Research Article

Microfluidic Generation of Microsprings with Ionic Liquid Encapsulation for Flexible Electronics

Yunru Yu, Jiahui Guo, Lingyu Sun, Xiaoxuan Zhang, and Yuanjin Zhao

State Key Laboratory of Bioelectronics, School of Biological Science and Medical Engineering, Southeast University, Nanjing 210096, China

Correspondence should be addressed to Yuanjin Zhao; yjzhao@seu.edu.cn

Received 21 February 2019; Accepted 20 May 2019; Published 19 June 2019

Copyright © 2019 Yunru Yu et al. Exclusive Licensee Science and Technology Review Publishing House. Distributed under a Creative Commons Attribution License (CC BY 4.0).

Inspired by helical or spiral veins, which endow plants with excellent flexibility and elasticity to withstand storms, we present novel hollow microsprings with ionic liquid encapsulation for flexible and stretchable electronics. The microsprings were generated by using a coaxial capillary microfluidic device to consecutively spin poly(vinylidene fluoride) (PVDF) presolution and an ionic liquid, which formed laminar flows in the coaxial injection microfluidic channels. The fast phase inversion of PVDF helps to form the core-shell structure of a microfiber and guarantees the in situ encapsulation of ionic liquid. The hybrid microfiber can then spiral and be further solidified to maintain the helical structure with increasing flow rates of the injection fluids. Because of the feasible and precise control of the injection fluids during the microfluidic spinning, the resultant microsprings have controlled core-shell structures, helical pitches, and corresponding electromechanical properties. By further embedding them into stretchable films, the simple paradigm of a flexible device shows great conductive performance in tensile tests and even motion cycles, which could be explored as a promising candidate in stretchable sensors, flexible electronics, and electronic skins.

1. Introduction

Soft and stretchable conductors are essential components in soft robotics as well as wearable, conformable, and deformable electronics [1–5]. Their interconnectors can work similarly to a blood vessel network to transport electrons throughout the whole devices. In comparison with rigid electronic materials, flexible, stretchable, and lightweight conductors are more appealing as they are capable of sensing and conducting under mechanical deformations like tensile strain, bending, and torsion [6–9]. In general, these conductors with different features can be fabricated by integrating conductive agents such as metals, semiconductors, carbonaceous materials, liquid metals, and ionic liquids, into supporting elastic materials, by structural conversion of established brittle and rigid inorganic materials, or directly based on highly stretchable and deformable textile structures [10–14]. Among those conductive agents, ionic liquids have been widely used due to their nonvolatility, low surface tension, relatively low Young’s modulus, and good flexibility [15–18]. However, current methods for integrating ionic liquids are based on the sophisticated processes of channel casting, filling, and sealing, which are time-consuming and difficult to realize at small scales. In addition, the resulting ionic liquid products usually have relatively simple structures and are weak at being integrated into flexible systems with much more complex three dimensional structures. Thus, the creation of good stretching and stable conducting materials integrated with ionic liquid is still anticipated.

In this paper, we present a novel hollow microspring with ionic liquid encapsulation using a microfluidic spinning approach for flexible and stretchable conduction (Figure 1). Helical or spiral veins can endow plants with excellent flexibility and elasticity to withstand storms and these interesting helical structures have stimulated researchers to fabricate materials with such structures. To achieve this goal, many approaches, including chemical assembly, photolithography printing, mechanical twisting, and microfluidics, have been developed to generate the helical structures at nano- or microscales [19–21]. Among these approaches, microfluidic technology has become an outstanding candidate because of its ability to generate materials with various morphologies and functions [22–27]. Feasibility and complexity of channel assembly and construction, precise control on small quantities of fluids in restricted cross-sectional channels, together with various choices of fluids enable the microfluidic
spinning technology to be versatile in the creation of microfibers with complex and tunable structures at small scales [28–30]. However, the recent microfluidic spinning approach has not fabricated microsprings with ionic liquid encapsulations, and their potential value in flexible and stretchable conductors remains unexplored.

Here, we employed a coaxial capillary microfluidic device with the function of consecutive generation of hollow poly(vinylidene fluoride) (PVDF) microsprings with ionic liquid encapsulation. Because of the low Reynolds number of the PVDF presolution and ionic liquid, they formed laminar flows in the coaxial injection channels of the microfluidic device. The fast phase inversion of PVDF helps to form the core-shell structure of the microfiber and guarantees the in situ encapsulation of ionic liquid. With increased flow rate of the fluid, the microfiber could then spiral and be further solidified to maintain the helical structure. Because the spinning and spiraling process could be precisely controlled by tuning the phase flow rates and the distance between the outlet tip and the bottom of the collection container, helical microfibers with desired morphologies and microstructures can be continuously generated. As a result, a series of microsprings with different conductivities could be generated and by embedding them into flexible films, they still had great flexibility and electronic performances, showing their practical value for flexible and stretchable conductors. These features proved that the generated microsprings are highly usable in smart flexible electronics.

2. Results

In a typical experiment, a coflow microfluidic system was assembled by coaxially inserting the inner spindle capillary into the outer tapered injection capillary. The generation of core-shell structured microfibers usually depends on the fast and simultaneous gelation along both the inner and outer layers of the precursor flow in microfluidic channels. During a typical spinning process, inner and sheath fluids are introduced into the microfluidic channels to form a coaxially laminar flow due to their small Reynolds numbers. The laminar flow can then be solidified at this stage via phase inversion. The fast reaction can keep the structure of the generated microfibers the same as the laminar flows; thus the microfiber with core-shell structures can be solidified and extruded continuously. Moreover, when there is a flow rate difference between the jetting stream and the surrounding fluid, the stream deforms and begins to spiral in the collection pool. Thanks to the continuous solidification of the extruded microfibers, this helical structure can be fixed and maintained and thus helical microfibers with core-shell structures can be spun continuously.

In this research, for flexible conduction ability, an ionic liquid (1-ethyl-3-methylimidazolium tetrafluoroborate, [Emim]BF4) was chosen as the core fluid. [Emim]BF4 is a widely used electrolyte and has advantages of nonvolatility, nonflammability. The nonvolatility makes it stable during the microfluidic spinning and spiraling processes together with the following embedding and stretching applications; the nonflammability proves the safe and stable encapsulation amounts of the ionic liquids in microsprings under different weather temperature or animal heats. The electrical conduction of our microfiber depends on the free ions in the encapsulated ionic liquids, which are relevant to their amounts. Thus, the unchangeable amount could stabilize the conductivity of the microsprings. To achieve a uniform shell structure for encapsulation, the sheath phase of PVDF dissolved in N,N-dimethylformamide (DMF) was chosen as the shell material of the microspring because of the rapid phase inversion of DMF induced fast solidification. The inner ionic liquid and sheath PVDF solutions were then simultaneously introduced into the relevant microfluidic channels (Figure 2(a)). The outlet of the microfluidic device was directly immersed into the collection pool containing deionized water. Owing to the hydrodynamic focusing effect and low Reynolds number of the fluids, the immiscible ionic liquid and PVDF fluids hardly mixed with each other. When the three-dimensional coaxial flow was pumping out, the solvent DMF from the PVDF precursor was transferred to the outer deionized water to facilitate the solidification of PVDF; thus the PVDF microfiber can be spun continuously (Figure 2(b)). Because the transfer is very fast, the generated microfibers can replicate the coaxial construction of the microfluidic
channels, and the encapsulation of the ionic liquid can take place simultaneously and successfully (Figure 2(c)). As a result, the core-shell structure of the microfibers can be easily tuned by changing the inner and sheath phase flow rates, as shown in Figure 2(d). When the sheath flow rate was fixed, an increase of the inner flow rate brought about a decrease of the shell thickness; conversely, an increased sheath flow rate brought about an increased shell thickness at a fixed inner flow rate. When the increase of the sheath flow continued, the microfiber began deforming and gradually formed a helical structure with a random direction of rotation (Figure 2(e)). The spiraling of the microfiber can be ascribed to the large velocity difference between the microfiber stream and the outer relatively static liquid environment. The outer collection pool also provides enough space for the DMF to transfer to the outer liquid from all directions, which would not interfere with the formation of the helical structure (Figure 2(f)). Finally, the solid microfiber can maintain the helical and core-shell structure owing to the simultaneous process of solidification, spiraling, and encapsulation.

To confirm their structure, the generated helical microsprings were collected for optical observation and dried for scanning electron microscope (SEM) characterization (Figures 2(g) and 2(h)). It was found that they had a free-standing helical geometry and a core-shell structure at the same time. It should be mentioned that the helical microfiber could be uncoiled and recover under strains (Figure S1). Although the dried microfibers under SEM cannot show the inner ionic liquid, it was convincing that the encapsulation was a success from the uniform shell and the continuous spinning and filling process. It could also be proved by the result of infrared spectra (Figure S2). The characteristic peaks of the microfiber (1574 cm$^{-1}$ (framework of aromatic ring), 2984 cm$^{-1}$(C-H), 1185 cm$^{-1}$(C-C)) suggested the successful encapsulation during the spinning process. By tuning the flow rates of the inner and sheath flow rates and the distance between the outlet of the microfluidic device and the bottom of the collection pool, the helical pitches of the solidified microfibers can also be precisely regulated (Figures S3 and S4). It can be concluded that the increased phase flow rates and the decreased distance contributed to the decreased helical pitches of the microfibers. Thus, by tuning proper flow rates and the distance between the outlet of the device and

![Image](https://via.placeholder.com/150?text=Figure+2%3A+Microfluidic+generation+of+microfibers.+%28a-h%29+Schematic+illustration+and+real-time+microscopy+image+of+microfluidic+spinning+of+straight+microfiber+with+ionic+liquid+encapsulation,+respectively.+(c) Optical+microscopy+image+of+core-shell+structured+microfiber.+%28d%29+Relationship+between+thickness+of+straight+microfiber+and+flow+rates.+%28e-f%29+Schematic+illustration+and+real-time+microscopy+image+of+microfluidic+spinning+of+microspring+with+ionic+liquid+encapsulation,+respectively.+%28g%29+Optical+microscopy+image+of+core-shell+structured+microspring.+%28h%29+SEM+image+of+the+core-shell+structured+microspring.+Scale+bars+are+200\+\mu\text{m}+in+%28c%29+and+%28h%29,+250\+\mu\text{m}+in+%28g%29,+respectively.)
the bottom of the collection pool, the core-shell structured helical microfibers with desired morphologies could be continuously spun from this microfluidic method.

It should be noted that microfibers with different morphologies can show different conductive performance; thus their relative resistances were firstly measured. The resistance of the straight microfiber conformed to the law of resistance \( R = \frac{\rho L}{S} \), where \( \rho \) is the resistivity of the ionic liquid, \( L \) is the length of the microfiber, and \( S \) is the cross-sectional area of the inner core (Figures 3(a) and 3(b)). The results directly indicated that the longer microfiber showed a larger resistance while a microfiber with a thinner shell showed a smaller resistance. When the shell thickness was fixed, the cross-sectional area was a constant and the increase of the length would bring about a larger resistance; when the length and the diameter of the microfiber remained unchanged, a decrease of the shell thickness resulted in an increased cross-sectional area of the liquid core, which results in a smaller resistance. The smaller resistance corresponded to the better conductivity of the microfiber. This also occurred on helical microfibers with different lengths and helical pitches (Figures 3(d) and 3(e)). For example, the resistance of a microspring with diameter of 100 \( \mu \text{m} \), length of 2 cm, and helical pitch of about 1200 \( \mu \text{m} \) is about 1.16 M\( \Omega \) and could be adjusted by further changing its diameter, length, and helical pitch (Figure S5). The resistance of the helical microfiber with a longer length and a smaller helical pitch will be even larger. There are many flexible, bendable, and stretchable textiles that exhibit lower resistance values and they could perform well in sensing applications. However, few of these filaments are generated from such a simple process by introducing conductive agents into microfluidic channels. Thus, the microfluidic generation method will offer inspirations to fabricate conductive microfibers. Also, it is believed that after optimization, the versatile microfluidic technology could generate ionic liquid encapsulated microsprings with much smaller resistance in the near future.

When the fabricated conductive microfibers were stretched, they were also able to respond, as explained by their obvious resistance change as a function of strain. The stiffness of the core-shell structured microfibers was first tested before recording their relative resistance change (Figure S6). It could be easily inferred from the curve that the straight microfiber can only achieve a stretchability of 80%. During the stretching process, the relative resistance change of the straight microfiber \( R/R_0 \), where \( R \) is the resistance in

![Figure 3: Conductivity performance of microfibers. (a) Relationship between relative resistance and the thickness of straight microfiber. (b) Relationship between relative resistance and the length of straight microfiber. (c) Relative resistance change when stretching the straight microfiber. (d) Relationship between relative resistance and the length of microspring. (e) Relationship between relative resistance and the helical pitch of microspring. (f) Relative resistance change when stretching the microspring with different helical pitches.](image-url)
real time and $R_0$ is the original resistance) was recorded, as shown in Figure 3(c). The resistance of the 50% stretched microfiber rose to twice as much as that in the original state. The underlying mechanism might lie in the deformation of the core ionic liquid encapsulated channel during the microfiber stretching process. Because the stretching process would cause an elongation of the microfiber and a shrinking of the cross-sectional area due to the Poisson ratio, the resistance underwent an obvious increase during this process. However, the increase of the resistance was not as much in the helical microspring stretching process, especially in those microsprings with smaller helical pitches (Figure 3(f)). This is because the elongation was minimal and negligible at the very beginning of the stretching process; the resistance change mainly depends on the variation of the cross-sectional area of the core ionic liquid encapsulated channel. Until the microspring was stretched to a straight, the increased strain would cause an elongation of the microfiber together with a shrinking of the cross-sectional area, bringing about a remarkable increase in the relative resistance change. Because the microspring with smaller helical pitch had a larger full straight length, the conduction was more reliable during the stretching compared with those microsprings with larger helical pitches. This indicated that the helical microsprings showed enhanced robustness and stretchability in electron conduction under deformation compared with the straight microfibers. The helical structure is the outcome of making full use of the space; thus the small diameter and helical structure guarantee the stable conduction when bended or stretched to a large extent. They will perform more flexibly than conventional bulk materials and are more likely to adapt strict situations like narrow gaps or flexible joints if they were constructed into more complex structures and shapes. They are thus promising candidates for applications in flexible electronics.

To explore the practical value of the generated conductive microsprings, they were then embedded in a flexible polymer film as a paradigm to prototype stretchable electronics (Figure 4(a)). By aligning the helical microfibers as flexible and conductive lines, covering a thin layer of polymer film and curing it, flexible electronics can be achieved (Figures 4(b)-4(d)). The thickness of the film could be adjusted to fit situations requiring thin cross-sections due to the adjustable radius of the microspring by tuning flow rates in microfluidic spinning process. Because the polymer is insulating, the hybrid film could then provide electrical conductivity along the microfiber and insulation along other directions to avoid short-circuiting during further applications. For those embedded microfibers with different helical pitches, their relative resistance change during the stretching process was also studied (Figure 4(e) and Figure S7). The results were nearly consistent with those of a single helical microfiber, which proved that the composition would not interfere with the conductivity of the helical conductors. Thanks to the good stretchability of the film and the helical microfiber itself (Figure S8), plus the low Young's modulus ionic liquid core, the strain-sensing ability functioned steadily even after 18 repeating cycles of stretching and recovering, which can be shown in the cycled stress-strain test (Figure 4(f)).

It also suggested that the maximum bending angle of the film was about 130°. Because the helical microfibers were embedded in the film, the elastomeric layer could prevent the embedded microspring from irrecoverable transformations under cycled surpass strains and there was no possibility for the ionic liquid leaking; thus the stability of the helical microwires inside and the potential that the composite could be reused and perform like the original was convinced. This hypothesis can be verified from the cycle test of the relative resistance change of the composite (Figure 4(g)). All of the flexible electronics were treated with repeated strains for 15 times, and the results showed that they can respond reproducibly and reliably to those strains, indicating an outstandingly stable conductivity for practical applications. Additionally, because the ionic liquid is nonvolatile and nonflammable, the complex film could conduct stably at different temperatures (Figure S9).

Currently, smart flexible devices can display regular motions in real time. An increase in the signal-to-noise ratio is always an object of optimization because of the nonnegligible background noise, which may come from the resistance change of the interconnectors. Thus, the generated ionic liquid encapsulated microsprings showing stable conductivities would be promising candidates in these flexible electronics. To investigate this, the integrated flexible film of the microsprings was employed to detect finger, wrist, and elbow motions, which are common human motions (Figure 5). The film was first stuck on a finger of a glove and worn by a volunteer, and the two terminals containing the integrated microsprings were connected to a digital multimeter, which can record the resistance in real time (Figure 5(a)). When the volunteer bent and straightened his or her finger, the resistances showed corresponding increases and decreases in a very small range and there appeared no changes in a general view (Figure 5(d) and Figure S10(d)). It should also be noted that the resistance changes could manifest the different degree of the finger bending (Figures S10(a)-S10(b)), but there still appeared no changes in the general view. In addition to the fingers, the wrist and elbow are two of the most widely detected motion parts in flexible electronic studies (Figures 5(b), 5(c) and Figure S10(c)). By sticking the integrated film on a wrist or elbow support, the motions can be studied by the corresponding resistance changes (Figures 5(e), 5(f) and Figure S10(e), S10(f)). When showing real-time increases and decreases in a small range, it can still be found that the resistance almost stayed the same in the general view, which indicated that the microsprings can offer reliably relative conductions in active situations. This was because although bending the finger, wrist, and elbow would strain the film together with the inside integrated microsprings, the deformation at this stage was minimal and would not cause obvious change in resistance of the helical conductors benefiting from their excellent three-dimensional structures. As a result, these microsprings with stable conductivities could then be applied in strain sensors, wearable devices, and flexible electronic fields to decrease the background noise and thus increase the signal-to-noise ratio.
3. Discussion

In summary, we have generated conductive microsprings by using a facile microfluidic spinning approach. The coaxial aligning of the injection capillaries allowed for the formation of the helical microfibers with core-shell structures, which could encapsulate the ionic liquid as conductive agents.受益于对微流体通道中相的简单操纵，因此壳层厚度、螺旋距离和因此电磁性质可以轻松调整。Microfluidic生成过程引入了微流体通道中的导电液体，这在以前的研究中很少涉及。此外，人们相信这种螺旋微纤维可以制造出电阻值非常小的导电微纤维。下一步，受导电螺旋微纤维与可拉伸薄膜相结合的启发，开发了简单可伸缩电子学概念，展示其导电稳定性与可重复性。这些导电螺旋微纤维在不同运动测试中的适应性表明，导电螺旋微纤维是伸缩传感器和柔性电子学的有希望的候选者。此外，由于可选的材料组成，导电螺旋微纤维与离子液体的封装将非常适用于不同应用领域。构建到更复杂结构中，
4. Materials and Methods

4.1. Materials. Ionic liquid (1-ethyl-3-methylimidazolium tetrafluoroborate, EMIMBF4) was bought from Aladdin. Polyvinylidene fluoride (PVDF, Mw=534000) was from Sigma-Aldrich. N, N-dimethylformamide (DMF) was purchased from Sinopharm Chemical Reagent Co., Ltd (Shanghai, China). Solutions were all filtered before being pumped into glass microcapillary devices. Ecoflex® 00-30 was purchased from Smooth-On, Inc (Macungie, PA). Water with a resistivity of 18.2 MΩ•cm⁻¹ was acquired from a Millipore Milli-Q system. All other chemical reagents were of the best grade available and used as received.

4.2. Microfluidics. The capillary microfluidic device was constructed by coaxially aligning one glass capillary with a spindle tip and one with a tapered tip on glass slide. Two capillaries with outer and inner diameters of 1.0 mm and 800 μm (World Precision Instruments) were firstly tapered and sanded to the desired orifice. The orifice of the spindle tip was about 80 μm and that of the tapered tip was about 100 μm. The spindle capillary was then assembled coaxially into the tapered one as the inner flow channel. A transparent epoxy resin was used to seal the capillaries where necessary.

The inner phase was ionic liquid (1-ethyl-3-methylimidazolium tetrafluoroborate, EMIMBF4) and the sheath fluid is PVDF dissolved in DMF at a volume-to-volume concentration of 1:10. A glass vial containing deionized water was put under the outlet of the microfluidic device keeping the liquid covering its tip. The inner flow rates could be changed in 0.1-0.5 mL/h, and the sheath flow rates could be 2-7 mL/h to generated microsprings with different morphologies.

4.3. Preparation of Flexible Films Integrated with Conductive Microspring. Firstly the generated microfibers were cut into desired lengths, which were longer than the final tested length to prevent the leaking of the inner ionic liquid. PDMS is also one of the widely used and outstanding elastomeric matrixes, however, in the case of the microfiber with extremely small helical pitch, and could be stretched more than 120%, in which PDMS may not be qualified. As a result, Ecoflex® 00-30 was chosen to produce the elastomeric layer. The two compartments of Ecoflex® 00-30 were mixed in equal volume and homogenized, and the mixture was uniformly coated on the flexibly arranged microfibers. The composites were solidified at room temperature for about 1 hour.

4.4. Characterization. The microfluidic spinning process was observed by a fast camera (F032B, Pike, Germany). Bright-field microscopic images were snapped by a stereomicroscope (JSZ6S, Jiangnan novel optics) equipped with a CCD camera.
silver glues, covered by conductive tapes, and then connected. The cross-sectional areas of the films were first coated with the same amount of silver glues to increase the contact area with the probe. A vernier caliper was used to record the strain of the tested microfibers during the stain process. The real-time resistance change was recorded by a digital multimeter (KEITHLEY, USA). Similar to the stain process, the conductivity tests of the generated microfibers were carried out by using a traditional two-probe technique on a semiconductor characterization system (4200-SCS, KEITHLEY). Because the cross-sectional areas of the microfibers were small, the two ends of the microfibers were all firstly coated with the same amount of silver glues, covered by conductive tapes, and then connected to the digital multimeter.

Conflicts of Interest
The authors declare no competing financial interests.

Authors’ Contributions
Yuanjin Zhao conceived the idea and designed the experiment; Yunru Yu carried out the experiments; Yunru Yu and Yuanjin Zhao analysed data and wrote the paper; Jiahui Guo, Lingyu Sun and Xiaoxuan Zhang contributed to scientific discussion of the article.

Acknowledgments
This work was supported by the National Key Research and Development Program of China [2017YFA0700404], the NSAF Foundation of China [Grant No. U1530269], the Natural Science Foundation of Jiangsu [Grant No. BE2018707], the Scientific Research Foundation of Southeast University, and the Scientific Research Foundation of the Graduate School of Southeast University [Grant No. YBJJ1779].

Supplementary Materials
Figure S1. The uncoiling and recovery process of microspring. Figure S2. The infrared spectra of ionic liquid, PVDF, and the ionic liquid encapsulated microspring. Figure S3. Relationship between flow rates and helical pitch of microspring. Figure S4. Relationship between height and helical pitch of microspring. Figure S5. The U-I curve of a helical microspring. Figure S6. Stress-strain test of the core-shell structured fiber. Figure S7. Stretching process of the flexible film integrated with microspring. Figure S8. Stress-strain test of the flexible film integrated with microspring. Figure S9. Relationship between the resistance of the flexible film and the temperature. Figure S10. Conductivity response to various human motions in real time. (Supplementary Materials)

References
[1] B. C.-K. Tee, A. Chortos, A. Berndt et al., “A skin-inspired organic digital mechanoreceptor,” Science, vol. 350, no. 6258, pp. 313–316, 2015.
[2] D. P. Dubal, N. R. Chodankar, D.-H. Kim, and P. Gomez-Romero, "Towards flexible solid-state supercapacitors for smart and wearable electronics," Chemical Society Reviews, vol. 47, no. 6, pp. 2065–2129, 2018.
[3] M. Su, F. Li, S. Chen et al., “Nanoparticle based curve arrays for multirecognition flexible electronics,” Advanced Materials, vol. 28, no. 7, pp. 1369–1374, 2016.
[4] W. Xu, Z. Lu, X. Sun, L. Jiang, and X. Duan, “Superwetting electrodes for gas-involving electrocatalysis,” Accounts of Chemical Research, vol. 51, no. 7, pp. 1590–1598, 2018.
[5] J. Jiang, B. Bao, M. Li et al., “Fabrication of transparent multilayer circuits by inkjet printing,” Advanced Materials, vol. 28, no. 7, pp. 1420–1426, 2016.
[6] B. Zhang, J. Lei, D. Qi et al., “Stretchable conductive fibers based on a cracking control strategy for wearable electronics,” Advanced Functional Materials, vol. 28, no. 29, Article ID 1801683, 2018.
[7] C. Yang, Z. Liu, C. Chen et al., “Reduced graphene oxide-containing smart hydrogels with excellent electro-response and mechanical property for soft actuators,” ACS Applied Materials & Interfaces, vol. 9, no. 18, pp. 15758–15767, 2017.
[8] N. Hu, T. Wang, H. Wan et al., “Synchronized electromechanical integration recording of cardiomyocytes,” Biosensors and Bioelectronics, vol. 117, pp. 354–365, 2018.
[9] H. Liu, M. Li, C. Ouyang, T. Lu, F. Li, and F. Xu, “Biofriendly, stretchable, and reusable hydrogel electronics as wearable force sensors,” Small, vol. 14, no. 36, Article ID 1801711, 2018.
[10] W. Zeng, L. Shu, Q. Li, S. Chen, F. Wang, and X.-M. Tao, “Fiber-based wearable electronics: a review of materials, fabrication, devices, and applications,” Advanced Materials, vol. 26, no. 31, pp. 5310–5336, 2014.
[11] W.-W. Liu, X.-B. Yan, J.-W. Lang, C. Peng, and Q.-J. Xue, “Flexible and conductive nanocomposite electrode based on graphene sheets and cotton cloth for supercapacitor,” Journal of Materials Chemistry, vol. 22, no. 33, pp. 17245–17253, 2012.
[12] L. Guan, A. Nilghaz, B. Su, L. Jiang, W. Cheng, and W. Shen, “Stretchable-fiber-confined wetting conductive liquids as wearable human health monitors,” Advanced Functional Materials, vol. 26, no. 25, pp. 4511–4517, 2016.
[13] R. Matsuzaki and K. Tabayashi, “Highly stretchable, global, and distributed local strain sensing line using gainsn electrodes for wearable electronics,” Advanced Functional Materials, vol. 25, no. 25, pp. 3806–3813, 2015.
[14] K. S. Egorova, E. G. Gordeev, and V. P. Ananikov, “Biological activity of ionic liquids and their application in pharmaceutics and medicine,” Chemical Reviews, vol. 117, no. 10, pp. 7132–7189, 2017.
[15] Y. Wang, S. Gong, S. J. Wang, G. P. Simon, and W. Cheng, “Volume-invariant ionic liquid microbands as highly durable wearable biomedical sensors,” Materials Horizons, vol. 3, no. 3, pp. 208–213, 2016.
[16] S. Sun, Z. Duan, X. Wang et al., “Cheap, flexible, and thermal-sensitive paper sensor through writing with ionic liquids containing pencil leads,” ACS Applied Materials & Interfaces, vol. 9, no. 34, pp. 29140–29146, 2017.

[17] D. G. Seo and H. C. Moon, “Mechanically robust, highly ionic conductive gels based on random copolymers for bending durable electrochemical devices,” Advanced Functional Materials, vol. 28, no. 14, Article ID 1706948, 2018.

[18] S. Chen, H. Liu, S. Liu et al., “Transparent and waterproof ionic liquid-based fibers for highly durable multifunctional sensors and strain-insensitive stretchable conductors,” ACS Applied Materials & Interfaces, vol. 10, no. 4, pp. 4305–4314, 2018.

[19] Y. Forterre and J. Dumais, “Generating helices in nature,” Science, vol. 333, no. 6050, pp. 1715-1716, 2011.

[20] N. Hu, T. Wang, H. Wan et al., “Hierarchically arranged helical fiber actuators driven by solvents and vapors,” Nature Nanotechnology, vol. 198, no. 12, pp. 1077–1084, 2015.

[21] Y. Yu, L. Shang, W. Gao, Z. Zhao, H. Wang, and Y. Zhao, “Microfluidic lithography of bioinspired helical micromotors,” Angewandte Chemie International Edition, vol. 56, no. 40, pp. 12127–12131, 2017.

[22] X. Hou, Y. S. Zhang, G. T. Santiago et al., “Interplay between materials and microfluidics,” Nature Reviews Materials, vol. 2, no. 5, Article ID 17016, 2017.

[23] L. Shang, Y. Cheng, and Y. Zhao, “Emerging droplet microfluidics,” Chemical Reviews, vol. 117, no. 12, pp. 7964–8040, 2017.

[24] W. Wang, M.-J. Zhang, and L.-Y. Chu, “Functional polymeric microparticles engineered from controllable microfluidic emulsions,” Accounts of Chemical Research, vol. 47, no. 2, pp. 373–384, 2014.

[25] T. Y. Lee, T. M. Choi, T. S. Shim, R. A. M. Frijns, and S.-H. Kim, “Microfluidic production of multiple emulsions and functional microcapsules,” Lab on a Chip, vol. 16, no. 18, pp. 3415–3440, 2016.

[26] H. Wang, Z. Zhao, Y. Liu, C. Shao, F. Bian, and Y. Zhao, “Biomimetic enzymes cascade reaction system in microfluidic electrospray microcapsules,” Science Advances, vol. 4, no. 6, Article ID eaat2816, 2018.

[27] S. Lee, T. Y. Lee, E. Amstad, and S.-H. Kim, “Microfluidic production of capsules-in-capsules for programed release of multiple ingredients,” Advanced Materials Technologies, vol. 3, no. 5, 2018.

[28] L. Shang, Y. Wang, Y. Yu et al., “Bio-inspired stimuli-responsive graphene oxide fibers from microfluidics,” Journal of Materials Chemistry A, vol. 5, no. 29, pp. 15026–15030, 2017.

[29] J. Wang, M. Zou, L. Sun et al., “Microfluidic generation of Buddha beads-like microcarriers for cell culture,” Science China Materials, vol. 60, no. 9, pp. 857–865, 2017.

[30] Y. Yu, F. Fu, L. Shang, Y. Cheng, Z. Gu, and Y. Zhao, “Bioinspired helical microfibers from microfluidics,” Advanced Materials, vol. 29, no. 18, Article ID 1605765, 2017.