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Three-Dimensional Large-Scale Fused Silica Microfluidic Chips Enabled by Hybrid Laser Microfabrication for Continuous-Flow UV Photochemical Synthesis

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Abstract: We demonstrate a hybrid laser microfabrication approach, which combines the technical merits of ultrafast laser-assisted chemical etching and carbon dioxide laser-induced in situ melting for centimeter-scale and bonding-free fabrication of 3D complex hollow microstructures in fused silica glass. With the developed approach, large-scale fused silica microfluidic chips with integrated 3D cascaded micromixing units can be reliably manufactured. High-performance on-chip mixing and continuous-flow photochemical synthesis under UV irradiation at ~280 nm were demonstrated using the manufactured chip, indicating a powerful capability for versatile fabrication of highly transparent all-glass microfluidic reactors for on-chip photochemical synthesis.

Keywords: ultrafast laser direct writing; chemical etching; carbon dioxide laser processing; 3D glass microfluidics; fused silica; continuous-flow photochemical synthesis

1. Introduction

Continuous-flow synthesis, in which a chemical reaction is performed in a flowing and confined environment (e.g., microchannels) other than conventional batch methods based on stirring, has attracted extensive attention in the past decade due to its unique advantages such as enhanced mass transfer, high throughput, low waste, and high safety for manufacturing pharmaceuticals and fine chemicals [1–6]. In the continuous-flow synthesis, reliable fabrication of the microchannel reactors is vital for practical industrial applications. As one of the most popular substrate materials, glass has many advantages such as high optical transparency, excellent chemical inertness, and low thermal expansion coefficients compared to other materials (e.g., polymers, silicon, etc.) for microreactor applications [7–12]. Particularly, fused silica has a wide transmission window from deep ultraviolet to infrared regions (200–3500 nm), which is an ideal substrate material for the versatile fabrication of photochemical microchannel reactors. To fabricate a closed glass-based microchannel reactor, the bonding of patterned glass workpieces is usually unavoidably employed using conventional fabrication methods such as lithographic methods and precision molding [12]. The validity and stability of bonding performance significantly affect the long-term operation of those microreactors, which increases the requirements of high-level surface quality and of the bonding...
environment of the glass workpieces. Currently, there is an increasing demand for the fabrication of microchannel reactors with three-dimensional (3D) configurations, which allows enhancement of the efficiency and the throughput of the device [13–16]. However, to form such a device with conventional methods will further increase the cost of the multiple-step bonding process. Therefore, developing advanced fabrication technologies of 3D microchannel reactors in a bonding-free manner is highly desirable.

Ultrafast laser-assisted chemical etching of glass based on nonlinear multiphoton absorption provides a unique approach for the 3D fabrication of hollow glass microchannel structures [17–19]. In this technique, ultrafast laser direct writing in glass allows in-volume processing of 3D, selectively modified regions under proper pulse energy. The modified regions can be further removed by etching solutions such as diluted HF solutions and hot KOH solutions to form the hollow channel structures [20–22]. In general, when the required lengths of microchannels are less than several centimeters, this technique can perform controllable fabrication of homogeneous microchannels without any bonding process. However, there are some technical barriers for fabricating long and large-volume microchannels with controllable feature sizes using this technique due to the inherent limitations of ultrafast laser-induced etching selectivity of glass [20–22]. Previously, to fabricate 3D centimeter-scale, high-throughput micromixers in fused silica glass, a new bonding process was developed [23]. To beat the limitation of the etching selectivity, the introduction of extra-access ports has been proposed before by several groups [24–27]. Recently, we have demonstrated that 3D all-glass microfluidic channels with arbitrary lengths can be fabricated using a hybrid laser microfabrication scheme based on a combination of ultrafast laser microfabrication and CO	extsubscript{2} laser irradiation of glass [28]. In the proposed hybrid scheme, ultrafast laser direct writing was employed to induce spatially selective modification inside the glass for subsequent chemical etching of hollow glass microstructures, including 3D microchannels and extra-access ports which were used for improving etching homogeneity of microchannels; and while defocusing CO	extsubscript{2} laser irradiation was employed to seal the etched extra-access ports on the glass surface to form a closed microfluidic structure with a few inlets and outlets. In this work, we demonstrate further progress on 3D large-scale fused silica microfluidic chips with integrated cascaded micromixing units that can be reliably manufactured using the improved hybrid scheme. With the improvement of CO	extsubscript{2} laser processing parameters, nearly perfect sealing of these ports has been realized controllably. Furthermore, high-performance on-chip mixing of dye solutions was demonstrated using the fabricated chip. Furthermore, as for proof-of-concept demonstration, continuous-flow on-chip UV photochemical synthesis based on a photocycloaddition reaction has been realized using the same chip, indicating a powerful capability for versatile manufacturing of highly transparent all-glass microfluidic systems for on-chip photochemical synthesis.

2. Materials and Methods

2.1. Fabrication of 3D Large-Scale Glass Microchannels

As illustrated in Figure 1a, the fabrication of 3D large-scale microchannels in fused silica consists of three main steps: (i) ultrafast laser direct writing, (ii) selective chemical etching, and (iii) defocusing CO	extsubscript{2} laser irradiation. For the ultrafast laser direct writing, fused silica glass plates (JGS2) with various sizes (e.g., 155 mm × 125 mm × 2 mm shown in Figure S1) were used for processing substrates and a laser amplifier system (Pharos 20 W, Light Conversion, Vilnius, Lithuania) with a central wavelength of 1030 nm, a pulse duration of 10 ps, and a repetition rate of 250 kHz was used for laser sources, respectively [29]. First, the glass plates were fixed on a 3D air-bearing stage system (Aerotech 3D X-Y-Z stage, Aerotech, Pittsburgh, PA, USA) with a positioning precision of 0.5 μm. Then, the laser beam was focused inside glass using a microscope objective (M Plan Apo NIR 10X, Mitutoyo, Kawasaki, Kanagawa, Japan) with a numerical aperture of 0.26 and a transmission rate of 80%. To create 3D modified micropatterns in glass, the laser direct writing was first performed through line-by-line scanning in a single layer from the bottom of the patterns and then moved upward in a layer-by-layer manner to the top
of the patterns (see Figure S2). The writing speed, line-by-line spacing, and layer-by-layer spacing were set at 65 mm/s, 30 μm, and 37.5 μm, respectively. As illustrated in Figure 1b, the laser-modified micropatterns include a 3D microchannel combined with an array of extra-access ports which connect the glass surface and the top of the channel. As previous works demonstrated [24–28], the introduction of extra-access ports provides a more homogeneous etching of the whole channel to beat the limitation of the inherent etching selectivity of glass. For the chemical etching, a mixed aqueous solution with a concentration of 10 M KOH and 1 M NaOH was used. The laser-modified glass samples were immersed in the etching solution in an ultrasonic bath with a temperature of ~90 °C. After chemical etching, the 3D hollow glass channel structures were obtained, and the etched extra-access ports were through from the glass surface to the inside of the 3D channel. For a glass plate with a size of 155 mm × 125 mm × 2 mm (see Figure S1), the whole etching time was ~48 h. In addition, the distance between the top of the channel and the surface reduced from 1 mm to 700 μm, and the height of the channel increased from 375 μm to 500 μm. For defocusing CO2 laser irradiation, the etched glass plates were placed in a 2D stage (OneXY-500-500-AS-CMS1, Coretech, Wuxi, Jiangsu, China). The openings of the extra-access ports were irradiated by the CO2 laser beam (FSTi100SWC, Synrad, Mukilteo, WA, USA) with a wavelength of 10.6 μm and a repetition rate of 20 kHz one after another in a defocused manner using a ZnSe lens (LA7028-E3, Thorlabs, Newton, NJ, USA) with a measured focal length of ~159 mm and finally sealed to form a closed 3D large-scale microchannel with several inlets and outlets. With a defocusing distance of 21 mm, the irradiated diameter of the CO2 laser beam on the glass surface was ~1 mm. Different laser processing parameters such as CO2 laser power and irradiation time were investigated to optimize the sealing.

Figure 1. Schematic of the fabrication procedure for a 3D large-scale microchannel chip with two inlets and one outlet in fused silica. It consists of three main steps: (i) ultrafast laser direct writing of 3D modified micropatterns in glass; (ii) selective chemical etching of hollow glass microstructures including 3D microchannels, micromixing units, and open extra-access ports; and (iii) defocusing CO2 laser irradiation for sealing the extra-access ports. (b) Close-up view of a 3D micromixing unit with several extra-access ports indicated by a dashed rectangle at each step in (a).
2.2. Continuous-Flow on-Chip Photochemical Synthesis

For on-chip photochemical reactions, a fabricated fused silica microfluidic chip with a size of 100 mm × 100 mm × 3 mm was closely arranged in front of a UV LED light source with a wavelength of ~280 nm and an illuminated area of 90 mm × 75 mm. To perform high-performance on-chip mixing, 158 micromixing units with a cross-section of 2 mm × 1 mm were integrated into a microchannel inside the microfluidic chip. The length and thickness of the fabricated microchannel are ~390 mm and 1 mm, respectively. All reagents for on-chip continuous-flow synthesis were commercially available (Shanghai Aladdin Biochemical Technology Co., Ltd., Shanghai, China) and used without any purification. Reaction solutions were first prepared by dissolving maleimide (0.1 M) and 1-hexyne (0.15 M) into acetonitrile (MeCN), respectively. Then, both prepared solutions were simultaneously pumped into the 3D microfluidic chip through two inlets with the same flow rate ranging from 0.03 to 0.1 mL/min, respectively. The power of the UV LED light source was set at 160 mW. The products were collected from the outlet of the chip into an opaque glass bottle for subsequent nuclear magnetic resonance (NMR) measurements.

2.3. Characterization

The morphologies of the laser-fabricated glass structures were recorded by a polarized optical microscope (BX53, Olympus, Tokyo, Japan). Additionally, 1H NMR spectra of the on-chip continuous-flow synthesized products were recorded at 500 MHz (AVANCE III HD500, Bruker, Swiss). For NMR characterization, the products were solved in the deuterated chloroform, and tetramethylsilane (TMS) was used as an internal standard.

3. Results and Discussions

3.1. Controllable Sealing of Extra-Access Ports on Laser-Fabricated Glass Microchannels

Figure 2a shows cross-sectional optical micrographs of a laser-fabricated embedded microchannel structure with a ~240 µm diameter extra-access port (see Figure S2) after different defocusing CO₂ laser irradiation conditions. By using defocusing CO₂ laser irradiation-induced glass in-situ melting, the extra-access port can be fully sealed to form a combined multilayer structure, including a surface crater, a sealed layer, and a residual port from top to bottom, as indicated in the top-left panel of Figure 2a. The height of the sealed layer and residual port can be well-controlled by tuning the laser power and irradiation time of the CO₂ laser. As shown in the top part of Figure 2a, when the laser power was set at 29.7 W, with the increase in irradiation time from 4 s to 10 s, the height of the sealed layer increased while the height of the residual port decreased, indicating the enhancement of time-dependent glass in-situ melting effects. With further increases in laser power to 31.3 W (the middle part of Figure 2a) and 32.6 W (the bottom part of Figure 2a), the height of the sealed layer continuously increased while the height of the residual port continuously decreased when the irradiation time was prolonged. Particularly, when the irradiation time was 10 s at a laser power of 32.6 W, the height of the residual port was nearly eliminated in which there was almost no dead-end (nearly perfect sealing) in the fabricated microchannel structure.

To further quantitively analyze the sealing performance of CO₂ laser processing of extra-access ports, Figure 2b,c plot the dependence of the height of sealed layer and residual port on irradiation time at different laser powers, respectively. As clearly presented in Figure 2b,c, the extension of the irradiation time could enable the increase in the height of the sealed layer and the reduction of the height of the residual port, which exhibited the same tendency as the optical micrographs of Figure 2a and Figure S3a. Particularly, in the case of 10 s irradiation at 32.6 W, a ~320 µm height sealed layer was obtained while the corresponding residual port was nearly eliminated. However, for a large-scale (e.g., centimeter-scale) microfluidic chip that includes many extra-access ports to be sealed, reducing the sealing time for a single port may promote the processing efficiency for the whole chip. Therefore, there is a tradeoff between irradiation time and sealed performance. On the one hand, the port must be sealed with a properly sealed height (thickness) and a
small residual volume in a certain amount of time for ensuring pressure-resistant microfluidic applications. On the other hand, the irradiation time must be kept in a short duration for rapid manufacturing.

![Figure 2.](image-url)

**Figure 2.** (a) Cross-sectional optical micrographs of a microchannel structure with an extra-access port after defocusing CO$_2$ laser irradiation at different laser powers (29.7, 31.3, and 32.6 W) and irradiation times (4, 6, 8, and 10 s). After CO$_2$ laser irradiation, an open extra-access port can be sealed to form both a sealed layer while a surface crater is created and a part of the port residues depending on specific conditions. Scale bar represents 1 mm. (b) Height of sealed layer versus CO$_2$ laser irradiation time under different laser powers. (c) Height of residual port versus CO$_2$ laser irradiation time under different laser powers.

Besides investigating the size dependence of sealed ports on CO$_2$ laser processing parameters, polarized microscopic observation of the sealed ports was also performed. As shown in Figure 3 and Figure S3b, the polarized micrographs reveal the distribution of anisotropic birefringence around the ports after defocusing CO$_2$ laser irradiation. With the same laser power, the extension of irradiation time leads to the reduction of anisotropic birefringence, which promotes the homogeneity of the distribution of residual stress. Meanwhile, in the same irradiation time (6–10 s), the increase in laser power creates a similar tendency of stress control around the port. Particularly, in the case of 10 s at
32.6 W, there is almost no anisotropic birefringence at the periphery of the port, indicating low residual stress in the condition of nearly perfect sealing as indicated in the bottom-right panel of Figure 2a.

![Figure 3](image_url)

**Figure 3.** Front-view polarized optical micrographs of a microchannel structure with an extra-access port after defocusing CO\textsubscript{2} laser irradiation at different laser powers (29.7, 31.3, and 32.6 W) and irradiation times (4, 6, 8, 10 s). After CO\textsubscript{2} laser irradiation, the stress distribution of a sealed port exhibits differently depending on irradiation conditions. Scale bar represents 0.5 mm.

### 3.2. Bonding-Free Fabrication of 3D Large-Scale Micromixing Fused Silica Chip

To demonstrate the capability of the proposed approach for 3D large-scale fabrication of fused silica microfluidic chips, a T-shape centimeter-scale glass microchannel chip which included a string of 3D micromixing units was fabricated (channel length: ~51 mm). Figure 4a,b show a schematic of fabricated 3D micromixing glass chips and photos of fabricated glass structures at each step, respectively. As illustrated in the inset of Figure 4a, the 3D micromixing unit is based on Baker’s transformation, which consists of splitting, routing, and converging the fluids for high-efficiency mixing in 3D spaces [13,14,30]. Figure 4c shows cross-sectional optical micrographs of a 3D micromixing unit connected with an extra-access port at each fabrication step, as described in Figure 4b. One can see in the left panel of Figure 4c that after ultrafast laser direct writing, 3D laser-written tracks which include the designable patterns of a 3D embedded microchannel beneath the glass surface and an extra-access port connected with the glass surface and the channel were created in a spatially selective manner. After selective chemical etching, the laser-modified micropatterns were transferred into a hollow, glass-channel structure including a 3D micromixing unit connected with an open port (see the middle panel of Figure 4c). Finally, the port can be sealed by defocusing CO\textsubscript{2} laser irradiation (4 s, 29.7 W) to form a 3D closed micromixing unit, and then those mixing units can be cascaded to form a T-shape, high-efficiency mixing chip with two inlets and one outlet in a robust and reproducible fashion.
left panel of Figure 4c that after ultrafast laser direct writing, 3D laser-written tracks which include the designable patterns of a 3D embedded microchannel beneath the glass surface and an extra-access port connected with the glass surface and the channel were created in a spatially selective manner. After selective chemical etching, the laser-modified micro-patterns were transferred into a hollow, glass-channel structure including a 3D micromixing unit connected with an open port (see the middle panel of Figure 4c). Finally, the port can be sealed by defocusing CO$_2$ laser irradiation (4 s, 29.7 W) to form a 3D closed micromixing unit, and then those mixing units can be cascaded to form a T-shape, high-efficiency mixing chip with two inlets and one outlet in a robust and reproducible fashion.

In principle, with the introduction of extra-access ports and subsequent port sealing, an all-glass microchannel structure with nearly arbitrary length and uniform size can be fabricated in glass using the proposed approach. However, regarding the inherent nature of chemical etching and the controllable port sealing, the available width and thickness of a channel structure and spatial spacing between channels vary depending on specific configurations. To promote the throughput of the microchannel chip, increasing the channel thickness to several mm is attainable; however, it will increase the etching time and the sealing difficulties. In addition, decreasing the spatial spacing between channels to a short distance (e.g., ~0.1 mm) is also possible when the layout and sizes of ports and the etching and sealing process are applicable. For instance, to further improve the etching, possible methods include optimization of the extra-access port design (spacing between adjacent ports, spatial shape, size, etc.), laser processing parameters (writing scheme, pulse energy, writing speed, repetition rate, etc.) and etching conditions (temperature, etchants, ultrasonic control, etc.), which will be explored in the future.

3.3. On-Chip UV Photochemical Synthesis

To verify the mixing performance of the fabricated fused silica chip, yellow and blue dye solutions were pumped into a fabricated fused silica chip (workpiece size:
100 mm × 100 mm × 3 mm) with a flow rate of 20 mL/min through two inlets shown in the left panel of Figure 5a, respectively. With the embedded mixing units (the same as Figure 4a) in the chip, the color of the mixing fluids quickly changed into green at the beginning of the entrance of the chip, indicating its high-performance mixing capability with a high throughput manner due to 3D configuration of the chip for manipulation of fluids. The capability of the high-throughput efficient mixing can also be identified by a laser-fabricated 3D mixer with a similar configuration in the previous report [23]. Moreover, by replacing the dye solutions with red and blue solutions, the purple solutions can be quickly obtained, as shown in the right panel of Figure 5a.

Figure 5. (a) Demonstration of high-efficiency mixing in a fused silica microfluidic chip with a size of 100 mm × 100 mm × 3 mm using different dye solutions. Left: yellow and blue dye solutions were pumped into two inlets with a flow rate of 20 mL/min, respectively; Right: red and blue dye solutions were pumped into two inlets with the same flow rate, respectively. (b) a [2 + 2] photocycloaddition reaction for continuous-flow photochemical synthesis. (c) Photo of an assembled fused silica microfluidic chip and a UV LED light source for on-chip photochemical reaction. (d) The 1H NMR spectra of the synthesized product using continuous-flow on-chip synthesis with and without UV (~280 nm) irradiation.
Regarding superior optical transmission properties of fused silica ranging from 200 nm to 3500 nm, on-chip UV photochemical synthesis using the same glass chip was performed. Cycloaddition reactions are reliable and effective in the synthesis of polycyclic compounds which are important for pharmaceutical engineering \[31–33\] and fine chemical engineering \[34,35\]. These reactions are usually driven by heat or light. Among the cycloaddition reactions, \([2 + 2]\) photocycloaddition, which is driven by economical light irradiation, is widely used in the synthesis of natural products and medicines such as Taxol \[36\], one of the most commonly used anticancer agents, and ent-kaurane diterpenoids \[37–39\], an important group of natural products with antibacterial and anti-inflammatory activities. Therefore, as for proof-of-concept demonstration, a \([2 + 2]\) photocycloaddition reaction (see Figure 5b) for continuous-flow on-chip photochemical synthesis was investigated.

As illustrated in Figure 5b, maleimide and 1-hexyne react under UV irradiation (\(\lambda \approx 280 \text{ nm}\)). To perform highly effective UV irradiation, the assembled fused silica chip in Figure 5a was first closely attached to the illumination surface of a UV LED light source (\(\approx 280 \text{ nm}\)) (see Figure 5c) and performed the on-chip photochemical synthesis. Besides ensuring on-chip mixing performance, a controllable residence time of the mixed fluids in the illuminated area is indispensable regarding possible limitations of the reaction rates and the completeness of the reactions, especially for high-throughput synthesis. To ensure a certain residence time for the photochemical reaction, most of the 3D microchannel (length: \(\approx 390 \text{ mm}\)) was located in the illuminated area (90 mm \(\times\) 75 mm). The synthesized product was collected and dried under vacuum at 25 \(^\circ\)C, then redissolved in deuterated chloroform for further NMR characterization. The 1H NMR spectra of the products obtained without and with UV irradiation are shown in Figure 5d. A singlet peak which was remarked as No. 4 hydrogen represents the sign of the target molecule. The reaction happens under UV irradiation since the singlet peak at 1.9 ppm emerged after irradiation, confirming the formation of the target molecule. In contrast, no reaction happens without UV irradiation. This result confirmed that a \([2 + 2]\) photocycloaddition reaction can work in the fabricated fused silica microfluidic chip under UV irradiation at \(\approx 280\) nm.

4. Conclusions

Controllable fabrication of large-scale fused silica microfluidic chips with 3D configurations has been demonstrated using the hybrid strategy based on a combination of ultrafast laser microfabrication and CO\(_2\) laser microprocessing. The introduction of extra-access ports connected with 3D microchannels allows the bonding-free manufacture of microfluidic chips with a workpiece size at a centimeter-scale level and a feature size at several hundreds of microns by beating the inherent limits of laser-induced etching selectivity of glass. Nearly perfect sealing of the ports has been stably obtained by optimizing the defocusing CO\(_2\) laser irradiation. Further, fabrication of centimeter-scale microfluidic glass chips with cascaded micromixing units for high-efficiency and high-throughput mixing has been achieved, and the function of on-chip UV photochemical synthesis at \(\approx 280\) nm has been verified. Besides the applications of on-chip high-throughput mixing and UV photochemical synthesis, the proposed approach holds great potential for advanced manufacturing of 3D large-scale, high-precision biomimetic microfluidic structures with complex geometries (e.g., organs-on-chips) and large-scale, high-density integration of high aspect ratio microchannels (e.g., high-resolution micro-separation chips) on glass substrates. Regarding the unique capability of ultrafast laser microfabrication, the proposed approach provides a versatile route for the high-performance fabrication of large-scale fused silica microfluidic chips with monolithically multifunctional integration such as optofluidics and electrofluidics \[20\] for state-of-the-art sensing and detection, which will pave the way for rapid manufacturing of “all-in-one” glass-based microfluidic smart microsystems.
Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/mi13040543/s1. Figure S1: Photos of a fused silica glass plate with a size of 155 mm × 125 mm × 2 mm at each fabrication step, Figure S2: Top-view (a) and side-view (b) optical micrographs of a microchannel with an extra-access port after ultrafast laser modification and after chemical etching; Figure S3: (a) Cross-sectional optical micrographs and (b) front-view polarized optical micrographs of a microchannel structure with an extra-access port after defocusing CO2 laser irradiation at different laser powers (29.7, 31.3, and 32.6 W) and irradiation times (3 and 5 s).

Author Contributions: Conceptualization, Y.C. and J.X.; methodology, J.X. and M.H.; software, A.Z., Y.S., J.Q. and W.C.; validation, A.Z., J.X., Y.L., M.H., Z.L. (Zijie Lin) and Z.L. (Zhaoxiang Liu); investigation, A.Z., Y.L., Z.L. (Zijie Lin), Y.S., J.Q., W.C. and Z.L. (Zhaoxiang Liu); resources, J.X., M.H. and Y.C.; writing—original draft preparation, A.Z., J.X. and Y.C.; writing—review and editing, J.X., M.H. and Y.C.; supervision, Y.C.; funding acquisition, J.X. and Y.C. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the National Natural Science Foundation of China (Grant Nos. 12174107, 61991444, 11933005, 11874060, and 11734009), National Key R&D Program of China (Grant No. 2019YFA0705000), and Science and Technology Commission of Shanghai Municipality (Grant No. 21DZ1101500).

Conflicts of Interest: The authors declare no conflict of interest.

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