

Quantifying cellular forces and biomechanical properties by correlative micropillar traction force and Brillouin microscopy

Stefano Coppola,^{1,4} Thomas Schmidt,¹ Giancarlo Ruocco,² and Giuseppe Antonacci^{2,3,5}

¹Physics of Life Processes - Kamerlingh Onnes-Huygens Laboratory, Leiden Institute of Physics, Leiden University, Leiden, The Netherlands

²Center for Life Nano Science @Sapienza, Istituto Italiano di Tecnologia, Rome, Italy

³Photonics Research Group, Ghent University - imec, Ghent, Belgium

⁴stefano.coppola87@gmail.com

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⁵giuse.antonacci@gmail.com

Abstract: Cells sense and respond to external physical forces and substrate rigidity by regulating their cell shape, internal cytoskeletal tension, and stiffness. Here we show that the combination of micropillar traction force and noncontact Brillouin microscopy provides access to cell-generated forces and intracellular mechanical properties at optical resolution. Actin-rich cytoplasmic domains of 3T3 fibroblasts showed significantly higher Brillouin shifts, indicating a potential increase in stiffness when adhering on fibronectin-coated glass compared to soft PDMS micropillars. Our findings demonstrate the complementarity of micropillar traction force and Brillouin microscopy to better understand the relation between cell force generation and the intracellular mechanical properties.

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1. Introduction

Cells experience a myriad of mechanical and physical cues within their three-dimensional microenvironment, and in turn, they respond by exerting forces, regulating their shape, internal cytoskeletal tension, and elastic modulus [1–5]. Over the past decade, several studies demonstrated that such physical forces are sufficient to differentiate mesenchymal stem cells [6], initiate transcriptional programs [7], drive morphogenesis [8], and direct cell migration [9]. Disruption of the cellular forces, as well as variations in subcellular mechanical properties, can lead to altered pathophysiological conditions and to the onset of diseases [10–12]. As a result, assessing the cellular forces and mechanical properties is of crucial importance to provide a better understanding of disease mechanisms. Methods to simultaneously gather this information involved the use of picosecond ultrasonics in fixed cells with a μ m-scale lateral resolution [13–15]. In this work, we demonstrate the combination of micropillar traction force and noncontact Brillouin microscopy to non-invasively evaluate both cellular forces and biomechanical properties with a sub-micron spatial resolution.

Microfabrication methods allow the production of cell substrates consisting of patterned arrays of elastic micropillars, usually made of polydimethylsiloxane (PDMS) [16–21] or polyacrylamide [22, 23]. The dependence of the effective substrate stiffness on micropillar geometry (i.e. radius and length) permits the generation of heterogeneous mechanical environments without altering material properties (e.g. cross-linker concentration for PDMS micropillars) [24]. In addition, magnetic actuators can be inserted in individual pillars to provide active mechanical stimuli [25, 26]. By a selective coating with extracellular matrix (ECM) proteins (e.g. fibronectin) through soft lithography techniques, cells adhere and exert forces only at the pillar tips whose deflections report cellular forces at subcellular level. This approach has a few advantages over traction microscopy methods that are based on the measurement of displaced beads embedded

into continuum substrates [27]. These include (i) the capability to calculate displacements by exploiting the undeflected pillar positions in the patterned (e.g. hexagonal) grid, without the need for a reference image and (ii) the simpler and less computationally intensive force calculation due to the discrete adhesive surface (i.e. deflections of a given pillar only depend on the force applied to that pillar). However, compared to continuum substrates, such discrete adhesive surface affects the morphology of cell-ECM adhesions and the ability to reproduce the wide variations in stiffness found in *in vivo* tissues due to the restricted stiffness range (approximately one order of magnitude).

Brillouin microscopy is an emerging approach showing increasing biological interest because it enables an all-optical, label-free and three-dimensional assessment of the subcellular biomechanical properties at a submicron resolution [28, 29]. In Brillouin imaging, a sample is scanned while a narrowband laser source probes point-by-point the spontaneous thermally-activated acoustic waves that locally propagate in the medium at the acoustic velocity [30]. A high-resolution (sub-GHz) spectrometer is thus used to analyze the spectrum of the scattered light [31, 32]. The latter contains an elastic (Rayleigh) peak of the same frequency as the illumination laser beam, and two sidebands referred to as the Stokes and Anti-Stokes Brillouin peaks thinly shifted by few GHz from the central peak [33]. The frequency and the linewidth of the Brillouin peaks are directly related to the high-frequency complex Longitudinal modulus (M), which is indicative of the viscoelastic properties of the material analyzed [34]. Recently, water content has been shown to dominate shifts in the Brillouin frequency of highly hydrated (>90%) gels [35], as expected by the compressibility nature of the longitudinal modulus [36]. Such a scenario is however rather unrealistic in biological samples where water content is significantly lower (<70%) and naturally contribute to the overall stiffness of the material investigated [37].

The noncontact, label-free and high-resolution capabilities of Brillouin microscopy have stimulated a wide range of applications, including the assessment of the cellular mechanical properties and their response to external stimuli [38–41], the mapping of the spinal cord stiffness in zebrafish larvae [42] and the investigation of the Alzheimer's plaque viscoelasticity [43]. Moreover, Brillouin microscopy holds promise to become a potential diagnostic instrument for diseases such as atherosclerosis [44], keratoconus [45], cancer [46], meningitis [47] and amyotrophic lateral sclerosis [48,49]. To this aim, significant research efforts have also focused on the instrumental advancement to achieve a high spectral contrast [38,50,51] and extinction ratio [52,53], as well as to decrease the data acquisition time [54,55], which are key ingredients to enable *in-vivo* measurements of living biosystems.

2. Materials and methods

2.1. Cell biology

3T3 fibroblasts were cultured in high-glucose Dulbecco's Modified Eagle's Medium (DMEM) supplemented with 10% calf serum (Thermo Scientific), 2 mM glutamine and 100 μ g/ml penicillin/streptomycin. Cells were seeded at single cell density directly on the micropillar array or on a glass (#1.5) bottom 35 mm dish (Ibidi). Cells were allowed to spread for 24 hours before fixation in 4% Paraformaldehyde for 15 minutes. After membrane permeabilization with 0.1% Triton-X (Sigma) for 10 minutes followed by a blocking step with 1% bovine serum albumin (BSA, Sigma) in PBS for 1 hour, actin filaments and nuclei were stained with Alexa-Fluor 532-conjugated Phalloidin and DAPI (Life Technologies), respectively. Micropillar arrays were subsequently inverted onto glass (#1.5) bottom 35 mm dish (Ibidi) with imprinted 50 μ m cell location grid and immersed in PBS. Glass weights were used to prevent the arrays from floating.



Fig. 1. Correlative micropillar traction force and Brillouin microscopy. (A) Schematic of the optical fluorescence ('SETUP 1') and Brillouin ('SETUP 2') independent setups. Cells adhere to a hexagonal patterned elastic micropillar array. The fluorescence setup ('SETUP 1') was used to collect the fluorescence signal emitted by the micropillars and therefore to measure the cellular adhesion forces from the pillar displacements [21]. The same sample was then moved to the Brillouin imaging setup ('SETUP 2') where the scattered signal was collected in a backscattering geometry and delivered to the background-deflection (BD) spectrometer. This is a single-stage VIPA spectrometer integrating a diffraction mask (DM) to deflect the elastic background signal from the dispersion axis [48]. (B) Sensitivity histogram of the Brillouin setup. N=1000 spectra were acquired on a PDMS sample. System sensitivity was evaluated to be $\sigma = 0.025$ GHz.

2.2. Force measurement on micropillar arrays

A hexagonal array of poly-di-methyl-siloxane (PDMS, Sylgard 184, Dow Corning) micropillars of 2 µm diameter, 4 µm center-to-center distance and with a height of 6.9 µm were produced using replica-molding from a silicon wafer (for details see [21]). The pillar arrays were flanked by integrated 50 μ m high spacers to allow the inversion onto glass bottom dishes, without compromising the already limited working distance of a high-NA objective on an inverted microscope. The tops of the micropillars were coated with a mixture of Alexa Fluor 647-labeled and unlabeled fibronectin (1:5, Life Technologies) using micro-contact printing. The position of the pillar tops was observed by fluorescence confocal microscopy at 647 nm excitation (Fig. 1A) and determined down to 30 nm accuracy using custom software (Matlab, Mathworks). Forces were obtained by multiplying the pillar deflections by the array's characteristic spring constant (16.7 nN/m determined by finite element modeling) [21]. Such spring constant corresponds to an equivalent Young's modulus in continuous substrates of ~ 11.6 kPa, calculated as proposed in [56]. Only pillars at the cell perimeter and with a deflection above a given threshold (~50 nm) were considered for the calculation of the average force per pillar and total cellular forces. Such threshold was determined for each confocal image as the 75th percentile of the displacements of pillars outside the cell area (i.e. not bent by the cells).

2.3. Confocal microscopy

3T3 fibroblasts adherent on fibronectin-coated glass and soft PDMS micropillar arrays were imaged with an inverted Olympus IX83 microscope, equipped with an UPLSAPO 60XW/1.2 NA

water immersion objective (Fig. 2A-B). The collinear light beams from 405 nm, 546 nm, and 647 nm laser diode light sources were injected into the microscope via a FV1200 MPE laser scanning confocal device. The 2048×2048 pixel fluorescence images (51 nm pixel size) were collected in line sequential mode to reduce the cross-talk among the fluorescence channels.

2.4. Brillouin microscopy

After fluorescence imaging, the same cells were transferred to the Brillouin confocal microscope (Fig. 1A) to investigate their biomechanical properties. The Brillouin microscope is composed of a standard scanning confocal imaging setup working in a backscattering geometry [29] where cells are illuminated by a single longitudinal mode laser beam (λ =532 nm, Coherent Verdi). The scattered light was collected by a microscope objective of NA=1.3 and a single mode fiber working as a confocal pinhole were used, providing a spatial resolution of ~ 0.3x0.3x1.1 μ m³ and a flexible beam delivery to a single-stage Virtually Image Phased Array (VIPA) spectrometer. The latter integrates a rhomboidal-shaped diffraction mask to deflect the background elastic light from the dispersion axis, in turn providing a spectral contrast of ~ 70 dB in a single-stage arrangement, as previously reported in [48]. Brillouin spectra were acquired at each scanning location across the cells and detected by a CCD camera (Photometrics Prime) with a data acquisition (dwell) time of 100 ms. The resulting spectral profiles were fitted by Lorentzian functions to evaluate the frequency shift, which is related to the real part of the Longitudinal compressive modulus *M*' (i.e. a measure of stiffness).

Preliminary measurements were performed to characterize the system sensitivity. To this aim, N=1000 Brillouin spectra of bulk PDMS were acquired and fitted using a Lorentzian function. Fig. 1B shows a histogram of the measured Brillouin shift counts, which gave a mean value of $v_B = 6.237$ GHz and a system sensitivity associated with the uncertainty of $\sigma = 0.025$ GHz.

Representative Brillouin images of 3T3 cells are shown in Fig. 2C-D. Normalized probability density functions (PDFs) of Brillouin shifts were calculated by pooling together the acquired Brillouin images and considering the sensitivity of the system (i.e. $\sigma = 0.025$ GHz) (Fig. 3). The highest peaks in Fig. 3A correspond to the contribution of PBS. To uncouple the buffer component from the cellular contribution, as shown in Fig. 3A, we performed a Gaussian fit (black dashed line) of the highest peak from glass (red solid line) and we defined the Gaussian mean plus its standard deviation as the Brillouin shift threshold (i.e. $v_B^{T}=7.466$ GHz).

3. Results and discussion

We measured in a correlative fashion the contractile forces and the intracellular mechanical properties of 3T3 fibroblast cells adherent on substrates of controlled stiffness (elastic micropillars and glass) and biochemical composition (i.e. fibronectin coating) (Fig. 1). Fig. 2 shows representative fluorescence confocal images of single cells adherent on an elastic micropillar substrate (Fig. 2A) and a standard glass coverslip (Fig. 2B). The associated Brillouin images are shown in Fig. 2C-D. The micropillar deflections allowed us to quantify the forces exerted by 3T3 cells on a soft substrate (Fig. 2A). The Brillouin images show a significant increase in the Brillouin frequency shifts v_B across the nucleus with respect to the cytoplasm, as previously reported [38,39]. Remarkably, the average v_B significantly increased from 7.51±0.05 GHz to 7.57±0.08 GHz (p<0.05) as a function of substrate stiffness (Fig. 2C-D).

To further exploit the subcellular resolution of the Brillouin microscope, we analyzed the distribution of v_B for adherent cells on soft micropillars and glass (solid blue and red lines in Fig. 3A, respectively). The highest peaks correspond to the contribution of the buffer content (i.e. PBS). We then calculated a threshold value (v_B^T =7.466 GHz) above which the v_B values belonged to the subcellular components (see Methods in 2.4). It is interesting to note that the left tail in the distribution of v_B on soft micropillars corresponds to the Brillouin signal of the PDMS micropillars themselves, whose positions can be clearly detected (Fig. 2C). In Fig. 3B



Fig. 2. Correlative high-resolution contractile forces and biomechanical properties of 3T3 fibroblasts. (A) Representative fluorescence confocal image of a 3T3 fibroblast cell (green and blue for actin and nucleus staining, respectively) adhering on fibronectin-coated (red) elastic micropillar arrays. The white arrows indicate the contractile forces quantified by multiplying the pillar displacements by the spring constant. The total cellular force was measured to be 60.8±0.9 nN. (B) Representative fluorescence confocal image of a 3T3 fibroblast cell (green and blue for actin and nucleus staining, respectively) adhering on fibronectin-coated glass. (C-D) High-resolution confocal Brillouin images of the representative cells in (A-B). The reported average cell Brillouin shifts v_B of 7.51±0.05 GHz and 7.57±0.08 GHz are significantly different (p-value<0.05 from a two-sample t-test, N=3146 and 4715 Brillouin shift values respectively). The cell Brillouin shifts were determined by using the triangle method [57]. Scale bar: 10 μm .

we show the multimodal distributions of v_B after thresholding (i.e. across cells) (solid blue and red lines) and the results of a non-linear least-squares fit with three independent Gaussian components (light blue and red lines) (see Fig. 6 in the Appendix for the choice of such model). We focused our further analysis and discussion exclusively on the Gaussian mean values rather than on the amplitude and variance because we believe that the latter present higher dependency on the biological variability and sample size. The obtained parameters in Table 1 interestingly suggest the existence of three major subcellular components with characteristic v_B values. The highest mean Brillouin shift values (i.e. $\mu_3=7.688\pm0.003$ GHz and $\mu_3=7.71\pm0.05$ GHz, for micropillars and glass, respectively) describe the contribution of the nuclei, whereas the lowest values (i.e. $\mu_1=7.489\pm0.001$ GHz and $\mu_1=7.489\pm0.001$ GHz, for micropillars and glass, respectively) describe the cytoplasmatic component, presumably characterized by a more

buffer-like content (i.e. v_B closer to the buffer values, Fig. 3A). Remarkably, these high and low Brillouin components were not significantly different as a function of substrate stiffness (Table 1). However, a different behaviour was found for the intermediate component of the Brillouin shift distribution, which significantly increased from μ_2 =7.544±0.005 GHz to μ_2 =7.608±0.002 GHz (p-value<0.001) in response to the substrate stiffening (Table 1).



Fig. 3. Intracellular Brillouin frequency shifts increase as a function of substrate stiffness. (A) Normalized probability density function (PDF) of Brillouin shifts v_B for cells on soft PDMS micropillars (solid blue line) and glass (solid red line). The Brillouin shifts components for PDMS, PBS buffer and cells are indicated by the arrows. A Gaussian fit of the 'Buffer' peak (black dashed line) is performed to set a threshold value v_B^T =7.466 GHz (see Methods in 2.4). (B) PDF plot of cellular Brillouin shift $v_B > v_B^T$. A three-Gaussian fit (solid lines) is performed for Brillouin shift profiles of cells on both soft PDMS micropillars (solid red line). Results are reported in Table 1. The choice of such model is clarified in Fig. 6 in the Appendix.

Table 1. Parameters from a non-linear least squares fit with three independent Gaussian functions of the Brillouin shifts PDF reported in Fig. 3B. A_* , μ_* and σ_* are the amplitude, mean and standard deviation (±95% CI) of the Gaussian functions. The p-values were obtained from a t-test for which the significance level was set to p-value<0.05 (N=5 cells and 6 cells for soft PDMS micropillars and glass, respectively).

Gaussian Fit	Soft PDMS micropillars	Glass	P-value
A ₁ (%)	39±5	29±1	< 0.001
μ_1 (GHz)	7.489 ± 0.001	7.489 ± 0.001	>0.05
σ_1 (GHz)	0.029 ± 0.001	0.031 ± 0.001	<0.01
A ₂ (%)	40±2	58±8	< 0.001
μ_2 (GHz)	$7.544 {\pm} 0.005$	$7.608{\pm}0.002$	<0.001
σ_2 (GHz)	0.048 ± 0.002	0.062 ± 0.003	< 0.001
A3 (%)	21±1	13±5	<0.01
μ3 (GHz)	7.688 ± 0.003	7.71±0.05	>0.05
σ_3 (GHz)	0.089 ± 0.002	0.09 ± 0.02	>0.05

Tee et al. [4] have previously demonstrated that the cell cortical stiffness increases as a function of substrate stiffness. The cell cortical stiffness was measured by Atomic Force Microscopy (AFM) in regions far from lamellipodia and nucleus, obtaining an ensemble average over both cortical actin and stress fibers (if close enough to the dorsal cell surface). In light of these findings, we hypothesized that the intermediate Brillouin shift values that we introduced above refer to the contribution of actin stress fibers and therefore provide a measure of their mechanical properties.

To validate our hypothesis, we performed a high-resolution Brillouin line scan far from the nucleus crossing actin stress fibers, which were detected in a correlative fashion by fluorescence confocal imaging (dashed red line in Fig. 4). The actin stress fibers could not be spatially resolved by the Brillouin microscope due to a size that typically falls below 100 nm in diameter. Nevertheless, the Brillouin shift line profile had an effective (mean) value close to the intermediate v_B value of Table 1 (μ_2 =7.608±0.002 GHz, dashed grey line). In addition, it is interesting to note a decrease in the Brillouin shifts towards μ_1 (dashed grey line) at the two ends of the line profile, i.e. in the proximity of the lamellipodia. These results support our hypothesis that we could (i) identify the Brillouin shift component potentially associated with the actin stress fibers and (ii) indirectly quantify changes in the intracellular biomechanical properties as a function of substrate stiffness.



Fig. 4. Label- and contact-free Brillouin imaging of actin stress fibers. (A) Confocal image of 3T3 fibroblast adhering on fibronectin-coated glass. Actin filaments are labeled with AF532-conjugated phalloidin (grayscale). The dashed red line indicates where the Brillouin line scan was performed. Scale bar: 10 μm . (B) Brillouin shifts v_B from the line scan across the cell (dashed red line in A) measured with an increased data acquisition time of 500 ms per spectrum. The dashed grey lines report the Brillouin shift values μ_1 , μ_2 , and μ_3 from Table 1.

We further investigated the relation between the actin Brillouin shifts and the mechanical forces generated by 3T3 fibroblasts on soft PDMS micropillars (Fig. 5). The average Brillouin shifts of the intermediate component (i.e. $|\nu_B - \mu_2| < \sigma_2$ from Table 1) increased with the average force per pillar by 0.003±0.004 GHz/nN. This relation provides an estimate of the saturating force per pillar of ~ 23 nN by using the actin Brillouin shift obtained from cells on glass (μ_2 =7.608±0.002 from Table 1). Such estimated force values corroborate data of 3T3 fibroblasts ([56] and unpublished data from our group) on stiffer micropillars (137-150 kPa) for which the average force per pillar saturated at ~ 12-25 nN. Hence, the correlative elastic micropillars and Brillouin microscopy approach allowed us to suggest a potential relation between the forces generated by actin stress fibers and their mechanical properties.

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Fig. 5. Actin Brillouin shifts and force per pillar of cells on soft PDMS micropillars. Data are reported as mean \pm SD. A linear fit (solid red line) using the York method [58] gives a slope of 0.003 \pm 0.004 GHz/nN and an intercept of 7.54 \pm 0.01 GHz (R²=0.14).

4. Conclusion

In this work, we showed a correlative combination of micropillar traction force microscopy and Brillouin imaging methods to quantify cellular forces and intracellular biomechanics as a function of substrate stiffness. The label- and contact-free nature of the Brillouin imaging approach allowed us to go beyond the limitations present in AFM assessments and to quantify the mechanical properties of actin stress fibers with unprecedented spatial resolution. A fully integrated and high-resolution fluorescence and Brillouin microscope with live cell imaging capabilities will grant us simultaneous access to fundamental biomechanical information to better understand cell mechanotransduction.

Appendix



Fig. 6. Comparison of non-linear least squares fits with one, two or three component Gaussian functions for soft PDMS micropillars (A) and Glass (B) conditions. The optimal model (i.e. three-component Gaussian fit) was chosen after the Akaike information criterion (AIC) values that are reported in the plots. The AIC values were determined as $AIC=2k+n\ln(RSS)$, where k is the number of parameters (e.g. k=3 for one-component Gaussian), n the sample size, and RSS the residual sum of squares (i.e. $\sum_i (y_i - f(x_i))^2$).

Funding

AXA Research Fund Post-doctoral grant (16-AXA-PDOC-004).

Acknowledgments

We thank Simone De Panfilis for his support in the fluorescence confocal microscopy.

Disclosures

The authors declare that there are no conflicts of interest related to this article.

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