

# Microfluidic chambers using fluid walls for cell biology

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Many proofs of concept have demonstrated the potential of microfluidics in cell biology. However, the technology remains inaccessible to many biologists, as it often requires complex manufacturing facilities (such as soft lithography) and uses materials foreign to cell biology (such as polydimethylsiloxane). Here, we present a method for creating microfluidic environments by simply reshaping fluids on a substrate. For applications in cell biology, we use cell media on a virgin Petri dish overlaid with an immiscible fluorocarbon. A hydrophobic/fluorophilic stylus then reshapes the media into any pattern by creating liquid walls of fluorocarbon. Microfluidic arrangements suitable for cell culture are made in minutes using materials familiar to biologists. The versatility of the method is demonstrated by creating analogs of a common platform in cell biology, the microtiter plate. Using this vehicle, we demonstrate many manipulations required for cell culture and downstream analysis, including feeding, replating, cloning, cryopreservation, lysis plus RT-PCR, transfection plus genome editing, and fixation plus immunolabeling (when fluid walls are reconfigured during use). We also show that mammalian cells grow and respond to stimuli normally, and worm eggs develop into adults. This simple approach provides biologists with an entrée into microfluidics.

microfluidics | fluid walls | tissue culture | sessile drops | interfacial tension

In icrofluidics addresses the manipulation of tiny volumes, typically less than 1  $\mu$ L. Despite many proofs of concept involving common protocols in cell biology, uptake of microfluidics by biologists is limited. Some well-known and documented reasons include the cost and complexity of manufacture of microfluidic devices (which might involve soft lithography and clean rooms), concerns regarding biocompatibility (materials used for fabrication like polydimethylsiloxane and the associated solvents are not traditionally applied in cell biology), and the inaccessibility of cells after being introduced into enclosed spaces within devices. This prompts the development of many alternatives, including "open" and "paper-based" microfluidics (1).

Microtiter plates are widely used during liquid handling; each is essentially an array of miniature test tubes with 96, 384, or 1,536 wells in a uniform footprint, where wells have working volumes of ~100 to 400, ~15 to 150, or ~3 to 10  $\mu$ L, respectively. Arrays with more wells and volumes down to a few femtoliters have been developed (2, 3). Arrays of aqueous drops sitting on flat (usually patterned) surfaces and overlaid with an immiscible liquid to prevent evaporation have also been fabricated (4–9); in these, liquid walls/ceilings confine the aqueous phase. The burgeoning field of droplet-based microfluidics also uses fluid walls to confine liquids (10–12). However, compared with the widespread use of microtiter plates, few of these alternatives are incorporated into workflows in cell biology (13); consequently, most still involve volumes of many microliters.

A recent method termed Freestyle Fluidics allows fabrication of microfluidic circuits by dispensing cell media in a desired pattern on a Petri dish and overlaying it with an immiscible liquid (14). The aqueous phase is bounded by fluid walls—the interface between water and immiscible liquid. One of the resultant circuits was used in a chemotaxis experiment with bacterial cells, and several benefits compared with traditional circuits were

demonstrated. Here, we also created microfluidics patterns with fluid walls. However, instead of depositing the aqueous phase in the desired pattern on the substrate and then overlaying the immiscible liquid, we simply reshaped the two fluids already on the substrate and allowed interfacial forces to build fluid walls accurately, reproducibly, and immediately. At the microscale, these fluid walls prove to be strong, pliant, and resilient; they morph above unchanging footprints when nanoliter volumes are added/removed. Although any 2D pattern can be made, we demonstrate the method and its versatility by creating analogs of a familiar experimental platform in cell biology, the microtiter plate. We show that mammalian cells grow and respond to stimuli normally and that worm eggs develop into adults. We also demonstrate many basic manipulations involved in cell biology (i.e., cell feeding, replating, cloning, and cryopreservation), plus some common downstream workflows (i.e., fixation/immunolabeling, cell lysis/RT-PCR, transfection/genome editing). Furthermore, we go beyond what is possible with existing microfluidics and reconfigure the fluidic structures in real time. We suggest that this method provides biologists with an easy entrée into microfluidics, without the usual expertise/equipment requirements, while also providing the freedom to create and reconfigure designs on demand.

#### **Significance**

Despite improvements in our ability to manipulate eversmaller volumes, most workflows in cell biology still use volumes of many microliters. We describe a method for creating microfluidic arrangements containing submicroliter volumes. It exploits interfacial forces dominant at the microscale to confine liquids with fluid (not solid) walls. We demonstrate many basic manipulations required for cell culture and some widely used downstream workflows. The method eliminates many problems associated with the fabrication of conventional microfluidic devices, thereby providing a simple on-demand approach for fabrication of microfluidic devices using materials familiar to biologists.

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#### Results

**Methodology.** Fig. 1.4 illustrates the fabrication of a  $2 \times 2$  grid. The bottom of a standard polystyrene Petri dish is completely covered with cell medium, excess medium is removed, and the residual thin film is overlaid with an immiscible liquid. This overlay can be less dense than water, like a hydrocarbon. Perhaps counterintuitively, it can be denser, like FC40—a transparent

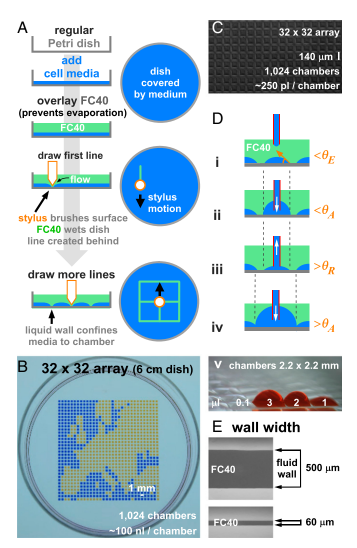


Fig. 1. Reverse printing. (A) Principle. A Petri dish is filled with DMEM plus 10% FBS, most medium is removed, and the residual thin film is overlaid with FC40 (shown in green). A hydrophobic stylus is lowered to touch the dish and to bring FC40 to the surface. As the stylus moves horizontally, medium is pushed aside and FC40 takes its place. This creates a track of FC40 pinned to the substrate and a liquid wall of FC40 dividing the aqueous layer. Drawing more lines creates a grid. (B) A  $32 \times 32$  grid made in  $\sim 4$  min. After printing, 70 nL of yellow or blue dye is added to each chamber. (C) A high-density grid made with a thin stylus (73% surface covered by medium). (D) Adding and subtracting medium. (i) The contact angle is  $\langle \theta_F \rangle$  (~70°). (ii) Medium can be added without altering the footprint until  $\theta_A$  is reached  $(\theta_A > \theta_E)$ . (iii) Medium can be removed without altering the footprint until  $\theta_R$ is reached ( $<3^{\circ}$ ), (iv) If  $\theta_A$  is exceeded, the pinning line breaks and chambers merge. (v) Within limits imposed by  $\theta_A$  and  $\theta_E$ , grids are used like conventional plates; aqueous liquids are pipetted into (or out of) chambers through FC40 instead of air. Here, 1 to 3  $\mu L$  of dye was added to chambers initially containing 0.1  $\mu L$  of medium. Note that the maximum contact angle in the square drop with 3  $\mu L$  was >70°. (E) Stylus width determines wall width. Lines were made using styli with wider and narrower tips than in B, and regions between chambers were imaged.

fully fluorinated liquid (density 1.855 g/mL) that is widely used in droplet-based microfluidics; at the microscale, effects due to gravity and buoyancy become negligible, and interfacial forces pin the aqueous phase to the plastic. A hydrophobic and fluorophilic stylus with a conical tip made of polytetrafluoroethylene (Teflon) and held by a three-axis traverse (a "printer") is then lowered through both liquids until it just touches the dish. Because FC40 wets Teflon and polystyrene better than water, the tip (now coated with FC40) brings fluorocarbon down to wet the substrate. When the stylus moves laterally, the aqueous liquid is displaced from the surface to leave a track of FC40 pinned to plastic by interfacial forces. Drawing more lines creates a grid.

Grids with few or many chambers can be made in minutes (Fig. 1 B and C, SI Appendix, Fig. S1, and Movie S1); for example, chamber density in the grid in Fig. 1C is analogous to that of a microplate with 393,216 wells. Colored dyes are often pipetted into chambers to aid visualization; they play no role in stabilizing liquid structures. Individual chambers are used much like wells in conventional microplates; liquids are simply pipetted into (or removed from) them through FC40 instead of air (Fig. 1D). This can be achieved without altering the footprint on the dish. Consider a sessile water drop in air sitting on a standard polystyrene Petri dish. The drop is shaped like the cap of a sphere, and its footprint depends on the equilibrium contact angle (the angle  $\theta_E$ , at the airwater-substrate interface) (15, 16). When tissue culture medium without serum replaces water,  $\theta_E$  is ~50°, and  $\theta_E$  increases to ~70° if FC40 replaces air (14). Slightly more medium can now be added without increasing the footprint, up to a limit determined by the advancing contact angle,  $\theta_A$  ( $\theta_A > \theta_E$ ); once  $\theta_A$  is breached, footprint area increases. Similarly, when medium is removed, the footprint shrinks once the receding contact angle,  $\theta_R$ , is reached. However,  $\theta_R$  is <3°, so at least 95% of a 5- $\mu$ L drop of medium can be removed without altering the footprint (14). Hereafter, medium with serum will generally be used, and  $\theta_A$  becomes >70°. The significant difference between  $\theta_A$  and  $\theta_R$  allows the addition and removal of liquids above unchanging footprints (Fig. 1 D,  $\nu$ ). The spacing between chambers can also be varied using styli with wider or narrower tips (Fig. 1E).

FC40 plays several additional roles. Fluorocarbons like FC40 were developed during the Manhattan Project as materials that could resist attack by highly reactive uranium hexafluoride; consequently, they are arguably the most inert liquids known. They are also the carrier fluid of choice in droplet-based microfluidics. In addition, FC40 carries the vital gases (O<sub>2</sub> and CO<sub>2</sub>) so effectively that it has been used as a blood substitute (17), and FC40's close relatives have been used for liquid ventilation of human preterm neonates (18, 19). FC40 also prevents the underlying aqueous layer from evaporating (the solubility of water in FC40 is <7 ppm by weight at room temperature) while isolating each chamber from others in a dish (and from the surroundings), thereby preventing contamination and communication between adjacent chambers. For example, if bacteria are pipetted manually through FC40 into every second chamber in a grid, bacteria grow only in inoculated chambers; the rest remain sterile (Fig. 24). Moreover, FC40 stabilizes pinning lines sufficiently so that grids can be shaken without altering footprints (Movie S2). While we generally use FC40, the method works equally well with silicone and hydrocarbon oils; glass surfaces can also be used, as can polystyrene coated with cell-friendly materials (Fig. 2 B–D).

Varying Initial Chamber Volume. The volume of liquid initially in a chamber can be controlled in various ways. In one, different amounts of media are added during the first step; however, the interface has curvature and, hence, the fluid does not have an equal height across the dish. Therefore, it is convenient to use the approach in Fig. 2E: Three 6-cm dishes are covered with thin films plus blue dye, FC40 is added, and four lines are drawn in each to create three large central chambers (with volumes of

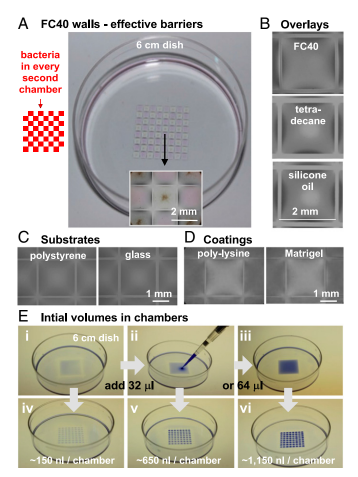


Fig. 2. Some different ways of making grids. (A) FC40 isolates chambers effectively. A 1.5  $\mu$ L portion of medium  $\pm$  *E. coli* was pipetted manually into every second chamber in an 8  $\times$  8 grid (pattern shown in cartoon), with 2  $\times$ 2 mm chambers. After incubation (24 h; 37 °C), a phase-contrast image was collected. Bacteria grew only in inoculated drops (seen as aggregates in chambers containing exhausted, slightly yellow media), and the rest remained sterile (slightly-pink chambers). (Inset) Aggregates and granularity indicate presence of bacteria. (B-D) Various overlays and substrates yield stable pinning lines. Films of media on polystyrene (uncoated except in D) or on glass (C) were overlaid with 2 mL of FC40 (or other oils in B), grids were created, and phase-contrast images were collected. (E) Preparing grids with different starting volumes (2.25  $\times$  2.25 mm chambers). (i-iii) Three dishes were covered with medium plus blue dye, FC40 was added, four 20-mm lines were drawn to create one central square in each dish (volume  $\sim 10 \text{ µL}$ ), and 0, 32, or 64 μL was pipetted into squares. (iv-vi) Subdivision yields grids.

 $\sim 10 \mu L$ ) to which more media plus dye are added (i.e., 0, 32, or 64 μL); lastly, the now-different volumes are split into three grids. Each of the 64 chambers in the three dishes ends up with  $\sim$ 150,  $\sim$ 650, or  $\sim$ 1.150 nL.

Essentially, all grids illustrated here are made by passing the stylus down the centerline of a large square or rectangle to divide them symmetrically into two; next, repeated symmetrical divisions (SI Appendix, Fig. S2 A, i) yield chambers with equal footprints and volumes. This is demonstrated using fluorescein instead of blue dye: chambers in a grid end up with similar fluorescence intensities, so volumes (chambers in SI Appendix, Fig. S2 A, ii) have a volume of 850 nL  $\pm$  4% (SD) (see SI Appendix, Fig. S2 for additional results, the calibration method, and reproducibility in a repeat experiment). However, if the stylus passes asymmetrically through a chamber offset from the centerline, the resulting subchambers can end up with very different volumes (in SI Appendix, Fig. S2B, they have 100 to 1,000 nL in an  $8 \times 8$  grid).

Adding and Removing Liquids. Liquids can be delivered to conventional wells and to our chambers in the same way: by immersing a pipet tip into a recipient well/chamber (Fig. 1 D, ii). However, when repeating this process with the same pipet, liquids can be carried over from one well/chamber to the next, so pipets are usually washed or replaced between deliveries. Therefore, we investigated methods not requiring washing or replacement. In our printer, the pipet is a stainless steel tube mounted next to the stylus on a three-axis traverse. In one method, the pipet tip was positioned in the FC40 above a recipient chamber, and liquid was ejected into it through a transient "liquid bridge" (Fig. 3A). Because the pump driving ejection starts and stops and the tip moves down, up, and across, we call this "discontinuous" delivery. This method has been used to deliver one liquid to many chambers without carryover (20). Fig. 3B and Movie S3 illustrate a liquid bridge without detectable upward transfer of red dye from recipient chamber to delivering pipet that could cause carryover. We

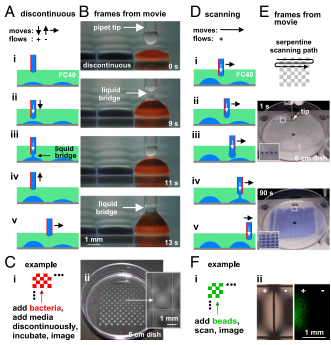


Fig. 3. Adding liquids discontinuously and continuously. (A) Discontinuous. (i-iii) The pipet is lowered (black arrow), and the pump ejects liquid (white arrow) that eventually merges with the chamber. (iv, v) After stopping the pump, the pipet is raised and moves to the next chamber. (B) Frames from Movie S3 illustrating that no red dye moved up through a liquid bridge to the pipet. Note that contact angles at the centers of chamber edges are  $\sim$ 90°. (C) Lack of carryover of bacteria during discontinuous delivery (2  $\times$ 2 mm chamber, ~150 nL). (i) A 300-nL portion of LB with or without ~20,000 E. coli was added manually to every second chamber, and then 500 nL of LB was added to all chambers by discontinuous delivery. (ii) After 3 d at 20 °C, imaging shows that bacteria (white) grew only in inoculated chambers. (D) Continuous (scanning). (i-iii) The pipet maintains a constant height (450 µm) above the substrate as it traverses (black arrow), continuously ejecting liquid (white arrow). (iv) Liquid is delivered to the chamber. (v) Continuing traverse breaks the liquid bridge (maximum final chamber height 380 µm). (E) Frames from Movie S4. The tip scans (15 mm/s) along a serpentine path, delivering ~70 nL of medium plus blue dye to each chamber. Insets show 4× magnifications. (F) No carryover of fluorescent beads between chambers during scanning (16  $\times$  16 grid, 2  $\times$  2 mm chambers, ~150 nL). (i) A 300-nL aliquot of medium with (+) or without (-) 9,000 fluorescent beads (1-µm diameter) was added manually to every second chamber in the grid. A total of 500 nL of medium was added by scanning to each chamber. (ii) Phase-contrast (Left) and fluorescent (Right) images showing that no beads were carried over between chambers.

demonstrated lack of carryover in another way. Bacteria were inoculated into every second chamber in a grid, and medium was delivered discontinuously into all chambers using the same tip (Fig. 3 C, i); on incubation, bacteria grew only in inoculated chambers while the others remained sterile (Fig. 3 C, ii). This confirms that bacteria are not carried over when medium is delivered, and that FC40 provides a sterile barrier between chambers.

With grids, there are no solid walls, and so no need to raise or lower the pipet as it traverses or to start and stop a pump. Consequently, one liquid can be delivered to many chambers by a pipet as it moves at constant speed and height, ejecting liquid continuously (Fig. 3D). Such "scanning" can deliver  $\sim$ 70 nL to each of 1,024 chambers in 90 s (Fig. 3E and Movie S4). The variation in delivery to a 16 × 16 grid (measured as in SI Appendix, Fig. S2) was 27 nL; it was 10 nL to a 32 × 32 grid (SI Appendix, Materials and Methods). Such delivery occurs without detectable carryover, exemplified by failure to transfer fluorescent beads between chambers (Fig. 3F). This illustrates another advantage of fluid walls: one liquid can be delivered through one tip speedily to many chambers without carryover because FC40 always remains in contact with the tip.

Our chambers have square footprints, and contact angles vary around the footprint (*SI Appendix*, Fig. S3). Therefore, we estimate maximum volumes that can be added to chambers based on scaling relative to a sessile circular drop with  $\theta_A$  of 70° and  $\theta_R$  of 3° (values for media without serum) and diameter equal to chamber width. Thus, a 1-mm square chamber in Fig. 1B is limited to minimum and maximum volumes of 4 and 120 nL, respectively (*SI Appendix*, Table S1 gives working volumes for chambers with different footprints; all are conservative estimates, as adding serum to medium increases  $\theta_A$ ).

Biocompatibility—Cells and Organisms. Grids are made using bioinert components: tissue-culture media and polystyrene dishes used by biologists, plus FC40. To confirm biocompatibility, a grid was prepared, NM18 cells (a line of mouse mammary tumor cells) were plated by scanning, and the dish was placed in a conventional CO<sub>2</sub> incubator (O<sub>2</sub> and CO<sub>2</sub> exchange freely through FC40); cells grew normally (Fig. 4A), as have all cells tested to date. Manipulations used in conventional tissue culture are easily adapted to grids, including feeding, trypsinization, replating (Fig. 4 A and B), and cloning (see Example of a Complex Cell-Based Workflow—Genome Editing). Cells can be grown on polystyrene or glass, and on polystyrene dishes coated with polylysine, Matrigel, collagen, fibronectin, and laminin (Fig. 2D and SI Appendix, Fig. S4 A and B). Cells can be adherent or nonadherent (SI Appendix, Fig. S4 B and C). They can also be cryopreserved in grids (Fig. 4C), as in conventional microfluidic devices (21, 22); this shows that these pinning lines are stable, which is remarkable, given the complex effects occurring when oilwater emulsions are frozen and thawed (23-25). Thus far, fluid walls have been made before adding cells; however, they can also be built around preplated cells (Fig. 4D). Consequently, fluid walls can be built around cells of particular interest already growing in a dish, so those with a characteristic morphology or expressing a particular fluorescent marker can be isolated. Note that here, the microfluidic pattern was modified during use, which is impossible with most microfluidic chips. Note also that FC40 provides an additional line of defense against contamination, as it is impermeable to bacteria (Fig. 3C). These experiments show that the standard procedures required for mammalian cell culture can be carried out in grids using submicroliter volumes.

We next demonstrate that whole organisms develop normally. *Caenorhabditis elegans* is a roundworm ~1 mm long that swims by undulatory locomotion; dorsal and ventral muscles contract alternately to generate waves along the worm's axis (26). Worms have been studied in conventional microfluidic devices (27) and droplet-based systems (28). We wished to see whether pinning

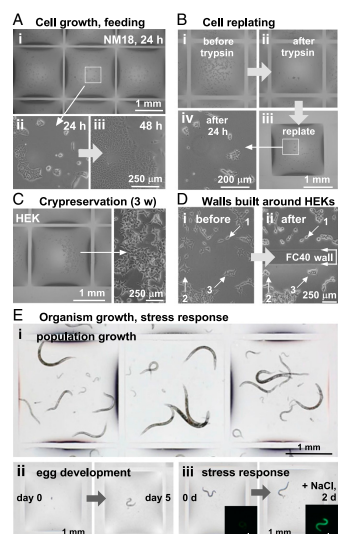


Fig. 4. Biocompatibility (2  $\times$  2 mm chambers, initial volume 150 nL). (A) Growth and feeding. (i) A total of 250 NM18 cells in 800 nL were added to chambers by scanning, grown (24 h), fed (800 nL removed and 800 nL fresh medium added), and imaged. (ii) High-power view of selected region. (Magnification: 20x objective.) (iii) Cells were regrown (24 h) and reimaged, revealing that the cells grow normally. (B) Trypsinization and replating. (i) A total of 500 HEK cells in 500 nL are plated by scanning, grown (24 h), and imaged. (ii) After trypsinization, 1 µL is retrieved, and the chamber is reimaged; most cells have been removed. (iii) Retrieved cells were deposited in a new chamber, and this chamber was imaged. (iv) After growth (24 h), imaging confirmed successful cell transfer. (C) Cryopreservation. HEK cells are cryopreserved in grids for 3 wk at -80 °C, thawed, and regrown (24 h). Pinning lines and cells survive freezing and thawing. (D) Creating fluid walls around growing cells. (i) HEK cells  $(2 \times 10^6)$  are plated, grown (24 h), and imaged. (ii) A grid was prepared around these adherent cells, and the same region of the dish was reimaged. A fluid wall divides the field; cells 1, 2, and 3 remain in their original positions as others in the path of the stylus are moved. (E) C. elegans develops and responds to osmotic stress normally. (i) Frame from Movie S5 illustrating worms living in a grid. (ii) Single eggs in 500 nL of S medium were deposited manually in chambers and fed daily; after 5 d, the resulting adult was imaged. (iii) Individual trauma-sensitive worms (strain CB7317) in 500 nL of S medium were deposited manually in chambers, and 500 nL of medium plus bacteria with or without 600 mM NaCl was added; after 2 d, worms were imaged. This strain expresses GFP in response to osmotic stress, and worms exposed to 300 mM NaCl fluoresce green. (Insets) Fluorescence images.

lines are strong enough to withstand swimming forces: they are (Fig. 4 *E*, *i* and Movie S5). After pipetting individual eggs manually into chambers, followed by food (i.e., bacteria), eggs

developed normally into adults (Fig. 4 E, ii). Worms also responded as expected to stress. Individuals of a strain (29) that expresses the green fluorescent protein (GFP) in response to trauma were grown with or without 300 mM NaCl; osmotic stress induced GFP expression (Fig. 4 E, iii). These experiments confirm biocompatibility, plus the strength and excellent optical properties of fluid walls.

Reconfiguring Fluid Walls—Antibody Labeling. We next show that fluid walls can be removed and rebuilt in the same place (Fig. 5 A, i). After plating in grids, NM18 cells were fixed, and the dish was removed from the printer, emptied of FC40, and washed with PBS; this destroys wall integrity, although some FC40 can remain pinned along initial footprints. After mounting the dish back on the printer, fluid walls were rebuilt in their original positions, although they can be built in different places if required.

Because breaking/making fluid walls is so easy, we incorporated it into an immunolabeling workflow. NM18 cells were induced by transforming growth factor β1 (TGF-β1) to reorganize their cytoskeleton and undergo the epithelial-to-mesenchymal transition (EMT; ref. 30). NM18 cells in some chambers were treated with TGF-β1 and fixed, and then the workflow involved cycles of destruction of fluid walls (when cells in all chambers are batchwashed, and, in one case, permeabilized) and wall rebuilding (so different reagents can be added to selected chambers; SI Appendix, Fig. S5). After finally remaking walls, cells were imaged. Cell morphology and fluorescence were as expected: TGF-β1 increased actin bundling (detected by phalloidin labeling) and vimentin expression (detected by immunolabeling)—two markers of the EMT (Fig. 5 A, ii). This confirms the self-healing properties of walls (they were pierced and resealed eight times) and demonstrates that immunolabeling can be performed in grids. This approach can be scaled to screen hundreds of antibodies with considerable cost savings while accelerating the workflow.

Pinning Lines Can Withstand Detergents—RT-PCR. A common cellbased workflow involves lysing cells with detergents, followed by RT-PCR to measure mRNA levels. We expected this would be impossible, as detergents reduce interfacial tension and destabilize pinning lines. However, it turns out that footprints in chambers made with medium plus serum are stable (Fig. 5 B, i and ii). We illustrate this using HEK cells that encode a GFP reporter gene controlled by a promoter switched on by tumor necrosis factor alpha (TNF- $\alpha$ ): When exposed to the cytokine, cells fluoresced green (14) (Fig. 5 B, iii) and RT-PCR confirmed levels of GFP mRNA increase (Fig. 5B, legend). This shows that pinning lines can withstand destabilizing effects of detergents, presumably due to denaturation/cross-linking of serum components.

Example of a Complex Cell-Based Workflow-Genome Editing. We next performed a complex workflow: editing the Casp6 gene using CRISPR-Cas9 (31, 32) (Fig. 6A). Here, NM18 cells growing in grids were transfected with an empty plasmid or with plasmid 880 or 881, which encodes Cas9 and a puromycin-resistance gene plus a guide RNA targeting a region of Casp6. After adding puromycin, cells were grown for 4 d; puromycin killed untransfected cells (which lack the resistance gene), whereas roughly half of the transfected ones remained alive (indicative of high transfection rates; SI Appendix, Fig. S6B). Cells were then transferred from chambers to conventional plates and expanded, and single cells were plated in new chambers. Some chambers received one cell (Fig. 6B), others received two, and some received none (based on the Poisson distribution). Many single cells grew into colonies (cloning efficiency 71%; Fig. 6B legend). The printer then picked clones and transfered them to microcentrifuge tubes. After expanding clones conventionally, followed by DNA amplification and sequencing, clones were found to have deletions at appropriate target sites (Fig. 6C); they also no longer expressed

#### Breaking, remaking fluid walls; immunolabeling 500 cells (500 nl) grow break make grid remake (24 h) walls, (150 nl) walls fix wash grey boxes indicate fluid walls present ii phalloidin + DAPI anti-vimentin + DAPI - TGF<sub>β1</sub> + TGF<sub>β1</sub> - TGF<sub>β</sub>1 + TGF<sub>β</sub>1 Lysing cell walls: RT-PCR 500 cells $TNF\alpha$ lyse make grid grow RT-PCR grey box indicates fluid walls present

after lysis

before

Fig. 5. Rebuilding fluid walls and destroying cell walls  $(2 \times 2 \text{ mm chambers})$ The 6-cm dishes were mounted in positioning rings (tightly fitting circular sleeves enabling a dish plus ring to be removed from a printer and remounted so that walls can be rebuilt in the same place). (A) Breaking and remaking fluid walls during immunolabeling. (i) NM18 cells were plated by scanning, grown, and fixed; fluid walls were broken, all cells batch permeabilized with Triton X-100, and walls remade. Fluid walls survive fixation but are destroyed by emptying the dish of FC40 and washing with PBS (some FC40 remains pinned to the dish). After returning the dish to the printer and overlaying FC40, fluid walls are rebuilt. (ii) Cells respond to TGF-β1 as expected. The workflow involves cycles of making and destroying fluid walls (SI Appendix, Fig. S5); when walls are present, individual chambers are treated differently, and when walls are absent, all cells in all of the different parts of the dish are treated similarly. Cells were seeded by scanning in chambers, grown with (+) or without (-) TGF-β1, and then fixed. Next, walls were destroyed, fixative was washed away, cells were permeabilized, walls were rebuilt, phalloidin or anti-vimentin (both conjugated with Alexa 488) was added to selected chambers, walls were destroyed, cells were washed, walls were rebuilt, DAPI was added, walls were destroyed, cells were washed, walls were rebuilt, and phase and fluorescent images were collected. TGF-β1 induces actin bundling and increases vimentin expression (detected by phalloidin labeling and immunolabeling, respectively). (B) Lysing cells and RT-PCR. (i) HEK-293 reporter cells were plated by scanning in chambers and then grown (24 h), medium with (+) or without (–) TNF- $\alpha$ (10 ng/mL) was added, cells were regrown (24 h) to allow the cytokine to induce GFP expression before washing and lysis, and lysates were transferred from chambers to microcentrifuge tubes for assessment of levels of GFP mRNA by RT-PCR. (ii) Pinning lines survive lysis. (iii) Phase-contrast (Top) and fluorescence images (Bottom) show TNF- $\alpha$  induces GFP expression. RT-PCR also showed that TNF- $\alpha$  increases GFP mRNA levels 205-fold (lower and upper bounds: 174- and 241-fold), normalized relative to the control (lower and upper bounds: 0.63- and 1.6-fold; n = 4).

- TNFα

Casp6 protein (SI Appendix, Fig. S6C). Here, walls were pierced 16 times, and pinning lines survived treatment with transfection reagent. Significantly, fluid walls provide excellent optical clarity; there are no solid walls to obscure the view (in contrast to cloning in conventional flat microplates, where cells often sit

RT-PCR

increases GFP mRNA

shows

TNFα

levels

205-fold

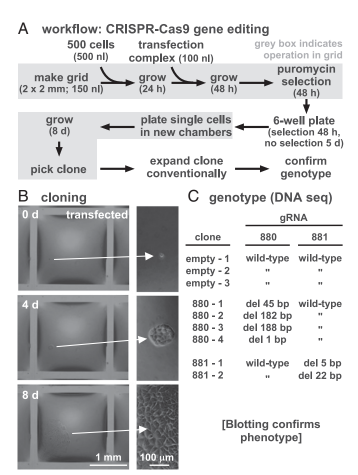


Fig. 6. A complex workflow: deriving a NM18 cell clone with a mutated Casp6 gene using CRISPR-Cas9. (A) Workflow. Cells were transfected with empty vector (encoding Cas9 plus puromycin resistance) or with plasmid 880 or 881, which additionally encode a guide RNA (gRNA) targeting different parts of Casp6. (B) Cloning. After transfection, puromycin selection (SI Appendix, Fig. S6), and expanding the cells conventionally, the printer delivered ~0.2 cells (by scanning) to each chamber; images show the single cell divides over 8 d. Initially, 42 of 256 chambers (40 expected from Poisson distribution) contained one cell; after 8 d, 30 had colonies (71% cloning efficiency). (C) Genotyping. After picking colonies (as in Fig. 4B), the printer delivered the cells to a microcentrifuge tube. Cells were then expanded conventionally, DNA was purified, the Casp6 region was amplified by PCR and cloned in bacteria, and genotypes were determined by sequencing. Clones derived from transfections with plasmids 880 and 881 have deletions (del) in expected regions, unlike those receiving empty vector. Immunoblotting confirmed the phenotype (SI Appendix, Fig. S6).

against surrounding walls and "edge effects" prevent them from being seen clearly). Moreover, when cells are seeded centrally in grids, fluid dynamics and chamber geometry ensure that few end up near pinning lines; preexisting (cell-free) medium is forced by incoming cell-containing medium toward edges (*SI Appendix*, Fig. S7). Consequently, users can have confidence in monoclonality. Since growing clones can be identified earlier, they can be picked sooner (in Fig. 6C, after 8 d, which contrasts with >14 d in conventional plates).

### Discussion

We describe a microfluidic platform for miniaturizing workflows in cell biology. Grids are made by covering the surface of a Petri dish with a thin layer of medium, overlaying FC40, and using a Teflon stylus to reshape the aqueous phase into an array of individual chambers; each chamber is isolated from neighbors by liquid walls of FC40 (Fig. 1 A-C). At the microscale, effects due to gravity and buoyancy are negligible, and the aqueous phase

remains pinned to the plastic; consequently, such grids can be used much like conventional microplates—liquids are simply pipetted through FC40 instead of air (Fig. 1D).

During fabrication, this platform has many advantages compared with other methods. It does not require a dedicated facility or specialized equipment beyond a syringe pump and an automated positioning system to drive the stylus; thus, fluid walls/ceilings are built accurately, reproducibly, and immediately by interfacial forces (e.g., our stylus generally traverses at 25 mm/s and builds 256 and 1,024 chambers in a 6-cm dish in ~2 and ~4 min, respectively; Movie S1). Building is scalable; for example, the array in Fig. 1*C* has a chamber density equivalent to a microplate with 393,216 wells. High-density grids are created efficiently (drawing twofold more lines yields fourfold more chambers), and ~90% of the surface area is available for cell culture in contrast to ~40% in a conventional 1,536-well plate (*SI Appendix*, Fig. S1*B*, calculated assuming fluid walls are 100 μm wide).

During use, there are additional advantages. First, pinning lines are stable, and fluid walls pliant and resilient. They withstand agitation (Movie S2), they morph above unchanging footprints when liquids are added/removed (Fig. 1 D, v), and self-heal when pipets are inserted through them (in Fig. 6, walls were pierced 16 times). Second, uniform volumes can be added to many chambers quickly and easily; because there are no solid walls, there is no need to raise/lower and/or wash the delivering pipet or to start/stop the pump. Instead, one pipet can scan through FC40 at constant speed and height, ejecting liquid continuously to feed chambers through transient liquid bridges. Such scanning can deliver 70 nL to each of 1,024 chambers in 90 s without detectable carryover (Fig. 3E and Movie S4). Third, all points in the aqueous phase are accessible. Fourth, walls confining the aqueous phase can be reconfigured during use; for example, new walls can be added (Fig. 4D) and existing ones destroyed and then rebuilt in the same place (Fig. 5A).

The platform has another significant advantage: It is biocompatible if used with the culture media and dishes (including coated ones; SI Appendix, Fig. S4 A and B) familiar to biologists, along with what is arguably the most bio-inert immiscible liquid available—FC40 (17-19). Cells (from bacteria to humans) grow normally in grids (Figs. 2A, 3C, and 4 A and B), worm eggs develop into adults (Fig. 4 E, ii), and cells/organisms respond as expected to stimuli (stress, puromycin, TGF-β, and TNF-α; Figs. 4 E, iii and 5). We also miniaturize the basic manipulations involved in mammalian cell culture (e.g., feeding, replating, cryopreservation, cloning; Figs. 4 A-C and 6B) and some widely used downstream workflows (fixation and immunolabeling; cell lysis and RT-PCR; and transfection and genome editing; Figs. 5 and 6). Because walls can be built around living cells of interest (Fig. 4D)—perhaps ones with characteristic morphologies or expressing particular fluorescent markers—selected cells can easily be isolated from others (for subsequent growth or analyses) without touching them. We also anticipate that grids will prove especially useful for cloning. With conventional microplates, single cells often sit against surrounding walls and cannot be imaged clearly because of edge effects. In contrast, fluid walls lack obscuring walls and thus yield excellent optical clarity. In addition, because cells are deposited centrally in chambers, and fluid dynamics and geometry ensure that few end up at the edges of footprints (SI Appendix, Fig. S7), users can be confident about which chambers contain only one cell. Moreover, clones can be picked sooner (e.g., after 8 d in Fig. 6B). Additionally, FC40 provides an extra barrier to contamination, isolating each chamber from others in one dish and from the external world (Fig. 24).

As with any platform, ours has limitations. First, an immiscible liquid is required to limit evaporation (as is generally the case wherever submicroliter volumes are handled). Second, liquids and surfaces must be matched to ensure that pinning lines are stable. We use the following approach to test new combinations

rapidly: A drop of aqueous liquid is placed on the substrate and overlaid with FC40, and most of the aqueous phase is removed. If pinning lines do not retract, then the combination can probably be used. Third, there are limits to upper and lower volumes that chambers can accommodate without change in footprint. These limits depend on advancing and receding contact angles; for example, with the angles of  $70^{\circ}$  and  $3^{\circ}$ , respectively, a 2 × 2 mm chamber has working volumes of 35 to 1,100 nL, and a  $140 \times 140$  µm chamber has working volumes of 9 to 270 pL (SI Appendix, Table S1).

In summary, we have developed a versatile platform for fabricating microfluidic patterns, exemplified by making arrays of square chambers with nanoliter volumes. These grids are used like conventional microplates, except that liquids are pipetted through FC40 instead of air. We anticipate that they will prove especially useful in cell biology, as they are made with materials familiar to users, can be incorporated into common workflows, and provide considerable savings in consumables.

#### **Materials and Methods**

General Reagents and Equipment. FC40 was purchased from Acota. It is bioinert, not found in regulatory lists of dangerous organic chemicals (33), and it should not be confused with the volatile chlorofluorocarbons that release the chlorine radicals destroying the ozone layer. If grids are to be kept for days, extra FC40 should be added when needed (14). Evaporated FC40 has had no untoward effects on any of many different cell types grown conventionally in the same incubator at the same time over 3 y. All other fluids and materials were from Sigma-Aldrich unless otherwise stated. Where indicated, aqueous drops contain water-soluble dyes (e.g., Allura Red, toluidine blue, resazurin).

Most grids were fabricated, and small volumes delivered to them, using an isoCell (lota Sciences Ltd). This is essentially a tool-head driven by a three-axis traverse and appropriate software; the tool-head holds a stylus—a Teflon rod (3.8-mm diameter) with a conical tip (angle at tip  $\sim$ 50°)—and a stainless steel dispensing needle (width 0.5-mm o.d.) connected to a syringe pump. This Teflon stylus was used to make all grids, except for ones shown in Fig. 1 C and D, v, Fig. 3, and SI Appendix, Fig. S1 B and C, where the rod was replaced by a Teflon tube (o.d.  $\sim$ 750  $\mu m$ ; Cole Parmer). As the dispensing needle is hydrophilic, liquid can run up the outside instead of into a chamber as wanted, and this both makes accurate delivery of small volumes unreliable (34) and increases carryover contamination when delivering one liquid to many chambers in a grid from one dispensing tube. Therefore, a hydrophobic sleeve (a piece of Teflon tubing) is included around the tip of the dispensing tube to prevent runback (Movie S4). The central workplace holds a 6-cm dish, plus microcentrifuge tubes containing reagents (often tissueculture media, 70% ethanol for sterilization; Movie S4). The 6-cm dish is placed in a positioning ring—a tightly fitting circular sleeve bearing a protrusion that ensures the dish plus ring can be mounted, removed, and

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remounted in the isoCell workplace in the same orientation. Where sterility is required, the printer is placed in a bio-safety cabinet, and sterile procedures used throughout (e.g., the stylus tip is sterilized with 70% ethanol, and the software includes cycles for aspirating 70% ethanol into the tube connected to the dispensing needle and ejecting the ethanol into a waste tube). When printing grids on rectangular flat (one-well) polystyrene microtiter plates (127.7 × 85.5 mm; Nunclon; Thermo Fisher Scientific), the 3D-traverse system described by Walsh et al. (14) fitted with a stylus (a Teflon tube) was used.

**Printing and Operation of Grids.** Grids were generally fabricated using the isoCell and 6-cm polystyrene tissue culture dishes [60-mm Falcon TC-treated cell culture dish (product #353002) and 60-mm Corning TC-Treated Culture Dish (product #430166)]; these dishes have internal diameters of  $\sim$ 5 cm. The 6-cm dishes were coated, where stated, with polylysine, Matrigel (Corning), fibronectin, laminin, or collagen (Cell Applications Inc.) by covering the bottom of the dish with the liquid coating using the concentrations, times, and temperatures suggested by the manufacturer (except for collagen, see below); removing most of the liquid coating to leave a thin film on the surface; and (without allowing the coating to dry) immediately adding DMEM plus 10% FBS and creating grids. For collagen, a 0.5× dilution (2.5 µg/cm<sup>2</sup>) was used, as the thicker coating prevented FC40 from wetting the surface. Glass substrates were either glass microscope slides/coverslips in 6-cm dishes, or 35-mm glass bottom dishes (No. 0; MatTek). DMEM plus 10% FBS was used to make all grids described, except those for use with worms; when the term "medium" is used in the context of mammalian cell culture, it should be assumed that serum is present unless stated otherwise. Typically, 1 mL DMEM plus 10% FBS is pipetted manually into a 6-cm dish, medium swirled around so the bottom is covered completely when the dish is horizontal, the dish tilted, and 0.9 mL medium removed and discarded. The bottom is now completely covered by a thin film of medium. A 3-mL aliquot of FC40 is manually pipetted into the dish so that a layer of FC40 covers the medium. Steps before overlaying FC40 are carried out quickly if the grid is to be used with cells so that the pH of the medium remains unchanged. After placing the dish in the work area on the printer, the system uses in-built software to "home" the tool head, select the stylus, "draw" lines to create the grid (stylus speed typically 25 mm/s), deselect the stylus, and go "home". Additionally, the software can select a pipet (the "pen"), deliver nanoliter volumes to selected chambers (by switching on and off an in-built syringe pump), deselect the stylus, and go "home".

Detailed methods for individual figures are described in SI Appendix, Materials and Methods.

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Soitu et al. PNAS | vol. 115 | no. 26 | **E5933** 

Title: Microfluidic chambers using fluid walls for cell biology

SUPPLEMENTARY INFORMATION

SUPPLEMENTARY MATERIALS AND METHODS

**Methods used individual Figures** 

For the 32 x 32 grid in **Figure 1B**, 70 nl medium + yellow or blue dye were delivered to selected chambers to create the wanted pattern. For **Figure 1C**, the grid was made in a 6 cm dish using a stylus constructed from a Teflon tube. For **Figure 1E**, medium was added to chambers on each side of the FC40 wall to increase wall—chamber contrast.

For Figures 2A and 3C, Escherichia coli (HB101) were inoculated directly from frozen stocks into Luria-Bertani broth (LB), and used as described. For Figure 3C, 300 nl LB medium  $\pm \sim 20,000$  E. coli was added to every second chamber, and 500 nl LB was added (ejection rate 8.3  $\mu$ l/s; as in Fig. 3A) to each chamber (minimum tip—dish-surface distance  $\sim 0.6$  mm; maximum chamber height increases from  $\sim 0.34$  to  $\sim 0.63$  mm). In Figure 3E, the dispensing tip scans (15 mm/s) along a serpentine scanning path as it delivers  $\sim 70$  nl medium + blue dye per chamber (ejection rate 0.7  $\mu$ l/s; maximal chamber height increases from  $\sim 0.15$  to  $\sim 0.34$  mm). In Figure 3F, 300 nl media  $\pm 9,000$  fluorescent beads (1  $\mu$ m diameter) are added to each chamber as indicated, and then 500 nl media is now added by scanning (traverse rate 1.67 mm/s, ejection rate 0.4  $\mu$ l/s) to every chamber (tip—dish-surface distance  $\sim 0.65$  mm; maximum chamber height increases from  $\sim 0.34$  to  $\sim 0.63$  mm). For a 16 x 16 grid, the standard deviation found after delivering 280 nl by scanning to each chamber (measured as in Fig. S2) was found to be  $\pm 27$  nl, (and for 32 x 32 grid it was found to be 10 nl after delivering 60 nl). When larger volumes are delivered (up to the maximum final volume given in Table S1), the uncertainty in volume delivered should remain the same as the main source of variation results from differences in initial chamber height. [Figures 3-6 and S4 illustrate delivery of 70-1,000 nl by scanning to chambers of different size.]

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For **Figure 4A**, NM18 cells (1) were used. For **Figure 4B**, 500 HEKs in 500 nl are plated by scanning, grown (24 h), and imaged. Next, 800 nl medium is removed, cells washed with PBS (1  $\mu$ l added, 1  $\mu$ l removed after 1 min), 1  $\mu$ l trypsin (TrypLE, Gibco #12563011, Gaithersburg, MD) added, the dish incubated (5 min; 37°C) and transferred to a printer, 1  $\mu$ l retrieved, and the chamber re-imaged. Finally, the retrieved 1  $\mu$ l (containing suspended cells) is mixed with 10  $\mu$ l fresh medium (to neutralize trypsin) in a micro-centrifuge tube, 1  $\mu$ l of the mixture deposited in a new chamber, this chamber imaged, and – after growth (24 h) – re-imaged. For **Figure 4C**, 500 HEK cells in 500 nl were seeded in grids in DMEM + 20% FBS + 10% DMSO, and immediately put in a -80°C freezer in an expanded polystyrene box surrounded by paper tissues, stored for 3 w, chambers thawed by placing the dish at room temperature, chambers washed (700 nl removed, 700 nl fresh medium added) 3 times (some cells were lost), and cells re-grown (24 h).

For experiments involving *C. elegans* (strain CB3717<sup>29</sup>), worms were grown on agar plates using *E. coli* OP50 as food (2). The strain used has been genetically modified to express GFP in response to trauma; it also expresses RFP constitutively, but results using this marker are not illustrated here. Grids were made with S medium + 0.03 mg/ml BSA (added to stabilized pinning lines), and worms in them grown at room temperature. Prior to collection of still images, dishes were placed at -20°C for 10 min to temporarily immobilize worms, and sometimes worms were washed with fresh medium to remove feces and bacteria. For **Figure 4Eii**, single eggs in 0.5  $\mu$ l S medium were deposited manually in chambers, 0.5  $\mu$ l medium + bacteria added, and 0.8  $\mu$ l removed daily and replaced with an equal volume of fresh S medium + bacteria; after 5 d, the resulting adult was imaged. For **Figure 4Eii**, individual worms in 0.5  $\mu$ l S medium were deposited manually in chambers, and 0.5  $\mu$ l S medium + bacteria  $\pm$  600 mM NaCl added; after 2 d, worms were imaged. Average fluorescence intensities/pixel over 5 worms treated  $\pm$  NaCl for 2 d were 1  $\pm$  0.1 and 4  $\pm$  1.5 arbitrary units ( $\pm$  SD).

For **Figure 5Ai**, NM18 cells (500 in 500 nl) were plated by scanning in each chamber (16 x 16 grids, 2 x 2 mm chambers, 150 nl initial volume) in a 6 cm dish mounted in a positioning ring, grown (24 h), and the dish remounted on the isoCell. After removing 650 nl, cells were fixed by adding 500 nl 4% paraformaldehyde in PBS, and one chamber imaged. After fixation (15 min), the dish (still in the positioning ring) was removed from the isoCell, and fluid walls destroyed by manually emptying the dish of FC40 and washing with PBS. The

same chamber was reimaged. The dish was then remounted on the isoCell, FC40 added, and the grid recreated in the same place as before. Finally, the same chamber was imaged. For Figure 5Aii, NM18 (500 in 500 nl) were plated by scanning in each chamber (16 x 16 grids, 2 x 2 mm chambers, 150 nl initial volume) in a dish mounted in a positioning ring, grown in a CO2 incubator (24 h), the dish returned to the isoCell, 400 nl media ± TGFβ1 (Peprotech, London, UK) added (final concentration 5 ng/ml), cells regrown (24 h), fed on the isoCell (by removing 800 nl medium, and replacing 800 nl media ± TGFβ1), regrown, refed on the isoCell (by removing 800 nl and adding 800 nl media ± TGFβ1, and cells regrown (24 h). After fixation and destruction of fluid walls (as above), cells in all chambers were batch permeabilized manually by adding PBS plus 0.1% Triton X100 to the dish. After washing with PBS to remove detergent, the dish was remounted in the IsoCell, FC40 added, and fluid walls rebuilt in their original places. Next, 500 nl PBS ± phalloidin-iFluor 488 (1/1000 dilution; Abcam; Cambridge, UK) or ± an anti-vimentin antibody conjugated with Alexa 488 (1/800 dilution; Cell Signaling Technology, Leiden, The Netherlands) were added to selected chambers, and the dish incubated (4°C, 15 h). Next, fluid walls were destroyed, the dish washed with PBS, fluid walls remade in the same place (all as before), 500 nl DAPI added (final concentration 1 μg/ml) and the dish incubated (20°C, 30 min). Finally, fluid walls were destroyed (as above), the dish washed with PBS, walls remade to facilitate identification of particular cells, and individual chambers imaged.

For Figure 5B, HEK-293 reporter cells (NF-kB/293/GFP-Luc<sup>™</sup> Transcriptional Reporter Cell Line; System Biosciences, catalogue number TR860A-I) were grown as recommended by the manufacturer in DMEM plus 10% FBS. They encode a GFP gene under the control of the minimal cytomegalovirus promoter downstream of four copies of the NF-κB consensus transcriptional-response element, and GFP expression can be induced by TNFα. 500 cells in 500 nl were plated by scanning in chambers, grown (24 h), 400 nl medium ± TNFα (10 ng/ml; Peprotech, London, UK) added, cells regrown (24 h) and imaged, washed with PBS (by removing 1 μl medium, adding 0.8 μl PBS, and retrieving 0.8 μl), and lysed by adding 0.5 μl RNA lysis buffer (Zymo Research, Orange, CA); the same chamber as before was now re-imaged. Next, levels of GFP mRNA were assessed by real-time RT-PCR. 0.5 μl cell lysate was transferred from a chamber to a microcentrifuge tube containing 100 μl RNA lysis buffer and levels of GFP mRNA assessed by RT-PCR. Total cell RNA was isolated using the Quick-RNA MicroPrep kit (Zymo Research), and cDNA synthesis and RT-PCR performed

using the SuperScript® III Platinum® One-Step qRT-PCR kit (Thermo-Fisher Scientific) with DNase treatment, according to the manufacturer's instructions. Gene-specific primers for GFP were synthesized commercially (Sigma-Aldrich; forward primer GATGGGCTACGGCTTCTACC, reverse primer GTACTTCTCGATGCGGGTGT).

Each mRNA value was normalized to levels of transcripts of the house-keeping gene, 5s rRNA (forward primer TACGGCCATACCACCCTGAA, reverse primer GCGGTCTCCCATCCAAGTAC). Cycling profiles (Eppendorf realplex² master cycler; melting-curve analysis) were: 50°C for 5′, then 95°C for 5′, followed by 40 cycles of 95°C for 15″ and 60°C for 30″, and finally 40°C for 1′. Results are reported as relative gene expression. The fold-change in gene expression relative to the control was calculated by the 2<sup>-ΔΔCT</sup> method (3).

For Figure 6, the Casp6 gene in NM18 cells was edited using the CRISPR-Cas9 system and plasmids 880 and 881; transfections involving no plasmid, and plasmids lacking inserts, provided controls. The guide RNA in plasmid 880 targeted the sequence TCCATCTTGTACTGCTCGGCTGG, and that in 881 targeted CGTTGGTGCCCCGCCTCTCTGGG. These plasmids were constructed by inserting oligonucleotides into pSpCas9(BB)-2A-Puro (PX459) V2.0 (Addgene, Cambridge, MA). After making a grid (16 x 16, chambers 2 x 2 mm, 150 nl), cells (500 in 500 nl) were delivered by scanning (Fig. 3D) to each chamber, and grown (24 h) in a CO<sub>2</sub> incubator. Transfection complexes were formed in a micro-centrifuge tube by manually pipetting 100 μl serum-free DMEM plus 1.5 μl TransIT-X2 (Mirus Bio LLC, Madison, WI) and ~2 μl plasmid DNA (1 μg), briefly vortexing, and then transferring 50 μl into a well on a glass slide (Nexterion Slide Glass B MPX 16N; Schott, Jena, Germany) contained in a rectangular container (custom made on a 3D printer) that fits into the station for micro-centrifuge tubes at the front of the isoCell. The slide in the container was now overlaid with FC40 and incubated (30 min, room temperature) to allow complexes to form. Next, the container plus slide was transferred to the isoCell. After placing the dish containing cells on the isoCell, the printer added 100 nl complexes to each chamber in the grid followed by 250 nl medium (so total chamber volume becomes 1 μl). Cells were regrown (48 h), fed with medium plus puromycin (the isoCell removed 800 nl media and added 800 nl fresh media ± 1 µg/ml puromycin to kill wild-type cells and select for ones expressing a transfected puromycin-resistance gene), grown in a CO<sub>2</sub> incubator (24 h), and refed ± puromycin (as above). After further growth (24 h; Fig. S6B provides images of cells growing in grids ± puromycin at this stage), the isoCell retrieved cells from chambers containing puromycin (as Fig. 4B), and transferred them to micro-centrifuge

tubes. Cells were now manually transferred to conventional 6-well plates, grown (48 h) in medium + puromycin, and regrown (5 d) without selection. Cells were harvested, counted, and diluted to 400 cells/ml, and the isoCell then transferred ~0.2 cells in 500 nl into each chamber in a new 16 x 16 grid (2 x 2 mm, 150 nl) followed by 200 nl medium containing serum (giving a total of 1 μl). Chambers containing one cell were imaged and recorded, the grid transferred to a CO₂ incubator for 8 days, and chambers re-imaged. Cells derived from a single clone were then transferred from a chamber to a micro-centrifuge tube by the isoCell, expanded conventionally, and genomic DNA purified and amplified by PCR, followed by TOPO TA cloning (Thermo Fisher Scientific and Life Technologies, Carlsbad, CA) and sequencing. Phenotypes of edited clones were also monitored by immuno-blotting (Fig. S6). Cells grown conventionally were harvested in Pierce RIPA buffer (25 mM Tris-HCl, pH 7.6, 150 mM NaCl, 1% NP-40, 1% sodium deoxycholate, 0.1% SDS; Fisher Scientific, Loughborough, UK) containing protease inhibitor cocktail (Fisher Scientific), protein concentrations determined using the Bradford method and kit (Sigma-Aldrich, Dorset, UK), and whole cell lysates resolved by 12% SDS-PAGE and transferred to ImmunoBlot PVDF membranes (Bio-Rad Laboratories, Herts, UK). Next, membranes were incubated overnight with primary antibody in 5% w/v non-fat dry milk, 1X TBS plus 0.1% Tween 20 at 4°C with gentle shaking. Antibodies used were rabbit polyclonal anti-Caspase-6 (Cat# 9762) and anti-rabbit IgG, HRP-linked antibody (Cat #7074; Cell Signaling Technology, Beverly, MA). After stripping membranes with Restore Plus Membrane Stripping Buffer (Thermo Scientific), membranes were re-probed with mouse monoclonal IgG GAPDH (G-9) (sc-365062) and goat anti-mouse IgG-HRP (sc-2005 F0115) from Santa Cruz Technology (San Diego, CA).

## **Imaging**

Images of dishes were taken using a digital SLR camera (Canon 600D). Bright-field, phase-contrast, and fluorescence microscope images of grids with and without cells were collected using either a camera (AxioCam MRm) attached to a microscope equipped for live-cell imaging (Zeiss Axioskop 40; Olympus LWD A20 PL 20x lens) or a zoom lens and digital SLR camera (Nikon D7100 DSLR) connected to an epi-fluorescent microscope (Olympus IX53; 1.25X, 4X, 10X, 25X objectives) with translation stage and overhead illuminator (Olympus IX3 with filters). For **Figure S2**, the objective used was an Olympus UPlanFl 4x (0.13 NA, 17 WD). Fluorescence intensities in **Figure 4Eiii** were quantified using ImageJ; for each image, background was

subtracted, the area containing the specimen selected, and the average intensity per pixel determined and normalized relative to the control (non-treated specimen).

## **SI References**

- 1. Deckers M, et al. (2006) The tumor suppressor Smad4 is required for transforming growth factor beta-induced epithelial to mesenchymal transition and bone metastasis of breast cancer cells. Cancer Res 66:2202–2209.
- 2. Stiernagle T (2006) Maintenance of C. elegans. WormBook, ed. The C. elegans Research Community, WormBook, 10.1895/wormbook.1.101.1.
- 3. Livak KJ, Schmittgen TD (2001) Analysis of relative gene expression data using realtime quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods 25:402–408.

## **SUPPLEMENTARY TABLES**

**Table S1.** Some characteristics of conventional microplates and grids.

Microplates (solid walls)				Grids (fluid walls)			
well number	inter-well spacing (mm)	working volume (μl)		inter-chamber spacing (mm)	working volume (μl)		example
		min	max		min	max	
96	9	~50	~360	9	3.6	113	
384	4.5	~15	~100	4.5	0.44	14	
1,536	2.25	~3	~10	2.25	0.05	1.6	Fig S1A
				2	0.035	1.1	Fig 2A
				1	0.004	0.12	Fig 1B
393,216	0.140			0.140	9 x 10 <sup>-6</sup>	270 x 10 <sup>-6</sup>	Fig 1C

Values for microplates are from manufacturers' data sheets. For grids, inter-chamber spacing includes the width of a 100  $\mu$ m FC40 wall except for the highest-density one where the width was 20  $\mu$ m; minimum (min) and maximum (max) working volumes – which are proportional to the cube of the characteristic length scale (the width of the chamber) – are conservatively estimated by scaling relative to a sessile drop with circular footprint with diameter equal to chamber width using  $\vartheta_A$  of 70° (serum-free medium) and  $\vartheta_R$  of 3° (both values from ref 14 for serum-free media). Maximum and minimum values for media plus serum are higher and lower than indicated, respectively.

## **SUPPLEMENTARY FIGURES**

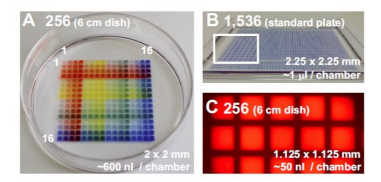


Figure S1. Grids of different sizes are made easily.

**A.** 16 x 16 grid which could be used for a combinatorial drug screen. 16 different dyes (representing 16 different drugs) were added in all pairwise combinations to 2 x 2 mm chambers (200 nl/chamber). Dye 1 (400 nl) was delivered to each chamber in row 1 and column 1, dye 2 to row 2 and column 2, and so on. Consequently, chambers on the diagonal (top-left to bottom-right) contain one dye, whilst others contain two in all pair-wise combinations (duplicates on each side of the diagonal). The stability of this grid is illustrated in **Movie S2**.

**B.** A 48 x 32 grid made in a 1-well microplate of standard area (i.e., a rectangular polystyrene Petri dish, 127.76 x 85.48 mm). Center-to-center spacings of chambers are 1/4 of those in a 96-well microplate, and like those in a 1,536-well microplate. After printing, blue dye was added to each chamber to aid visualization. Inset: 2x zoom.

**C.** Fluorescence micrograph of 16 x 16 grid with 256 chambers made in a 6 cm dish using medium + blue dye (which fluoresces red). Center-to-center spacings are 1/8 of those in a 96-well microplate.

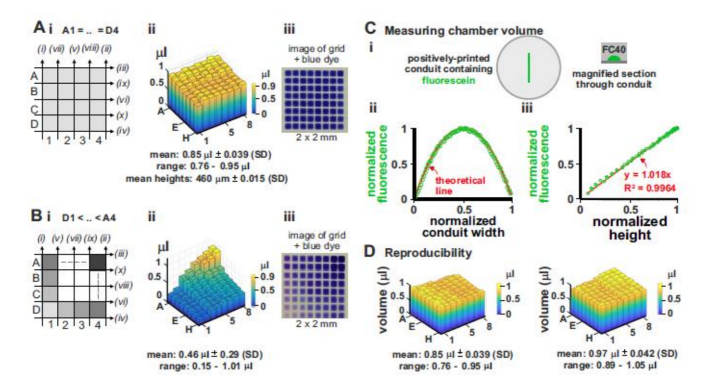


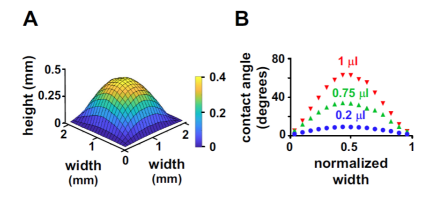
Figure S2. Creating chambers with the same or different starting volumes by altering stylus path (chambers 2 x 2 mm). Two dishes are covered with media + fluorescein, FC40 added, four 20 mm lines drawn to create one central square in each dish (volume  $\sim 10~\mu$ l), and 50 or 32  $\mu$ l (in **A** and **B**, respectively) pipetted into each square (as **Fig. 2Eii**). The stylus tip is a cone that displaces equal amounts of liquid to each side as it moves across the substrate at 25 mm/s, and it creates grids by symmetrical or asymmetrical division. The fluorescence intensity/chamber (which reflects volume) is measured by microscopy.

**A.** Symmetrical division. (i) Stylus path (arrows – stylus direction; roman numerals – drawing sequence). Lines (i)-(iv) create a square, and then (vi)-(x) a 4 x 4 grid as successive lines pass through centers of pre-existing aqueous bodies. Additional lines (not shown) create an 8 x 8 grid in which chambers have similar volumes (volume A1 = B1 ... C4 = D4). (ii) Measurement of fluorescence intensity/chamber shows chambers contain similar volumes. (iii) Image of grid created using blue dye instead of fluorescein.

**B.** Asymmetrical division. Chambers with different volumes created as in (**A**) except the stylus follows a different path. (**i**) Lines (*i*)-(*iv*) create a square (as before), but (*v*)-(*x*) scan progressively from lower-left to upper-right to divide pre-existing aqueous bodies asymmetrically (except for last two); this progressively shifts the center of mass of the larger body of media up and to the right, and chamber volume D1 < C1 ... B4 < A4. Additional lines (not shown) create an 8 x 8 grid. (**ii**) Measurement of fluorescence intensity/chamber shows chambers contain different volumes. (**iii**) Image of grid created using blue dye instead of fluorescein; chambers at lower left clearly contain less dye.

**C.** Method used to measure chamber volume. (i) A straight 30 mm conduit containing media + 10% FBS supplemented with fluorescein is positively printed in air on a 6 cm dish (0.5 mm diameter pen) using a syringe pump connected to a hollow stainless-steel dispensing needle, and immediately overlaid with FC40 to prevent evaporation (14). As the cross-section of this aqueous conduit is shaped like the cap of a sphere, it has a known radius, maximum height, and width. (ii) Normalized fluorescence intensity across the conduit. Using a fluorescence image of a central section of the conduit, the fluorescence intensity (arbitrary units per 50 x 50 pixels across conduit) is measured (green circles). The red curve is the one expected if the conduit section is shaped like the cap of a sphere, and if intensity is proportional to cap height. All image processing steps were carried out in Matlab. (iii) Relating fluorescence to height. Data from (ii) is replotted by relating intensity to height.

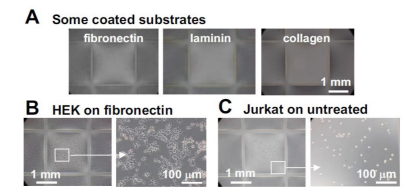
**D.** Two grids were made by symmetrical division, and chamber volumes calculated (as in **C**). Chambers have similar fluorescence intensities – and so volumes. Left-hand plot prepared using data from (**Aii**).



**Figure S3.** Chamber shape.

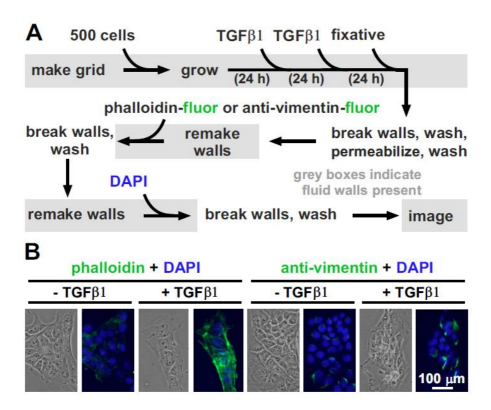
**A.** Overall shape. 2 x 2 mm chamber was made (as in **Fig. 2E**) using medium plus fluorescein; its volume was 850 nl (measured as in **Fig. S2C**). The (maximum) contact angle (measured as in **B**) at the midlines of edges is 52°. A droplet of the same volume with a spherical footprint of 2 mm diameter has a contact angle of 54°; consequently, calculations based on the use of droplets with spherical footprints in **Table SI** provide good estimates of working volumes.

**B.** Contact angles along a pinning line of a 2 x 2 mm square chamber holding 0.2, 0.75 and 1  $\mu$ l medium plus fluorescein. After preparing height profiles as in (A), average heights of units of 150 x 150 pixels along a chamber edge were determined (2,250 pixels per 2 mm edge), and contact angles given by the middle of the edge of that unit (zero height) and the middle of each unit (average height) calculated (using Matlab).



**Figure S4.** Phase-contrast images of grids prepared on dishes with different coatings, and using adherent and non-adherent cells (delivered by scanning).

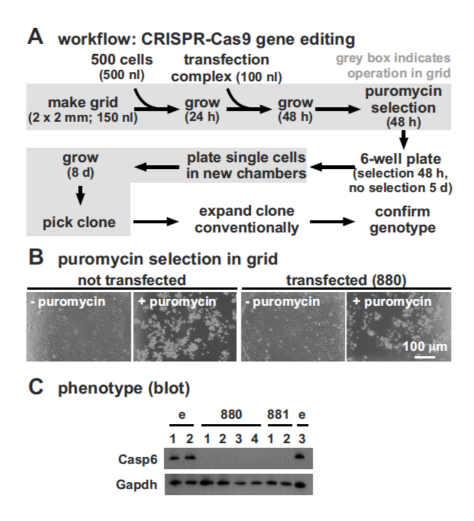
- **A.** Pinning lines are stable on dishes coated with various substrates (2.25 x 2.25 mm chambers; no cells added).
- **B.** Adherent HEKs grow normally on fibronectin. 800 cells in 1  $\mu$ l were delivered to each chamber, grown (24 h), and imaged (zoom shows region of chamber).
- **C.** Non-adherent Jurkat cells (an immortalized line of human T lymphocyte cells) grow in suspension. 660 cells in 1  $\mu$ l were delivered to each chamber, grown (24 h), and imaged (zoom shows region of chamber).



**Figure S5.** Breaking, remaking fluid walls during immuno-labeling. This Figure provides the workflow used in Figure 5Aii.

**A.** The workflow. NM18 are plated by "scanning", grown, treated twice with  $\pm TGF\beta 1$ , and fixed. Fluid walls were now broken (as in Fig. 5Ai), and all the fixed cells in the dish simultaneously batch-washed and permeabilized with Triton X100. Fluid walls were now remade, and phalloidin-Alexa488 or anti-vimentin-Alexa488 added to individual chambers. Next, fluid walls were re-broken, all cells in the dish simultaneously batch-washed. Walls were now remade, and DAPI added to chambers. After breaking walls, and batch washing, walls were remade, and central areas of different chambers imaged.

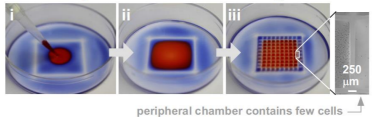
**B.** Phase and fluorescent images show TGF $\beta$ 1 induces actin bundling and increases vimentin expression (detected by phalloidin- and immuno-labeling respectively). These images are reproduced from Figure 5Aii for convenience.



**Figure S6.** Data supporting the complex workflow illustrated in Figure 6 (deriving a NM18 clone with a mutated *Casp6* gene using CRISPR-Cas9).

- **A.** Workflow (reproduced from Fig. 6A for convenience).
- **B.** Puromycin selection. Views of chamber centers after puromycin selection (48 h). Untransfected cells are refractile and rounded (indicative of cell death); in contrast,  $\sim$ 50% transfected cells remain non-refractile and attached (and viable).
- **C.** Phenotyping picked clones. After expanding the picked clones conventionally, immuno-blotting confirms clones derived from transfections with 880 and 881 lose Casp6 (loading control Gapdh), unlike those with empty vector (e).

## A Added dye / cells remain in center of large chamber



# B Added cells remain in center of small chamber

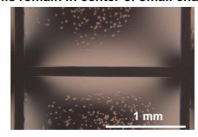


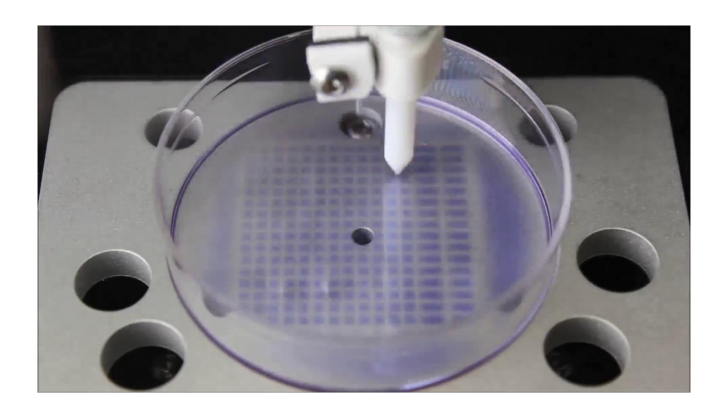
Figure S7. Cells tend to be found centrally in chambers (6 cm dishes).

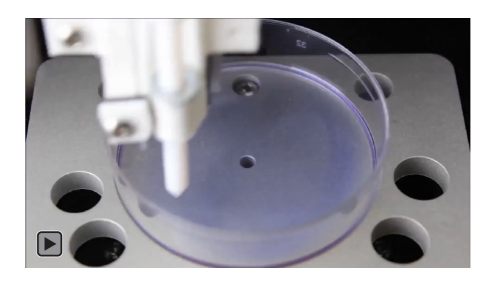
**A.** Large chamber. Medium plus blue dye is added to a dish, most medium removed to leave a thin film covering the bottom, and one  $28 \times 28$  mm printed in the center of a dish (as **Fig. 2E**). (i)  $30 \mu$  red dye is now pipetted into the middle of the square. (ii) Red dye spreads to push blue dye to the edges of the square. (iii) After splitting the central square into a  $10 \times 10$  grid (2.25 x 2.25 mm chambers), peripheral chambers mainly contain blue dye, and those in the central  $8 \times 8$  grid mainly red dye. Inset: when blue dye is omitted, and 20,000 HEKs replace red dye, few cells are found in a peripheral chamber.

**B.** Cell distributions in small chambers. NM18 were seeded in 2 x 2 mm chambers, and a bright-field image collected immediately. There are no cells at the edge of the chamber close to the pinning line.

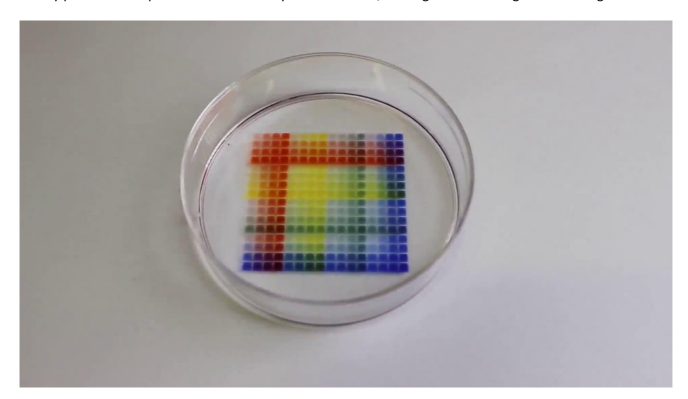
## **SUPPLEMENTARY MOVIES**

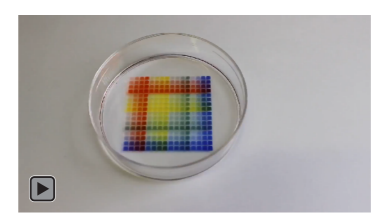
**Movie S1.** Printing a 16 x 16 grid in a 6 cm dish (the movie runs in real time for 4 s, and then is speeded up 10x for the remainder). The dish initially contains a thin layer of medium plus blue dye overlaid with FC40, and the white Teflon stylus draws a grid.



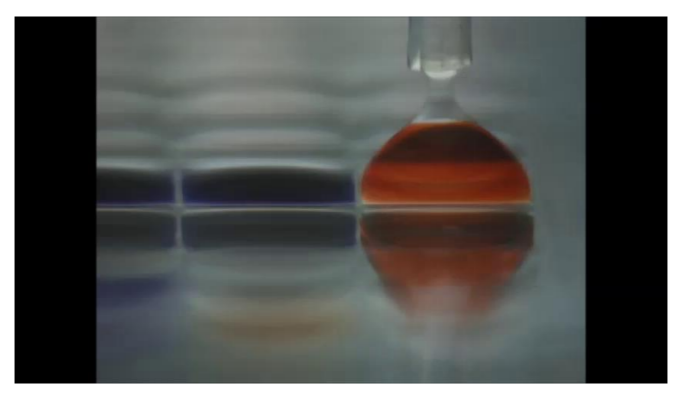


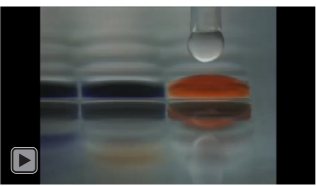
**Movie S2.** A grid overlaid with FC40 can survive violent agitation (the movie runs in real time). The grid was printed on a 6 cm dish, 16 different dyes delivered in all pairwise combinations to chambers (see **Fig. S1A**), the dish placed on a shaker, and the lid of the dish removed. The movie begins when the shaker was started; as the speed of shaking increases, waves form in the FC40 and increase in height. Footprints of chambers remain unchanged throughout. Of course, if chambers made with serum have contact angles of >90° and are closely packed and separated from others by thin fluid walls, shaking can cause neighbors to merge.



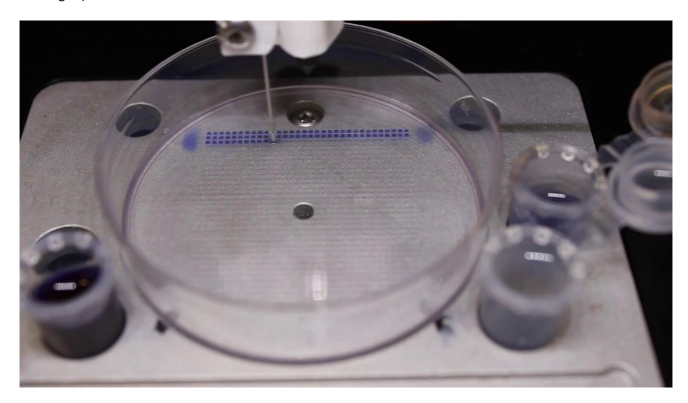


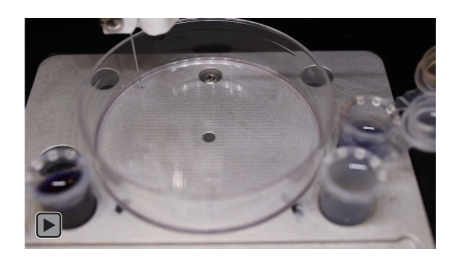
Movie S3. Adding fluid via a liquid bridge to a chamber (the movie runs in real time, and frames from it are reproduced in Fig. 3B). A 16 x 8 grid (2.25 x 2.25 mm chambers, rectangular dish) was made using medium, and red and blue dyes delivered to chambers. Between 0-13 s, medium is continuously ejected from the pipet tip, which is immersed in FC40 and positioned above the recipient chamber containing red dye. The pipet is a hydrophilic tube with a hydrophobic sleeve at the end. At 0 s, ejected medium begins to form a bubble at the end of the tip. By 9 s, the bubble has grown sufficiently to touch the surface of the chamber, and a liquid bridge forms. Note that there is no detectable transfer upwards of red dye through this bridge. By 11 s, the bridge has broken, and no red dye can be detected in the bubble on the tip. After 13 s, a new bridge has formed, and again no dye is being transferred upwards. The tip will now move to a new position above another chamber, and deliver medium to it. This movie indicates that essentially no red dye will be transferred between chambers during this process.





**Movie S4.** Delivering liquid to a grid by scanning along a serpentine path (the movie runs in real time, it is truncated after delivering to 3 rows, and frames from it are reproduced in **Fig. 3E**). A 32 x 32 grid (initially with  $\sim$ 40 nl per 1 x 1 mm chamber) was made in a 6 cm dish using medium. The movie begins as a hollow stainless-steel dispensing needle with a Teflon sleeve at its tip dispenses blue dye as it scans along a serpentine path delivering  $\sim$ 70 nl to each of 1,024 chambers in 90 s (some dye is added to the left and right of the grid).





**Movie S5.** Fluid walls withstand forces exerted by living worms at room temperature (a frame from this movie is illustrated in **Fig. 4Ei**). An 8 x 8 grid (chambers 2 x 2 mm) was made using S medium + 0.03 mg/ml BSA, bacteria added as food, dishes left at room temperature, worms washed in S medium and transferred to new chambers, and living worms imaged.

