Nanogenerator-Based Self-Charging Energy Storage Devices

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HIGHLIGHTS

• The progress of nanogenerator-based self-charging energy storage devices is summarized.
• The fabrication technologies of nanomaterials, device designs, working principles, self-charging performances, and the potential application fields of self-charging storage devices are presented and discussed.
• Some perspectives and problems that need to be solved are described.

ABSTRACT One significant challenge for electronic devices is that the energy storage devices are unable to provide sufficient energy for continuous and long-time operation, leading to frequent recharging or inconvenient battery replacement. To satisfy the needs of next-generation electronic devices for sustainable working, conspicuous progress has been achieved regarding the development for nanogenerator-based self-charging energy storage devices. Herein, the development of the self-charging energy storage devices is summarized. Focus will be on preparation of nanomaterials for Li-ion batteries and supercapacitors, structural design of the nanogenerator-based self-charging energy storage devices, performance testing, and potential applications. Moreover, the challenges and perspectives regarding self-charging energy storage devices are also discussed.

KEYWORDS Nanomaterial; Nanogenerator; Energy storage device; Self-charging
1 Introduction

With the rapid development of economy and society, microelectronic devices are playing an increasingly important role in our daily lives. Usually, these devices can be powered using lithium-ion batteries or supercapacitors, which require external power sources to periodically charge them due to their limited capacities [1–3]. Moreover, it will cost a significant quantity of manpower, financial resources, and time, especially in remote areas. For power supply, scientists are researching new methods of scavenging clean energies from the surrounding environment [6–10]. Based on this background, the piezoelectric and triboelectric nanogenerators were invented by Wang et al. in 2006 and 2012, respectively [4, 5], which can effectively convert small mechanical energy into electrical energy in the ambient environment, such as wind energy [6, 7], wave energy [8], droplet energy [9], and other mechanical energies [10]. They are clean or wasted energies in our surrounding environment. Nanogenerators not only can effectively scavenge the mechanical energies mentioned above, but also have several advantages such as simple, small, light, low cost, no auxiliaries, and convenient. They can be applied to wireless sensors and microelectronics devices. Currently, research and development of self-powered electronic devices have become a hot topic among scientists [11–28]. In particular, remarkable progress has been made in the field of self-charging power textile for wearable electronics [29–33]. Thus, it is important to investigate self-charging energy storage devices that can effectively integrate energy harvesting and storage units in one device for powering some small electronic devices with sustainable energy supply.

This review focuses on the progress of nanogenerator-based self-charging energy storage devices in recent years. The fabrication technologies of nanomaterials, device designs, working principles, self-charging performances, and the potential application fields of self-charging storage devices are presented and discussed here. Moreover, some perspectives and problems that need to be solved are also described, which can pave the path for practical applications.

2 Nanomaterials

Due to the large specific surface area and excellent energy storage characteristics, nanomaterials demonstrate a reversible capacity higher than that of the commercial products. New nanomaterials have fundamental advancements regarding energy storage and conversion devices, both of which are important to satisfy the challenges of the finite nature of fossil fuels and environmental problems. Over these years, scientists have conducted a wide range of research and achieved a series of experimental progresses.

Nanomaterials can not only be used as positive and negative electrode materials, but also be used as special electrolytes or separators. In 2017, He et al. fabricated a novel all-solid-state self-charging power cell (SCPC) using mesoporous polyvinylidene difluoride (PVDF)–LiPF$_6$ film as piezoelectrolyte [11]. The morphology and microstructure of SiO$_2$ and PVDF film are presented in Fig. 1a, demonstrating that the average diameters of SiO$_2$ are approximately 200 nm, and the pore size and thickness of PVDF film prepared using SiO$_2$ as template are 200 nm and 1 μm, respectively. Figure 1b presents that a PVDF–PZT (Lead zirconate titanate) nanocomposite film and multi-walled carbon nanotubes (MWCNTs) have been used as a piezo-separator and negative material in SCPC [12]. It can be distinctly observed that the thickness of the PVDF–PZT
A nanocomposite film is approximately 60–110 µm; the MWCNTs have lengths of 5–15 µm and diameters of 40–60 nm, respectively. Kim et al. designed a SCPC consisting of LiCoO₂ nanomaterials as the cathode and artificial graphite as the anode with a porous ZnO-free PVDF separator [13], as illustrated in Fig. 1c. Figure 2a displays that the nano-polyaniline/carbon nanotube (PANI/CNT) and nano-polyaniline (PAn)/CNT were utilized to fabricate a hybrid electric power device (HEPD) [14]. The porous multilayer structures of laser-induced

![Fig. 1](image-url)
graphene (LIG) electrodes were used to fabricate the TENG and micro-supercapacitor, as depicted in Fig. 2b [15]. As illustrated in Fig. 2c, ZnO nanowire (NW) arrays and PVDF mesoporous nanostructured films have been used in a piezo-driven SCPC [16]. Ramadoss et al. fabricated a piezoelectric effect-driven self-charging supercapacitor power cell (SCSPC) using MnO$_2$ NWs as positive and negative electrodes (Fig. 2d) and a PVDF–ZnO film as a separator [17]. Luo et al. successfully applied 3D Au@MnO$_2$ nanocomposites into the transparent and flexible self-charging power film (SCPF), as exhibited in Fig. 2e [18]. Figure 2f displays the morphology of LiMn$_2$O$_4$ precursor NWs and carbon NWs by electrospinning and annealing calcinations, where the LiMn$_2$O$_4$ precursor NWs exhibit smooth surfaces with diameters ranging from 150 to 260 nm and lengths of approximately

Fig. 2  a SEM images of PANI/CNT, unmodified or PtNP-modified PPy/CNT surfaces. Reprinted with permission from Ref. [14]. b Cross-sectional SEM image of a double-sided laser-engraved PI substrate with both sides of LIG, SEM image of the LIG thin film, and HRTEM image obtained from the edge of a LIG flake. Reprinted with permission from Ref. [15]. c Cross-sectional SEM images of ZnO NW arrays and PVDF mesoporous nanostructured film. Reprinted with permission from Ref. [16]. d FESEM image of MnO$_2$ nanostructure. Reprinted with permission from Ref. [17]. e SEM images of the 3D Au@MnO$_2$ nanocomposites (scale bar, 500 nm). Reprinted with permission from Ref. [18]. f SEM images of the LiMn$_2$O$_4$ precursor NWs and the obtained carbon NWs. Reprinted with permission from Ref. [19]
The polyvinylpyrrolidone NWs were oxidized under air atmosphere and then were carbonized under Ar protection at 850 °C to obtain conductive carbon NWs as the anode of Li-ion batteries (LIBs) [19]. Zhao et al. developed a new strategy for fabricating a metal–organic framework (MOF) for the template-directed growth of hierarchically well-oriented NW arrays based on carbon nanotube fibers (CNTFs) for electrochemical supercapacitors, where the corresponding SEM images of the MOFs, MOF derivatives, and CNTFs materials are illustrated in Fig. 3a [20]. Researchers have prepared CuO nanoflake arrays, hierarchical NiCoOH nanoplies and reduced graphene oxide (RGO) nanosheets for energy storage devices. As displayed in

**Fig. 3**  
(a) SEM images of ZnCo$_2$O$_4$@Zn-Co-S HA, as-prepared CNTFs@Zn/Co-Zif, ZnCo$_2$O$_4$@ZnCo$_2$O$_4$ HA, CNTFs@H-Co$_3$O$_4$ NA, CNTFs@H-Co$_3$O$_4$@ZIF-67 HA, and CNTFs@H-Co$_3$O$_4$@CoNC HA. Reprinted with permission from Ref. [20].  
(b) FESEM images of CuO nanoflake arrays grown on Cu foil, hierarchical NiCoOH nanoplies electrodeposited on CuO@Cu foil and RGO nanosheets. Reprinted with permission from Ref. [21].  
(c) SEM images of a CNF-coated yarn and a PEDOT: PSS/CNF-coated yarn (scale bar, 10 μm). Reprinted with permission from Ref. [22].  
(d) SEM images of the TiO$_2$ nanotube arrays on the Ti wire (scale bar, 100 nm). Reprinted with permission from Ref. [23].
Fig. 3b, a uniform and homogeneous growth of CuO nanoflake arrays with an average thickness of 35–45 nm on Cu foil substrate and the 3D architectures of NiCoOH nanoplates with an average wall thickness of 50–70 nm were successfully deposited on CuO@Cu foil, and the wrinkled sheets of RGO with an average thickness of 30–40 nm were evident [21]. The all-solid-state symmetric yarn SC was fabricated using dip-coating carbon nanofiber (CNF) and poly(3,4-ethylenedioxythiophene)-poly-(styrenesulfonate) (PEDOT: PSS) on a carbon fiber (CF) bundle, as depicted in Fig. 3c [22]. Wen et al. used highly ordered TiO$_2$ nanotube arrays in F-DSSC, as illustrated in Fig. 3d [23]. The SCPC was fabricated using PVDF/0.5(Ba$_{0.7}$Ca$_{0.3}$)TiO$_3$–0.5Ba(Zr$_{0.2}$Ti$_{0.8}$)O$_3$ (BCT–BZT) nanocomposite as piezo-separator [36]. Two-dimensional materials have excellent properties, such as open ion diffusion channels and abundant active surfaces that enable fast transport and storage ions. Therefore, it is very important for self-charging energy storage devices. Pazhamalai et al. fabricated the SCSPC using 2D-MoSe$_2$ nanosheets as the electrodes, PVDF-co-HFP/TEABF$_4$-based ionogel as the electrolyte, and the electrospun PVDF/NaNbO$_3$ nanofibrous mat as piezo-separator [37].

3 Li-ion Batteries and Supercapacitors

Currently, LIBs and supercapacitors are widely utilized as the main electrochemical energy storage devices. They can be used as the energy supply units for powering mobile phones, personal wearable devices, micro-electronic devices, etc. The reported self-charging energy storage devices are mainly based on LIBs and supercapacitors. These devices can collect and convert mechanical energy into electric energy in the surrounding environment, and then store the scavenged energy as chemical energy. Energy scavenging function of the devices can be realized by piezoelectric nanogenerators or triboelectric nanogenerators. Figure 4a–d depicts the self-charging energy storage devices based on the piezoelectric effect [11, 17, 21]. The mesoporous PVDF–LiPF$_6$ film (Fig. 4a, b) and PVDF–ZnO film (Fig. 4c) are used in the device, where a PVA–KOH gel electrolyte-soaked perforated fish swim bladder (Fig. 4d) was used as piezoelectric separator. Meanwhile,
the flexible self-charging energy storage devices using piezoelectric nanogenerator have been developed. Yuan et al. reported a paper-based flexible SC using PANI/Au/paper as electrodes, which can be charged by a piezoelectric generator [38]. The voltage of six SCs connected in series can be charged to 2.6 V in approximately 11 h; they could power a blue LED for 5 min. As presented in Figs. 5a–f and 6a–d, the LIBs and supercapacitors were based on triboelectric nanogenerators [18–28]. Figure 5a displays the schematic diagrams of all-solid-state transparent and flexible supercapacitors (TFSCs) based on interdigital electrodes of 3D Au@MnO2 nanocomposites, which were connected to form an array on the backside of the nanogenerator [18]. The flexible LIBs based on electrospun LiMn$_2$O$_4$ NWs as cathode and carbon NWs as anode are presented Fig. 5b [19].

Dong et al. fabricated the all-solid-state symmetric yarn supercapacitor (SC) using dip-coating carbon nanofiber (CNF) and poly(3,4-ethylenedioxythiophene)-poly(styrenesulfonate) (PEDOT:PSS) on a carbon fiber (CF) bundle, as presented in Fig. 5d [22]. Figure 5e illustrates that the fiber-shaped SCs (F-SCs) were fabricated using RuO$_2$·xH$_2$O-coated carbon fibers and PVA/H$_3$PO$_4$ gel electrolyte [23]. Gao et al. reported a novel strategy of implanting a solid Li-ion battery (SLB) into a triboelectric fluorinated ethylene propylene (FEP) film of TENG to simultaneously scavenge and store wind energy as

![Image](image_url)
chemical energy, where the SLB was prepared using the counter rolling mechanism in the order of the anode, PEO–LATP membrane, and cathode, as displayed in Fig. 5f [25]. Figure 6a presents that three EP-TENGs connected in parallel were integrated with three EP-SCs connected in series to create a self-charging power system, where the conductive carbon paper acted as capacitive materials, while the PAN paper served as separator [24]. The flexible paper-based supercapacitor is displayed in Fig. 6b, where the graphite, H$_3$PO$_4$/PVA, and kirigami were utilized as the active material, electrolyte, and separator, respectively [26]. Liu et al. reported a convoluted power device by internally hybridizing a TENG and an SLB by sharing common electrodes. The SLB consists of TiO$_2$ nanotubes as anode, the polyethylene oxide-Li$_{(1+x)}$Ti$_{(2-x)}$Al$_x$(PO$_4$)$_3$ (PEO–LATP) as solid electrolyte, and the LiMn$_2$O$_4$ nanoparticles as cathode (Fig. 6c) [27]. Figure 6d presents the schematic diagram of the all-paper-based cut-paper self-charging power unit (PC-SCPU), where it can be distinctly observed that the supercapacitor is composed of graphite as active material and H$_3$PO$_4$/PVA as electrolyte [28].

4 Self-Charging Principles

There are two typical charging principles of self-charging energy storage devices. One is based on piezoelectric potential-driven electrochemical oxidation and reduction reaction. For piezoelectric nanogenerator-based self-charging lithium-ion batteries, the piezoelectric field created by the PVDF piezo-separator forms along the z-axis from the cathode side of the system when compressive force is applied to the battery surface. The piezoelectric potential of the porous PVDF separator causes Li$^+$ ions to migrate from the cathode to the anode through ion-conducting separator and electrolyte, resulting in the incorporation of Li$^+$ ions into the anode electrode. An electron can take the same pathway to maintain charge neutrality, where this migration represents complete conversion of mechanical energy into electrochemical energy, as displayed in Fig. 7a [13]. For piezoelectric nanogenerator-based self-charging supercapacitors, since no pressure force is applied to the device, there is no electrochemical reaction due to the electrochemical equilibrium between the electrodes and electrolyte. A compressive stress can be caused by
the polarization of the separator (PVDF–ZnO or BPES, Fig. 7b, c) due to the piezoelectric effect. Moreover, the separator produces a potential difference along the thickness direction. These positive and negative piezoelectric potentials were generated at the top and the bottom of the film, driving the electrolytic ions to move
toward the positive and negative electrodes. The piezo-electric field induced cationic movement in the electrolyte to screen the generated piezopotential across the separator. This ionic movement induces an electrochemical imbalance in the electrolyte and the positive/negative electrodes. As a result, the positive and negative potentials were developed on the upper and lower surface of the separator that can drive the electrolyte ions toward the different electrodes. The charging process can be continued to accomplish equilibrium of the two electrodes with the generated piezoelectric potentials. The devices directly convert the mechanical energy into electrochemical energy in the SCs, as presented in Fig. 7b, c [17, 21], respectively.

Another charging principle is based on triboelectrification. Firstly, a TENG can be utilized to convert mechanical energy into electric energy, and then a bridge rectifier is used to convert the generated AC current of TENG into direct current signals before charging the energy storage devices, as displayed in Fig. 8a [23]. The electrical outputs generated by TENGs have the characteristics of high voltage and low current signals. Usually, a transformer is used to decrease the voltage and increase the current before rectifying, as displayed in Fig. 8b–d [19, 25, 27] or by adding a power management circuit between the TENGs and energy storage devices to reduce the voltage, increase the current, and turn the AC into DC signals [34]. More importantly, Fig. 8c, d displays the self-charging process of the LIBs that can simultaneously generate and store electric energy by itself, which is an important step toward next-generation LIBs for pushing the practical applications in self-powered electronic devices.

Fig. 8  a Circuit diagram of the self-charging powered textile for wearable electronics (WE). Reprinted with permission from Ref. [23]. b Implanting an SLB into a triboelectric nanogenerator. Reprinted with permission from Ref. [25]. c Schematic diagram of self-charging process for the Li-ion batteries, distinctly showing the flow of electrons in the circuit. Reprinted with permission from Ref. [19]. d Schematic illustration of the electric storage process from the external capacitor to the convoluted power device as an SLB. Reprinted with permission from Ref. [27]
Pu et al. reported a method to solve the problem of the impedance match between the TENG and a battery with appropriate design of transformers [35]. It provides an effective method to solve the problem of impedance match between TENGs and energy storage devices.

5 Self-Charging Performances

The mechanical energy scavenged from environment using nanogenerators can be converted into electricity, which can be then stored in the energy storage devices such as lithium-ion batteries or capacitors using a management circuit. In this process, energy conversion efficiency and self-charging performance are undoubtedly critical parameters. Figure 9a presents the self-charging curves of the all-solid-state SCPC under periodically applied compressive deformations. When a compressive force of 30 N was applied onto the SCPC at the frequency of 1 Hz, the voltage of the SCPC increased from 25 to 473 mV within 240 s, which can be discharged back to its original voltage in 85 s under a constant current of 5 μA. The obtained corresponding capacity of the SCPC is approximately 0.118 μAh [11]. With a compressive force of 10 N applied to the SCPC based on the PVDF–PZT nanocomposite film at a frequency of 1.5 Hz, the voltage of the device can be increased from 210 to 297.6 mV in 240 s. The device was discharged back to its original voltage in 37 s under a constant discharge current of 1 μA, as depicted in Fig. 9b [12]. Figure 9c presents the voltage change curves of the coin cell in the self-charging and discharging processes. When a compression mechanical energy of 282 mJ was applied on the cell at a frequency of 1 Hz, the voltage of the device can be increased from 1.2 to 1.4 V in approximately 200 s. After achieving the self-charging process, the cell was discharged at a constant current of 0.01 mA, resulting in the decrease in voltage to 1.2 V, where the corresponding

![Fig. 9](image_url)
discharge capacity was 0.4 μAh [13]. Figure 9d displays the charging/discharging curves of HEPD as a SC with a specific capacitance of approximately 0.5 F/cm², when the fuel (ascorbate) and oxidant (O₂) were absent in the anolyte and catholyte, respectively [14]. The device was externally charged and then discharged, realizing excellent reproducibility for several cycles. The charging curves of the MSC array component are displayed in Fig. 10a, where the stored charge can be increased steadily with increased charging time, and the potential can be approximately 3 V in 117 min [15]. Figure 10b illustrates that the voltage of the SCPC can be increased from 160 to 299 mV in 250 s when the device is under a compressive force of 34 N with a working frequency of 1.8 Hz. After the self-charging process, the device was discharged with a constant current back to its original voltage in 207 s under a discharge current of 3 μA [16]. The self-charging capability of the five SCSPCs in series was tested under the continuous human palm contact to all devices, as displayed in Fig. 10c. Under continuous self-charging process, the open-circuit voltage can be increased from 160 to 280 mV (120 mV charged) in 350 s and is then sustained in 250 s, even after removing the deformation [17]. The self-charging and discharging process of the fabricated device under continuous human finger imparting (at F ≈ 16.4 N, f = 1.65 Hz) for 80 s is illustrated in Fig. 10d. Although an initial voltage of approximately 130.1 mV was perceived, the device can be charged up to 281.3 mV from its initial voltage after repeated finger imparting. As a result, the device exhibits a distinct voltage increase of 151.2 mV. After removing finger imparting, the device was discharged back to its initial voltage in 145.5 s under a constant discharge current of 10.5 µA [21]. More importantly, Pazhamalai et al. reported that the MoSe₂ SCSPC device can be charged from 85 to 708 mV under an applied compressive force.

![Fig. 10](image1)

**Fig. 10** a Charging curve of the MSC array component. Reprinted with permission from Ref. [15]. b A typical self-charging process under periodic compressive stress of a SCPC fabricated using PVDF mesoporous nanostructured film as piezo-separator. Reprinted with permission from Ref. [16]. c Self-charging process of the serially connected five SCSPCs under periodic compressive strain provided by human palm impact to the whole devices. Reprinted with permission from Ref. [17]. d The self-charging-discharging profile under continuous human finger imparting. Reprinted with permission from Ref. [21]
of 30 N within 100 s [37]. By comparison, it is observed that the MoSe₂ SCSPC device has better self-charging performance than the devices depicted in Figs. 9a and 10b. The excellent self-charging performance of the fabricated MoSe₂ SCSPC devices might be related to intercalative type 2D MoSe₂ energy storing electrode. As displayed in Fig. 11a, the SCPF was charged using different types of finger motions. For finger tapping, the SCPF was charged from 0 to 2.5 V within 6102 s (blue line) and discharged at 1 μA for 1639 s. When the finger motions are slow sliding (0.5 m s⁻¹) and fast sliding (0.8 m s⁻¹), the charging times can be shortened to 3518 s (green line) and 2094 s (red line), respectively [18]. The following corresponding discharging at 1 μA can still be sustained in approximately 1630 s. Figure 11b presents the self-charging performance of the top Li-ion battery; it can be easily charged from 1.5 to 3.5 V in approximately 3 min under the vibrations of the FEP film induced by a wind blowing at a speed of approximately 10 m s⁻¹ though the device [19]. Figure 11c displays the voltage change of the F-DSSC, where it can be charged from 1.8 to 3.5 V in approximately 33 s using F-TENG [23]. Figure 12a depicts the charging curves of the self-charging power system under the stable and fixed frequency of 5 Hz for the practical application. The circuit schematic diagram of the self-charging system is displayed in the inset of Fig. 4l. It requires approximately 2150 s to charge the EP-SCs to 2 V [24]. The SLB can be easily charged from 1.5 to 3.6 V in approximately 55 s under the vibrations of the FEP film induced by a wind blowing with the speed of approximately 24.6 m s⁻¹ though the device, as presented in Fig. 12b [25]. The convoluted power device in the human shoe under the fast walking condition can charge a Li-ion battery to a higher voltage of approximately 1.12 V in approximately 5 min as compared with that (1.05 V) under the slow walking condition, as displayed

![Charging curves of four TFSCs in series charged by the TF-TENG at fast sliding, slow sliding, and pressing. Reprinted with permission from Ref. [18].](image1.png)

![Self-charging and discharging curves of the top Li-ion battery. Reprinted with permission from Ref. [19].](image2.png)

![Charging curve of the F-DSSC and the F-TENG, where the light blue-shaded area corresponds to the charging curve of the F-DSSC and the light red-shaded area corresponds to the charging curve of the F-DSSC-F-TENG hybrid. Reprinted with permission from Ref. [23].](image3.png)
in Fig. 12c [27]. Figure 12d presents the voltage–time curve of a PC-SCPU when charging and instantaneously working (charging frequency: 3 Hz), where it requires charging for approximately 60 s for wireless remote operation [28]. The above research results indicate that the nanogenerator-based self-charging storage devices have good self-charging performances, which can push the practical applications of self-charging devices.

6 Applications

The fabricated nanogenerator-based self-charging energy storage devices can be utilized as a power source for powering certain electric devices. The all-solid-state SCPC can power smartwatch, sports bracelet, and LEDs, as illustrated in Fig. 13a [11]. Figure 13b presents that two lighted LEDs can be powered by the self-charging miro-supercapacitor power unit (SCMPU), where the SCMPU was inserted in the insole of a shoe to drive a commercial hygrothermograph [15]. Operation of green light-emitting diode uses integrated SCSPCs in series as the power source, as displayed in Fig. 13c [17]. Figure 13d illustrates that a calculator can be driven by the SCPF [18]. Figure 13e presents that the twisted Li-ion battery can be used to light up a green LED [19]. Moreover, it can be used as a power supply for small electronic wrist watch, commercial digital calculator, mobile LCD screen, and portable speaker (Fig. 13f) [21]. As displayed
Fig. 13  
(a) The all-solid-state SCPC can power smartwatch, sports bracelet, and LEDs. Reprinted with permission from Ref. [11].
(b) Circuit diagram of the energy supply mode and photographs of showing two LEDs being powered by the SCMPU, the SCMPU inserted in the insole of a shoe and using the SCMPU to drive a commercial hygrothermograph. Reprinted with permission from Ref. [15].
(c) Operation of green light-emitting diode using serially connected SCSPCs as the power source. Reprinted with permission from Ref. [17].
(d) Photograph of a calculator driven by the SCPF. Reprinted with permission from Ref. [18].
(e) Photographs of the twisted Li-ion battery to light up a green LED. Reprinted with permission from Ref. [19].
(f) Photographs of the fabricated devices can instantaneous lightning of four commercial red LEDs upon repeated finger imparting and lightning of a green LED light with adequate intensity. Meanwhile, power-up small electronic wrist watch, commercial digital calculator, mobile LCD screen, and portable speaker. Reprinted with permission from Ref. [21].
in Fig. 14a, LED light signals of “FIB” can be lighted up by tapping a TENG bracelet wrapped around the wrist. Two SCs in series can be used to power the temperature–humidity meter and calculator [22]. Figure 15a displays a self-charging power system, which can be used to power an electronic watch and a calculator at 5 Hz [24]. The fabricated all-in-one shape-adaptive self-charging power package in conventional wearable electronics and the integrated KP-SC (6 units, 3 devices in series) light up a single commercial green LED under cycling stretching movement, as demonstrated in Figs. 14b and 15b, respectively [26]. Figure 15c displays a green LED that can be lighted up by the convoluted power device as an SLB [27], where a PC-SCPU-driven wireless remote

![Image](https://doi.org/10.1007/s40820-019-0251-7)
control can be observed. Meanwhile, a PC-SCPU can be used as a power source for driving a digital electric watch and a temperature meter, as demonstrated in Fig. 15d [28]. Thus, these devices have a wide range of potential applications in electronic devices and wireless sensors.

7 Summary and Outlook

This review is focused on the recent progress of nanogenerator-based self-charging energy storage devices. The major achievements in this field can be summarized as follows: (1) Various self-charging devices have been developed to scavenge the mechanical energy and store it in themselves, which can be used to power some small electronic devices. (2) Self-charging principles of the energy storage devices have been investigated in details, where piezoelectric and triboelectric effects have been used to explain the working mechanisms of these devices. (3) Substantial practical applications of self-charging energy storage devices have been demonstrated, ranging from wearable electronics, sports monitoring, wireless sensors, to daily electronics.

Although significant improvements have been achieved, some problems regarding the investigations of nanogenerator-based self-charging energy storage devices are needed to be addressed: (1) Device life is a very critical matter in practical application. More attention needs to be focused on the device life to realize long-life devices. It may be necessary to start with material selection, structure design, etc. (2) LIBs or supercapacitors can be used for self-charging energy storage devices; the capacities and impedances of them should match the output of the energy-harvesting systems for higher conversion efficiency. Moreover, cost, safety, and easiness to integrate these devices are also one of the prior concerns. (3) The development of self-charging energy storage devices in
future should follow the trend of miniaturization, diversification, integration, and portability.

In summary, developing nanogenerator-based self-charging devices is one of the effective methods to solve issues of continuous energy supply of the next-generation microelectronic devices. A series of research results regarding scavenging and storing the mechanical energy has been obtained, but there are still several problems to be solved such as other energy scavenging and storage systems, device life. Owing to the unremitting efforts by a large number of researchers around the world, we believe that the self-charging storage devices will be extensively applied in our daily life in the near future, especially for wearable electronic devices and self-powered systems.

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