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Development of pressure sensors which display high sensitivity, are flexible, inexpensive, and easy to manufacture has drawn significant interest due to diverse applications such as tactile skin sensors (e.g. "electronic skin"), pulse detectors, speech recognition elements, and others. While varied technologies and molecular constructs have been demonstrated for pressure sensing, considerable conceptual and technical challenges still hamper broad implementation of many such systems. A novel flexible piezoresistive sensor comprising a conductive Au-coated elastomeric polymer sponge is presented. The piezoresistive sponge is prepared through a simple chemical route in which Au nanoribbons are spontaneously grown upon amine-functionalized polyurethane framework. The Au nanoribbon layer coats the internal surfaces within the polymer pore network, resulting in electrical current modulation upon pressure application / release through changes in the overall contact areas between the conductive surfaces. The Au-polyurethane piezoresistive sensor exhibits excellent functionalities, including enhanced sensitivity, low detection threshold, high fidelity, and physical stability. Application of the sensor is demonstrated for high resolution monitoring of wrist arterial pulses.

Introduction

Sensitive, easy to manufacture, and inexpensive pressure sensors have considerable scientific, technological, and commercial impact as they are used in many practical applications and devices, such as tactile skin sensors (e.g. "electronic skin"), speech recognition elements, and others.¹⁻⁸ Varied molecular systems and technologies have been developed for pressure sensing including field effect transistor devices, ⁹⁻¹² capacitance-based sensing, ¹³⁻¹⁵ piezoelectric sensing, ¹⁶⁻²³ and piezoresistive sensing²⁴⁻²⁷. Nevertheless, considerable technical challenges still exist. Specifically, in many instances, sensing devices exhibit complex design and/or elaborate, expensive manufacturing routes, limiting the practical and commercial potential.

Elastomeric polymers coupled with conductive elements have been recently shown to constitute useful platforms for chemically-based pressure sensors²⁸⁻³⁰ Such assemblies rely on the intrinsic flexibility of the polymer frameworks to produce piezoresistive sensors in which the compression / release of the matrix modulates the electrical resistance.³¹⁻³³ Porous polymers or "sponges" have been particularly attractive in that regard since such materials provide readily utilized and inexpensive host matrixes for embedding the

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conductive layers.^{34,35} Pressure sensors based upon elastomeric polymer sponges encapsulating carbon nanotubes,³⁶ graphene,³⁷ or metals³⁸⁻⁴⁰ have been reported. However, despite the promise of such "bottom-up" strategies, pressure sensors constructed through such schemes have displayed in many cases relatively low sensitivity, and their production generally comprise of multistep and complex chemical processes.

We present a novel, sensitive piezoresistive sensor comprising a conductive Au-coated elastomeric polymer sponge. The piezoresistive sponge was prepared through a simple two-step chemical route based upon self-assembly of Au nanoribbon thin film onto an amine-functionalized polyurethane (PU) framework. The Au-PU piezoresistive sensor exhibited excellent pressure sensing parameters, including low detection threshold, signal reproducibility, and physical stability. We demonstrate application of the sensor for monitoring and waveform analysis of wrist arterial pulses.

Experimental Section

Materials

HAuCl₄·3H₂O, KSCN, branched polyethylenimine (b-PEI) with average Mw ~25,000 by LS average Mn ~10,000 by GPC, N-Hydroxysuccinimide (NHS), 2-(N-morpholino)ethane sulfonic acid (MES) and Conductive silver paint were purchased from Sigma-Aldrich. H₂SO₄were purchased from Bio Lab (Israel). 1-(3-dimethylaminopropyl)-3-ethylcarbodiimide (EDC) 98%, and Chromium trioxide were purchased from Alfa Aesar (USA). All above listed reagents were used as received. Water used in the experiments was doubly purified by a Barnstead D7382

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water purification system (Barnstead Thermolyne, Dubuque, IA), at 18.3 $m\Omega$ cm resistivity.

Synthesis of the KAu(SCN)₄ Complex

1 mL of an aqueous solution of $HAuCl_4 \cdot 3H_2O$ (24 mg/mL) was added to 1 mL of a solution of KSCN in water (60 mg/mL). The precipitate formed was separated by centrifugation at 4000 g for 10 min. The supernatant was decanted, and the residue was dried at room temperature.

Amine modification and fiber fracture of the PU sponges

Commercially available PU sponge (average pore size 0.1-0.3 mm; porosity of around 93%) was cut into cuboids and cleaned in deionized water (DIW), ethanol and acetone successively. In order to perform surface amine modification and fiber fracture of the PU sponges, the following steps were performed: carboxylation of PU sponges and fiber fracture of the PU sponges were achieved at the same time by immersion of PU sponges in a solution containing CrO_3 (100 g/ L) and H_2SO_4 (98 wt %) (100 g/L) for 90 seconds. Subsequently, the sponges were washed with water (10 times). Amine grafting were thereafter carried out under the following experimental procedure: the PU sponges were immersed in a solution of 2 g b-PEI in 20 ml of 0.1 M MES buffer, pH=6, containing 105mM EDC and 100mg of NHS for 24 h at 23 °C under shaking. Afterwards the PU sponges were washed with water with continuous shaking 5 times.

Growth of Au films on amine-modified PU sponges

An aqueous solution of Au(SCN)₄ (3 mg/ml) was prepared in doubly distilled water. The amine-modified PU sponges were immersed in the solution and kept at 4°C for 1 days. After the gold growth had reached completion, the sponges were taken out of the growth solution and washed thoroughly with water to remove non-reacted materials and subsequently dried at room temperature. In order to enhance the conductivity of the Au/ PU sponges, the Au/ PU sponges were inserted to a plasma cleaner, PDC - 32G, Harrick plasma, and were exposed to plasma for 3 min at high RF (radio frequencies) and power of 18W.

Fabrication of the pressure sensing device and arterial pulse sensor

Two pieces of copper sheets connected with copper wires were placed onto the top and bottom surfaces of as-prepared Au/PU sponge. The size of the pressure sensor was 2 mm × 2 mm × 1mm. The copper electrodes and the as-prepared Au/PU sponge were connected with conductive silver paste. Pressure was applied or released by adding or reducing different weights onto the device. For fabrication of pulse sensor device, a piece of Au/Pu sponge (1cm×1cm×0.5cm) was sandwiched between 2 copper electrodes which is connected with copper wires, then it is fixed onto the radial artery of the test person by adhesive tape.

Instruments and characterization

Scanning electron microscopy (SEM) was conducted with a JEOL (Tokyo, Japan) model JSM-7400F scanning electron microscope equipped with EDS (Thermo Scientific). Powder X-ray diffraction (XRD) measurements were carried out on a Panalytical Empyrean powder diffractometer equipped with a parabolic mirror on the incident beam providing quasimonochromatic Cu K α radiation ($\lambda = 1.54059$ Å) and an X'Celerator linear detector. X-ray photoelectron spectroscopy (XPS) analysis was conducted using a Thermo Fisher ESCALAB 250 instrument with a basic pressure of 2× 10–9 mbar. The samples were irradiated in two different areas using monochromatic Al K α , 1486.6 eV X-rays, using a beam size of 500 µm. Electrical measurements were performed on a Keithley 2400 sourcemeter.

Results and Discussion

Fabrication of the Au nanoribbon-coated polymer pressure sensor

Fig. 1A depicts a scheme of the two-step assembly process for construction of the Au/PU piezoresistive sensor (detailed protocols are provided in the Experimental Section). The starting material was PU *sponge* – an inexpensive, highly deformable and flexible porous polymer which was previously employed as a platform for piezoresistive pressure sensors.⁴¹ The PU matrix was functionalized with amine (NH₂) moieties through reaction with polyethileneimide (PEI; many other routes for amine functionalization of polymer networks are also available⁴²). ζ -potential measurements confirm coating of the PU sponge with amine moieties following functionalization (see Fig. S1 in ESI⁺).

The amine-functionalized PU substrate was then immersed in an aqueous solution of gold thiocyanate, which produced Au-coated PU sponge after drying and brief air-plasma treatment⁴³ (Fig. 1B). Spontaneous reaction between watersoluble Au(SCN)₄ and amine-displaying surfaces has been recently demonstrated as a versatile and effective technology for creation of conductive Au nanostructured films.⁴⁴⁻⁴⁶ This simple chemical process is based upon electrostatic attraction between the Au(SCN)₄ complex and surface-displayed amine residues, and subsequent reduction of the Au³⁺ ions by the thiocyanate ligands, which ultimately produce conductive "gold nanoribbon" networks.⁴⁴ Notably, this synthesis route employs readily-available and inexpensive reagents, and does not require co-addition of reducing agents nor nanostructure templates. As depicted in Fig. 1C, the resultant Au/PU sponge comprised the core of the piezoresistive sensor, as its electrical resistance is modulated through compression / release of electrodes placed at either side (Fig. 1C).

Characterization of the Au/PU system

Table S1 provides the mechanical properties of the PU matrix before and after gold deposition. The details outlined in Table S1 indicate that after chemical oxidation, the rigid PU sponge became softer because of fiber fracture (i.e. decrease of Young's modulus).

However, the deposition of gold coating did not significantly alter the mechanical properties of the sponge. The Au-coated PU



Fig. 1 Preparation of the Au-coated polyurethane piezoresistive sensor. A) Schematic illustration of the fabrication process. The PU sponge is functionalized to display amine residues. Subsequent incubation with Au thiocyanate results in thin gold coating of the polymer framework. B) Photograph of the Au/PU sponge. C) Schematic illustration of the operating principle of the piezoresistive sensor comprising the Au/PU sponge.

sponge was characterized by several analytical techniques (Fig. 2). Fig. 2A depicts scanning electron microscopy (SEM) images of the material. The SEM image shows the fractured porous framework of the PU matrix; fracturing of the polymer scaffold was apparent following carboxylation (acid treatment) required for amine functionalization. While fracturing of the PU framework is not essential for achieving piezoresistive properties, it is believed to contribute to the flexibility and high sensitivity response to pressure changes.⁴¹ Indeed, we have found that fracturing improved the mechanical and sensing properties of the Au/PU matrix. Similar enhanced pressure sensitivity was recently reported, accomplished through hydrothermal/mechanical PU fracturing. It should be noted, however, that the 90 second acid treatment reported here is simpler and significantly less time-consuming than the reported process.41

The magnified area shown in the SEM image in Fig. 2A (right image) shows a dense coating of *Au nanoribbons*; the corresponding Energy-dispersive X-ray spectroscopy (EDS) spectrum in Fig. 2B further confirms that gold deposition occurred within the porous framework of the PU sponge. The presence of *metallic* crystalline gold was verified by x-ray photoelectron spectroscopy (XPS, Fig. 2C) and x-ray diffraction (XRD, Fig. 2D) experiments. Specifically, the XPS spectrum in Fig. 2C indicates that the nanoribbon coating comprised of Au⁰, giving rise to peaks at 84.4 eV and 88.1 eV.



Fig. 2 Characterization of the Au films grown on the polyurethane sponge. A) Scanning electron microscopy (SEM) images of Au/PU sponge in different magnifications. B) The energy-dispersive X-ray spectroscopy (EDS) spectrum corresponding to the Au/PU sponge shown in A. C) X-ray photoelectron spectrum (XPS) of the Au/PU sponge; the two Au⁰ peaks in the (4f) region are highlighted. D) X-ray diffraction (XRD) pattern of the Au/PU sponge; the reflections corresponding to Au crystal planes are apparent.

The XRD pattern in Fig. 2D corroborates the XPS result and further underscores the *crystalline* organization of the gold nanoribbons. Indeed, the XRD pattern in Fig. 2D reveals the distinctive reflections corresponding to crystal planes of metallic Au. Together, the spectroscopy data in Fig. 2 confirm the occurrence of crystallization / reduction of gold, resulting in deposition of crystalline metallic Au⁰ through spontaneous reaction between the amine-functionalized PU sponge surface and Au(SCN)₄⁻. Importantly, no reducing agents were co-added to the reaction mixture – in contrast to the significant majority of experimental schemes reported for controlled synthesis of Au nanostructures.⁴⁷ This distinction reflects the contribution

of the *thiocyanate ligands* as the electron-donating entities in the $Au(SCN)_4$ system.⁴

The role of plasma treatment in the fabrication process of the new piezoresistive sensor is twofold. First, plasma treatment was designed to remove the organic residues (-SCN, -CN) in the gold film. The oxygen species created in the plasma (O_2^+ , O_2^- , O_3 , O, O^+ , O^- , ionized ozone, metastable excited oxygen) and free electrons react with organic contaminants to form H₂O, CO, CO₂, and lower molecular weight hydrocarbons. These compounds have relatively high vapour pressures and are evacuated from the chamber during processing. Second, the electrons produced in the plasma gas react with the deposited gold cations, reducing them to



Fig. 3 Pressure-dependent electrical properties of the Au/PU sponge. A) Currentvoltage (I-V) curves of Au/PU sponges recorded at different pressures. B) Pressuredependent current recorded at different pressures (indicated). C) Pressure-response resistance plots for the Au/PU sponge pressure sensor. R_p corresponds to the resistance at pressure p, while R₀ is the pressure without pressure applied. The indicated slopes reflect the pressure sensitivities of the system (rates of resistance change upon pressure modulation). D) Stability of the sensor is demonstrated by repeatedly applying and releasing a pressure of 1 KPa in 1000 cycles.

crystalline gold species. Powder x-ray diffraction and X-ray photoelectron spectroscopy data (Fig. S2, ESI⁺) confirm that plasma treatment effectively reduced the Au cations.

Pressure-dependent electrical properties of the Au/PU sponge of the Au/PU system

Fig. 3 highlights the piezoresistive properties of the newly assembled Au/PU sponges. Fig. 3A depicts the current-voltage (I-V) curves recorded upon placing the sponge between two flat copper electrodes and applying different pressures (according to the experimental scheme in Fig. 1C). The linear I-V curves and milliamper current range shown in Fig. 3A attest to the excellent conductivity attained in the Au/PU sponge system. The calculated conductivity of the Au/PU sponge was about 0.24S m⁻¹ (no applied pressure). Furthermore, Fig. 3A shows that the slopes of the I-V curves were dependent upon the pressures applied, reflecting the relationship between the sponge resistance and pressure exerted between the two planar electrodes. The pressure- dependent electrical conductivity shown in Fig. 3A is explained by the increased

contact areas among the Au nanoribbon films attained upon compressing the sponge (e.g. contracting the internal pores), thereby enhancing electron transport and reducing the overall resistance of the Au/PU sponge.^{36,41} The opposite process (*reduced* internal contact area) occurs when *less* pressure is applied.

The electrical current values of the Au/PU piezoresistive sponge recorded upon application of different pressures are outlined in Fig. 3B. Importantly, Fig. 3B demonstrates that the output signals generated upon on/off pressure applications are sharp, uniform, and reversible. Furthermore, the pressure threshold obtained (e.g. sensor sensitivity) is lower than 10 Pa, a sensitivity threshold that is on par or better than many reported pressure sensors (which, by comparison, are mostly more difficult to fabricate).^{36,48} To quantitatively assess the pressure sensitivity of the new Au/PU system we plotted R_p/R_0 [in which R_p is the resistance at applied pressure P, and R_0 is the resistance without applied pressure] vs the applied pressure P (Fig. 3C). The two linear fits in Fig. 3C and calculated slopes (S1 and S2 indicated on the plot) reflect the distinct pressure regimes usually observed in chemicallyassembled pressure sensors.^{28,41} Specifically, a steeper slope is apparent upon initial pressure applied - in which even slight changes in pressure significantly affect the contact area between the Au-coated polymer fibers and consequently alter the corresponding resistance. In the high pressure regime (above ~2 kPa, Fig. 3C), modulation of the internal contact area within the sponge is less pronounced upon changing the pressure, giving rise to a lower slope.

The pressure sensitivities obtained in the Au/PU system and highlighted in Fig. 3C are similar or better than previously reported pressure sensors and attest to the excellent response parameters of the Au/PU system.^{36,48} Fig. 3D underscores the stability of the Au/PU sponge as a pressure sensor. The graph in Fig. 3D shows the current values recorded upon application of a 1000 on/off pressure cycles; no significant drift is observed in recorded current. The *response time* of the new piezoresistive sensor was lower than 10 msec (Fig. S4), further supporting practical sensing applications.

Arterial pulse sensing by the Au/PU sponge

Fig. 4 presents an example of a practical use of the Au/PU sponge as a physiological sensing platform (other pressure sensing applications are presented in Fig. S3 in ESI⁺). Specifically, the graphs in Fig. 4 depict the current variations induced by blood pulses recorded upon placing the sponge upon the wrist artery (as shown in the photograph in the inset, Fig. 4A). The high fidelity and sensitive pulse measurements by the piezoresistive Au/PU sensor are apparent in the variation of current with time depicted in Fig. 4A. Furthermore, the pressure sensor clearly distinguishes between pulse profiles recorded in rest and following physical activity (Fig. 4A, right panel). The rapid response of the Au/PU sensor to pressure modulation enables high resolution monitoring of the arterial pulses and illuminate even small variations in blood pressures. (As shown in Fig. S4 in ESI⁺, the response time of this Au/PU sensor is less than 10 ms.) Specifically, Fig. 4B depicts high temporal resolution current plots of several

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pulses recorded in rest. Importantly, Fig. 4B demonstrates that the piezoresistive Au/PU sensor is capable of distinguishing fine features of the arterial line waveform, including the steep systolic uptake and gradual decline, the dicrotic notch, and diastolic runoff

(indicated, respectively, by the Roman numerals in Fig. 4B).⁴⁹



Fig. 4 Blood pulse sensing by the Au/PU sponge. A) Photograph of the pulse sensor placed upon the wrist artery, and signals of heartbeat measurement at rest (72 beats min⁻¹) and after physical activity (114 beats min⁻¹). B) High resolution monitoring of blood pressure in arterial pulses. The Au/PU sensor is capable of distinguishing the systolic uptake (i), systolic decline (ii), dicrotic notch (iii), and diastolic runoff (iv).

Conclusions

In conclusion, we developed a new sensitive pressure sensor based upon a simple, easy to produce construct comprising amine-functionalized polyurethane sponge coated with a thin layer of Au nanoribbons. The Au/PU sponge exhibits excellent pressure sensing parameters, including stability and reproducibility, high sensitivity, and low response threshold. We demonstrate application of the new pressure sensor for monitoring arterial pulses. In particular, the fast response time and sensitivity made possible high resolution analysis of the pulse waveform features. The straightforward fabrication route using readily available and inexpensive chemical components, favourable physical properties and operation parameters of the Au/PU sponge system underscore the potential implementation of the new piezoresistive platform in varied pressure sensing applications.

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Colour graphic: maximum size 8 cm x 4 cm



A new piezoresistive sensor comprising Au nanoribbon-coated elastomeric polymer