



AFRL-RX-WP-JA-2018-0139

LAYERED LIQUID CRYSTAL ELASTOMER ACTUATORS (POSTPRINT)

**Tyler Guin, Michael Settle, Benjamin A. Kowalski, Anesia D. Auguste, Richard Beblo,
Gregory Reich, and Timothy J. White**

AFRL/RX

**8 MARCH 2018
Interim Report**

**Distribution Statement A.
Approved for public release: distribution unlimited.**

© 2018 NATURE

(STINFO COPY)

**AIR FORCE RESEARCH LABORATORY
MATERIALS AND MANUFACTURING DIRECTORATE
WRIGHT-PATTERSON AIR FORCE BASE, OH 45433-7750
AIR FORCE MATERIEL COMMAND
UNITED STATES AIR FORCE**

REPORT DOCUMENTATION PAGE

Form Approved
OMB No. 0704-0188

The public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Department of Defense, Washington Headquarters Services, Directorate for Information Operations and Reports (0704-0188), 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302. Respondents should be aware that notwithstanding any other provision of law, no person shall be subject to any penalty for failing to comply with a collection of information if it does not display a currently valid OMB control number. **PLEASE DO NOT RETURN YOUR FORM TO THE ABOVE ADDRESS.**

1. REPORT DATE (DD-MM-YY) 8 March 2018			2. REPORT TYPE Interim		3. DATES COVERED (From - To) 23 August 2016 – 8 February 2018	
4. TITLE AND SUBTITLE Layered Liquid Crystal Elastomer Actuators (Postprint)			5a. CONTRACT NUMBER FA8650-16-F-5419			
			5b. GRANT NUMBER			
			5c. PROGRAM ELEMENT NUMBER 62102F			
6. AUTHOR(S) Tyler Guin, Michael Settle, Benjamin A. Kowalski, Anesia D. Auguste, Richard Beblo, Gregory Reich, and Timothy J. White - AFRL/RX			5d. PROJECT NUMBER 4348			
			5e. TASK NUMBER			
			5f. WORK UNIT NUMBER X14W			
7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) AFRL/RX Wright-Patterson AFB Dayton, OH 45433			8. PERFORMING ORGANIZATION REPORT NUMBER			
9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES) Air Force Research Laboratory Materials and Manufacturing Directorate Wright-Patterson Air Force Base, OH 45433-7750 Air Force Materiel Command United States Air Force			10. SPONSORING/MONITORING AGENCY ACRONYM(S) AFRL/RXAP			
			11. SPONSORING/MONITORING AGENCY REPORT NUMBER(S) AFRL-RX-WP-JA-2018-0139			
12. DISTRIBUTION/AVAILABILITY STATEMENT Distribution Statement A. Approved for public release: distribution unlimited.						
13. SUPPLEMENTARY NOTES PA Case Number: 88ABW-2018-1160; Clearance Date: 23 Feb 2018. This document contains color. Journal article published in Nature Communications, Vol. 9, 28 June 2018. © 2018 Nature. The U.S. Government is joint author of the work and has the right to use, modify, reproduce, release, perform, display, or disclose the work. The final publication is available at www.nature.com/naturecommunications DOI: 10.1038/s41467-018-04911-4						
14. ABSTRACT (Maximum 200 words) Liquid crystalline elastomers (LCEs) are soft, anisotropic materials that exhibit large shape transformations when subjected to various stimuli. Here we demonstrate a facile approach to enhance the out-of-plane work capacity of these materials by an order of magnitude, to nearly 20 J/kg. The enhancement in force output is enabled by the development of a room temperature polymerizable composition used both to prepare individual films, organized via directed self-assembly to retain arrays of topological defect profiles, as well as act as an adhesive to combine the LCE layers. The material actuator is shown to displace a load >2500× heavier than its own weight nearly 0.5 mm.						
15. SUBJECT TERMS Liquid crystalline elastomers; shape memory alloys (SMAs); musculo-skeletal systems; mechanical anisotropy						
16. SECURITY CLASSIFICATION OF:			17. LIMITATION OF ABSTRACT: SAR	18. NUMBER OF PAGES 9	19a. NAME OF RESPONSIBLE PERSON (Monitor) Timothy White 19b. TELEPHONE NUMBER (Include Area Code) (937) 255-9551	
a. REPORT Unclassified	b. ABSTRACT Unclassified	c. THIS PAGE Unclassified				

ARTICLE

DOI: 10.1038/s41467-018-04911-4

OPEN

Layered liquid crystal elastomer actuators

Tyler Guin^{1,2}, Michael J. Settle^{3,4}, Benjamin A. Kowalski^{1,2}, Anesia D. Auguste¹, Richard V. Beblo^{3,4}, Gregory W. Reich³ & Timothy J. White¹

Liquid crystalline elastomers (LCEs) are soft, anisotropic materials that exhibit large shape transformations when subjected to various stimuli. Here we demonstrate a facile approach to enhance the out-of-plane work capacity of these materials by an order of magnitude, to nearly 20 J/kg. The enhancement in force output is enabled by the development of a room temperature polymerizable composition used both to prepare individual films, organized via directed self-assembly to retain arrays of topological defect profiles, as well as act as an adhesive to combine the LCE layers. The material actuator is shown to displace a load >2500× heavier than its own weight nearly 0.5 mm.

¹Air Force Research Laboratory, Materials and Manufacturing Directorate, Wright-Patterson Air Force Base, OH 45433, USA. ²Azimuth Corporation, 4027 Colonel Glenn Hwy, Beavercreek, OH 45431, USA. ³Air Force Research Laboratory, Aerospace Systems Directorate, Wright-Patterson Air Force Base, OH 45433, USA. ⁴University of Dayton Research Institute, 1700 S Patterson Blvd, Dayton, OH 45469, USA. Correspondence and requests for materials should be addressed to T.J.W. (email: timothy.white.24@us.af.mil)

Responsive materials are currently subject to intense research, motivated in part by end-use applications in robotics¹. An emerging research frontier is materials systems that inherently emulate the motion, dexterity, and force output of natural musculo-skeletal systems². An increasingly common approach is to locally vary the organization or other materials properties such that “the material is the machine³.” Realization of real world devices will require further innovation and development of both hard and soft materials.

In some implementations, shape reconfigurability will be an important aspect of robotic control. Stimuli-responsive shape change of monolithic elements is exhibited by a range of material platforms, including shape memory alloys (SMAs)⁴. SMAs achieve large force output but limited deformation, and are found in end-use applications in medicine, automobiles, and aerospace⁵. Recent explorations focus on soft materials in which the mechanical response can be localized and potentially programmed, at the expense of output force⁶.

Natural musculo-skeletal systems employ anisotropy to optimize function, as well as grade the interfacial interaction of stiff and soft elements. Liquid crystalline elastomers⁷ have been studied for nearly 50 years, traceable to original predictions from de Gennes⁸. Uniquely, the molecular orientation of these materials can be programmed pixel-wise with micron-scale resolution. Robust and high-throughput patterning is enabled by exploiting directed self-assembly (both spatial and hierarchical) onto a patterned template surface⁹. The molecular orientation governs the anisotropy of macroscopic mechanical response, and monolithic elements composed of these materials can be permanently programmed to exhibit reversible, stimuli-responsive shape transformations. A wide range of shapes can be realized such as origami folds^{9–11}, arrays of cones¹², or arbitrary curvatures, such as paraboloids¹³. Notably, these materials are continuous in composition and absent of multimaterial interfaces. Mechanical responses in these materials can be triggered by exposure to heat^{14, 15}, light^{16, 17}, electrical fields^{18, 19}.

The tremendous shape transformation of LCEs can create useful work. LCEs with uniform orientation (via mechanical stretching) exert muscle-like contractile force generating strains of up to 400%²⁰. A number of recent reports detail a comparatively distinctive approach to generating force. LCE sheets with spatially patterned orientation can act as out-of-plane lifters, using shape change to generate considerable work over a large stroke, with a work capacity of as much as 2.6 J/kg⁹ (from a soft material of 50 μm thickness). The extraordinary work capacity of these materials is attributable to the fundamentals of the shape transformation. The spatial variation in the director profile dictates that the material must emanate into a third dimension, via stretch. It is predicted in ref. ²¹ that force outputs should correlate to increasing the film thickness. However, the achievable thickness of LCEs prepared by surface anchoring is limited. For cell thicknesses exceeding roughly 50 μm , the patterned alignment surface can no longer effectively prescribe alignment through the entire cell, due to finite anchoring energies of surface interactions²².

Here we present a novel method to create arbitrarily thick LCE films that are continuous in composition and maintain complex director orientations, prescribed into the material via directed self-assembly by photoalignment. Our approach is to laminate as many as six LCE films bonded with interfacial layers of the same composition. Critically, these adhesive layers take on the residual orientation of the adjacent LCE layers. To enable this process, we develop a LCE composition with a room temperature nematic phase. The laminates maintain the shape transformation of single layer LCE films when heated. However, the laminated films exhibit extraordinary lifting forces to nearly 20 J/kg (as much as

2500 \times the weight of the mechanical system). We illustrate that these materials are now capable of withstanding significant positive pressure, which could open up end uses in aerospace and other application domains.

Results and discussion

Materials preparation and characterization. The LCE films examined here were formulated by mixing mesogenic diacrylates (RM82 and RM257) with a dithiol chain-transfer agent (Fig. 1a). As detailed in ²³, the dithiol additive reduces the crosslink density of the polyacrylate via chain transfer (primary) and chain extension (secondary). The concentration of RM82 and RM257 in the mixture was selected to suppress the nematic-crystallization phase transition²⁴, producing a supercooled mixture which was meta-stable (>1 h) to $-20\text{ }^\circ\text{C}$. The broad phase range enabled processing and photopolymerization to occur at room temperature. The LCE films are optically clear (Fig. 1b). The glass transition temperature (T_g) of the LCEs prepared from this composition was $26\text{ }^\circ\text{C}$, (Supplementary Figure 1) similar to prior reports¹³. Local organization of the monomeric mixture was directed by photoalignment cells (PAAD-22, BEAM Co.)²⁵. Here we impose various director profiles into the material to localize the orientation of the liquid crystalline monomers into topological defects, which are subsequently retained after photopolymerization.

Dating to de Gennes' original predictions⁸, LCEs have been discussed as synthetic candidates to emulate the tremendous force output of muscle fiber¹⁵. Examinations of LCEs in a planar (i.e., monodomain, single crystal) orientation have reported force outputs of 250 kPa¹⁵. Recent examinations of LCEs imprinted with complex topologies, such as the examination here, have reported specific work capacities of 2.6 J/kg⁹. The focus of this work is to explore and demonstrate approaches and considerations to substantially increase the work capacity of LCEs while maintaining (or extending) the large stroke.

An approach to increase the work capacity of these material systems is to increase the thickness of the LCEs²⁶. However, the physics of surface anchoring²⁷ and the anchoring strength of the photoalignment layer²⁸ limit the maximum thickness for retention of surface-induced director profiles to $\sim 50\text{ }\mu\text{m}$ ²². To circumvent, we report here a newly developed method to laminate LCE films (illustrated in Supplementary Figure 2). After polymerization and under crossed polarizers, two LCE films of identical composition and director profile were registered. Leveraging the room temperature nematic phase of the monomer composition employed to prepare the films, the samples were coated with a thin layer of the monomer mixture (same composition as above). After registry, the two LCEs were sandwiched together and gently clamped. The samples and monomer mixture were briefly heated and slowly cooled, to allow the adhesive layer to take on the residual surface alignment of the films. The films were cured by UV light exposure. From this two layer laminate, additional layers can be added to realize up to 300 μm thick LCE laminates examined here. Due to the consistency in the materials chemistry acting as both the LCE layer and adhesive, we observed no delamination in any of our experiments.

The orientation of the laminated films was confirmed with polarized microscopy²⁹ to confirm the registry of the LCE layers and that the adhesive layers are taking on the order of the LCE surfaces (Supplementary Figure 3). The thermally induced contraction (Fig. 1c) of the uniaxially aligned LCEs is nearly identical for a single layer, double layer, and four layer laminates. The contraction measured in Fig. 1c was determined from dimensional changes observed in the LCE films upon heating.

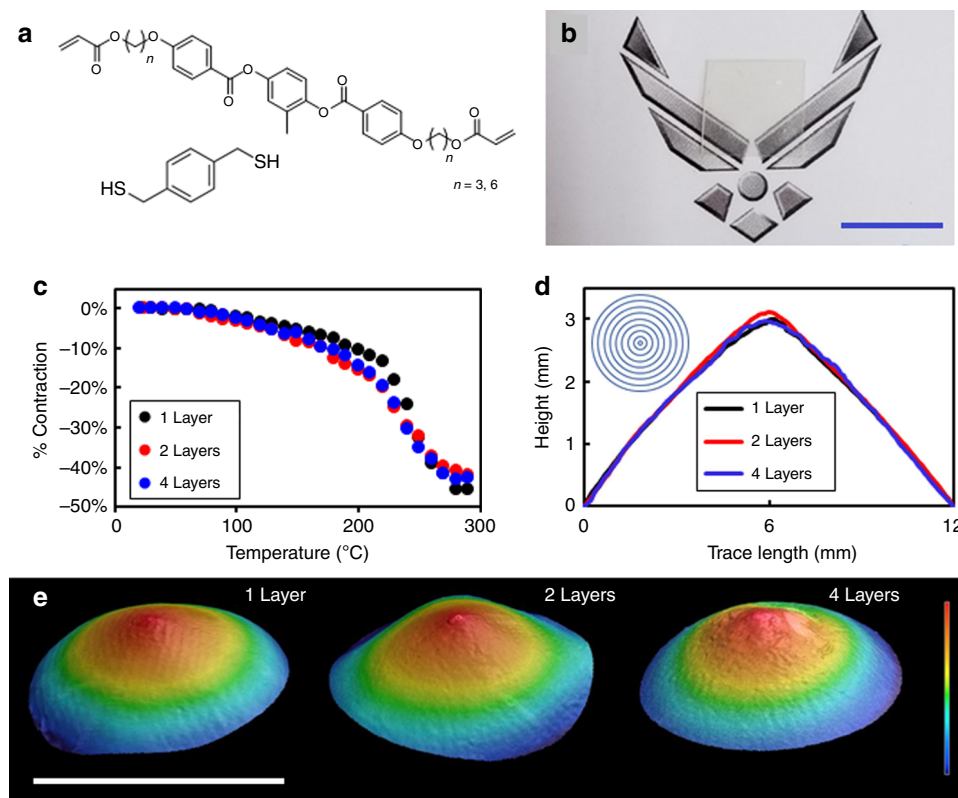


Fig. 1 Unloaded deformation of LCE laminates. **a** Chemical structures of: (1) the liquid crystal monomers (RM257, RM82) and (2) the dithiol additive. A mixture of 7:2 wt:wt of RM257:RM82 formed the basis of the liquid crystal elastomers (LCEs). **b** A 4 layer LCE laminate is optically clear. Scale bar is 1 cm. **c** The deformation of one, two, and four layer LCE laminates was measured in homogeneously aligned LCE films. 50% strain is observed in all three samples. **d** The height profiles for the single layer, double layer, and four layer LCE laminates are nearly identical over the 12 mm diameter of the films. Horizontal scale bar is 12 mm, vertical scale bar is 3.4 mm. **e** Height profiles of LCE laminates in which the individual layers were subject to directed self-assembly to organize into radial +1 topological defects. The LCE laminates were heated to 140 °C and imaged with an optical profilometer to quantify the deformation to thermal stimuli

Deformation of laminated LCEs. When heated, nematic LCEs reversibly contract along the liquid crystalline director and expand in the orthogonal directions³⁰. This anisotropic contraction, when subject to spatial variation dictated by the directed self-assembly of localized surface alignment, can result in dramatic out-of-plane shape deformation^{14, 31}. Here we employ a well-understood and predictable director profile, the azimuthal +1 topological defect. The director profile is inset in Fig. 1d, where the mesogens organize in concentric rings around a central region (point defect). This pattern was predicted³¹ and experimentally confirmed¹⁴ to deform into a cone upon heating.

The deformation of single layer, double layer, and four layer LCE laminates was quantified by structured-illumination optical profilometry (Keyence VR-3000). The LCE film and laminates actuate into cones upon heating (Fig. 1d). The amplitude of the peak height (3.4 mm, ~70× the film thickness) and the angle of the cone tip are nearly identical among films of 50 μm (single layer), 102 μm (double layer), and 210 μm (four layer) thickness (Fig. 1e). The increased thickness of the laminates does not diminish the shape transformation. The insensitivity of the shape-morphing to film thickness is in agreement with a prediction of³² that the deformation of an LCE sheet into a cone should be largely independent of sheet thickness, except for slight deviation near the tip. The agreement evident in Fig. 1d, e are strong, indirect evidence that each LCE layer as well as the adhesive interfaces are cooperatively deforming.

Using the force. In a previous report, a 2 × 2 array of +1 topological defects in a 50 μm thick LCE film was shown to lift up to 150× its weight with a stroke of 1 mm⁹. The stroke/force output of the single layer LCE films translates to a specific work capacity of 2.6 J/kg. Informed by the results in Fig. 1, we extend this examination to characterize and assess the potential actuation force of the LCE laminates. In Fig. 2, 1 × 1 cm LCE films patterned into a 2 × 2 arrays of radial +1 topological defects (director profile illustrated in Fig. 2a) were prepared. Single, double, and four layer LCE laminates were heated with a resistive heating element. A piece of glass was placed on top of the films, which was loaded with weight. As illustrated in the representative photograph in Fig. 2b, the deformation of the films is observable under load (28.7 g in Fig. 2b, see also Supplementary Movie 1). The deformation under load for one, two, and four layer LCE laminates are presented in Fig. 2c. Similar to the unloaded case in Fig. 1e, the stroke of the LCEs under load is relatively unaffected with increasing thickness. However, the increase in thickness increases the output force dramatically. The four layer LCE laminate (210 μm thick) produces 280 mN of force at a stroke of 1.6 mm. As illustrated in Fig. 2d, the specific work can reach nearly 19 J/kg. The four layer LCE laminate with the director configuration described in Fig. 2a can lift over 1100 times the weight of the film itself, a 100× improvement in specific work when compared to a single layer LCE.

When selecting actuators³³ both force output and stroke length are important considerations. So-called piezoelectric stacks can be

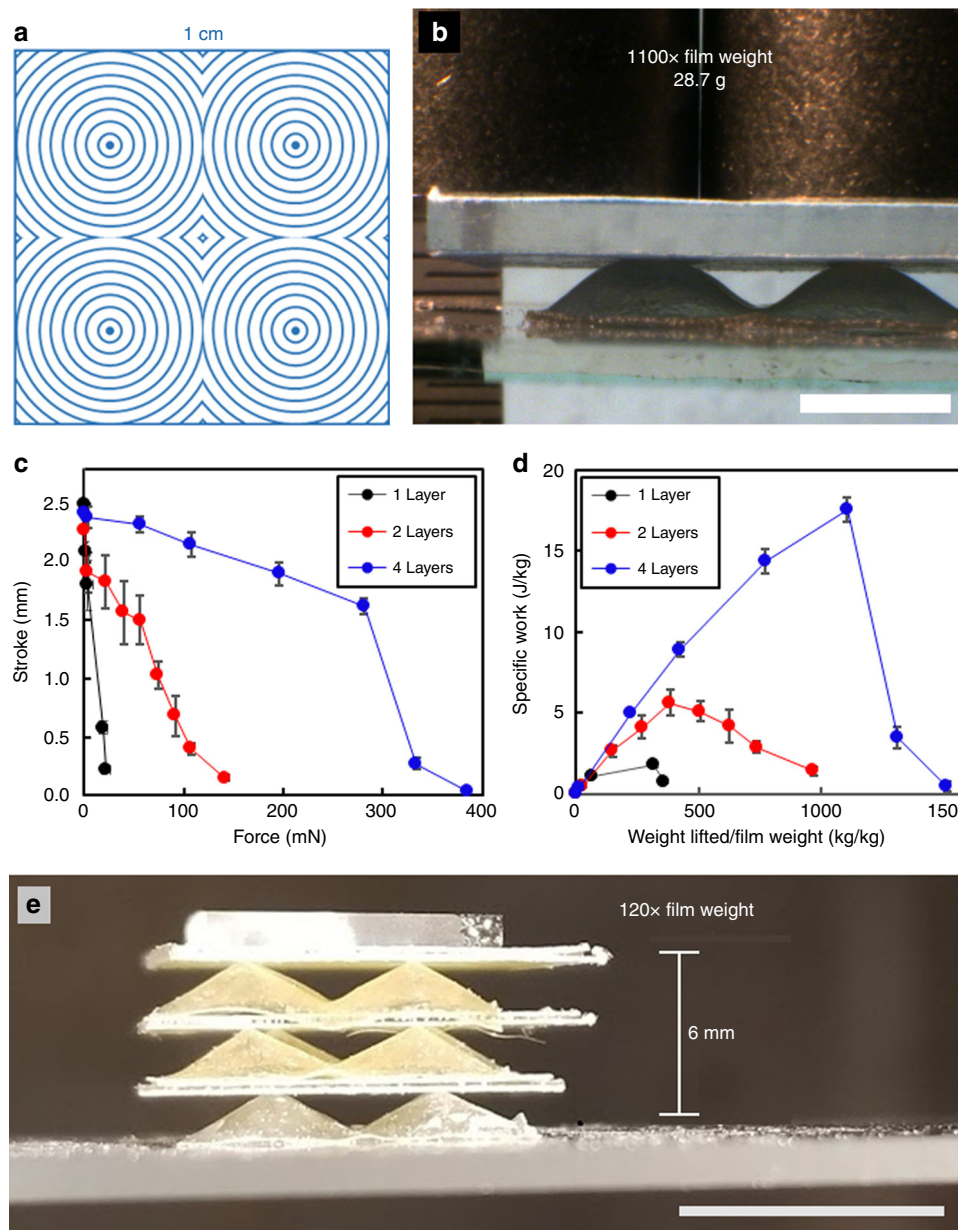


Fig. 2 Soft weightlifting. **a** Prescribed director profile, to prepare a LCE films with a 2×2 array of radial +1 defects. **b** Deformation of a four layer LCE laminate lifting 28.7 g > 2 mm. Scale bar is 3 mm. The LCE laminate was heated to 180 °C. **c** Stroke of LCE laminates under load. **d** The specific work of the LCE laminates is contrasted against normalized weight. Error bars represent statistical experimental error. **e** Three 1×1 cm LCE films are stacked with interfacial glass sheets, emulating a piezoelectric stack. A total of 1 g was lifted over 6 mm. Scale bar is 1 cm

emulated³⁴. Evident in Fig. 2e, a large stroke actuator is demonstrated in which the films are stacked on top of each other, separated by a rigid substrate (glass coverslips). With three LCE layers, a stroke exceeding 6 mm is achieved while still lifting 120 times the weight of the assembled device.

A distinguishing characteristic of LCEs in contrast to other shape-changing polymeric systems is their excellent reversibility and resistance to fatigue³⁵. Shape memory effects in polymers must be reprogrammed after each actuation^{36, 37}. The robustness of the actuation of the LCE films examined here is illustrated in Fig. 3. Figure 3a, b contrasts the deformation of an LCE film under the load of nearly 1 g. There is little distinguishable difference in stroke or shape of LCE film after 10 thermal cycles. Figure 3c is representative of the flat state reached by the LCE films after each cycle. Figure 3d summarizes the consistency in the stroke observable in 11 thermal cycles.

More is better. We initiated this study by examining a 2×2 array of +1 topological defects in a 1 cm^2 film. The force output onto the loaded substrate in Fig. 2 should be sensitive to the number of contact points. To illustrate this, LCE films with a 3×3 array of +1 defects were prepared in 1 cm^2 films. By increasing the number of defects (contact points) from four to nine, the total force output increases from 300 (Fig. 2c) to 560 mN (Fig. 4a). Evident in Fig. 4b, the four layer LCE laminate composed of the 3×3 array of +1 defects is able to lift 2150 times the weight of the film itself. A four layer LCE film can lift 56 g nearly 0.5 mm (Fig. 4c).

Evident in Fig. 4a, the stroke is significantly decreased for the 3×3 arrays in the 1 cm^2 film when compared to the 2×2 arrays in Fig. 2c. The direct comparison of 2×2 and 3×3 arrays in Fig. 4a, b are from samples in which the dimensions of each defect region was $0.33 \times 0.33 \text{ cm}^2$. These LCE

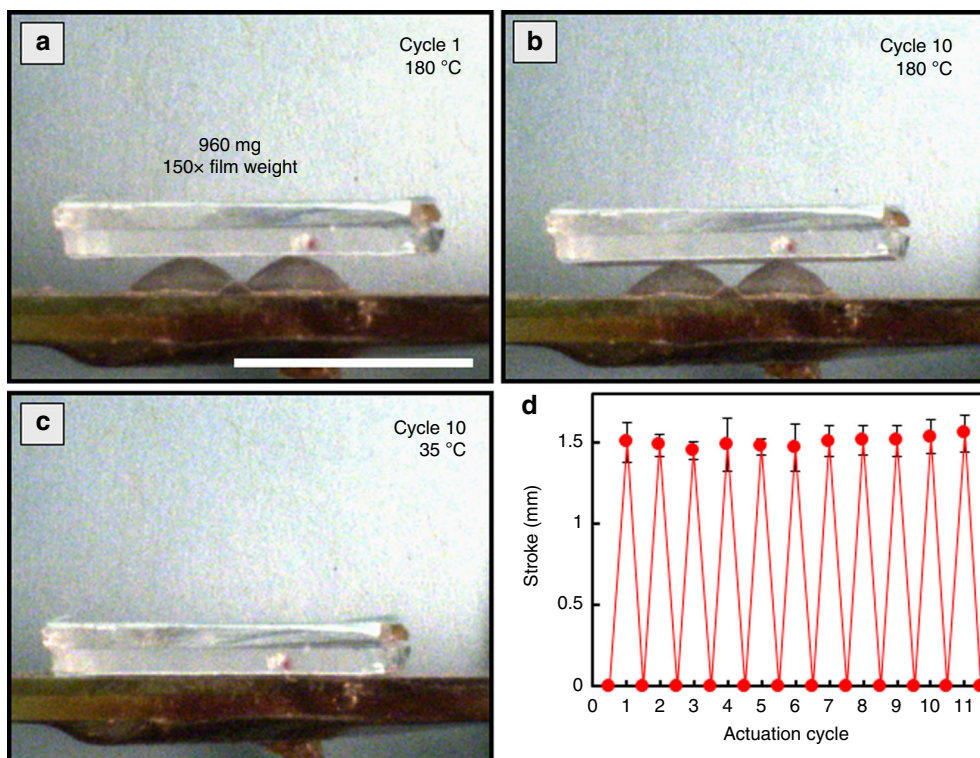


Fig. 3 Actuation cycles. Four layer LCE laminate composed of layers with 2×2 array of +1 defects. Scale bar is 1 cm. Deformation of the four layer LCE laminate under 960 mg load was monitored **a** after 1 cycle, **b** 10 cycles (heating step), and **c** 10 cycles (cooling to 35 °C). **d** Stroke vs thermal cycle for 11 actuation cycles. Error bar is statistical accuracy of measurement

laminates exhibit identical stroke lengths. However, evident in Fig. 4a, the output force is considerably increased by increasing the number of contact points. Conceivably, employing large area patterning techniques and substrates not available to us in our laboratory could allow for preparing larger area films composed of 1000s of contact points.

Deformation under pressure. Numerous end-use applications of LCEs have been discussed including irises³⁸, biomimetic actuators³⁵, valves³⁹, and shape-changing lenses^{40, 41}. One potential aerospace application is to prepare reconfigurable topographical surface features to manipulate flow⁴². We conclude this examination by measuring the deformation of the LCE laminates under positive pressure. LCE films were once again patterned with radial +1 topological defects. The deformation of a single defect subsumed in the center of a 12 mm diameter film was examined. The LCE laminates were placed in a pressure chamber where the back (or bottom) of the film was maintained at ambient pressure while the front (or top) is subjected to positive pressure. The entire chamber was heated and then the shape of the film is measured via optical profilometry. Figure 5 presents the optical scans of single, double, four, and six layer LCE laminates. Direct heating of the films in ambient pressure conditions results in the expected conical deformations (leftmost column of Fig. 5). Notably, the deformation of the films is less than that observed in free standing films, largely attributable to film anchoring. However, upon adding even slight positive pressure, the single layer film (50 μm) immediately loses its shape, compressing into the pressure chamber. The two layer LCE laminate behaves similarly. Informed by the prior results, the four layer LCE laminate (210 μm) withstands much higher loads and is able to maintain a cone-like shape at 1.5 kPa of pressure. A six layer LCE laminate (320 μm) is able to withstand over 7 kPa (>1 psi) and still

maintain a conical shape. Profiles of actuated samples are shown in Supplementary Figure 4 and Supplementary Figure 5.

Here we have demonstrated an approach to realize thick LCE films capable of large force output and stroke. Upon exposure to thermal stimulus, the LCE laminates deform into the expected shapes. Notably, despite the increase in film thickness in the LCE laminates, the deformations of the materials (e.g., the stroke) remain constant. The increase in thickness allow the laminates to impart work on objects >2000 times heavier than the laminates themselves. End-use applications in aerospace, such as reconfigurable topographical surfaces, require deformation to positive pressure. Six layer LCE laminates are shown to deform up to 7 kPa pressure and retain the expected conical deformation. In this way, this work is a critical step forward in opening up new opportunities for the implementation of these materials in applications ranging from aerospace, automobiles, and consumer goods.

Methods

Materials synthesis. RM82 (1,4-Bis-[4-(6-acryloyloxyhexyloxy)benzoyloxy]-2-methylbenzene) and RM257 (1,4-Bis-[4-(3-acryloyloxypropyloxy)benzoyloxy]-2-methylbenzene) were purchased from the Synthon Chemicals, and recrystallized from methanol before use. LCE formulations were prepared by adding 69 wt% RM82, 20 wt% RM257, 11 wt% BDMT (benzenedimethanethiol, Sigma Aldrich), with 1 wt% Irgacure 651 (BASF), and 0.5 wt% butylated hydroxytoluene to a glass vial and thoroughly mixing. A total of 50 μm thick liquid crystal cells^{9, 16} were filled via capillary action at 90 °C in the isotropic state, and then cooled slowly to 25 °C (5 min). The cells were then exposed to 365 nm UV (120 mW/cm²) light for 20 min to initiate photopolymerization. After curing, the films were collected by soaking the cells in deionized water for 2 h. Care was taken to deconstruct the cells so the LCE films remain adhered onto one of the glass substrates, to prevent wrinkling. Two films, both adhered to a single glass substrate, were placed on a hot plate at 50 °C. A drop of the LCE formulation was placed on one of the films, and the films clamped together. The films were briefly heated using a 100 °C heat gun until the adhesive layer was no longer hazy, and then allowed to cool to 25 °C over 5 min. The adhesive was cured until 365 nm UV light for 20 min. This process was repeated until the desired number of layers was achieved.

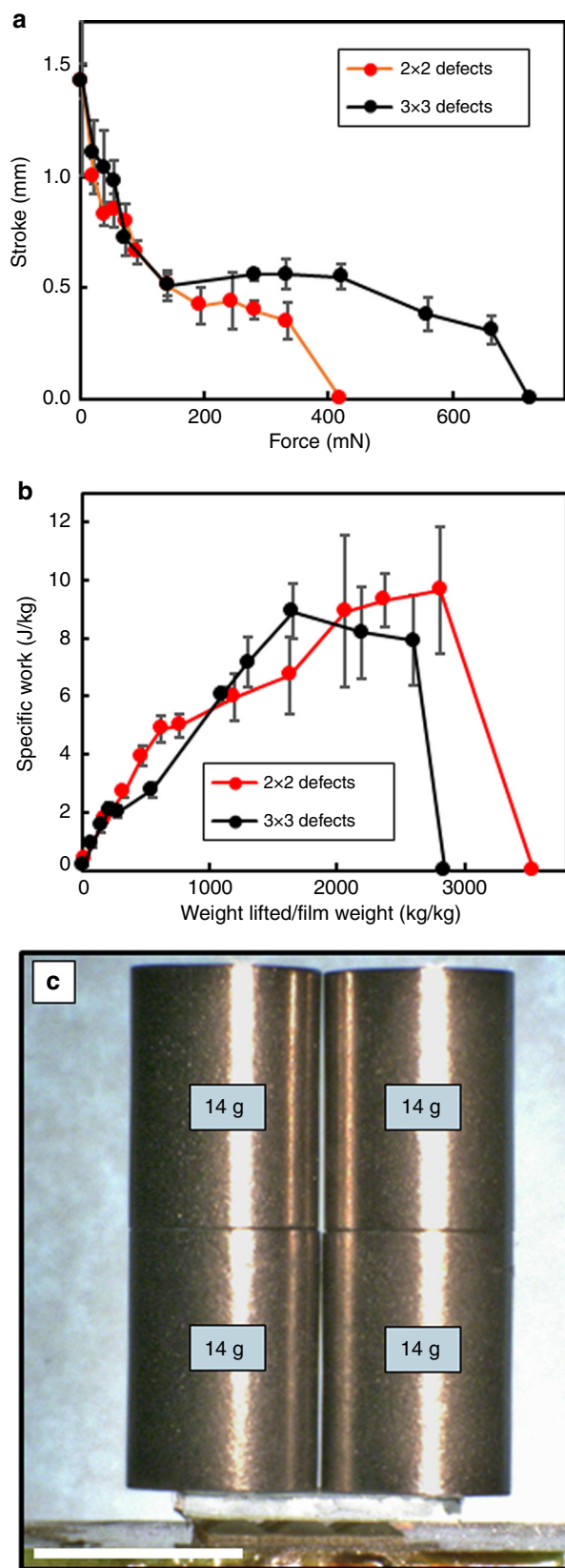


Fig. 4 Force enhancement. **a** Stroke as a function of load for 1×1 cm films patterned with 2×2 or 3×3 +1 defect arrays. **b** Specific work of the 2×2 and 3×3 arrays. Error bars represent tilt of the lifted weight and range of three experiments. **c** Illustration of the deformation of a 1×1 cm, four layer LCE laminate (26 mg) lifting over 56 g of load over 0.4 mm. Scale bar is 1 cm

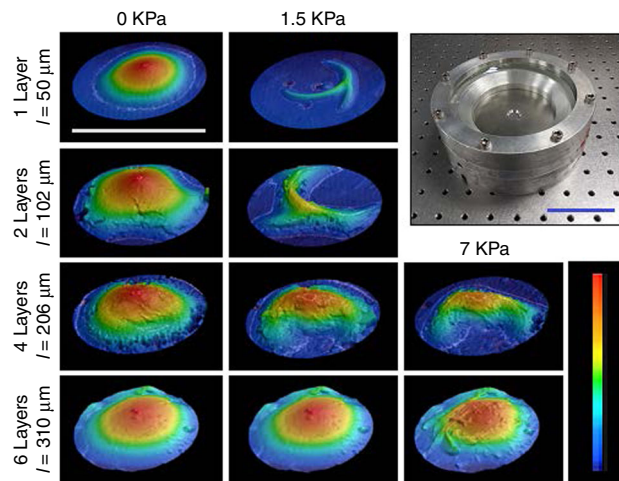


Fig. 5 Shape deformation under positive pressure. LCE films and laminates (10 mm diameter) patterned with +1 topological defects subsumed in the center were heated to 100°C in a pressure chamber (top right). Air pressure is applied on one side of the films, and the deformation measured via optical profilometry. The horizontal scale bar is 1 cm, while the height scale is 2.4 mm. The scale bar of the pressure chamber is 5 cm

Materials characterization. Phase transitions, birefringence, and film quality were measured via polarized optical microscopy (POM) (Nikon) in transmission mode, and the temperature was controlled by a Mettler Toledo HS82 heat stage. Contraction and areal change of homogenous planar films, floating on silicone oil and $5\ \mu\text{m}$ glass spacers, as a function of temperature was also determined using POM. Differential scanning calorimetry (DSC) (TA Instrument Q1000) was performed under nitrogen from -40°C to 120°C for monomer mixtures and 0°C to 300°C for cured films in hermetically sealed pans at $2^\circ\text{C}/\text{min}$. The nematic transition determined from the peak of the heat flux trace on second cooling, and the glass transition was determined from the peak of the derivative of the heat flux trace. Shape of actuated samples was measured through structured-illumination optical profilometry (Keyence VR-3200).

To measure lifting force and stroke of the actuation, 1×1 cm samples were placed on a resistive heater, loaded with weight, and heated to 180°C . The samples were loaded with successively heavier tungsten weights and/or glass slides after each test, and the height was measured by a CCD camera at the plane of the films. A ruler was always imaged frame and plane to calibrate the distance, and the film displacement was measured using ImageJ. All tests were performed by loading the sample and then heating.

Actuation under pressure was measured using a homebuilt pressure chamber, and the resultant shape change monitored in situ via optical profilometry. The sample was placed in the central hole, with a series of small holes cut out under the sample to allow the back to be exposed to ambient pressure. An exploded drawing is displayed in Supplementary Figure 6. The chamber was heated to 100°C using resistive heating silicone elements, and allowed to equilibrate for 30 min at 100°C . The chamber was then pressurized until the films collapsed. The shape change was monitored every 0.2 psi.

Data availability. The authors declare that all data supporting the findings of this study are available within the paper and its Supplementary Information.

Received: 14 March 2018 Accepted: 29 May 2018

Published online: 28 June 2018

References

- Sun, L. et al. Stimulus-responsive shape memory materials: a review. *Mater. Des.* **33**, 577–640 (2012).
- Lepora, N. F., Verschure, P. & Prescott, T. J. The state of the art in biomimetics. *Bioinspir. Biomim.* **8**, 013001 (2013).
- Bhattacharya, K. & James, R. D. The material is the machine. *Science* **307**, 53–54 (2005).
- Jani, J. M., Leary, M., Subic, A. & Gibson, M. A. A review of shape memory alloy research, applications and opportunities. *Mater. Des. (1980-2015)* **56**, 1078–1113 (2014).

5. Dye, D. Shape memory alloys: towards practical actuators. *Nat. Mater.* **14**, 760 (2015).
6. Rus, D. & Tolley, M. T. Design, fabrication and control of soft robots. *Nature* **521**, 467 (2015).
7. White, T. J. & Broer, D. J. Programmable and adaptive mechanics with liquid crystal polymer networks and elastomers. *Nat. Mater.* **14**, 1087 (2015).
8. De Gennes, P. Réflexions sur un type de polymères nématiques. *CR Acad. Sci. Ser., B* **281**, 101–103 (1975).
9. Ware, T. H., McConney, M. E., Wie, J. J., Tondiglia, V. P. & White, T. J. Voxellated liquid crystal elastomers. *Science* **347**, 982–984 (2015).
10. Gimenez-Pinto, V., Ye, F., Mbanga, B., Selinger, J. V. & Selinger, R. L. Modeling out-of-plane actuation in thin-film nematic polymer networks: From chiral ribbons to auto-origami boxes via twist and topology. *Sci. Rep.* **7**, 45370 (2017).
11. Fuchi, K. et al. Topology optimization for the design of folding liquid crystal elastomer actuators. *Soft Matter* **11**, 7288–7295 (2015).
12. de Haan, L. T. et al. Accordion-like actuators of multiple 3D patterned liquid crystal polymer films. *Adv. Func. Mater.* **24**, 1251–1258 (2014).
13. Kowalski, B. A., Mostajeran, C., Godman, N. P., Warner, M. & White, T. J. Curvature by design and on demand in liquid crystal elastomers. *Phys. Rev. E* **97**, 012504 (2018).
14. de Haan, L. T., Sánchez-Somolinos, C., Bastiaansen, C. M., Schenning, A. P. & Broer, D. J. Engineering of complex order and the macroscopic deformation of liquid crystal polymer networks. *Angew. Chem. Int. Ed.* **51**, 12469–12472 (2012).
15. Thomsen, D. L. et al. Liquid crystal elastomers with mechanical properties of a muscle. *Macromolecules* **34**, 5868–5875 (2001).
16. McConney, M. E. et al. Topography from topology: photoinduced surface features generated in liquid crystal polymer networks. *Adv. Mater.* **25**, 5880–5885 (2013).
17. Liu, Y., Wu, W., Wei, J. & Yu, Y. Visible light responsive liquid crystal polymers containing reactive moieties with good processability. *ACS Appl. Mater. Interfaces* **9**, 782–789 (2017).
18. Guin, T. et al. Electrical control of shape in voxellated liquid crystalline polymer nanocomposites. *ACS Appl. Mater. Interfaces* **10**, 1187–1194 (2018).
19. Courty, S., Mine, J., Tajbakhsh, A. & Terentjev, E. Nematic elastomers with aligned carbon nanotubes: new electromechanical actuators. *Europhys. Lett.* **64**, 654 (2003).
20. Wermter, H. & Finkelmann, H. Liquid crystalline elastomers as artificial muscles. *e-Polymers* **1**, 1 (2001).
21. Modes, C., Bhattacharya, K. & Warner, M. Gaussian curvature from flat elastica sheets. *Proc. R. Soc. A* **467**, 1121–1140 (2011).
22. Kowalski, B. A., Tondiglia, V. P., Guin, T. & White, T. J. Voxel resolution in the directed self-assembly of liquid crystal polymer networks and elastomers. *Soft Matter* **13**, 4335–4340 (2017).
23. Godman, N. P., Kowalski, B. A., Auguste, A. D., Koerner, H. & White, T. J. Synthesis of elastomeric liquid crystalline polymer networks via chain transfer. *ACS Macro Lett.* **6**, 1290–1295 (2017).
24. Broer, D., Crawford, G. P. & Zumer, S. *Cross-linked liquid crystalline systems: from rigid polymer networks to elastomers*. (CRC Press, Boca Raton, FL: 2011).
25. Kowalski, B. A., Guin, T. C., Auguste, A. D., Godman, N. P. & White, T. J. Pixelated polymers: directed self-assembly of liquid crystalline polymer networks. *ACS Macro Lett.* **6**, 436–441 (2017).
26. Spillmann, C. M. et al. Stacking nematic elastomers for artificial muscle applications. *Sens. Actuat. A: Phys.* **133**, 500–505 (2007).
27. Sato, Y., Sato, K. & Uchida, T. Relationship between rubbing strength and surface anchoring of nematic liquid crystal. *Jpn. J. Appl. Phys.* **31**, L579 (1992).
28. Yaroshchuk, O. & Reznikov, Y. Photoalignment of liquid crystals: basics and current trends. *J. Mater. Chem.* **22**, 286–300 (2012).
29. Wu, S.-T., Efron, U. & Hess, L. D. Birefringence measurements of liquid crystals. *Appl. Opt.* **23**, 3911–3915 (1984).
30. Küpfer, J. & Finkelmann, H. Nematic liquid single crystal elastomers. *Makromol. Chem: Rapid Commun.* **12**, 717–726 (1991).
31. Modes, C. D. & Warner, M. Blueprinting nematic glass: systematically constructing and combining active points of curvature for emergent morphology. *Phys. Rev. E* **84**, 021711 (2011).
32. Modes, C. & Warner, M. Shape-programmable materials. *Phys. Today* **69**, 32–38 (2016).
33. Ashby, M., Brechet, Y., Cebon, D. & Salvo, L. Selection strategies for materials and processes. *Mater. Des.* **25**, 51–67 (2004).
34. Sherrit, S. et al. Multilayer piezoelectric stack actuator characterization. *Proc. Soc. Photo-Opt. Ins.* **6929**, 692909 (2008).
35. Agrawal, A. et al. Electromechanically responsive liquid crystal elastomer nanocomposites for active cell culture. *ACS Macro Lett.* **5**, 1386–1390 (2016).
36. Hager, M. D., Bode, S., Weber, C. & Schubert, U. S. Shape memory polymers: past, present and future developments. *Prog. Polym. Sci.* **49–50**, 3–33 (2015).
37. Zhao, Q., Qi, H. J. & Xie, T. Recent progress in shape memory polymer: new behavior, enabling materials, and mechanistic understanding. *Prog. Polym. Sci.* **49–50**, 79–120 (2015).
38. Zeng, H., Wani, O. M., Wasylczyk, P., Kaczmarek, R. & Priimagi, A. Self-regulating iris based on light-actuated liquid crystal elastomer. *Adv. Mater.* **29**, 30 (2017).
39. Sánchez-Ferrer, A. et al. Liquid-crystalline elastomer microvalve for microfluidics. *Adv. Mater.* **23**, 4526–4530 (2011).
40. Xu, S. et al. Adaptive liquid lens actuated by photo-polymer. *Opt. Expr.* **17**, 17590–17595 (2009).
41. Petsch, S., Grewe, A., Köbele, L., Sinzinger, S. & Zappe, H. Ultrathin Alvarez lens system actuated by artificial muscles. *Appl. Opt.* **55**, 2718–2723 (2016).
42. Lachmann, G. V. *Boundary layer and flow control: its principles and application*. (Pergamon Press, Oxford, England: 1961).

Acknowledgements

This work was financially supported by the Air Force Office of Scientific Research and the Air Force Research Laboratory.

Author Contributions

Initial research directions were identified by T.G., R.V.B., G.W.B., and T.J.W. Experimental examinations were undertaken by T.G. with assistance from M.J.S., B.A.K., and A.D.A. All authors contributed to the writing of the manuscript.

Additional information

Supplementary Information accompanies this paper at <https://doi.org/10.1038/s41467-018-04911-4>.

Competing interests: The authors declare no competing interests.

Reprints and permission information is available online at <http://npg.nature.com/reprintsandpermissions/>

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>.

© The Author(s) 2018