ORIGINAL RESEARCH



Shape memory polymer solar cells with active deformation

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Abstract

The reliable and lightweight deployable solar arrays require the capability of large deformation for packaging and the ability to actively deform for deployment. To satisfy such demands, the shape memory polymer solar cells (SMPSCs) are fabricated and demonstrated in this paper by using flexible and conductive silver nanowire/shape memory copolyamide (AgNW/SMPI) composite film as the transparent substrates. The AgNW/SMPI composite film has good optical transparency (~73% at the wavelength of $450 \sim 1100$ nm), smooth surfaces (average RMS: ~3.32 nm), good shape memory performances (shape fixation ratio > 98%, shape recovery ratio > 98%), and can maintain excellent conductivity (~10 Ω/\Box) after mechanical deformations with large strain. Owing to the shape memory effect of the substrate, SMPSCs can be deformed into arbitrary shape and actively recover to the original shape upon heating. The power conversion efficiency of SMPSC (2.94%) is lower than that of ITO-based solar cells with the same structure (3.44%), due to the relatively lower optical transparency of SMPI. However, SMPSCs can maintain good photovoltaic performances after 50 bending-recovery cycles or few shape recovery cycles, demonstrating better flexibility and durability than ITO-based solar cells. The SMPSCs have the potential to be used in areas of sensors, medical probes, and displays.

Keywords Shape memory electrode · Flexible solar cells · Active shape changing · Durability · Shape memory copolyamide

1 Introduction

Deployable solar arrays are vital components for spacecraft. Most of the current deployable solar arrays have complicated structures and utilize non-controllable, non-testable, and non-reusable electro-explosive devices for deployment, which may lead to high impact and the risk of malfunction. To fabricate reliable and lightweight ultra-large deployable solar arrays with simple structures, the capability of large deformation for packaging to save the space before launching and the ability to actively deform for deployment will be required. Shape memory polymers (SMPs) are typical smart polymers that can response to external stimuli [1–14] and demonstrate huge application potential in the field of

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² Department of Astronautical Science and Mechanics, Harbin Institute of Technology, Harbin 150001, Heilongjiang, China deployable aerospace structures [1]. SMP structure can be deformed from original shape into complex shape due to the low modulus of SMPs at temperature higher than the transition temperature. After cooling, the stiffness of SMPs increases significantly, and such complex shape can be fixed. When being heated again, the structure can actively recover from different pre-deformed shapes to the original shape. Owing to the above features, SMPs are suitable for the substrate of deployable solar arrays. In reports, shape memory function has been endowed with some flexible electronic devices when employing shape memory polymers (SMPs) as substrates [13–16]. Accordingly, the flexible solar cells with shape memory properties are feasible.

Since SMPs are not conductive, the electrode material is a challenge for fabricating the shape memory solar cells. Although extensively utilized as the electrode in electronic devices [17–20], indium tin oxide (ITO) cannot satisfy the requirements of large deformation due to its intrinsic brittleness. Even coated on a polymer film, ITO still breaks easily at relatively low bending strain or under repeated bending-recovery cycles, which results in severe deterioration

in electrical conductivity [21]. Novel electrode materials have been developed to improve the flexibility of electronic devices [22-37], including graphene [22-24], carbon nanotubes [25–27], metal grids [28–30], random metal nanowires [16, 21, 31, 32], conducting polymers [33], and their combinations [34-36]. Among these electrode materials, silver nanowires (AgNWs), fabricated by drop-casting [16, 34, 38], Meyer rod coating [39, 40], spray coating [41, 42], or vacuum filtering [43] are promising due to their low resistivity and high optical transparency. Moreover, the AgNWs could remain conductive while bending. However, the uneven surface may lead to shorts across the functional layers, and the poor adhesion to the substrates makes AgNW electrodes unsuitable for direct employment in flexible electronics [38, 40]. The adhesion to polymers and the surface quality of the AgNWs can be effectively improved through embedding AgNWs into polymers [44, 45]. These composite electrodes have good conductivity and optical transparency, low surface roughness, and low conductivity reduction after mechanical deformation [44, 45] and have been utilized in some flexible electronic devices like solar cells and light-emitting diodes [16, 32, 37].

In this paper, we demonstrate the shape memory polymer solar cells (SMPSCs) which utilize the transparent and conductive AgNW/shape memory polyimide (SMPI) composite as the substrates and electrodes. The transparent SMPI is chosen as the substrate because it has application potential in the field of aerospace and flexible electronics due to its excellent mechanical strength, good optical transparency, high-temperature and radiation resistance, light weight, chemical inertness, and low thermal expansion coefficient [2, 3, 46–48]. The glass transition temperature (T_o) of the SMPI used in this work is 182 °C, and other basic properties of the SMPI film are listed in Table 1 and shown in the Supporting Information. The transparent electrodes were fabricated through embedding AgNW into synthesized SMPI substrates using a simple and low-cost fabrication method. The AgNW/SMPI electrodes could remain good optical transparency and conductivity after bending-recovery cycles. Due to the shape memory effect of the AgNW/SMPI electrode, SMPSCs can be deformed into arbitrary shape and recover to the original shape upon heating. The SMPSCs demonstrate better flexibility and durability than ITO/glass-based solar cells, while the power conversion efficiency (PCE) of the SMPSCs is just slightly lower than that of ITO/glass-based solar cells.

2 Experimental section

2.1 Materials

Poly(3-hexylthiophene) (P3HT, 99.99%), phenyl-C61-butyric acid methyl ester (PC61BM, 99.99%), and bisphenol A dianhydride (BPADA, 97%) were obtained from Sigma-Aldrich Co. LLC. (USA). 1,3-bis(3-aminophenoxy)benzene (BAB, 98%) was achieved from Tokyo Chemical Industry Co., Ltd. (TCI, Japan). Lithium fluoride (LiF), aluminum (Al), and 4,4'-(1,1'-Biphenyl-4,4'-diyldioxy)dianiline (BAPB, 97%) were bought from Aladdin Bio-Chem Technology Co., Ltd. (Shanghai, China). Dimethylacetamide (DMAc) was purchased from Kemiou Chemical Reagent Co., Ltd. (Tianjin, China) and distilled with activated 4 Å molecular sieves for 24 h under reduced pressure. Poly(3,4-ethylenedioxythiop hene):poly(styrenesulfonate) (PEDOT:PSS, Clevios P VP AI4083) were purchased from Heraeus Materials Technology Shanghai Ltd. (China). ITO/glass and ITO/PEN with the sheet resistance of ~9 Ω/\Box and ~35 Ω/\Box was afforded by Yingkou You Xuan Trade Co., Ltd. (Liaoning, China). Silver nanowires (average diameter: \sim 70 nm, average length: \sim 40 µm) were provided by Seashell Tech. (California, USA).

2.2 Fabrication of shape memory silver nanowire/ polymer composite electrodes

The fabrication processes of shape memory AgNW/SMPI composite electrodes are schematically shown in Fig. S1. Before the spin-coating of AgNW solution, the glass substrate was ultrasonically cleaned in detergent, deionized water, acetone, ethanol, and isopropanol in turn. AgNW solution, which has been diluted to the required concentration with isopropyl alcohol, was spin-coated onto glass substrates at 500 rpm for 9 s and then 1100 rpm for 30 s. After drying the AgNW/glass substrates at 80 °C for 30 min, the pre-prepared poly(amic acid) solution (precursor of SMPI, details in Supporting Information) was drop-coated onto

Table 1Performancescomparison of the substrates fordevices

Substrates	Thickness (mm)	$R_s(\Omega/\Box)$	RMS (nm)	Transmittance at 450~1100 nm (%)	Shape fixity ratio (R _f) (%)	Shape recov- ery ratio (R _r) (%)
ITO/glass	~1.10	~9	<10	>86	-	-
SMPI	~0.120	_	-	>86	98.8	100
AgNW/SMPI	~0.125	~10	3.32	>73	98.7	100
AgNW/SMPI	~0.125	~1	1.28	> 35	98.8	100

AgNW and then experienced step-wised curing processes of which the temperature was successively kept at 80 °C, 120 °C, 180 °C, 210 °C, and 250 °C, and each temperature was kept for 2 h. The smooth and flexible shape memory AgNW/SMPI composite electrode was achieved after being peeled off glass substrates in hot deionized water. The thickness of the AgNW/SMPI composite electrodes is determined by the amount of PAA solution and was controlled at around 120 µm in this work.

2.3 Preparation of shape memory polymer solar cells

Before the fabrication of SMPSCs, AgNW/SMPI composite electrodes were ultrasonically cleaned successively in detergent, deionized water, acetone, ethanol, and isopropanol, and then thermally treated at 120 °C for 1 h on vacuum dry chamber. After pre-ultrasonicated for 30 min, PEDOT:PSS solution, without any dilution, was spin-coated onto electrodes at 650 rpm for 9 s and then 3500 rpm for 60 s, and subsequently annealed at 120 °C for 20 min. The thickness of PEDOT:PSS film, measured by a Dektak 6 M stylus profilometer, was around 40 nm. The electrode/PEDOT:PSS was then transferred into a nitrogen-filled glove box where the oxygen and moisture levels are below 1 ppm. Before spin-coating active layer, P3HT and PC61BM, which were mixed in a weight ratio of 1:1, were co-dissolved in 1,2-dichlorobenzene and stirred at 60 °C for 24 h inside the glove box. The P3HT:PC61BM solution (22 mg mL⁻¹) was then spin-coated on the electrode/PEDOT:PSS at 550 rpm for 9 s and then 1100 rpm for 30 s to form a 210-nm-thick film and subsequently annealed at 135 °C for 5 min. A 1-nm-thick LiF thin film and a 100-nm-thick Al electrode were then consecutively thermally evaporated on electrode/ PEDOT:PSS/P3HT:PCBM to obtain the SMPSCs. The active area of the SMPSCs was 12 mm² which is determined by the shadow mask. Lastly, the whole device was thermally annealed at 120 °C on hot plate for 30 min under nitrogen atmosphere to improve the performances of solar cells. The solar cells with the same structure except for the ITO/glass electrodes were also fabricated as the control devices.

2.4 Performance testing

Ultraviolet–visible (UV–Vis) transmittance of 120-µm-thick SMPI film and AgNW/SMPI composite electrode at the wavelength from 200 to 2000 nm was tested on a Perkin-Elmer Lambda 950 spectrometer, respectively. The sheet resistance (R_s) of the films was tested by a four-probe sheet resistance/resistivity tester (RG-7C, NAPSON). The rootmean-square (RMS) roughness was scanned by an atomic force microscope (AFM, Digital Instrument, NS4/D3100CL, Germany) with the scanning area of 10×10 µm² and was analyzed with Nanoscope 1.50 software. The surface images of samples were taken from Quanta 200FEG-scanning electron microscopy (SEM) at an acceleration voltage of 15.0 kV. The photovoltaic performances of solar cells were characterized under a 1.5 solar illumination at 1000 W m⁻² (1 sun). The current–voltage (*J-V*) characteristics of solar cells can be monitored and recorded by computer, from which the open-circuit voltage (V_{oc}) and short-circuit current density (J_{sc}) can be directly read. Fill factor (FF) and PCE can be calculated by the Eqs. S5 and S6 according to Fig. S5, respectively.

3 Results and discussions

3.1 Properties of the AgNW/SMPI composite electrodes

The fabrication process of the shape memory AgNW/SMPI composite electrodes is illustrated in Fig. S1 and detailed in Sect. 2.

The SEM and AFM images of the AgNW/SMPI composite electrodes are shown in Fig. 1a–f. From the SEM images, we can clearly see that AgNWs were uniformly distributed and embedded in the SMPI substrate to form the conductive pathway. The RMS roughness of the shape memory AgNW/ SMPI composite electrode with ~10 Ω/\Box and ~1 Ω/\Box were 3.32 nm and 1.28 nm, respectively, which are comparable to that of ITO (~9 Ω/\Box , <10 nm). However, the RMS roughness of the conductive AgNW networks that were directly spin-coated onto SMPI substrates was greater than 100 nm. This is consistent with other reported results [32, 34, 38, 40]. Embedding AgNW into SMPI film can effectively improve the surface quality of the composite electrodes since SMPI will fill out the void inside AgNW networks, which is beneficial to improve the photovoltaic performances of solar cells.

The AgNW/SMPI composite electrodes had high conductivity and optical transparency. The sheet resistance (R_s) of AgNW/SMPI composite electrodes was dependent on the concentration of the AgNW solution and R_s values of about 100 Ω/\Box , 10 Ω/\Box , and 1 Ω/\Box could be obtained when the concentration is respectively 1.5 mg/mL, 4 mg/mL, and 10 mg/mL. Figure 2a illustrates the UV–Vis transmittance at different wavelength of the AgNW/SMPI composite electrode with different R_{s} . The increasing of the conductivity is the result of the increased AgNW density, which could enhance the light scattering. Therefore, as can be seen from Fig. 2a, the optical transparency decreased with the increase of conductivity of AgNW/SMPI composite electrode. The AgNW/SMPI composite can also be used as a heater due to its good electrical conductivity. Figure 2b shows the temperature changes of AgNW/SMPI composite electrodes (R_s : ~10 Ω/\Box) when the external voltage was applied for 600 s and

Fig. 1 a, b SEM morphologies of the AgNW/SMPI composite electrode, showing that AgNW was embedded inside the SMPI substrates. The 2D atomic force microscope (AFM) image (c) and 3D AFM image (d) of AgNW/SMPI composite electrode with the R_s of ~1 Ω/\Box . The 2D AFM image (c) and 3D AFM image (f) of AgNW/SMPI composite electrode with the R_s of ~10 Ω/\Box



then switched off for another 600 s. That means it is possible to trigger the shape recovery through Joule heating [6].

The AgNW/SMPI composite electrodes also demonstrated excellent flexibility and durability while maintaining high conductivity. We compared the changes in R_s of AgNW/SMPI composite electrode and commonly used ITO/ PEN after a number of bending-recovery cycles. The configuration of a rectangular AgNW/SMPI film after bending is illustrated in Fig. S6 (the bending angle is close to 180°), and the results are shown in Fig. 2c. The R_s of ITO/ PEN increased by more than one hundred times of its initial value (~35 Ω/\Box) after only 10 bending-recovery cycles, while the R_s of AgNW/SMPI was almost unchanged after 100 bending-recovery cycles. In addition, the AgNW/SMPI composite electrode had good shape memory performances and could remain good conductivity after 100 bending shape recovery cycles, as shown in Fig. 2d. As shown in Fig. 2e, f, the AgNW/SMPI film could be used as the conductive wire to light the LED when the electrode is in deformed state (bending angle: 90°) and recovered state after 100 shape memory cycles.

Table 1 summarizes the performances of ITO/glass, SMPI, and AgNW/SMPI with R_s of ~10 Ω/\Box and ~1 Ω/\Box . To avoid the decrease of the light absorption of photoactive layers due to low transmittance, the AgNW/SMPI with sheet resistance of ~10 Ω/\Box was used in the fabrication of SMPSCs.

3.2 Performance of the shape memory polymer solar cells

The complete layer stack of the SMPSCs is illustrated in Fig. 3a, where PEDOT:PSS, P3HT:PC61BM, LiF served as the hole-transporting layer, active layer, and electron-transporting

Fig. 2 a UV–Vis transmittance spectra of the shape memory AgNW/SMPI composite electrodes with different $R_{\rm c}$. **b** The temperature change of AgNW/ SMPI composite electrode $(R_{\rm s}: \sim 10 \ \Omega/\Box)$ under different applied voltages. The resistance changes of c AgNW/SMPI and ITO/PEN electrodes after bending-recovery cycles and d the resistance change of AgNW/ SMPI composite electrode after shape memory cycles. The AgNW/SMPI composite electrode was used as the conductive wire to light the LED when the bending angle is 90° (e) and 180° (**f**)



layer, respectively, and the band energy diagram for SMPSCs is illustrated in Fig. 3b. The J-V characteristics of the solar cells fabricated on ITO (~9 Ω/\Box) and AgNW composite electrodes (~10 Ω/\Box) are presented in Fig. 3c, d, respectively. The inset photographs show the solar cells fabricated on ITO electrodes and AgNW/SMPI composite electrodes, respectively. The FF and V_{oc} of two devices were close, and the J_{sc} of solar cells with AgNW electrodes (~7.85 mA/cm²) was lower than that of solar cells with ITO electrodes (~9.36 mA/cm²). Correspondingly, the PCE of SMPSCs (2.94%) was lower compared to that of the ITO-based solar cells (3.44%). However, SMP-SCs with AgNW/SMPI composite electrodes demonstrated comparable photovoltaic performance to some reported solar cells with AgNW electrode [32, 37]. One possible reason is that the lower optical transparency of AgNW/SMPI electrodes (~73% at wavelength of $450 \sim 1100$ nm) than that of ITO electrodes (~88% at wavelength of 450~1100 nm) reduces the light adsorption of the P3HT:PC61BM and thus leads to the decrease of electron–hole pairs. This can be validated by the rough estimation that the J_{sc} of the solar cell is in proportion to the transmittance of the electrode, since the solar cells we compared had the same structure except for the electrode. The SMPI is chosen in consideration of the high temperature and radiation resistance required for aerospace applications [49–54] and the photovoltaic performances of SMPSCs can be enhanced by using SMPs with higher transparency [16] in other application scenarios. Furthermore, the efficiency of the SMPSCs can be improved by using AgNW/SMP composite electrode in flexible solar cell with high efficiency, like flexible perovskite solar cells [55–57].

The 3D AFM surface images of SMPI/AgNW/PEDOT:PSS, glass/ITO/PEDOT:PSS, SMPI/AgNW/PEDOT:PSS/P3HT:PCBM, and glass/ITO/PEDOT:PSS/P3HT:PCBM films are illustrated in Fig. 4, and the 2D AFM images are illustrated in Fig. S7. The corresponding RMS roughness is 1.13 nm, 1.63 nm, 4.15 nm, and 3.54 nm, respectively. The

Fig. 3 a The complete layer stack of SMPSCs. **b** Band energy diagram for P3HT:PC61BM blend solar cells with SMPI/AgNW composite electrodes and Al electrodes. *J-V* characteristics for devices with ITO electrodes (**c**) and AgNW composite electrodes (**d**), and the inset photographs are the fabricated solar cells



results indicate that functional layers fabricated on the AgNW/ SMPI electrode had similar surface quality to that fabricated on ITO/glass. The smooth surfaces are beneficial to reduce interface contact resistance and improve carrier flow. Furthermore, the grain-like composition of the surfaces guaranteed the good photovoltaic performances of solar cells due to the phase separation and the generation of macroscopic domains of a 1:1 blend of P3HT and PC61BM [58].

The performances of SMPSCs after different numbers of bending-recovery cycles and bending shape memory cycles were investigated. Figure 5a is the schematic of the shape memory cycle of SMPSCs. The SMPSC is firstly bent into the deformed shape at a temperature higher than the T_g of SMPI and then cooled while the deformed shape is maintained. The temporary shape can be fixed after the external mechanical load is removed. In the experiments, the shape memory cycle of SMPSCs was conducted inside the nitrogen-filled glove box. The SMPSC can recover to the original flat shape at 200 °C in 20 s (Movie S1).

From Fig. 5b, c, SMPSC demonstrated good flexibility and durability, and the photovoltaic performance of the SMPSC reduced slightly after 50 bending-recovery cycles. The *J-V* characteristics of SMPSCs before and after shape memory cycles are shown in Fig. 5d and the inset

Fig. 4 3D surface images of a SMPI/AgNW/PEDOT:PSS after annealing at 120 °C for 20 min, b glass/ITO/PEDOT:PSS after annealing at 120 °C for 20 min, c SMPI/AgNW/PEDOT:PSS/ P3HT:PCBM after annealing at 135 °C for 5 min, and d glass/ ITO/PEDOT:PSS/P3HT:PCBM after annealing at 135 °C for 5 min





Fig.5 a The bending shape memory cycles of SMPSCs. **b** *J-V* characteristics of SMPSCs after bending-recovery cycles, and **c** the effects of bending-recovery cycles on V_{oc} , J_{sc} , PCE, and FF. **d** *J-V*

characteristics of SMPSCs after bending shape memory cycles, and the inset photographs are one shape memory cycle of real products

photographs shows the original state, deformed state, and recovery state of SMPSC in one shape memory cycle. The corresponding results of V_{oc} , J_{sc} , PCE, and FF are summarized in Table 2. The value of V_{oc} and FF was similar to the simulation results in reference [59]. However, the PCE of SMPSCs decreased from 2.94 to 2.00% after 10 shape memory cycles and dropped to 0.72% after 30 shape memory cycles. From Table 2, J_{sc} reduced most significantly after 30 shape memory cycles, which results from the decrease of electron–hole pairs. One reason is that the resistance of the AgNW/SMPI film increases by several times after 30 shape memory cycles, as shown in Fig. 2d. Another reason is that defects possibly generated in layers or interfaces [15] after multiple deformations at high temperature.

From the aforementioned results, the SMPSCs have the potential to be used in deployable solar arrays in the future. The shape memory property makes it possible to package or roll the ultra-large solar arrays based on SMPSC for reducing the space before launching and to actively deploy the solar array in space. The variable stiffness of SMP is also beneficial for the shape fixation after deployment. For the deployable solar arrays, limited shape memory cycles are usually required. In such case, the performance deterioration can be avoided, and the photovoltaic performance of SMPSCs has only slight reduction.

Table 2 Comparison of	
photovoltaic performances	s for
devices after shape memor	ry
cycles	

Substrates	$R_s\left(\Omega/\Box\right)$	Shape memory cycle numbers	$V_{\rm oc}$ (V)	$J_{\rm sc}$ (mA/cm ²)	FF (%)	PCE (%)
AgNW/SMPI	~10	0	0.560	7.85	67	2.94
AgNW/SMPI	~10	1	0.566	7.58	61	2.61
AgNW/SMPI	~10	10	0.543	6.77	55	2.00
AgNW/SMPI	~10	30	0.506	2.66	54	0.72

4 Conclusion

In this paper, we demonstrate shape memory polymer solar cells with the structures of SMPI/AgNW/PEDOT:PSS/ P3HT:PCBM/LiF/Al, of which AgNW/SMPI served as the transparent and conductive substrates. Due to the flexibility and durability of AgNW/SMPI composite electrode, SMPSCs were highly flexible and could be repetitively deformed with large strain. Moreover, owing to the shape memory property of the substrate, SMPSCs could be deformed into arbitrary shape and actively recover to the original shape upon heating. Though the PCE of SMPSC is lower than that of the control device fabricated on ITO/glass substrate, the photovoltaic performances of SMPSCs just had slight reduction after 50 bending-recovery cycles or few shape memory cycles. However, the photovoltaic performances deteriorated severely after large number of shape memory cycles. The SMPSCs have the potential to be used in deployable solar arrays in the future, and the transparent AgNW/SMPI composite electrodes may also be used in areas of sensors, medical probes, displays, and so on.

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Availability of data and material Data are available from the authors upon reasonable request.

Declarations

Competing interests The authors declare no competing interests.

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