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A customizable, low-cost 3D-printed device for live cell confinement imaging

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Physical confinement profoundly shapes cell functions from migration to cancer invasiveness. Although multiple *in vitro* confinement platforms have been designed to replicate physical constraints while enabling simultaneous live-cell imaging, broad adoption across the biology community has been limited because most available platforms require specialized fabrication, offer limited design flexibility, or are prohibitively costly. Here, we introduce an inexpensive, fully customizable confinement platform fabricated using standard 3D printers and readily available materials. The confinement device uses a PDMS pillar to uniformly compress two coverslips against polystyrene spacer beads to define precise confinement heights of 3, 7, or 12 μm . We show that this platform reliably confines both adherent and suspension cells, generates graded morphological changes, and maintains high cell viability for at least 24 hours under live-cell imaging conditions. By combining tunability, reproducibility, and broad accessibility, this device provides a practical alternative to existing confinement methods and enables wider exploration of how mechanical restriction modulates cell function across physiological and pathological contexts.

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1. Introduction

In vivo, cells routinely experience physically restricted environments, such as tight interstitial spaces, dense extracellular matrices, and crowded multicellular tissues.^{1,2} The extent of this confinement varies widely across tissues, from the soft, porous tissue environment of the brain to the densely packed architecture of epithelial layers, and becomes even more pronounced during aging, inflammation, or tumor progression.^{3–9} These physical constraints profoundly influence cellular behaviors, including migration,^{10–12} stem cell differentiation,^{13–15} immune cell activation,^{16,17} and epithelial repair during wound healing.^{18–20} In cancer and other pathologies, physical confinement is not simply a byproduct of tissue crowding but an active mechanical stimulus that influences disease evolution.^{21–24} For example, studies have shown that solid tumors are characterized by extreme cell crowding and elevated interstitial pressure which can suppress healthy cell proliferation, dampen anti-tumor immune responses,^{25–28} and increase leader-cell phenotypes amongst cancer cells which drive metastasis.^{23,28,29} Collectively, these findings highlight the need for tools that

enable systematic investigation into how the cellular physical microenvironment impacts cell fate and function.

Despite broad interest, current *in vitro* approaches for studying cell functions under physical confinement remain limited as few tools allow systematic and well-controlled confinement during live-cell investigation. In addition, many of these existing tools involve technical barriers that restrict accessibility and hinder widespread adoption (summary in Table S1). Tools such as AFM provide exquisite force control but are restricted to single-cell measurements and require specialized instrumentation.^{26,27,30} Micropillar arrays, microfluidic chambers, and photolithography-based devices offer controlled confinement at the population level,^{12,23,31–34} but rely on fabrication facilities and expertise unavailable in many biology laboratories. Microporous scaffolds and crosslinked alginate systems can impose defined pore sizes^{16,19,22} and even allow tunable compression,^{7,22} but involve synthetic steps that further limit accessibility. Weighted agarose pads provide a simpler alternative.²⁸ However, their passive loading produces highly non-uniform pressures that depend on gel mass, gel uniformity, and cell morphology, resulting in variable and difficult-to-standardize confinement. Consequently, there remains a critical need for devices that deliver precise, reproducible confinement while being low-cost, easy to assemble, and compatible with standard laboratory workflows.

Here, we present a 3D-printed confinement chamber that directly overcomes these longstanding impediments by providing a simple, customizable, and reproducible platform

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for applying well-defined mechanical confinement to cells. The device incorporates two coverslips which compress cells, polystyrene bead spacers that set confinement height, and a PDMS (polydimethylsiloxane) pillar to apply uniform, gentle compression. This confinement device can also be modified for brightfield imaging and reagent infusion. Crucially, the device can be fabricated using widely available benchtop 3D printers and inexpensive, commercially sourced materials, thus eliminating the need for microfabrication facilities or specialized expertise.

We demonstrate the versatility and compatibility of the confinement device by confining adherent (RAW 264.7/HEK 293T cells) and suspension cells (Jurkat T cells) to different heights for many hours. By enabling rapid, consistent, and scalable confinement assays at minimal cost, this platform makes mechanobiology experiments accessible to a much wider community. More broadly, it establishes a versatile tool for probing how physical confinement shapes cellular behavior across diverse biological contexts.

2. Materials and methods

2.1 Confinement chamber fabrication

Cell confinement device was designed using FreeCAD 0.21 (design detail in SI), exported into UltiMaker Cura 5.9, then printed on an UltiMaker S Series 3D printer extruding tough polylactic acid (PLA). Cura print settings: resolution, 0.1 mm; shell thickness, 1.2 mm; support type, tree; support placement, everywhere; and adhesion, on.

2.2 PDMS pillar fabrication

PDMS elastomer base and curing agent (NC9285739, Fisher Scientific) were mixed at a 16:1 w/w ratio and stirred for 10 min. The mixture was then degassed under vacuum for 20 min. To mold the pillar, an acrylic tube (20 mm height \times 18 mm internal diameter) was placed on top of a coverslip, filled entirely with PDMS, and capped with a 150 g metal weight. Capping both ends of the tube ensured both ends of the PDMS pillar emerged flat. The assembly was cured overnight in an oven at 80 °C. After curing, the PDMS pillar was pushed out of the tube; if stuck, the pillar edges were loosened, and ethanol was added to hasten removal. Cured pillars were sonicated in ultrapure water (18 M Ω cm) for 5 min and sanitized with UV before use.

2.3 Spacer bead solution preparation

Polystyrene particles were diluted in ultrapure water to specific concentrations at given particle sizes (Fig. S1): 6.75 \times 10⁶ particles per mL for 3 μ m polystyrene particles (A37368, ThermoFisher), 2.60 \times 10⁵ particles per mL for 7 μ m polystyrene particles (ABPS-0700, Abvigen), and 7.95 \times 10⁴ particles per mL for 12 μ m polystyrene particles (ABPS-1200, Abvigen). The diluted particle suspensions were sonicated for 1 min and immediately added to poly-L-lysine (PLL) coated coverslips.

2.4 Confinement coverslip preparation

18 mm coverslips (CS-R18-100, AmScope) were sonicated in ethanol for 3 min and rinsed three times with ultrapure water. Cleaned coverslips were dried in an oven at 80 °C. A 0.1% PLL solution (25988-63-0, MilliporeSigma) was prepared by mixing 100 μ L of stock solution with 900 μ L of ultrapure water. Next, 330 μ L of the diluted PLL solution was applied to each coverslip and incubated at 20 °C for 20 min. Coated coverslips were washed three times with ultrapure water and dried again at 80 °C. Finally, 350 μ L of spacer solution was added to each coverslip and the coverslips were incubated at 20 °C for at least 1 h.

2.5 Cell culture and seeding

RAW 264.7 GFP-actin macrophages (generous gift from Dr. Sergio Grinstein, The Hospital for Sick Children Research Institute) and GFP-Fc γ RII 293T cells (generous gift from Prof. Peter Cresswell, Yale University) were cultured in Dulbecco's Modified Eagle Medium (12491015, ThermoFisher) supplemented with 10% fetal bovine serum (MT35010CV, Corning), 2 mM L-glutamine (25030081, ThermoFisher), 100 units mL⁻¹ penicillin-streptomycin (MT30001CI, Corning) at 37 °C in a humidified incubator with 5% CO₂. Jurkat E6.1T cells (ATCC) were cultured in RPMI-1640 medium (30-2001, ATCC) supplemented with 10% FBS (fetal bovine serum), 1 units mL⁻¹ penicillin-streptomycin, and 2 mM L-glutamine at 37 °C in a humidified incubator with 5% CO₂. For passage, RAW 264.7 GFP-actin cells were washed with PBS (phosphate-buffered saline) then removed with a cell scraper. 293T cells were unadhered from flasks with 0.25% trypsin-EDTA (25200056, ThermoFisher). For experiments, 0.5 million RAW 264.7, 0.2 million GFP-Fc γ RII HEK 293T cells, or 0.4 million Jurkat E6.1T cells were seeded on piranha solution etched 30 mm coverslips (30-1313-03192, Bioprotechs). Lower seeding densities were used for larger cells resulting in approximately 50% confluency at time of confinement.

2.6 Confinement device assembly

Cell confinement device assembly is depicted in Fig. 1 and shown in Video S1. First, the top of the confinement device was inverted, and a PDMS pillar was inserted into the central groove of the chamber lid. Next, an 18 mm coverslip containing spacer beads was placed onto the pillar end with the beads facing outward. After that, a 30 mm coverslip with cells was secured in the imaging chamber, which was then pressed into the base of the confiner. Finally, the top and bottom halves of the device were aligned, pressed together gently to apply confinement, then secured with two pins. This assembly process takes approximately 20 s. For cleaning, the device was disassembled and coverslips were discarded. Then, each component of the device was sonicated in soapy water, rinsed thoroughly, and UV sanitized.

2.7 Confocal imaging of confined cells

Cells not expressing GFP were stained with cholera toxin subunit B-Alexa Fluor 488 (C22841, ThermoFisher) for visualization of



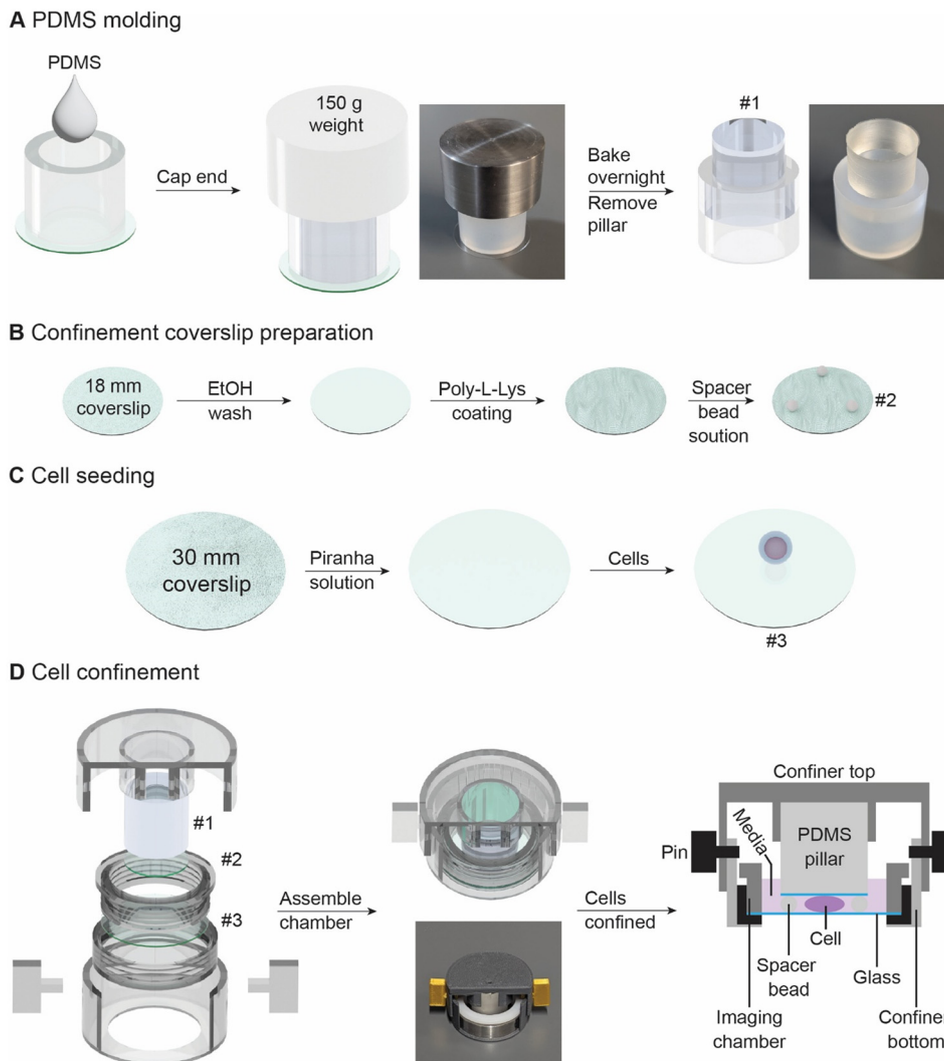


Fig. 1 Fabrication and assembly of the 3D-printed cell confinement device. (A) PDMS pillar fabrication: an acrylic tube was placed on a coverslip, filled with PDMS, capped, baked, and then the cured PDMS pillar was pushed out. (B) Confinement coverslip preparation: coverslips were cleaned with ethanol, coated with PLL, and seeded with a small number of spacer beads. (C) Cell seeding: cells were plated onto a piranha-etched coverslip. (D) Device assembly: the PDMS pillar was inserted into the top of the chamber (#1), the confinement coverslip was attached to the bottom of the pillar (#2), the imaging chamber with cells was pushed into the base of the device (#3), and the two halves were secured with pins. A cross-section of the assembled confinement device is depicted on the right.

cell morphology. All cells were stained with $12.5 \mu\text{g mL}^{-1}$ Hoechst (H21486, ThermoFisher) in cell media for 10 min to label nuclei, followed by a single wash with cell media. Immediately after staining, cells were imaged in FluoroBrite Dulbecco's modified Eagle medium (A1896701, ThermoFisher), then confined and imaged again. Fluorescence channels were acquired as follows: GFP (Ex 488/Em 525), Hoechst (Ex 405/Em 450), and Alexa Fluor 488 (Ex 482/Em 520). Images were acquired using a Leica Stellaris 8 FALCON equipped with a $63\times$ water immersion objective. Confocal images were analyzed using Leica Microsystems's Las X Life Science microscope software platform.

2.8 Absolute confinement measurement

30 mm coverslips were etched in piranha solution then coated with PLL per the protocol used in confinement

coverslip preparation. Next, 30 mm coverslips were placed in imaging chambers and 1 mL PBS + 1 μL ($0.2 \mu\text{m}$) FluoSpheres carboxylate, yellow-green (F8811, ThermoFisher) + 50 μg CF640R (92108, Biotium) were added after brief sonication. 18 mm (confinement coverslips) were prepared as detailed in confinement coverslip preparation except 1 μL of FluoSpheres was added to 1 mL spacer bead solution. After allowing the particles to settle for 1 h, confinement devices were assembled using the FluoSphere coated coverslips and confinement coverslips.

2.9 Raw 264.7 membrane and nucleus morphology measurements

RAW 264.7 GFP-actin macrophages were used to visualize cell spreading. Their nuclei were stained with $2 \mu\text{g mL}^{-1}$ Hoechst



33342 in culture media 30 min before confinement. Images were captured using an 100× oil-immersion objective lens and filter sets for GFP (Ex 488 nm/Em 525 25 nm) and Hoechst (Ex 402 nm/Em 450 nm) at three time points: before confinement, immediately after, and 1 h post-confinement. Cell and nuclear areas were analyzed using ImageJ *via* a multistep process. Cells were segmented *via* thresholding, converted to binary images, then had boundaries refined using the watershed algorithm. Morphological parameters, including area, major diameter, and minor diameter, were quantified using the ‘analyze particles’ function.

2.10 Cell viability assay

Cell viability was assessed using propidium iodide (P4170-10MG, Sigma-Aldrich), a membrane-impermeable dye that fluoresces upon DNA intercalation. Propidium iodide cell staining indicates a loss of membrane integrity. For this assay, cells were seeded on 30 mm coverslips. After cells adhered, 5 μg mL⁻¹ propidium iodide was added to cell media. Static images of PI-positive cells were captured using a Nikon Eclipse-Ti inverted fluorescence microscope equipped with a 40× dry objective lens and GFP/PI filter set (ex 488/ex 565). Fluorescence images were captured before confinement and 3, 6, 24, 48, and 72 hours post confinement. For each sample, three replicates were imaged, then random fields of view per replicate were analyzed. Non-viable cells (PI-positive) and total cells (GFP) were counted using ImageJ (v1.54f). Percent cell viability was calculated using the following formula.

$$\text{Cell viability (\%)} = \left(1 - \frac{\text{PI-Positive cells}}{\text{Total cells}}\right) \times 100$$

2.11 Device modification for reagent infusion

To enable reagent infusion, the confiner top was redesigned to include an off-center port accommodating an 18G needle (305196, BD). The needle tip was sanded flat to prevent damage to the chamber surfaces and ensure smooth insertion. Confining coverslips were cut from sterile Petri dish lids to maintain sterility and compatibility. Assembly proceeded as follows: the PDMS pillar was inserted into the modified confiner top, and the 18G needle was advanced through the confiner top and the PDMS pillar. The coverslip was then positioned at the base of the pillar, and the contact point of the needle was marked. A corresponding hole was drilled at this marked location in the coverslip to create a continuous fluidic channel. The hole in the polystyrene confining surface was made using a 1 mm drill bit. Upon final assembly, the needle tip was aligned flush with the end of the PDMS pillar and precisely matched with the hole in the coverslip to ensure direct access to the confined chamber. Reagent infusion was performed using an NE-1000 syringe pump to provide controlled and reproducible flow.

2.12 Device modification for brightfield imaging

To enable brightfield compatibility, the confinement device top was fabricated using transparent polyethylene

terephthalate glycol-modified (PETG; 32101, Bambu Lab). The component was printed on a Bambu Lab P1S series 3D printer using manufacturer-recommended settings and a 0.4 mm nozzle. Prior to printing, the build plate was coated with adhesive (Bambu Lab glue stick) to promote proper first-layer adhesion and ensure print stability.

2.13 Statistical analysis

Nuclear and cellular deformation under confinement data was analyzed using a one-way ANOVA with Tukey's HSD (honestly significant difference) *post hoc* test. Statistical significance: ns, $P > 0.05$; *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$, ****, $P < 0.0001$.

3. Results

3.1 Device design and operation

Reproducible cell confinement can be technically challenging because the confining surface must be steadily positioned within 3–12 μm of the substrate without compromising cell viability. Some existing approaches, such as the weighted agar gel method, struggle to meet this requirement. The confinement system developed here addresses these challenges through several key design features (Fig. 1). This device is comprised of a rigid 3D-printed chamber body, a PDMS pillar to apply a uniform compressive load, and two glass coverslips with surface-deposited polystyrene spacer beads to define confinement height. The confinement coverslip was coated with (PLL) to promote the electrostatically driven adsorption of polystyrene beads at sufficiently low density that provided enough space for cell culture while maintaining confinement. These spacer beads set the distance between coverslips (Fig. 1B), while the PDMS pillar ensured even pressure across the cellular plane (Fig. 1D). The entire chamber was laterally stabilized by overlapping chamber walls and dual fastening pins to ensure cell viability was not compromised *via* sliding coverslips. We chose three specific confinement heights to induce distinct mechanical regimes, primarily based on the expected deformation of RAW 264.7 cells: slight membrane compression (12 μm), intermediate membrane compression (7 μm), and nuclear compression (3 μm). This selection is consistent with previous confinement studies.^{16,27–30}

3.2 Absolute height and cellular confinement

To evaluate device performance, we quantified the absolute height between the confining coverslip and the imaging chamber coverslip. Both coverslips were labeled with adsorbed 0.2 μm fluorescent beads, and CF640R in PBS was introduced to visualize the inter-coverslip gap. Reconstructed confocal Z-stacks and line-scan intensity profiles demonstrated that the spacer beads maintained precise separations of 12, 7, and 3 μm (Fig. 2A and B; more plots in Fig. S3). Next, we confined three representative cell types: adherent RAW 264.7 macrophages and HEK 293T cells, and



non-adherent Jurkat T cells, and successfully achieved confinement across all cell types (Fig. 2C–E). Confocal Z-stack reconstructions verified that apical-basal cell heights of those cells precisely matched the spacer-defined confinement levels of 12, 7, and 3 μm . Cell morphology and nuclear shape were visualized by staining cells pre confinement using actin-GFP or cholera toxin subunit B-Alexa Fluor 488, with nuclei labeled by Hoechst. In all cases, Hoechst staining clearly revealed nuclear deformation under increasing compression. At 12 μm , nuclei remained largely spherical with only slight deformation, whereas increased confinement produced pronounced flattening, culminating in oblate, disk-like nuclei at 3 μm .

3.3 Quantification of confined cell morphology

To quantitatively validate the morphological changes we observed in confined cells (Fig. 3A and B), we measured the cell spread area and nuclear dimensions ($n \approx 1000$ cells) of GFP-actin RAW 264.7 cells. GFP-actin RAW 264.7 cells with Hoechst-labeled nuclei were imaged one hour after confinement (Fig. 3C). As anticipated, 12 μm confinement produced no significant change in cell spread area,

consistent with minimal mechanical restriction (Fig. 3D). In contrast, 7 μm and 3 μm confinement increased spread area by 13.5% and 44.3%, respectively (Fig. 3D). Membrane blebbing emerged under 7 μm confinement and became more pronounced under 3 μm confinement (Fig. 3C). Blebbing was likely a consequence of increased hydrostatic pressure in the cytoplasm which has been previously reported to cause detachment of the plasma membrane from the cortical cytoskeleton.^{35,36} These blebs can either be resolved if intracellular pressure decreases or a precursor to cell rupture if pressure increases.^{35,37} Nuclear morphology followed similar trends. Nuclear area increased only modestly from $\sim 80 \mu\text{m}^2$ in unconfined cells to a slightly larger value at 12 μm , but expanded markedly at 7 μm and reached $\sim 120 \mu\text{m}^2$ at 3 μm confinement (Fig. 3E). The nuclear major axis length increased by 4.7%, 11.6%, and 16.7% at 12, 7, and 3 μm confinement, respectively, while the minor axis length exhibited slightly greater increases of 4.0%, 13.9%, and 19.5% (Fig. 3F and G). To assess the consistency of the confinement height across the sample chamber, in addition to the data in Fig. 3D obtained from RAW 264.7 cells expressing actin-GFP, we measured cell spreading in RAW

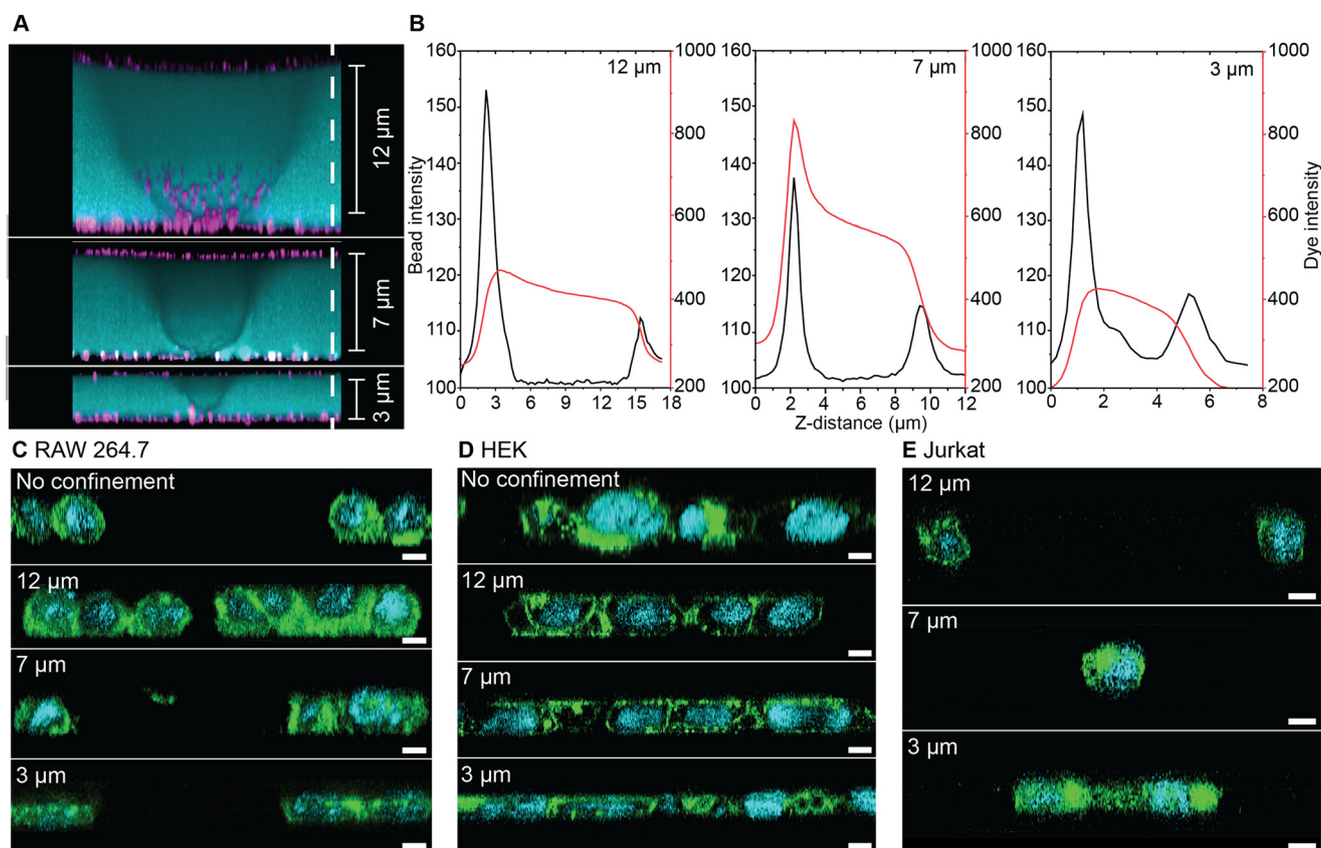


Fig. 2 Confocal Z-stack reconstructions illustrating absolute confinement, adherent cell confinement, and non-adherent cell confinement. (A) Confinement heights of 3, 7, and 12 μm , visualized using coverslips coated with 0.2 μm fluorescent beads (magenta) and CF640R in PBS filling the chamber (cyan). (B) Line-scan intensity profiles of surface-adsorbed beads on coverslips and fluorescent dye in PBS, indicating confinement heights. (C–E) RAW 264.7, HEK 293T, and Jurkat T cells under different levels of confinement. Cell-specific labels: (B) RAW 264.7: F-actin (GFP, green) and nuclei (Hoechst, cyan); (C) HEK 293T cells: Fc γ RII (GFP, green) and nuclei (Hoechst, cyan); (D) Jurkat T cells: membrane (cholera toxin B-Alexa Fluor 488, green) and nuclei (Hoechst, cyan). Scale bars 5 μm .



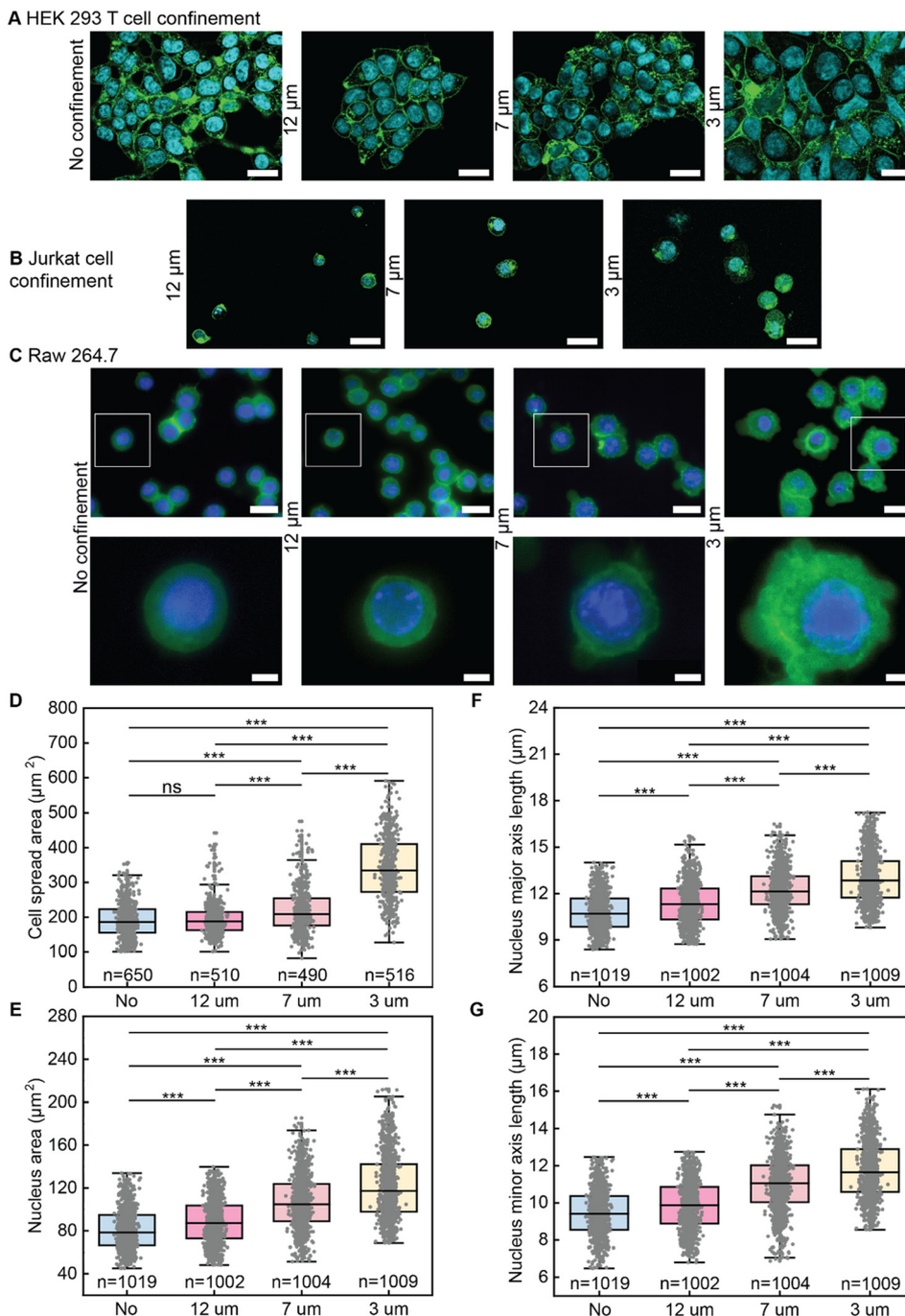


Fig. 3 Visualization and quantification of 2D nuclear morphology and cell spread area under confinement. (A) Epifluorescence x,y slices of HEK 293 T $Fc\gamma RII$ (GFP, green) and nuclei (Hoechst, cyan) under 12, 7, and 3 μm confinement. (B) Epifluorescence x,y slices of Jurkat T cells: membrane (cholera toxin B-Alexa Fluor 488, green) and nuclei (Hoechst, cyan) under 12, 7, and 3 μm confinement. (C) Representative epifluorescence images of RAW 264.7 macrophages expressing GFP-actin (green) with Hoechst-stained nuclei (blue) before and after 1 h of confinement at 12, 7, and 3 μm . (D) Cell spread area, (E) nuclear area, (F) nuclear major axis length, and (G) nuclear minor axis length for confined RAW 264.7 cells. Scale bar: 20 μm ; zoomed-in regions: 5 μm . Box plots show median (thick line), first and third quartiles (box edges), and whiskers representing 1.5 \times interquartile range; scatter points represent individual cells. Data are pooled from about 5000 cells from $N \geq 3$ independent experiments. Statistical significance: ns, $P > 0.05$; ***, $P < 0.001$ (one-way ANOVA with Tukey's HSD *post hoc* test). Exact p -values are shown in SI Fig. S4.

264.7 cells expressing RelA-GFP (Fig. S5A), a reporter cell line used to monitor inflammatory NF- κ B responses. The results confirmed that both cell types exhibited comparable cell areas under confinement. In addition, when cell areas were

averaged from images acquired at multiple locations across the chamber and plotted (Fig. S5B), the values showed negligible variation between locations, indicating that the confinement height was uniform throughout the device.



Together, these quantitative measurements confirm that the device reliably produces characteristic increases in cell spreading and nuclear deformation associated with graded mechanical confinement.

3.4 Cell compatibility

Lastly, we evaluated cell compatibility, a critical attribute for any confinement device, by quantifying cell viability and proliferation. Viability was assessed using propidium iodide (PI) staining, which selectively labels cells with compromised membranes.³⁸ RAW 264.7 macrophage cells were confined at 12 μm , 7 μm , and 3 μm for durations ranging from 0 to 72 h (Fig. 4). Cell viability was calculated as the percentage of PI-positive cells relative to the total number of GFP-labeled cells. Confinement for up to 24 h maintained high viability

(>95%) across all groups, comparable to the unconfined controls, suggesting minimal impact on cellular integrity (Fig. 4B). Extended confinement (48–72 h), however, caused a progressive decline in cell viability inversely related to confinement height (Fig. 4B). After 72 h, unconfined controls showed 78% viability, similar to 74% under 12 μm confinement, which induces minimal membrane compression. In contrast, viability dropped to 67% and 55% under 7 μm and 3 μm confinement, respectively, reflecting increased mechanical stress. These results align with previous reports of reduced viability under extreme confinement.^{31,39} Collectively, these findings demonstrate that our 3D-printed confinement device is highly compatible with cells for up to 48 h, with longer durations leading to decreased viability due to combined effects of mechanical stress and nutrient limitations. In addition, we observed negligible cell proliferation under all three

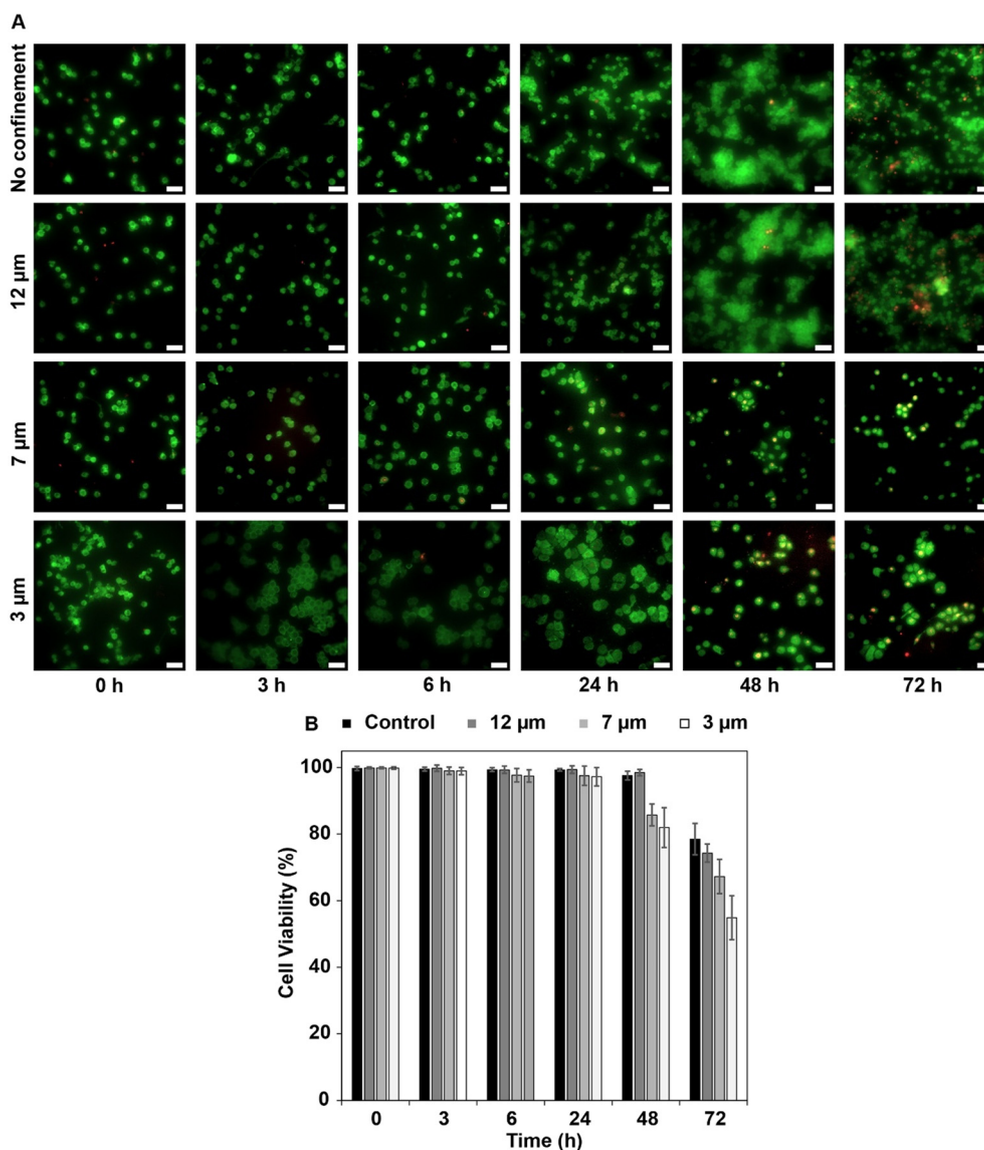


Fig. 4 Viability of RAW 264.7 GFP-actin cells under confinement. (A) Representative epifluorescence images showing GFP-actin (green) and propidium iodide-stained dead cells (red). (B) Quantification of cell viability over time at different confinement heights.



confinement conditions. This observation is consistent with literature showing that even modest confinement, with confined heights below 20 μm , can inhibit cell proliferation due to restrictions on cell rounding required for mitosis.^{27,31,40}

4. Discussion

Beyond device fabrication, cell confinement, and assessment of viability and proliferation, we next expanded the platform to enable *in situ* reagent infusion and exchange (Fig. 5A). To incorporate an infusion port, the top confining surface was fabricated from a sterile Petri dish lid, and a vertical channel was created by punching a hole through the device top, PDMS pillar, and confining coverslip. This channel accommodated an 18G needle (tip sanded flat), allowing controlled introduction of reagents into the confined chamber without disassembly.

To validate this modification, 0.2 μm fluorescent beads and 7 μm spacer beads were first adsorbed onto the imaging

chamber coverslip. The infusion port was loaded with CF640R in PBS, and the device was assembled under confinement. Two-channel confocal imaging showed that, upon initiation of injection, CF640R rapidly flowed across the bead-labeled surface (Fig. 5C). Quantitative analysis revealed that fluorescence intensity began increasing at approximately 50 s and reached a plateau by ~ 90 s (Fig. 5B), confirming efficient reagent exchange within the confined space. Using the same approach, we successfully stained confined cells with Hoechst (Fig. S6). Together, these results establish that reagents can be delivered reliably to confined cells, enabling time-resolved perturbation experiments.

In addition, the device can be adapted for brightfield compatibility when required. By fabricating the confinement top from transparent polyethylene terephthalate glycol-modified (PETG), we enabled brightfield imaging directly through the device (Fig. 5D).

We further systematically compared alternative methods for spacer bead immobilization, particularly in cases where

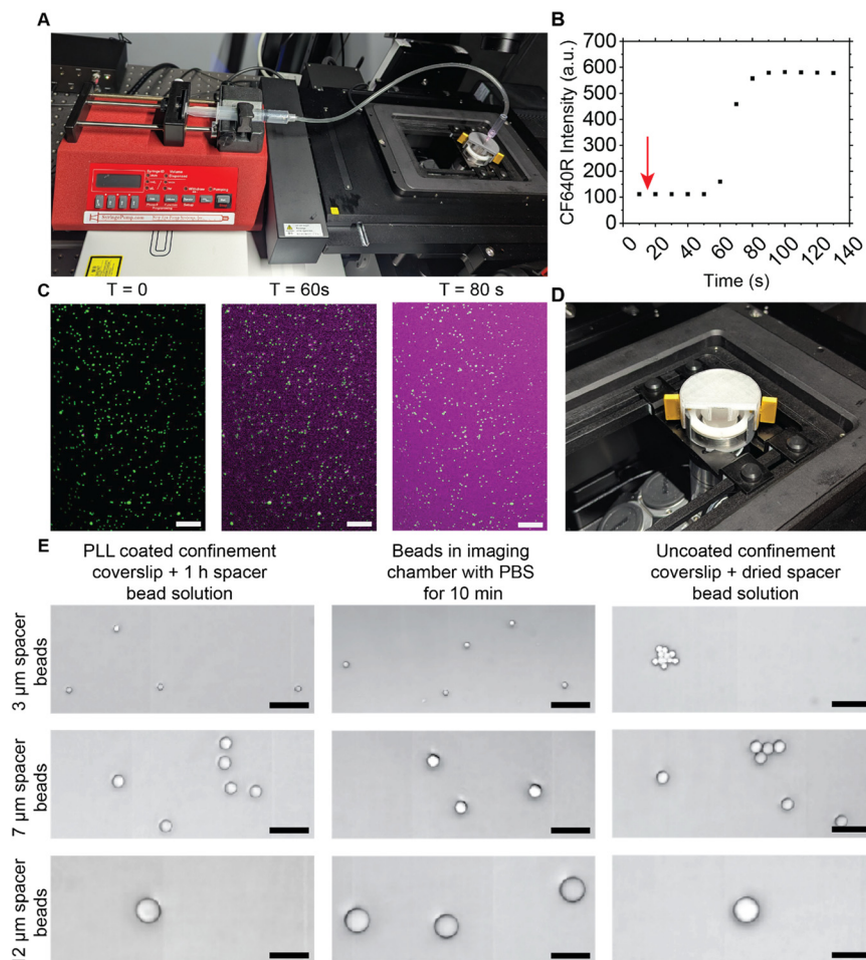


Fig. 5 Adaptivity of the confinement device for infusion and different spacer bead immobilization. (A) Confinement device with infusion ports mounted on the microscope stage. (B) Fluorescence intensity of infused CF640R dye plotted as a function of time following initiation of dye injection (red arrow). (C) Confocal images showing 0.2 μm fluorescent beads marking the imaging chamber surface and injected CF640R in PBS (magenta). Scale bar, 4 μm . (D) Brightfield-compatible confinement device mounted on the microscope stage. (E) Brightfield images showing spacer bead distributions under confinement using different bead incorporation methods: PLL coated coverslips (left column), adsorption without PLL (middle column), and evaporation-induced adsorption (right column). Scale bar 20 μm .



PLL coating is undesirable. The first method, which was used throughout the main study, involved incubating spacer bead suspensions on PLL-coated confinement coverslips for 1 h prior to assembly (Fig. 5E, left). To eliminate PLL, we tested a second method in which spacer beads were introduced into PBS-filled imaging chambers and allowed to adsorb for 10 min before confinement (Fig. 5E, middle). A third approach relied on evaporation: spacer bead suspensions were deposited onto uncoated confinement coverslips, allowed to dry, and then assembled (Fig. 5E, right). We found that 7 and 12 μm spacer beads could be incorporated effectively using all three methods, showing comparable distributions and maintaining the intended confinement heights. In contrast, 3 μm beads aggregated substantially under the evaporation method and therefore could not be reliably incorporated using this approach. Importantly, these findings demonstrate that PLL coating is not required for spacer bead immobilization, which may be advantageous in applications where PLL could influence cellular behavior.

Our PDMS pillar confinement device had two unsuccessful precursors. We previously attempted to confine cells using a weight-based confiner and a screw-based confiner (Fig. S7). These devices were unable to induce consistent, level confinement largely due to resolution limits of affordable 3D-printers and lateral coverslip motion. Thus, we incorporated two key design elements into our final device which ensured proper function. First, uniform compression was achieved by integrating a PDMS pillar. The moderate compressibility of PDMS compensates for minor inaccuracies in 3D printing, ensuring even force distribution across the confining coverslip. Second, lateral stability was enhanced through a nesting shell design fastened with dual pins. In addition, we were able to further improve our confinement device to overcome another limitation of many confinement systems: reliance on diffusion to deliver reagents to cells. Our device was modified to allow rapid delivery of reagents directly to confined cells by incorporating an injection channel through the PDMS pillar. Finally, we were able to achieve brightfield imaging by incorporating a transparent top. Collectively, these design refinements resolve key challenges encountered in earlier prototypes and provide a versatile, accessible, and reproducible platform for studying cellular responses under defined mechanical constraints.

5. Conclusions

Here we present a microscopy-compatible, 3D-printed cell confinement device that provides a robust, tunable, and accessible platform for quantitatively studying cellular responses to mechanical confinement. Our device was able to probe a range of cellular confinement levels from membrane perturbation to nuclear deformation. In addition, cell viability was preserved under short-term confinement (up to 48 h) at all tested heights (3–12 μm) which allows for characterization of both short- and long-term cellular responses to confinement. Prolonged confinement (48–72 h)

at 7 and 3 μm , reduced viability and inhibited proliferation, highlighting the impact of sustained mechanical stress on cellular homeostasis. Compared with existing confinement platforms, this device offers advantages in affordability and adaptability, as demonstrated by its tunable confinement range, ease of operation, compatibility with reagent infusion, and support for brightfield imaging.

Author contributions

H. R. conceptualized, designed, and fabricated cell confinement devices. H. R. and J. O. conducted experiments. H. R., J. O., and M. K. analyzed data. H. R. drafted the manuscript. H. R. and Y. Y. revised the manuscript.

Conflicts of interest

The authors declare no competing interests.

Data availability

Data related to the manuscript and confinement device CAD models are provided in the main text and supplementary information (SI) section. Additional raw data is available at Figshare: <https://doi.org/10.6084/m9.figshare.30862697>.

Supplementary information is available. See DOI: <https://doi.org/10.1039/d5lc01146a>.

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