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High energy conversion efficiency conducting polymer actuators based on PEDOT:PSS/MWCNTs composite electrode†

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Conducting polymer actuators have attracted enormous attention due to their capacity to be driven by low-voltage, their large deformation, and ability to work in air. However, they usually present low conversion efficiency, cycle stability and response actuation speed which severely hamper their practical application. Here, we report a high performance conducting polymer actuator based on a PEDOT:PSS/MWCNTs electrode, which displays high actuation performance, including high electromechanical conversion efficiency (1.04%), large actuation strain (0.64%), high strain rates (8.83% s⁻¹), excellent cycle stability (100 000 cycles), and good generated blocking force (1.43 mN) under 2.5 V. The key to the higher performance lies in the porous structure, high electrical conductivity (153.8 S cm⁻¹) and mechanical strength (918.3 MPa) of the PEDOT:PSS electrode coupled with carboxylic MWCNTs, which result in a great improvement of ion storage capacity in the actuator. Our studies demonstrate the important role of MWCNTs in enhancing the performances of conducting polymer actuators, which will guide the development of next generation electroactive polymer actuators.

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1. Introduction

In past decades, ionic electroactive polymer (EAP) actuators based on ionic polymer metal composites (IPMC), conducting polymers and carbon nanomaterials have been widely studied due to their impressive actuation performance.^{1–3} Among them, conducting polymers, such as polypyrrole, polythiophene and polyaniline, have been regarded as promising candidates for the preparation of lightweight, flexible and biocompatible EAP actuators. Actuators based on CPs, which dimensional changes resulting from electrochemical doping, transportation of dopant ions between the polymer matrix and the surrounding electrolyte solution, have the following advantages: combination of low driving voltage (typically 1–3 V), large deformation and fast response speed compared with traditional IPMC.^{4–6}

Poly(3,4-ethylenedioxythiophene):poly(styrenesulfonic acid) (PEDOT:PSS) is considered as one of most successful commercially available conducting polymers owing to its high transparency, high flexibility, and excellent thermal stability. However, its low conductivity and mechanical strength have been limiting factors for practical applications. The electrical

conductivity of PEDOT:PSS can be improved by various of approaches. Recent studies have found that the conductivity of PEDOT:PSS can be significantly enhanced treating by thermal, organic solvents, ionic liquids, surfactants and acids.^{7–11} Kim *et al.* reported the impressive high conductivity of PEDOT:PSS up to 4380 S cm⁻¹ by H₂SO₄ treatment, which achieved over four orders of magnitude enhancement compared with untreated PEDOT:PSS.¹² Besides the above mentioned treatment method, it is highly desirable to add an additional additive into PEDOT:PSS. Such an additive is required to own excellent conductivity and mechanical rigidity to improve the properties of pristine PEDOT:PSS. Recently, carbon nanotubes, which first discovered by Iijima,¹³ are regarded as a good additive for PEDOT:PSS due to their superb electrical, mechanical and chemical properties.^{14–16} Terasawa *et al.* developed PEDOT:PSS/SWCNT/IL actuator which combining electrostatic double-layer and faradaic capacitors that exhibited more superior electrochemical characteristics and actuation performance compared with PEDOT:PSS/IL actuator.¹⁷

In this work, PEDOT:PSS electrode doping with various content of carboxylic MWCNTs were prepared by solution-casting method and characterized by means of a four-point technique, FE-SEM and tensile test *etc.* The doping of carboxylic MWCNTs generates conjugate effect and hydrogen bonding with PEDOT:PSS, which not only obviously changes polymer structure but also enhances the electrical conductivity and mechanical properties of the electrodes. Furthermore, we fabricated three-layer structure actuator (as shown in Fig. 1)

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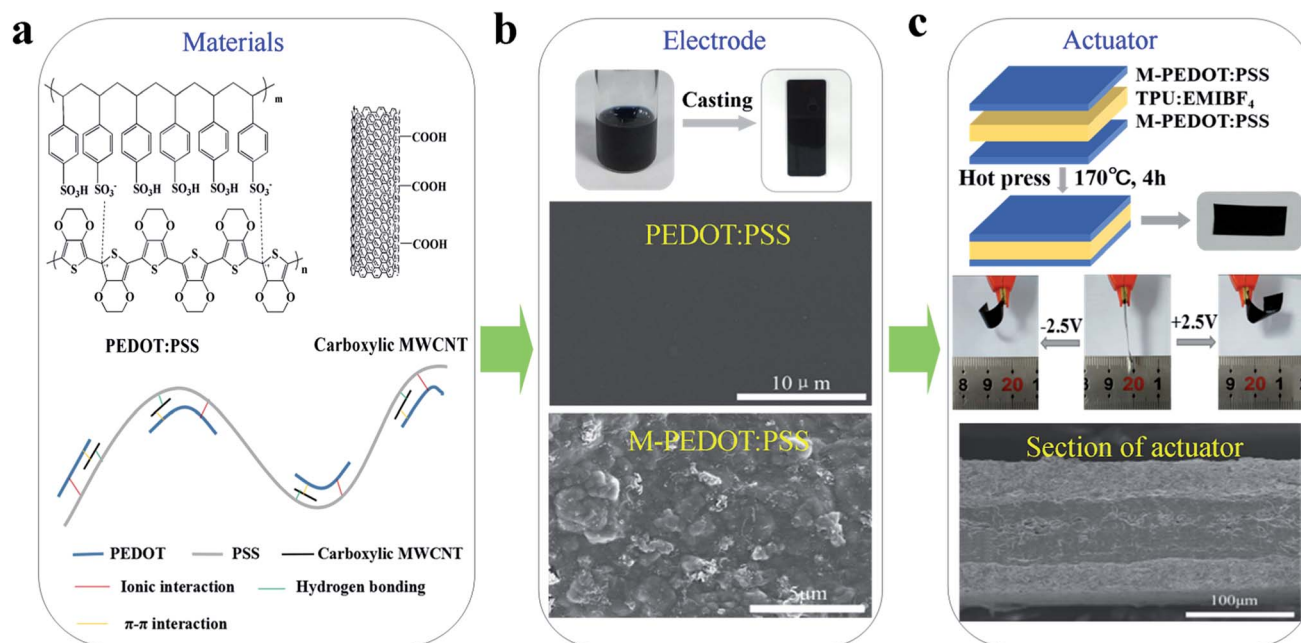


Fig. 1 Schematic illustration of actuator based on M-PEDOT:PSS electrode and fabricated by hot press.

taking TPU/EMIBF₄ as supporting polymer. Benefit from the porous structure and high electric conductivity of composite electrode, the obtained actuators based on PEDOT:PSS/MWCNTs (M-PEDOT:PSS) electrode displayed a greatly improved actuation performance compared with pristine PEDOT:PSS actuator. The actuator based on M-PEDOT:PSS electrode were capable of producing large bending deformation (up to the strain of 0.64%), high strain rates (8.83% s⁻¹), high electromechanical conversion efficiency (1.04%) and enhanced force generation (1.43 mN) at 2.5 V.

2. Experimental section

2.1 Materials

PEDOT:PSS (Clevios, PH1000, 1.3 wt%) was commercially available in the form of water dispersion form Heraeus. The carboxylic MWCNTs (95%, XF NANO, Inc.) had an -COOH content 3.86 wt%, average length of 10–30 μm , and surface area 500 m² g⁻¹. Fig. 1a shows the chemical structures of PEDOT:PSS and carboxylic MWCNTs. Thermoplastic polyurethane (TPU) was obtained from DuPont™. 1-Ethyl-3-methyl imidazolium tetra-fluoroborate (EMIBF₄, 99%) purchased from Shanghai Cheng Jie Chemical Co. *N,N*-Dimethylformamide (DMF) were obtained from Sinopharm Chemical Reagent Co., Ltd and deionized water was purified through Ultrapure Milli-Q system.

2.2 Preparation of electrodes layers

The carboxylic MWCNTs was dispersed in the PEDOT:PSS solution by a magnetic stirring for 6 h to obtain a homogeneous solution. The content of carboxylic MWCNTs in PEDOT:PSS was 10–30 wt%, and it was then homogenized for 2 h by bath sonication using a KQ-300DA sonicator at room temperature

(Fig. S1a, ESI†). Finally, 3 mL mentioned above homogeneous solution was casted on a glass substrate (2.5 cm $\times$ 7.5 cm size) and dried at 60 °C for 1 day to obtained PEDOT:PSS/MWCNT (M-PEDOT:PSS) electrodes film (Fig. S1b†). The PEDOT:PSS electrodes film without carboxylic MWCNTs was fabricated in the same method. The thickness of the resulting electrodes film was about 25 μm .

2.3 Preparation of the electrolyte layers

1 g TPU and 1 g EMIBF₄ was dissolved in 30 mL DMF at 80 °C overnight to obtain uniform solution. Then 2 mL TPU/EMIBF₄ solution was casted on a glass substrate (2.5 cm $\times$ 7.5 cm size) and dried at 80 °C for 2 days to obtained TPU/EMIBF₄ electrolyte film.

2.4 Assembly of the sandwiched actuators

Two piece of M-PEDOT:PSS electrode films were laminated on TPU/EMIBF₄ electrolyte film and heat pressed at 170 °C for 4 hours and naturally cooled to room temperature. After that, the film was carefully peeled from the glass substrate to get a free-standing sandwiched membrane. The thickness of obtained actuator membranes were about 110 μm . Then the actuator membranes were cut into strips with a standard dimension (5 mm in width and 20 mm in length). The pristine PEDOT:PSS electrode based actuator was constructed by the same process.

2.5 Characterization

FE-SEM was recorded by a Hitachi S-4800. The samples were dried in the vacuum oven overnight before capturing the surface and cross-sectional images. The electrical conductivity of electrode was tested by a multifunction digital four-probe tester (ST-2258C). The mechanical properties were evaluated



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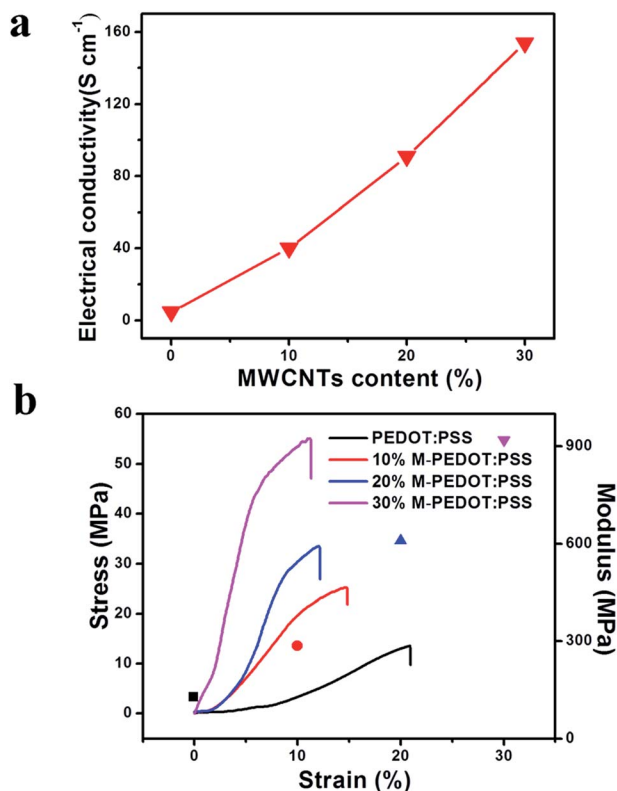


Fig. 3 (a) The electrical conductivity of composite films as a function of weight fraction of carboxylic MWCNTs. (b) Stress and modulus as a function of strain for the electrode membrane.

in M-PEDOT:PSS, therefore the charge becomes more delocalized on the PEDOT chains, which will be helpful to the charge transfer and conductivity enhancement of composite electrode films.²² The other reason for enhancing of electrical conductivity is a highly conductive MWCNTs network is formed in the polymer matrix with the MWCNTs weight fraction reaches a certain content.¹⁵

3.4 Tensile test

In order to expound the effect of carboxylic MWCNTs on mechanical properties of the electrode and actuator films, tensile test was performed and presented in Fig. 3b and S2 (ESI).[†] Fig. 3b shown the stress–strain curves of the pristine PEDOT:PSS and M-PEDOT:PSS electrode film with weight percentages of from 10% to 30%. Pristine PEDOT:PSS electrode film fractured at maximum stress of 13.54 MPa, whereas M-PEDOT:PSS film with weight percentages of 10% and 30% fractured at maximum stresses of 25.14 MPa and 54.89 MPa respectively. The calculated tensile modulus is listed in Table S2 (ESI).[†] The tensile modulus of the M-PEDOT:PSS electrode film (918.27 MPa) with a weight percentage of 30% is seven times that of the pristine PEDOT:PSS electrode film (127.64 MPa). It is noted that the elongation at break of the 30% M-PEDOT:PSS electrode film attained only 11.27%, which is almost half the PEDOT:PSS electrode film (20.93%). According to the results we can see that the mechanical strength and stiffness of the flexible

PEDOT:PSS polymer are greatly enhanced by the addition of the rigid carboxylic MWCNTs. The greatly enhancements in tensile strength and modulus of the M-PEDOT:PSS electrode have a significant impact on the mechanical properties of obtained actuators. As shown in Fig. S2,[†] the 30% M-PEDOT:PSS electrode based actuator has a tensile strength of 7.75 MPa, which is 73.65% higher than that of the pristine PEDOT:PSS electrode based actuator. The calculated tensile modulus of the 30% M-PEDOT:PSS electrode based actuator is almost two times as much as the pristine PEDOT:PSS electrode based actuator (Table S2[†]).

3.5 Electrochemical properties

To evaluate the actuators' charge storage behavior, cyclic voltammetry (CV) was employed to display it. The thickness of the actuator is well controlled within the range between 108 μm and 118 μm . Fig. 4a shown the electrochemical properties of the pristine PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators examined by CV with a potential window from -2.5 to 2.5 V and a scan speed of 100 mV s^{-1} . The shape of the CV curve for the pristine PEDOT:PSS electrode based actuators expressed as typical Faraday capacitor (FC) behavior, and the specific capacitance value which calculated from the integrated area of CV curves is 26.95 F g^{-1} as presented in Fig. 4b, which is close to the value reported by Terasawa.¹⁷ The 30% M-PEDOT:PSS electrode based actuators performed more electrical double layer

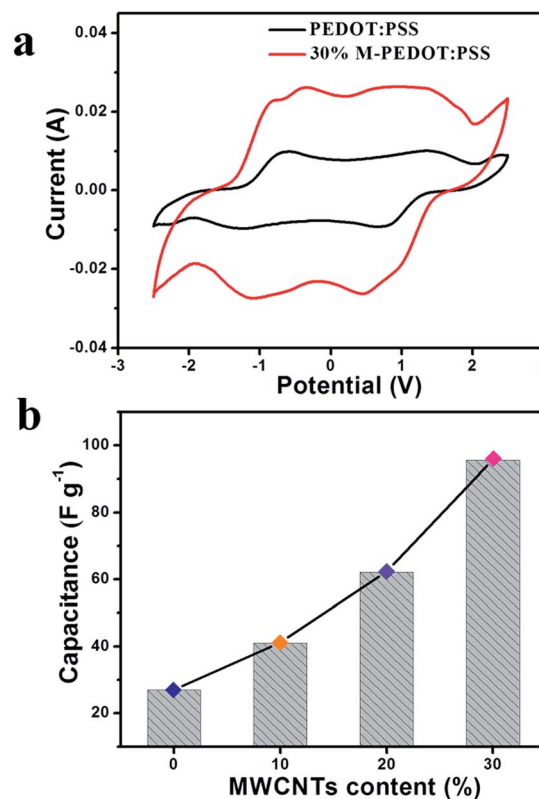


Fig. 4 (a) Cyclic voltammetry (CV) curves for the PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators at 100 mV s^{-1} sweep rate. (b) The calculated capacitances for actuators.



capacitor (EDLC) behavior except FC behavior owing to the addition of carboxylic MWCNTs. Because carbon nanotubes are regarded as ideal materials for electrochemical actuators based on the double-layer electrostatic mechanism.²³ And the calculated specific capacitance value 30% M-PEDOT:PSS electrode based actuators increases to 95.63 F g^{-1} , with the EDLC mechanisms providing the greatest contribution. In general, the significantly improvement in capacitance is attributed to the addition of carboxylic MWCNTs. On the one hand, the 30% M-PEDOT:PSS electrode owns high electrical conductivity which is helpful to acquire higher electrochemical energy storage capacity. On the other hand, the porous structure which formed after the addition of the carboxylic MWCNTs provides larger surface area and pore volume for adsorbing and accommodating ions, thus creating higher electrochemical performance. The electrochemical energy storage capacity plays a key role in emerging large bending deformation and blocking force of the actuator.

3.6 Actuation performance

The actuation performances of the actuators were examined with various conditions of harmonic electric inputs. The 30% M-PEDOT:PSS electrode based actuator performed symmetrical

and large deformation (optical images in Fig. 1c) under the input voltage of $\pm 2.5 \text{ V}$ at frequency of 0.1 Hz . Fig. 5a introduced the responses of the two types actuators under the input voltage of $\pm 2.5 \text{ V}$ at an frequency of 0.1 Hz . The peak to peak displacement of the 30% M-PEDOT:PSS electrode based actuator is up to 15.7 mm , which is about 2 times of PEDOT:PSS electrode based actuators. The high bending performance of the actuators is as a result of the high electrical, electrochemical and mechanical properties of the 30% M-PEDOT:PSS electrode. Additionally, peak-to-peak displacements were measured under different step voltage from ± 0.5 to 2.0 V peaks at 0.1 Hz frequencies. As shown in Fig. S3,[†] The degree of bending deformation is roughly affected by the applied voltage. Since ionic mobility is highly dependent on the electric potential field in the polymer matrix. The peak-to-peak displacements of the 30% M-PEDOT:PSS electrode based actuator attained at $\pm 2 \text{ V}$ is more than 3 times that at $\pm 0.5 \text{ V}$. Moreover, the peak-to-peak strain of the actuators were also measured and calculated along with various excitation frequencies range from 0.1 to 10 Hz . As shown in Fig. 5b, the bending performance of the 30% M-PEDOT:PSS electrode based actuator is impressively higher than pristine PEDOT:PSS electrode based actuator under all frequency conditions from 0.1 to 10 Hz . The inset graph of Fig. 5b revealed the harmonic responses of the 30% M-

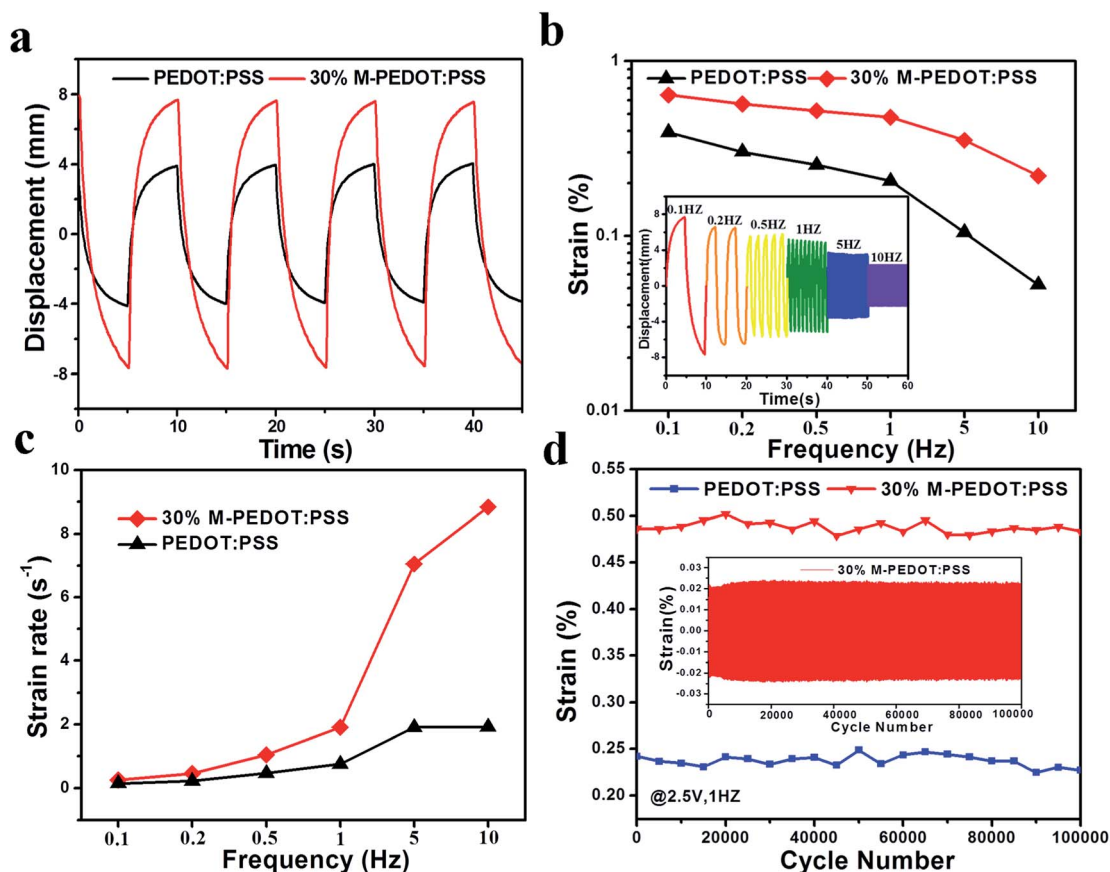


Fig. 5 Bending actuation performances of the PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators (a) displacement under 2.5 V at 0.1 Hz (b) strain under 2.5 V from 0.1 – 10 Hz (c) strain rate under 2.5 V from 0.1 – 10 Hz (d) durability results of harmonic response for 100 000 cycles under 2.5 V at 1 Hz .

PEDOT:PSS electrode based actuator with different excitation frequencies from 0.1 to 10 Hz. As the frequency decreased, the tip displacement of the 30% M-PEDOT:PSS actuator gradually increased due to the sufficient time to move the inner electrolytes of the TPU polymer toward the electrodes. Fig. 5c was the calculated strain rate of two types actuators at different frequency from 0.1 to 10 Hz. The strain rate of 30% M-PEDOT:PSS electrode based actuator is superior to that of the PEDOT:PSS electrode based actuators at high frequencies especially at 10 Hz ($8.83\% \text{ s}^{-1}$). In short, the porous structure of 30% M-PEDOT:PSS electrode led to easy and fast ion transport between the interfaces, thereby offering the capability to improve the bending actuation performance. Furthermore, the durability performance of the actuator was measured as seen in Fig. 5d. The 30% M-PEDOT:PSS electrode based actuator demonstrated hardly any significantly performance degradation for 100 000 cycles under an input voltage of $\pm 2.5 \text{ V}$ at 1 Hz. The strain of the 30% M-PEDOT:PSS electrode based actuator is maintained at as high as 99.4% compared to initial strain after 100 000 cycles, while that of the PEDOT:PSS electrode based actuator is only about 93.8%. This can be explained by the mechanical stability of electrode materials. In the process of actuation, the structure of PEDOT:PSS suffered certain damage due to frequent migration of ions. Therefore the strain of actuator decreased after 100 000 cycles. But for M-PEDOT:PSS electrodes, the porous structure and synergy effect between PEDOT:PSS and carboxylic MWCNTs gave them high mechanical stability, which greatly improved cycle stability of actuator. As shown in the inset of Fig. 5d. The 30% M-PEDOT:PSS electrode based actuator behaves excellent durability without significantly performance degradation over 100 000 cycles. The excellent cycle stability of 30% M-PEDOT:PSS electrode based actuator is also attributed to the tightly integrated between the electrode layer and electrolyte layer by hot press method. The 30% M-PEDOT:PSS electrode of the actuator is completely bonded on the surface of the TPU membrane without delamination after 100 000 cycles as shown in the cross-section SEM images in Fig. 1c.

The actuation performances enhancements of the M-PEDOT:PSS electrode based actuators can be explained by actuation mechanism. As we all know, the dimensional changes of actuators based on CPs resulting from electrochemical doping, transportation of dopant ions between the polymer matrix and the surrounding electrolyte solution.²⁴ When a electric field is applied between two conducting polymer PEDOT:PSS electrode layers based actuator, the $\text{PEDOT}^+\text{PSS}^-$ ion pairs can be formed *via* the electrochemical doping induced by the potential difference between the positive and negative electrodes.²⁵ And the cations or anions of the dissociated ionic liquid in TPU matrix will be absorbed into or excluded from the PEDOT:PSS electrode. In the positive electrode, the PEDOT network is oxidized and anions BF_4^- is incorporated inside the electrode according to the number of positive charges formed. Simultaneously, cations EMI^+ move and accumulate to the negative electrode driven by the electric field. Eventually, the actuator resulting in bending motion to the positive direction owing to the molecular size of cations EMI^+ is far larger than

anions BF_4^- . For 30% M-PEDOT:PSS electrode based actuator, benefit from rough and porous structure of electrode, more ions in TPU matrix can be incorporated and accumulate into it when electric field is applied. It will lead to actuation performance enhancements of M-PEDOT:PSS electrode based actuator compared to PEDOT:PSS electrode based actuator.

3.7 Efficiency of the conversion

Energy conversion efficiency is an important parameter to evaluate performance of the actuator. In order to attain high conversion efficiency, the actuator need to convert a large proportion of the input electrical energy into mechanical work. According to previous reports, the electromechanical conversion efficiency of an actuator is mainly determined by the mechanical and electrical losses within the material.²⁶ Liang *et al.* proposed a much detailed and convincing method to calculate the energy conversion efficiency of the actuator.²⁷ The energy conversion efficiency from the electrical to mechanical ($\eta_{\text{transduction}}$) of the actuator is calculated by the following equation:

$$\eta_{\text{transduction}} = \frac{0.5 Y \epsilon^2 f \times \text{Vol}}{V \times I_{\text{charge}}}$$

where Y is the Young's modulus, ϵ is the bending strain, f is the frequency of actuation, and Vol is the volume of the actuator, $V \times I_{\text{charge}}$ is the electric input power.

Fig. 6 was electromechanical conversion efficiency of the PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators at different frequencies. According to equation, the calculated $\eta_{\text{transduction}}$ are mainly affected by the bending strain and tensile modulus of the actuators. Benefit from the higher bending strain, the conversion efficiency of the 30% M-PEDOT:PSS electrode based actuators are greatly improved compared to PEDOT:PSS electrode based actuator. As the frequency increases from 0.1 to 10 Hz, the conversion efficiency of the 30% M-PEDOT:PSS electrode based actuator significantly increased and reached to the maximum value (1.04%) at 5 Hz, which is almost 55 times compared to pristine PEDOT:PSS electrode

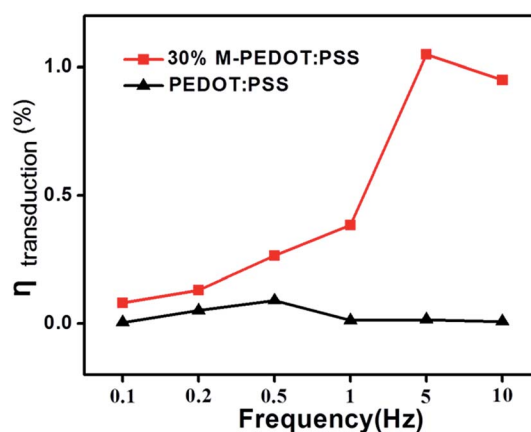


Fig. 6 Energy conversion efficiency of the PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators at different frequencies.



based actuator at 5 Hz. It is also noted that the conversion efficiency of pristine PEDOT:PSS electrode based actuator achieves the maximum value at 0.5 Hz. This is because the 30% M-PEDOT:PSS electrode based actuator displays more slowly strain attenuation compared with pristine PEDOT:PSS electrode based actuator along with the increase of frequency (as shown in Fig. 5b). The greatly improvements of conversion efficiency are attributed to porous structure of M-PEDOT:PSS electrode, which accelerate the migration of ions and further promote the conversion from electric energy to mechanical energy. Our results shown the higher electromechanical conversion efficiency compared with the IPMC and other conducting polymer actuators.^{27–31}

3.8 Blocking force

The blocking force is an important parameter of the actuators because it is crucial for future engineering applications. It has been reported that actuator with higher stiffness (higher tensile strength and modulus) can generate higher blocking force.³² Fig. 7a shown the blocking force of the pure PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators under the excitation of ± 2.5 V at frequency of 0.1 Hz. The 30% M-PEDOT:PSS electrode based actuator has the highest blocking force 0.54 mN, which is 193% enhancement than that of the pure PEDOT:PSS electrode based actuator (0.15 mN). Fig. 7b was the

blocking force of actuators under the excitation of 2.5 V DC. The blocking force of the 30% M-PEDOT:PSS electrode based actuator is up to 1.43 mN, which is three times that of the pure PEDOT:PSS electrode based actuator. The results demonstrates that the addition of carboxylic MWCNTs into PEDOT:PSS matrix significantly improve the blocking force of the actuator. We attributed the highly enhanced output force of the 30% M-PEDOT:PSS electrode based actuator to the high tensile modulus and electrochemical properties caused by M-PEDOT:PSS electrodes.

4. Conclusion

In this paper, we developed a high-performance electrochemical actuator based on M-PEDOT:PSS electrode. The good compatibility of PEDOT:PSS with MWCNT through π - π interaction and hydrogen bond provided much higher stiffness and strength for the M-PEDOT:PSS electrodes. In addition, the porous structure and high electric conductivity of M-PEDOT:PSS electrode could facilitated ion transmission and accumulation into it, which played a key role in improving actuation performance. The obtained M-PEDOT:PSS electrode based actuators possessed superior electro-chemo-mechanical properties and exceptionally high actuation performance compared with pure PEDOT:PSS actuators. It is found that the M-PEDOT:PSS electrode based actuator performed higher tensile modulus (82.18 MPa) and specific capacitances (95.63 F g^{-1}) with increasing the MWCNT content from 0 wt% to 30 wt%. Furthermore, the 30% M-PEDOT:PSS electrode based actuator are capable of producing large bending deformation (up to the strain of 0.64%), high strain rates (8.83 s^{-1}) and greatly force generation (1.43 mN) at 2.5 V. Beyond that, superb energy conversion efficiency (1.04%) of electrical input to mechanical output is obtained for the M-PEDOT:PSS electrode based actuators, which shows great promise in the development of next generation electroactive polymer actuator and practical application.

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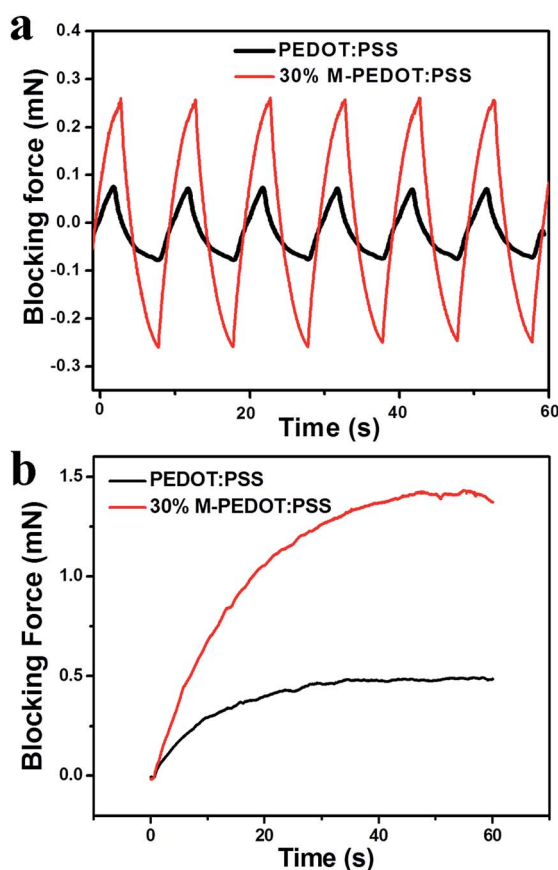


Fig. 7 Blocking force of PEDOT:PSS and 30% M-PEDOT:PSS electrode based actuators (a) under 2.5 V at 0.1 Hz. (b) under 2.5 V DC.



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