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Synthetic microfluidic paper: high surface area and high porosity polymer micropillar arrays†

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We introduce Synthetic Microfluidic Paper, a novel porous material for microfluidic applications that consists of an OSTE polymer that is photostructured in a well-controlled geometry of slanted and interlocked micropillars. We demonstrate the distinct benefits of Synthetic Microfluidic Paper over other porous microfluidic materials, such as nitrocellulose, traditional paper and straight micropillar arrays: in contrast to straight micropillar arrays, the geometry of Synthetic Microfluidic Paper was miniaturized without suffering capillary collapse during manufacturing and fluidic operation, resulting in a six-fold increased internal surface area and a three-fold increased porous fraction. Compared to commercial nitrocellulose materials for capillary assays, Synthetic Microfluidic Paper shows a wider range of capillary pumping speed and four times lower device-to-device variation. Compared to the surfaces of the other porous microfluidic materials that are modified by adsorption, Synthetic Microfluidic Paper contains free thiol groups and has been shown to be suitable for covalent surface chemistry, demonstrated here for increasing the material hydrophilicity. These results illustrate the potential of Synthetic Microfluidic Paper as a porous microfluidic material with improved performance characteristics, especially for bioassay applications such as diagnostic tests.

Introduction

High aspect ratio polymer micropillars have received a lot of research attention recently due to their unique mechanical, optical, and surface properties. They have been utilized as, e.g., biomimicking dry adhesives,¹ airflow sensors,² and superhydrophobic surfaces.³ Within microfluidics, micropillar arrays have been used as capture surfaces for cells⁴ and biomolecules,⁵ in capillary pumping,⁶ for particle sorting,⁷ and for studying cell behavior.⁸⁻¹⁰ At high aspect ratio and high density, micropillar arrays are prone to collapse due to adhesive and capillary forces during liquid or humid operation or during manufacturing schemes involving liquids, such as lithographic processes. Micropillars collapse towards each other when the capillary force is larger than the opposing elastic force exerted by the pillars and the effect is irreversible when the adhesive forces that hold the pillars together exceed the elastic forces that pry them apart.¹¹ While capillary micropillar collapse has been used to create novel functional structures,¹² it is generally undesired due to unpredictable material properties on the microscale. Micropillar arrays of high density are desired in applications where a large relative surface area is sought, e.g. for surface chemical interactions or for high capillary pumping pressure. Micropillar arrays with small pillar diameter and large height are desired in applications where high flow throughput is sought, e.g. for filtration or for high capillary flow.

In previous work, increased aspect ratios have been achieved by strengthening the mechanical properties of the material, e.g. by reinforcing pillars with stiff materials¹³ or using highly cross-linked polymers as the constituting material.¹⁴ However, these approaches still have limits on aspect ratio, and also exclude utilizing low modulus polymer materials, which often are of special interest in microfluidics for use in flexible devices¹⁵ and self-folding structures.¹⁶ Moreover, low modulus polymer materials open up for new chemical groups, not readily available in highly cross-linked materials. Examples of such materials include photocurable PDMS,¹⁷ polyacrylate hydrogels,¹⁸ and the recently introduced Off-Stoichiometry-Thiol-Ene (OSTE),¹⁹ a polymer system that enables excellent surface control due to inherently available functional groups on its surface.

In contrast to current micropillar array designs, we realize capillary collapse resistance through geometrical design by slanting and interlocking the constituting micropillars in truss-like structures. We explore the geometrical limits of capillary collapse for traditional straight micropillar arrays
and for slanted interlocked micropillars with the same pitch and diameter, which are manufactured in the same low modulus polymer. We show the increase in surface area and in porous fraction enabled by the new geometry. We call this novel material Synthetic Microfluidic Paper and demonstrate its use as a capillary pump.

Material design

Fig. 1 illustrates the design and manufacturing of Synthetic Microfluidic Paper using slanted interlocked micropillar geometry. The material consists of four sets of interlocked pillars, all sets inclined with an angle of $\phi \approx 40^\circ$ with respect to the vertical axis, and the four sets being in azimuthal directions $0^\circ$, $90^\circ$, $180^\circ$, and $270^\circ$, thus causing the pillars in different sets to interlock. The geometry is further determined by the interpillar pitch ($p$), the pillar diameter ($d$), and the vertical height of the structures ($h$). The number of interlocking points for every pillar depends on the height and interpillar pitch. We chose lithographic manufacturing, instead of 3D-printing or laser writing, because of its scalability towards volume manufacturing and the ease with which sub 100 micron features are replicated. All devices are manufactured in Off-Stoichiometry-Thiol-Ene (OSTE), which previously has demonstrated excellent lithographic properties and easy surface modification. To realize the slanted pillar geometry, we utilize multidirectional UV-lithography, previously used for stoichiometric Thiol-Enes and Thiol-Ene-Epoxies, which is here realized in a single exposure using an angled mirror setup as illustrated in Fig. 1c (see Fig. S2 in the ESI† for photo of the setup).

Materials and methods

Manufacturing

The manufacturing process is performed in four steps: (multidirectional) photopatterning, developing, flood curing, and drying. Straight micropillar arrays are formed by vertical illumination through a photomask containing circular openings; slanted interlocked micropillars structures in Synthetic Microfluidic Paper are formed by multidirectional lithography through the same mask. Multidirectional lithography is achieved by reflecting the vertically collimated light to angles of 60 using four aluminum mirrors. OSTE polymer consists of tetraallyloxyethane (Tokyo Chemical Industry Co., Ltd., Japan) and tris[2-(3-mercaptopropionyloxy) ethyl] (BOC Sciences, NY, USA) with 40% thiol group excess, 1% Irgacure 184 (BASF Corp., USA) and 1% effective weight of Q1301 (Wako Chemical Inc., USA) in a saturated toluene (Sigma-Aldrich, USA) solution. Development is done in butyl acetate (VWR, USA).

Capillary collapse testing

The micropillar geometries tested feature pillar diameters of $d = 10–100$ μm, pitch $p = 30–400$ μm, and height $h = 50–450$ μm. The Young’s modulus of a fully cured OSTE material slab was measured to be 5.5 MPa using dynamic mechanical and thermal analysis (DMTA) (DMA Q800, PerkinElmer, Waltham, USA). To test the structural resistance against capillary collapse, drying of the butyl acetate developer under a fume hood during the last manufacturing step is used, followed by visualization of the resulting pillar array geometry using microscopy. The contact angle of butyl acetate, having a surface energy of 0.0251 N m$^{-1}$, to the OSTE surface was measured to be 5.8$^\circ$ by the sessile drop method using Attension Theta Lite (Biolin Scientific, Sweden).

Capillary collapse classification

All tested array geometries were inspected by light microscopy using 4–20× long-range objectives (Leica, Germany) from the top, from an angle, and from a teared-off cross-section when necessary. For arrays difficult to classify (mostly for small geometries), inspection by SEM was also conducted. Pillars were defined as collapsed when they were stuck to
each other or on the bottom surface. Pillars were defined as non-collapsed when adherence to each other or to the bottom surface was not observed. For arrays containing both collapsed and non-collapsed parts, the most prevalent feature defined the classification. Examples of this classification can be seen in Fig. 2c–f. Manufacturing imperfection prevented studying the collapse of some of the smallest structures. These cases were classified as failed manufacturing. Overcuring, which was the main failure mode, was detected in polymer structures extending between pillars and blocking the void between interlocked pillars, i.e. in the masked regions.

**Solvent uptake in OSTE**
The mass of a fully cured OSTE slab, 1 mm × 11 mm × 27 mm, was measured with a scale (Kern ABS, Germany), before and after incubation in butyl acetate for 48 h. Change in mechanical properties of the soaked sample was studied by manual manipulation of the sample using tweezers.

**Hydrophilic surface treatment**
The fabricated OSTE devices prepared for capillary filling experiments was rendered hydrophilic by UV grafting of HEMA. A solution of 5% hydroxyethylmethacrylate (HEMA) (Sigma-Aldrich, USA), 0.1% Lucirin TPO-L (BASF Corp., USA), and 0.1% benzophenone (BP) (Sigma-Aldrich, USA) in isopropyl alcohol (IPA) (SunChem AB, Sweden) was prepared. A glass slide was placed above the device, with 150 μm spacing between the top surface of the interlocked pillar array and the glass slide, where the solution was injected. Collimated UV-light was

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**Fig. 2** Capillary collapse resistance of straight micropillar arrays compared to Synthetic Microfluidic Paper (slanted interlocked micropillar arrays), showing that Synthetic Microfluidic Paper allows geometries beyond the practical and theoretical limitations of straight micropillar arrays. Test structures are of 150 μm height, fabricated using a photolithography mask with a square pattern of openings with diameter \( d \) and pitch \( p \). (a–b) Plots of the results for different opening diameters and pitches for (a) straight pillars and (b) slanted interlocked pillars. The diameters of the pillars tested are 10, 20, 30, and 50 μm. The solid red line in (a) indicates the theoretical capillary collapse limit for straight pillars; the dotted red line in (a) indicates the limit of ground collapse observed, and the dashed black line in (a) and (b) indicates where pillars overlap \( p > d \). (c–f) SEM images of (c) a non-collapsed straight pillar array at \( d = 30 \mu m, p = 105 \mu m \), and \( h = 150 \mu m \), (d) a collapsed straight pillar array at \( d = 20 \mu m, p = 60 \mu m \), and \( h = 150 \mu m \), (e) a non-collapsed slanted interlocked micropillar array at \( d = 10 \mu m, p = 110 \mu m \), and \( h = 150 \mu m \), and (f) a collapsed slanted interlocked micropillar array at \( d = 10 \mu m, p = 110 \mu m \), and \( h = 150 \mu m \). Scale bars are 30 μm.
illuminated onto the solution and the pillar array through the glass slide for 300 seconds. After the exposure, the pillar array was rinsed with pure IPA, soaked and incubated with pure IPA for 300 seconds, and dried at room temperature.

Capillary filling

Three types of capillary driven devices \((d, p, h) = (100, 150, 150), (100, 100, 150),\) and \((20, 40, 150)\) μm were prepared in 5 mm wide strips and subjected to a hydrophilic surface treatment. For blood imbibition, 30 μl of human blood (sample was collected from a healthy human subject, after informed signed consent was obtained) was applied onto one end of a device with a pipette. For water imbibition, the devices were put into contact with a water meniscus defined between 2 glass slides, 1 mm apart. Filling of the devices was recorded by a camera (EOS 500D, Canon, Japan) and analyzed in Matlab using a threshold algorithm for one image per second of the recorded movie.

Results and discussion

Capillary collapse resistance

Fig. 2 shows the result of the capillary collapse resistance testing, which is performed by evaporating the solvent from interlocked pillar arrays and from straight pillar arrays, each of 150 μm height. Fig. S3 (see the ESI†) shows the corresponding results for arrays of 50, 300, and 450 μm height. We first discuss the comprehensive results in Fig. 2 and then see that the additional data in Fig. S3† confirms our conclusions.

For straight pillar arrays, indicated by filled circles in Fig. 2a and imaged in Fig. 2c, collapse resistance is achieved when the pillar diameter, \(d\), and pitch, \(p\), are not too small. The pillars, 10 μm in diameter, collapse even at a high pitch, which can be explained by ground collapse, i.e. the interaction between the pillars and the bottom surface.24 The limit of ground collapse, which is independent of the pitch, can here be estimated between \(d = 10\) μm and \(d = 20\) μm and is indicated by a red dotted line in Fig. 2a.

The theoretical limit of capillary collapse for straight pillars is plotted as a solid red line in Fig. 2a, based on an approximation of the theory from Chandra and Yang11 for OSTe as the constituting material and butyl acetate as the liquid medium (\(E_{\text{OSTe}} = 5.5\) MPa, contact angle \(\theta = 5.8^\circ\), and interfacial energy, \(\gamma_{\text{liquid-vapour}} = 0.025\) N m\(^{-1}\); see the ESI† for details). Theory and experimental results are in good agreement for pillar diameters \(d \geq 50\) μm, whereas for smaller pillar diameters, the pillars collapse at a much larger pitch than that predicted by theory. We speculate that this might be caused by the influence of the solvent on the mechanical and geometrical properties of the pillars: experiments reveal 8% weight increase and a significant decrease in E-modulus of the pillars when exposed to the solvent. The exact E-modulus of the pillars during evaporation is difficult to measure; however, the effect of the solvent can be expected to be more severe for small pillar diameters due to their increased surface-to-volume ratio, which would explain the deviation from theory for these geometries. Additionally, the propensity for capillary collapse of small diameter straight pillars emphasizes the need for capillary collapse resistant alternatives.

Non-collapsing slanted interlocked micropillars, indicated by filled squares in Fig. 2b, are achieved for a substantially larger range of pitches and diameters than for straight pillar arrays. Slanted interlocked micropillars only collapse when diameter \(d = 10\) μm and pitch \(p \geq 60\) μm. Interlocked pillars thus withstand collapse for pitches and diameters below the theoretical limits of straight pillars. Part of the explanation for this can be found in the mechanical nature of the interlocked micropillars. Whereas freely suspended straight pillars collapse by bending11 (seen in Fig. 2d), interlocked pillar collapse by buckling (see Fig. 2f). Moreover, the interlocking points partition each pillar in shorter sections, which are more difficult to deform and which have a section length:

\[
L = \frac{p - d}{2\sin\theta}
\]

that decreases with the interpillar pitch and is independent from the overall structure height, \(h\). Collapse occurs when the capillary force, \(F_c\), is larger than the restoring elastic force of the pillar, \(F_e\). The dependence of these forces on the geometrical parameters is

\[
F_c \propto \frac{d^2}{p - d}
\]

\[
F_{e,\text{straight+bending}} \propto \frac{d^4}{h^2}
\]

\[
F_{e,\text{interlocked+buckling}} \propto \frac{d^4}{(p - d)^2}
\]

i.e., with decreasing interpillar distance, \(p - d\), the capillary force, \(F_c\), increases linearly; the mechanical strength of straight pillars against bending, \(F_{e,\text{straight+bending}}\), remains unaltered; but the critical buckling strength of interlocked pillars, \(F_{e,\text{interlocked+buckling}}\), increases quadratically. Consequently, straight pillar arrays collapse with decreasing interpillar distance, whereas slanted interlocked pillar arrays become increasingly more collapse resistant. The current limit in decreasing the interpillar distance is the manufacturing scheme: we observe overcuring, manifested as pillar broadening, which depends on diffraction of light and on the exact UV-exposure time, and which could be optimized in future work.

During capillary collapse testing, an interesting dynamic behavior is observed while the solvent evaporated from the slanted interlocked pillars. Fig. 3 shows how, first, the capillary force causes part of the slanted interlocked micropillars to collapse down towards the substrate. While the solvent evaporated, the energy stored in the buckled pillars restored...
the structure to its original non-collapsed geometry. This phenomenon was observed for a few structures with a relatively large pitch and small diameter ($d = 20 \mu m$ and $p = 100 \mu m$, for $h = 150$ and $300$), but not for larger diameters and smaller pitches. This self-restoring property only applies to elastomeric or low modulus materials and could potentially be utilized in self-folding micro- or macrostructures in future developments.

**Increased surface area and porous fraction**

Due to their capillary collapse resistance, the slanted interlocked micropillar arrays of Synthetic Microfluidic Paper provide two specific benefits over straight pillar arrays: increased surface area and increased porous fraction.

The increase in solid surface area, $A_p$, compared to the device footprint area, $A_{fp}$,

$$\eta = \frac{A_p}{A_{fp}}$$

is a measure of the potential of the device for surface capturing assays, such as cell or biomolecular capturing. Fig. 4a illustrates that the area increase scales inversely with the pillar diameter and pillar pitch. The maximum increase for straight pillars is 7.5, whereas the best area increase for slanted interlocked pillars is 45, i.e. a six-fold increase. This is a direct result of the capillary collapse resistance, which allows thinner and denser pillar arrays. Moreover, each $p^2$ unit of the footprint area contains one straight pillar, but four slanted interlocked pillars. The porous fraction for lateral flow, $f_p$, is the pore area (i.e. the interpillar area) per total area, i.e.

$$F_p = \frac{p - d}{p}$$

Fig. 4b shows the relative increase of the porous fraction of slanted interlocked pillars compared to straight pillars for different values of the interpillar distance $p - d$. The relative porous fraction of slanted interlocked pillars increases with decreasing interpillar distance, and is three times larger for 10 $\mu m$ interpillar distance arrays. This increase is the result of the interlocking design, which allows for a smaller pillar diameter at a given interpillar distance. The relative increase in porous fraction for Synthetic Microfluidic Paper is expected to allow for an equal increase in flow throughput in capillary pumping and particle sorting or filtering applications, for the same capillary pressure and filter cut-off values, respectively.

**Capillary pumping**

To demonstrate its practical use in microfluidic applications, 5 mm wide strips of Synthetic Microfluidic Paper, consisting of capillary pumps were prepared. Geometries of the slanted
interlocked micropillar arrays were chosen to demonstrate pumped volume and flow rates relevant for lateral flow assays, i.e. similar to previous lateral flow assay materials. First, the free surface thiol groups of the OSTE polymer are utilized to bind hydrophilic HEMA molecules, resulting in a lowered surface contact angle of 25° for DI-water. A drop of human blood is placed on top of the strip, resulting in imbibition of the blood into the Synthetic Microfluidic Paper. Fig. 5a and b show the capillary flow of blood in Synthetic Microfluidic Paper with geometries \( (d = 50 \, \mu m, p = 150 \, \mu m, h = 150 \, \mu m) \), \( (d = 50 \, \mu m, p = 100 \, \mu m, h = 150 \, \mu m) \), and \( (d = 20 \, \mu m, p = 40 \, \mu m, h = 150 \, \mu m) \), demonstrating volumetric pumping capacities of 16 \( \mu l \), 6 \( \mu l \), and 9 \( \mu l \) in one minute, respectively. Least mean square regression fitting shows that the blood liquid front propagation scales approximately \( (R^2 = 0.99, 0.98, \text{ and } 0.97, \text{ respectively}) \) with \( t^{1/2} \), as expected from the Washburn equation, with liquid front velocities of 1.7 mm s\(^{-1/2}\), 1.8 mm s\(^{-1/2}\), and 2.0 mm s\(^{-1/2}\), respectively. This flow behavior indicates that the material is suitable as a capillary pump, and that the pumping rate makes it a potential alternative to other porous microfluidic materials, such as paper, nitrocellulose or straight micropillar arrays. Comparable flow rates in other materials have been reported previously, e.g. 4 mm s\(^{-1/2}\) in a polymer micropillar array\(^6\) for serum (serum having 2–3 times lower viscosity then blood) and 1.6 mm s\(^{-1/2}\) for blood in nitrocellulose.\(^{26}\) The slight deviations from the Washburn equation follow mainly from our experimental measurement setup and the difficulty of exactly determining the starting point of propagation from an imprecisely placed blood drop.

Fig. 5c shows water imbibition tests from a well-defined water meniscus, on 4 devices of Synthetic Microfluidic Paper with identical geometries \( (d = 50 \, \mu m, p = 150 \, \mu m, h = 150 \, \mu m) \), resulting in a liquid flow speed of 9.9 mm s\(^{-1/2}\), i.e. 4 cm in 16 s, with a standard deviation of 2%. This is a considerable improvement over commercial nitrocellulose substrates, which feature a liquid pumping speed of 75 s for 4 cm for the fastest nitrocellulose substrate, with a standard deviation of 8%.\(^{27}\)

**Other potential applications**

The successful covalent surface modification of the slanted interlocked micropillars indicates the versatility of the material and motivates other future potential microfluidic applications, such as high area/volume biosensing surfaces.

Although not experimentally tested, we can speculate on the use of slanted interlocked pillars for other applications. The surface area increase makes our novel geometry especially beneficial for microfluidic applications relying on high capture surface.\(^4,5\) The combination of controlled surface properties, capillary evaporation, and the possibility of using low E-modulus polymers provides new possibilities for flexible devices and bioinspired self-folding materials. Our slanted interlocked pillars may be suitable for use as biomimicking dry adhesives, which rely on sub-micrometer pillar features, with as high as possible number of pillars capable of attaching to a surface \((-p^{-2}\)) and whose particular geometry \((h, d)\) is of less importance, as long as the pillars are flexible enough to attach to uneven surfaces without breaking, curling or tangling.\(^1\) The slanted pillar design introduced here may allow increase in the pillar density, although this would require further development of the 3D manufacturing to the sub-micrometer scale. The structure is indeed superhydrophobic\(^1\) for its native surface, but airflow sensing,\(^5\) relying on flexible pillars, can be expected to have an inverse effect on sensitivity at low flow rates. For particle sorting applications, the increased porous fraction is beneficial. Moreover, the 3D nature of this material could open up particle sorting in two dimensions instead of only one, as for straight pillars.\(^7\)
Conclusions

We introduce Synthetic Microfluidic Paper, consisting of slanted interlocked micropillar arrays, as a novel porous substrate material in microfluidics. We studied this novel microfluidic material as an alternative to traditional straight micropillar arrays and to nitrocellulose. We show that, compared to straight micropillar arrays, Synthetic Microfluidic Paper features higher resistance to capillary collapse, even beyond the fundamental limitations of capillary collapse for straight micropillar arrays. This results in a material with up to six-fold increased surface area and up to three times higher porous fraction compared to straight micropillar arrays. The latter is expected to allow increase in flow throughput in capillary pumping and particle sorting or filtering applications. We also demonstrate the use of Synthetic Microfluidic Paper for reliable capillary pumps. Compared to nitrocellulose, Synthetic Microfluidic Paper shows a wider range of pumping speeds and four times lower device-to-device variation. We foresee future applications of the material, e.g. in biosensing applications and self-folding structures.

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