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Microstructured Nematic Liquid Crystalline Elastomer Surfaces with Switchable Wetting Properties

Zi Liang Wu, Axel Buguin, Hong Yang, Jean-Marie Taulemesse, Nicolas Le Moigne, Anne Bergeret, Xiaogong Wang, and Patrick Keller*

Inspired by the lotus leaf, scientists have developed many superhydrophobic surfaces, some of which show remarkable switching between hydrophobic and hydrophilic state under external stimuli. However, the switch usually relies on the change of chemical properties rather than on the modification of the topographic structure of the surface. In this paper, the roughness-change-related switchable wetting properties of microstructured responsive surfaces made of nematic liquid crystalline elastomers (LCEs) is reported. First, various carbonate LC monomers and side-on LCEs are synthesized with low nematic-to-isotropic transition temperature, T_{NI} . Then, LCEs prepared from 3'-vinylcarbonyloxypropyl 2,5-di(4'-octyloxybenzoyloxy)benzoate monomer, with T_{NI} of 76 °C and contraction of 34% are used to construct a surface covered with micropillar arrays by using a replica molding technique. The contraction of the micropillars induces a reversible roughness change of the microstructured surface. Water contact angle of this microstructured surface changed with temperature, indicating a successful approach at building a surface with switchable wetting properties.

1. Introduction

Nature has developed myriad functional materials, which serve as excellent paradigms for scientists to create new materials and devices.^[1,2] The functionalities of the natural materials are very

often associated with specific micro- and nanostructured surfaces,^[2] such as for the self-cleaning properties of the lotus leaves or the vivid colors of the butterfly wings.^[2a-c] For example, inspired by lotus leaf, many organic as well as inorganic structured surfaces with superhydrophobicity have been prepared.^[3,4] Furthermore, some of them, fabricated or coated with responsive polymers or inorganic nano-materials, showed switchable properties from hydrophobic to hydrophilic under suitable stimuli.^[4] However, the switching mainly originated from the change of chemical properties rather than from the change of topographic structure of the surface.

In this paper, we present our first attempt at making microstructured responsive surfaces for which large surface roughness changes could be induced by external stimuli. This approach might give us the opportunity to better control

the physical properties of the surfaces, such as the wetting and the adhesion.^[5]

Nematic liquid crystalline elastomers (LCEs) as a classic responsive soft material have been recognized as one of the best systems to develop artificial muscles and soft actuators.^[6] These materials combine the anisotropic structure of liquid crystals and the elasticity of the elastomers, exhibiting reversible, anisotropic shape change under external stimuli, such as light and temperature. The motor for the contraction is the reversible shape change of the macromolecular chain, from stretched to coiled, that occurs at the nematic-to-isotropic (N-I) phase transition.^[6b,7] A considerable amount of work aiming at preparing macroscopic LCEs and using them to create different kinds of soft actuators and sensors has been published.^[6,8] A seminal two-stage procedure to prepare monodomain nematic polysiloxane LCEs developed by Finkelmann et al.^[8a] has been used in hundreds of papers up to now. The orientation is achieved under strain by a simple extension before the material is fully crosslinked. Thus obtained LCEs usually exhibit large contractions up to ~300% and low nematic-to-isotropic phase transition temperatures, T_{NI} , due to the flexible siloxane backbone used. However, this method is limited to the preparation of macroscopic LCE samples. An alternative method is to synthesize LCEs by thermal or photo-initiated radical polymerization of liquid crystalline (LC) monomers.^[6b,8b] Magnetic field or surface effects are usually used to align the mesogens

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before the polymerization and crosslinking reaction. Compared with the method of two-step crosslinking, photo-initiated radical polymerization, which can be combined with microfluidics, soft-lithography, and photolithography, has overwhelming advantages to create micrometer-sized LCE actuators.

Several groups have developed new approaches to produce micron-size actuators based on LCEs.^[9–11] Ohm et al. used a microfluidic device coupled with UV lamp for photopolymerization to prepare LCE micro-particles, which changed their shape from sphere to ellipse when heated above the T_{NI} of the LCE.^[10] We applied a replica molding technique to produce surfaces covered with micrometer-sized pillars based on side-on and main-chain LCEs, which showed reversible shape changes.^[11] However, until now, the photopolymerization approach has produced thermotropic LCEs with relatively high T_{NI} , over 120°C,^[9–11] greatly limiting the applications of these micrometer-sized LCEs in microfluidics or responsive surfaces. Therefore, keeping in mind our objective to build microstructured responsive surfaces with adjustable wetting properties, we designed and synthesized LC monomers, which could give nematic LCEs with low T_{NI} and large contraction.

In this article, we report the synthesis of vinyl carbonate and allyl carbonate side-on nematic monomers and LCEs with low T_{NI} , up to 70 °C, and the construction of responsive structured surfaces based on these LCEs. Water contact angle measurements as a function of temperature have been performed for the microstructured surfaces, showing significant reversible changes above and below the T_{NI} of the LCEs.

2. Results and Discussion

Following the synthetic route of **Figure 1**, we successfully obtained the target carbonate LC monomers, as confirmed by ¹H NMR spectra and elemental analysis (Figure S1, Figure S2, and Table S1 in the Supporting Information). Although all the new synthesized monomers showed nematic phase in a certain temperature range, the properties of the monomers needed to be further investigated regarding whether they were photopolymerizable to prepare LCEs, as well as for the quality of the resultant LCEs, i.e., their T_{NI} and contraction. We first studied the effect of several chemical structure variations on the physical properties of the carbonate monomers, and then selected the most promising ones to prepare the responsive structured surfaces.

The monomers are noted as VC x ($n = \gamma$) or AC x ($n = \gamma$), where V or A indicates vinyl or allyl monomer, x and γ are the number of carbon in the end alkyl chains and in the linked alkyl chain (between the benzoate group and carbonate group), respectively.

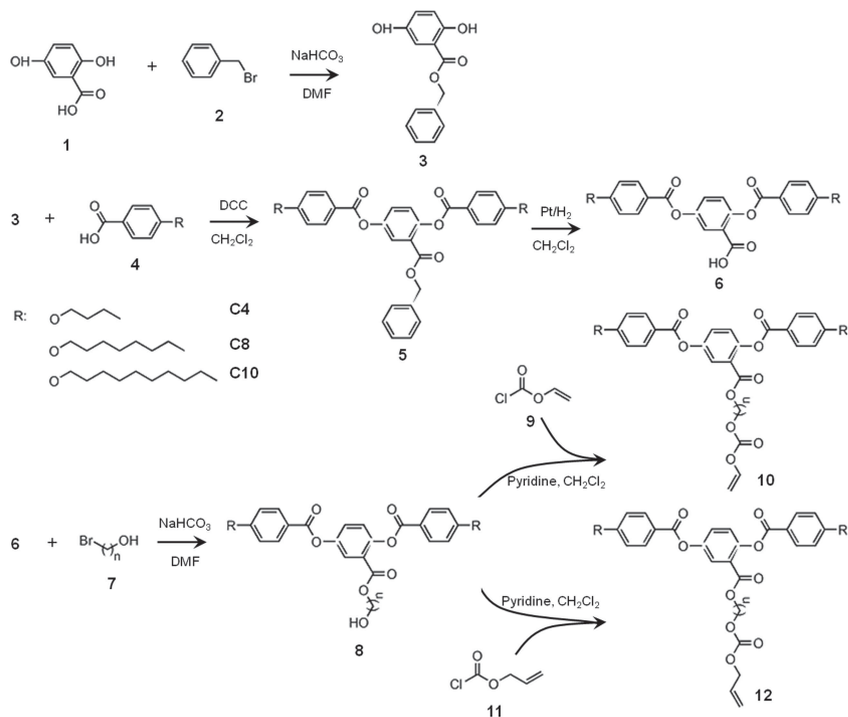


Figure 1. Synthesis of vinyl and allyl carbonate monomers containing di(4'-alkyloxybenzoyloxy)benzoate as mesogens.

Liquid crystalline polymers (LCPs) were also synthesized by photopolymerization of the monomers in absence of chemical crosslinker to compare their T_{NI} with those of LC monomers and LCEs. LCE and LCP samples are noted with “E” and “P” in front of the name of LC monomers, respectively.

2.1. Alkylene Carbonate Monomers and Their LCEs

2.1.1. Effect of the Carbonate Group on T_{NI} of LC Monomers

Using a typical side-on LC monomer 4'-acryloyloxybutyl 2,5-di(4'-butyloxybenzoyloxy)benzoate (A444, molecular structure shown in **Figure 2a**) as the starting point and the reference,^[8b] we introduced a carbonate, instead of an ester group to link the reactive C–C double bond and the mesogenic part, in an attempt at decreasing the T_{NI} of the LC monomers.

A vinyl carbonate monomer, 3''-vinylcarbonyloxypropyl 2,5-di(4'-butyloxybenzoyloxy)benzoate (VC4 ($n = 3$); molecular structure shown in **Figure 2b**) was synthesized, which has the same mesogen as the acrylate monomer A444. Even though the spacer of the carbonate monomer VC4 ($n = 3$) was somehow

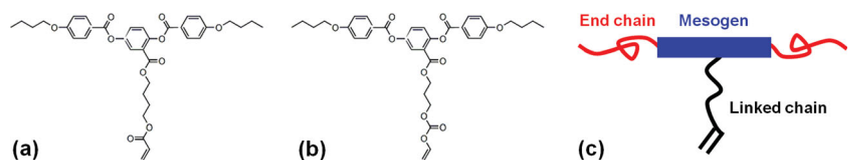


Figure 2. a, b) Molecular structure of A444 (a) and VC4 ($n = 3$) (b). c) Typical structure of synthesized monomers. After polymerization, they form side-on LCEs.

Table 1. Phase transition temperatures of the synthesized alkylene carbonate monomers.

Sample	Phase transition temperature ^{a)} [°C]	ΔH_{NI} ^{b)} [J/g]	T_{NI} ^{c)} [°C]
VC4 ($n = 2$)	K 80.5 N 99.3 I	1.74	100
AC4 ($n = 2$)	K 59.1 N 91.3 I	2.11	92.9
VC4 ($n = 3$)	K 63.8 N 89.2 I	1.53	91
AC4 ($n = 3$)	K 57.6 N 86.8 I	1.54	87.7
VC4 ($n = 6$)	K 45.6 N 79.1 I	1.13	79
AC4 ($n = 6$)	K 39.7 N 70.8 I	1.35	71.3
VC8 ($n = 3$)	K 59.9 N 83.1 I	1.59	83.2
AC8 ($n = 3$)	K 38 N 72.3 I	1.38	73.8
VC10 ($n = 3$)	K 55.7 N 80 I	1.76	80
AC10 ($n = 3$)	K 48.6 N 72.3 I	1.44	72.5

^{a)}Transition temperatures were obtained from the second heating scan of differential scanning calorimetry (DSC) thermograms of the monomers; heating rate: 5 °C/min. K: crystalline; N: nematic; I: isotropic; ^{b)}Enthalpy associated with the N-I phase transition of the monomers, ΔH_{NI} , was calculated from the corresponding peak of the DSC thermograms; ^{c)} T_{NI} was measured by observation of the N-I phase transition under polarizing optical microscope.

shorter than that of A444 ($n = 4$), the T_{NI} was 89.2 °C (Table 1), ~10 °C lower than that of A444. The allyl carbonate monomer, 3''-allylcarbonyloxypropyl 2,5-di(4'-butyloxybenzoyloxy)benzoate (AC4 ($n = 3$)) had a lower T_{NI} of 86.8 °C. This result indicated that the introduction of a carbonate group can effectively decrease the T_{NI} of the LC monomers, compared with acrylate LC monomer. The carbonate group introduces more decoupling between the rigid C–C double bond and the poly(methylene) spacer, bringing in more disorder, thus weakening the intermolecular interactions between mesogenic groups and reducing the stability of the nematic mesophase.

The vinyl carbonate LC monomers could be successfully photopolymerized to form LCPs, or copolymerized with chemical crosslinker to form LCE films, whereas the allyl ones could not, even after UV irradiation during extended time. All obtained LCE films showed reversible contraction at the N-I phase transition; as shown in Figure 3, E-VC4 ($n = 3$) with 10 mol% crosslinker had a contraction of 37%, comparable to 33% of E-A444 (Table 2). When the crosslinker concentration was increased to 20 mol%, the contraction of E-VC4 ($n = 3$) decreased to 28%.

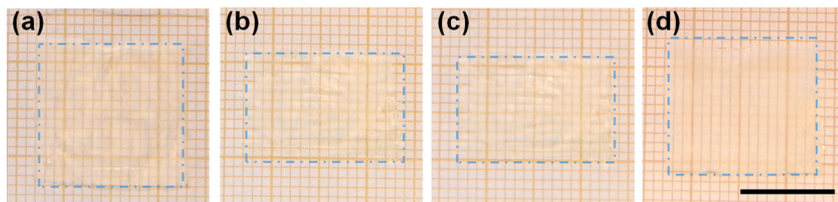


Figure 3. Images of LCE film (E-VC4 ($n = 3$), 10 mol% crosslinker) demonstrating the reversible contraction. The LCE film, floating on silicone oil, was heated to a) 50 °C, b) 95 °C, c) 110 °C, and then cooled back to d) 40 °C. The dotted line rectangles which outline the shape of the LCE film are drawn as a visual guide. Scale bar: 1 cm.

Table 2. Contraction and T_{NI} of LCEs and T_{NI} of LCPs.

Sample	Contraction	$T_{NI,LCE/LCP}$ ^{a)} [°C]	ΔH_{NI} [J/g]	$T_{NI,LCE/LCP} - T_{NI,M}$ ^{b)} [°C]
E-A444	33%	124.7	1.68	26.4
E-VC4 ($n = 3$)	37%	90.1	1.62	0.9
E-VC4 ($n = 2$)	41%	97.1	1.75	-2.2
E-VC4 ($n = 6$)	27%	85.7	1.44	6.6
E-VC8 ($n = 3$)	34%	76	1.21	-6.9
E-VC10 ($n = 3$)	16%	70.8	0.93	-9.2
P-A444	–	128.7	3.23	30.4
P-VC4 ($n = 3$)	–	94	2.2	4.8
P-VC8 ($n = 3$)	–	79.3	1.19	-3.6
P-VC10 ($n = 3$)	–	73.6	1.01	-6.4

^{a)} T_{NI} of LCEs or LCPs, noted as $T_{NI,LCE/LCP}$, was measured by DSC with heating rate of 5 °C/min; ^{b)} T_{NI} of the monomer was noted as $T_{NI,M}$ here to distinguish it from $T_{NI,LCE/LCP}$.

The effect of carbonate group on T_{NI} is even more evident in LCPs and LCEs. The T_{NI} of E-VC4 ($n = 3$) is 90.1 °C, just 0.9 °C higher than for the monomer. By contrast, T_{NI} of E-A444 is 124.7 °C, 26.4 °C higher than that of monomer (Table 2). A similar situation is found for the glass transition temperatures, T_g , of the LCPs, P-A444 and P-VC4 ($n = 3$), which have the same mesogen and similar spacer length. P-VC4 ($n = 3$) has a T_g of ~26 °C, much lower than that of P-A444, T_g of ~50 °C (Supporting Information Figure S3), indicating that P-VC4 ($n = 3$) possesses a relatively flexible polymer backbone.^[12,13]

2.1.2. Effect of the Spacer and of the Terminal Chain Lengths on the Stability of the Nematic Mesophase

The synthesized monomers share a common structure, a rod-like mesogen with two end grafted chains and a middle grafted chain which includes the reactive C–C double bond (Figure 2c). Either an increase in the flexible end chain length or the middle grafted chain length should bring more molecular disorder; thus, the intermolecular interaction should be weakened, decreasing the transition temperatures.

We first studied the effect of the spacer length, i.e., the number of CH₂ group, n , between the carbonate group and the benzoate group, on the T_{NI} of the LC monomers. Three kinds of vinyl carbonate monomers, VC4 ($n = 2, 3, 6$), were synthesized with the same mesogen part, as well as the same end chains grafted on the mesogen. As n increased, both the melting temperature, T_m , and T_{NI} decreased (Figure 4a and Table 1) (DSC thermograms for VC4($n = 2$) and VC4 ($n = 3$) show an extra exotherm peak at temperature around 20 and 10 °C, respectively. Those peaks are related to the partial recrystallization on heating of the compounds, which remain in the nematic supercooled phase for extended times). T_{NI} of VC4 decreased from 99.3 °C for $n = 2$ to 79.1 °C for $n = 6$. This effect was also found in the acrylate LC

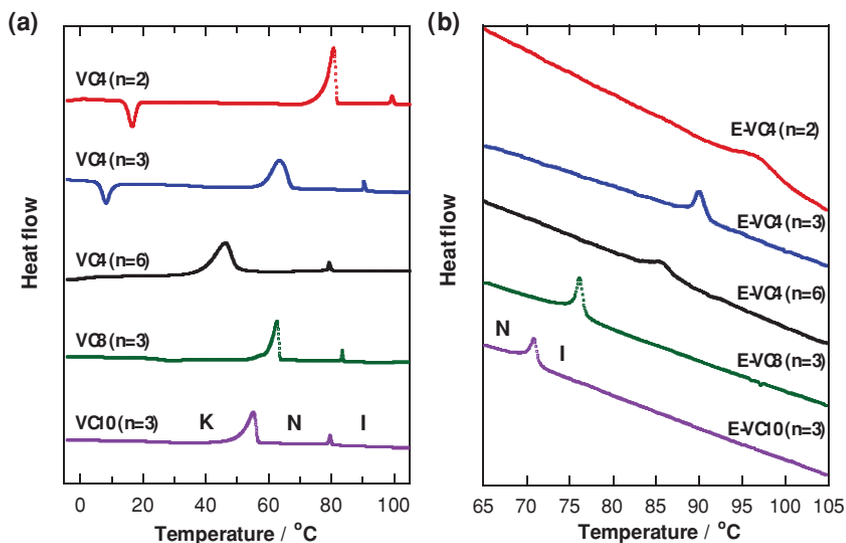


Figure 4. a) DSC thermograms of five vinyl carbonate LC monomers and b) their LCEs. Heating rate: 5 °C/min.

monomers.^[12c,14] The same tendency was observed in the corresponding LCEs (Figure 4b).

We should point out that a large n is detrimental to the coupling of the side-on mesogens and the backbone in the LCEs after polymerization. For side-on LCEs, the length of the spacer, which connects the backbone and the laterally grafted mesogens, is crucial for the extent of the reversible contraction of LCEs. The motor for contraction of LCEs is the conformational change occurring at the level of the polymer backbone at the N-I phase transition: due to the influence of the nematic order on the polymer chain, the backbone adopts an anisotropic conformation.^[7] If this coupling is weakened, the related LCE will have a smaller contraction. Indeed, the E-VC4 ($n = 6$) has a contraction of 27%, much lower than 41% for E-VC4 ($n = 2$) (Table 2). Considering the effect of n on T_{NI} and the contraction of the LCEs, we set $n = 3$ and studied the effect of the terminal chain length on the T_{NI} of the vinyl carbonate monomers.

We synthesized monomers with butyloxy, octyloxy, and decyloxy as the end chains. As expected, both T_m and T_{NI} decreased with the increase in the length of end chains (Figure 4a and Table 1). 3''-vinylcarbonyloxypropyl 2,5-di(4'-decyloxybenzoyloxy) benzoate, VC10 ($n = 3$), had a T_{NI} as low as 80 °C. Increase in the end chains also weakened the intermolecular interactions, and therefore decreased the T_{NI} of monomers, as well as T_{NI} of the LCEs (Figure 4b and Table 2).^[15]

Based on the above results, it was clear that (i) the introduction of carbonate group instead of ester group could effectively decrease T_{NI} of the related side-on LCEs, lower than 100 °C, because the C–O bond of carbonate group provided more free rotation; (ii) the spacer length increase weakened the coupling between mesogen and backbone and decreased the contraction of LCEs, although it could lower the T_{NI} of LCEs; (iii) lengthening of the chains which were grafted on two ends of the mesogen could lower the T_{NI} of LCEs, without decreasing too much the contraction of LCEs.

These side-on LCEs with considerable contraction and low T_{NI} are ideal materials for developing artificial muscles and soft

actuators. They are perfectly suitable for our goal of preparing responsive surfaces whose roughness could be adjusted by an external physical stimulus.

2.2. Responsive Surfaces with Adjustable Roughness

Microtextured surfaces were prepared by a technique combining the specific properties of a nematic mesophase (such as alignment by a magnetic field) with a soft-lithography approach to prepare the PDMS molds, following a protocol described in ref. [11] After removing the mold, well-arranged micropillars were left on the surface of LCEs, as shown in Figure 5. For the structured surface of E-VC4 ($n = 3$) (Figure 5a,b), the pillars had dimension of diameter = 20 μm , length = 75 μm , and spacing = 60 μm . Under external stimulus, i.e., raising the temperature above

T_{NI} , the pillars on the LCE surface contracted along their long axis and in a direction perpendicular to the surface. To better observe their contraction, the pillars were cut off from the surface with a razor blade. During the heating-cooling cycles, the pillars showed a reversible contraction, as shown in Figure 6. When the temperature was above T_{NI} of the E-VC4 ($n = 3$), 90.1 °C, the birefringence of the pillars disappeared, indicating that the contraction was related to the N-I phase transition of LCE pillars. A LCE film covered with well-arranged micropillars, when heated up to its N-I phase transition, should show

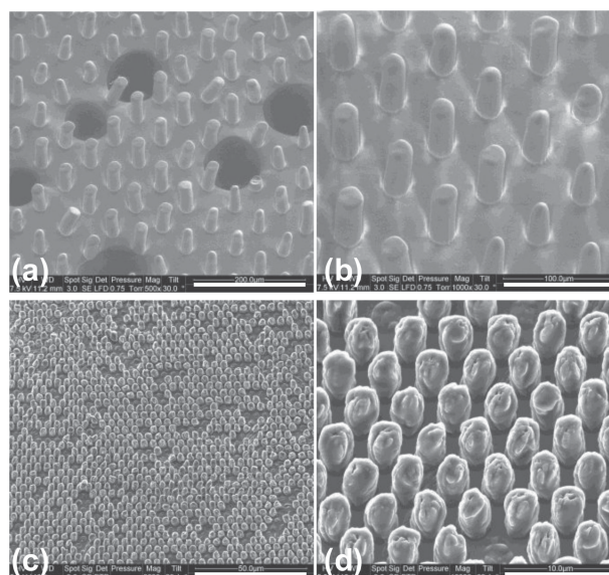


Figure 5. SEM images of structured surface with hexagonally-arranged micro-pillars. a,b) surface based on E-VC4 ($n = 3$), pillar size: diameter = 20 μm , length = 75 μm , spacing = 60 μm ; c,d) surface based on E-VC8 ($n = 3$), pillar size: diameter = 3 μm , length = 6 μm , spacing = 1.5 μm . Scale bar: (a) 200 μm ; (b) 100 μm ; (c) 50 μm ; (d) 10 μm .



Figure 6. Micrographs of pillars (E-VC4 ($n = 3$), 10% crosslinker) observed at different temperatures. The pillars were cut off from the structured surfaces and suspended in silicone oil to observe the shape change with temperature under polarizing optical microscope. The pillars were heated at a) 30 °C, b) 96 °C, and then cooled back to c) 48 °C. The dotted lines outline the shape of the pillars to guide the eye. Scale bar: 100 μm . A: Analyzer; P: Polarizer.

a change of surface roughness due to the contraction of the pillars. This responsive surface should enable us to control the physical properties of the surfaces, such as the wetting (Figure 7a).

We chose VC8 ($n = 3$) LC monomer to prepare the responsive surface, because the corresponding LCE had relatively low T_{NI} and large contraction, i.e., 76 °C and 34%, respectively. As shown in Figure 5c,d, microstructured surfaces were successfully obtained, which were covered by hexagonally-arranged pillars with dimensions of diameter = 3 μm , length = 6 μm , and spacing = 1.5 μm . The pillars were deformed to some extent because E-VC8 ($n = 3$) was softer than E-VC4 ($n = 3$) (much softer than E-A444);^[16] the deformations might have been generated when the mold was peeled off. At room temperature, this LCE with structured surface had a water contact angle, WCA, of 127°. Theoretically, according to the Cassie model, WCA (noted as θ^*) should be 125° , with $\cos \theta^* = -1 + \Phi_s (\cos \theta + 1)$,^[17] in which Φ_s was the area fraction of the asperities, θ was the WCA of the flat surface (here $\Phi_s = 0.4$, $\theta = 87^\circ$). On the other hand, the WCA for E-VC8 ($n = 3$) with a flat surface was 87°. When

the temperature was gradually raised to above the T_{NI} of the LCE (76 °C), the WCA of the flat surface just slightly decreased, reaching 84° at 85 °C. In contrast, for the first heating cycle, the WCA of the microstructured surface decreased from 127° at 40 °C to 86° at 81 °C (Figure 7b).

When the responsive surface was cooled back to room temperature, WCA was 114°, not fully back to ~127°. However, in the following thermal cycles, good reversibility was observed; WCA changed from ~110° to 86° when the structured surface was subjected to 35 °C and 80 °C, respectively (Figure 7c).

Much larger amplitude between the two states could be obtained by optimizing the size and spacing of the pillars.^[4g,18]

We should emphasize that the observed hydrophobicity change for the responsive surface made of LCE resulted from the roughness change of the surface owing to the contraction of the micropillars, which is different from the change of chemical properties used in the previously developed systems.^[4] Our work clearly proves that the physical properties of the surface such as wetting could be well tuned by reversibly changing the surface roughness. Responsive surfaces with biomimetic microstructures and smart response to external stimuli should provide a general means to design novel functional materials and devices.

3. Conclusions

By introducing carbonate group instead of the ester group, we have successfully synthesized side-on nematic LC monomers and LCEs with low T_{NI} , lower than 100 °C. Moreover, extending

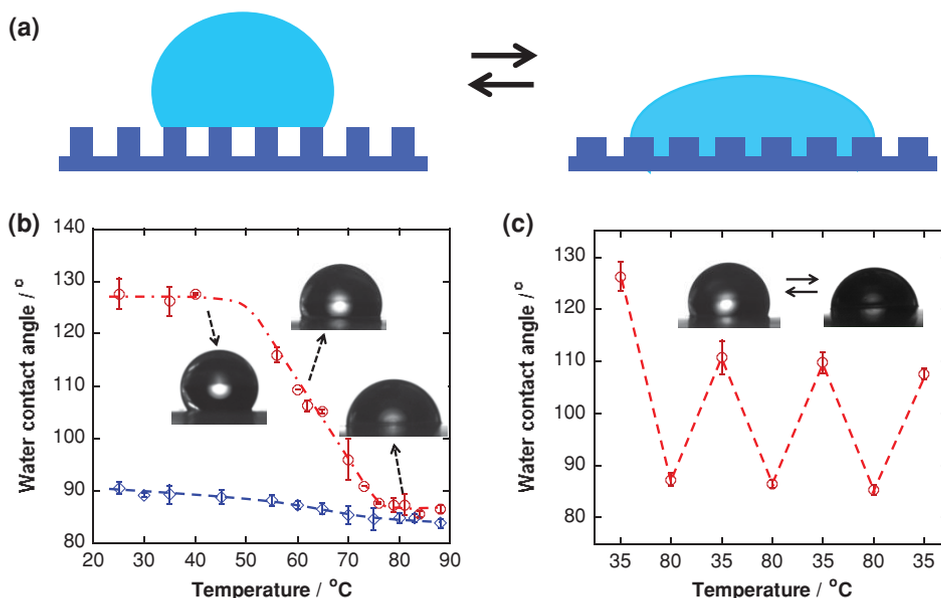


Figure 7. a) A scheme describing the control of the wetting properties by tuning the surface roughness. b) Hydrophobicity changes for the responsive surface, E-VC8 ($n = 3$) covered with micro-pillars, at different temperatures. structured surface; flat surface. c) Cyclic measurements of the WCA for a structured surface switching between 35 °C and 80 °C. Insets are representative profiles of a water droplet on the surfaces.

the length of the terminal chains, which were grafted on two ends of the mesogenic unit, can further lower the T_{NI} of the LCEs up to ~ 76 °C. These LCEs have been used to create micro-textured surfaces, covered by well-arranged micropillars, by using a replica molding technique. Each pillar behaved as a single micro-actuator and reversibly contracted along the axis of the cylinder, resulting in a responsive change of the surface roughness that gave us the opportunity to control the physical properties of the surfaces, and in particular the wetting properties. The water contact angle (WCA) of the structured surface could be switched between 110° and 86° , when the responsive surface underwent cyclic temperature changes between 35 and 80 °C. The functionality of the surfaces relied on both the microstructure and the responsiveness of the materials, which should serve as a paradigm to develop novel functional systems.

4. Experimental Section

Synthesis of Vinyl and Allyl Carbonate LC Monomers: The monomers have been prepared using the synthetic routes described in Figure 1. The synthesis of 3''-vinylcarboxypropyl 2,5-di(4'-butyloxybenzoyloxy)benzoate as a representative example is presented below.

The 2,5-di(4'-butyloxybenzoyloxy)benzoic acid (**6**, **C4**) was synthesized by a typical process, as described elsewhere.^[8b] We simply present here the reaction protocol. To a stirred solution of 2,5-dihydroxybenzoic acid (**1**) in dimethylformamide (DMF) was added solid NaHCO_3 . The mixture was heated and stirred at 70 °C for 1 h. Benzyl bromide (**2**) was added, and the mixture was heated for 7 h. After the purification process, benzyl 2,5-dihydroxybenzoate (**3**) was obtained. A solution of **3**, 4-butyloxybenzoic acid (**4**, **C4**), *N,N*-dicyclohexylcarbodiimide (DCC), and pyrrolidinopyridine in dichloromethane was stirred at room temperature for 12 h. The target product, benzyl 2,5-di(4'-butyloxybenzoyloxy)benzoate (**5**, **C4**), was then hydrogenated to produce 2,5-di(4'-butyloxybenzoyloxy)benzoic acid (**6**, **C4**).

(3''-hydroxypropyl) 2,5-di(4'-butyloxybenzoyloxy)benzoate (**8**, **C4**, $n = 3$) was synthesized as follows. To a solution of **6** ($R = \text{butyloxy}$; 2 g, 3.9 mmol) in DMF (15 mL) was added solid NaHCO_3 (1 g, 11.8 mmol). The mixture was heated and stirred at 70°C for 1 h. 3-bromo-1-propanol (**7**, $n = 3$) (0.7 mL, 7.74 mmol) was added, and the mixture was heated for 7 h at 70°C . The reaction mixture was cooled, diluted with water (100 mL), and extracted twice with 100 mL of a 50:50 (volume ratio) hexane/ethyl acetate mixture. The organic phases were washed with water (100 mL) and dried over Na_2SO_4 . After evaporation of the solvents, the crude product (3''-hydroxypropyl) 2,5-di(4'-butyloxybenzoyloxy)benzoate (**8**, **C4**, $n = 3$) was recrystallized twice from absolute ethanol (20 mL). ^1H NMR spectra of the compound is given in Figure S1 (Supporting Information).

3''-vinylcarboxypropyl 2,5-di(4'-butyloxybenzoyloxy)benzoate (**10**, **C4**, $n = 3$). (3''-hydroxypropyl) 2,5-di(4'-butyloxybenzoyloxy)benzoate (**8**, **C4**, $n = 3$) (2 g, 3.47 mmol) was dissolved in dichloromethane (40 mL). The solution in a flask was stirred and cooled in an ice bath. The air in the flask was removed by bubbling argon gas for 10 min. Under argon, to the solution were added successively pyridine (0.4 mL) and vinyl chloroformate (**9**) (0.4 mL, 4.38 mmol) via syringe.^[9] After 1 h, the ice bath was removed, and the reaction was continued for another 5 h. The reaction mixture was diluted with dichloromethane (200 mL), and extracted with water (200 mL). The organic phase was dried over Na_2SO_4 . After evaporation of the solvents, the crude product of 3''-vinylcarboxypropyl 2,5-di(4'-butyloxybenzoyloxy)benzoate (**10**, **C4**, $n = 3$) was recrystallized 3 times from absolute ethanol (15 mL). ^1H NMR spectra is presented in Supporting Information Figure S2a. Elemental analysis: calculated for $\text{C}_{35}\text{H}_{38}\text{O}_{11}$: C, 66.18; H, 5.99. Found: C, 66.29; H, 6.01.

(3''-allylcarboxypropyl) 2,5-di(4'-butyloxybenzoyloxy)benzoate (**12**, **C4**, $n = 3$) was synthesized by a similar procedure, using allyl chloroformate (**11**) instead of **10**. The related ^1H NMR spectra is shown in Supporting Information Figure S2b. Elemental analysis: calculated for $\text{C}_{36}\text{H}_{40}\text{O}_{11}$: C, 66.60; H, 6.17. Found: C, 66.60; H, 6.18.

Following this protocol, series of vinyl and allyl carbonate monomers were synthesized by varying the spacer length ($n = 2, 3$ and 6 in **10** and **12**) and the length of the terminal alkyl chains, which are grafted on the mesogens (R in **10** and **12**). Most of them exhibit a nematic mesophase with T_{NI} lower than 100 °C. The carbonate LC monomers are noted VC_x ($n = y$) or AC_x ($n = y$), where V or A indicates vinyl or allyl monomer, x and y are the number of carbon in the end alkyl chains and in the spacer (between the benzoate group and carbonate group), respectively.

Preparation of Monodomain LCE Films: Mixtures of carbonate LC monomer (90 mol%), divinyl sebacate (chemical crosslinker, 10 mol%), and 2-benzyl-2-(dimethylamino)-4-morpholinobutyrophenone (photoinitiator, 0.3 mol% of the total reactants) were used in the preparation of aligned LCE films. Mixtures, heated in their isotropic phase, were filled by capillarity in rubbed polyimide-coated glass cells of 20 μm gap (commercial Instec LC cells). The filled cells were slowly cooled down at -1 °C/min to the nematic phase of the mixture to achieve planar alignment. The cooling and the subsequent UV irradiation were carried under a stream of nitrogen (using a zip-lock bag),^[11] to ensure an inert atmosphere during the radical polymerization. After UV irradiation (365 nm, 30 mW/cm²) for 5 h, the free-standing, aligned LCE films were obtained by dissolving the glass windows of the cells with 40% aqueous hydrofluoric acid solution.^[20]

LCE samples are noted with "E" in front of the name of LC monomers. Liquid crystalline polymers (LCPs) were synthesized by photopolymerization of the monomers directly in the crucible used for differential scanning calorimetry (DSC) in absence of chemical crosslinker to compare their T_{NI} with those of LC monomers and LCEs. LCP samples are noted with "P" in front of the name of LC monomers.

Synthesis of LCE-Based Structured Surfaces: Responsive surfaces based on LCEs were prepared by photopolymerizing the same LC mixtures used above for the LCE films. Using a set-up described previously,^[11] LCE surfaces covered with micrometer-sized pillars of various shapes and sizes were obtained.

In practice, a small amount of the LC mixture including crosslinker and photoinitiator was heated to the isotropic phase on a microscope slide positioned atop a rare earth permanent magnet ($= 1$ T NdFeB rare earth magnet). The PDMS soft mold, prepared through a soft-lithography technique,^[11a,21] was then gently pressed down on the melted sample, which filled the inner structure of the mold. To keep an inert atmosphere for the radical polymerization, the whole setup was then sealed in a zip-lock bag under a nitrogen flow. The temperature was then slowly decreased (-1 °C/min) down to the temperature at which the mixture is in its nematic phase. During the cooling process, the applied magnetic field ensured the alignment of the nematic director parallel to the long axis of the pillars. The sample was then irradiated through the mold using a UV lamp for 5 h to promote the photopolymerization and crosslinking reaction of the monomer mixture. After cooling to room temperature, the PDMS mold was peeled off, leaving a thin glassy polymer film covered by a regular array of micropillars.

Characterization: ^1H NMR spectra were recorded on a Bruker 300 MHz spectrometer to confirm the molecular structures of the synthesized alcohols (**8**), and final monomers (**10** and **12**). All spectra were run in chloroform-d (CDCl_3) solution. The mesomorphic properties of the monomers, LCPs, and LCEs were studied by thermal optical polarizing microscopy (Leitz Ortholux microscope equipped with a Mettler FP82 hot stage) and differential scanning calorimetry (Perkin-Elmer DSC7). The heating and cooling rates were 5 °C/min. The contraction of macroscopic LCE films and micron-size pillars were measured by the dimension change of the samples in the alignment direction of side-on mesogens. The contraction was calculated by $(L - L_0)/L_0$, in which L and L_0 are the length of the samples at room temperature and above T_{NI} , respectively. The scanning electron microscopy (SEM) images were obtained at room temperature for samples tilted at a 30° angle, on a Quanta 200

FEG high-resolution environmental scanning electron microscope (FEI company). It might be noticed that the small pillars were sputter coated with carbon to avoid any degradation during observation. Water contact angles (WCAs) of the responsive surfaces at a given temperature were measured on a Digidrop contact angle meter (GBX, France). 4 μ L MilliQ water droplets were dropped carefully onto the surfaces of LCEs, which were pre-heated to the required temperature.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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