

Review

Two-Photon Polymerization: Functionalized Microstructures, Micro-Resonators, and Bio-Scaffolds

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Abstract: The direct laser writing technique based on two-photon polymerization (TPP) has evolved considerably over the past two decades. Its remarkable characteristics, such as 3D capability, sub-diffraction resolution, material flexibility, and gentle processing conditions, have made it suitable for several applications in photonics and biosciences. In this review, we present an overview of the progress of TPP towards the fabrication of functionalized microstructures, whispering gallery mode (WGM) microresonators, and microenvironments for culturing microorganisms. We also describe the key physical-chemical fundamentals underlying the technique, the typical experimental setups, and the different materials employed for TPP.

Keywords: direct laser writing; ultrashort laser pulses; two-photon polymerization; functional microdevices; whispering gallery mode microresonators; scaffolds for biological applications



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1. Introduction

The processing of materials by ultrashort lasers, which allows high precision-patterning of 2D and 3D micro/nanostructures, has become a cornerstone technology for a wide range of areas related to academic research and engineering [1]. An ultrashort laser-based fabrication technique that has been widely used is direct laser writing via two-photon polymerization (TPP) [2,3]. As opposed to the most conventional micro/nanofabrication techniques, TPP is a maskless patterning tool and does not require harsh processing conditions nor a cleanroom facility [4]. Furthermore, this technique is not limited to the processing materials in the powder form, such as selective laser sintering (SLS) [5–8], another laser fabrication method that allows sculpting macroscopic structures with micrometric resolution. TPP leverages the nonlinear nature of two-photon absorption to fabricate 3D polymeric microstructures with arbitrary geometries and sub-diffraction features [9]. Moreover, a large variety of materials can be employed as photoresist, or be incorporated into it, which allows customizing the microstructures’ physical, chemical or biological properties to meet specific applications [10,11]. For example, improvements in the spatial resolution and the amount of dry-related shrinkage undergone by the microstructures can be accomplished by varying the photoresist components proportion [1,12–14].

The TPP technique has found applications in numerous fields, including photonics [15–19], biosciences [20–22], and micromechanical systems [23–25]. Precise photonic building blocks, such as optical microresonators [26], photonic crystals [27], and waveguides [28], as well as photonic wire bondings [29] and free-form coupling elements [16], have been successfully fabricated by TPP. Active photonic structures have also been fabricated [30,31]. In the field of biological sciences, TPP has mainly been used to create microscaffolds suitable for culturing cells, tissues, and microorganisms [21,32,33], and microneedles [22] and porous microstructures [20] for drug delivery. TPP has also been

a remarkable tool for fabricating optically driven micromachines that allows grabbing or probing nanoscale objects [34]. Microstructures containing movable parts, such as nanotweezers [23], microgears [24], and micropump setups [35], have been demonstrated.

This paper summarizes the advances of TPP towards three of its major applications: functionalized microstructures, whispering gallery mode (WGM) microresonators, and microscaffolds for culturing microorganisms. The favorable properties of TPP, such as fine-tuning of the structure dimensions, the integration of the fabricated structures into a range of substrates, and its material flexibility, have pushed forward the design and potential of applications of WGM microresonators. Moreover, the material flexibility of TPP, along with its 3D capability and gentle processing conditions, has made it well suited for the fabrication of bioactive microscaffolds with spatially varying properties.

The further sections of the present review are organized as follows: In Section 2, we briefly describe the physical-chemical fundamentals underlying the TPP technique. We subsequently show the typical experimental setups and different materials employed in TPP. Section 3 provides an overview of the general applications of TPP. Emphasis is given to the fabrication of WGM microresonators and 3D microscaffolds to study microorganism's development. Finally, Section 4 presents the overall conclusions and prospects of TPP in the near future.

2. Two-Photon Polymerization (TPP) Fundamentals

2.1. Two-Photon Absorption

The spatial confinement of TPP, which allows fabricating 3D microstructures with arbitrary geometry and submicrometric features, arises from the nonlinear nature of the two-photon absorption (TPA) phenomenon. The latter refers to an electronic transition excited by the combined action of two photons [2,36], such that the sum of the energy of the individual photons must be resonant with the transition energy [37]. When the photons have the same energy, as indicated in Figure 1b, the process is called degenerated [38]. TPA can also be observed with photons of different energies since the sum of their energies matches the energy gap.

In a quantum mechanics framework, TPA is described by considering the existence of a virtual intermediate state, which takes part in the electronic transition [36,39]. In this picture, a transition to a real state is possible only if the time interval between the two-photon is shorter than the lifetime of the virtual state [39]. Because the virtual state lifetime is extremely short, in the order of femtoseconds [37], TPA is usually described as the “simultaneous” absorption of two photons.

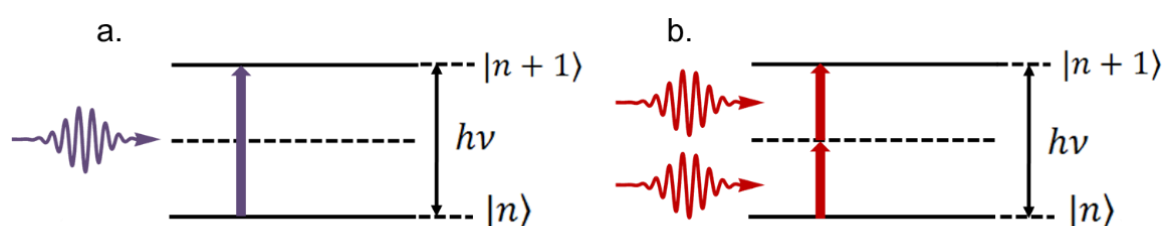


Figure 1. Energy diagram representing the absorption of (a) one and (b) two photons with the same energy to promote the electronic transition between the states $|n\rangle$ e $|n+1\rangle$. Reprinted with permission [40]. Licensed under a Creative Commons Attribution 4.0 International license.

The TPA is a nonlinear optics phenomenon that can occur in the presence of high-intensity light, i.e., when the magnitude of the light electric field is comparable to the interatomic electric field ($E_{int} \approx 10^{11} \text{ V}\cdot\text{m}^{-1}$) [36]. In this regime, the electric field induced in the material responds nonlinearly with the electric field of light, which can prompt unusual effects such as light generation at frequencies other than that of the incident light and absorption of light in the frequency range of transparency of the material [36]. When

high-intensity light is employed, the absorption becomes dependent on the excitation intensity according to [36,38]:

$$\alpha(I) = \alpha_0 + \beta I, \quad (1)$$

in which, α_0 and β are, respectively, the linear absorption and TPA coefficient, and I is the light intensity. β is directly associated with the TPA cross-section (σ_{TPA}) as stated in Equation (2) [38]:

$$\beta = \frac{N\sigma_{TPA}}{h\nu}, \quad (2)$$

in which, N , h , and ν are the concentration of entities that constitute the material (e.g., atoms, molecules, etc.), the Planck constant, and the pump frequency, respectively. For most molecules, the TPA cross-section (σ_{TPA}) is in the order of $10^{-50} \text{ cm}^4 \cdot \text{s} \cdot \text{photon}^{-1}$ or 1 GM. Such unit is a reference to the Nobel-laureate physicist Maria Göppert-Mayer [38,41].

As light propagates throughout the material, its intensity is attenuated exponentially [37]. For light in the frequency range of transparency of the material, for which $\alpha_0 = 0$, the light dissipation rate along the propagation direction (z) is described by [37]:

$$\frac{dI(z)}{dz} = -\beta I^2(z). \quad (3)$$

From Equation (3), we see that the dissipation rate for a TPA process depends quadratically on the light intensity. The nonlinear dependence of TPA on the intensity of light, together with the relatively small TPA cross-section of materials, allows attaining spatial confinement of the excitation when focused high-intensity lasers are used, such as Ti:sapphire lasers [38]. An illustration comparing the light dissipation produced by one and two photons is presented in Figure 2. For the linear absorption, the light intensity is exponentially attenuated as it propagates through the medium. In contrast, for the TPA case, attenuation is confined to the laser focal volume.

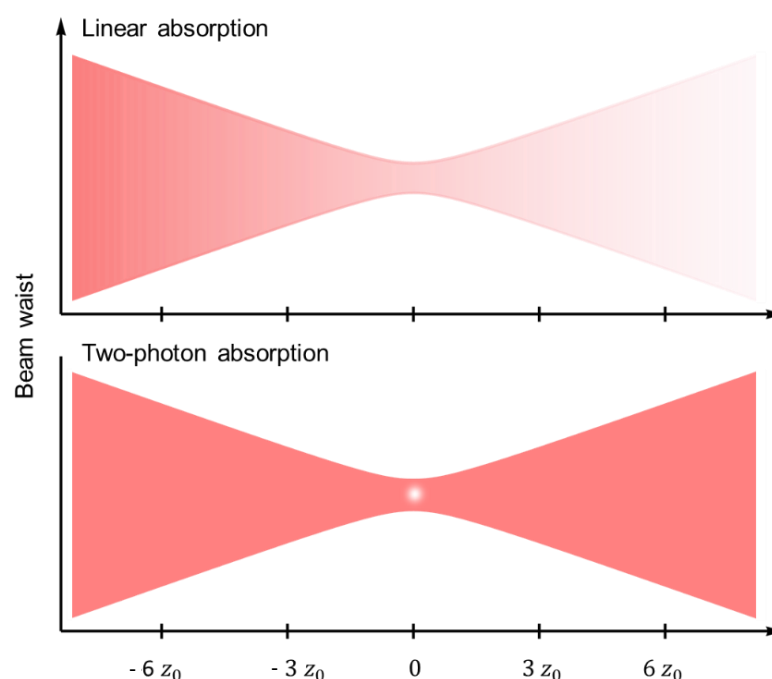


Figure 2. Attenuation of a focused laser beam prompted by the absorption of one (above) and two (below) photons. The white color indicates attenuation. The horizontal axis is given in units of the confocal parameter (z_0): the distance from the focal plane at which the beam waist increases by a factor of $\sqrt{2}$. Reprinted with permission [40]. Licensed under a Creative Commons Attribution 4.0 International license.

The spatial confinement of the excitation promoted by TPA allows to restrict the polymerization reaction to the focal volume of a high-intensity laser, which confers 3D capability and high resolution to the TPP technique [2]. Besides, the quadratic dependence of TPA on the light intensity pushes the resolution down by a factor of $\sqrt{2}$, which allows fabricating sub-diffraction features [42].

2.2. Two-Photon Polymerization (TPP)

Since the 1990s, TPP technique [3] has been widely used in the development of different technological devices due to a varied range of advantages. In this technique, the utilization of high-intensity pulse lasers allows the observation of nonlinear optical phenomena in the irradiated sample. Besides that, the polymerization of small volume (called voxels) [43] occurs only near the focal spot, giving the sculpted structure finer resolution (below to the diffraction limit) [9]. Moving the laser beam into the sample, it is possible to fabricate three-dimensional structures without shape limitation, with complex contours, including manipulable devices [9,44]. Samples employed in TPP usually are composed of monomers, hybrid sol-gels materials, hydrogels, or composite matrices [45,46]. These compounds are mixed into a photoinitiator, an organic molecule responsible for absorbing the laser energy and starting the polymerization procedure. The use of different monomers or elements in the employed sample allows matrices with specific features (Section 2.4), which can be important for desired applications.

An essential optical element to TPP technique is the microscope objective, which is used to focus the laser beam into the sample. Figure 3 shows a schematic diagram for different objectives [38].

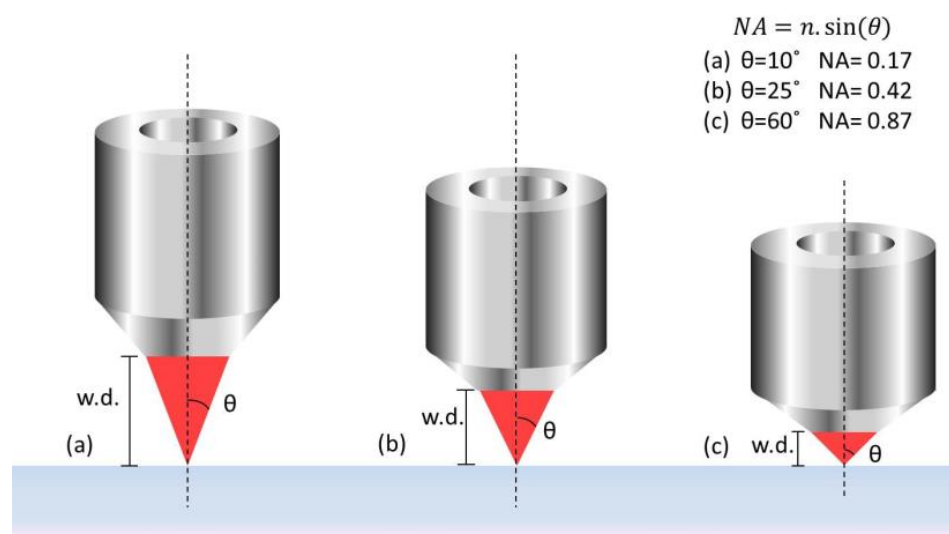


Figure 3. Light focusing through a microscope objective lens with different numerical aperture (NA) and working distance (w.d.). (a) NA = 0.17, (b) NA = 0.42 and (c) NA = 0.87. Reproduced with permission [38]. Licensed under a Creative Commons Attribution 4.0 International license.

The greater the numerical aperture, the smaller the working distance, i.e., the distance between the focal plane and the surface of the objective. The smallest spot size achieved is given by Abbe's expression [47]:

$$r = \frac{0.61\lambda}{NA} \quad (4)$$

in which, r is the focal spot radius, λ is the light wavelength, and NA is the numerical aperture of the microscope objective. The numerical aperture is defined by Abbe as:

$$NA = n \cdot \sin(\theta) \quad (5)$$

in which, n is the index of refraction in the focusing medium, and θ is the convergence angle of the beam.

However, most pulsed lasers have an approximately Gaussian beam profile. Besides that, two-photon polymerization setups use microscope objectives with high numerical aperture, and the minimum beam waist can be expressed by:

$$w_0 = \frac{\lambda}{\pi \cdot NA} \sqrt{n^2 - NA^2} \quad (6)$$

where w_0 is the beam waist at the focal plane. The beam waist is defined as the distance from the center of the Gaussian beam up to the point where the amplitude of the electric field falls to $1/e$ of its maximum.

The photopolymerized structure resolution can also be improved when the fabrication procedure is performed around the polymerization threshold. When the laser beam using energy below the polymerization threshold value irradiates the sample, no polymerization occurs (Figure 4). The more the peak power gets closer to the threshold power, the smaller is the voxel size.

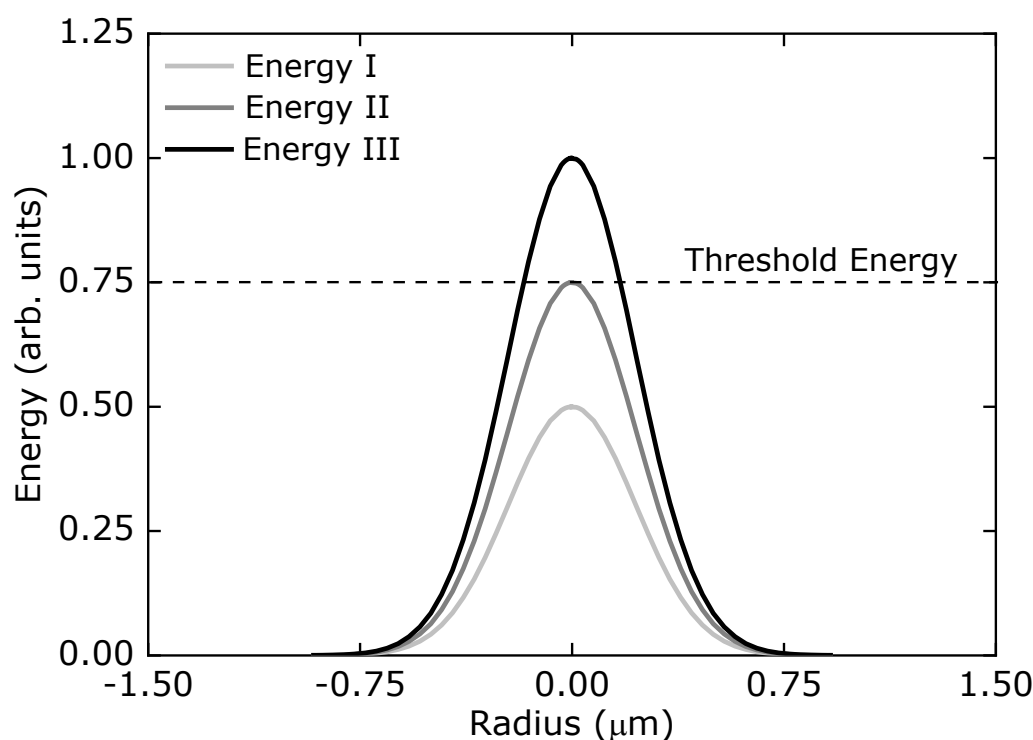


Figure 4. Threshold power relative to various Gaussian profiles. Polymerization is not observed if peak power is below the threshold power.

In the next section, we discuss the experimental features of TPP technique.

2.3. Experimental Aspects of TPP

The optical systems used for microfabrication via TPP vary in different aspects, including the laser system itself and the optical components. A schematic showing the basic components of a TPP system is presented in Figure 5. The laser beam comes from a mode-locked Ti:sapphire oscillator and is focused on the photoresist using an objective lens. The pulse energy can be adjusted by using a half-wave plate and a polarizer. A set of two movable mirrors is used to scan the laser beam transversally in the photoresist. A telescope is used to expand the laser beam to match the objective aperture. As previously mentioned, the choice of the objective lens is essential, as it determines the focal volume, allowing control of the size and thickness of the fabricated structures. A translational stage

is used for the sample positioning, and a mechanical shutter controls the laser exposure. Control programs are used to manage the shutter, movable mirrors, translation stage, exposure time, and laser scanning speed. The combination of movable mirrors with the 3D translational stage allows the fabrication of the microstructures over a large area in the order of millimeters. The TPP writing process is performed in a successive layer approach and can be monitored in real-time with a CCD camera and a backlight illumination, while the refractive index of the polymerized structure is slightly changed during the process. After the TPP fabrication, the sample is immersed in an appropriate solvent, to remove the unpolymerized material.

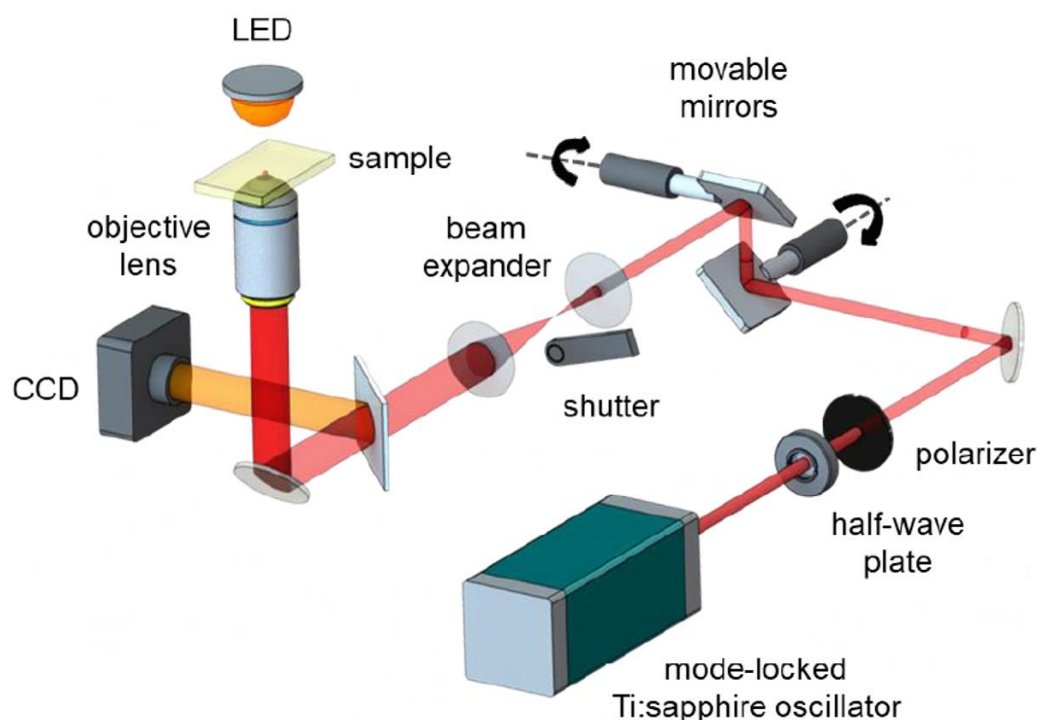


Figure 5. Representation of the femtosecond laser microfabrication via TPP setup, showing the basic components of the system. Reprinted with permission [40]. Licensed under a Creative Commons Attribution 4.0 International license.

2.4. Materials Used for TPP

In polymerization systems initiated by light absorption, there are two classes of photoresist materials that can be employed, and they are denominated as positive or negative photoresists. A great advantage of TPP technique is the maskless fabrication, due to the focalization of the laser beam into the sample. When exposed to light, positive photoresists, such as S1800 [48], become soluble and, in the non-irradiated region, the material solidifies. In the case of negative photoresists, such as SU-8 [49], the process is reversed, and the region that will be solidified is the one that is exposed to light.

A variety of materials can be used in the TPP experiments. The matrices used in this laser fabrication technique should be selected considering the application of interest. For instance, acrylate monomers are widely used in TPP [34,50]. These compounds usually present high optical quality and are biocompatible with different microorganisms [51,52]. Besides that, these resins can be easily functionalized with several materials, giving specific optical or biological features [53–56]. The combination of two acrylate monomers, for example, can improve the hardness and prevent the shrinking of the final structure [50]. Epoxy-based compounds, such as the SU-8, are also attractive for the TPP experiment, allowing the manufacture of structures with a high aspect ratio [57]. Hybrid materials also are noteworthy in TPP experiments [58,59]. These matrices combine the advantages of organic compounds (for instance, flexibility and chemical resistance) with those of inorganic

components, improving different properties of the final device (such as mechanical stability and optical quality in several wavelengths) [60].

Negative-tone resins produced by Nanoscribe GmbH & Co. (Eggenstein-Leopoldshafen, Germany) also are applied in the TPP setups [61–63]. These resins, called IP Photoresins, can be used to fabricate several technological devices, such as optical elements for the biological scaffolds. Other commercial matrices, such as Ormocer[®] and Ormocomp[®], have been used to sculpt devices for photonics, optoelectronics, and biological areas [48,64,65]. However, many groups have developed their own samples, directing it to the desired application. For instance, some researchers have expended efforts to develop photocrosslinkable gelatin hydrogels for biofabrication applications [66–70]. For tissue engineering applications, photoreactive resin based on poly(trimethylenecarbonate) (PTMC) has been modified through chemical reactions, aiming at the fabrication of highly porous and biodegradable scaffolds [71].

Still, regarding the employed samples in the TPP experiments, photoinitiator compounds are essential to the technique. Many groups have been working on the development or testing of new photoinitiators for TPP experiments [72,73]. Usually, they look for compounds with a high value of TPA cross-section (σ_{TPA}).

After absorbing the laser pulse energy, radicals are generated. Then, these radicals can react with other sample components (monomers) and start the photopolymerization procedure. There are two mechanisms through which the photoinitiator can trigger the polymerization reaction in the TPP: polymerization by free radicals generation and cationic polymerization [74].

The active species in the polymerization by free radicals are neutral, and their incorporation with the monomers is not influenced by the polarity of the monomer or the solvent. However, molecular oxygen can inhibit its action, which is often advantageous for better resolutions in manufactured devices [9]. In free radical polymerization, photoinitiators can be classified into two groups, according to the mechanism of radical formation. The photoinitiators classified as Norrish Type I, when absorbing one or two photons, are brought to an excited state and undergo breakdown (or chemical fission), producing radicals, in general, equally reactive. Among the Norrish Type I photoinitiators, the commercial compound named Lucirin TPO-L [75] is a typical material employed in TPP fabrication. The photoinitiators classified as Norrish Type II, for example, benzophenone [76], do not generate radicals independently, and the use of a co-initiator is necessary. Such photoinitiators undergo a bimolecular reaction with a co-initiator to create radicals. Thus, as the bond cleaves homolytically after excitation, two radicals are generated, and at least one of them will initiate the polymerization reaction [77].

In cationic photoinitiation systems, an acid is generated after laser irradiation. Unlike photopolymerization initiated by free radicals, sometimes cationic polymerization can occur without chain growth control. However, some photoinduced methods allow better control of the cationic polymerization [78]. Also, cationic photopolymerization is not inhibited by oxygen. Different classes of monomers, such as epoxy, vinyl esters, and siloxanes can be polymerized by a cationic mechanism.

The variety of materials that can be employed in the TPP experiments, added to the ease of sophisticated structure fabrication, makes this technique very attractive for developing the device. In the next section, we present some applications of the sculpted structures by means of TPP technique.

3. Devices Fabricated via TPP: General Applications

3.1. Functionalized Structures

One of the most attractive advantages of the TPP technique is related to the production of functionalized devices, which can be applied in several technological fields. Usually, these devices are designed with specific properties according to the application need. There are different methods able to realize the device's functionalization. The mixture of the functionalizing compound with the host sample (used in the TPP experiment) is the

simplest method to achieving devices with special properties. For optics and photonics applications, organic dyes can easily be added to the pure matrices only by mixing the desired elements [53,55,79]. Several works were performed to manufacture microstructures doped with fluorescent dyes, such as Rhodamine B, Disodium Fluorescein, and Stilbene 420 [53]. Specifically, the use of these three dyes allows the fabrication of RGB-like syntonized devices. Other optically active compounds of considerable interest in photonics are the azobenzene-based dyes, such as Disperse Orange 1 (DO1), Disperse Red 13 (DR13), or Sudan Black B (SBB). This kind of dye can also be used as a photoinitiator in TPP experiments, keeping their original optical properties even after all microfabrication procedures [80]. Figure 6 shows some 3D structures fabricated using azobenzene-based dyes as a photoinitiator.

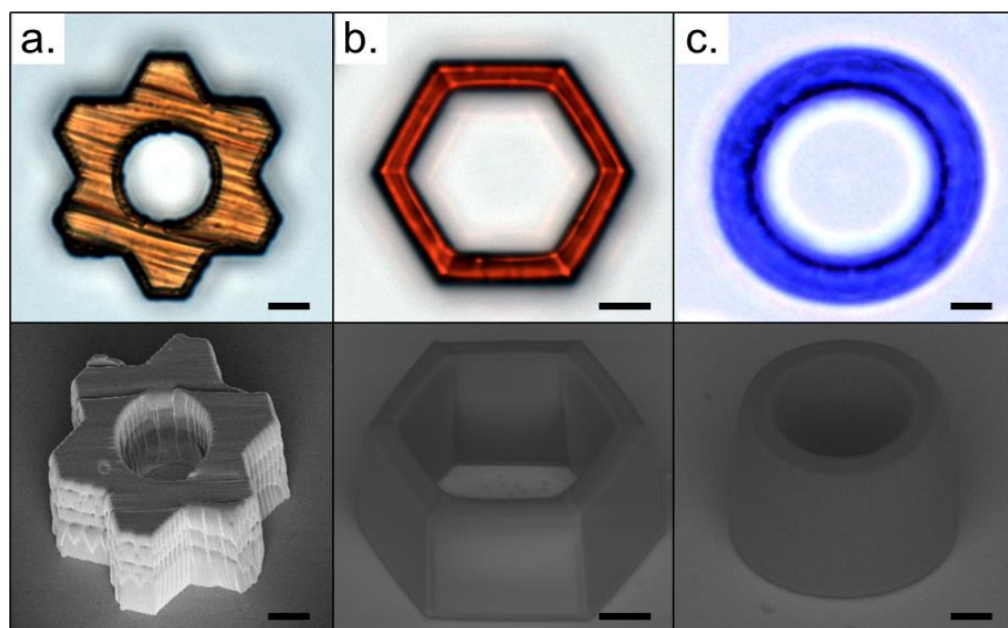


Figure 6. Optical microscopy (**top**) and SEM images (**bottom**) of 3D structures fabricated using azobenzene-based dyes as a photoinitiator in a concentration of 1.00 wt%: (a) Disperse Orange 3 (DO3), (b) Disperse Red 13 (DR13), and (c) Sudan Black B (SBB). The scale bar is 20 μm .

As can be seen in Figure 6, structures fabricated through the free radical generation in azoaromatic compounds present good structural integrity. The final structure color is similar to the pristine dye solved in ethanol (the solution used to incorporate the dye into the pure resin). Moreover, due to a high dye concentration used as a photoinitiator, a significant amount remains unchanged in the produced structures, and some optical properties, such as optically induced birefringence, can be monitored [80].

Polymeric materials sometimes have poor physical-chemical properties, which makes difficult their application in microdevices. An excellent alternative to solve this problem is, for example, the development of polymeric materials reinforced with carbon nanotubes, a material widely known to have interesting electrical, thermal, optical, and mechanical properties [81–84]. Even a low concentration of carbon nanotubes dispersed into the polymeric matrices (0.01 wt%) can significantly improve the device's properties [85].

Some groups have performed the fabrication of 2D and 3D structures using polymeric matrices reinforced with single-walled or multi-walled carbon nanotubes (SWCNT or MWCNT) [85–89]. Figure 7 shows some examples of different structures fabricated via TPP, using SWCNT/polymer composites.

Still, regarding microstructures fabricated using SWCNT/polymer composites, researchers have demonstrated that the carbon nanotubes are uniformly distributed into the sample [85], and also, they can stay aligned inside the photopolymerized structures,

according to the laser scanning direction [88,89]. Besides that, structures fabricated with MWCNT/polymer matrices, grown upon two pairs of Au electrodes, had different electrical conductivity responses according to the laser fabrication direction. For instance, structures fabricated with parallel scanning (in respect to the Au electrodes) were 1000 times more conductive than those fabricated with perpendicular scanning [89]. An increase in the electrical conductivity of over 11 orders of magnitude can be observed in acrylate-based devices functionalized with 0.2 wt% MWCNT [89].

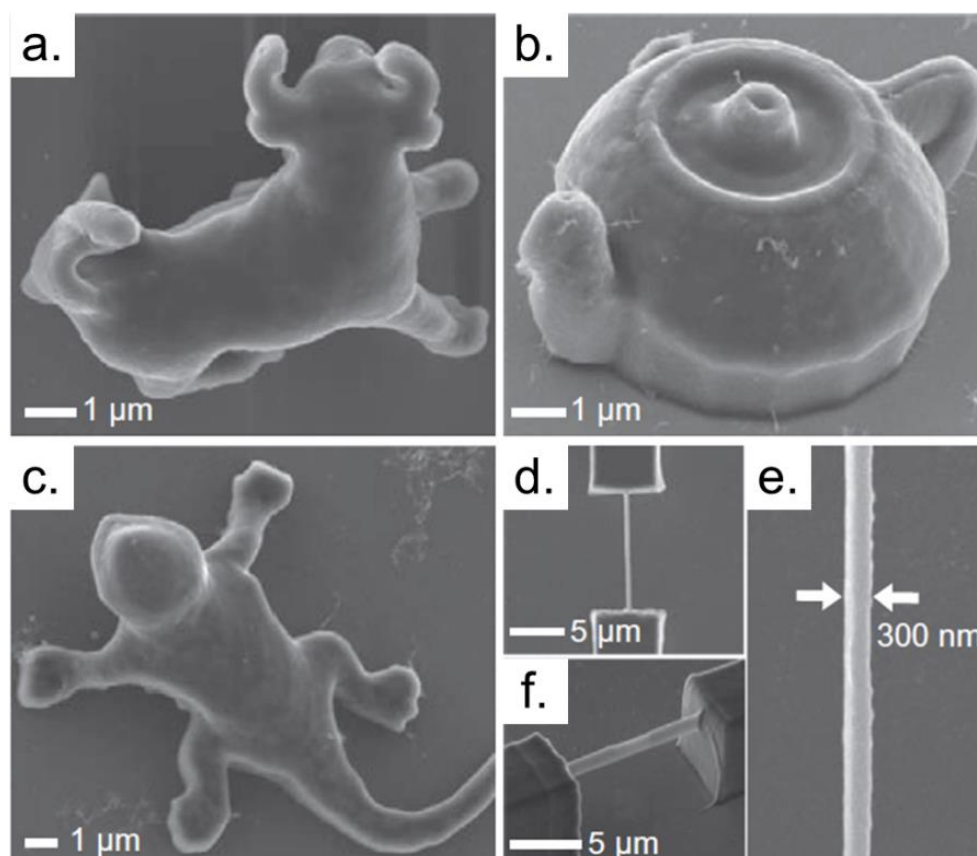


Figure 7. 3D micro/nano structural SWCNT/polymer composites are fabricated by using the TPP lithography. The structures in (a–f) are a 8-lm-long micro bull, a micro tea pod, a micro lizard, a nanowire suspended between two micro boxes, magnified image of (d), and a perspective view of the nanowire, respectively. Reproduced with permission [87]. Copyright © 2013, Elsevier.

Mechanical properties are also improved after CNT incorporation in acrylate resins. For structures reinforced with 1 wt% of SWCNT, the change of the modulus of elasticity and viscoelasticity is notable [85]. If the MWCNT concentration increases in the acrylate material, the shrinkage of the photopolymerized structures decreases [89].

The device functionalization also can be performed after the complete laser fabrication procedure [65,90]. Figure 8 shows 3D metallic structures fabricated in a large area by TPP, using for this purpose a microlenses array that allows the fabrication of hundreds of devices simultaneously [90]. The structures' metallization was realized by depositing thin films of small silver particles via electroless plating. Chemical changes of the resin and the hydrophobic coating of the glass substrate avoid metal deposition and confine the adhesion on the polymer [90].

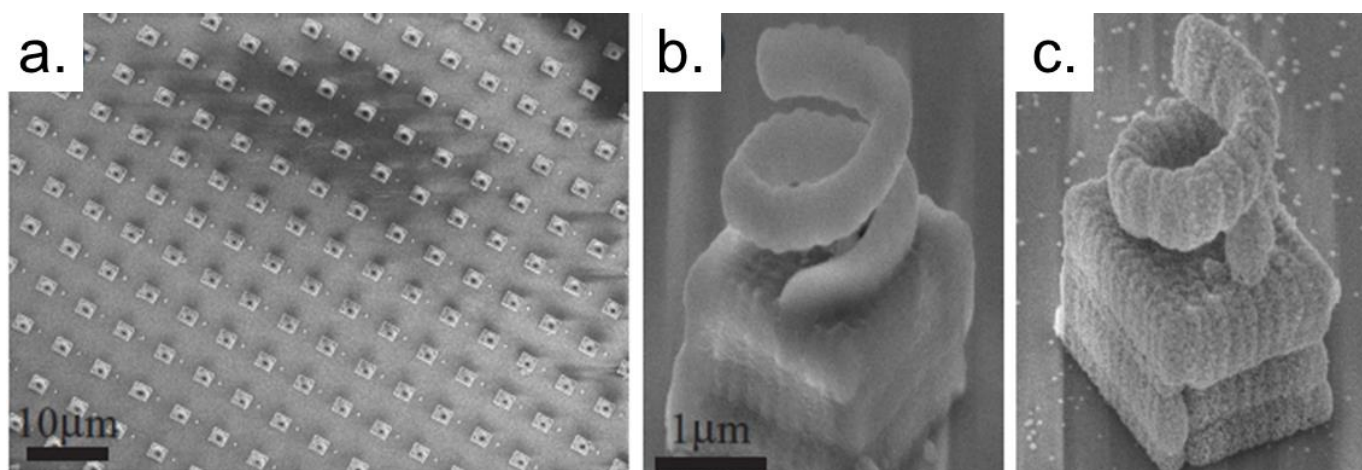


Figure 8. (a) $78 \times 58 \mu\text{m}^2$ SEM image of a 3D periodic silver-coated structure fabricated on a hydrophobic coated glass surface. (b) Tilted magnified view of an individual uncoated polymer structure composed of a cube ($2 \mu\text{m}$ in size) holding up a spring (height $2.2 \mu\text{m}$, inner diameter $1 \mu\text{m}$). (c) SEM image of an individual silver coated structure after electroless plating. Reproduced with permission [90]. Copyright © 2006, The Optical Society.

The use of TPP combined with other laser fabrication techniques, such as Laser-Induced Forward Transfer (LIFT), allows selective structure functionalization or even selective microorganisms inoculation [91]. Figure 9 shows a microstructure fabricated via TPP and selectively coated with a conducting polymer (PPV). It is important to highlight that the transferred polymer keeps its original features, even after the laser irradiation on the donor material. Besides, a material can be LIFTed several times upon the photopolymerized device [92], expanding its use to functionalize microstructures.

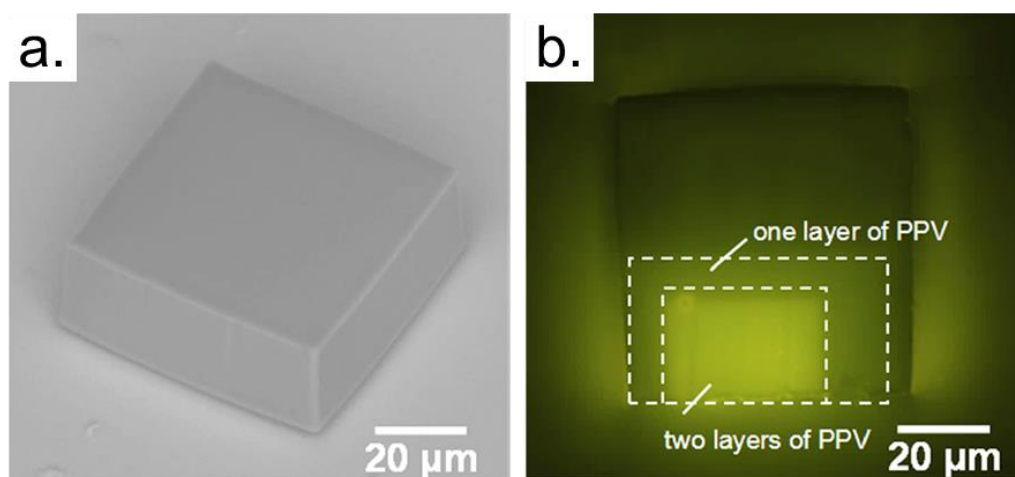


Figure 9. (a) SEM image of a polymeric microstructure fabricated by fs-laser writing via 2PP. (b) Fluorescence microscopy image of the microstructure shown after it was selectively coated with PPV via fs-LIFT. This image was obtained with excitation at 450–490 nm. Reproduced with permission [92]. Copyright © 2021, Springer Nature.

Structures that can be manipulable are very attractive to various fields, mainly for microfluidics or biomedical research [44,93,94]. The structure's movement can be achieved through the use of optical tweezers or magnetic manipulation. An example of manipulable structures is presented in Figure 10.

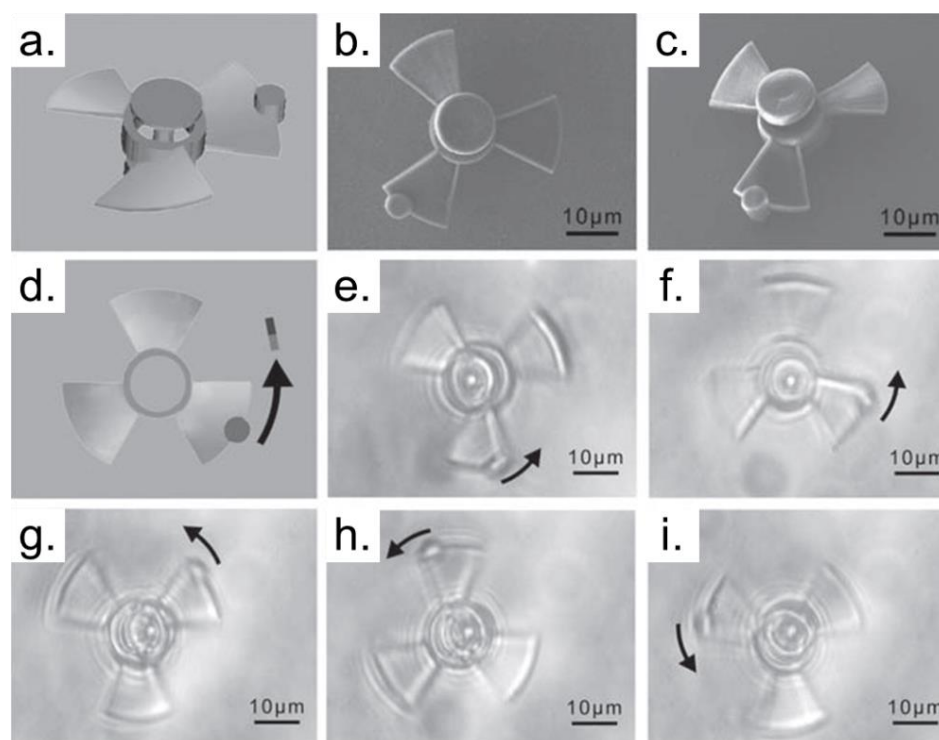


Figure 10. Remote control of the micro-turbine in acetone. (a) Model of the micro-turbine. (b) and (c) SEM images of the micro-turbine. (d) Top view scheme model for circumgyration. (e–i) Optical microscopy images of the microturbine in a circumgyration cycle. For remote control and observation of the micro-turbine, a piece of ferromagnet was placed on a vortical device around the objective lens. Reproduced with permission [93]. Copyright © 2010, John Wiley and Sons.

In Figure 10, a photopolymerizable ferrofluid was developed, and 3D micro-turbines were fabricated via TPP. The micro-turbine size is approximately 35 μm in diameter with a central axletree and three blades [93]. These kinds of devices can be moved and controllable aided by an external magnet. Moreover, these devices can turn clockwise and anticlockwise. Based on these results, the importance of manipulable devices is evident, and is possible thanks to the TPP process.

Functionalized devices are essential for several applications. One of the most significant areas is the whispering gallery mode microresonators, which we discuss in the next section.

3.2. Whispering Gallery Mode Microresonators

Optical microresonators are microstructures that strongly confine light, both in the time and space domains [95]. They have been widely exploited for basic studies on the light–matter interaction [96,97], and for numerous applications, ranging from lasers to sensors [98–100]. In particular, optical microresonators supporting whispering gallery modes (WGM) are noteworthy due to their unique features, such as high temporal confinement of light, high sensitivity to the environment, and their ease integration in photonic systems [95]. A light beam can travel over 10^6 roundtrips inside a WGM microresonator, which significantly enhances light–matter interactions and makes it possible to achieve a plethora of scientific discoveries and technological breakthroughs [101–108].

WGMs are electromagnetic waves that propagate along the rim of circular structures, e.g., cylinders and spheres, by total internal reflection (TIR) [109]. In a geometric optics framework, the WGMs can be seen as light rays that perform a close trajectory after a round trip in the circular-shaped structure, as shown in Figure 11d. The theoretical analysis of the WGMs in a given geometry is obtained by solving Maxwell’s equations with the proper boundary conditions [40,110]. To provide an insight on the theoretical aspects of WGM

microresonators, we will take dielectric microstructures with cylindrical symmetry as an example. For this particular case, the WGMs are electromagnetic fields that propagate in the azimuthal direction and have a zero axial propagation factor [111].

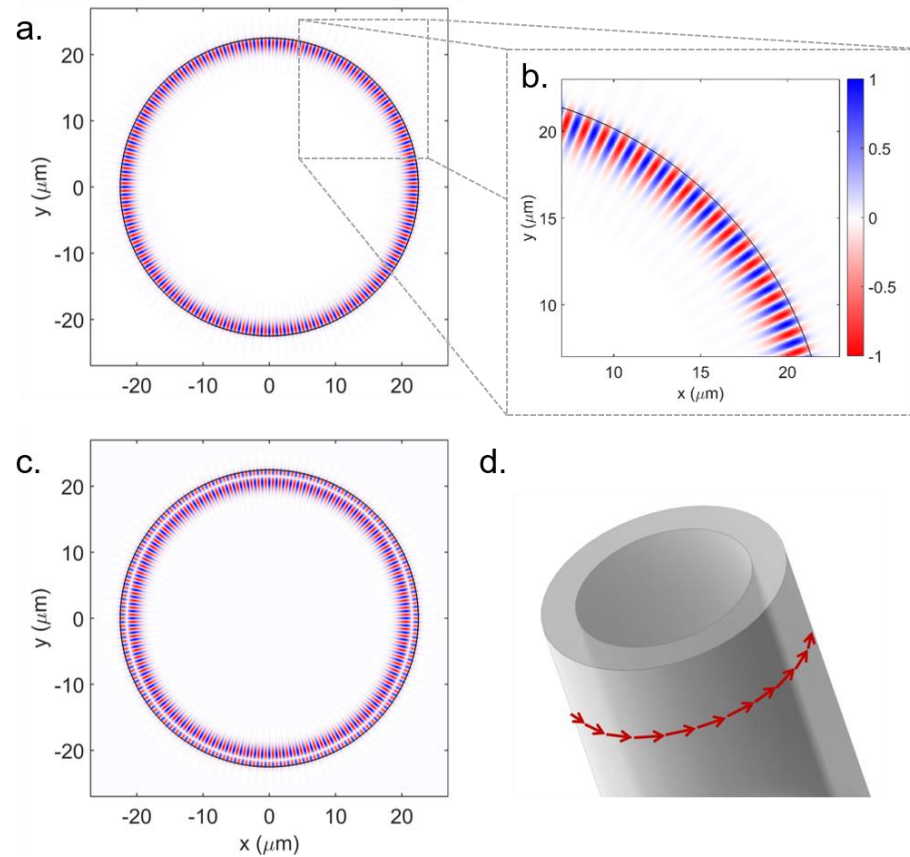


Figure 11. E_z field amplitude over a transverse plane of the cylinder for the azimuthal order $m = 120$ and the following radial orders: (a,b) $l = 1$ ($\lambda_{120,1} = 1668.2$ nm) and (c) $l = 2$ ($\lambda_{120,2} = 1579$ nm). Item (b) shows a zoomed-in view of the field distribution displayed in (a), along with the color bar for all the field distributions. The circumference corresponds to the cylinder's edge. To obtain these field profiles, the refractive indices of both media and microcylinder's radius were set to $n_1 = 1.51$, $n_2 = 1.0$ and $a = 22.5$ μm in Equation (9). (d) Geometric Optics picture of the WGMs propagating along the rim of a microcylinder. Adapted with permission [40]. Licensed under a Creative Commons Attribution 4.0 International license.

The resonance takes place when the fields interfere constructively after a roundtrip in the structure, i.e., [40]:

$$\vec{E}(\rho, \phi) = \vec{E}_0(\rho)e^{i\beta\phi} = \vec{E}_{\text{after a roundtrip}}(\rho, \phi) = \vec{E}_0(\rho)e^{i\beta(\phi+2\pi)}. \quad (7)$$

$$\therefore \beta = m; \quad m = 1, 2, 3, \dots$$

where (ρ, ϕ, z) are the radial, azimuthal, and longitudinal coordinates; $\vec{E}_0$ is the electric field amplitude; and β is the propagation constant in the azimuthal direction. The condition given by Equation (7) takes into account the selectivity of the fields allowed in the microstructure. The fields for which β is an integer constitute the modes of the structure.

However, there is a spectrum of modes for each polarization of the electromagnetic field, which are expressed as $\text{TM}^{(z)}$ modes when the magnetic field in the longitudinal direction is null ($H_z = 0$), and $\text{TE}^{(z)}$ modes when the electric field in the longitudinal

direction is null ($E_z = 0$) [40]. For a microcylinder, the longitudinal component of the electric field for a $TM^{(z)}$ mode is given by [110]:

$$E_z(\rho, \phi, t) = \begin{cases} a_m J_m(k_1 \rho) e^{i(m\phi - \omega t)}, & \rho < a. \\ b_m H_m^{(2)}(k_2 \rho) e^{i(m\phi - \omega t)}, & \rho > a. \end{cases} \quad (8)$$

for which: $k_i = n_i \frac{2\pi}{\lambda_0}$, $i = 1, 2$.

In the Equation (8), a is the microcylinder's radius, $J_m(k_1 \rho)$ and $H_m^{(2)}(k_2 \rho)$ are Bessel function of first kind and Hankel function of second kind, respectively, m is the azimuthal order, a_m and b_m are the field amplitude coefficients, n_i is the refractive index of each medium, and λ_0 is the wavelength of light.

The boundary conditions of the tangential fields at the dielectric interface are applied to obtain the characteristic equation for the $TM^{(z)}$ mode [110]:

$$\frac{1}{n_1} \frac{J_m(k_1 a)}{J'_m(k_1 a)} = \frac{1}{n_2} \frac{H_m^{(2)}(k_2 a)}{H_m^{(2)'}(k_2 a)}; \quad k_i = n_i \frac{2\pi}{\lambda_0} \quad (i = 1, 2). \quad (9)$$

For each azimuthal order (m), Equation (9) gives a set of complex wave numbers that constitute the $TM^{(z)}$ modes of the cylinder. Their real part gives the resonant wave numbers, whereas their imaginary part accounts for radiation losses resulting from the cylinder's curvature [40,110,111].

The total internal reflection imposes an additional boundary condition that constrains the allowed modes to the interval [110]:

$$k_2 a < m < k_1 a, \text{ in which } k_i = n_i \frac{2\pi}{\lambda_0}. \quad (10)$$

Thus, the WGMs are the roots of the characteristic equation (Equation (9)) that meet the TIR constraint.

As an example, in Figure 11a–c it is shown the E_z distributions over the transverse plane of a microcylinder, obtained for the WGMs of azimuthal order $m = 120$ and different radial orders ($l = 1, 2$). The azimuthal order corresponds to the number of field amplitude maxima along the azimuthal direction, while the radial order corresponds to the number of field amplitude maxima along the radial direction. As shown in Figure 11a, the field distribution of fundamental radial modes is more confined to the microcylinder's surface. Albeit, higher-order radial modes (Figure 11c) exhibit field distributions spread out inside the resonator, making them more susceptible to losses and rendering lower peak-intensity. The electromagnetic field confined in the microcylinder exhibits an exponentially decaying tail that extends to the surrounding medium, with a decay length of a few hundred nanometers. This characteristic allows light coupling in and out of the structure, being exploited, for instance, in the development of sensors.

For fundamental radial WGM modes ($m \gg 1, l = 1$), the resonance condition can be described by [110]:

$$\lambda_0 = \frac{2\pi}{m} n a \quad (11)$$

where λ_0 is the resonance wavelength, m is the azimuthal order, and n and a are the refractive index and the radius of the microresonator, respectively. Note that, since the fundamental radial WGM modes are strongly confined to the microcylinder's surface, their effective index can be approximated to the microresonator's refractive index ($n_{eff} \sim n$). Equation (11) is helpful to estimate the spectral position of the WGM resonances without solving the characteristic equation. From Equation (11), the free spectral range (FSR),

which is the frequency/wavelength spacing between consecutive azimuthal modes, can be estimated by [98,112]:

$$FSR = \frac{\lambda_0^2}{2 \pi a n}, \quad (12)$$

One of the most important parameters used to measure the time confinement of microresonators is the quality factor (Q), which is given by the ratio between the total energy stored and the power dissipated in the resonator in one cycle of the field [112,113]:

$$Q = \omega_0 \frac{\text{stored energy}}{\text{dissipated power}} = \frac{\omega_0 \tau}{2} = \frac{\nu_0}{\Delta \nu}, \quad (13)$$

where ω_0 is the angular resonance frequency, τ is the time for the stored energy to decay by a factor of $1/e$, and ν_0 and $\Delta \nu$ are, respectively, the resonance frequency and its corresponding linewidth.

The energy inside the microresonator dissipates through many routes, being intrinsic or external to the resonator. The overall losses can be written as [114]:

$$Q^{-1} = Q_{in}^{-1} + Q_{ext}^{-1} = (Q_{rad}^{-1} + Q_{mat}^{-1} + Q_{surf}^{-1}) + Q_{ext}^{-1}, \quad (14)$$

in which, Q_{in}^{-1} (Q_{ext}^{-1}) are the intrinsic (extrinsic/coupling) losses, and Q_{rad}^{-1} , Q_{mat}^{-1} and Q_{surf}^{-1} are the major contributions to the intrinsic losses due to radiation, material, and surface scattering losses, respectively. Moreover, power dissipation in the microresonator can also occur due to the presence of surface contaminants (e.g., water adsorption) [115,116], and through nonlinear processes, such as multiphoton absorption [117], prompted by the high intensities achieved in WGM microresonators. The overall Q-factor is limited by the dominant source of loss.

WGM microresonators with high Q are able to confine light for a relatively long time, such that high circulating intensities are achieved even from moderate excitation levels, because of the extremely small volume of WGMs in these structures [99]. WGM microresonators with Q in the range of 10^3 – 10^6 are called high-Q, and those surpassing 10^7 are named as ultra-high-Q [96].

The materials used for fabricating WGM microresonators are diverse and depend on the application and fabrication technique. In particular, polymeric microresonators have become increasingly competitive with microresonators made of other materials due to their structural flexibility, ease of processing and functionalization, and low cost [99]. Moreover, there are polymers with a wide range of refractive indices and with electro-optic properties available [118].

Various techniques have been used to fabricate WGM polymeric microresonators, including direct lithography patterning [118,119], molding from a master structure [120,121], surface-tension-based approach [122], solution-printing technique [123], and femtosecond laser writing via two-photon polymerization (TPP) [17,51]. In the past two decades, the latter technique has attracted significant interest due to its 3D fabrication capability, fine-tuning of the structure dimensions, and the integration of the fabricated structures into a range of substrates [2,9].

Polymeric microresonators with several geometries have been fabricated by TPP, including toroids [124], disks [125–127], cylinders [17,128], rings [26,51,129], and spheres [130]. Moreover, the flexibility of geometry afforded by this technique allows the fabrication of suspended parts (Figure 12a,c), varying the height of structures on the same chip (Figure 12b), and wire bonding fiber-to-chip coupling elements (Figure 12d and Ref. [18]), which opens up new degrees of freedom for microresonator designing. For example, the microresonators displayed in Figure 12b, which were incorporated with a photoresponsive deformable material, are vertically coupled to integrated waveguides [131]. Owing to the 3D integration scheme, the microresonator spectral response can be dynamically tuned not only through its refractive index variation but also through its mechanical deformation by a

remote optical stimulus. Another example is the integrated microresonator displayed in Figure 12d, which is coupled to a fiber coupler through a photonic bonding wire [129]. In addition to compactness, this architecture avoids alignment difficulties and reduces the losses related to fiber-to-chip coupling [18,29].

The integration of the microresonators into different substrates represents a great advantage of TPP over the other techniques. Figure 12c,e,f respectively shows microresonators fabricated inside a microfluidic channel, onto a silicone-based substrate, and onto the polished facet of a multicore optical fiber. The interface between microresonator systems and microfluidic channels/optical fibers gives rise to portable and easy-to-handle sensing units, which are promising tools for medical care, chemical manufacturing, and environmental monitoring [26,132]. Besides, the integration of microresonators into multicore fibers allows light coupling into and out through the different fiber cores, thus making the microresonator interface with laboratory instrumentation very convenient. The use of silicone as a substrate can enable various photonic applications by allowing reversible and precise tuning of the gap distance between pairs of microresonators through mechanical stretching [133,134]. It has been shown that this method affords a tuning precision comparable to a nano-positioning piezo stage but with the significant advantage that the microresonators are on the same substrate [133].

Another convenience of the TPP technique for the fabrication of microresonators is its material flexibility. Several polymeric formulations have been employed as photoresists, such as acrylic [51,135] and epoxy resins [130,132], and various materials have been incorporated into them to meet specific applications. Polymeric microresonators fabricated by TPP have been incorporated with organic dyes to realize optically active microdevices [125,130,134], with liquid crystal networks to confer elasticity for optical tunability [131,136], and with nanodiamonds to enable quantum photonics applications [135], among other materials [137].

WGM microresonators fabricated by TPP can achieve Q-factors, in the near-infrared, from 10^2 until 10^6) [124,125,138–140]. Since the WGMs are strongly confined to the microresonator's edge, the Q-factor in such structures is primarily limited by light scattering at the microresonator surface. Therefore, the major challenge behind accomplishing a high-Q microresonator by TPP lies in ensuring optically smooth surfaces. As described in Section 2, the laser writing process gives rise to polymerized volumes (voxels), which overlap to create the geometrical shapes that constitute the microstructure. In order to produce microstructures with smooth surfaces, the exposure and laser scanning parameters must be adjusted to ensure that the voxels are properly overlapped, i.e., the surface roughness associated with the individual characteristics of the voxels is significantly reduced.

Figure 13a shows one of the highest Q-factor WGM microresonators fabricated by TPP thus far [17]. It is a 22.5- μm -radius hollow microcylinder fabricated from an acrylate polymer using a 0.25 NA objective lens, 0.1 nJ pulses from a Ti:sapphire oscillator, and 40 $\mu\text{m}/\text{s}$ of laser scan speed. Such parameters resulted in a longitudinal and transversal voxel overlap of 85% and 90%, respectively, translating into an average surface roughness as small as 2 nm. A morphology chart of the microresonator sidewall surface is shown in Figure 13b. The microresonator was interrogated by evanescent coupling through a 2 μm -waist tapered optical fiber, as depicted in Figure 13c. Its broadband spectral response around 1550 nm (Figure 13d) contains resonances accounting for different polarization states and different azimuthal and radial order WGMs. The TE and TM fundamental radial order WGMs and their corresponding FSR are highlighted in Figure 13d. The microresonator loaded Q-factor was found to be 1×10^5 , which is very close to the fundamental limit imposed by the acrylate material losses ($Q_{\text{mat}} = 1.9 \times 10^5$). Such Q-factor is competitive with the current semiconductor platforms and holds great prospects for several applications in photonics.

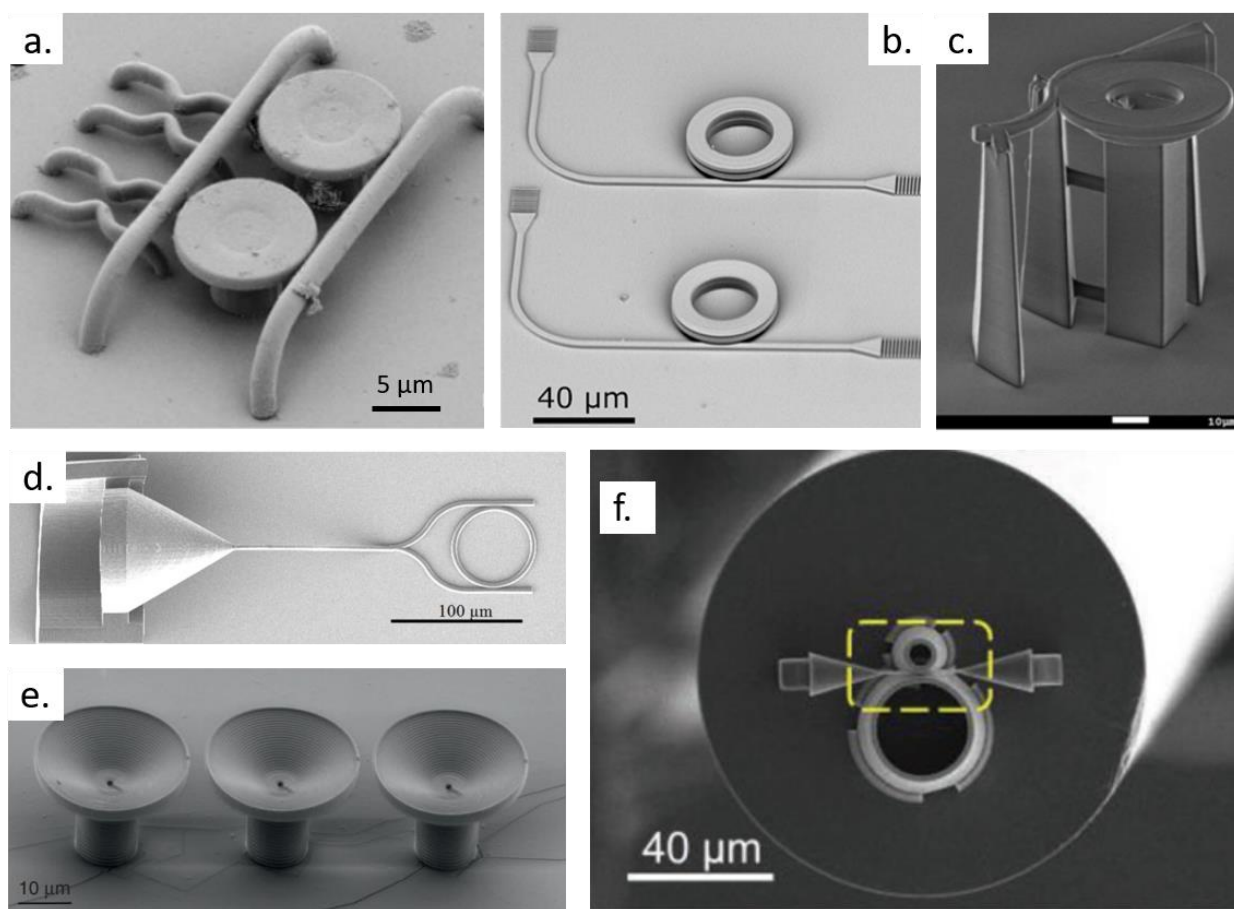


Figure 12. Polymeric microresonators fabricated by femtosecond laser writing via TPP. (a) microdisk resonators fabricated from a nanodiamond photoresist to enable quantum photonics applications, (b) two vertically-coupled microring resonators incorporated with liquid crystal networks and an azo dye to afford light-driven tunability, (c) microring resonator inside a glass microfluidic chip for biosensing, and (d) integrated microring resonator used for ultrasound detection. (e) Goblet-shaped microresonators fabricated on a PDMS elastomer substrate for coupling tunability, and (f) microring resonators fabricated on the facet of a multicore optical fiber for vapor sensing. Figure 12a Reproduced with permission [135]. Licensed under a Creative Commons Attribution 4.0 International license. (b) Reproduced with permission [131]. Copyright © 2018, American Chemical Society. (c) Reproduced with permission [132]. Copyright © 2019, The Royal Society of Chemistry. (d) Reproduced with permission [129]. Copyright © 2017, The Optical Society. (e) Reproduced with permission [133]. Licensed under a Creative Commons Attribution 4.0 International license. (f) Reproduced with permission [26]. Copyright © 2019, John Wiley and Sons.

The microresonators with cylindrical geometry do not support light confinement in the vertical direction, which brings alignment difficulties to the coupling and leads to a more spread out mode volume. However, it can be challenging to accomplish more confined geometries, such as spheres, toroids, or even disks and integrated microrings, with smooth surfaces by TPP. Indeed, the Q-factor of microresonators with spherical [130] and toroid-like [124] geometries is in the order of 10^4 or lower, and the Q-factor of integrated microrings is in the range $10^2 - 10^3$ [51,129,138,141]. Microdisks and integrated microrings suffer from additional scattering losses at their top surface, which is usually rougher than their sidewall surface due to the pointed shape of the voxel tips. One of the strategies used to reduce light scattering in a microdisk consists of designing its edge in the shape of a wedge and engineering the wedge angle to minimize the mode overlap with the top and sidewall surfaces [139].

One of the most studied applications of microresonators fabricated by TPP is lasing. The unique combination of Q-factor and tight mode confinement provided by WGM mi-

microresonators enables strong interaction between the light and the gain medium, which makes it possible to achieve low threshold and narrow linewidth microlasers [99]. Moreover, its small size and the possibility of on-chip integration make them promising as bright sources for integrated photonics applications.

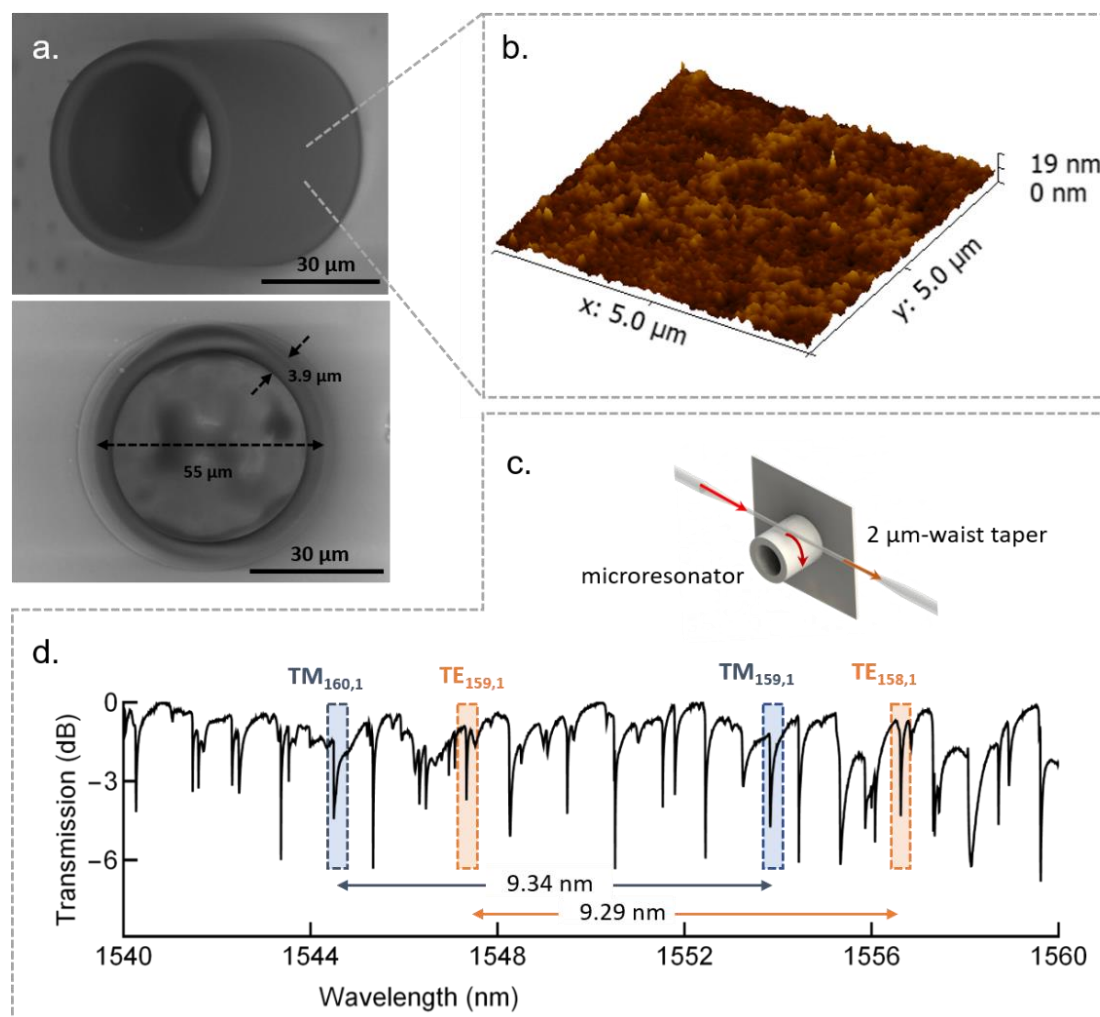


Figure 13. Polymeric microcylinder resonator fabricated by TPP. (a) Scanning electron micrographs of the microresonator. (b) Atomic force microscopy of the microresonator outer sidewall surface. (c) Schematics of the evanescent light coupling to the microresonators through the use of tapered fibers. (d) Spectral response of the microresonator. The attenuation peaks represent the WGMs accounting for different azimuthal and radial orders. Highlighted in the figure are the TE and TM fundamental radial WGMs, with their corresponding FSR. Adapted with permission [17]. Copyright © 2017, John Wiley and Sons.

To realize microlasers, the microresonators are incorporated with organic dyes, such as Rhodamine B (RhB) [125] and Pyrromethene 597 [134]. The microresonators displayed in Figure 14a were fabricated from an acrylic-based photoresist incorporated with RhB [31]. Their hollow cylindrical geometry was strategically chosen since it supports a smaller number of high-order radial WGMs that would otherwise compete with the most fundamental WGMs for the pump energy. A 3D reconstructed confocal fluorescence micrograph confirmed the homogeneity of the dye distribution throughout the microresonator (Figure 14b), and absorbance/fluorescence analyses showed the neutrality of the polymeric matrix, i.e., it was shown not to affect the emission properties of the dye significantly.

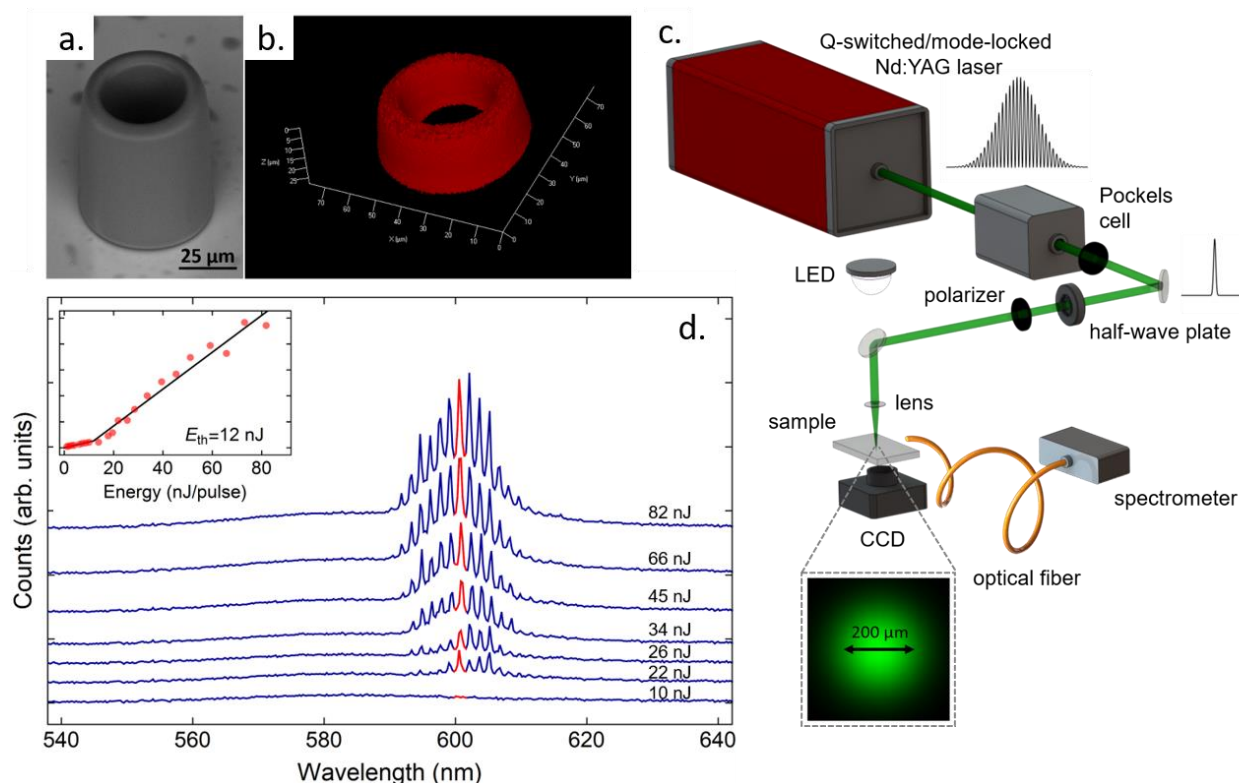


Figure 14. Rhodamine B-doped WGM polymeric microlaser fabricated by TPP. (a) Scanning electron, and (b) 3D reconstructed confocal fluorescence micrographs of the microlaser. (c) Setup used to excite the microlasers and collect their emission. (d) Emission spectra of a microlaser for the excitation energy levels indicated on the right hand side of the graph. To make the visualization easier, an offset was applied to the y-axis of each curve. The first curve represents the spectrum below lasing threshold. The inset shows a graph of the amplitude of the peak highlighted in red as a function of excitation energy. The graph is plotted on a linear scale. The threshold energy, which was obtained by fitting the experimental data with a bilinear curve (black line), is indicated in the graph. Adapted with permission [31]. Licensed under a Creative Commons Attribution 4.0 International license.

The setup used to pump and collect the emission of the RhB-doped microlasers is shown in Figure 14c. In this setup, 100 ps-pulses from a Q-switched and mode-locked Nd:YAG laser centered at 532 nm are loosely focused on the microresonator top surface, and the microresonator emission is collected with a multimode optical fiber connected to a 250-pm-resolution spectrometer. Most of the other polymeric microlasers are also pumped with pulsed lasers with a time duration in the order of 10 – 100 ps [125,127,142]. The pulsed excitation prevents population transfer to the triplet state of the dye molecules, which would hinder its fluorescence quantum yield [143]. Overall, since the fluorescence lifetime of organic dyes is in the order of nanoseconds [144], the excitation being performed in the ps regime leads to less molecular cycles, thus extending the device lifetime.

The emission spectra of the RhB-doped microlasers feature a set of well-defined peaks within a narrow spectral range around 600 nm. The lasing threshold was measured at 12 nJ of pulse energy, which is well within the energy levels afforded by conventional ps-lasers. Besides, such a low lasing threshold is comparable to that which has been achieved for polymeric microlasers fabricated by other techniques [123,145,146], thus reinforcing the potential of TPP in fabricating active microresonators.

Some strategies have been developed to improve directionality, monochromaticity, and tunability for WGM microlasers fabricated by TPP. One of the strategies that has been used to overcome the lack of directionality of such microlasers consists in breaking their rotational symmetry. This can be accomplished by designing a spiral-shaped microdisk, which has been shown to exhibit directional lasing emission with a far-field divergence of

about $20^\circ - 40^\circ$ [142,147]. To ensure single-mode operation without significantly degrading the microlaser Q-factor, coupled WGM microlasers have been employed. It has been demonstrated that the coupling of two microdisks with different diameters suppresses some resonances and increases the effective FSR, which allows accommodating only one mode within the gain region [142,148]. Laser tunability has been achieved by using elastomer-based materials either as the substrate [134] or integrated into the microlasers [136]. For example, it has been reported that the lasing modes of a WGM microlaser integrated with photo-responsive liquid crystal elastomers reversibly reach a 0.51 nm- shift for every increase in the pump energy by 0.14 mJ/pulse.

Another important application of WGM microresonators fabricated by TPP is in sensor technology. The operational principle of WGM sensors is to monitor changes in their spectral properties prompted by environmental changes. Sensing of refractive index [128], temperature [141], volatile organic compounds [26], ultrasound waves [129], and the label-free detection of biological specimens [132] have been accomplished with these structures. The sensing mechanism usually relies on resonance wavelength shifts and/or resonance linewidth variations. An example of WGM sensor fabricated by TPP is shown in Figure 15. The multicore fiber containing the WGM polymer microresonator (Figure 15a) is placed in a vapor environment produced by an aqueous solution of propylene glycol monomethyl ether acetate (PGMEA) [26]. As shown in Figure 15b, by increasing the concentration of PGMEA, the WGMs are shifted towards longer wavelengths in response to microresonator swelling and refractive index variation of its surrounded medium. Besides PGMEA, the WGM sensor showed in Figure 15a was used to detect isopropanol and alcohol vapors, exhibiting a sensitivity of 21.7, 3.38, and 3.87 pm/ppm for each of these vapors in the low concentration range [26].

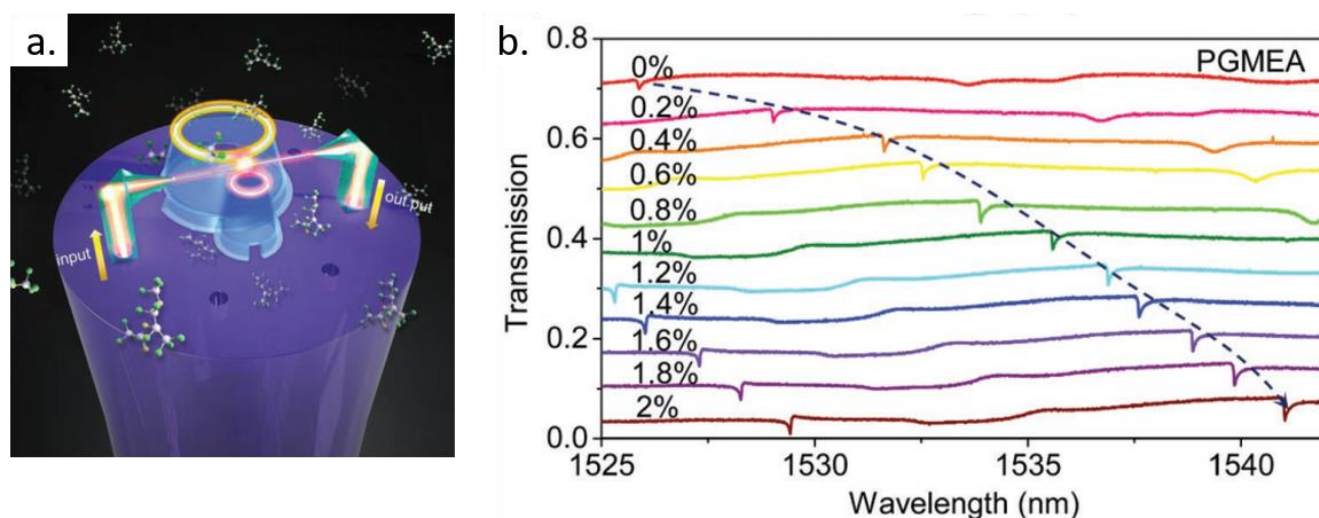


Figure 15. Microring resonators fabricated by TPP on the facet of a multicore optical fiber for vapor sensing. (a) Illustration of the light propagation in the photonic setup. (b) Spectral response of the microresonator when the fiber tip is placed in the vapor environment produced by an aqueous solution of propylene glycol monomethyl ether acetate (PGMEA) with the concentration varying from 0% to 2%, in steps of 0.2%. Adapted with permission [26]. Copyright © 2019, John Wiley and Sons.

In the context of WGM sensors, it is desirable to have a clean resonance spectrum to prevent distortions associated with the overlap of multiple resonance peaks. One proposed solution to achieve mode cleaning is by incorporating a lossy element in a low concentration into the microresonator. It was shown that the incorporation of graphene oxide (GO) to WGM microresonators led to the suppression of a number of resonances in the spectrum without significantly reducing the Q-factor of the most prominent resonances [137]. Another route to mode cleaning is by decreasing the microresonator diameter to a few

microns [149]. In this case, radiation losses would be considerably raised for all the modes, but the higher-order radial WGMs would be the most affected and ultimately the ones to be filtered out. The most loss-free strategy towards mode cleaning consists of reducing the sidewall thickness of hollow microcylinders or microrings to dimensions comparable to light wavelengths [121,149]. Such thin sidewall microresonators would only support the fundamental radial WGMs. However, it can be challenging to achieve, by the TPP technique, microresonators with such a small sidewall thickness that still maintain their structural integrity.

3.3. Three-Dimensional Scaffolds for Biological Investigations

At present, a significant part of the TPP sculpted devices is dedicated to biomedical applications due to the free form fabrication achieved by this technique, combined with ease functionalization of the microenvironment. The design and fabrication of biological scaffolds can be optimized to obtain platforms with biomimetic shapes and responses. Several researchers have investigated the geometry (or porosity) influence of photopolymerized scaffolds in the microorganisms' development [21,150,151]. Besides, the mechanical and chemical resistance are also important features that should be characterized before applying these scaffolds in biological studies. Usually, to optimize the scaffold production, many parameters have been defined, and computational simulation is required. Figure 16 shows an example of three types of human cells profiles, red blood cell—Figure 16a–d, smooth muscle cell (SMC)—Figure 16e–h, and ciliated columnar epithelial cell (CEC)—Figure 16h–l, which were produced by TPP technique, with a negative-tone commercial acrylic photoresist from Nanoscribe [152].

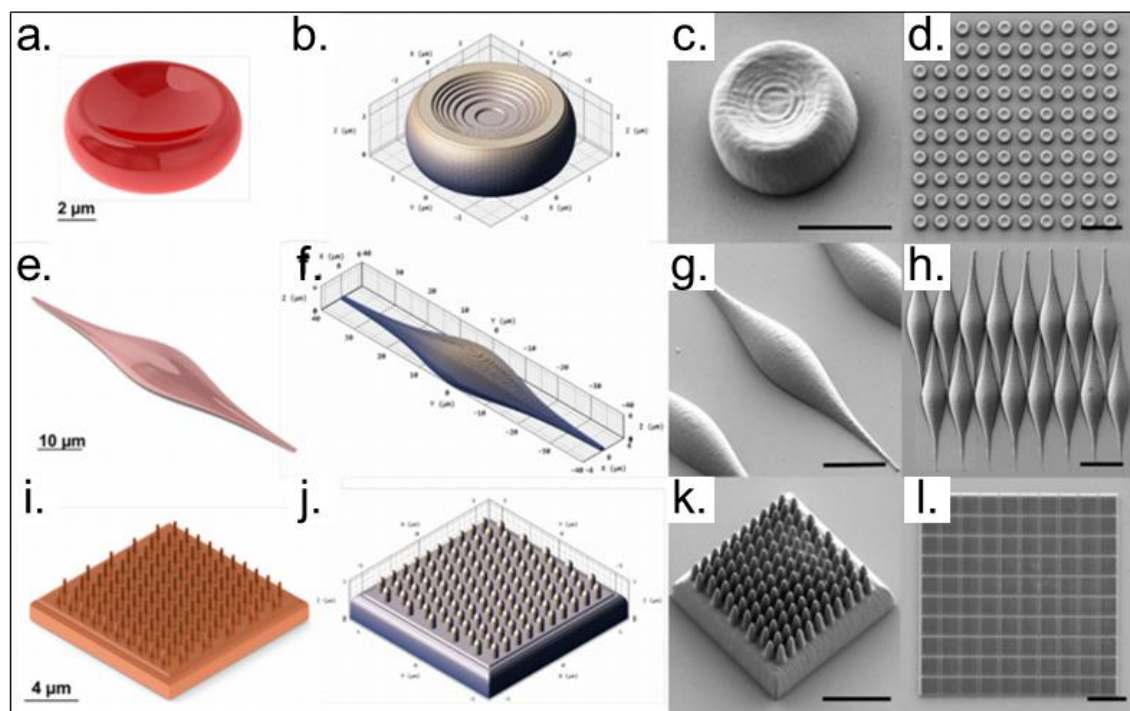


Figure 16. Images showing the designed and fabricated cell scaffolds using additive manufacturing with optimized parameters. (a–d) Red blood cell (RBC) model. (e–h) Smooth muscle cell (SMC) model. (i–l) Ciliated columnar epithelial cell (CEC) model. (a,e,i)—Rendered images of the 3D designs for a single cell. (b,f,j)—preview of a single cell generated in the DeScribe software after defining the parameters for the additive manufacturing fabrication process. (c,g,k)—Scanning electron micrographs of the fabricated individual cells, acquired using a 30° stage tilt, scale bar is 5 μm . (d,h,l)—120 $\times$ 120 μm^2 cell arrays, scale bar is 25 μm . Reproduced with permission [152]. Licensed under a Creative Commons Attribution 4.0 International license.

The growing interest in the tissue engineering and regenerative medicine fields has motivated the development of several bio-applicable materials. For instance, the use of a highly reactive photopolymer in TPP experiments, with their fabrication parameters optimized, allows the production of scaffolds with large size (in the order of millimeters in the three dimensions), with relevant size for clinical application [71].

Scaffolds sculpted from a biodegradable and biocompatible poly(trimethylene-carbonate) (PTMC)-based material by TPP (Figure 17a) supported the adhesion and proliferation of human adipose-derived stem cells (hASCs) [71]. Two days post-seeding, hASCs attached to the scaffold exhibited elongated filopodia (Figure 17b). Over 28 days of culture, the hASCs colonized the entire scaffold and formed a dense cellular layer on the surface of the scaffold (Figure 17c,d).

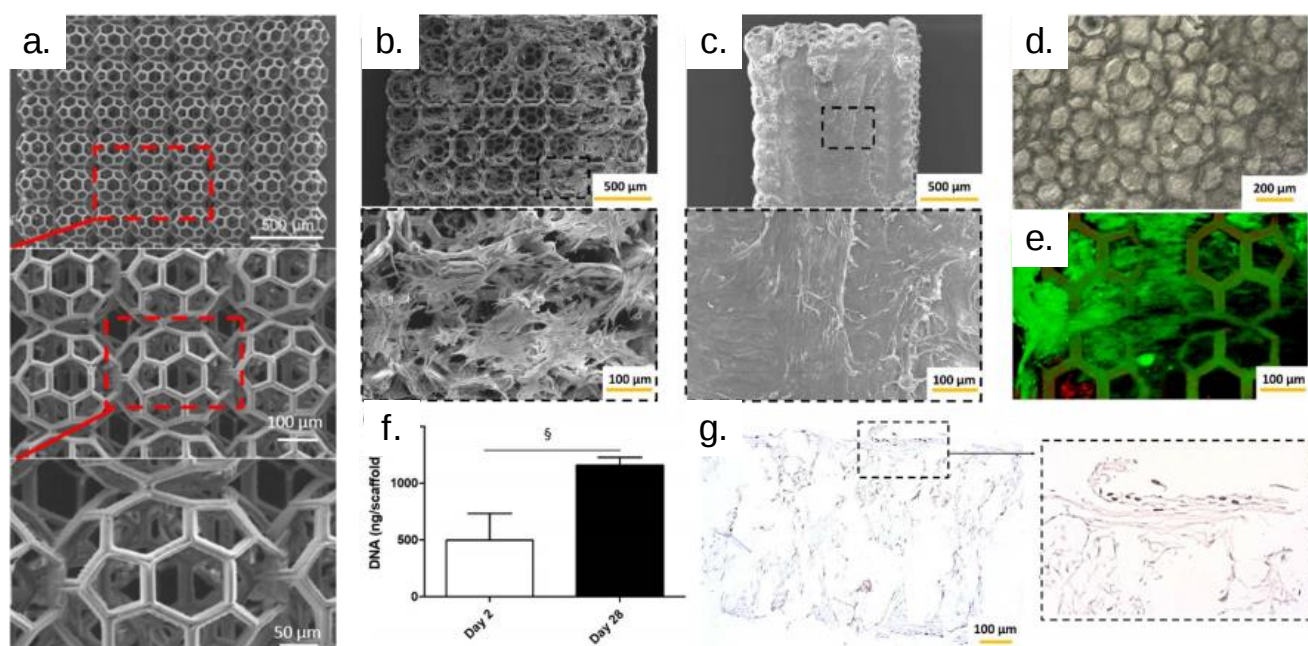


Figure 17. TPP-produced PTMC-based scaffolds support cell invasion and proliferation: (a) SEM images of the tri-layered PTMC-based scaffold at various magnifications ($\times 50$, $\times 150$ and $\times 300$). SEM images of cell-seeded scaffolds day 2 (b) and day 28 (c) post seeding. (d) Bright field and (e) live/dead illustrations of the colonized scaffold at day 28. (f) Cellular proliferation quantified by DNA assay (* indicates statistical significance) and (g) Ki67 immunostaining (nuclei of positive proliferative cells are stained in black). Adapted with permission [71]. Licensed under a Creative Commons Attribution 4.0 International license.

Figure 17e presents live/dead staining conducted after 28 days, showing a high cellular viability. The proliferation of the hASCs was validated by the significant increase of DNA (Figure 17f) and the presence of ki67-positive cells (Figure 17g) [71].

Cultured neuronal networks are the subject of many biological studies because they allow researchers to investigate neuronal activity in a controlled environment, in comparison to in a live organism. These experiments provide, for instance, a better understanding of the primary disease mechanisms and investigating the response to medicines [153]. TPP can be employed to produce 3D scaffolds, with high-resolution, using biocompatible and nondegradable photoreactive resins [153]. Figure 18 shows examples of 3D scaffolds fabricated using Dental LT Clear (DClear) resin employed in 3D neural cell culture and their characterization.

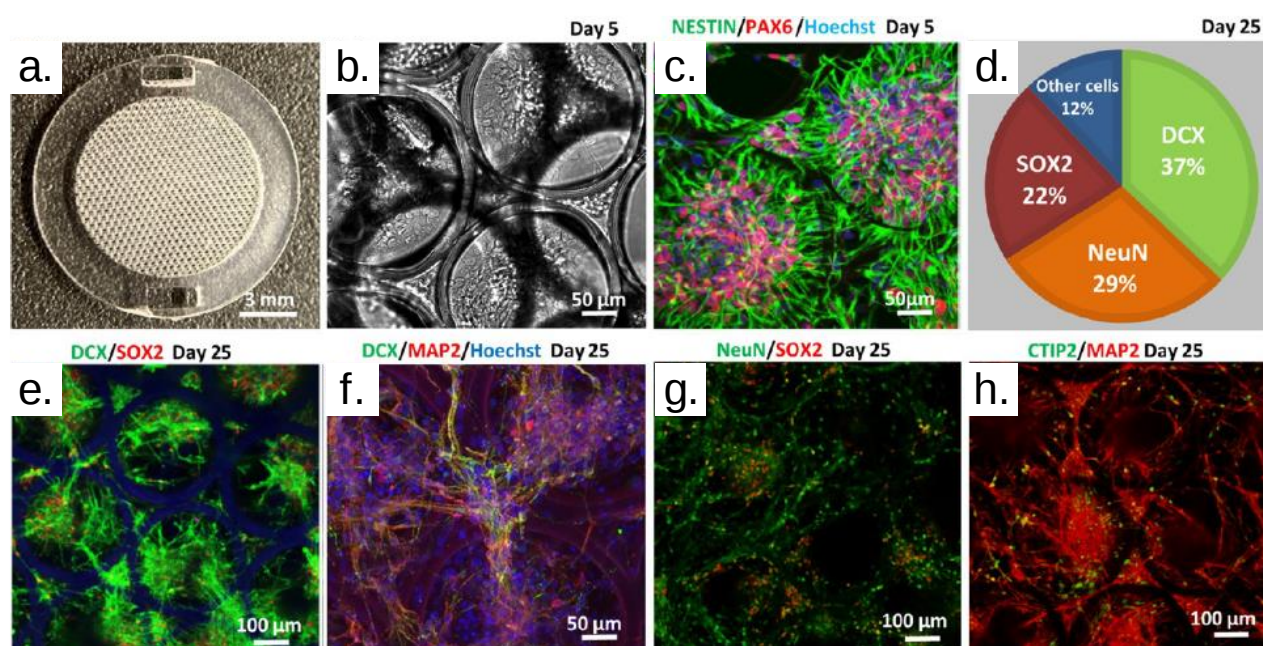


Figure 18. Generation of a 3D neuronal cell culture using laser-fabricated scaffolds. (a) General view of the scaffold structure (“honeycomb”) used for 3D neuronal culture development. (b) Phase-contrast image showing uniformly distributed cells within the scaffold 5 days after seeding. (c) Immunofluorescence image depicting well-accommodated NSCs in the pores of the scaffold. The labeled proteins are Nestin and PAX6, characteristic of cortical stem cells. (d) Quantitative ratio of young (DCX+) and mature (NeuN+) neurons as well as SOX2+ multipotent neural progenitor cells in the scaffold at day 25 of culture/differentiation. (e–g) Expression of young and mature neuronal markers DCX/MAP2/NeuN and SOX2, indicating a healthy formation of a neuronal network. (h) Differentiation of early-born CTIP2+ cortical neurons from neural stem cells. Reproduced with permission [153]. Copyright © 2021, American Chemical Society.

Figure 18a presents a 3D honeycomb-like scaffold fabricated via TPP, while Figure 18b shows a phase-contrast image demonstrating uniformly distributed cells within the scaffold 5 days after seeding. Still, in Figure 18b the scaffold design allows a noninvasive high-resolution visualization of real-time cell growth and neuronal network formation in a 3D environment. Immunofluorescence analysis showed that the fabricated scaffold provides a suitable 3D environment for the rapid development of neural stem cells (NSCs), facilitating the growth and long-term viability of neuronal networks (Figure 18c) [153], which may be related to the increased cell density in the cylindrical areas of the scaffolds that leads to strengthened neurons communications. Figure 18d represents a quantitative ratio of young (DCX+) and mature (NeuN+) neurons as well as SOX2+ multipotent neural progenitor cells in the scaffold at day 25 of culture/differentiation. In Figure 18, the expression of young and mature neuronal markers DCX/MAP2/NeuN and SOX2 indicate a healthy formation of a neuronal network. Finally, on day 25 of 3D culture, a large proportion of neurons already expressed the early-born cortical neuronal marker CTIP2 (Figure 18h) [153].

Microenvironments functionalized with biological compounds can also be sculpted using TPP. To evaluate the *E. coli* bacteria development, microenvironments were developed with selective functionalization using ciprofloxacin antibiotic. The inhibition of bacterial growth was primarily observed around the doped element. By analyzing images obtained from the microenvironment, it was possible to determine the maximum range of inhibition (12 μm), as well as to quantitatively determine the bacterial density in the inhibition halo [56].

The natural biopolymer named bacterial cellulose (BC) is an attractive material for biomedical experiments because it presents uniform structure and morphology, besides unique features, such as high purity, good mechanical properties, chemical stability, and others [154,155]. This polymer is completely biocompatible and can be produced in practi-

cally any shape due to its high moldability. Figure 19 shows 3D acrylate-based structures fabricated via TPP used to evaluate the BC formation through the development of *Komagataeibacter xylinus* bacteria [156]. In Figure 19, it is possible to observe the biofilms when bacteria are inoculated in the microstructure for 6 h (Figure 19a) and 24 h (Figure 19c). As displayed in Figure 19a, even after a short incubation time, it is possible to observe a network of nanofibers around the structures. After 24 h, it can be seen a thin film covering the microstructures. Here, it is important to highlight that the BC grown in microenvironments, such as the one presented in the Figure 19, have the same chemical composition as BC grown in macroscale. The use of BC as a flexible substrate in TPP experiments is very interesting for the fabrication of versatile devices. The natural polymer was not altered by the laser irradiation, such that its original properties were kept until the end of the photopolymerization. The structures well adhered to the natural substrate, even after washing the sample in heated ethanol (a procedure required to remove the unpolymerized resin). BC flexible substrates are potential platforms for liquid diffusion mechanisms in microenvironments, opening, for example, novel approaches for drug delivery and tissue engineering.

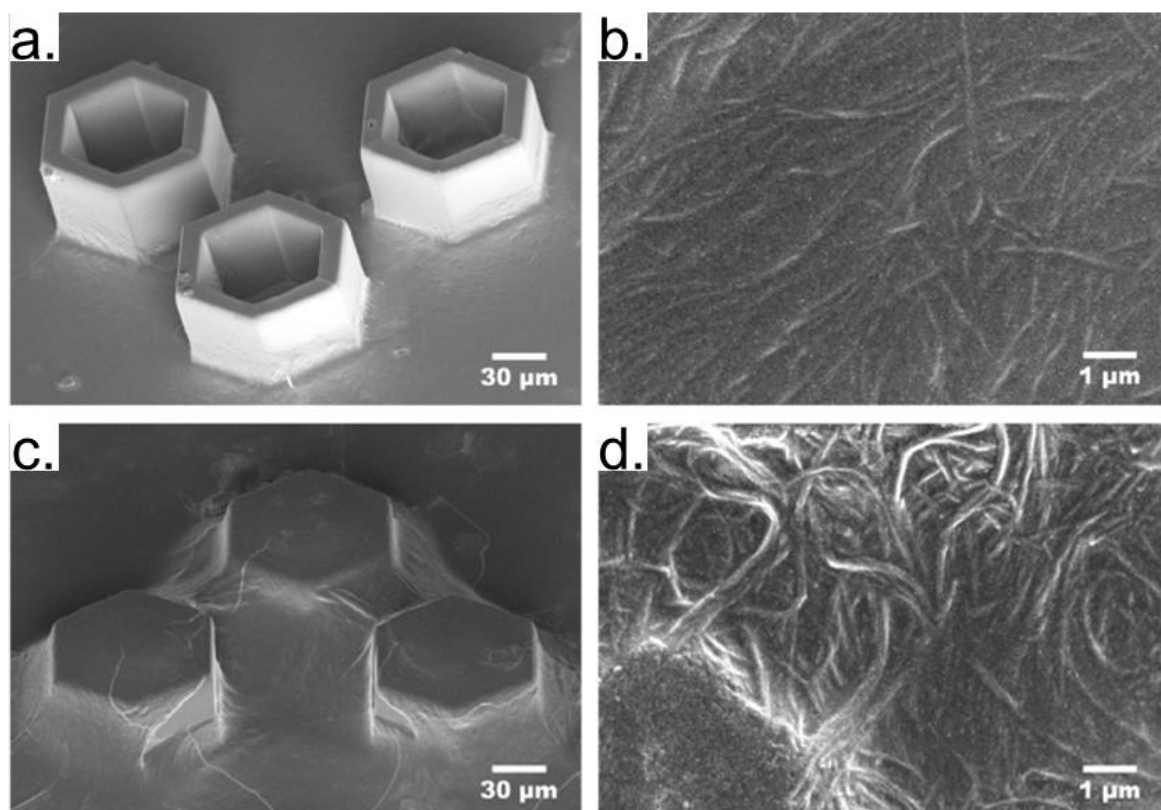


Figure 19. Bacterial cellulose (BC) grown on polymeric materials. (a) BC film formed after 6 h of bacteria incubation and (b) SEM image of the BC network grown in these structures. (c) BC film formed after 24 h of bacteria incubation and (d) SEM image of the BC network grown in these structures. Reproduced with permission [156]. Licensed under a Creative Commons Attribution 4.0 International license.

Often, biological studies require several simultaneous analyses, and the TPP experiments (in their most straightforward setup) are not attractive to the microenvironments fabrication. However, the TPP technique can be used as a tool to create the main environment, which can be replicated several times later. As an example, Figure 20a,b shows SEM images of the master nanograss and Figure 20c,d nanograss replica [157].

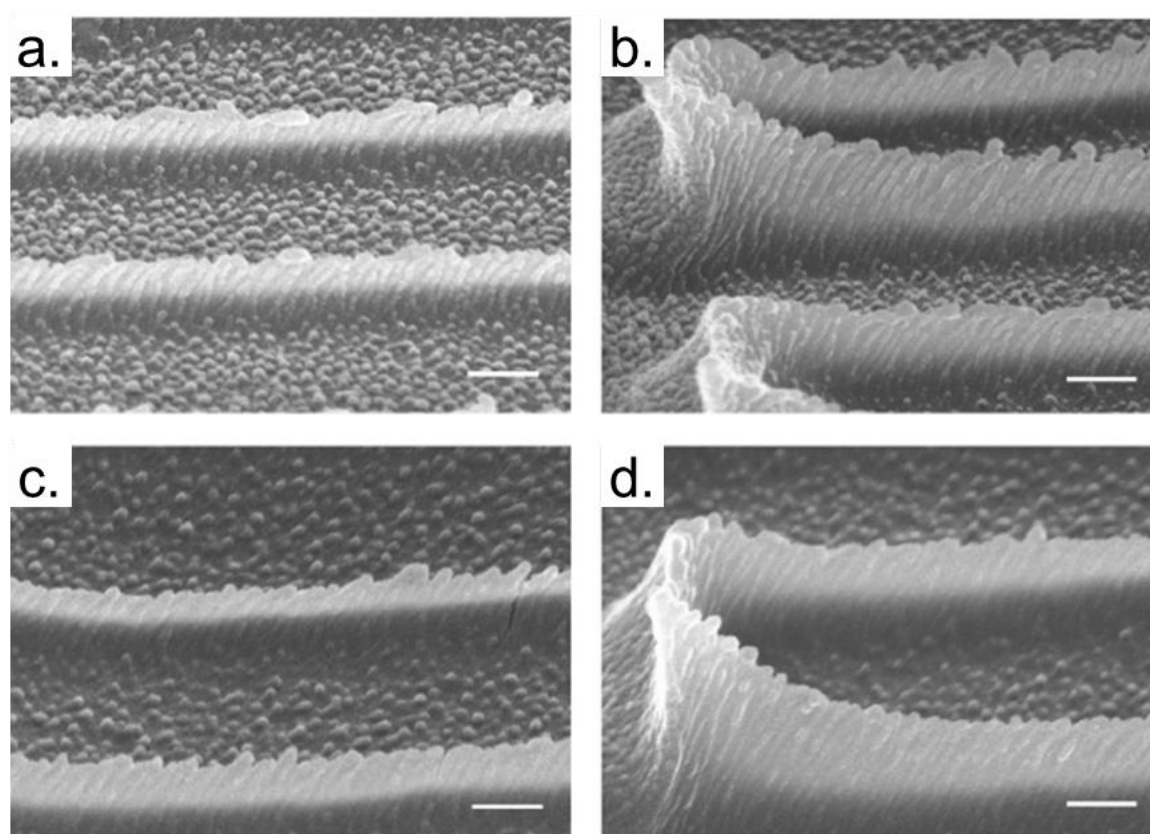


Figure 20. Replication of nanoglass. (a,b) SEM images of the master nanoglass. (c,d) nanoglass replica. Scale bars, 200 nm. Reproduced with permission [157]. Licensed under a Creative Commons Attribution 4.0 International license.

The mass production of nanotopographic surfaces opens new possibilities to study the mechanism related to cell migration and development. Besides, several replica experiments provide more accurate results, which can ease the understanding of microorganisms growth.

Based on the few examples presented in this review, it is possible to see the substantial importance of TPP technique to device manufacturing. The versatility of this fabrication method allows several advantages for sophisticated technological applications.

4. Final Remarks and Perspectives

This review discussed the technological potential of the laser micromachining method known as Two-Photon Polymerization (TPP), which has a unique set of advantages. Several materials have been used in TPP experiments, such as polymeric matrices, composites, and gelatins. The search for new materials that can be used in this technique is one of the major efforts of different current researches. Structures fabricated by this technique have great applicability in optics, photonics, and biology, mainly due to the possibility of functionalization with several compounds of interest or the manipulable devices fabrication. Combining TPP technique with other laser-processing methodologies, such as Laser Induced Forward Transfer (LIFT), allows the selective functionalization of the sculpted structures, creating new approaches to functionalize the devices. Even living microorganisms can be transferred for 3D scaffolds, without any damage. The wide variety of 3D structures that can be fabricated using TPP is one of the attractive advantages of this manufacturing technique. The growth of microdevices manufactured via TPP is notable, principally regarding photonics devices and bio-platforms development. For instance, high-Q WGM microresonators with distinct geometries can be fabricated by TPP in few steps, achieving performance comparable to resonators fabricated using other fabrication methods. Bioscaffolds also can be sculpted using TPP and replicated several times, keeping their original features. Many microenvironments have been created for studies and

monitoring of microorganisms development and can be easily functionalized with biological compounds of interest. Certainly, TPP is a consolidated laser fabrication technique with large applicability in the most diverse technological fields and will continue to be employed in sophisticated devices fabrication.

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