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Technical Innovation

Low cost lab-on-a- chip prototyping with a consumer grade 3D printer

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Versatile prototyping of 3D printed lab-on-a-chip devices, supporting different forms of sample delivery, transport, functionalization and readout, is demonstrated with a consumer grade printer, which centralizes all critical fabrication tasks. Devices cost 0.57 US$ and are demonstrated in chemical sensing and micromixing examples, which exploit established principles from reference technologies.

Classical microfabrication techniques support the complete range of lab-on-a-chip (LOC) configurations and functions, however, they involve specialized resources and skills beyond the scope of many potential LOC users. Simplified processes have been explored, leading to the elimination of photolithographic masks and even to the use of routine microscopes as microfabrication platforms. Still these approaches require dedicated skills and clean room services, for complementary processes.

Direct fabrication of 3D structures can be subtractive, such as in laser writing and computer numerical control milling, or additive like in thermoplastic extrusion systems and micro stereo lithography platforms. Relative advantages of these and other existing 3D structuring technologies have been reviewed and refer to type and variety of materials they can employ as well as the platform cost, achievable resolutions, surface roughness, complexity, cost of materials and maintenance.

LOC microfabrication with 3D printers was initially demonstrated with custom-made systems, and later extended to more affordable additive printing systems able to combine different materials for reactionware and millifluidics. Recently, we demonstrated micro fabrication of templates for classical PDMS on glass LOC (PDMS-LOC) using consumer grade 3D micro stereo lithography (MS) printers (2290 US$, Miicraft). The technique is compatible with usual LOC dimensions and delivers sub-micrometric surface roughness; however, further exploration of 3D printing possibilities could entirely eliminate templates and transfer all fabrication specialized tasks to the printer, thus completely disengaging design complexity from fabrication skills. The implications entail custom LOC access with minimum infrastructure and highly versatile microfabrication compatible with flexible prototyping and immediate exploitation.

The value of a prospective fabrication platform can be gauged by its ability to accommodate different designs and principles using essentially the same tool. This work demonstrates fabrication of LOC devices, designed around a 3D printed single monolithic body, or unibody LOC (ULOC), which can be easily completed with simple complementary materials and procedures.

The versatility of ULOC is demonstrated through the integration of printed connectors, complex geometric features to facilitate functionalization, the incorporation of lateral flow transport capabilities, and the inclusion of glass micro beads in continuous flow mixers.

This first demonstration of ULOC illustrates passive and active transport as well as other accessory functions that can be directly migrated from PDMS-LOC, lateral flow and micro beads supported systems. The devices are compatible with fluorescence microscopy and color imaging readout, and were applied to H$_2$O$_2$ and glucose detection.

Devices in this work cost in average 0.57 US$/device, and can be fabricated within 30 min, thus offering an affordable and agile breadboard for LOC optimization.

ULOC devices described in this work are possible due to the resolution presently achievable with consumer grade MS 3D printers. Previous demonstrations of 3D printed fluidics, made with affordable thermoplastic extrusion systems, imposed restrictions on resolution and surface roughness, which prevented regular LOC channel geometries and simple sealing procedures.

In contrast, last generation of affordable MS 3D printers, aided by digital micro mirror devices (DMD), deliver resolutions in the 50µm range, and surface roughness under 182nm (see ESI), which enables efficient sealing with regular adhesive tape or PDMS films. For similar resolutions other advanced additive systems are about 10 to 100 times more expensive that the Miicraft printer used in this work. On the other hand, those systems are capable of combining different materials and can print larger objects.

The printer software transforms computer-aided designs (CADs) into bitmap image collections (black and white png format), corresponding to cross-sections of the 3D design, spaced at 50µm steps in the vertical direction. Each of these images is sequentially displayed in the DMD element to cure each layer of the 3D printout. These bitmap images can be edited and provide an easy way to modify the design with common image retouching software, such as Microsoft Paint or Photoshop, instead of...
Lab on a Chip

Accordingly, different channel sectors must be able to seal independently, which in this case was solved with a 100µm thick roofed channel (detail Fig. 1a) bridging the space between sector1 and sector2. Sector2 was sealed with a PDMS film separately functionalized for H₂O₂ fluorescent detection, and the long part of the micro channels, under sector1, introduces a time delay that permits controlled delivery of fixed sample volumes, when the device is used for manual delivery. Sealing of regions 1, 3 and the backside, were completed with adhesive tape (3M Ruban Adhesive Scotch® Nastro Adhesive).

Defining features of a 3D design, such as channels, can be made along the z-axis, or on the xy-plane with the z-axis displacement configuring the thickness of the planar design. The first alternative is the natural mode of additive 3D printing, and in the case of thermoplastic extrusion systems there is not much difference in the final result. The situation drastically changes with MS 3D printers, where an entire low roughness xy-plane is exposed every 7s. To exploit this characteristic, unibodies are made planar, resulting in minimal roughness that allows complementary tape sealing. As a consequence long channels can be created open, not sacrificing reliability and with minimal texture. This use of the printing direction also permits to exploit the resolution limits, since 50µm deep channels are easier to clean from uncured resin than 500µm deep elements.

The same printer used in the conventional fabrication direction takes several hours to produce similar size prototypes, restricts the minimum channel cross section compatible with resin release to 1000µm×500µm, and leaves the quality surface finishing unexploited.

In ULOC, unexposed resin is removed by 20s sonication in industrial grade ethanol after which the unibody is air-dried and ready for assembly. Uncured resin can be problematic to evacuate from confined spaces, such as long roofed channels, and the 2mm long section separating sector1 and sector2 was a conservative compromise between the required functionality and fabrication reliability.

The blue resin composition is proprietary information, and the material safety data sheet only reports that is a modified acrylate oligomer and monomer in combination with an epoxy monomer, a photo initiator, and additives. UV-curable formulations use oligomers or prepolymer consisting of reactive sites based on functional acrylate groups to limit volatile emissions, whereas the use of epoxy enables versatile tailoring of properties such as viscosity, harness and chemical compatibility.

The surfaces of numerous commodity polymeric materials that are suitable with microfabrication are not inert and interfere with chemical detection or cell viability, thus requiring specific surface conditioning. In this work chemical functionalization is carried out in well-described substrates such as PDMS in the case of H₂O₂ detection and cellulose powder for glucose measurements. The ULOC assembly allows migrating these established solutions to the unibody. Printed structures are hydrophobic (93.4° contact angle in Fig. 1c), which complement well with the confinement of aqueous solutions and for the cellulose passive transport configuration (Fig. 2).

The use of printed templates for the fabrication of PDMS LOC showed that templates can be exposed to 65°C for several
hours and still be reused, suggesting sufficient thermal stability, although for the purpose of this work the entirely fabrication and characterization was made at room temperature.

A newer clear type of resin, which is reported with the same composition as the blue resin, has been shown compatible with droplet extraction using a decanol organic phase, and with electrophoretic separation. Custom formulations of epoxy acrylates are known to be compatible with LOC uses, and eventually tailoring the resin chemistry could expand the range of compatible uses, meanwhile the unibody offers a board for to integrate proven materials and solutions within a communal assembly.

PDMS was functionalized for H₂O₂ detection (Fig. 1e). H₂O₂ concentration in blood and urine has been suggested as a biomarker for oxidative stress in humans, which is associated with the development of numerous serious conditions such as cancer and heart failure.

For H₂O₂ detection with ULOC, connectors were assembled with silicon tubing (Fig. 1d), and 3µL of control solution (C), 0.1µM (S) and 1µM (H) H₂O₂ in saline solution were pipetted into the tubing, and later pumped into the device using an air loaded 5mL syringe. These concentrations test the lowest clinically relevant limit of H₂O₂ in urine for oxidative stress monitoring.

The device was designed for fluorescence microscope readout and the inset shows the three separately sealed channels in the detection zone (insert in Fig. 1e). Time-lapse acquisition, at 10s intervals, captured the assay time response from which a sensitivity of 15.22 intensity units/µM, with a resolution of 0.098µM, evaluated within the [10,12] min interval could be estimated.

Protocols for H₂O₂ detection in aqueous solution were adapted to ULOC from established configurations (see ESI). In this assay, horseradish peroxidase (HRP) catalyzed the de-N-acetylation and oxidation of non-fluorescent Ampliflu-Red by H₂O₂ to highly fluorescent resorufin. Resorufin is also a substrate for HRP, which leads to the total consumption of the fluorescent product after a while (Fig. 1e). Time response features depend on the initial concentration of H₂O₂, and can be adapted to the desired detection range with the assay proportions, as has been shown elsewhere.

To demonstrate the versatility of ULOC to accommodate different configurations, a lateral flow device was implemented around the unibody concept, for the detection of glucose. Monitoring of glucose in blood (glycemia) is central for control of diabetes mellitus, which affects 2% of the world population and 6% of adult population in the western world, and is the most representative biosensing target involving 85% of the biosensing market.

The ability to modify a core design, without extensive knowledge of CAD operation, is a bonus for biosensor developers whose focus is beyond microfabrication. With the 3D printer the exposure patterns are bitmap images that can be edited with image retouching software. In the present case, the first device (Fig. 1) was transformed into the lateral flow example (Fig. 2) using only Photoshop edition. The possibility to image edit exposure patterns can significantly shorten the workflow during the development phase, since it avoids CAD re-edition, slicing and control procedures, by just correcting the few bitmaps that define the design.

Fig. 2a shows the fabricated device with channels filled with cellulose powder paste, which were locally functionalized for glucose detection. The image corresponds to the final response after the sensing surface has been exposed to a control (C) and two glucose concentrations (H and S) within the diagnostic range.

Cellulose powder is a well-established material for thin layer chromatography and provides passive capillary transport in this lateral flow configuration. The hydrophobic channels in the printed unibody, naturally confine the flow within the cellulose powder, which can be easily delivered into the channels by simple procedures (see ESI).
with the sample, and once dried, a 10 min interval was allowed for response stabilization, after which the average response in the last 15 min was used for quantification, rendering the observed linear behavior (Fig. 2c). From these measurements a sensitivity of 3.9 intensity units/mM with a resolution of 0.56 mM (or 0.50 mg/dL) could be estimated.

The assay uses a colorimetric detection principle (see ESI). In the presence of glucose and oxygen, glucose oxidase (GOx) catalyzes the oxidation of glucose to glucono delta-lactone and hydrogen peroxide, which is used by HRP to catalyze the oxidation of colorless iodide to brown iodine monitored in the measurement.

The final LOC functionality tested here in ULOC configuration is the integration of micromixers. Mixing is a common preparatory stage in chemical analyses, and accordingly, an important function for LOCs that must be supported by any prospective fabrication method. Microfluidic dimensions favor laminar flow regimes, where mixing occurs as a slow diffusional process, and different designs exist to favor turbulence and better mixing performance. Chaotic mixers are one possibility to improve performance, and have been demonstrated on PDMS-LOC created on 3D printed templates.

A diffusional mixer (Fig. 3b), without the through hole, was used to compare performances, and shows 2 unmixed streams at the distal end of the device (highlighted by a white cross). When the through hole feature was added (Fig. 3c), the mixing behavior changed and the flow divided in three apparent streams, almost entirely mixing (single color) at the end of the device (~80% of the channels length), whereas with the beads (Fig. 3d), only one color stream could be distinguished at ~40% of the channel length, thus providing a compact possibility to double the mixing efficiency.

Results in this work demonstrate the possibility of 3D printed LOC devices compatible with different sample delivery, transport and detection approaches. In all cases, migration from established solutions, in classical LOC, was shown possible and simple, whereas ULOC fabrication demands minimal efforts and infrastructure.

The roofed channel used to separate independent sealed sectors (Fig. 1) also illustrates the advantage of direct additive printing respect to replicas from templates. Independently of the technology used to create the template, a seemingly trivial roofed element cannot be produced in a single manufacturing step. On the other hand, templates are reusable, whereas direct printing implies reprinting each new device. Unibody optimizes additive prototyping by minimizing the device thickness and by exploiting the planar dimension, where the printer can deliver sub micrometric surface roughness and reliable resin removal in small features over long distance ranges.

**Conclusions**

In this work, affordable 3D printed LOC devices have been demonstrated. Complex geometries, directly created in 3D printed structures, enable transference of demanding fabrication tasks to the printer, thus maximizing reliability and removing the influence of user fabrication skills from the prototypes.

These capabilities have been investigated with devices serving H2O2 and glucose detection, using passive and active transport, as well as configurations suitable with fluorescence and colorimetric readout. In addition with continuous flow micro mixers, these results demonstrate a versatile set of essential configurations for LOC systems.

Consumer grade MS 3D printers not only simplify access to custom LOC designs, but can also accommodate sophisticated features only accessible in advanced LOC fabrication techniques. These aspects were shown possible, at a convenient development cost of 0.57 US$/device, and for prototype fabrication times under 30 min.

3D printed ULOC devices offer a breadboard for experimentation that is compatible with fast and affordable prototype optimization, which can exploit established LOC solutions.

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Notes and references

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