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Abstract- We report a plasmonic self-powered UV detector based on triboelectric nanogenerators, for the first time. The proposed structure benefits from plasmon-assisted photoresponse in reduced graphene oxide/Ag nanoparticles heterostructure, serving as a high sensitive photoconductive layer in the fabricated triboelectric nanogenerator. Ag nanoparticles enhance the contacting surface area of the triboelectric nanogenerator, and improve the output open circuit voltage by a factor of about 5. The presented triboelectric nanogenerator operates as an active self-powered UV detector, so that the open circuit voltage is modulated by the UV illumination. The open circuit voltage sensitivity ($\Delta V_{oc}/V_{oc}$) is measured as about 50 % in response to UV illumination, with illumination power density of 500 $\mu$W/cm$^2$.

Index Terms: Triboelectric nanogenerator, plasmon, reduced graphene oxide, Ag nanoparticles, self-powered photodetector;

I. INTRODUCTION

Emergence of the miniature and low-power electronic systems has highlighted the importance of generating the required electric power from environmental energies such as sun light, motions and vibrations. On the other hand, using battery power involves environmentally unfriendly materials, and restricts the lifespan, stability of power, and scalability of the systems. Thus mechanical energy harvesting has attracted massive interests during the last decade. Among various principles for harvesting mechanical energy [1-3], triboelectric nanogenerators (TENGs) have shown promising properties and wide applications [4-11], since their first invention in 2012. TENGs have shown high energy conversion efficiencies [12], low cost fabrication process, and integration potential [13]. In this field of research, self-powered devices and sensors have been proposed as one of the most attractive applications of TENGs [14-17]. However, there are limited reports on self-powered photodetectors based on TENGs [18-20]. Here, for the first time, we propose a plasmonic self-powered UV detector, benefiting from reduced graphene oxide (rGO)/Ag nanoparticles (Ag NPs) heterostructure. The proposed UV detector is a self-powered active sensor that can generate electrical signal as a response to UV-illumination and environmental tapping energies.

II. FABRICATION PROCESS AND OPERATION PRINCIPLE

First, Ag NPs are synthesized by solution method, in which silver nitrate ($\text{AgNO}_3$) is the precursor and sodium borohydride ($\text{NaBH}_4$) serves as the reducing agent and stabilizer [21]. Then, graphene oxide (GO) is synthesized by modified Hummer’s method [22], from graphite powder. The resultant GO powder is dissolved in DI water and sonicated for 30 minutes. To prepare polydimethylsiloxane (PDMS) layers, silicone elastomer base and the curing agent\(^1\) (Sylgard184 from Dow Corning Company) are mixed at the ratio of 10:1, for 10 minutes. The mixture is degassed from the created bubbles after a 10-minute period of relaxing. Then, the prepared PDMS is spin-coated on a pre-cleaned indium tin oxide (ITO) coated glass, and cured at 100°C for 35 minutes. Fig. 1(a) displays the fabrication process of the proposed TENG, schematically. The PDMS surface on the ITO electrode (Fig. 1.a) is treated by an oxygen plasma (Fig. 1.b) in order to be rendered hydrophilic, which is necessary for the uniform coating of Ag NPs from its aqua solution. RF plasma power is 50W, oxygen flow rate is 100sccm, and plasma time duration is about 1 minute in this step. Then the prepared solution of Ag NPs is drop casted on the surface of the PDMS layer, and dried at 60°C. This cycle is repeated for six times (Fig. 1.c). Then the synthesized rGO sheets are deposited on the sample by dip coating, and dried (Fig. 1.d). The schematic operation principle of the fabricated TENG in contact/separation mode is illustrated in parts (e-i) of Fig. 1. This operation is based on coupling between electrostatic effect and triboelectric effect. As shown in Fig. 1.e, rGO and steel electrodes are two contacting surfaces in the fabricated TENG. When these two surfaces come into contact, a chemical bond is formed between them and charge injection occurs from metallic electrode to the rGO surface, due to their electrochemical potential difference (Fig. 1.f). Hence, a positive charge is formed on steel, while a negative charge is formed on the surface of rGO [23]. When the contacting surfaces are released, some of the bonded atoms keep extra

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1. Sylgard184 from Dow Corning Company

A. Keshvari (arvin.keshvari@gmail.com) and S. Darbari (s.darbari@modares.ac.ir) are both with Nano Sensors and Detectors Lab., and Nano Plasmonphotonic Research Group, Faculty of Electrical and Computer Engineering, Tabriat Modares University, Tehran 1411713116, Iran. M. Taghavi (majid.taghavi@bristol.ac.uk) is with Department of Engineering Mathematics, University of Bristol, Bristol, BS8 1UB, UK.
electrons, while some have a tendency to give them away, which is believed to be the reason behind creating tribo charges. After separation, a lower potential is established at the back electrode (ITO). Thus, electrons are driven from the ITO electrode to the steel electrode (Fig. 1.g), until the charges on the steel plate are completely screened (Fig. 1.h). By applying force during the next pressing step, electrons will flow in opposite direction to rebalance the charges (Fig. 1.i). After a complete contact, charges on two surfaces will be equal once again (Fig. 2.f), and the presented cycle be repeated.

III. RESULTS AND DISCUSSIONS

Fig. 2.a illustrates the TEM image of the synthesized Ag NPs with an average size of about 10 nm. We drop casted and dried the prepared Ag NPs on a glass substrate for 6 cycles, then achieved absorption spectrum, which is shown in Fig. 2.b. The plasmonic peak relating to the Ag NPs is observed at 400 nm. Fig. 2.c presents the TEM image of the prepared rGO sheets. The Raman spectroscopy result of the rGO sheets is shown in Fig. 2.d, where the typical trace of rGO sheets at 1312 cm⁻¹ and 1580 cm⁻¹ are shown by arrows, corresponding to D band and G band, respectively [24]. It is well established that G band is attributed to the size of sp² carbon-type domains, while the D band refers to the structural imperfections in the hexagonal graphitic layers, including hydroxyl or epoxide groups in rGO sheets [24]. Fig. 2.e shows the absorption spectrum of rGO/Ag NPs heterostructure, where the absorption peak of Ag NPs at about 400 nm is detectable through the rGO absorption spectrum and is shown by an arrow. Fig. 3.a and Fig. 3.b display the measured short circuit current (Isc) and open circuit voltage (Voc) of a bare TENG with PDMS thickness of 20μm before coating Ag NPs. In Fig. 3.c, effect of adding Ag NPs on Voc of the TENG is studied. An enhanced open circuit peak-to-peak voltage is observed, by a factor of about 5 in comparison with the bare sample (Fig. 3.b). This enhancement is attributed to the increased effective area of the contact surfaces in the presence of nanoparticles [25]. Fig. 3.d highlights the impact of varying PDMS thickness (tPDMS) on the output open circuit peak-to-peak voltage (Vp-p) of the bare TENGs without Ag NPs. It can be observed that increasing tPDMS from 20μm to 230μm increases the open circuit peak-to-peak voltage from 3.5V to 10V. This observation can be attributed to the modulation of electrostatic induction by changing thickness of the PDMS layer, while tribo charges on its surface do not change. In other words, a fixed amount of generated tribo charges causes higher (lower) potential difference across a capacitor with lower (higher) capacity, for the case of thicker (thinner) PDMS layer. This observation is in agreement with the previous theoretical studies of Niu [26].

Fig. 3.e shows the output Voc signals, corresponding to different tPDMS values, investigated in part (d). Fig. 3.f displays Voc signal of the bare TENG with tPDMS=230μm, in response to continuous pressing/releasing cycles, revealing an acceptable stability for the fabricated TENG. At the next stage, we present the fabricated self-powered UV-detector, based on rGO/Ag NPs heterostructure. First we investigate the photoresponse (∆G/G₀) of the rGO/Ag NPs layer (Fig. 4.a) in response to a Sun-Kraft UV lamp, revealing an enhancement factor of 2.6, comparing with the pristine rGO layer.

Fig. 4. Photoresponse of (a) rGO/Ag NPs, and (b) rGO layers to Sun-Kraft UV source. (c) Photoconductive sensitivity of rGO/Ag NPs layer in response to different visible wavelengths.
layer (Fig. 4.b). This improved photoresponse is believed to be related to the excitation of localized surface plasmons in Ag NPs that can result in a kind of plasmon-assisted photodesorption of residual oxygen groups from the surface of rGO sheets. Similarly, Tan et al. have previously proved that this photodesorption effect in rGO/Ag NPs heterostructures dominates the original photogeneration in the bare rGO sheets [27]. This plasmon-assisted desorption is believed to be originated from lowering of the adsorption strength and activation energy barrier for desorption, at the presence of an external electric field [28]. Moreover, Fig. 4.c presents the photoconductive sensitivity of rGO/Ag NPs layer in response to different visible wavelengths, which agrees with the reduction of Ag NPs’ absorption spectrum in visible range (Fig. 2.b). Here, we benefit from the plasmon-assisted enhanced UV-response of rGO/Ag NPs heterostructure for realizing self-powered UV-detector. For this purpose, we incorporate the rGO/Ag NPs heterostructure as an UV-sensitive photoconductor in the configuration of fabricated TENG. Fig. 5.a indicates the applied UV-sensitive configuration, in which external force is exerted on the top steel plate and UV source is exposed from the bottom. Samples with 20μm-thick PDMS layers are investigated and a Sun-Kraft UV lamp (model A-1) has been utilized as the source. As depicted in Fig. 5.a, we have electrically connected the top steel electrode to the rGO/Ag NPs layer in the UV sensing structure in order to effectively incorporate the rGO/Ag NPs photoresistor in electrical configuration of the fabricated TENG. Fig. 5.b exhibits the response of the investigated UV-sensitive TENG based on rGO/Ag NPs, to UV illumination. As shown in this figure, the observed open circuit peak-to-peak voltage is near 2.5V before UV-illumination, however it is reduced to about 1.25V when illuminated, and returns to the initial value by turning the UV source off. The incident light power is about 500μW/cm² in this experiment. Fig. 5.c manifests a simplified equivalent circuit model for the presented UV-sensitive TENG, in which c₁ and c₂ represent the capacitances between the rGO/Ag NPs layer with the top steel electrode and bottom ITO layer, respectively. The capacitive value of c₁ varies during the pressing/releasing cycles, while c₂ is a fixed capacitor. The photoresistor (Rph) represents the UV-sensitive rGO/Ag NPs layer which plays as a shunt resistance for c₁. The output VOC is measured between the top steel electrode and the bottom ITO electrode, as shown in Fig. 5.a. Regarding the equivalent circuit in dark condition, a high shunt resistance results in the typical operation of the TENG as discussed in the Fabrication Process and Operation Principle section. However, when the sample is exposed to UV illumination, the reduced shunt resistance causes the tribo charges on rGO layer to discharge and VOC to decrease. Fig. 5.d illustrates the open circuit sensitivity (=ΔVp-p/Vp-p), in response to different incident light powers of the utilized Sun-Kraft UV lamp. It can be observed that increasing incident illumination power from 70μW/cm² to 500μW/cm² results in raising sensitivity from rom 10% to about 50%. To elaborate the visible response of the realized self-powered UV detector, the open circuit sensitivity has been measured in response to three different monochromatic visible sources with the same illumination powers. Fig. 5.e illustrates the photoresponse spectrum, confirming that the output sensitivity decreases when the incident wavelength increases that is consistent with the plasmonic absorption spectra of Ag NPs (Fig. 2.b). Benefiting from the plasmonic behavior of Ag NPs in this approach, we have realized a high sensitive self-powered UV detector, which is blind to Deep-UV and visible range, and possesses UV-selective photoresponse. Furthermore, tunability of the UV-sensitive range of this detector may be achieved by controlling size distribution of Ag NPs, which can be proposed as another potential for the presented UV detector.

IV. CONCLUSION

We reported a plasmonic self-powered UV detector based on TENG, by taking advantage of rGO/Ag NPs heterostructure as the UV-sensitive layer. It is proved that the rGO/Ag NPs layer shows a plasmon-assisted improved photoresponse to UV-illumination. Presence of Ag NPs improves the output VOC of the triboelectric nanogenerator by a factor of about 5, which can be related to the enhanced surface area of the contacting electrodes. When exposed to UV illumination with light power of 500 μW/cm², the output VOC of the fabricated triboelectric nanogenerator responds with a sensitivity (ΔVp-p/Vp-p) of about 50%. In summary, we have achieved a high sensitive and UV-selective self-powered plasmonic detector based on TENGs, for the first time.

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