells (HUVECs; by Lonza, Basel, Switzerland) on the gels, performed TIRF microscopy on them, and combined it with traction force microscopy by tracking fluorescent beads on surfaces of the gels.

**Materials and Methods**

To prepare the gel layers, the components of the HRI gel pre-polymers, parts A and B of QGel 920 and parts A and B of QGel 903 (both by Quantum Silicones LLC, Richmond VA; refractive index of 1.49 when cured), were mixed in various proportions (Table 1) and coated onto 25 mm no. 1 round cover glasses using a home-built spin-coater rotating at 1920 rpm. Each cover glass was baked at 100°C for 2 hr to create a layer of cured gel on it with a thickness, \( \xi \approx 34 \) μm. After baking, the gels on the cover glasses were treated with 3-aminopropyl trimethoxysilane for 5 minutes and incubated for 10 minutes at room temperature under a suspension of 40 nm carboxylated far-red fluorescent beads (excitation/emission 690/720 nm, by Invitrogen, Carlsbad, CA) in a 100 μg/ml solution of 1-Ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) in water to covalently link beads to the gel surface. This technique made it possible to have all beads in one plane corresponding to the surface of the gel. Therefore, the beads could be imaged under wide-field (epi-fluorescence illumination) with minimal background and their displacements reflected the deformation of the very top of the substrate. To promote cell adhesion, fibronectin (FN) was covalently linked to the gel surface by incubation in 50 μg/ml of FN with 100 μg/ml EDC in PBS, pH 7.4 for 30 min at room temperature.

The elastic modulus (Young’s modulus), \( E \), of the gels was evaluated by applying a known hydrodynamic shear stress, \( \tau \), to the gel surface using a custom-built microfluidic device, measuring the resulting bead displacement, \( \Delta x \), calculating the shear of the
gel, $\gamma = \Delta x/\xi$, and applying the equation $E = 2(1 + \nu)\tau/\gamma$, where $\nu$ is the Poisson ratio, as explained in detail elsewhere [26]. Because the Poisson ratio of silicone gels is nearly equal to 0.5 [27], the equation was reduced to $E = 3\tau/\gamma$. To measure the gel thickness, $j$, a small amount of the 40 nm far-red fluorescent beads was deposited on the cover glass surface before it was coated with the gel pre-polymer. The fluorescence microscope was first focused on beads on the glass surface and then on those on the gel surface and the difference in the readings of the nosepiece ($z$-axis) knob was recorded (with a correction for the mismatch between the refractive indices of the gel and immersion liquid), resulting in $\sim 1 \mu m$ accuracy. The shear, $\gamma$, was found to be a zero-crossing linear function of $\tau$ for $\Delta x$ of up to at least 3 $\mu m$ (greater than $\Delta x$ produced by HUVECs; see below) for all gels, with no sign of plastic deformations (see also [26]), thus validating the use of the equation $E = 3\tau/\gamma$, which applies to linear materials. Furthermore, measurements of $\Delta x$ vs. $\xi$ at different constant values of $\tau$ resulted in linear dependencies, indicating homogeneity of mechanical properties of the gel layers.

### Results and Discussion

TIRF and regular epi-fluorescence microscopy were performed on a Nikon Eclipse Ti-E microscope with a 60x/1.49 TIRF objective. To test the TIRF imaging, a suspension of fluorescent beads was pipetted onto a gel-coated cover glass. A TIRF movie (Movies S1 and S2) showed beads randomly appearing and disappearing in various locations without going out of focus, as expected for beads diffusing in and out of the thin layer above the substrate, which is illuminated under TIRF microscopy. The TIRF background of a gel-coated cover glass (with no beads) was nearly as low as the background of a blank cover glass (Fig. S1).

For live cell microscopy, HUVECs expressing lifeact-eGFP [28] were plated on the FN-coated gel substrates and incubated at 37$^\circ$C in 5% CO$_2$ for 1 hour. In early experiments, we observed toxic effects of freshly prepared gel substrates, which were likely due to low-molecular organic compounds present in the gels. The toxic effects disappeared, after the protocol was changed, and...
substrates were soaked in a buffer for ~1 hour prior to the cell plating. A likely factor facilitating rapid elution of harmful compounds from the gels was their relatively small thickness. HUVECs were cultured on the gels for up to 3 days.

An epi-fluorescence image of a typical cell (Fig. 2A) showed a developed F-actin network, whereas in a TIRF image of the same cell, only separate fluorescent patches could be seen, corresponding to F-actin in the substrate adhesion points (Fig. 2B). Far-red fluorescent beads deposited on the substrate were imaged under epi-fluorescence illumination (Fig. 2C), first, when the cell was intact and later after it was lysed by the application of 3% Triton X-100. We note that in spite of the high surface density (6 ± 1.7 per a 3 × 3 μm square), beads only protrude by 40 nm above the gel surface, occupy only ~0.1% of the surface area, and are thus expected to have a minimal effect on cellular adhesion (less than the imprinted micro-ridge patterns proposed before [22]).

A vector map of the bead displacement due to cell traction was constructed by calculating differences between the bead positions in the first and second image using a code in Matlab (Fig. 2D) [29,30]. The maximal bead displacement was 0.15 μm, whereas the noise level calculated as a root-mean-square of bead displacement in an area far away from the cell was 0.015 μm. A detailed reconstruction of traction forces from the substrate deformation is mathematically involved, and multiple alternative numerical protocols have been suggested [13]. Here, we used an implementation of the Boundary Element Method (BEM) [31] to convert the bead displacement map in Fig. 2D into a traction force map (Fig. 2E). The conversion was facilitated by relatively high surface density and even distribution of the tracking fluorescent beads (Fig. 2C). In addition, the placement of the beads on the surface of the gel, rather than in the bulk, as often practiced with polyacrylamide gels [32], reduced the fluorescence background and eliminated the uncertainty regarding the vertical position of the tracer particles, resulting in more reliable conversion of displacement into stresses. (See the Supporting Information, Appendix S1, for further details on how the maps in Fig. 2D and 2E were constructed.) Superposition of a contour plot of traction stress magnitude with the TIRF image of the cell (Fig. 2F) showed many F-actin localization points in regions with high traction stresses.

The experiment was repeated with gel substrates with $E$ of 0.4, 18, and 130 kPa (Fig. 3), which were prepared by mixing the components of QGel 920 and QGel 903 (that is harder than QGel 920) in different proportions (Table 1). TIRF microscopy of 4–6 cells was successfully performed on each substrate. Characteristic deformations were not visibly different for the 3.7, 18 and 130 kPa substrates, whereas characteristic traction stresses greatly increased with $E$ of the substrate, in general agreement with a previous report [33]. Deformations of the gel with $E=0.4$ kPa were substantially greater than those of the other gels (Fig. 3F vs. Figs. 3A and 3B). In addition, in contrast to cells on gels with higher $E$, cells on the gel with $E=0.4$ kPa had substantially less actin bundles, more rounded shapes, and many F-actin localization points at the substrate (as seen with TIRF) in the central area of the cell (Fig. 3E vs. Figs. 3A, 3B and 2).

To summarize, we formulated and characterized a series of silicone gels with elastic moduli covering nearly the entire physiological range and with a refractive index of 1.49 that enables taking full advantage of the high NA oil-immersion TIRF objectives for performing TIRF microscopy of adherent mammalian cells. TIRF microscopy is expected to reduce phototoxicity, allowing improved time resolution, and because of the low background level, better resolve small adhesion points, potentially enabling the detection of single fluorescent proteins. The combination of TIRF microscopy with traction force microscopy that we demonstrated will enable studies on the contributions of various components of cell adhesion machinery to generation of substrate traction forces and on the connection between these forces and membrane trafficking events. TIRF imaging of cells grown on the silicone gels can also make it possible to relate variations in membrane trafficking events and molecular composition of adhesion points to changes in the rigidity of the substrate in a range corresponding to a variety of living tissues and pathological environments.

Supporting Information

Figure S1 TIRF background of a gel coated coverslip vs. blank coverslip. (a) and (b) TIRF images of the surface of a blank coverslip and a coverslip coated with high-refractive index silicone gel, respectively, with 300 × 300 pixels in each image. The images were taken at identical illumination and acquisition conditions using a 100 × /1.46 Olympus TIRF objective and a cooled Hamamatsu camera, with a 1 sec exposure time and a maximal gain. The images are not completely dark because of a combination of the read-out noise and dark current of the camera.
(amplified by the gain), the incomplete blockage of the excitation light by the fluorescence filters (combined with the reflection and scattering of light from the glass and gel surfaces), and autofluorescence of the coverslip and gel. Small dots seen on the surface of the gel are possibly due to scattering of light at small defects in the gel and residual autofluorescence of microparticles stuck to the gel surface during its preparation. (c) Histograms of the pixel values of the images in (a) and (b) are shown by red and blue dots, respectively. The mean values of the pixels were 59.5 and 64.7 for the blank and gel-coated coverslip, respectively. Therefore, the TIRF background of the gel-coated coverslip was <9% higher than that of the blank coverslip.

Appendix S1 Reconstruction of the traction force map from the bead displacement map.

Movie S1 Real-time movie of a high refractive index gel substrate with a suspension of 100 nm red fluorescent beads (excitation/emission 580/605 fluorescent spheres by Invitrogen, Carlsbad, CA) above the gel taken with TIRF microscopy using a 60x/1.49 TIRF objective. Bright objects are either steady dots, which are beads stuck to the gel surface, or less bright dots that disappear immediately after appearing (blinking), as expected for beads entering and exiting (due to their Brownian motion) an ~100 nm deep region of evanescent illumination above the gel.

(MOV)

Movie S2 Real-time movie of the high refractive index gel substrate with the suspension of the 100 nm red fluorescent beads above the gel (same as in Supporting movie S1) taken under wide-field fluorescence (epi-fluorescence) illumination using the 60x/1.49 TIRF objective. In addition to steady bright dots, which are beads stuck to the gel surface, one can see two slowly moving large bright objects, which are likely aggregates of fluorescent beads, and a large number of relatively bright dots that remain visible for extended time intervals, as expected when Brownian motion of 100 nm particles is visualized under wide-field fluorescence illumination.

(MOV)

Author Contributions
Conceived and designed the experiments: EG ET AG. Performed the experiments: EG ET. Analyzed the data: EG ET AB GD AG. Contributed reagents/materials/analysis tools: AB PS KL GD MHG. Wrote the paper: EG ET MHG AG.

References
1. Nemir S, West JL (2009) Synthetic materials in the study of cell response to substrate rigidity. Ann Biomed Eng 38: 2–20.
2. Brown AE, Disher DE (2009) Conformational changes and signaling in cell and matrix physics. Curr Biol 19: R701–709.
3. Paznek MJ, Zahir N, Johnson KR, Lakini JN, Rosenzweig GI, et al. (2005) Tensile homeostasis and the malignant phenotype. Cancer Cell 8: 241–254.
4. Inger DE, Folkman J (1989) Mechanochanical switching between growth and differentiation during fibroblast growth factor-stimulated angiogenesis in vitro: role of extracellular matrix. J Cell Biol 109: 317–330.
5. Reilly GC, Engler AJ (2010) Intrinsin extracellular matrix properties regulate stem cell differentiation. J Biomech 43: 55–62.
6. Engler AJ, Sen S, Sweeney HL, Disher DE (2006) Matrix elasticity directs stem cell lineage specification. Cell 126: 677–689.
7. Disher DE, Mooney DJ, Zandstra PW (2009) Growth factors, matrices, and forces combine and control stem cells. Science 324: 1673–1677.
8. Zaman MH, Trapani LM, Seminski AL, Mackellar D, Gong H, et al. (2006) Migration of tumor cells in 3D matrices is governed by matrix stiffness along with cell-matrix adhesion and proteolysis. Proc Natl Acad Sci U S A 103: 10891–10894.
9. Kruzda D, Schmid H, Eismann JL, Lother U, Adler G, et al. (2010) Substrate stiffness and the receptor-type tyrosine-protein phosphatase alpha regulate spreading of colon cancer cells through cytoskeletal contractility. Oncogene 29: 2724–2738.
10. Isenburg BC, Dinnilla PA, Walker M, Kim S, Wong JY (2009) Vascular smooth muscle cell durotaxis depends on substrate stiffness gradient strength. Biophys J 97: 1313–1322.
11. Dubin-Thaler BJ, Hofman JM, Cai Y, Xie H, Spielman I, et al. (2008) Quantification of cell edge velocities and traction forces reveals distinct motility modules during cell spreading. PLoS One 3: e3735.
12. Guo WH, Frey MT, Burnham NA, Wang YL (2006) Substrate rigidity regulates the formation and maintenance of tissues. Biophys J 90: 2213–2220.
13. Sahass B, Gardel ML, Waterman CM, Schwarz US (2008) High resolution traction force microscopy based on experimental and computational advances. Biophys J 94: 207–220.
14. Axelrod D (2001) Total internal reflection fluorescence microscopy in cell biology. Traffic 2: 764–774.
15. Hu K, Ji L, Applegate KT, Danuser G, Waterman-Storer CM (2007) Differential transmission of actin motion within focal adhesions. Science 315: 111–115.
16. Pelkmans L, Zerial M (2005) Kinase-regulated quantal assemblies and kinesin-recycling of caveolae. Nature 436: 128–133.
17. Reichert WM, Truskey GA (1990) Total Internal-Reflection Fluorescence (Tirf) Microscopy. I. Modeling Cell Contact Region Fluorescence. Journal Of Cell Science 96: 219–230.
18. Sundd P, Gutierrez E, Pospieszalka MK, Zhang H, Grosman A, et al. (2010) Quantitative dynamic footprinting microscopy reveals mechanisms of neutrophil rolling. Nature Meth 7: 821–824.
19. Pelham RJ, Wang YL (1997) Cell locomotion and focal adhesions are regulated by substrate flexibility. Proceedings Of The National Academy Of Sciences Of The United States Of America 94: 13661–13665.
20. Harris AK, Stopak D, Wild P (1981) Fibroblast Traction As A Mechanism For Collagen Morphogenesis. Nature 290: 249–251.
21. Burton K, Taylor DL (1997) Traction forces of cytokinesis measured with optically modified elastic substrate. Nature 385: 450–454.
22. Balaban NQ, Schwarz US, Riveline D, Goichberg P, Tzur G, et al. (2001) Force and focal adhesion assembly: a close relationship studied using elastic micropatterned substrates. Nat Cell Biol 3: 466–472.
23. Schwarz US, Balaban NQ, Riveline D, Berchadsky A, Geiger B, et al. (2002) Calculation of forces at focal adhesions from elastic substrate data: The effect of localized force and the need for regularization. Biophysical Journal 83: 1308–1394.
24. Iwadate Y, Yamura S (2008) Actin-based propulsive forces and myosin-II-based contractile forces in migrating Dictyostelium cells. Journal Of Cell Science 121: 1314–1324.
25. Iwadate Y, Yamura S (2008) Molecular dynamics and forces of a motile cell simultaneously visualized by TIRF and force microscopes. Biotechniques 44: 739–44.
26. Gutierrez E, Grosman A (2011) Measurements of elastic moduli of silicone gel substrates with a microrheological device. Submitted for publication.
27. Mark JE (1999) Polymer Data Handbook. Oxford: Oxford University Press.
28. Riedl J, Crevenna AH, Kessenbrock K, Yu JH, Neukirchen D, et al. (2008) Lifeact: a versatile marker to visualize F-actin. Biophysical Journal 83: 150–167.
29. Sundd P, Gutierrez E, Pospieszalka MK, Zhang H, Grosman A, et al. (2010) Quantitative dynamic footprinting microscopy reveals mechanisms of neutrophil rolling. Nature Meth 7: 821–824.
30. Pelham RJ, Wang YL (1997) Cell locomotion and focal adhesions are regulated by substrate flexibility. Proceedings Of The National Academy Of Sciences Of The United States Of America 94: 13661–13665.
31. Harris AK, Stopak D, Wild P (1981) Fibroblast Traction As A Mechanism For Collagen Morphogenesis. Nature 290: 249–251.
32. Burton K, Taylor DL (1997) Traction forces of cytokinesis measured with optically modified elastic substrate. Nature 385: 450–454.
33. Schwarz US, Balaban NQ, Riveline D, Berchadsky A, Geiger B, et al. (2002) Calculation of forces at focal adhesions from elastic substrate data: The effect of localized force and the need for regularization. Biophysical Journal 83: 1308–1394.
34. Iwadate Y, Yamura S (2008) Actin-based propulsive forces and myosin-II-based contractile forces in migrating Dictyostelium cells. Journal Of Cell Science 121: 1314–1324.
35. Iwadate Y, Yamura S (2008) Molecular dynamics and forces of a motile cell simultaneously visualized by TIRF and force microscopes. Biotechniques 44: 739–44.
36. Gutierrez E, Grosman A (2011) Measurements of elastic moduli of silicone gel substrates with a microrheological device. Submitted for publication.
37. Mark JE (1999) Polymer Data Handbook. Oxford: Oxford University Press.
38. Stricker J, Sabass B, Schwarz US, Gardel ML, et al. (2007) Lifact: a versatile marker to visualize F-actin. Nat Methods 5: 605–607.
39. Feral GC, Zijlstra A, Tkachenko E, Prager G, Gardel ML, et al. (2007) CD98hc (SLCUA2) participates in fibroactin matrix assembly by mediating integrin signaling. J Cell Biol 176: 701–714.
40. J Li, Danuser G (2005) Tracking quasi-stationary flow of weak fluorescent signals by adaptive multi-frame correlation. Journal Of Microscopy-Oxford 219–220.
41. Stricker J, Sahass B, Schwarz US, Gardel ML (2010) Optimization of traction force microscopy for micron-sized focal adhesions. Journal Of Physics-Condensed Matter 22.
42. Saiz A, Bugain A, Silberzan P, Ladoux B (2005) Is the mechanical activity of epithelial cells controlled by deformations or forces? Biophys J 89: L52–54.