Development of a conductive photoresist with a mixture of SU-8 and HCl doped polyaniline

Uma M Annaiyan*, Kouros Kalantar-Zadeh, Qiang Fang, Irena Cosic

Abstract— A novel electrically conductive photoresist has been formulated to fabricate microcomponents. The developed conductive photoresist is based upon SU-8 photopolymer, an insulating negative-tone epoxy, in which protonically doped polyaniline (PANI) nanoparticles have been dispersed to enhance the electrical properties. The characteristics of this new conductive photoresist have been studied via electrical measurements. The process for preparing the conductive films from the combination of EB (Emeraldine base) PANI, SU-8 and NMP (N-methyl-2-pyrrolidinone) will be presented. Different weight percentages (wt %) of SU-8 in the above combination have been prepared and spin coated to form thin-films which have been protonated with HCl. The conductivities of the thin-films were measured using a four point probe. The highest conductivity achieved was approximately 1.6 S/cm for the mixture of 10wt% of SU-8 25 with EB-PANI mixed with NMP. The morphology of the thin-films was studied using a Scanning Electron Microscopy (SEM).

Index Terms— SU-8, photoresist, polyaniline, conductive polymer, protonation, biosensors.

I. INTRODUCTION

Current microfabrication processes are used to develop devices with dimensions in the sub-micrometer to millimeter range. Such processes the development of devices such as micro-sensors and micro-actuators. Micromachining techniques for fabricating three-dimensional structures differ from conventional, sequential machining processes, such as drilling or milling with mechanical tools [1]. Integrated circuit fabrication processes are the most prevalent example of microfabrication [2-4]. Microfabrication also presents the possibility of batch processing, the optional integration of electronics and the reproducibility of sensor characteristics due to the accurate geometric control in the fabrication process [1]. The four basic microfabrication techniques are: deposition, patterning, doping and etching. These techniques are combined to build up a device layer by layer. Generally thin-films are deposited on top of the substrate as active or passive components. Microfabrication lithography techniques rely on radiation-sensitive polymers called resists. The resist is irradiated through a mask containing the pattern to be transferred, resulting in cross-linking, which creates a difference in solubility between the irradiated and unexposed areas of the polymer [5]. Finally, the pattern is transferred from the photoresist layer to the substrate by etching or lift off techniques. After removing the remaining photoresist, the next layer is deposited and structured. Doping of a semiconductor material can be undertaken directly after photolithography or after patterning an implantation mask. Doping techniques are typically undertaken to modify the electrical conductivity of materials which are patterned. Hence it is a key step in the fabrication of semiconductor devices such as diodes and transistors.

Each of the previously mentioned steps, thin-film deposition, photolithography and doping, increases the cost of fabrication. Having a photoresist with a controlled conductivity will dramatically reduce the fabrication process time as the photoresist layer can be directly implemented as the semiconducting material. Utilisation of a conductive photoresist eliminates the etching process and simplifies the microfabrication process. Additionally, such a material can also be of tremendous benefit in the fields of sensors and actuators where chemical, electrical and mechanical properties are the key parameters in the development of micro-devices. There are other advantages that this material can offer. For instance for the fabrication of high aspect ratio devices such as spikes which have many applications for biosensing measurements. The silicon etching process currently used to develop such structures has inherent deficiencies such as: slow etch rate, difficulties in applying external connections, lack of control in the conductivity of structures and incompatibility with layered device fabrication. All these problems could be solved with a conductive photoresist with controllable conductivity.

This work was supported in part by the Royal Melbourne University of Technology, Australia under RIF Grant.

Uma M. Annaiyan, is with the School of Electrical and Computer Engineering Department, Royal Melbourne Institute of Technology, Melbourne, VIC 3001, AUSTRALIA, Royal Melbourne Institute of Technology, Melbourne, VIC 3001, AUSTRALIA (Corresponding author : phone: 61-3-99259533; fax: 61-3-99252007; email: s2118335@student.rmit.edu.au).

* Kouros Kalantar-Zadeh, Lecturer, is with the Royal Melbourne Institute of Technology, Melbourne, VIC 3001, AUSTRALIA (e-mail: kourosh.kalantar@rmit.edu.au).

Qiang Fang, Lecturer is with the School of Electrical and Computer Engineering Department, Royal Melbourne Institute of Technology, Melbourne, VIC 3001, AUSTRALIA (e-mail: john.fang@rmit.edu.au).

Irena Cosic, Professor and Head of School, is with the School of Electrical and Computer Engineering Department, Royal Melbourne Institute of Technology, Melbourne, VIC 3001, AUSTRALIA (e-mail: irena.cosic@rmit.edu.au).
In order to develop a conductive photoresist a novel approach of mixing a conductive polymer with the photoresist is presented in this paper. SU-8 is used as the base photoresist. To improve the functionality of the SU-8, it was doped with a conductive polymer. The goal is to load the SU-8 epoxy with conductive particles in order to enhance its electrical conductivity. In this paper, we use polyaniline (PANI), a conductive polymer. A different approach to develop conductive SU-8 was initiated by Jiguer et al [6] where they combined silver nano-composites with SU-8.

SU-8 is a negative tone, chemically amplified, near UV photoresist which was first developed at IBM Yorktown, USA in the late 1980s [7]. The specific properties of SU-8 enable ultra-thick layers of high uniformity with low edge bead, high transparency at wavelengths above 340 nm, low optical absorption [8] and smooth surface up to 1 mm by a single coating step. Near UV exposure with low doses yields patterns with nearly vertical sidewalls of high aspect ratio >50 [9-11]. Ultra thick resists with high aspect ratios are essential for several MEMS, bioMEMS and MOEMS applications, such as electrostatic sensors and actuators, fluidic channels, nanobiosensors and micro robots [12], microelectronics, micromechanics, microfluidics, packaging, magnetics, microoptics, at panel displays, and microwave devices [5]. SU-8 is also sensitive to electron beam exposure and has several advantages in applications when compared to the most widely used e-beam resist, polymethyl methacrylate (PMMA) [13]. Firstly SU-8 has a much higher sensitivity which greatly reduces the time required to expose a given pattern. Secondly, cross linked (fully exposed) SU-8 is an optically transparent epoxy that is chemically and physically durable. For many applications these properties eliminate the need for a dry-etch pattern transfer onto a more durable substrate [13]. The SU-8 material consists of an epoxy called EPON SU-8, a solvent called gamma-butyrolactone (GBL) and a photoinitiator taken from the family of the triarylmethyl-sulphonium salts [14].

It is possible to create conductive polymers with a diverse range of properties [15]. The material class of conducting polymers is used in actuators, transistors, wires, super capacitors, strain gauges [15], electromagnetic interference (EMI) shielding, electrostatic discharge (ESD) protection, metallization, lithography—(e.g., charge dissipaters, conducting resists), corrosion protection of metals [5], fibers, antistatic and other coating, films, diodes, electroluminescent devices, displays, printed circuit boards, electrochromic windows in houses and cars, conductive fabrics [16], optoelectronic devices [17] and much more. The use of a polymer as a component material for the biosensing applications provides the sensors with very high sensitivity and convenient mechanical properties. Conductive polymers such as PANI are increasingly applied in biosensor construction for the sensing process [5].

Polyaniline (PANI) is one of the most popular and chemically stable polymers [18] under ambient conditions, having high conductivity [19]. PANI has been described in many papers for the past 120 years [20-23]. It is most promising because of its straightforward polymerization [24,25]. The conductivity of PANI depends on various factors such as oxidizing agents, aniline/oxidant mole ratio (k), reaction temperature and time of polymerization [26], the percentage of protonation and the degree of water and solvent content [27]. The term 'PANI' is widely used to indicate any polymeric material obtained by the oxidation of aniline [28]. These materials are generally doped with protonic acids such as aqueous hydrochloric acid (HCl) to provide conductivity in the order of 1S/cm [20].

PANI is composed of aniline repeat units connected to form a backbone of alternating nitrogen atoms and benzene rings; it exists in a variety of forms that differ in chemical and physical properties [29]. It also comes in three classifications depending upon the degree of oxidation of the nitrogen atoms; the leucoemeraldine base (LB; fully oxidized form); the emeraldine base (EB; half oxidized form) and the pernigraniline base (PNB; fully oxidized form) [30,31].

PANI can be doped by protonation. The unusual properties of PANI arise because it is A-B nature, since most other conducting polymers are A-A type [15]. Of the three different states of oxidation, the one that can be doped to the highly conducting state is called "polymeraldine". It consists of amine (-NH-) and imine (=N-) sites in equal proportions (Fig. 1). The imine sites are protonated to generate the bipolaron form (Fig. 2) [27].

![Figure 1: Emeraldine base (nonconductive) form of PANI -type polymer.](image-url)
The EB can be transformed into a conducting state upon doping with protonic acids, such as HCl to yield the emeraldine salt (ES) [20, 27]. It has been shown that protonation of PANI in the emeraldine oxidation state leads to a dramatic change in conductivity, from approximately $10^{10}$ S/cm for the unprotonated polymer to a maximum conductivity of $10^4$ - $10^4$ S/cm when doped with aqueous acids [32, 33]. The EB PANI has electrons spread throughout after doping [34, 31].

The present work reports the development of a conductive photoresist. The process of the development of films from the combination of EB PANI, SU-8 and NMP (N-methyl-2-pyrrolidinone) is presented. These thin-films are studied in terms of their conductivity and morphology.

II. EXPERIMENTS

EB PANI (Sigma Aldrich) with a Mw of 214.27 was vacuum dried [35] for 12 hours in a to remove any moisture. 1-Methyl-2-pyrrolidone (NMP) of 99% purity from Sigma Aldrich was used as the solvent for the vacuum dried EB powder because NMP has non- chlorinated or chlorinated hydrocarbons [36] and is also commercially used as a SU-8 remover. It was reported by Kang et al. [37] that any polymer which is soluble in NMP can be blended with the NMP solution of EB.

The chemical polymerisation is carried out by dissolving various molecular weights of EB in NMP [5,20,34,38,28,31]. Initial tests for conductivity, were carried out with very low concentrations of EB (~ 0.01g) in 5ml of NMP (mixture 1). Three different SU-8 viscosities, SU-8 2, SU-8 25 and SU-8 50 from Microchem were considered for experiments. The resultant mixture was passed through a syringe filter (pore size 0.2 m). Concentrations from 0.05g/ml to 0.08g/ml of SU-8 2 in "mixture 1" were spin coated on glass substrates at 3000rpm for 40 sec.

After curing on a hotplate for 5 minutes at 55°C, the samples were soaked in 1M hydrochloric acid (HCl) (33% purity) aqueous solution for 2 hours at room temperature. Because of the insulating nature of EB PANI, this soaking process protonates the polymer film. Therefore, the obtained chemical composition consists of emeraldine hydrochloride [20]. The resistance of the samples was measured using an ohm-meter (Fig. 3). The measured resistance was within the k to M range. After protonation, the samples changed colour from dark blue to green. It was found that the greener the sample, the higher the conductivity. Thin films were sufficiently transparent for photolithographic applications.

It was concluded from the initial experiments that the resistance is measurable but the thickness of the polymer thin film deposited over the substrate was uneven because of low viscosity of the mixture. It was decided to increase the viscosity by using a higher concentration of SU-8 and EB.

"Mixture 2" was prepared by mixing 0.4g of EB in 5ml of NMP. Greater percentage of SU-8 25 was added to increase the viscosity of samples. 10 samples with the volume in the range of 0.1ml to 1.0 ml of SU-8 were mixed with 1.0 ml of "mixture 2" in an ultrasonic bath for 72 hours. After spin coating and curing the samples, they were soaked in 1M HCl (37% purity) solution for 2 hours. The samples were air dried and measured for resistance using four-point probe technique as shown in Fig. 4 [39,40].

The four probe technique is a very efficient method for measuring the sheet resistance. It measures the sample resistance by measuring the current that flows for a given applied voltage. The outer two pins force a current through the sample and the inner two pins measure the voltage drop. In the case when the Im thickness is much smaller than the
thickness of the substrate, the resistivity can be calculated using [39]:

$$\rho = \frac{\pi d}{\ln(2)} \frac{V}{I} \Omega \text{cm}$$

(1)

Where, t is the thickness of the thin film, V is the measured voltage and I is the forced current.

![Figure 5: Average conductivity versus SU-8 volume percentage](image)

Fig. 5 shows the average conductivity in all 10 samples of "mixture 2". The conductivity range was between 1.6 S/cm and $2.5 \times 10^{-3}$ S/cm for volume percentage of SU-8 in the range of 0.1 to 0.6, respectively (Fig. 5).

![Figure 6: SEM Micro-graph of conductive thin film (0.4Mmol SU-8 and 1Mmol of "mixture 2")](image)

Fig. 6, which shows low concentration of SU-8 (0.4 Mmol SU-8 and 1Mmol of "mixture 2"), has particle dimensions ranging from approximately 50 nm to 200 nm. It is observed that these particles are embedded in a continuous polymeric media. The film conductivity is approximately 0.03 S/cm.

Fig. 7, which shows high concentration of SU-8 (0.9 Mmol SU-8 and 1Mmol of "mixture 2"), has particle dimensions ranging approximately from 100 nm to 400 nm. The materials are isolated clusters of particles with different dimensions.

These clusters are also located in a polymeric media. The thin film appears to be highly resistive, above the operating range of the measurement apparatus. It is suggested the formation of separate particles in this case reduces the continuity of PANI chains which reduces the conductivity.

III. CONCLUSIONS AND FUTURE WORK

A process for preparing chemically polymerized EB PANI with photoresist polymer SU-8 25 was developed. It was seen that for 0.1 Mmol SU-8 in 1 Mmol of the mixture of EB PANI, and NMP, a conductivity as high as 1.6 S/cm can be obtained. It was observed that increasing the weight percentage of EB in the mixture increases the conductivity. Morphology of the conductive and non-conductive thin films have been studied. In non-conductive thin films the media consists of separate clusters whilst the conductive thin films consists of nanoparticles embedded in a polymeric media. Properties of the thin films such as adhesion, spread viscosity
and transparency of the material make the material suitable for applications such as biosensors. The novel photoresist could play an important role in developing structures such as conductive channels, gas sensors, biosensors and many more applications in the field of semiconductor device fabrication. Such a novel mixture can also be used for the development of high aspect ratio structures such as micro-needle biosensors which are generally developed using silicon material with a lengthy fabrication process.

In future work, it is intended to study the photolithographic properties of these thin films. The developed technique will be applied to produce thin layers of conductive photoresist for fabrication of various devices. It is also planned to develop high aspect ratio semiconductor structures.

ACKNOWLEDGMENT

This research is supported by a RIF grant from RMIT University.

REFERENCES


