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Cyclic performance of viscoelastic dielectric elastomers with solid hydrogel electrodes

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Hydrogels containing electrolyte can work as ionic conductors to actuate dielectric elastomer (DE) artificial muscles. Based on a popular acrylic elastomer and an elegant design of a circular actuator, we study both theoretically and experimentally the cyclic performance of DE with solid hydrogel electrodes. The elasticity of solid electrodes constrains the maximum strain that is attainable for one cycle of triangular voltage, and it also diminishes the accumulated relaxation after many cycles of loadings.

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Dielectric elastomer (DE) has emerged as one of the competitive candidates for the realization of soft robotics, adaptive optics, actuators and sensors, and generators due to its unique features such as low cost, ease of production line manufacture or even 3D printing, silentness and large deformation capability. Among various commercial dielectric elastomers, the VHB series acrylic elastomers by 3M company, are extensively used for prototyping or real-purpose device fabrication. This type of elastomer, however, has long been identified to be viscoelastic. Viscoelasticity of DE would affect the performance of DE devices and should be taken into account during the design stage. This aspect is crucial for applications involving dynamic input voltages, such as loudspeaker or vibrator. Cyclic behavior is also closely related to the lifetime of DE. Several independent theoretical and experimental studies regarding the viscoelastic or dynamic responses of DE parallel in the literature, whereas rigorous correlation between theory and experiment is lacking.

Conventional DE devices mainly use carbon grease compliant electrodes to drive the elastomer. Other fluidic or solid electrodes have also been developed. All these electrodes are electronic conductors. Very recent work reported that hydrogels containing electrolyte can work as ionic electrodes to actuate a DE loudspeaker capable of operation at frequencies beyond 10 kilohertz and voltages above 10 kilovolts. Motivated by this new discovery, this letter presents both experiment and analysis of cyclic viscoelastic behavior of DE actuators with solid hydrogel electrodes.

We adopt the circular configuration to investigate the performance of commercial VHB 4910 (thickness 1mm) from 3M company. Figure 1(a) and (b) shows the schematics of the top
and side views of the experimental setup. A pair of solid hydrogel electrodes was glued on the center of the prestretched circular actuator and connected to external circuits. Cycles of triangular waveform voltages as shown in Figure 1(c) were supplied. The hydrogel electrode was synthesized following a standard procedure reported elsewhere by dissolving Acrylamide (AAm) monomer and lithium chloride (LiCl) powder into deionized water, and crosslinking AAm by N,N’-methylenebisacrylamide (MBAA). The obtained PAAm hydrogel containing LiCl is transparent, stretchable and conductive. The Young’s modulus of the hydrogel is 1.8kPa, the maximum rupture stretch is over 20, and the measured conductivity is about $10^2$ S/m (to be measured). Figure 1(d), (e) and (f) present the snapshots of DE with hydrogel electrodes corresponding to voltage-off state, voltage-on states for cycle 1 and cycle 30, respectively. As a comparison, Figure 1 (g), (h) and (i) present the snapshots of DE with carbon grease electrodes corresponding to voltage-off state, voltage-on states for cycle 1 and cycle 30, respectively.

The applied voltage given in Figure 1(c) ramps up from zero to its peak and then goes down to zero again to complete one cycle. During this process, the areal strain of the actuator increases firstly to reach its maximum strain and then decreases gradually. At the instant of voltage becomes zero, the elastomer retains a residual strain due to the viscoelasticity. The measured areal strain is recorded up to 30 cycles and three typical cycles are shown in Figure 2 (a). Given a fixed ramping velocity of applied voltage, say $V_{ramp} = 250$V/s in this case, the process in Figure 2(a) can be converted equivalently into a time history shown in Figure 2(b), whereby open blue circles denote experiment and red solid line is simulation. Figure 2(a) and (b) indicate that both residual and maximum strains increase as number of cycles increases, and eventually
they reach their steady-state values.

The simulation result in Figure 2(b) is obtained by adopting the previously developed rheological models of DE\textsuperscript{43} but with incorporation of elasticity of solid electrode.\textsuperscript{33} DE is approximated by a rheological model with two parallel units: one unit consists of a spring with shear modulus $\mu^\alpha$, and another unit consists of a spring with shear modulus $\mu^\beta$ and a dashpot with viscosity $\eta$. The solid hydrogel electrode is treated as a hyperelastic spring with shear modulus $\mu^\gamma$. The Gent model is used to represent all springs, with $J^\alpha_{\text{lim}}$, $J^\beta_{\text{lim}}$, $J^\gamma_{\text{lim}}$ being three corresponding extension limits, respectively.\textsuperscript{44} The original thickness of DE is $H_0$, and it becomes $H_i = H_o / \lambda_p^2$ after an equibiaxial prestretch, $\lambda_p$. The overall thickness of hydrogel electrodes are $H_2$, then the volume fraction of DE and electrodes are defined as $\phi_1 = H_i / (H_i + H_2)$ and $\phi_2 = 1 - \phi_1$, respectively. Incorporating the effects of electrode elasticity and assuming a homogeneous deformation state, the constitutive relation of the DE-hydrogel composite system reads

$$\lambda_p^{-1} \sigma_p \lambda + \phi_1 \varepsilon E_0 V^2 \lambda^3 = \phi_1 \left[ \frac{\mu^\alpha (\lambda^2 - \lambda^{-4})}{1 - (2 \lambda^2 + \lambda^{-4} - 3) / J^\alpha_{\text{lim}}} + \frac{\mu^\beta (\lambda^2 \xi^{-2} - \lambda^{-4} \xi^{-4})}{1 - (2 \lambda^2 \xi^{-2} + \lambda^{-4} \xi^{-4} - 3) / J^\beta_{\text{lim}}}ight] + \phi_2 \left[ \frac{\mu^\gamma (\lambda^2 \xi^{-2} - \lambda^{-4} \xi^{-4})}{1 - (2 \lambda^2 \xi^{-2} + \lambda^{-4} \xi^{-4} - 3) / J^\gamma_{\text{lim}}}ight],$$

(1)

where $\varepsilon = 4.11 \times 10^{-11} \text{F/m}$, the permittivity of DE; $\lambda$, the overall stretch of DE (the same as the stretch of spring $\alpha$); $V$, applied voltage; $\sigma_p = \frac{\mu^\alpha (\lambda_p^2 - \lambda_p^{-4})}{1 - (2 \lambda_p^2 + \lambda_p^{-4} - 3) / J^\alpha_{\text{lim}}}$, the stress needed to attain a prestretch, $\lambda_p$, and set as a constant in simulation; $\xi$, the stretch in the dashpot. The stretch of spring $\beta$ is determined by adopting a multiplication rule as $\lambda / \xi$. The dashpot is modeled as a Newtonian fluid, and the rate of deformation is related to stress via
\[
\frac{d\xi}{dt} = \frac{1}{\eta} \left( \mu^\beta \left( \frac{\lambda^2 \xi - \lambda^4 \xi^4}{\xi} \right) \right) \left( 2\lambda^2 \xi^{-2} + \lambda^4 \xi^{-4} - 3 \right) / J_{\text{lim}}^\beta.
\]

Equation (1) recovers the relation for a DE with compliant electrode, e.g., carbon grease, by letting \( \phi_2 = 0 \) and \( \phi_1 = 1 \), where electrode thickness and the constraining effects of electrodes are negligible. The parameters used in our simulation is as follows: \( \mu^\alpha = 24.5 \text{kPa,} \) \( \mu^\beta = 40 \text{kPa,} \) \( \mu^\gamma = 0.6 \text{kPa,} \) \( J_{\text{lim}}^\alpha = 110, \) \( J_{\text{lim}}^\beta = 55, \) \( J_{\text{lim}}^\gamma = 500, \) \( \eta = \tau_\nu \cdot \mu^\beta, \) \( \tau_\nu \) is a parameter reflecting the relaxation time, \( \tau_\nu = 400s. \)

It is intriguing to compare the cyclic performance of DE with hydrogel electrodes to that of DE with carbon grease electrodes. This comparison gives clearly the effects of elastomeric electrodes on the viscoelastic deformation of DE. Figure 3(a) plots both simulation and experimental results of DE with hydrogel and carbon grease electrodes. The ramping velocity is 250V/s and the amplitudes of applied voltage are 4kV for 0.5mm thick hydrogel electrodes, and 3.5kV and 4kV for carbon grease electrodes. Even for a soft hydrogel electrode (Young’s modulus 1.8kPa herein), its constraining effect is obvious: the plateau of the steady-state maximum strain of DE with hydrogel electrode at 4kV is less than half of that with carbon grease electrode at the same voltage; it is even less than that with carbon grease at 3.5kV. Both experimental and simulation results prove this constraining effect. For not very large deformation, such as DE with hydrogel electrodes or DE with carbon grease electrodes working at not high voltage, e.g., less than 3.5kV, there is a good agreement between the experiment and model prediction. For DE with carbon grease electrode at 4kV, the deformation is pretty large. The constant approximation of mechanical loading, \( \sigma_p \), does not hold any more. That is the
reason why there is a discrepancy between theory and experiment. Rigorous calculation with a varying mechanical loading needs solution of a boundary-value problem, and the details can refer to reference.\textsuperscript{33}

Ramping velocity also affects the cyclic performance of DE, and systematic experiments for both hydrogel electrodes and carbon grease electrodes have been carried out. Figure 3(b) and (c) show the evolution of maximum strain for carbon grease electrodes and hydrogel electrodes, respectively. The peak value of the applied voltage is set at the same, 4kV, but the ramping velocities are varied as 250V/s, 500V/s and 1000V/s. The cyclic maximum strain evolves gradually, starting from the first maximum strain that is achieved after cycle 1, to its steady-state value indicated by the plateau. The shift from the first maximum strain to the plateau of maximum strains depends on ramping velocity and property of electrodes. For carbon grease electrodes, the shifts from the first to the steady-state maximum strains are 0.26, 0.17 and 0.15, respectively, corresponding to ramping velocities 250V/s, 500V/s and 1000V/s. The decrease of shift to steady-state plateau with increasing ramping velocity is due to the relaxation of elastomer: If ramping velocity is low, the elastomer has enough time to relax when one cycle is completed, and the accumulated relaxation will become larger after many cycles; when applied voltage ramps up and down very quickly, the elastomer does not have enough time to response such that the relaxation does not fully develop and exhibits smaller shift. For hydrogel electrodes, however, the shifts are about 0.06~0.07, approximately constant for the three ramping velocities. This implies that the elasticity of solid electrodes diminishes the viscoelastic deformation of DE.
In summary, we study both theoretically and experimentally the cyclic performance of an acrylic elastomer. Polyacrylamide hydrogels containing LiCl salt work as ionic conductors to actuate the dielectric elastomer. The elasticity of the solid electrodes and the ramping velocity of applied triangular voltages are found to influence the residual and maximum strains of a circular actuator. Compared with carbon grease electrodes, solid electrodes constrain electro-actuation deformation and diminish the viscoelastic relaxation of elastomer. Good correlation between experiment and simulation is achieved. These results help the analysis and assessment of performance of dielectric elastomer actuators subject to dynamic electrical loadings.

ACKNOWLEDGEMENTS

This research is supported by Natural Science Foundation of China (grants 61025002, 11072185, 11372239 and 11021202). ZS acknowledges the support of NSF MRSEC (DMR-0820484) and visiting appointment at the International Center for Applied Mechanics.
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Metal Ion Implantation for the Fabrication of Stretchable Electrodes on Elastomers

Ordered Zigzag Stripes of Polymer Gel/Metal Nanoparticle Composites for Highly Stretchable Conductive Electrodes


Self-healing electrodes for dielectric elastomer actuators


Flexible and stretchable electrodes for dielectric elastomer actuators


42Baohong’s paper.


A New Constitutive Relation for Rubber
Figure 1. Experimental setup of a circular DE actuator. (a) and (b): schematics of the top and side views of the setup; (c): applied cyclic triangular waveform voltage in experiment; (d), (e) and (f): photos of DE with hydrogel electrodes corresponding to voltage-off state, voltage-on states for cycle 1 and cycle 30, respectively; (g), (h) and (i): photos of DE with carbon grease electrodes corresponding to voltage-off state, voltage-on states for cycle 1 and cycle 30, respectively. The hydrogel electrodes are stretchable and transparent. The white scale and the ruler are used for calculating the areal strain.
Figure 2. Viscoelastic performance of a circular DE with hydrogel electrodes under supply of a series of cyclic voltages. (a): Areal strain versus voltage for three typical cycles (cycles 1, 2 and 30, respectively; ramping velocity: 250V/s); (b): Comparison of experiment and simulation of the time history of areal strain (open blue circle: experiment, red solid line: simulation; ramping velocity: 250V/s).
Figure 3. Effects of elastomeric electrodes on the maximum areal strain of DE actuators. (a): Comparison of cyclic maximum strains for carbon grease electrodes and 0.5mm thick hydrogel electrodes. The applied ramping velocity is fixed to 250V/s. The markers are experimental results while the red solid line represents the simulation result; (b) The effects of ramping velocity on the evolution of cyclic maximum strains. The amplitude of applied voltage is fixed to be 4kV and the ramping velocities are varied to be 250V/s, 500V/s and 1000V/s, respectively. The marked shift is the difference between the first and the steady-state maximum strain. HYG
denotes hydrogel electrodes while CG denotes carbon grease electrodes.