**Patterning of worm-like soft polydimethylsiloxane structures using a TiO2 nanotubular array**

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ABSTRACT: Patterned polydimethylsiloxane (PDMS) is an important structure for soft lithography. Various materials have been deployed as mould for patterning PDMS. Anodised nanotubular array has been sought after as cost-effective alternative for textured silicon. An array of TiO2 nanotubes with characteristic diameter ≈140 nm and the length of ≈1.5 microns, created by anodic oxidation of a titanium substrate, was used here as a template for soft PDMS moulding. The optimal moulding process was developed by a combination of silanisation, use of solvent, application of a vacuum and hydraulic pressing. The silanisation was confirmed by Fourier transform infrared spectroscopy and contact angle measurements while the PDMS structure was examined by scanning electron microscope and energy dispersive X-ray spectroscopy. Hydraulic pressing significantly improved the infiltration of PDMS into the pores of nanotubular array resulting in formation of PDMS nano-bumps after separation of the polymer from the template. Complete infiltration of PDMS precursor into the cavity of nanotubes was observed on the hydraulic-pressed sample without toluene impurities. The hydraulic-pressed samples exhibited higher adhesion strength than non-pressed ones. The adhesive strength was measured by a simple experimental arrangement, in which the PDMS layer was stuck on a vertical glass surface followed by pulling it downwards.



Graphical Abstract

**1. Introduction**

Soft lithography, an alternative method to photolithography, provides low cost and rapid fabrication of micro- and nanostructures by eliminating the need for intense radiation equipment 1. Such nanofabrication technique is rapidly advancing many areas of research such as tissue engineering 2, protein printing 3, anti-biofouling 4, dry adhesive 5, and microfluidic 6. The process begins with fabrication of patterned mould or stamp which is deployed to replicate the micro- or nanostructure. Polydimethylsiloxane (PDMS) is often used as an elastomeric stamp or mould due to its low shrinkage rate 7, hence the name “soft” in soft lithography. The stamp or mould with patterned structure is a key aspect to prototype micro- and nanostructure 8. The patterned surface shows promise for applications in microelectronics and life sciences. In microelectronics, patterned PDMS is usually used as stamp to fabricate thin film of semiconductor, metal, or insulating materials for microelectronic devices 9. In life sciences, textured PDMS is often utilised as substrate for microfluidic devices especially for studying the cell adhesion and protein adsorption 10. It also can be used as dry electrode sensor patch in wearable sensor 11. The development of an elastomeric stamp or mould on a submicrometre or nanometre scale is crucial.

A PDMS stamp can be achieved either by masking the flat PDMS followed by etching with ultraviolet (UV) light 12, or by moulding of PDMS using textured surfaces from natural or synthetic materials. Natural materials such as plant leaves 13–15, shark skin 16, and sea urchin 17 have micro-patterned surface that can be deployed for PDMS mould. Geometrically complex surfaces from nature may provide the opportunity to enhance understanding of microbial attachment on a unique topography. However, the low stability of the natural products prevents high curing temperature of PDMS which may alter the surface texture of the natural mould. Low curing temperature (40-80oC) is usually used for natural mould yielding relatively low elastic modulus of PDMS even after a longer curing time. For high curing temperature, synthetic materials such as micro/nanofabricated silicon 18,19 , SU-8 epoxy 20, CuO nanowire 21 porous nickel oxide 22, and alumina nanotubes array 23–25 can be utilised.

Photolithography is needed to synthesise patterned silicon, SU-8 epoxy, and CuO nanowire while a relatively simpler and cost-effective technique such as anodising and electrodeposition can be used for ceramic microfabrication. Nickel particles were electrodeposited from a nickel chloride aqueous electrolyte (pH 4) on the stainless steel cathode using a DC power supply (3 A) 22. Hydrogen gas evolution at the cathode resulted in pore formation in the nickel layer. The deposited nickel particles mutually coalesced on sintering at 600‒800oC while larger pores left cavities (5 µm in diameter) in the film. An uncured PDMS mixture was poured into the cavities, cured then peeled from the nickel film, creating a microfabricated PDMS pillared film of 3‒5 µm diameter and 5‒10 µm long.

An anodic aluminium oxide (AAO) nanotubes template with a diameter of 25‒300 nm was attainable by controlled anodising of aluminium in presence of fluoride ions 26. In place of a negative replica of nanotubes (e.g., pillars, fibres), cone shaped bumps with a diameter of ≈100 nm were fabricated by moulding soft PDMS (Sylgard 184) into 100‒200 nm anodised aluminium oxide (AAO) nanotubes 23,24. When using aluminium as a substrate for the aluminium oxide layer, the heating rate required carefully control to avoid crack formation between aluminium and aluminium oxide, since the coefficient thermal expansion (CTE) of aluminium and aluminium oxide are 13.1x10-6 °F-1 and 4.5x10-6 °F-1 27. The CTE values of titanium and titania are 8.41x10-6 °C-1 and 9.0x10-6 °C-1 so titanium has less thermal mismatch and is suitable for heating 27. Moreover, the surface wetting properties of silanised titania shows can be modified from hydrophobic to hydrophilic on UV irradiation 28 and annealing 29, providing a versatile surface for moulding polymers.

An accurate replica of soft PDMS with high aspect ratio and good quality using a pores smaller than 150 nm is still challenging task 19. A higher fidelity of PDMS at a smaller diameter (≈100 nm) was achieved on the silicon template, with composites of two different PDMS layers 30. A stiffer polymer consisting of vinyl and hydrosilane end-linked polymers (hard PDMS, h-PDMS) was used as the first layer which improved the quality of the replica due to its lower viscosity. Whilst soft PDMS was deployed as second layer to give a flexible support of PDMS composite film. However, only nanobumps of soft PDMS with a height ≈ 25 nm and ≈ 80 nm in diameter were observed in a high density AAO nanotube array 25.

Anodising is a relatively simple and cost-effective method which is promising for lithography. To the best of our knowledge, only soft PDMS bumps with low aspect ratio (<2) were produced by replicating anodised sample 23–25. An intensive research is needed to improve the aspect ratio of PDMS replica. In this work, the facile approach of anodising was used to fabricate a titanium oxide or titania nanotubular template. Titania can be produced in controlled morphologies, including nanotubes, nanofibres, and nanosheets 31,32. Grimes *et al.* 33 introduced the first generation of titania nanotubes up to 500 nm long by anodising in hydrofluoric acid (HF) aqueous solution. During anodising, titania nanotubes were also simultaneously etched by the acid, limiting the length of nanotubes. As an alternative, non-aqueous solutions (e.g., glycerol, ethylene glycol) containing fluoride ions can be used as electrolyte 34,35. The limited solubility of titanium fluoride in ethylene glycol-based electrolytes manifested a high aspect ratio and ultra-long nanotubes up to 720 µm which are comparable to hard anodising of aluminium. A more viscous solution, such as glycerol, exhibited a moderate growth rate of up to 7 µm for 13 h. Here, environmentally friendly glycerol was used to produce a pronounced nanotube length. Appropriate control of water content and applied voltage can improve diameter and length of the tubes while the duration of anodising affects their length 36. However, water may distort the shape of the ordered tubes array, so the water content was kept at 2 vol.% in this work.

Patterned PDMS was produced using only one layer of soft PDMS (Sylgard 182) for simplicity. Sylgard 182 is similar to Sylgard 184 with a longer working time at 25oC 37 allowing more time for PDMS to fill the pores of the template. An optimal moulding process was achieved by a combination of silanisation, the use of solvent, the application of a vacuum, controlled peeling rate, and hydraulic pressing. The importance of hydraulic pressing and slow peeling rate was observed by scanning electron microscope imaging, elemental analysis and contact angle measurement. A simple adhesion test was deployed in this work to introduce potential applications of worm-like PDMS structures.

**2. Experimental Section**

**2.1. Synthesis of the titania nanotubular template**

Titania nanotubes array was synthesised by potentiostatic anodising of 0.25 mm thick titanium foil (99.5% metals basis purity, Alfa Aesar) in fluoride contained glycerol electrolyte 38. The foil was cut into 4 cm x 1.5 cm and wet polished using distilled water with 320, 800, 1200, 4000 grades of SiC, respectively. These were followed by rinsing with water and acetone. Anodising was performed using a DC regulated power supply in a two-electrode configuration with titanium foil as anode and graphite as cathode at 1.2 cm separation. An area of 2 cm x 1.5 cm of titanium foil was immersed in an electrolyte containing 0.25 wt.% NH4F and 2 vol.% DI water in glycerol (≥98.5%, Sigma Aldrich). A constant voltage (60 V) was applied for 6 h anodising at 22±2.5oC with a stirring speed of 150 rpm to synthesise a uniform titania nanotubular array. After anodising, samples were taken out, rinsed with ethanol, dried in a N2 stream followed by bath ultrasonication in acetone for 1 min.

**2.2. Silanisation of the titania nanotubes array**

Titania nanotubes array were treated with fluoroalkyl silane solution to create self-assembled monolayer coating 39,40. The nanotubes array were sequentially immersed for 1 h in methanol solution of 1.0 wt.% 1H,1H,2H,2H-perfluorooctyltrichloro silane (FOTS, Sigma Aldrich), rinsed with pure methanol and dried in N2 stream. The array was subsequently heated at 140oC for 1 h.

**2.3. Moulding of polydimethylsiloxane**

Several coated titania nanotubular templates were soaked in toluene for 10 min or directly deployed for PDMS moulding. Commercially available PDMS, Slygard 182 elastomer kit (Dow Corning) contains silicon elastomer base and curing agent. Both components were mixed in a ratio of 10 to 1 (elastomer base to curing agent). Toluene was added to dilute the mixture, creating either 50 vol.% or 33.33 vol.% of PDMS. 500 µL of the mixed solution was casted onto 2 cm x 1.5 cm of titania nanotubes array and left under vacuum for 4 h to remove solvent and gas from the pores of the nanotubes, which generated during curing of PDMS. 8.33 MPa of pressure were applied to some samples by hydraulic pressure. For pressurised samples, 500 µL of the PDMS mixture was added as the backing layer. All samples were cured in an oven at 140oC for 20 min and carefully peeled off (≈1 mm s-1) from the nanotubular template, except for sample 6 when rapid peeling (>3 mm s-1) was conducted. A control sample was made by casting 500 µL of 50 vol.% PDMS in toluene, on soda lime glass, followed by vacuum treatment for 4 h. It was cured at 140oC for 20 min and peeled carefully.The process is illustrated in Figure 1 and the nomenclature of samples is provided in Table 1.

Figure 1

Table 1

**2.4. Characterisation**

The morphology of titania nanotubes and PDMS analysis were carried out by field emission scanning electron microscopy (FESEM, JEOL JSM 6500F) in which elemental analysis was performed using energy dispersive X-ray spectroscopy (Oxford Inca system). To estimate the deviation of nanostructures morphology, 50 measurements were made on each sample via Microsoft Visio software using FESEM images. The cross section of titania nanotubes was inspected by peeling away some of the nanotube array using 3M scotch tape. PDMS samples were sputtered with thin layers of gold altering its surface electrical conductivity to enable FESEM characterisation. To ensure the silanisation of TiO2, Fourier transform infrared spectroscopy (FTIR) was deployed using Nicolet 380 FT-IR Spectrometer, Thermo Scientific. The contact angle of DI water on samples was measured using a drop shape analyser (DSA100). A droplet of deionized water (1 μL) was deposited on the sample surface using a syringe clamped on the column of the apparatus. The images were recorded with a charge-coupled device (CCD) camera and transmitted the information to the computer. DSA3 software was deployed to perform automatic calculation of contact angle for three measurements on each sample.

**3. Results and Discussion**

**3.1. Fabrication of surface modified titania nanotubes template**

The nanotubes length and diameter are essential as moulding template in which large nanotubes, up to 200 nm in diameter, can be produced by anodising 41. Electrolyte composition, applied voltage, and duration of anodising exerted significant influence on nanotubes length and diameter. Anodising of titanium foil in a glycerol-based solution (0.25 wt.% NH4F, 2 vol.% water in glycerol) at a cell voltage of 60 V for 6 h formed a nanotubes array with 140±20 nm diameter (Figure 2 a) and ≈1.5 µm length (Figure 2 b).

Insert Figure 2

Insert Figure 3

To facilitate separation of PDMS from the template, the nanotubular array was coated with fluoroalkyl chlorosilane to lower its surface energy and adhesion 42. Perfluorooctyltrichloro silane (FOTS) was adsorbed and hydrated onto the surface of nanotubes. Figure 3 shows the FTIR spectra of anodised TiO2 and silanised TiO2. Both samples exhibited characteristic Ti‒O and Ti‒O‒Ti peaks at 980 and 820 cm-1 43. Nevertheless, several changes in the FTIR peaks were observed after silanisation. The ‒OH and H‒O‒H peaks of anodised TiO2 disappeared due to the reaction with silane group and transformation into a hydrophobic surface. The attachment of fluoroalkyl silane groups was shown by Si‒O-C, C‒F, ‒CH2 and ‒CH3 peaks at 1145, 1250, ≈1600, and ≈2800 cm-1 44. Heat treatment at 140oC for 1 h ensured condensation and polymerisation of a self-assembled monolayer (SAM) coating on the surface. Helmy *et al.* 45 studied the effect of silane head groups on its attachment onto titania surface. SAMs from hydrolysis of FOTS were grafted significantly faster and denser than SAMs of triethoxy and dimethylchlorosilane as described by first order kinetics reaction. Although the combination of silane with various chain lengths was proven to yield the highest hydrophobic properties, the increment from 157.1o to 158.7o 46 was small. Here, only one silane (FOTS) was deployed to alter the surface of nanotubes providing a cost-effective route to functionalisation. Contact angle measurements in Figure 2 c showed that the wetting properties of titania nanotubes array switched from hydrophilic (≈0o) to hydrophobic (128.7±4.5o). The hydrophobic nature of nanotubes after treatment implied adequate attachment of fluoroalkyl chlorosilane. The procedure improved the wetting of toluene and PDMS during the critical toluene soaking step.

**3.2. Moulding of polydimethylsiloxane**

In a vertical cylinder, it has been postulated that time required for liquid to penetrate the pore are proportional to the viscosity of the liquid, which can be described by Lucas-Washburn equations as 47

(1)

(2)

(3)

where *t* is the time required to fill the tube, is the density difference between the penetrating liquid and the fluid (e.g., gas, other liquid) inside the porous medium, *µ* is dynamic viscosity of liquid, *z* is the depth of the tube, *r* is the radius of the tube,  is surface tension at liquid/air interface, **is contact angle of liquid. The addition of 60 wt.% toluene to the PDMS may reduce the viscosity of PDMS by 1 order magnitude 48. For silicon nanoholes, the filling depth was increased exponentially by >40 wt.% addition of toluene 48. Hence, the weight percentage of PDMS in toluene as well as the use of toluene soaking was studied in this work. The contact angle of the liquid was correlated with the wetting of the nanotubular template. The states of wetting on rough/porous surfaces, which in this case the surface was patterned by nanotubular array, can be described by two states, Wenzel 49 and Cassie-Baxter 50 state. The Wenzel model is used for fully wetted surface when the liquid contacts the surface area of the sample without any gap. The Cassie-Baxter model is deployed when a sessile drop sits on the top of the surface while substances (e.g., gas or immiscible liquid) entrapped beneath the drop 51. The Cassie-Baxter state only occurs when the value of the roughness is small enough compared to the size of the droplet.

Figure 4 a show a flat surface of PDMS which casted on soda lime glass without any treatment as sample 1 or the control sample. In sample 2, 3, and 4, the nanotubes were filled by toluene to the base of the tubes (Wenzel state) after soaking the nanotubular template in toluene for 10 minutes, which may ultimately enhance the surface wetting of the tube by PDMS 19. However, only microroughness observed on diluted PDMS sample 2 and 3 (Figure 4 b & c) indicating improper infiltration of PDMS. More defined micro-bumps were observed in sample 3 (as shown in Figure 4 c and d) indicating deeper infiltration of 33.33 vol.% PDMS. Without stirring, toluene in nanotubes (from toluene soaking process) may not mixed well with PDMS at 25oC and hold the PDMS mixture on top of the tube, in which external energy was needed to overcome the energy barrier of Cassie-Baxter state. Energy barrier increased as the diameter of tubes decreased which exerted significant effect on nanosized sample 52. Additional pressure on the sample improved the infiltration of PDMS. Figure 4 e & f showed bumps from PDMS sagging.

Insert Figure 4

**3.3. Hydraulic pressing**

After hydraulic pressing, the filling depth of PDMS was 221.64±52.76 nm in sample 4 (Figure 5 a), which far less than the length of nanotubes (≈1500 nm). Toluene left inside the nanotubes after toluene soaking impeded further infiltration of PDMS. The removal of toluene soaking in sample 5 may increase probability of trapped air inside nanotubes. Conversely, air is more compressible compared to liquid toluene during hydraulic pressing because air is a mixture of gases. For air, the pressure needed for filling the nanotubes was only 0.101 MPa, which can be calculated from the following equations.

(4)

(5)

*P* is the total pressure required, *Pair* is atmospheric pressure (0.101 MPa), *ρ* is the density of Slygard 182, which is 1049.97 kg m-3, and *g* is acceleration due to gravity (9.81 N kg-1), while *h* is the depth of the tube, 1.5 x 10-6 m. Since the pores are parallel, the pressure needed to fill all the pores was the same, at 0.101 MPa. However, 8.33 MPa of pressure was applied to overcome wall friction or debris that might have hampered penetration.

The filling depth was increased by this method (Figure 5 b & c). The height of worm-like PDMS were similar to the nanotube length in which the average length of worm-like PDMS were 1172±193 nm. The length frequency histograms of sample 4 and 5 were provided in Figure 6. Although the sample 5 had a significantly longer length (i.e., 1.17 micron) compared to sample 4 (221±52.76 nm), sample 5 showed a broad distribution with a deviation of 193 nm. PDMS may swell during the moulding process due to crosslinking of PDMS, yielding larger PDMS diameter (≈200 nm) compared to a titania mould (≈140 nm). This result was in accordance with the swelling study conducted by Vogt *et al.* where the PDMS was swollen by around 30% for a 10:1 wt.% base to curing agent ratio of PDMS 53.

The peeling rate was important in ensuring complete separation of PDMS and nanotubes. The peeling rate was ≈1 mm s-1 for sample 2, 3, 4, and 5. In Figure 5 e and f, rapid peeling rate (approx. >3 mm s-1) on sample 6 produced either flat PDMS or composites of PDMS and TiO2 nanotubes. The presence of TiO2 was shown by energy dispersive X-ray (EDX) spectroscopy, where the Ti atoms were absent in Figure 5d but evident in Figure 5f. The swollen PDMS in TiO2 nanotubes exhibited mechanical interlocking between PDMS nanostructures and inner wall of nanotubes, hampering the detachment of PDMS structures.

Insert Figure 5

Insert Figure 6

Contact angle measurement in Figure 7 a & b showed enhancement from 100.3±1.2o to 109.6±1.1o for flat and patterned PDMS sample 5, respectively due to the improvement in the surface roughness. When tilted up to around 45o, the 1 µL water droplet remained pinned to the surface rather than rolled over it, indicating a Wenzel state of wetting. It was expected that the Wenzel state would be the stable wetting state of rough PDMS in which the transition to Wenzel state occurred at a Wenzel limit of 110o 54. Ground collapse and collision of PDMS may diminished the roughness of sub-micron textured on PDMS sample. Ground collapse and collision of PDMS can be attributed to low modulus of soft PDMS (7.3 MPa for Sylgard 182) 37 and adhesive force to the ground (base of PDMS) 55. The critical aspect ratio of PDMS to avoid collapses are around 2, for tube diameter less than 500 nm 56. Such collapse was expected in this work considering high aspect ratio of PDMS (≈ 6). Stiffer materials such as epoxy and polyurethane can be used to prevent wall collapse. In worm-like PDMS, the structure can be recovered to the upright position by ultrasonication in a low surface energy solution 57. However, high aspect ratio worm-like PDMS are metastable and easily disturbed by moisture or electron beam during SEM characterisation 55.

Insert Figure 7

**3.4. Adhesive strength**

PDMS nanostructures can be used in climbing robots 58, microfluidics 59 and dry dermal patch 11 devices. Here, an adhesion test was conducted to stimulate potential application as dry adhesive. The experimental setup was outlined in Figure 8.

Insert Figure 8

A PDMS of 1 cm2 area was attached to the pin using carbon tape. The patterned surface of PDMS was then pressed on the planar glass surface with prior cleaning by acetone. The glass, PDMS, and pin setup was positioned vertically. The weighing scale was hung on the tip of the pin. Silica powder was added gradually until the adhesive was peeled from the glass. The peeling force was calculated from the final mass, allowing the shear adhesive strength to be obtained. The adhesive strength of samples 3, 4, 5 was 0.009, 0.312, 0.29 N cm-2 while all the other samples were not sufficiently robust to be measured. A significant increase was seen on the hydraulic pressed sample (sample 4 and 5) compared to non-pressed sample (sample 3).

The adhesion mechanism of dry adhesive has been extensively studied to explain the adhesion mechanism used in gecko attachment to solid surfaces. It was found that the dry adhesion mainly relies on van der Waals forces rather than electrostatic or suction 60. Small and dense nanostructures (i.e., 14400 setae per mm2 in gecko) resulted in high adhesion energy from van der Waals forces. Sample 4 and 5 in this work exhibited a dense worm-like PDMS (≈9 worm-like structures per µm2) compared to sample 3 with approximately 3 bumps per µm2 as determined by SEM. Although the aspect ratio in sample 5 was higher than in sample 4, the structures were collapsed to the ground hence the effective height was only the diameter of the worm-like structure producing less adhesive energy compared to sample 4. The 0.312 N cm2 adhesion strength of sample 4 was comparable to fibrin glue that is ≈0.2‒0.5 N cm2, which could be useful as a medical dry adhesive 61.

**4. Conclusions**

Titania nanotubular array, created by anodising titanium foil at 60 V using 0.25 wt.% NH4F and 2 vol.% water in glycerol as electrolyte, has been used as a template for PDMS (Sylgard 182) moulding. It was found that application of hydraulic pressure during casting PDMS into TiO2 nanotubes array mould has significantly improved the infiltration of the polymer into the pores of nanotubes. Although toluene soaking is commonly used in the moulding protocol to improve template wetting, it probably hindered full penetration of PDMS during hydraulic pressing. Complete penetration of PDMS mixture was observed on the hydraulically pressed sample without toluene soaking. The curing of PDMS mixture inside the pores of the tubular array followed by peeling of the polymer film has resulted in formation of worm-like structure with typical diameter ≈200 nm and length ≈1.17 µm. However, obtained high aspect ratio worm-like PDMS structures were deformed and collapsed due to low modulus of soft PDMS. The resulting structure was used as a dry adhesive, providing an adhesive strength up to 0.312 N cm2.

**Abbreviations**

PDMS, polydimethylsiloxane; TiO2, titanium dioxide; FOTS, 1H,1H,2H,2H‒perfluorooctyltrichloro silane.

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Table Caption

**Table 1.** Nomenclature of samples

Table 1

|  |  |  |
| --- | --- | --- |
| Sample | Name | Description |
| 1 | Flat | Control sample cast on soda lime glass |
| 2 | TS-0.5P-NP | Soaked with toluene, 50 vol.% PDMS, not pressed |
| 3 | TS-0.3P-NP | Soaked with toluene, 33.33 vol.% PDMS, not pressed |
| 4 | TS-0.3P-P | Soaked with toluene, 33.33 vol.% PDMS, hydraulically pressed |
| 5 | NTS-0.3P-P | Not soaked with toluene, 33.33 vol.% PDMS, hydraulically pressed |
| 6 | NTS-0.3P-P-R | Rapidly peeled sample, not soaked with toluene, 33.33 vol.% PDMS, hydraulically pressed |

Figure Captions

**Figure 1.** The illustration of PDMS moulding with titania nanotubes array.

**Figure 2.** SEM images of (a) top view of titania nanotubular array, (b) cross-section of nanotubes, and contact angle measurement of (c) uncoated nanotubular array, (d) coated nanotubular array

**Figure 3**. FTIR spectra of (i) the as-anodised TiO2 nanotubular array, (ii) the silanised TiO2

**Figure 4.** SEM images of PDMS, (a) control sample 1, (b) sample 2 TS-0.5P-NP, (c) sample 3 TS-0.3P-NP, (d) magnified image of sample 3, (e) sample 4 TS-0.3P-P, (f) magnified and 30o tilted image of sample 4

**Figure 5.** SEM images of PDMS, (a) 30o tilted images of sample 4 TS-0.3P-P (b) top view of sample 5 NTS-0.3P-P, (c) magnified images of sample 5 viewed on top, (d) EDX spectra of sample 5, (e) rapid peeling on sample 6 NTS-0.3P-P-R, (f) EDX spectra of sample 6

**Figure 6.** Length frequency histogram of (a) PDMS sample 4, (b) PDMS sample 5

**Figure 7.** Contact angle measurement of (a) flat PDMS sample, (b) sample 5 patterned PDMS

**Figure 8.** Simple experimental setup to measure dry adhesive strength, (a) the attachment of PDMS to pin holder, (b) the measurement of adhesive force on glass by weight addition

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**Figure 1**





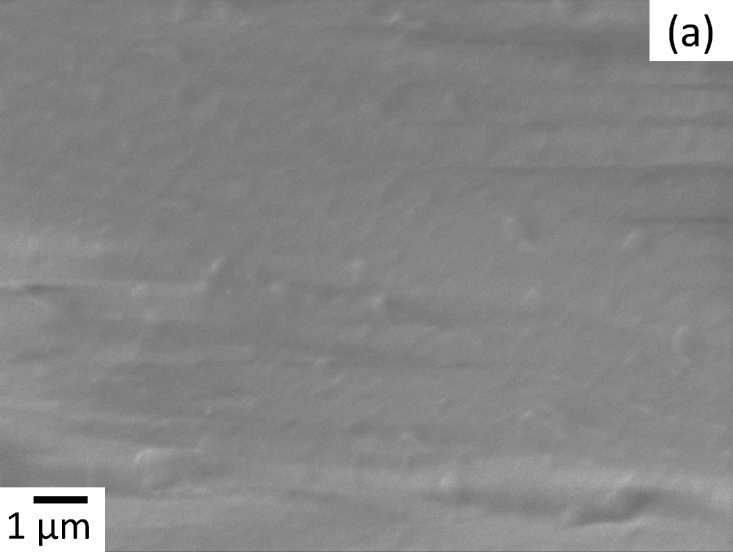
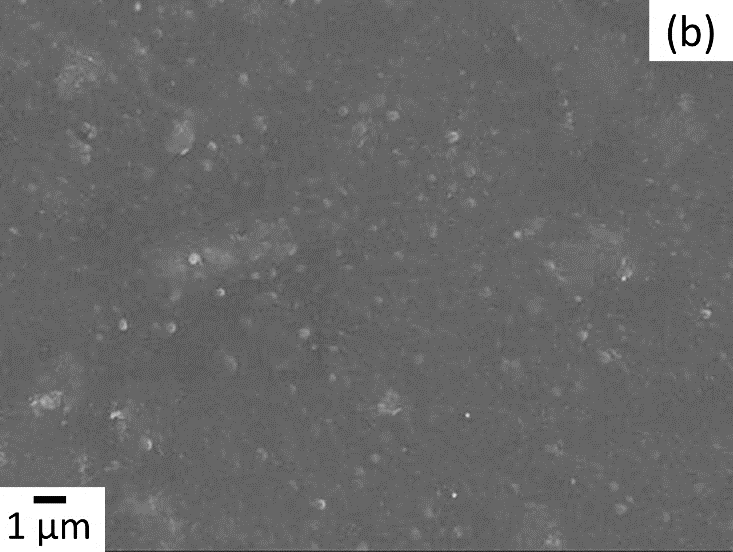


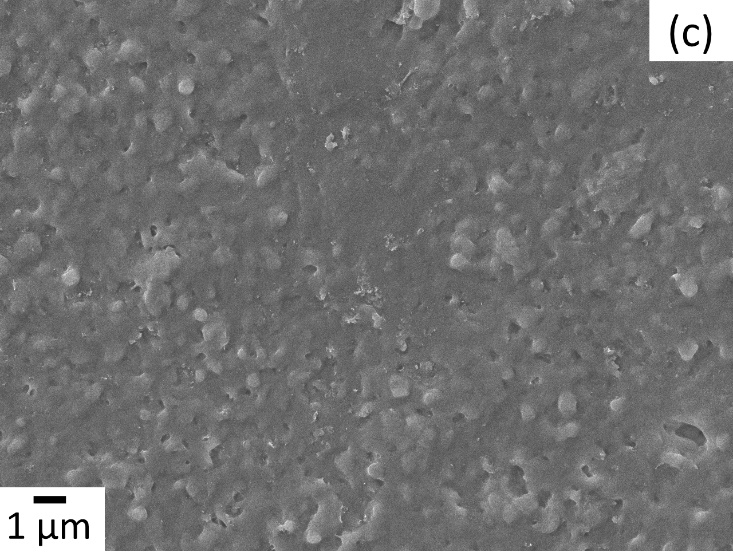


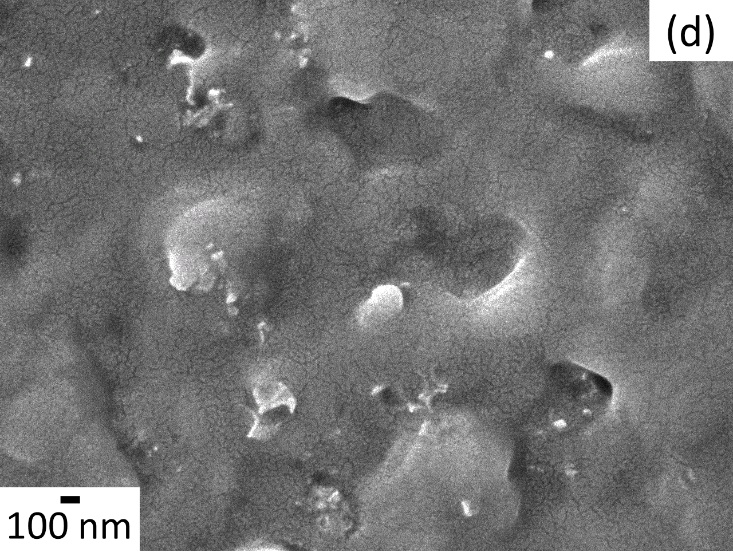
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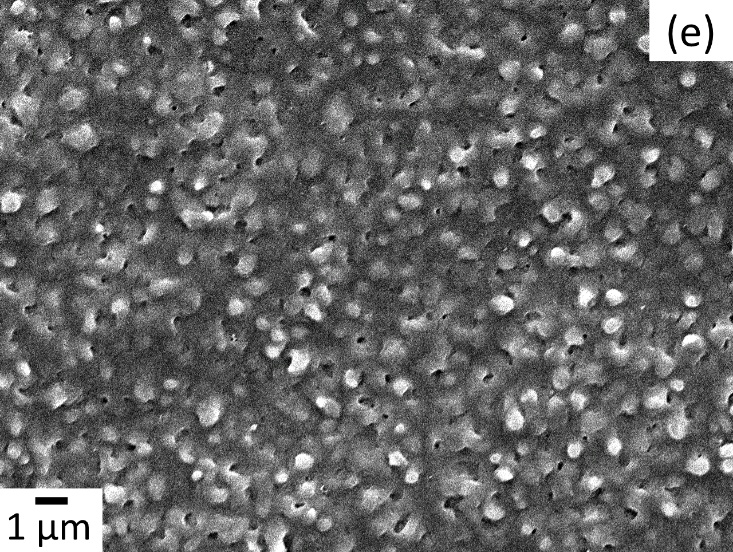
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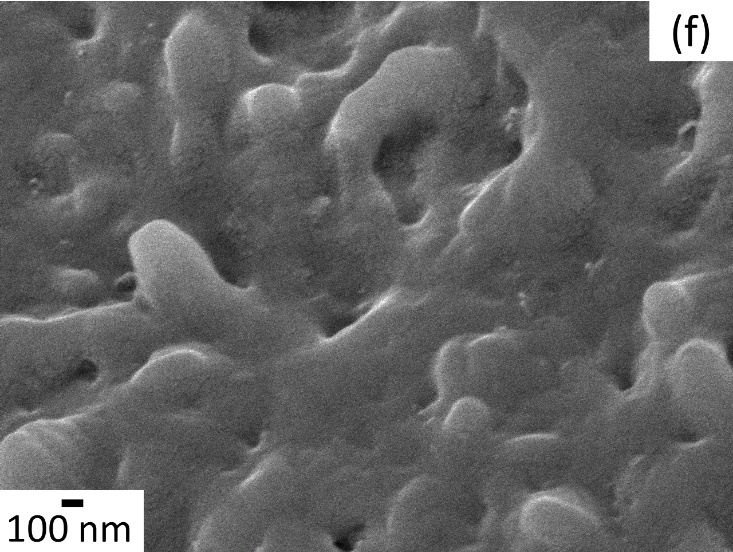
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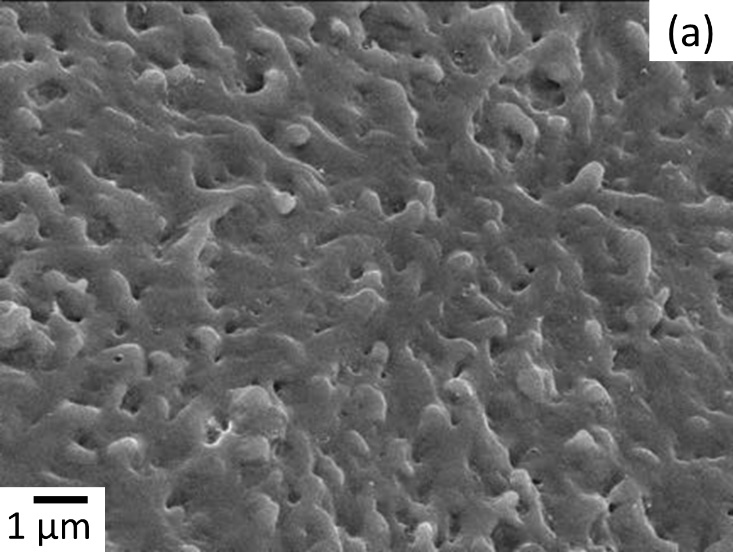


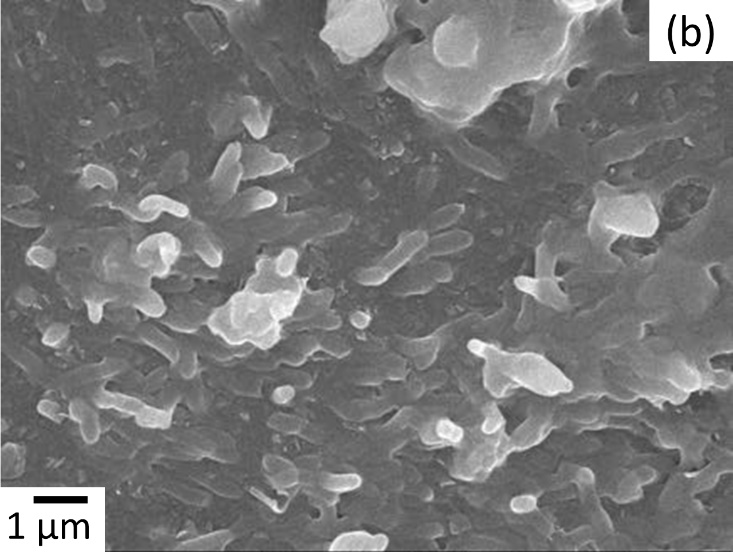


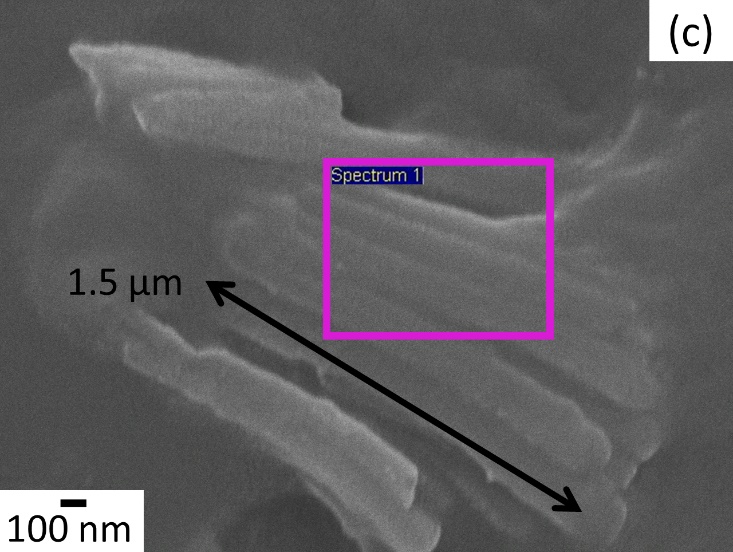
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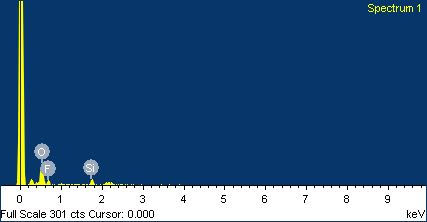
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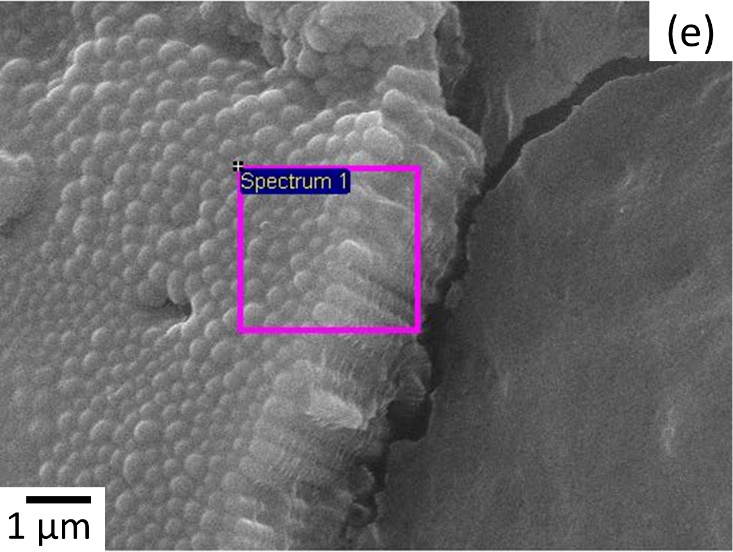


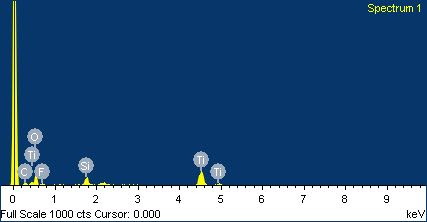






(d)





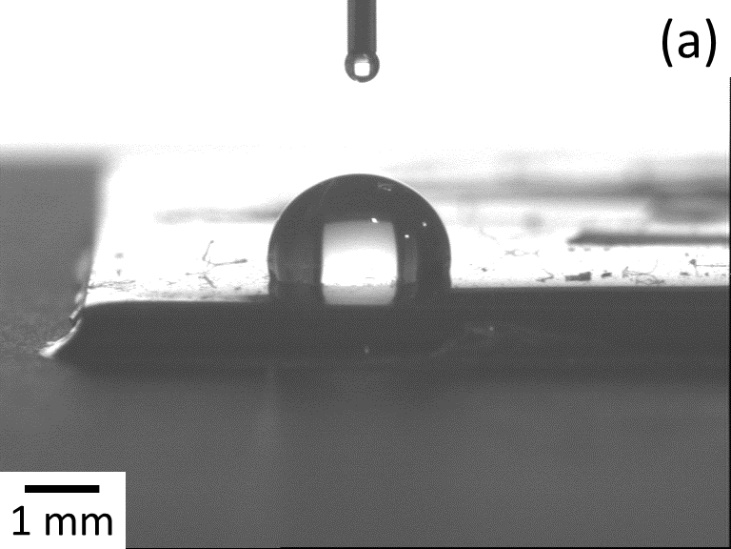
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**Figure 5**

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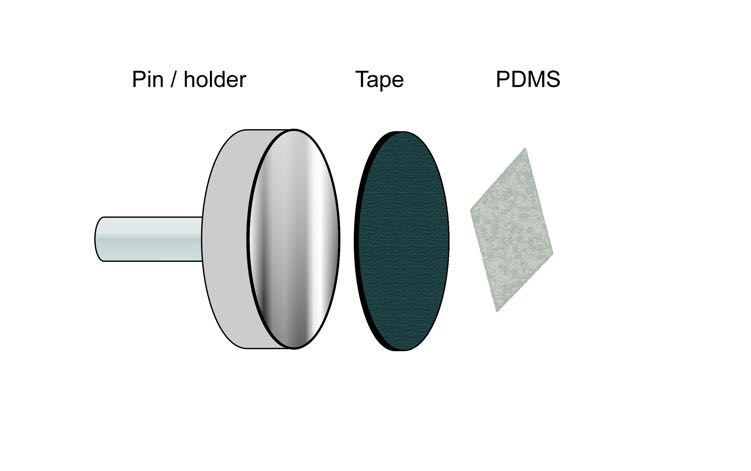
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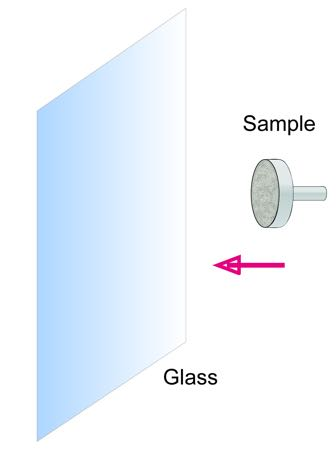
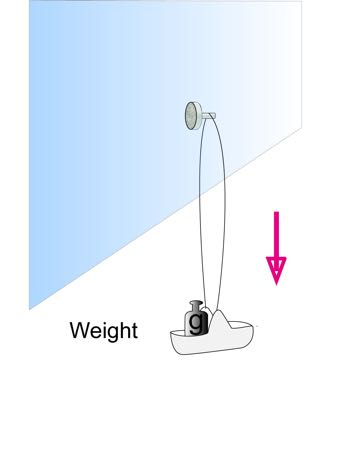




**Figure 7**



(a)



(b)

**Figure 8**