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Teach your microscope how to print: low-cost and rapid-iteration microfabrication for biology[†]

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The application of traditional microfabrication techniques to biological research is hindered by their reliance on clean rooms, expensive or toxic materials, and slow iteration cycles. We present an accessible microfabrication workflow that addresses these challenges by integrating consumer 3D printing techniques and repurposing standard fluorescence microscopes equipped with DMDs for maskless photolithography. Our method achieves micrometer-scale precision across centimeter-sized areas without clean room infrastructure, using affordable and readily available consumables. We demonstrate the versatility of this approach through four biological applications: inducing cytoskeletal protrusions *via* 1 μ m-resolution surface topographies; micropatterning to standardize cell and tissue morphology; fabricating multilayer microfluidic devices for confined cell migration studies; imprinting agar chambers for long-time tracking of *C. elegans*. Our protocol drastically reduces material costs compared to conventional methods and enables design-to-device turnaround within a day. By leveraging open-source microscope control software and existing lab equipment, our workflow lowers the entry barrier to microfabrication, enabling labs to prototype custom solutions for diverse experimental needs while maintaining compatibility with soft lithography and downstream biological assays.

Introduction

Microfabrication technologies using photolithography and soft lithography have enabled researchers to build cellular environments with micrometer precision. By shaping and patterning the geometry, topology and composition of the extracellular space at a precision that matches the intrinsic scale of cells, these technologies provide a powerful tool to study interactions between cellular systems and their environment.^{1–3} Microfabricated structures can be used in a wide array of downstream applications: to confine bacteria,⁴ eukaryotic cells,^{5–7} or microscopic animals;⁸ fabricate microfluidic devices⁹

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^b Graduate School for Cellular and Biomedical Sciences, University of Bern, Switzerland e.g. to study 3D migration,^{10,11} or mimic complex geometries to grow more realistic organ¹² and cancer^{13,14} models *in vitro*; to pattern surfaces to study 2D cell migration,15-19 measure forces exerted by cells onto their substrate,²⁰ and homogenize cell morphologies.²¹⁻²⁴ Furthermore, lab-on-a-chip devices use microfabrication to miniaturize systems, enabling massive parallelization.²⁵ Photolithography techniques for biological research were originally adapted from the semiconductor industry.^{26,27} These methods were designed to meet the submicrometer precision requirements of electronics manufacturing, whereas many biological applications do not require such precise spatial resolution. Bottlenecks in microfabrication processes are that they require clean room access^{28,29} and expensive single-purpose hardware such as mask aligners. Further, the required reagents and substrates can be costly, toxic, and difficult to source. The continued reliance on high-precision techniques for biological research has led to unnecessary complexity, creating barriers to accessibility.^{30,31} Traditionally photolithography uses a high resolution photomask that contains the design. Although commercially available photomasks are relatively inexpensive, their long production times hinder iterative design cycles. Recent maskless approaches and efforts to simplify protocols are improving iteration speed and accessibility of these

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methods.^{18,30,32–35} This approach has been commercialized by Alvéole is widely used.

In contrast, 3D printing has significantly expanded access to additive manufacturing and rapid prototyping, proving useful in many laboratories.^{36,37} A class of 3D printers recently becoming available for the consumer market, commonly known as "resin printers", employs photolithographic techniques to achieve x/y resolutions down to 50 µm,³¹ but they still fall short of producing geometric features at the cellular scale.

By combining elements of semiconductor microfabrication with consumer 3D printing, we present a

simplified microfabrication protocol tailored for biological applications, achieving micrometer precision at the centimeter scale while reducing time and procedural complexity. We drastically reduce the cost of consumables by replacing the commonly used SU-8 photoresist with 3D printing resin, and silicon wafers with standard microscope slides. Our approach repurposes an existing microscope setup used for targeted photostimulation as a maskless microfabrication system, streamlining the process from concept to fabricated structure within a day. Compared to commercial solutions, our approach relies on open-source software and does not require specific proprietary hardware



Fig. 1 Microfabrication method A protocol overview. A mask is designed using computer graphics software. Using a microscope, UV light is projected onto a glass slide coated with UV curable resin. The uncured resin is then washed away and a PDMS cast is formed from the 3D printed structure. B Our method can be applied across biological scales, from subcellular to cellular. Subcellular: A REF52 cell forming protrusions into microfabricated wells (Fig. 2, *lifeact::mNeonGreen*). Cellular: a single NIH3T3 cell on a circular fibronectin micropattern (Fig. 3, black: *ERK-KTR:: mRuby2*; blue: *H2B::miRFP670*). Multi cellular: sparsely seeded MCF10A cells on a circular fibronectin micropattern (black: *ERK-KTR::mRuby2*; blue: *H2B::miRFP670*). 2D gastruloid: human embryonic E9 stem cells growing on a circular Matrigel pattern (Fig. 5, purple: *eft-3p::mScarlet*). C Structures at the cm scale can be fabricated while maintaining μm resolution (here a microfluidic chip, Fig. 4). D–F The microfabrication technique can be used for a variety of applications: (D) patterning of surface topology with subcellular precision (Fig. 2), (E) patterning of surface chemistry to e.g. induce a specific cell morphology (Fig. 3), (F) microfluidic devices (Fig. 4), (G) imprinting other substrates like Matrigel or agar, using the PDMS mold as a stamp (Fig. 5).

or reagents, ensuring compatibility with existing equipment, reducing costs, and enabling customization.

We first describe the method, then demonstrate its application across various biological model systems and scales, ranging from subcellular to whole organisms, and from micrometers to centimeters. Specifically, we demonstrate:

1. Fabrication of μ m-scale pillar topologies to guide formation of cytoskeletal structures.

2. Surface patterning with adhesive and non-adhesive coatings to control and standardize fibroblast cytoskeletal organization and 2D gastruloid growth.

3. Manufacturing of microfluidic devices to study confined cell migration through constrictions.

4. Imprinting chambers into agar to confine *C. elegans* movement.

Results

Simplified rapid iteration microfabrication workflow

begin by fabricating structures maskless We via photolithography (Fig. 1A). In this process, an image mask designed in a computer graphics software is projected onto a thin film of UV-curable resin. Analogous to a standard video projector, the light pattern is shaped the light is shaped using a DMD, but instead of projecting the image onto a screen, the image is demagnified using a microscope objective and focused on the microscope slide. We can repurpose our microscope set up for targeted optogenetic photostimulation without any hardware modifications. In the areas exposed to UV light the resin hardens, while the unexposed regions remain soft and are washed away, leaving behind a mold that can be replicated with an elastomer such as polydimethylsiloxane (PDMS). The resulting PDMS copy serves as a foundation for downstream applications, such as microfluidics and stamping.

An overview of our method is provided here, with a detailed protocol and additional practical considerations outlined in the methods section of this paper. The required consumables and devices are provided in Tables 1 and 2.

Spin coating. The procedure begins by coating a standard microscope slide with 3-(trimethoxysilyl)propyl methacrylate (TMSPMA) to enhance the adhesion of the UV resin.³⁸ A thin layer of consumer-grade 3D-printing resin is then spincoated onto the slide. The z-layer height is controlled by the spin coating RPM, and can be calculated with a calibration test print and a simple formula described later.

UV projection. The prepared slide is placed onto a standard fluorescence microscope (Eclipse Ti, Nikon) equipped with a system capable of projecting UV light patterns. In our setup, we use a commercial DMD (Mosaic 3, Andor or Polygon 1000, Mightex) available as microscope attachment in combination with a 395 nm UV light source (Spectra X, Lumencor) to project patterns onto the resin. By adjusting the microscope objective, we can control the projected feature size: a 20× air objective offers a good tradeoff between field of view (FOV) size and resolution for most applications. For large features and thick

layers, we use a 4× objective because of its increased depth of field. Fig. 1B illustrates applications of microfabricated circles ranging from 1 to 1400 μ m. Using the microscope's x/y stage, we iteratively project images to create patterns larger than the FOV while maintaining spatial resolution (Fig. 1C). The footprint of the structure is thus limited by the carrier glass used (up to 50×70 mm minus border padding) or stage travel range. The UV exposure time is dependent on the layer height, objective and light source used, but can be empirically calibrated within 5 min (Fig. S1[†]). Due to the high UV sensitivity of the resin, illumination times never exceeded 1000 ms per field of view, even with a relatively low energy output measured at 2.96 mW. A microscope slide can thus easily be scanned and exposed within a few minutes. The resin is designed for the consumer market, optimized for use in home settings without requiring specialized environments like yellow light rooms typically needed in conventional photolithography.

Post-processing. After exposure, the unexposed resin is washed away, forming the initial resin mold. This mold is then post-cured using UV light and heat.³⁹ Once cured, PDMS is poured over the mold and hardened in an oven. The PDMS cast is then lifted off and can be used for various experiments, including controlling surface topology, micropatterning surfaces, fabricating microfluidic devices, and creating agar microchambers (Fig. 1D and F). We show results for these four applications in the next chapters.

Microscope programming. To automate the printing process, we control the microscope using custom code available on GitHub.[‡] We employ μ Manager⁴⁰ in combination with Pycromanager⁴¹ to control the hardware from Python, and Napari⁴² to visualize the camera feed and provide interactivity. An interactive notebook with step-by-step instructions simplifies the initial hardware setup and experiment execution, making it accessible even to researchers without coding experience.

Validation & performance. The z-layer height h is controlled by the spin coating RPM, and can be calculated with a calibration test print and the following formula:

$$\text{RPM}_{\text{desired}} = \text{RPM}_{\text{known}} \times \left(\frac{h_{\text{known}}}{h_{\text{desired}}}\right)^2$$

To calibrate for a large range of RPM (200–3200 RPM), we find that a slightly more complex model with a constant offset leads to a good fit of the data (R^2 : 0.96 with offset, R^2 : 0.90 without offset):

$$h(\text{RPM}) = \frac{a}{\sqrt{\text{RPM}}} + h_0$$

where *h* describes the film thickness as a function of the spin speed RPM, *a* is a proportionality constant and h_0 represents a constant offset (Fig. S2†). The theoretical maximal resolution of the projection for a certain optical setup can be calculated by

[‡] https://github.com/hinderling/fabscope.

Table 1 Consumables for microfabrication

Material	Product	Quantity	Cost (USD)
TMSPMA 3-(trimethoxysilyl)propyl methacrylate	Sigma Aldrich, 440159	100 ml	\$71
PDMS polydimethylsiloxane	SYLGARD 184 Silicone Elastomer Kit	0.5 kg	\$162
UV-printing resin	Copymaster3D Tough UV Resin Clear	500 ml	\$31
Methanol	_	_	_
Isopropanol	_	_	_
Ethanol	_	_	_
Microscope slides	_	_	_

Table 2 Equipment used in microfabrication

Device	Model
Microscope with DMD and	Nikon TiE, Andor Mosaic 3/Mightex
μManager	Polygon 100
UV light source	Lumencor Spectra X
Spin coater	Ossila 120–6000 RPM
Plasma cleaner	Femto Science, Cute
Oven 70 °C	—

measuring the FOV of the DMD divided by its resolution. For a 20× objective, we measure a projection width of 560 µm, divided by 800 px DMD x-resolution we get a resolution of ~ 0.7 μ m px⁻¹. In practice, diffraction artifacts from the DMD mirror edges and the optical path are reducing the resolution. Using a 20× objective, we can reliably print features sizes down to 5 µm (Fig. S3[†]), smaller features are possible depending on the design. For the 1 µm sized pits in the first example, we used a 100× oil immersion objective (see Table 3 for a list of optical properties of the objectives used in this study). In our tests, we found that exposure settings around 0.5 mJ mm⁻² lead to good results across a range of layer heights (Fig. S4A⁺). The achievable resolution also depends on the layer height, with thinner layers enabling higher resolution (Fig. S4B[†]). For stitching multiple FOVs, we performed a quality control experiment, printing regularly spaced lines across a 5 mm field by stitching 10 × 10 FOVs (20× objective). While minor horizontal misalignments were visually detectable, quantitative analysis revealed a pitch standard deviation of less than 2 µm in both the x and y directions (Fig. $S5^{\dagger}$) across the print.

Engineering surface topology with 1 μm size features to guide cell protrusions

Cells can sense and respond to the geometry and topology of their 3D extracellular matrix surroundings. Early approaches to creating artificial topologies used natural materials such as spider webs⁴³ and fibrin fibers extracted from plasma.⁴⁴ More recently, microfabrication techniques have been employed to achieve greater control over surface structures.^{45–47}

Here, we create an artificial surface topology featuring round pits or pillars with diameters as small as 1 μ m (Fig. 2A). This is achieved by printing pillars using backside illumination with a 100× oil objective through a coverslip. The printed structure is cast into a negative PDMS stamp, forming a surface with pits. This negative is then cast again into a positive PDMS stamp to produce pillar structures. For experimental use, either the pit or pillar structure is transferred by applying a drop of liquid PDMS to a coverslip and imprinting the topology using a passivated PDMS stamp on the uncured PDMS.

The coverslip is subsequently placed in an oven to harden the PDMS, after which the stamp is carefully removed, leaving behind a surface with well-defined pits or pillars. REF52 fibroblast cells expressing a fluorescent biomarker for F-actin,⁴⁸ are then seeded onto the patterned surface.

Live cell imaging reveals that the cells form actin-rich protrusions extending into the pits, indicating an active response of the cell to the engineered surface topology (Fig. 2B, Movie M1†). Timelapse imaging results of cells on pillar structures are shown in Fig. 2C and Movie M2.† Striking F-actin patches form at the leading edge of the cell, completely engulfing the pillars.

Patterning of surface chemistry to control cytoskeletal shape or colony growth

Micropatterning enables the modeling of cell and tissue microenvironments by chemically patterning surfaces. This technique allows researchers to control cell and tissue morphology by enforcing specific shapes, facilitating causal investigations into the relationship between morphology/ geometry and function, or reducing heterogeneity by standardizing cell shape.^{22,29,49}

Table 3Optical parameters and calculated depth of focus for microscope objectives used for printing in this study at 395 nm. Pixel size calculated withAndor Mosaic 3 ($600 \times 800 \text{ px}$)

Objective	NA	DMD pixel size (µm)	Depth of focus (μm)
Nikon CFI Plan Fluor 4× air	0.13	3.51	23.37
Nikon CFI Plan Apo Lambda 20× air	0.75	0.70	0.70
Nikon CFI Plan Apo Lambda 40× air	0.95	0.35	0.44
Nikon CFI E Plan Achromat 100× oil	1.25	0.14	0.38

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Fig. 2 Microtopology: A cells are seeded on a thin PDMS layer with pits or pillars with diameters of 1 μ m and μ m. Fluorescence imaging and intensity profiles show the regular spacing of actin clusters forming at pit locations (confocal z-slice, more data Fig. S8 and Movie M1†). B 3D imaging shows that cells form actin rich protrusions into 5 μ m pits (confocal z-stack). C Timelapse microscopy reveals rich actin dynamics of two cells migrating on 5 μ m pillars. Small pictures show selected frames of the movie with arrows indicating the migration direction, brightfield image shows PDMS surface with 5 μ m pillars. Last image shows temporal-color code of actin movie (Movie M2†). All images in this figure are single confocal z-slices.

The most widely used micropatterning method today is microcontact printing.⁵⁰ In this approach, extracellular matrix proteins such as fibronectin or laminin are stamped onto a glass slide using a PDMS stamp. The unstamped regions are then coated with PLL-*g*-PEG, a non-adhesive polymer that prevents cell attachment⁵⁰ (Fig. 3A). Commercially available slides with experiment-ready, standardized patterns simplify this procedure and enhance reproducibility; however, custom patterns are often required depending on the experiment or the adhesive properties of specific cell lines. Our method allows for the rapid prototyping of PDMS stamps with various designs. As a proof of concept, we designed variations of the commonly used anchor shape pattern (Fig. 3B) to identify the optimal size and design that allows optimal spreading of NIH3T3 fibroblasts. Instead of stamping the fibronectin with a PDMS stamp, we first coated the whole slide with fibronectin (or fluorescent laminin for quality control) then protected areas by placing the PDMS stamp on top. We then etched away the unprotected areas, with a plasma cleaner. The etched regions are subsequently



Fig. 3 Micropatterning: A glass is patterned with patches of matrix proteins like fibronectin which cells attach to, and PLL-*g*-PEG which is nonadhesive. The cells adopt the shape of the pattern. B Microfabricated PDMS stamp containing different designs and sizes to screen. C Fluorescent laminin showing precise surface patterning (full chip shown in Fig. S6†). D Cells attaching to the micropatterns. E Sample cells on three different shapes (anchor, umbrella, circle). The cells on the anchor pattern are more contracted compared to the umbrella shape. F Quantification of cells on different shapes shows that cells on umbrella patterns are significantly larger than on anchor patterns (t = 4.33, p < 0.001, df = 159). G The best patterns from the screen are selected to create large arrays of the same pattern. The morphology of hundreds of cells can be homogenized per experiment. Crops show cells with segmentation outline, automatically detected and filtered by morphological features. Brightfield images in B and G are high-pass filtered to reduce vignetting and out of focus artifacts.

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filled with PLL-g-PEG, ensuring that cells adhere exclusively to the fibronectin-coated regions while they can be washed away from the non-adhesive PLL-g-PEG areas. This left us with patterns with sharp edges, clearly reproducing fine structures of the shape down to a few μ m (Fig. 3C) across an area of a few mm (Fig. S6,† to which the cells attaches to well (Fig. 3D). For single cell patterns, coating the entire slide first and then using the stamp to protect specific areas, rather than coating the stamp and stamping onto the glass, has been more convenient and reproducible for us.

Our goal was to identify patterns that accommodate a single cell while promoting optimal spreading without inducing excessive retractions or cell detachment. For example, many cells on the anchor shape seemed to collapse on the sides, while on the umbrella shape they were more spread out (Fig. 3E). We automatically segmented the cells with Cellpose3 (ref. 51) and measured their area (anchor: $891.99^2 \pm 287.2$ (n = 79), umbrella: $1094.29 \ \mu\text{m}^2 \pm 304.68$ (n = 82), circle: $1007.31 \ \mu\text{m}^2 \pm 439.72$ (n = 80)), and found that it is significantly higher in the umbrella shape *versus* the anchor shape (independent *t*-test: t = 4.33, p < 0.001, df = 159) (Fig. 3F).

After screening different pattern variations, we selected the most suitable design and fabricated an array featuring the optimal shape and size for our experiments. This patterned grid enables the production of hundreds of cells with a standardized cytoskeletal organization, ensuring morphological uniformity (Fig. 3G).

Additionally, we demonstrate the value of our approach at the tissue scale to generate circular patterns that are routinely used in the stem cell field to generate 2D gastruloids of defined diameters, similar to a previous publication⁴⁹ (Fig. 1B and S7†). For stem cell colony formation with large diameters (250–1500 μ m), a basement membrane extract is stamped onto a glass slide using a PDMS stamp, followed by cell seeding. No non-adhesive coating is needed.

Multilayer microfluidic devices to study confined cell migration

Beyond classic 2D migration models, cells migrate in a 3D environment *in vivo*.⁵² Microfluidic devices with precisely

shaped constrictions have provided a way to study 3D migration under well-defined conditions. These devices have been used to investigate nuclear deformation⁵³ and the cytoskeletal mechanisms that generate the forces required for cells to squeeze and migrate through narrow gaps.^{10,11}

We demonstrate that our fabrication method can produce such microfluidic chips with customizable geometries (Fig. 4A-D). To enhance flow through the channels, we print larger layer heights for the regions leading to the constrictions. The device master mold is fabricated by iteratively spin-coating and exposing layers of 3D printing resin, aligning the layers precisely using a fiducial marker and the microscope camera. The fiducial marker is printed as part of the microfabricated structure (X-shape visible in top left corner Fig. 1C). After calibrating with a single layer and printing two stacked layers, we observed a 4.92% error from the expected layer thickness, indicating that spin coating resin directly on glass or on a previously printed layer does not significantly affect spin coating properties (Fig. S9[†]). A UV-free light source is used during alignment to prevent accidental polymerization of the resin.

REF52 cells expressing a biosensor for F-actin are seeded onto the PDMS device passivated with PLL-g-PEG and are allowed to settle for two hours. Spinning-disk confocal imaging enables clear visualization of actin patterns along the cortex, as cells migrate through the constrictions (Fig. 4, Movie M3[†]).

Imprinting agar chambers for long-term tracking of C. elegans

Regulating organ size during development is crucial, as even minor imbalances in growth rates can lead to significant deviations in organ proportions. Studies using *C. elegans* have demonstrated that organ size scaling remains remarkably consistent across individuals.^{54,55} To track growth over several days by imaging, individual worms were placed in agarose microchambers⁸ (Fig. 5A). While microfluidicsbased systems allow temporary immobilization of worms for imaging weak fluorescent signals,⁵⁶ the microchamber approach discussed here enables imaging of a larger number



Fig. 4 Microfluidics: A 2.5D structure is achieved by iteratively spin-coating and exposing 3D printing resin. Supply channels are printed with a larger layer height to increase fluid flow rate. The microfluidic device is built by punching inlet and outlet holes, then plasma-bonding the PDMS stamp onto a coverslide. B Complete microfluidic chip with adapters for syringes or automated pumping systems. Chip is filled with food coloring to visualize the channels. Grid lines for scale (small squares are 1 mm, large squares 1 cm). C Section of the chip photographed with a smartphone through a microscope ocular, manually adjusted white balance. D Brightfield image 20× objective. As the microscope camera used only provides grayscale images, multiple exposures with different filters are merged and color balance is adjusted to achieve an RGB image. E A different chip design, showing a timelapse movie of a cell migrating trough a constriction (Movie M3†).

of animals in parallel, is simpler to manufacture, and does not require vacuum or pressure pumps.

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Microchamber designs can be ordered from lithography companies. Multiple designs can be batched into a single order to reduce costs. Depending on the size of the design and the company, around 20 designs can be ordered in one go, resulting in an approximate cost of 60ε per design and a delivery time of a few weeks. Our method allows for customization and testing of different chamber patterns within a day. For instance, chamber size can be adjusted to match the microscope's field of view, optimizing space and food quantity for each worm while preventing it from moving out of frame. Our method reduces material costs to <1 ε per chip.

Here, we present circular microchambers arranged in a honeycomb pattern (Fig. 5B), enabling long-term tracking of many individual worms in parallel. For preparing worm imaging chambers, a PDMS master is first fixed to a glass slide using double-sided tape and is plasma treated to clean it and enhance hydrophilicity. The master is then used to cast chambers by sliding it into melted agarose gel on a glass slide. After a brief curing period, the agarose is trimmed to retain only the wells and surrounding space.

Bacteria are added to the wells as a food source. Worm eggs are collected from culture plates and individually placed into the wells using an eyelash pick, ensuring only eggs at the 2-fold stage are selected. The agarose wells are then inverted onto an imaging dish with a gas-permeable polymer bottom (Fig. 5C).

To seal the system, the dish is covered with low-melting agarose followed by a PDMS overlay, and parafilm to prevent evaporation. The PDMS cures at room temperature during image acquisition. A custom plate holder allows simultaneous imaging of six dishes on a single microscope. The worms are tagged with fluorescent proteins, marking the pharynx and somatic tissues (Fig. 5D). Timelapse imaging and automated image analysis allows to track the growth of individual worms over multiple days (Fig. 5E), capturing the complete lifecycle, from hatching to laying eggs (Fig. 5F, Movie M4[†]).

Discussion

Photolithography and soft lithography are widely used in biological research for microfabrication due to their high spatial resolution and versatility. However, these methods require specialized equipment, trained personnel, and the use of toxic chemicals, which can be a barrier for many laboratories. In contrast, 3D printing offers a low-cost and accessible alternative but lacks the spatial resolution needed for many microfabrication applications.

In this work, we repurpose a fluorescence microscope designed for targeted photostimulation for microfabrication. By combining 3D printing with established lithography techniques, we achieve micrometer-scale precision over centimeter-scale areas while maintaining rapid prototyping capabilities. This approach makes microfabrication more accessible and applicable to a wide range of research questions.

We now approach microfabrication much like conventional 3D printing: as a low-friction tool for rapidly designing custom solutions. When an idea arises, we often ask: could microfabrication be a solution here? The ease of use allows us to quickly fabricate a prototype, yielding results within a day. The low cost minimizes the risk of trying. This accessibility has led us to integrate microfabrication into numerous projects within the lab and institute, where we previously wouldn't have considered it, simply because it is now so straightforward to implement.

Previous efforts to simplify microfabrication workflows have focused on eliminating the need for clean room facilities,³² using glass³⁰ or polyethylene terephthalate



Fig. 5 Microchambers A agar microchambers for long-term tracking of *C. elegans* tagged with fluorescent markers expressed in the pharynx and somatic tissues. B Digital mask containing an array of microwells and photo of the corresponding structure made from 3D printing resin. From this structure a PDMS stamp is molded. C The PDMS stamp is used to imprint microwells into agar. Photo shows an array of microwells loaded with worms loaded in an ibidi dish for long term imaging. D Microscope image of a microchamber containing a worm. The brightfield channel shows the microchamber and bacteria as food source, fluorescent channels the pharynx (green, *myo-2p::GFP*) and somatic tissues (red, *eft-3p::mScarlet*). E The microchambers restrict the movement of the worms, allowing to track the individual growth over multiple days. F Timeseries showing the complete lifecycle of one worm (Movie M4†).

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(PET)⁵⁷ instead of silicon wafers as a substrate, and finding alternatives to the expensive photoresist SU-8.^{18,58} Maskless lithography systems utilizing scanning stages^{59,60} or projectors^{18,30,33–35,61} have been introduced to increase iteration speed. While many maskless systems rely on custom hardware setups, some labs have leveraged commercial DMD systems and microscopes.^{18,35} Another group has developed a method to facilitate the separation of PDMS structures without using chlorosilane coating,⁶² a widely used but highly toxic chemical that releases hydrochloric acid upon contact with water. Our approach integrates these scattered simplifications into a single workflow and builds upon them to further streamline the process.

Our microfabrication workflow only requires 3 nonstandard consumables: TMSPMA, PDMS and 3D printing resin (all listed in Table 1). The use of 3D printing resin as a substitute for SU-8 eliminates time-sensitive baking steps and reduces the need for extensive glass slide cleaning. Instead of using highly corrosive piranha solution, a short submersion in TMSPMA solution is sufficient to ensure strong bonding between the printed structures and the glass substrate. Leveraging microscope control software with Python scripting allows for seamless customization and automation of the printing process, which is particularly beneficial for iterative design processes.

Using μ Manager, our method is compatible with a wide range of existing microscope hardware, ensuring reproducibility across different lab settings. While this paper focuses on DMD-based patterning, we have also explored the use of a total internal reflection fluorescence (TIRF) and fluorescence recovery after photobleaching (FRAP) system (iLas 2, GATACA) with galvo mirrors. Although this system offered considerably slower scanning speeds per field of view, it was still viable for producing useful microfabricated structures.

For labs looking for a dedicated microfabrication microscope, there are complete hardware solutions available commercially (Primo Optical Platform, Alvéole) that should be compatible with the methods presented in this paper.

A key tradeoff in this workflow is the choice of microscope objective, which determines both lateral (*x*/*y*) resolution and axial (*z*) focus range. As outlined in the Methods section, the depth of focus can be approximated by $\Delta z = \frac{n \cdot \lambda}{NA^2}$, where n is the refractive index of the immersion medium, λ is the exposure wavelength, and NA is the numerical aperture of the objective. Table 3 lists the calculated depth of focus for each objective used in this study, illustrating that objectives with higher magnification and NA (*e.g.*, 100× oil) achieve finer lateral resolution but at the cost of a shallower *z*-range. Confocal fluorescence exclusion imaging (methods) provided the best available method for assessing axial resolution, but was still limited by optical artifacts, particularly at material interfaces (Fig. S9†), preventing accurate quantification of sidewall profiles.

However, we find that practical layer thicknesses exceed the theoretical focus range while still maintaining sufficient precision for most applications (*e.g.*, estimated focus range for $20 \times$ air is 0.7 µm, but good results can be achieved even with 400 RPM (Fig. S4†), which corresponds to 45 µm layer thickness (Fig. S2†)).

In practice, we find that layer thicknesses can exceed the theoretical depth of focus while still producing features with good structural accuracy. For example, although the estimated axial focus range for the $20\times$ air objective is only 0.7 µm, reliable results are achieved even at 400 RPM (Fig. S4†), which corresponds to a layer thickness of 45 µm (Fig. S2†). This demonstrates that even when the axial focus range is much narrower than the printed layer thickness (0.7 µm vs. 45 µm), reliable microfabrication is still possible.

The impact of layer thickness on lateral feature fidelity is shown in Fig. S4B,[†] where increased thickness leads to reduced precision. To circumvent this limitation, complex structures can be fabricated by stacking multiple layers (2.5D printing, see Fig. S9[†]). While this approach increases achievable aspect ratios, it also requires more labor-intensive alignment and sequential exposure steps.

Also certain cost-related limitations remain: plasma cleaners and DMD systems (see Table 2 for required hardware devices) and can be expensive if not already available in the lab, although DIY alternatives have been demonstrated at a fraction of the cost^{63,64} of commercial systems. Among the required materials, PDMS is the most expensive (\$162 per 0.5 kg), but the cost per fabricated device is low and the chips can be reused multiple times.

Looking ahead, future efforts will focus on developing lower-cost hardware solutions to further democratize access to microfabrication techniques. Digital light processing (DLP) printers contain much of the necessary hardware (UV lamp, driver board, DMD) at a fraction of the cost (Anycubic Photon Ultra is \$300). The maker community has already made significant progress in adopting such systems for microfabrication.⁶⁵ Integrating such components into lowcost modular microscope systems, such as UC2,⁶⁶ could provide an affordable alternative to high-end microscope projection photolithography setups.

Our Python-based workflow, along with interactive fabrication features that allow regions of interest to be selectively exposed in real time using live camera feedback, presents opportunities for intelligent automation. Potential applications include automatic alignment and exposure compensation. Closed-loop positioning with computer vision could compensate for low-precision x/y stages by digitally adjusting the stimulation mask to correct mechanical stage positioning errors. By mixing dark particles into the 3D printing resin, we can generate unique speckle patterns that serve as fiducial markers for image registration, which in the future might enable high-precision microfabrication despite Automatic image-guided stage limitations. exposure compensation, recently implemented to improve fine-feature resolution in hydrogel printing,67 could be adapted to our system and appears compatible with the existing hardware setup. Real-time feedback capabilities could also enable the

fabrication of UV-crosslinkable, cell-compatible hydrogels, allowing researchers to print structures *in situ* and capture cells of interest with high precision.

We hope this paper stands out by demonstrating the versatility of the technique across diverse biological applications. We believe the simplicity and versatility of our method will encourage broader adoption across the scientific community. Our detailed Methods section, which includes practical troubleshooting tips, should further facilitate reproducibility. The open-source nature of the project is expected to inspire further developments and novel applications across diverse fields.

Methods

Microfabrication protocol

All of the microfabrication applications start by creating a 3D printed mold from UV-curable resin, from which a PDMS cast is made. Standard microscope slides are used as a carrier for the microfabricated structure for their sturdiness and availability.

1. Prepare the glass slide for good adhesion. Clean the glass slide for 2 min in the plasma cleaner (other methods proposed in the literature are ultrasonic cleaning, cooking in deionized water, EtOH bath but led to worse results for us).

Coat the slide with 3-(trimethoxysilyl)propyl methacrylate (TMSPMA) by submerging it in a 2% solution of TMSPMA (solvent: 96% EtOH, 4% Keton) for 5 minutes. After incubation the slide is washed in a EtOH bath for 5 min, then dried in the oven for 10 minutes at 70 °C. To conserve TMSPMA solution, a small amount can be applied with a pipette instead of full submersion; due to the glass hydrophilicity from plasma activation, the solution spreads evenly across the surface.

2. Coat the glass slide with the UV-curable resin. The thickness of this layer determines the final height of the structure. The thickness is adjusted by varying the RPM of the spin coater and is dependent on the viscosity of the resin. The viscosity differs strongly between brands, and also increases as solvents from the resin evaporate. To minimize this, we recommend to prepare aliquots of resin in 50 ml falcon tubes instead of opening the storage bottle many times. We first deposit ~ 1 ml of resin onto the glass slide, a pipet can be used with a cut-off tip to facilitate flow of the viscous fluid. Be careful to not create any bubbles, as they result in uneven layer height. The resin is roughly distributed on the slide by tilting in in different directions before spinning. We recommend a two-step program for the spin coater, a slow first step with 1/2 of the final speed or 800 RPM (whichever is lower), for 10 seconds to distribute the resin evenly on the slide, then a second step for 20 seconds with the final speed to achieve the required thickness. Spinning with high RPM directly may result in the resin being slung off the glass slide. The slide is now ready for UV exposure. When carrying the slide between rooms, cover it to reduce contamination with dust particles.

3. Expose the pattern with UV light. The slide is placed into the microscope, resin side towards the objective. For inverted microscopes, make sure there is no excess resin that could drip into the objective. Find the focus plane, load the pattern and set the exposure time (described in detail below).

4. Remove uncured resin. Immediately after pattern exposure, the uncured resin is washed by placing the slide in an Isopropanol bath for 5–15 minutes, depending on layer thickness. Any remaining uncured resin, *e.g.* in narrow gaps, can be washed out efficiently by spraying Isopropanol from a pressure sprayer. When producing many slides, to reduce Isopropanol waste, multiple baths can be used, moving the slides sequentially from the dirtiest to the cleanliest. After washing, the glass and 3D print should be clear and not show any smears or diffuse residues.

5. Post-curing the resin. The resin is exposed under a homemade UV-light (see Calibrating the UV-exposure time below) for 5 min to activate all photoinitiators remaining in the resin and complete hardening of the pattern. To evaporate all volatile species that could cause curing inhibition (see below), keep the slide at 70 °C over night.

6. Creating a PDMS replicate. The 3D printed structure is used as a mold to cast a negative PDMS mold (see below for PDMS preparation). The PDMS is slowly poured from one side to avoid trapping pockets of air. To remove any air bubbles, the slide can be placed in a vacuum pump for 30 min or until no more bubbling is visible. The slide with the PDMS is then placed into the oven at 70 °C for around 2 hours or until the PDMS is hardened and non-sticky to the touch. Now the PDMS mold can be peeled of the glass slide with the 3D printed structure. If there are issues with adhesion between the 3D print and glass, the 3D printed structures can detach during this step and stick to the PDMS instead (check below for a trouble shooting guide).

7. Inverting the PDMS replicate. We recommend to use the negative mold as template to create any PDMS structures actually used in experiments, as they show greater durability than the 3D printed slide for repeated mold use (resistance to accidental scratches, ease of separation). To fabricate a positive PDMS structure from the negative mold, the surface must first be passivated; otherwise, the two PDMS layers will irreversibly bond. This is achieved by briefly plasmaactivating the negative mold (10 s), followed by immersion in a methanol or ethanol bath.⁶² After soaking, the methanol/ ethanol-coated mold is left at room temperature (5 min), then the excess solvent is discarded. The slide is allowed to dry fully, first at room temperature, then in an oven at 70 °C (5 min). PDMS can then be poured into the passivated mold, taking care to avoid air bubbles; if necessary, bubbles can be removed by degassing the mold in a vacuum chamber. The PDMS is cured in the oven at 70 °C for approximately 2 hours. Once fully cured and non-sticky to the touch, the positive replica can be separated from the negative mold. This replication process can be repeated multiple times to produce additional PDMS copies, for example to scale up the number of devices used in parallel experiments.

PDMS preparation. PDMS is mixed in a 1:10 ratio of crosslinker and elastomer in a large batch, then aliquoted into 50 ml falcon tubes and centrifuged to remove trapped air bubbles. Centrifugation works faster and produces less waste than removing air bubbles with the vacuum pump. Aliquots can be stored at -20 °C for months without curing. When using the PDMS, the desired quantity can be poured cold directly from the cold falcon tube.

Finding the focus plane. The slide is placed into the microscope, resin side towards the objective. For inverted microscopes, make sure there is no excess resin that could drip into the objective. To find the focus plane, a checkerboard is projected onto the glass slide with red light. Moving in the *z*-axis towards the slide, the first focus plane is resin surface, the second focus plane corresponds to the surface of the glass slide. We got best results focusing on the surface layer of the resin (1st focus plane) or slightly below.

Calibrating the UV-exposure time. Set the correct UV-light exposure time, which is dependent on the brand of resin, light source intensity, DMD, objective, and even the design of print. Very small or thin patterns may require increasing the exposure times. The multitude of factors makes calculating the exposure time difficult, and its easiest to just perform an exposures test-series. The test series can be performed automatically using code we provide. The quality can be inspected directly after cleaning with isopropanol, so the whole calibration procedure should not take more than 10 min. For our setup, at a layer thickness of 14 µm, we use 600 ms for 4× magnification objectives, 150 ms for 10×, 30 ms for 20×. We use a LED light source (Lumencor Spectra X) at 395 nm with a 395/25× excitation filter (Lumencor) and measure 2.96 mW at the 10× objective surface. The fraction of activated DMD pixels is linear to the power reaching the sample (see Fig. S10,[†] linear fit of the data shows a R^2 value of 1.00, slope 2.71 mW/100% ON pixels, intercept -0.02 mW).

Avoiding curing inhibition. One of the main difficulties when using 3D printing resins with PDMS is that unactivated photoinitiators in the resin can leach into the PDMS and prevent it from polymerizing, while remaining resin monomers can increase adhesion of PDMS to the 3D printed mold, making them difficult to separate.³⁹ They find that 11 out of 16 tested resins can be treated using a combination of 120 °C heat treatment and UV exposure with a total curing time below 135 min, which should be considered if protocol duration is essential. For us curing at 70 °C over night was a practical tradeoff, also preventing issues warping/detachment of the 3D print at high temperatures. We also have one oven running at 70 °C in the lab anyways, using it instead of running an additional oven at 120 °C is also more energy efficient. We built a UV-curing station by cladding a box with reflective aluminium foil and using a 405 nm LED unit (CR-6565-4CLED) marketed as replacement part for Epson flatbed printers. It comes packaged with an aluminum heatsink and cooling fan. Available online for <20 USD with different wavelengths (365 nm, 385 nm, 359 nm, 405 nm). The unit runs at 24 V/40 W, we use a AC85-265 V to DC24-36 V

converter (LED driver YJ-TG50W-1300) as power supply. We also observed curing inhibition caused by insufficient mixing of the PDMS elastomer and crosslinker during preparation. This typically appears as uncured "puddles" of PDMS unrelated to the printed structures, often found on the glass or top surfaces of the sample. Thorough mixing is essential to ensure complete curing and avoid such defects.

Plasma cleaning and activation. We perform all our plasma cleaning or activation steps with atmospheric gas mix at 3 torr pressure, only varying the timing. PDMS can start to break down if activating for too long, leading to a rough surface texture that looses its adhesive properties. This can be observed visually as PDMS appearance of an opaque surface layer.

Trouble shooting resin-glass adhesion. The composition of glass slides used can make a difference, which is difficult to figure out as not all brands are clear about the additives and surface coatings used. We had some experiments fail when using a freshly opened package of a very old stock of slides, which upon close visual inspection showed oily smeared surface. TMSPMA coating was integral to ensuring good adhesion. We tried coating the slides in batch and storing for later experiments, but they develop a visible layer of impurities if stored after some weeks which decreases adhesiveness. For best results use freshly coated slides.

Trouble shooting PDMS–PDMS separation. Duration of plasma activation is important for PDMS–PDMS separation. While we get good results with 10 s activation time, layers bonded much too strongly when activating 30 s or longer. Methanol incubation time did not seem to affect separation quality in either of the procedures.

3D imaging of the PDMS stamps for quality control. Liquid dye fluorescent under UV light is extracted from a yellow highlighter pen by breaking it open and squeezing the ink from the fibers into a solvent (water or ethanol). The negative PDMS stamp is plasma-activated and coated with the extracted fluorescein. It is then pressed against a coverslip, allowing excess fluorescein to be expelled from the sides. Z-stacks are acquired using a confocal microscope. Alternatively, a positive stamp can be pressed onto a coverslide, and the empty volume can be filled with fluorescein from the sides (florescence exclusion). We used the second method to measure z-layer height accuracy. We printed test patterns at different RPM (200, 400, 800, 1600, 3200), and replicated the structure in PDMS with double casting. Brightfield and fluorescence exclusion z-stacks of a sample pattern (400 RPM) are shown in Fig. S2.†

Micropatterning for single cell fibroblasts

1. Preparing the well plate. A 24-well plate is plasma activated for 30 seconds, then coated with 250 μ l of 10 μ g ml⁻¹ fibronectin (human plasma fibronectin purified protein, Merck) diluted in MilliQ water. Plasma activation increases the hydrophilicity of the glass, decreasing the volume of fibronectin solution required to coat the surface. The plate is

incubated for 1 hour at 37 °C or overnight at 4 °C. It is then washed thoroughly with PBS and stored at 4 °C. Before use, PBS is aspirated, and the wells are allowed to dry in the hood. The PDMS stamp is placed onto the coated wells. The well plate is plasma activated for 1 minute and 30 seconds. After activation, the PDMS stamps are removed, and the wells are washed with MilliQ water. A solution of 250 µl PLL-PEG (100 µg ml⁻¹) (PLL(20)-g[3.5]-PEG(5), SuSoS) or F-Pluronic (5%) (Pluronic F127, Bioreagent) is added and incubated at room temperature for 1 hour. F-Pluronic is cheaper but has decreased anti-adhesive properties for some cell lines. The solution is aspirated, followed by washing with PBS. The wells are covered with PBS and are ready to use. Alternatively, they can be stored at 4 °C for months.

2. Seeding cells. PBS is aspirated from the wells, and 300 μ l of growth medium is added. ~20000 cells are then seeded per well into growth medium, and are incubated at 37 °C and left to attach to the micropatterns for 3 h (varies depending on cell line, pattern, and cell density). Seeding density is checked, and the cells are gently washed with PBS until the desired density is achieved.

Reusing the PDMS stamp. The PDMS stamps can be cleaned with ethanol and re-used after drying. If soaked in ethanol for long, the stamp can acquire a opaque texture, which disappears after drying. Drying can be sped up by placing the stamp into a 70 °C oven.

Quality control micropatterns. The well plate is coated with 1 μ g ml⁻¹ fluorescent laminin (LMN01-A, Cytoskeleton) for 1 hour at 37 °C or overnight at 4 °C. It is then washed with MilliQ water and covered in MilliQ for storage at 4 °C. Before stamping, MilliQ is aspirated, and the well plate is dried in the hood. The pattern is stamped using the plasma etching technique described above, and the fluorescent pattern is imaged.

Micropatterning for stem cell 2D gastruloid

1. Preparing the PDMS stamps. Stamps are cut to appropriate size, so they can easily fit the well of a 24-well plate. Sterilize stamps in ethanol and then dry them under the hood.

2. Preparing the well plate. Coat a 24 well plate with (3-mercaptopropyl)trimethoxysilane by putting the plate open in the desiccator with 100 μ l of (3-mercaptopropyl) trimethoxysilane in a falcon tube lid. Pump is activated for 5 min and then turned off, stamps are left for another 30 min in the desiccator, then put in a 80 °C oven.

3. Stamping matrix. 1:100 Geltrex solution (\sim 50–100 µl per stamp) is added to dried stamps and put in the incubator for at least 30 min. Extra Geltrex is removed and stamps are left to dry (can be observed under the microscope). Once stamps are mostly dry, they are picked with up with tweezers sterilized in ethanol. Stamp is flipped and placed it in the center of a well with one motion. Stamp is gently pressed. This is repeated for all stamps. Stamps are then left for 20 min before being removed with tweezers with one motion. Wells are rinsed well with PBS.

4. Coating non-adhesive polymer. 400 μ l of 5% Pluronic-F 127 is added to each well for 1 h. Each well is then gently washed 3× for 5–10 min in PBS. Plate is now ready to use.

5. Seeding cells. 1 ml of StemPro Accutase is added to the cells to dissociate. Cells are put back in the incubator and regularly checked. When most cells are dissociated (after 3–5 min), they are thoroughly pipetted and put in a 15 ml falcon tube. 7 ml of E8 flex (A2858501, ThermoFisher) is added, then the tube is centrifuge at 1000 RPM for 1 min. Cells are resuspended in 1 ml E8 flex with 10 μ M ROCK inhibitor (A3008, Apex Bio Lubio). Cells are counted, ~400 000 cells are seeded per well in 400 μ I E8 flex with 10 μ M ROCK inhibitor. When cells adhere, change to E8 flex medium without ROCK inhibitor (after 2 h or next day).

Microfluidic chip

1. Preparing the PDMS stamp. The PDMS stamp containing the microfluidic channels is placed channel-side up on a cutting mat or a piece of cardboard. Access holes are created by punching through the chip using a biopsy punch, allowing connection to the channels from the outside.

2. Bonding the PDMS stamp to the glass slide. The PDMS stamp and a large coverslip (Matsunami micro cover glass 50×70 mm, 0.13–0.17 µm) are placed channel-side up in a plasma oven and activated for 30 seconds. After activation, the PDMS stamp is gently placed onto the glass slide. By applying light pressure, the surfaces are brought into contact. Flipping the chip and reflecting uniform light from below helps identify bonded and non-bonded areas–bonded areas appear as dark patches, while non-bonded areas reflect more light. Localized pressure can be applied using a finger or a pen to ensure full bonding; however, excessive force may collapse the channels, causing the ceiling to irreversibly bond to the glass bottom. Once all areas are bonded, the chip is placed in an oven at 70 °C for 15 minutes to complete the covalent bonding process.

3. Adding fittings to inlets and outlets. Fittings are lightly dipped into a small puddle of PDMS. Excess PDMS should be removed by gently pressing the stamp onto a piece of paper to prevent clogging of the punched channels. The fitting is then positioned over the pre-punched inlet or outlet hole. The liquid PDMS should form a continuous seal around the entire circumference of the interface between the adapter and the PDMS stamp. The chip is then placed back into the oven to ensure a strong bond between the chip and the fittings. Additional PDMS can be applied externally to reinforce the bond if needed.

4. Preparing the chip for the experiment. The microfluidic chip is plasma-activated for 1 minute to render the channels hydrophilic, facilitating the flushing process with medium. For the experiments presented in this paper, the channels are passivated with a non-adhesive coating by flushing the chip with a 0.5 mg ml⁻¹ solution of PLL-*g*-PEG in PBS, followed by incubation at 37 °C for 30 minutes. The PBS is then replaced with the experimental medium by adding it to the inlet port.

Negative pressure is applied at the outlet using a syringe, drawing out the PBS until it is completely replaced by the medium. The chip is now ready for cell seeding.

5. Seeding the cells. REF52 cells are washed twice with PBS and then detached using trypsin. The detached cells are transferred to a Falcon tube, and 5 ml of medium is added before centrifugation at 1000 RPM for 4 minutes. After centrifugation, the supernatant is removed, and 200 μ L of fresh medium is added to achieve a high cell density. A drop of this dense cell suspension is placed in the chip inlet. A syringe is attached to the outlet fitting, and under microscopic observation, negative pressure is gently applied by pulling the syringe to draw cells into the channels. Once the desired cell density within the channels is achieved, the syringe is removed, and the chip is ready for imaging.

To validate the layer height accuracy when stacking multiple layers, we use the 3D imaging procedure for quality control described above. We assume the measured film thickness follows the well-established relationship that h is inversely proportional to the square root of the spinning speed RPM:

$$h_{\rm f} \propto \frac{1}{\sqrt{\rm RPM}}$$

We print a circular pillar, one slide with one resin layer at 120 RPM and one slide with two iteratively printed layers at 210 RPM, aligning the pillar to stack its height. We plot a z-projection across the well border and measure the well depth using Fiji (Fig. S9†). We measure a depth of 43.93 μ m for the single layer, and a total depth of 69.77 μ m for the dual layer (34.885 μ m per layer). For each measurement h_i , RPM_{*i*}, we compute the constant:

$$k'_i = h_i \times \sqrt{\text{RPM}_i}$$

If the relationship holds, $k'_1 \approx k'_2$. We calculate a final error = 4.93% using:

error% =
$$\frac{|k'_1 - k'_2|}{\frac{k'_1 + k'_2}{2}} \times 100$$

The inverse square root relationship between spin speed and film thickness has proven useful in calibrating our setup, enabling us to achieve the desired layer thicknesses in a single attempt, provided the RPMs are not too far off, as discussed in the results section.

Agar microchambers for C. elegans

Method to create agar microchambers was previously described. $^{\rm 55}$

1. Preparing the PDMS stamp. The PDMS stamp is fixed to a standard glass slide using double sided tape, then plasma activated for 30–60 seconds to clean it and make it hydrophilic.

2. Creating the microchambers. Chambers are cast by sliding the PDMS stamp into a pool of molten 4.5% agarose

gel dissolved in S-basal on a glass slide. After curing for 2 minutes, the sides of the agarose are cut to be left only with the wells and some space on the side. As a food source, the bacterial strain OP50-1 was grown on NGM plates by standard methods, scraped off using a piece of 3% NGM agar without cholesterol and then filled into the wells of the agarose gel.

3. Placing the worms. Eggs are picked from plates previously grown and delivered to the spaces left on the side of the well array. Using an eyelash pick, wells are individually filled with eggs at 2-fold stage (younger eggs will not hatch).

4. Imaging. Wells are inverted onto a dish of 3.5 cm diameter with a high optical quality gas-permeable polymer bottom (ibidi). The remaining surface of the dish gets covered with 3% low melting temperature agarose dissolved in S-basal, cooled down to below 42 °C prior to application. The agarose gets topped with ~0.5 ml PDMS and the dish is sealed with parafilm to minimize water evaporation. PDMS is allowed to cure at room temperature on the microscope during the acquisition. Using a custom-made plate holder, six dishes can be imaged simultaneously on one microscope.

Analysis of microchamber experiments was done using a custom-made modular pipelining tool and an associated python package. Both are open source (BSD-3) and hosted on GitHub (https://github.com/spsalmon/towbintools_pipeline, https://github.com/spsalmon/towbintools).

Data analysis

UV energy dose estimation

To estimate the energy dose delivered during UV exposure, we assumed a linear relationship between the LED power setting and the optical output. A maximum output of 2.96 mW was previously measured at 100% LED power. For an exposure setting that performs well across different layer heights (identified in Fig. S5A†), the LED was operated at 46%/64% intensity for 83 ms, illuminating an area of 421.2 μ m × 561.6 μ m (DMD projection size with 20× objective). The power output at 46%/64% was estimated to be 1.36 mW/1.89 mW, yielding a total energy of 0.113 mJ/0.157 mJ. Dividing this by the illuminated area (0.237 mm²) results in an estimated energy dose of 0.478 mJ mm⁻²/0.665 mJ mm⁻².

Automated grid detection and quantitative alignment analysis

This section describes the image processing and quantitative analysis performed in Fig. S5.† To extract regularly spaced grid patterns from the image, we first upscaled the input, then binarized and skeletonized it. Line segments are detected using the probabilistic_hough_line function from the skimage.transform module. Detected segments are classified as horizontal or vertical based on their angles. To represent each physical bar with a single line, segments are grouped by the coordinate of their midpoint along the dominant axis (x for vertical lines, y for horizontal). Within each group, all endpoints are collected, and the pair with the greatest Euclidean distance is selected to define a merged line that spans the full extent of the bar. We then computed the orientation angles of all merged segments and calculate the mean and standard deviation for each direction to assess alignment accuracy. Grid pitch is estimated by measuring the spacing between the midpoints of adjacent lines, converted to physical units (μ m) using the known pixel size. The resulting pitch values are used to compute the mean and standard deviation along both the *X* and *Y* axes.

Estimation of depth of focus

The depth of focus for each microscope objective listed in Table 3 was estimated to provide an intuitive understanding of how the axial focusing range varies with objective choice. We used the standard approximation for the depth of focus in optical systems:

$$\Delta z = \frac{n \cdot \lambda}{\mathrm{NA}^2}$$

where Δz is the depth of focus, *n* is the refractive index of the immersion medium (1.0 for air, 1.515 for oil), λ is the exposure wavelength (395 nm), and NA is the numerical aperture of the objective.

Data availability

The software is open source (BSD-3) and hosted on GitHub: https://github.com/hinderling/fabscope.

Author contributions

Conceptualization, visualization: LH; methodology, writing – review & editing: all authors; investigation, data curation: LH, JF, RH, MK, BG, SP; formal analysis: LH, RH; software: LH, BG; funding acquisition, writing – original draft: LH, OP.

Conflicts of interest

The authors declare that they have no conflict of interest.

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