

Materials Science

Special Topic: Flexible Electronic and Micro/Nanomanufacturing

Achieving rapid actuation in liquid crystal elastomers

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Received 23 April 2024; Revised 16 May 2024; Accepted 3 June 2024; Published online 6 June 2024

Abstract: Liquid crystal elastomer (LCE) is one kind of soft actuating material capable of producing large and reversible actuation strain, versatile and programmable actuation modes, and high work density, which can be widely exploited for next-generation soft robots. However, the slow response speed and low power density in LCE-based actuators remain a challenge, limiting their practical applications. Researchers have been considering how to improve these performances. In this review, we discuss the fundamentals of the LCEs and emphasize the fast actuation strategies developed in recent years. Initially, we introduce conventional preparation strategies. Then, we describe typical actuation mechanisms of LCEs, discussing their features and limitations. Subsequently, we summarize several possible approaches as case studies to enhance the actuation performance of LCEs, including reducing physical sizes, introducing active heating-cooling mechanisms, utilizing mechanical instability, and developing dielectric LCEs. Finally, we discuss the future research opportunities and challenges for rapid actuation of LCEs.

Keywords: liquid crystal elastomer, soft robotics, rapid actuation

INTRODUCTION

Robotics made from soft materials exhibit salient features, such as body compliance, large and continuous deformation, and benign human-machine interaction, which stand in contrast to their rigid counterparts [1]. These unique features make soft robots a promising candidate for broad applications in space exploration, rehabilitative devices, prostheses, and dexterous manipulations. In recent decades, numerous works have been dedicated to developing soft robots with various locomotion, grasping or manipulation, etc. For instance, Tolley *et al.* developed an untethered pneumatic soft quadruped that could crawl under extreme and harsh environments (e.g., snowstorms, water, and fire) [2]. Inspired by the characteristics of deep-sea creatures, Li *et al.* developed an untethered, self-powered soft robot that could be used for deep-sea exploration [3]. Practical tests were conducted at the deepest point of 10,900 m in the Mariana Trench. This soft robot exhibited exceptional pressure resistance and swimming performance owing to its electrostatic mechanism. Wood *et al.* presented an ultragentle soft robotic actuator that could grasp delicate specimens of

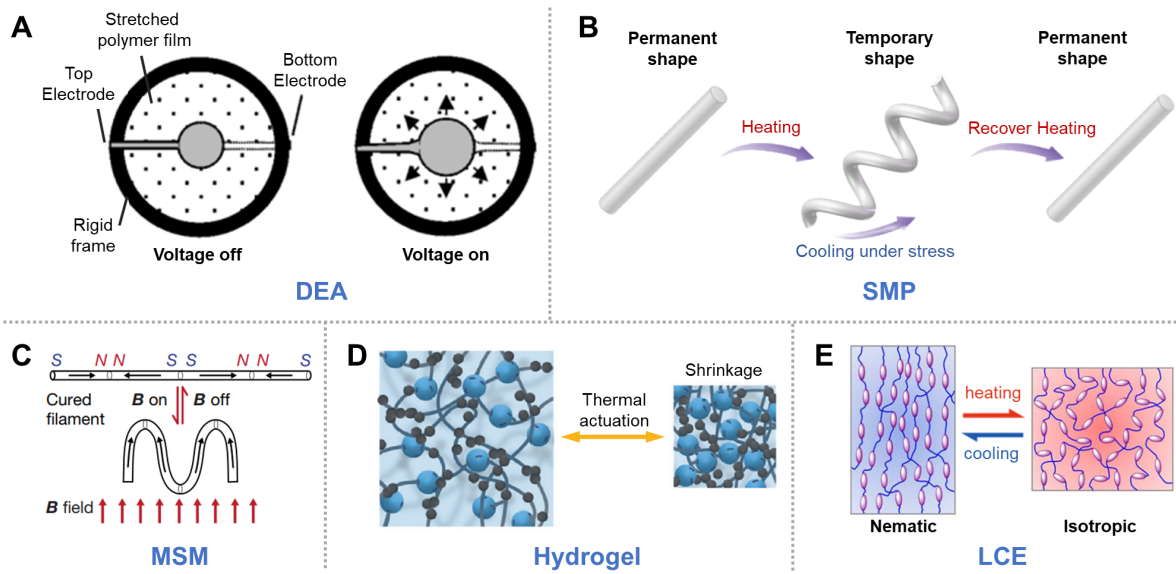


Figure 1 The actuation mechanism of typical soft actuating materials. (A) When subject to high voltage, a strong electrostatic interaction is generated between two electrodes, compressing the film in the thickness direction and expanding it in the area. Reprinted with permission from Ref [6]. (B) After being deformed at a high temperature and fixed into a temporary shape after cooling, shape memory polymer (SMP) can recover to its permanent shape upon heating. Reprinted with permission from Ref [9]. (C) Magnetic soft material (MSM) with programmed magnetic domains exhibits complex shape changes under external magnetic field stimulus. Reprinted with permission from Ref [10]. (D) Thermal stimulated isotropic volumetric change of poly(N-isopropylacrylamide) (PNIPAAm) hydrogel. Reprinted with permission from Ref [14]. (E) Upon thermal stimuli, LCE undergoes the nematic-isotropic phase transition, causing a large contraction in the alignment direction. Reprinted with permission from Ref [15].

gelatinous marine life. Three types of live jellyfish were successfully grasped and manipulated using the soft robotic gripper [4].

For all these examples, the soft actuating materials and deployable structures work as the “motor” to drive the locomotion and conduct advanced functions. They are compliant and capable of producing large deformation in response to external stimuli. Recently, researchers have devoted lots of effort to developing various soft actuating materials with different characteristics [5], including dielectric elastomer actuators (DEAs), shape memory polymers (SMPs), hydrogels, magnetic soft materials (MSMs), and liquid crystal elastomers (LCEs). The DEA is a deformable capacitor made by sandwiching a dielectric elastomer (DE) film between two compliant electrodes. When subjected to voltage, DEAs generate a strong electrostatic interaction between two electrodes, compressing the film in the thickness direction and expanding it in the area (Figure 1A) [6,7]. The DEAs can produce rapid actuation when a large electric potential of several kilovolts is applied. However, high voltage conditions increase the complexity of the control system and pose a potential safety threat to humans. SMPs can change morphology and stiffness when the temperature varies, owing to the phase transition or decrosslinking of temporary interaction (Figure 1B) [8,9]. Still, they usually have one-way actuation, which cannot recover to the initial shape after deformation. MSMs are made by dispersing magnetized or magnetizable microparticles in polymeric matrices (Figure 1C) [10,11]. These composites could deform wirelessly under an external magnetic field. It is fast and can be remotely controlled, which can be broadly used in biomedical applications. Nevertheless, the control of soft magnetic composites is limited by external magnetic fields. Responsive hydrogels, for example, poly(N-isopropylacrylamide) (PNIPAAm), can change into a hydrophobic state when the temperature is increased,

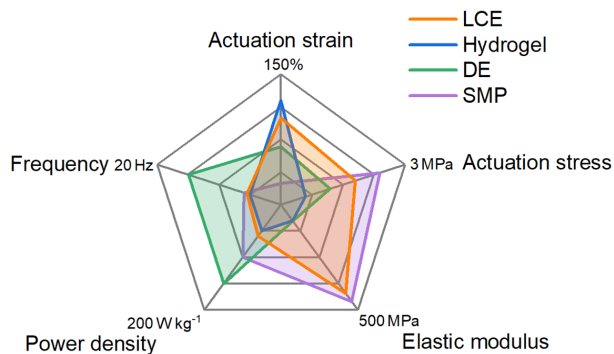


Figure 2 Actuation performances of different soft actuating materials.

leading to a volumetric shrinkage [12]. They have been widely studied in fields such as marine robots and biomedical tools (Figure 1D) [13,14]. However, the work mechanism of hydrogels relies on diffusion, causing a slow response speed.

Recently, LCEs have received significant interest due to their large reversible actuation strain, versatile and programmable actuation modes, and high work density. These distinctive characteristics make them an ideal option for achieving sensing and actuation functions in soft robots [16]. LCEs are a combination of liquid crystal mesogens and polymer networks, possessing anisotropy of mesogens and elasticity of polymer networks [17–19]. When subjected to external stimuli (e.g., temperature, light, non-polar solvents), LCEs undergo the nematic-isotropic phase transition, causing an anisotropic contraction (Figure 1E) [20–22]. By spatially patterning the mesogens, they can produce versatile in-plane or out-of-plane actuation modes, including bending, twisting, curling, and extension [23–25]. Once external stimuli are removed, LCEs fully recover to their initial shape, showing good reversibility. However, the actuation and recovery of the LCE are relatively slow compared to DEAs and conventional machines, significantly restricting their applications regardless of their large deformation, high work density, and easy actuating methods (Figure 2) [26–31]. Thus, it is emerging to develop strategies to produce rapid actuation in LCE materials.

In this review, we discuss the fundamentals of LCEs and highlight various strategies to improve their response speed and power density. We first introduce different synthesis strategies for LCEs. Then, we discuss typical actuation methods of LCEs. Subsequently, four possible approaches to improve the response speed are highlighted, including reducing the critical length scale of LCEs, introducing active heating and cooling systems in LCEs, utilizing instability, and developing dielectric liquid crystal elastomers. Finally, perspectives on challenges and opportunities for rapid actuation of LCEs are discussed.

SYNTHESIS AND ALIGNMENT OF LCES

The LCEs are made by integrating liquid crystal mesogens into a polymer network. The synthesis and processing of LCEs determine their actuation performance and application scenarios. The actuation behavior of LCEs is dependent on the alignment of mesogens. Without additional treatment during preparation, the LCEs are usually in a polydomain state, in which the alignments of mesogens in different domains are random. Therefore, the polydomain LCEs cannot produce macroscopic deformation upon external stimuli.

To achieve actuation capability, the mesogens should first be highly aligned and then fixed by a polymer network, namely monodomain LCEs. Several strategies have been explored in the past years to prepare the monodomain LCEs. They can be classified into three categories: one-step polymerization, two-step polymerization, and dynamic covalent network methods [32].

One-step polymerization method

In the one-step polymerization method, reactive liquid crystal molecules are used. When heated above the phase transition temperature, the small molecules have a low viscosity, which can be aligned by surface or external electric/magnetic field. Then, the aligned molecules are exposed to ultraviolet (UV) light or heat to form a polymer network, fixing the alignment of mesogens and producing the monodomain LCEs. The alignment patterns of the mesogens can be controlled through well-developed techniques, enabling versatile actuation modes and rich design space of LCEs. For instance, Zeng *et al.* prepared an LCE iris using photoalignment technology. The liquid crystal cell (20 μm thick) was constructed so that the mesogens adopted homeotropic alignment at the upper surface. In contrast, the bottom surface was coated with a photoalignment layer, resulting in radial alignment of the mesogens. Like the natural iris, the LCE could autonomously open and close in response to incident light intensity (Figure 3A) [33]. In another work, Yang *et al.* utilized a designed channel-pattern mold with a pattern feature size of 1 μm to prepare LCE films with programmable alignment [34]. Upon thermal activation, 2D LCE films could be deformed into desired 3D shapes, such as the complex human face.

In the surface alignment technique, the mesogen is usually aligned parallel to the surface, while the magnetic field can generate an alignment perpendicular to the substrate. For instance, Cui *et al.* utilized a pair of permanent magnets to prepare an LCE pillar micropattern, in which the mesogens were aligned along the pillar direction (Figure 3B) [35]. The pillar micropatterns could undergo reversible switching between adhesive and non-adhesive states in response to temperature changes. However, the thickness of the LCE film prepared by the surface alignment or magnetic/electric field-assisted alignment is limited to tens to hundreds of micrometers.

Two-step polymerization method

Unlike the one-step polymerization method, in the two-step polymerization method, the liquid crystal mesogens are first polymerized into linear oligomers or loosely crosslinked polymer networks with residual reactive groups inside. The mesogens are further aligned by mechanical force. Then, the alignment is fixed by forming a polymer network due to the crosslinking of residual reactive groups. The two-step polymerization method can be employed to prepare the LCE film on large scales. Finkelmann and coworkers first reported the two-step polymerization method for fabricating monodomain LCEs using two kinds of hydrosilylation reaction with different reaction rates [36]. Later, Yakacki *et al.* used a commercially available liquid crystal mesogen that could undertake a two-stage thiol-acrylate Michael addition and photopolymerization reaction, significantly improving the simplicity and reproducibility of the monodomain LCEs (Figure 3C) [37].

In the two-step polymerization method, the mesogens can be first polymerized into linear oligomers, which

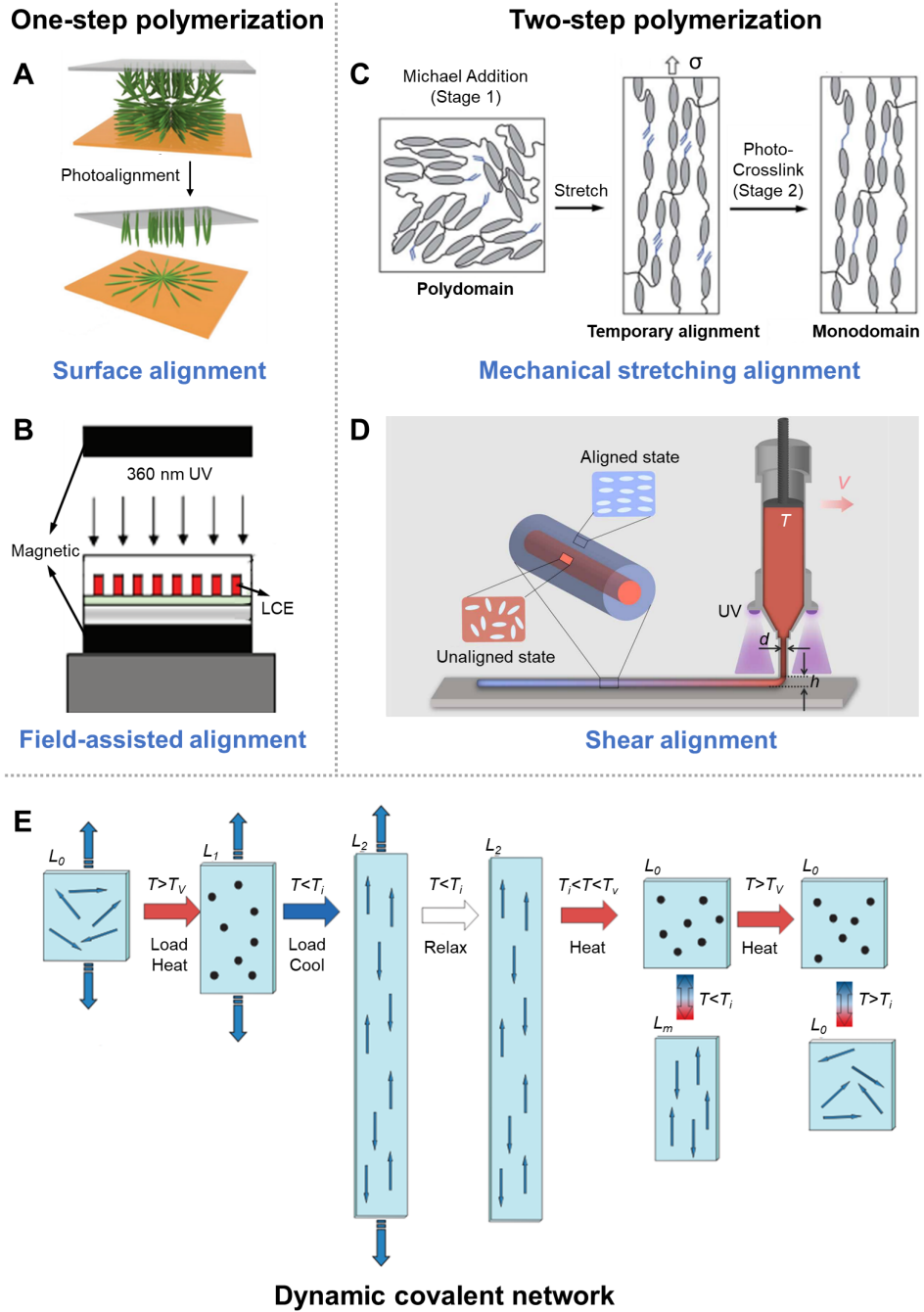


Figure 3 Synthesis and alignment of LCEs. (A) In the one-polymerization method with surface alignment technique, liquid crystal monomer mixture adopts a centrally symmetric splayed alignment directed by molecular orientation at the top and bottom surfaces. Reprinted with permission from Ref [33]. (B) The mesogens align along the longitudinal axis of the LCE pillars by the application of a magnetic field. Reprinted with permission from Ref [35]. (C) The process of a typical two-step polymerization method. The thiol-acrylate Michael addition and photopolymerization (TAMAP) reactions are used to synthesize monodomain LCEs. The polydomain LCE is prepared first and mechanically stretched to align the mesogens into a temporary monodomain. Then, the alignment is fixed through the second-step cross-linking process. Reprinted with permission from Ref [37]. (D) Schematic illustration of the DIW of LCE. The mesogens are aligned along the printing path by shear stress and then are fixed by post-curing. Reprinted with permission from Ref [48]. (E) The schematic illustration of preparing monodomain with dynamic covalent chemistry method. First, the polydomain LCE is uniaxial stretched under external forces and the exchange reaction of dynamic bonds is activated under external stimuli, leading to the rearrangement of the polymer network. As a result, the LCE is programmed into a monodomain state. Reprinted with permission from Ref [49].

are extrudable and thereby can be used as the ink in the direct ink writing (DIW) 3D printing technique, enabling the spatial alignment of mesogens by computer-aided design (CAD) [38–42]. In this way, LCEs can be fabricated in desired alignment patterns and shapes, thereby achieving customized deformation [43–45]. During the printing process, the mesogens are aligned along the printing direction via shear force and then are fixed by post-curing [46,47]. The degree of alignment can be tailored by printing parameters, such as the inner diameter of the nozzle, printing temperature, distance between the nozzle and substrate, etc. Wang *et al.* printed versatile active morphing structures with functionally graded LCEs by varying printing parameters, achieving the versatility of deformation and structure (Figure 3D) [48]. However, the spatial resolution of the alignment of mesogens is limited by the nozzle diameter ($> 150 \mu\text{m}$).

Dynamic covalent network method

The LCEs with regular covalent bonds prepared by one-step and two-step polymerizations are thermoset, and their actuation behavior cannot be reprogrammed, reprocessed, and recycled once cured. Ji *et al.* creatively introduced the dynamic covalent bonds (DCBs) into LCEs to make them reprogrammable and modulable (Figure 3E) [49]. The DCBs can undergo exchange reactions under external stimuli (such as heat or light), leading to the rearrangement of the polymer network. As a result, the LCE can be programmed into a monodomain state. The LCEs with DCBs exhibit reprocessing, reshaping, and self-healing abilities and can be assembled with modulable building blocks [50,51]. At present, numerous DCBs have been incorporated into LCEs, such as ester groups [52–57], disulfide bonds [15,58,59], boronic esters [60,61], Diels-Alder reactions [62], diselenide bonds [63], and so on.

ACTUATION METHODS OF LCEs

The nematic-isotropic phase transition of LCEs can be triggered via thermal or photochemical stimuli, depending on the types of mesogens. Azobenzene derivatives are typical photo-responsive mesogens. They can undertake trans-cis transition upon applying ultraviolet light [64–66]. Ceamanos *et al.* showed that an LCE strip containing azobenzene could generate photochemical actuation (Figure 4A) [67]. After being irradiated by UV light, the azobenzene molecules underwent a transition from trans to cis isomer, and the LCE strip bent towards the light. After being exposed to blue light (455 nm) for 30 min or left in darkness for 24 hours, the LCE strip returned to its original shape. However, the depth of light penetration limits the photochemical actuation of LCEs. Compared with photochemical-responsive LCEs, thermally responsive LCEs have a variety of types and can be prepared from commercially available materials. When heated above the phase transition temperature (T_i), they could deform into desired shapes. Changing mesogens or tuning the crosslinking density can easily adjust the phase transition temperature. Regarding the activating approaches of the LCE, environmental heating, electrothermal, photothermal, and magnetothermal heating have been utilized to trigger the deformation by structural design. In the following, we introduce several recently developed heating actuation methods.

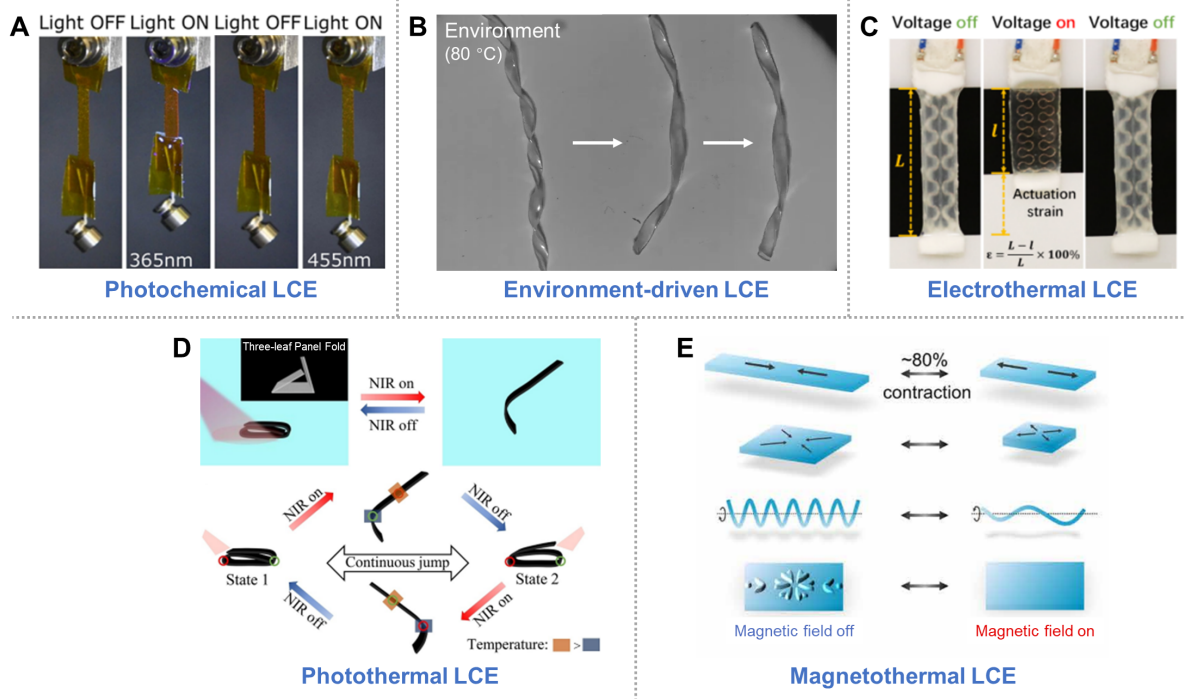


Figure 4 Actuation performance of LCEs with different actuation types. (A) Photochemical mechanism. Under UV irradiation, azobenzene molecules can transition from trans to cis isomerization, leading to the contraction of the LCE strip. After exposure to blue light (455 nm), the LCE strip returns to its original shape. Reprinted with permission from Ref [67]. (B) Environmental heat mechanism. A helical-shaped LCE self-rolling robot powered by environmental heat achieves autonomous locomotion. Reprinted with permission from Ref [72]. (C) Electro-thermal mechanism. Joule heat is generated when a voltage is applied to the metallic wire, increasing the temperature and causing LCE to contract in the longitudinal direction. After turning off the voltage, the LCE gradually recovers to its original shape. Reprinted with permission from Ref [85]. (D) Photo-thermal mechanism. When the CNT-LCE ribbon is exposed to NIR light, CNT can convert the light to heat, causing the temperature to increase and further inducing the deformation. Reprinted with permission from Ref [95]. (E) Magnetothermal mechanism. The Fe_3O_4 nanoparticle doped-LCE actuators can exhibit different motion modes under the alternating magnetic fields. Reprinted with permission from Ref [102].

Environmental heating actuation

Environmental heating could directly induce the deformation of the whole structure. In this way, the LCE-based robots could exhibit locomotion or manipulation with the sequential on/off of environmental heating. Ji *et al.* made a tubular LCE gripper that could autonomously grab an object under a high-temperature environment. The object was released automatically when the gripper moved to a cold place [68]. In addition, environments with a gradient temperature field could also be employed to drive the autonomous motion of the LCE structures [69]. For instance, Cai *et al.* reported that a simple LCE rod could roll on a flat and hot surface, exhibiting the capability of environmental autonomy [70]. By modifying the structure of the LCE rod, Yin *et al.* subsequently developed a helical-shaped LCE with self-navigation capability. This robot could autonomously escape a complex maze (Figure 4B) [71,72]. Furthermore, Daraio *et al.* designed an assembled LCE pentagonal prism that could perform a programmed response to fold and roll when subjected to thermal stimuli [73]. Nevertheless, this design has been only used to produce limited rolling locomotion. By integrating thermally responsive cables, non-responsive cables, and stiff rods, Wang *et al.* constructed a self-propelling hybrid tensegrity structure that enabled multimodal self-propelled motion [74]. For all these examples, the LCEs driven by environmental heat have relatively simple structures without the requirements

of onboard control systems and power supplies (e.g., batteries).

Electrothermal actuation

Embedding electrothermal elements into the LCEs offers a convenient approach for actuation control and system integration. Once the voltage is applied, Joule heat can increase the temperature of LCEs and result in the actuation. The heating patterns and heating sequence/algorithm could be well designed and controlled to accomplish more efficient thermal regulation. Various types of heating elements have been embedded into LCE, including nanoparticles (e.g., graphite and silver paste) and metallic wires (e.g., gold, copper, and liquid metal) [75–79]. The conductive nanoparticles are usually used as fillers and dispersed throughout the LCE precursors. While metallic wires are generally embedded into the LCEs as separate heaters through lamination or coaxial 3D printing, etc. For instance, Luo *et al.* demonstrated an electro-conductive LCE containing graphite/carbon black that produced 40% contraction under 150 V [80]. In parallel, stretchable electronics have also been integrated into LCE as heating elements and sensors. Yu *et al.* designed and fabricated ultrathin electronics innervated soft robots that could realize adaptable crawling locomotion through the conjugation of sensing and actuation [81]. Similar advanced functions have been demonstrated using liquid metal (LM) with high stretchability. Majidi and coworkers embedded LM micro droplets into the LCE matrix, forming an LM-LCE composite that could enhance thermal and electrical properties [82]. Additionally, LCE can be integrated via the 3D printing technique. Lewis *et al.* printed a core-shell LM-LCE fiber, which could be actuated by Joule heating and achieve closed-loop control based on changes in electric resistance [83]. Recently, Liu *et al.* created an LM-LCE actuator with programmable resistance regulated by gravity that exhibited a gravitropism response due to the dynamic interaction between the thermomechanical response of the actuator and the gravity [84]. Meanwhile, the electrical control methods provide a promising solution for making untethered soft robots at the system level. For example, He *et al.* created an untethered LCE robot by combining a lithium polymer battery, LCE tubular actuators, and a microcontroller, completely free from the restriction of an external power supply or control unit (Figure 4C) [85].

Photothermal actuation

The actuation of LCEs can also be stimulated via a photothermal mechanism. Compared to the photochemical mechanism, LCEs driven by the photothermal mechanism exhibit unique features, including large penetration depth, long-term stability, and diverse materials selections. Multiple types of photothermal agents, including carbon nanotubes (CNTs) [86–88], gold nanorods (AuNRs) [89,90], polydopamine (PDA) [91,92], and MXene [93,94], have been incorporated into the LCE matrix. When LCEs are exposed to light, these doped agents directly transform light into heat, causing the temperature to increase and further inducing the actuation. The photothermal actuation methods provide a remote control on the deformation of the actuators. For instance, Yang *et al.* designed a three-leaf panel folding structure prepared by CNT-LCE, which could generate jumping motion through energy storage and release (Figure 4D) [95]. Li *et al.* reported an MXene-LCE soft tubular actuator, which could exhibit adaptive phototropism and robust omnidirectional light-tracking capability in 3D space [96]. In addition to excellent photothermal conversion, MXene can be a sensor due to its high conductivity. Liu *et al.* produced a near-infrared-driven MXene-LCE bimorph

Table 1 The actuation performance of conventional LCEs and human skeleton muscle regarding actuation strain, stress, response time (actuation and recovery), work density, and power density

Materials	Actuation strain (%)	Actuation stress (MPa)	Actuation time (s)	Recovery time (s)	Work density (kJ m ⁻³)	Power density (W kg ⁻¹)	References
Environment-driven LCE	44	0.19	180	210	40	0.21	40
CNT-LCE	15	1.02	42	50	38	1.66	88
Heating wire-LCE	41	0.35	30	240	150	0.53	85
Heating wire-LCE	28	0.23	23	180	80	0.32	78
Heating wire-LCE	32	0.49	20	150	340	0.58	59
Heating wire-LCE	40	0.44	30	100	110	1.35	79
LM-LCE	50	0.15	15	125	180	0.54	82
Human muscle	50	0.35	—	—	40	50	103

membrane with self-sensing and feedback loop control functions. The microcontroller received the signal of the resistance change and issued a command to modulate the near-infrared (NIR) laser irradiation to realize the closed-loop control of the LCE actuation [97]. Meanwhile, under a constant light source, light-driven LCEs can produce self-oscillation by a self-shadowing effect during photo-thermal-mechanical actuation and self-regulate by the resulting built-in negative feedback loop. He *et al.* developed a candle soot (CS)-doped LCE/polydimethylsiloxane (PDMS) bilayer structure that could achieve sunlight-powered self-oscillation [98].

Magnetothermal actuation

Magnetic nanoparticles, such as Fe₃O₄ nanoparticle [99,100], can be introduced to endow LCEs with magnetothermal actuation. Under alternating magnetic fields (AMFs), magnetic nanoparticles convert the energy of AMF into heat due to the hysteresis loss, relaxation processes, and other factors, resulting in a temperature increase of LCEs [101]. The magnetothermal actuated LCEs can be controlled remotely in enclosed and confined spaces on account of the strong penetrating power of magnetic fields. Ji *et al.* developed a magnetothermal-responsive LCE actuator by dispersing Fe₃O₄ nanoparticles into LCE, which could exhibit different motion modes, including large contraction (~80%), biaxial shrinkage, complex 3D patterns, and so on (Figure 4E) [102]. Furthermore, by integrating LCEs with different contents of magnetic particles, local and sequential magnetic control has been achieved.

RAPID ACTUATION STRATEGIES FOR LCE ACTUATORS

As summarized above, the LCEs exhibit large deformation and high work density via different thermal activating methods. Nevertheless, the response times of LCEs, including the actuation and recovery, are limited to a few seconds. The slow response significantly limits the practical applications of LCE actuators in soft robots. This is because the response of thermally driven LCEs depends on the thermal conductivity of the LCE materials and the heat diffusion process during the actuation.

Table 1 summarizes the actuation performance of conventional LCE-based soft actuators and human skeleton muscles [103]. It can be seen that the actuation strain, stress, and work density of LCEs are all

comparable to human muscle. However, the response speed and power density are one magnitude lower than human muscle. In this context, multiple strategies have been developed to address this challenge in recent years. These approaches include reducing critical size, introducing an active heating-cooling system, utilizing mechanical instability, and developing other actuation mechanisms. In this section, we will discuss these strategies in detail. Similar strategies can be applied to other thermally driven materials, such as shape memory polymers, twisting nylon fibers, and hydrogels.

Reducing the critical length scale of LCEs

The reversible actuation depends on the contraction and recovery speed of LCEs, which is determined by the thermal diffusion process. According to the thermal capacity model, the thermal time constant is determined by the equation: $\tau = \frac{d\rho c}{h}$, where d is the characteristic thickness of the LCE, ρ is the mass density, c is gravimetric specific heat capacity, and h is the convective coefficient. The LCE materials determine the density ρ , specific heat capacity c , and convective coefficient h . Therefore, reducing the critical length scale of LCEs shortens the thermal time constant, thereby significantly enhancing the response speed of LCEs. Moreover, the power density can also be improved since it is proportional to the frequency of actuators ($P \sim \sigma \varepsilon f$, where σ , ε , and f are actuation stress, strain, and frequency, respectively.).

Fabricating LCE fibers by the spinning method

Researchers have used different manufacturing techniques to fabricate LCE fibers at the microscale [104,105]. The spinning method is widely adopted to convert liquid or solid precursor materials into continuous solid fibers with specific structures and properties. The commonly used methods to prepare LCE fibers include electrospinning, dry spinning, and multi-drawdown spinning techniques [106].

Electrospinning is a well-established method of producing microfibers by spraying and stretching polymer solution under electrostatic voltage [107,108]. For instance, using this technique, He *et al.* massively produced LCE microfiber actuators within a short period (Figure 5A) [109]. When a high voltage was applied between the LCE ink and the conductive target, the solvent was evaporated, and dry LCE fibers were obtained. These fibers could then be collected from the metallic target and exposed to UV light to undertake a cross-linking reaction. The diameters of LCE microfibers could be controlled in the range of 20 to 80 μm by adjusting spinning parameters, including ink concentration, voltage, and distance between the nozzle and collector. To precisely control the actuation in the micro-scale, the LCE fibers were coated with poly-dopamine (PDA). Based on the photothermal effect, it produced rapid actuation and recovery once the NIR was applied to the LCE. Quantitatively, these LCE microfiber actuators exhibited large actuation strain (~60%), a fast response speed (< 0.2 s), and high power density (400 W kg⁻¹). These performances were all comparable to human skeleton muscle. Moreover, these LCE microfiber actuators could be used in multi-scale robotic systems, such as micro tweezers, micro swimming robots, and light-power microfluidic impellent pumps. However, this work has two limitations. One is the non-uniformity of diameters of as-spun LCE fibers. Another challenge is that the mesogens of as-spun LCE microfibers are not aligned. Fibers always require external loads to align the mesogens along the axial direction, significantly limiting their

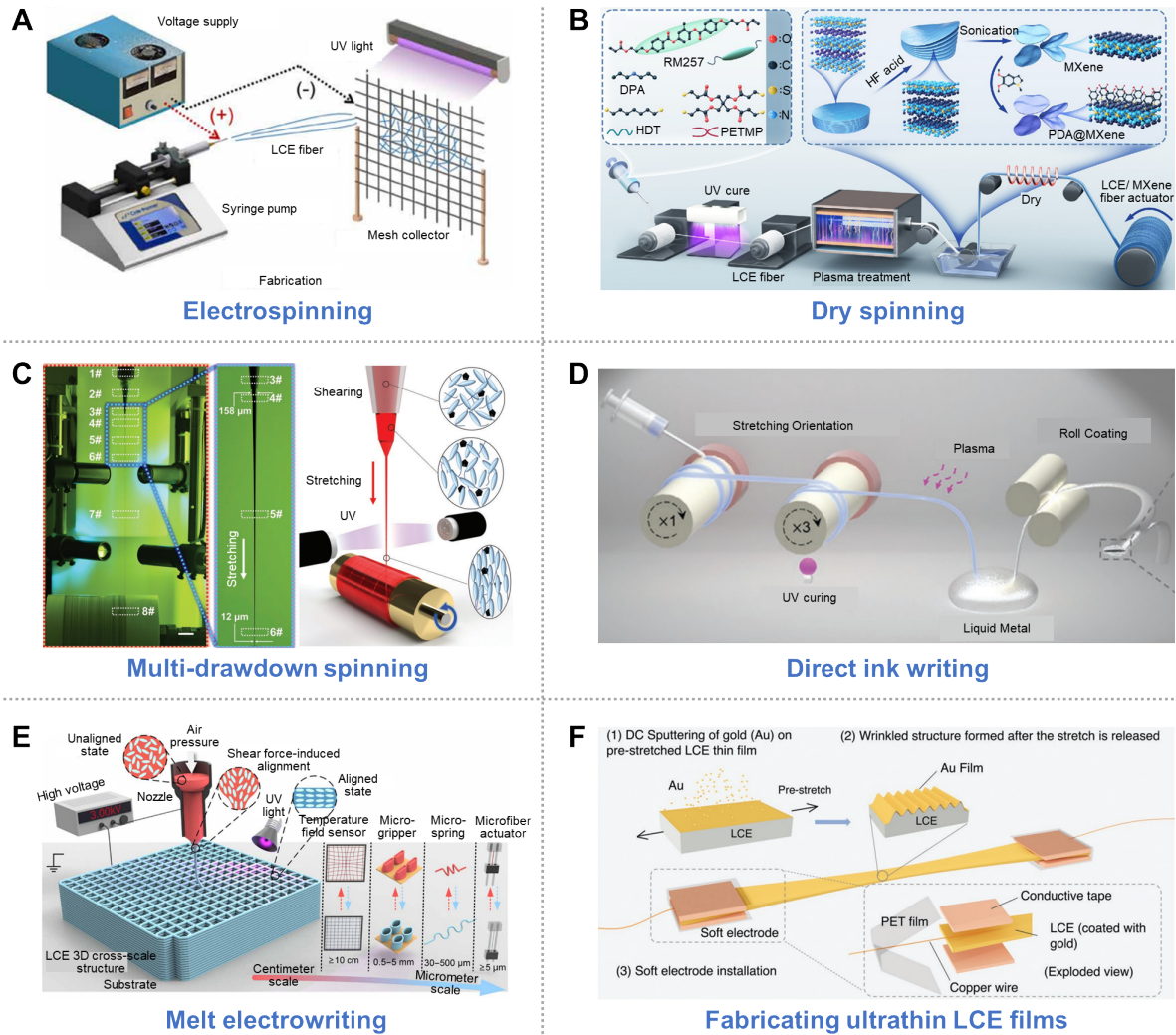


Figure 5 Different strategies to reduce the critical length scale of LCEs. (A) Schematic illustration of the electrospinning of LCE microfiber. Reprinted with permission from Ref [109]. (B) Schematic illustration of the preparation process of the PDA-modified MXene/LCE fiber by dry spinning. Reprinted with permission from Ref [111]. (C) Schematic illustration of fiber shaping and orientation using the multi-drawdown spinning technique. Scale bar: 1 mm. Reprinted with permission from Ref [112]. (D) Schematic diagram of the preparation of LM-LCE fiber via DIW printing. Reprinted with permission from Ref [117]. (E) Schematic illustration of 3D MEW-printed LCE microfibers and cross-scale structures. Reprinted with permission from Ref [121]. (F) Schematic illustration of fabricating ultrathin gold-film-coated LCE film. Reprinted with permission from Ref [122].

practical uses.

In dry spinning technology, a spinning solution is extruded into hot air through a spinneret, causing the solvent to evaporate rapidly and the material to solidify into fibers [110]. Wei *et al.* fabricated PDA-modified MXene/LCE fibers via a dry spinning technology and a two-step cross-linking strategy (Figure 5B) [111]. The LCE microfibers, as designed, possessed uniform fiber diameter ($710 \pm 27 \mu\text{m}$) and could generate large actuation strain (~60%) within 0.4 s. Furthermore, the LCE/MXene fibers could be further constructed as a smart circuit switch and a light-fueled adaptive photovoltaic smart window. The fibers produced by dry spinning are uniform, but their diameters are often large (hundreds of micrometers).

Inspired by the spiders who adopt unusual processes to produce fibers, Lv *et al.* developed a multi-

drawdown spinning technique that enabled continuous and rapid production of aligned LCE fibers with uniform controllable diameters (Figure 5C) [112]. Through internal and external drawdowns and alignments, thin, aligned LCE fibers (the minimum diameter of 2.6 μm) could be successfully prepared. Both actuation stress and mechanical properties have been significantly enhanced by incorporating graphite. It was worth mentioning that LCE's actuation stress reached 5.3 MPa, one magnitude higher than that of conventional LCE actuators and human muscle. As a result, the LCE fiber in this work exhibited high response frequency (50 Hz), unprecedented power density ($20,440 \text{ W kg}^{-1}$), and remarkable durability ($> 250,000$ cycles without breakage). The multi-drawdown spinning technology possesses the unique capability that enables continuous and high-speed production of uniform thin microfibers with the well-defined orientation of mesogens, which few processing technologies can achieve.

Fabricating LCE fibers by extrusion method

LCE precursors are highly compatible with 3D printing processes, so extrusion methods, including direct ink writing (DIW) and melt electrowetting (MEW), have also been used to fabricate LCE fibers.

The DIW technology uses a nozzle to print a single filament that can be viewed as fiber [113–115]. Qi *et al.* developed a method for fabricating reversibly actuatable LCE fibers using DIW printing (Figure 5D) [116]. The obtained fibers had a modulus of 2 MPa, 51% actuation strain, and a failure strain of well over 100%. Yang *et al.* subsequently coated LM on the LCE fibers prepared by DIW printing, achieving a programmable electrothermal actuation response [117]. Applying pulsed voltage electrical stimulation allowed LM-LCE fibers to contract at a maximum rate of $284\% \text{ s}^{-1}$ and produce a large contraction ratio of over 40%. Furthermore, the LM-LCE fibers exhibited high power density (367 W kg^{-1}) and work density (417 kJ m^{-3}).

Fiber diameters in hundreds of microns are typically achieved via DIW. On the other hand, the MEW method integrates principles from melt electrospinning (MES) with extrusion-based 3D printing technologies to accurately place ultrafine fibers and fabricate intricate polymer microstructures [118,119]. Somolinos *et al.*, for the first time, reported the successful electrowetting of photopolymerizable liquid crystal inks leading to LCE fibers with diameters ranging from several hundred nanometers to tens of microns [120]. In addition, complex fiber-based scaffolds with programmable and reversible deformation could be fabricated. In another work, using the MEW method, Wang and coworkers fabricated LCE-based microfiber actuators and various 3D actuators on the micrometer to centimeter scales with high resolutions (4.5 to 60 μm), actuation strains (10 to 55%) and a maximum work density of 160 J kg^{-1} . Moreover, based on a deep learning model, large-scale, real-time, LCE grid-based spatial temperature field sensors were designed, exhibiting a fast response time (< 42 ms) and a high precision (94.79%) (Figure 5E) [121].

Fabricating ultrathin LCE films

In addition to LCE fiber, ultrathin LCE films exhibit fast actuation and high-power density with a small thermal inertia. Wang *et al.* developed a compression-assisted mold-casting method to fabricate ultrathin LCE film (Figure 5F) [122]. The compressive loads could quantitatively control the thickness of the film. For instance, the minimal thickness of the LCE film could reach 10 μm under a load of 100 N. A thin layer of gold was sputtered on the LCE film as a heating element to trigger the actuation via an electro-thermal

mechanism. Subject to a voltage pulse, LCE thin film instantaneously contracted with remarkable 750% s⁻¹ strain rates and 1,360 W kg⁻¹ power density, several times beyond the human skeleton and insect flight muscles. Moreover, due to their electrical controlling method, the LCE thin films could be used as motor units and directly integrated into multiple dexterous artificial neuromuscular systems for rapid manipulation and closed-loop control. However, the ultrathin LCE films sacrifice the magnitude of the actuation force due to the small area of cross section. It can be improved by the assembly of multiple films.

The size reduction of LCEs significantly improves the response speed and power density. However, the output force of the LCE actuator will also be diminished. It is emerging to address this challenge for further exploration. One feasible solution is to have an assembly that arranges multiple fibers (or thin films) in parallel, forming a bundle. However, the distance between individual actuators is critical since it significantly affects the thermal process, which requires careful design.

Introducing active heating and cooling systems in LCEs

Conventional LCE-based actuators rely on passive cooling to reduce the temperature for recovery. The response time depends on the critical length of the specimen. As a result, the cooling time scale is several seconds with a thickness of millimeters, which is relatively slow. According to thermal diffusion, $\tau = \frac{d^2}{\alpha}$, the thermal time constant is proportional to the square of the characteristic length of LCE d , and inversely proportional to thermal diffusivity α (constant for LCE). Therefore, reducing the characteristic length of LCE shortens the response time. Researchers have introduced microfluidic channels to realize an active cooling method in LCE actuators with large scale (~mm). In this way, low-temperature fluids could internally and actively flow through microfluidic channels to accelerate the cooling process of LCE. He *et al.* developed a fluid-driven LCE actuator comprising two layers of LCE film and a middle layer patterned with a fluidic channel [123]. By injecting hot water (95°C) into its internal fluidic channel, the LCE actuator could contract by 40% of its initial length within 10 s. Adopting active cooling methods to reduce the temperature of the LCE actuator, cold water (23°C) was injected into the internal fluidic channel, and the LCE actuator recovered to its initial length of about 10 s, which was much faster than external and passive air cooling. However, the delamination between layers caused by weak bonding usually happened under cyclic actuation, resulting in the leakage and failure of the actuator.

The development of LCE materials offers a new opportunity to address the bonding issue. Recent work has shown that LCE with dynamic disulfide covalent bonds possesses self-healing, re-processing, and recycling capabilities. A disulfide exchange reaction occurs by heating above the reactive temperature, which could firmly bond all three layers (two thin film layers and a fluidic channel layer). As a result, the LCE actuator containing disulfide bonds can sustain the fluids with high temperatures and large flow rates without any leakages. In Figure 6A, by alternatively injecting hot (90°C) and cold water (20°C) into its internal fluid channel, it generated fast actuation as well as recovery [124]. The temperature and flow rates of the fluids regulated the actuation strain (~30%), stress (~0.3 MPa), and frequency (~1 Hz) of LCE actuators. Using the same principle, a bi-morph actuator-based soft gripper was further developed to handle objects with different features. The internal active heating and cooling system offers another opportunity for LCE to be operated in a broad range of temperatures. In a high-temperature environment, the LCE spontaneously deforms based on

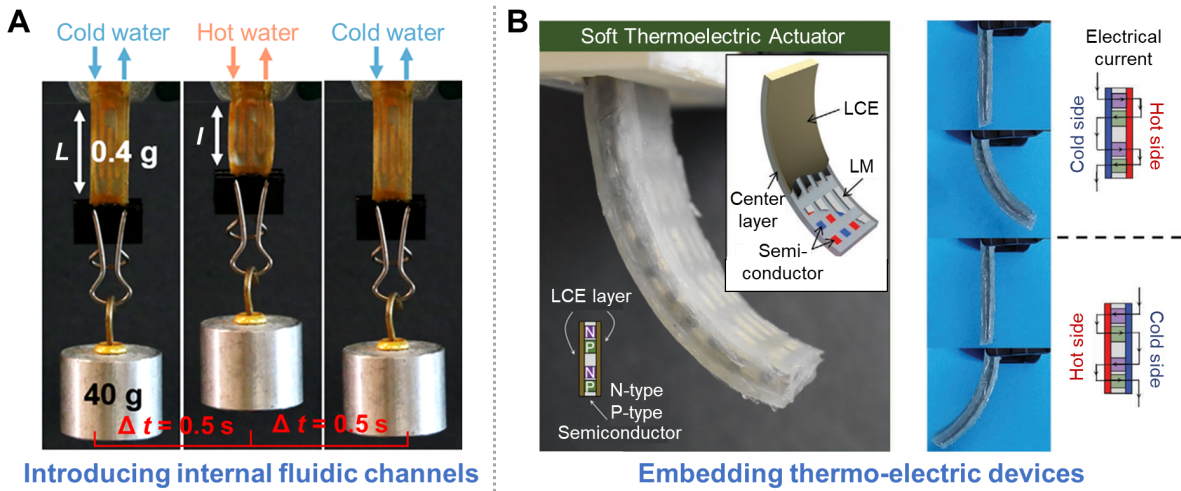


Figure 6 Internal and active heating and cooling systems in LCEs. (A) Introducing the microfluidic channel. The LCE actuator containing a microfluidic channel can produce fast actuation and recovery by alternately injecting hot water (90°C) and cold water (20°C). Reprinted with permission from Ref [124]. (B) Integrating thermal electric device. When a current is applied to a soft TED, the right side of the LCE contracts due to the heating by the Peltier effect, causing the TED-LCE to bend uniformly in the counterclockwise direction. Reversing the flow of current causes the previously heated right side to actively cool, and the left side of the device to heat up, resulting in a clockwise bending. Reprinted with permission from Ref [125].

its phase transition. Through actively injecting cold water into its internal fluidic channel, a temperature gradient will be generated, resulting in the actuation of LCE. However, the fluid regulations require multiple mechatronics, including microcontrollers, pumps, solenoid valves, and circuits. These components are quite complex and bulky, which limits the untethered actuation of LCE.

Thermoelectric devices realize the mutual conversion between thermal and electrical energy based on Peltier effects. As an electrical current passes through the devices, heat is absorbed or released at the joints of conductors, depending on the flow direction of the current. Using this principle, Majidi *et al.* embedded a soft, stretchable thermoelectric device (TED) into LCE, realizing active cooling driven by the voltages (Figure 6B) [125]. Specifically, when a current was applied to TED, a temperature gradient would be generated via the Peltier effect that caused the local contraction of LCE, resulting in the bending of the LCE in a specific direction (e.g., counterclockwise). Reversing the current flow caused the LCE to bend in the opposite direction (e.g., clockwise). The applied voltages determined the magnitude of bending angles and angular velocity of LCE. The maximum bending angle of 27° and angular velocity of 2.5° s⁻¹ could be achieved under the voltage of 2.9 V. Compared to conventional LCE bending actuators of the same size, TED-LCE is one order of magnitude faster due to its unique internal and active cooling approach. Additionally, TED provides more advanced functions, such as energy-harvesting capability.

Utilizing mechanical instability

Mechanical instability has recently been applied to diverse actuating materials for fast motion using different design principles [126]. In nature, biological structures and organisms instantaneously release their stored energy using mechanical instabilities to produce high-speed locomotion, fast grasping, and shape re-configuration. It has also been utilized to tackle the actuation speed of LCE actuators.

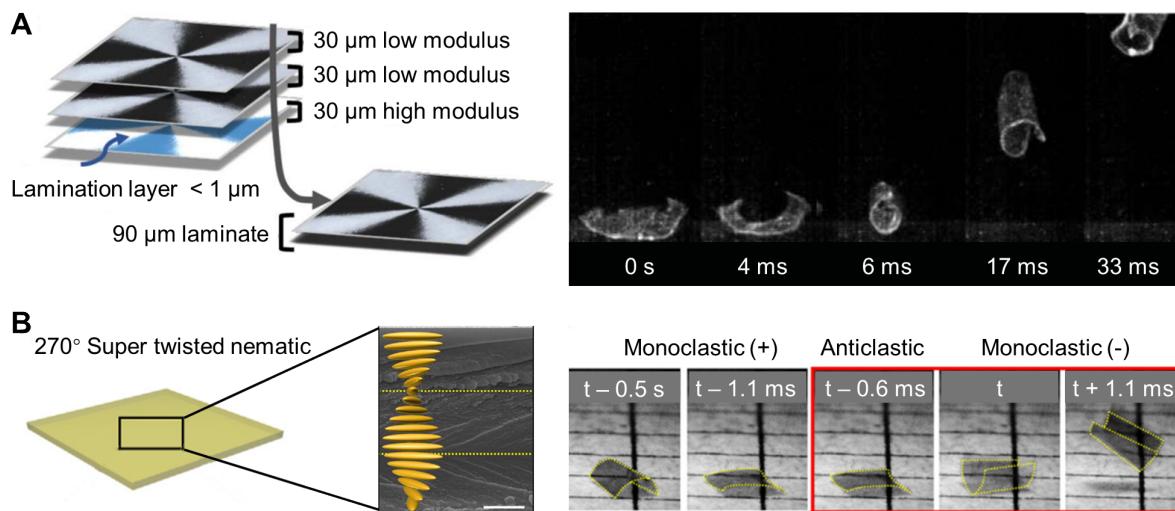


Figure 7 Utilizing mechanical instability of LCEs to achieve rapid actuation. (A) An LCE film is prepared by stacking patterned LCEs to create a modulus gradient through the thickness. Upon heating, the laminated LCE jumps from the hot surface due to a snap-through instability. Reprinted with permission from Ref [127]. (B) An LCN monolith with 270° super twisted nematic alignment of mesogens undergoes mechanical snap-through instability, resulting in the jumping motion. Scale bar: 2.5 mm. Reprinted with permission from Ref [128].

White *et al.* introduced snap-through instability into LCEs, achieving rapid jumping locomotion (Figure 7A) [127]. The LCEs with spatial variation of the nematic director of +1 topological defects were prepared, which could deform into a cone during heating. Two LCE films with different moduli were prepared by adjusting the crosslinking density. They laminated two soft LCE films with lower modulus and one stiff LCE film with higher modulus, ensuring that defect centers and edges of the squares were aligned to obtain an LCE device with the required modulus gradient. When the LCE device was placed on a hot surface (160°C) with the high modulus side downwards, only the bottom high modulus surface was initially heated and underwent strain. The film first deformed upwards into a cone, thereby storing elastic energy. Subsequently, due to the larger deformation on the lower modulus side of the upper layer, the curvature of the LCE film was reversed through mechanical instability, resulting in a snap-through instability in the center of the device, thereby forming an inverted cone. The LCE device underwent rapid jumping due to this snap-through deformation, converting the stored elastic energy into kinetic energy. The force and acceleration of this transition caused the LCE to jump from the surface in less than 6 ms, reaching a height over 200 times the material thickness and 2.5 times the width of the LCE device.

Wie *et al.* reported a continuous photomechanical jumping of azobenzene-liquid crystal polymer network (LCN) monolith with on-demand height and angle programmability (Figure 7B) [128]. The LCN monoliths were designed as the spring-like 270° super twisted nematic (STN) molecular architecture to impart macroscopic bi-stability reliably. The perpendicular molecular alignment at the top and bottom of the LCN monoliths provided for the effective accumulation of photogenerated stress from the modified energy barrier between nonisometric structures, realizing jumping by instantaneously releasing energy via snap-through. Remarkably, the maximum jumping height reached 15.5 body length (BL) with a maximum instantaneous velocity of 880 BL s⁻¹. By varying macroscopic geometry (the aspect ratio) and light intensity profile, the jumping height and angle of the LCN monoliths could be programmed. Furthermore, by selective switching

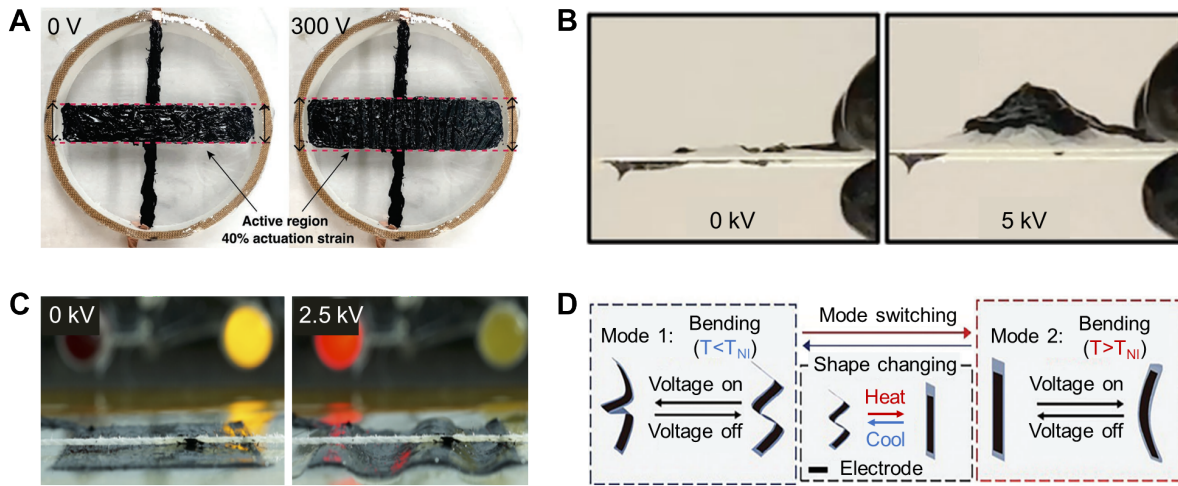


Figure 8 Developing dielectric liquid crystal elastomers. (A) The dielectric LCE film achieves fast and large actuation at a low voltage. Reprinted with permission from Ref [129]. (B) A flat DLCE film exhibits a three-dimensional cone deformation under Maxwell stress when a voltage is applied. Reprinted with permission from Ref [130]. (C) A DLCE device generates 2D to 3D shape change with nonlocal Gaussian curvature at a voltage of 2.5 kV. Reprinted with permission from Ref [131]. (D) The actuation mode switching of DLCE is based on the reversible LCE shape-changing. Reprinted with permission from Ref [132].

of the light irradiation direction according to the post-landing film curvature, continuous jumping motions could be reliably implemented with a single LCN monolith. Four continuous and directional jumping sequences were demonstrated within 5 s to overcome an obstacle. The LCE actuators based on mechanical instability for rapid actuation typically generate jumping locomotion. However, achieving more generalized behaviors in LCEs remains a challenge.

Developing dielectric liquid crystal elastomers

Dielectric elastomer actuators (DEAs) have been widely discovered due to their large actuation strain, fast response, high energy efficiency, and energy density. Multiple materials and structures have been used as DEAs, including 3M VHB tape, polydimethylsiloxane (PDMS), and hydraulically amplified self-healing electrostatic (HASEL). When subjected to high voltage (> 5 kV), a compressive Maxwell stress will be produced along the thickness direction, causing a rapid expansion of DE-film in the other two directions. Recently, an electro-mechanical actuation mechanism of dielectric liquid crystal elastomers (DLCEs) has been developed that combines the desirable characteristics of both DEAs and LCEs, exhibiting fast and highly efficient actuation and shape programmability.

Cai *et al.* developed a novel DLCE actuator that relatively low voltages could drive compared with traditional DEAs (Figure 8A) [129]. The soft elasticity and the pre-stretch ratio of DLCEs produced the desirable stress-strain relation in terms of low modulus, early strain-stiffening, and high dielectric permittivity. These properties enabled the DLCEs to generate large actuation strain (~40%) at 1 Hz under a low voltage of 300 V. Additionally, the electromechanical actuation mechanism results in higher energy conversion efficiency (20%) than thermally-driven LCEs (< 1%). Moreover, versatile rapid actuation modes can be achieved by spatially programming the alignment of mesogens and local cross-linking density of DLCE. For instance, White *et al.* demonstrated a three-dimensional cone deformation of DLCE under Maxwell

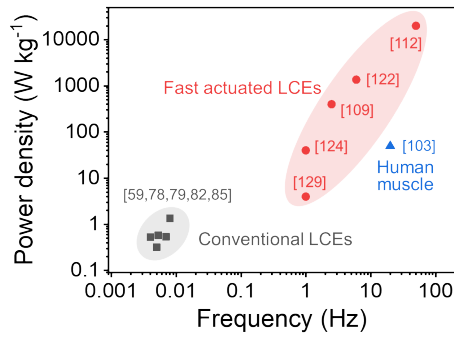


Figure 9 Comparison of frequency and power density between conventional and fast-actuated LCEs. By adopting the strategies mentioned in this paper, the response speed and power density of LCEs have been greatly improved by two orders of magnitude, which are comparable to human skeletal muscle.

stress (Figure 8B) [130]. In this example, the orientation of mesogens in DLCE was in a circumferential direction. The material was soft in the radial direction but stiff along the circumferential direction. Therefore, this DLCE expanded radially upon applying electrical potential, resulting in a 3D cone shape. In another work, Yang *et al.* also demonstrated programmable 2D to 3D shape changes (e.g., nonlocal Gaussian curvature) with large actuation strain ($> 1,800\%$) and fast response time (~ 1 s) (Figure 8C) [131]. Recently, Xie *et al.* reported a low e-field ($8 \text{ V } \mu\text{m}^{-1}$) driven multimodal DLCE actuator (Figure 8D) [132]. Specifically, the temporary and permanent shapes through the LC phase transition possessed different bending stiffness, which led to distinct actuation modes under an electric field, achieving mode-switching upon temperature change. Moreover, the temporary and permanent shapes could be reprogrammed, so the multimodal actuation could be further diversified on demand. Regarding practical applications, various DLCE-based devices have been developed recently, including artificial muscles, tunable smart lenses, and haptic devices. DLCEs can achieve rapid actuation while requiring high input voltage due to the limitation in electrostatic mechanisms.

CONCLUSION AND OUTLOOK

Figure 9 compares the fast-actuated LCE with conventional LCE actuators regarding frequency and power density. Adopting the abovementioned four strategies has greatly enhanced the response speed and power density by two orders of magnitude, comparable to human skeleton muscle. The frequency of the fast actuated LCEs can reach 1 Hz or even higher. Despite significant progress in fast-actuated LCEs, several challenges still exist.

Although the actuation performance has been enhanced, one-dimensional LCE microfibers produce limited output force. To magnify this force, multiple fibers have to be arranged in parallel to form a fiber bundle. However, the distance between individual fibers is critical as it greatly affects heat transfer. Additionally, LCE microfiber and ultrathin film can only generate contractile actuation once triggered. Converting 1D or 2D LCE in-plane contraction into diverse three-dimensional deformation is challenging. As demonstrated in previous work, one possible way is to combine LCE fiber with fabrics to induce complex 3D deformation [133,134]. For the 2D film, a feasible approach is to program the arrangement of mesogens leveraging

advanced manufacturing techniques (e.g., two-photon polymerization). Moreover, the regulation of the fluids requires external control systems. It still faces challenges in constructing an untethered robotic system for locomotion and manipulation. Furthermore, the LCE actuators based on mechanical instability for rapid actuation typically generate jumping locomotion. Achieving broader actuation in LCE remains a challenge. Lastly, DLCEs have to be activated by high voltage inputs, which requires the voltage amplifier and associated circuits to achieve.

The design of high-performance LCE actuators is a multidisciplinary subject that blends knowledge across various fields, including materials science, applied mechanics, additive manufacturing, and robotics. We hope our review could inspire soft robotics design and novel devices using LCEs.

Data availability

All data needed to evaluate the conclusions in the paper are presented in the paper.

Funding

This work was supported by the National Key Research and Development Program of China (2023YFB3812500), the “Pioneer” and “Leading Goose” R&D Program of Zhejiang (2024SSYS0082), the National Natural Science Foundation of China (52105003), the Beijing Municipal Natural Science Foundation (2222058), the CUHK Academic Equipment Funding (4937214), and CUHK Direct Grant (4055217).

Conflict of interest

The authors declare that they have no conflict of interest.

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