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Fabrication of highly electrically conducting fine patterns via substrate-independent inkjet printing of mussel-inspired organic nano-material

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We report the inkjet printing of an aqueous suspension of synthesized mussel-inspired poly(dopamine) nanoparticles. Fine lines of printed nanoparticles were deposited on both glass and polyethylene terephthalate (PET) substrates by exploiting the coffee ring effect. Deposited particles were then used for site-selective silver metallization via a simple electroless plating process at controlled temperature (30°C) and plating time (12 min to 120 min). The resulting narrow lines of silver exhibited a resistivity 10× that of bulk silver.

Lines on PET retained good electrical and adhesion performance even after prolonged bending tests. This technique satisfies general requirements of flexible electronics manufacturing – low temperature, low cost, small feature size and good electrical conductivity independent of substrate material.

Electrically conducting patterns are essential structures for emerging flexible electronics. Drop-on-demand inkjet printing is a material-conservative deposition process compatible with the low temperature requirements of flexible polymer substrates.

The implementation of printing within a roll-to-roll (R2R) infrastructure enables continuous, high-speed and large-scale manufacturing. The functional materials suitable for inkjet printing are generally available as nanoparticle (NP) dispersions and precursor solutions. The NP ink is comprised of functional cores surface-coated by a stabilizing agent and uniformly dispersed in a volatile liquid; precursor inks are metal salts dissolved in suitable solvents. To transform printed deposits into a highly electrical conducting phase, a post-printing process is generally required either to remove the adsorbed stabilizing agent and sinter the functional cores (for NP inks) or to precipitate metal structure in-situ (for precursor inks). The standard approach is the application of heat in a controlled ambient environment. The thermal sensitivity of low cost flexible substrates (such as polyethylene terephthalate (PET)) has led to extensive research into alternative non-thermal techniques such as laser,[2] UV[3]/IR[4] light, electrical field[5], chemical[6] and plasma[7,8]. However, these post-printing processes generally require sophisticated and expensive equipment. In this letter, we report a printing process for patterned metal deposition which employs site-selective electroless plating (ELP) of silver (Ag) on printed patterns of organic NPs.

Electroless plating generally uses a solution of metal salt, reducing agent, a complexing agent, and additive(s) (such as bath stabilizer and plating rate adjusting agent)[9]. Metal nucleates on the cataclytically active surface and continues to promote further metal reduction and growth. This is the defining characteristic of ELP’s autocatalytic nature.[10] A pre-patterned catalyst layer on the target substrate will promote site-selective deposition. Hidber et al.[11] utilized microcontact stamping to pattern palladium (Pd) colloids which yielded copper (Cu) patterns after ELP. Harkness et al.[12] used photolithography to pattern a Pd-bonded-organic seed layer (hydrogen silsesquioxane) and achieved site selective ELP of Cu and Ag. The prohibitively large cost of photolithography has prompted research into the use of inkjet printing to directly pattern the catalyst[13-20]. Most of the inkjet printing studies exploited Pd-based ink[14-18], while the investigation of other ink materials has been less focused[19,20]. Furthermore, the application of Pd-based inks is limited in versatility because ink-substrate adhesion is substrate dependent[14-20]. Finally, although Pd-based catalyst is useful for electroless plating of a wide range of metals, its high cost prohibits its use for many applications.[21]

A mussel-inspired organic polymer poly(dopamine) (PDA) was recently found capable of triggering metal ion reduction[22] leading to its implementation as an ELP catalyst[23,24]. PDA also shows universal adhesion on a wide range of organic and inorganic materials[22]. PDA can be synthesized as either continuous coatings by triggering dopamine polymerization in pure water[22] or as spherical nanoparticles in water-alcohol mixtures at controlled pH[22]. Studies show that polydopamine nanoparticles (PDA-NP) retain the same surface chemistry as continuous PDA coatings.[22,25]

Inkjet printing as a method of creating fine features suffers from the limitation of the individual drop size, which can range from 20-80μm[26-29]. We have recently reported a technique that exploits the so-called “coffee-ring” phenomenon studied in sessile drops[30]. In a printed line of dilute particle ink, evaporative deposition of particles at the contact line can produce a pair of parallel lines with minimal solid deposit between the lines[31]. The resulting line widths can be controlled to less than 10% of the...
In this study, we report a novel patterned metal deposition technique that combines twin line inkjet printing of aqueous PDA nanoparticle ink with site-selective ELP of Ag on both glass and PET substrates. To the best of our knowledge, the inkjet printing of PDA-based nanoparticle ink for patterned metal deposition has not been reported. We report the effect of plating time on morphology, crystalline structure, and electrical resistivity of lines of plated silver. The reliability of Ag lines on flexible PET substrates subjected to cyclic bending and adhesive tape tests is also reported.

Polydopamine nanoparticle (PDA-NP) powder was synthesized according to a published protocol. Briefly, PDA-NPs were generated by initiating dopamine self-polymerization in a water-isopropanol mixture which was buffered by Tris (See ESI†). The average size of PDA-NP was measured with a dynamic light scattering system (Zetasizer-Nano, Malvern) to be $338 \pm 55$ nm (See ESI†). The as-synthesized PDA-NP powder was dispersed in distilled deionized (DI) water at 1% wt by 10 min agitation by ultra-sonication. The viscosity and surface tension of the PDA-NP ink were measured as $1.0 \pm 0.1$ mPas and $72.3 \pm 0.1$ mN/m using a cone-plate rheometer (AR1000, Texas Instrument) and bubble pressure tensiometer (BP100, Kruss), respectively.

A single-nozzle inkjet printer was used to print the PDA-NP suspension onto plasma-treated glass substrates (2947, Corning) and PET substrates (Melinex ST506, Dupont) to form arrays of five lines (Scheme 1 and ESI†). Evaporative-driven transport in the presence of a pinned contact line leads to potential deposition of particles at the contact lines. The result is a twin line particle pattern with minimal interline particle deposition (Fig. 1a and ESI†). Each of the twin lines measured $24 \pm 1$ µm and $10 \pm 1$ µm in width for the glass and PET substrates, respectively.

Ag deposition onto the printed pattern can be achieved by simple immersion into a silver salt solution such as silver nitrate ($\text{AgNO}_3$). However, an ELP process that includes an additional reducing agent is necessary to ensure the rapid formation of a continuous and sufficiently thick metal layer. A potential mechanism of PDA-induced ELP has been suggested to involve the electrostatic interaction of metallic ions with surface catechol groups of PDA followed by catechol oxidation to quinone and reduction of metallic ions to elemental metal. The newly deposited elemental metal further catalyzes the redox reaction for continuous deposition. In this study, Ag ELP was performed by immersing the printed PDA patterns in a chemical bath at 30°C for different durations (See ESI†).

The morphology of the printed twin line region before and after the plating process was characterized by optical microscopy (Axio Imager, Zeiss) and scanning electron microscopy (SEM) (Supra 55VP, Zeiss). Top and cross-sectional views show that for the as-deposited lines, each twin line typically consists of a 1-2 particles thick PDA-NP structure (Fig. 1a, SEM). At early time Ag ELP, small Ag grains nucleate on top of each twin line. The grains grow in size and merge to form a slightly granular but continuous Ag layer with extended ELP time. The Ag continuity and grain size increase with extended ELP time. No obvious structural differences in particle deposits on glass and PET substrates were observed. The SEM images of the interline region can be found in the ESI†.

Cross-sectional SEM images were also analyzed to measure the Ag thickness on each twin line as a function of metal plating time (Fig. 2(a)). In general, the Ag growth rate is linear with ELP time and exhibits no apparent differences between glass and PET substrates.
substrates.

The electrical resistance of Ag-plated lines was measured using a line scratching technique\(^\text{19,30}\). Briefly, a pair of Ag electrodes was sputtered through a mask over each line array so that the lines spanned the distance (4mm) between electrodes (ESI†). Four-point-probe measurements were carried out to determine the total resistance of the five lines. Each line was carefully cut by a needle attached to a micro-positioner and the new resistance value recorded. Under the assumption that each line array is a parallel resistor network, this method allowed measuring the average resistance of each cut line. As Fig. 2(b) shows, single line resistance decreases rapidly within 24min (62Ω and 212Ω for glass and PET, respectively) of ELP, followed by a slower decay on a longer time scale to reach the lowest value of 12Ω for glass and 33Ω for PET at 120min. It was possible to determine whether the Ag in interline region contributed to the measured resistance by disconnecting all the twin lines. It was found that for plating times longer than 58min (glass) and 96min (PET), an electrically conductive path forms in the interline region. However (as Fig. 2(b) shows), this structural transition did not correlate with a measured transition in line resistance, which implies that the resistance of the interline region is much larger than that of each twin line.

![Fig. 2 Deposited structural characterization: (a) thickness (t) evolution at twin region; (b) individual-line resistance as a function of silver electroless plating time; (c) electrical resistivity change at twin regions from 12min to 56min silver electroless plating time.](image)

The electrical resistivity of Ag plated on each twin line was estimated for ELP duration of 56min or shorter, i.e., before the transition to inter-line conductivity on each substrate (see ESI†). A twin-line structure consisting of two parallel conductors with no electrical contribution from the inter-line region was assumed. The resistivity ($\rho$) can be expressed as $\rho = 2RtW/L$, where $R$ the measured resistance of a twin line, $t$ the Ag thickness, $W$ the width of each twin line and $L$ the length between two electrodes. Fig 2(c) plots $\rho$ as a function of ELP time for both glass and PET substrates. At the same Ag ELP time, resistivity is similar for both substrates. Within the first 24min of plating, $\rho$ reaches a value which is about 10.4× and 15.2× larger than the resistivity of bulk Ag, for glass and PET, respectively. After longer plating time, $\rho$ decreases slowly. The lowest values for $\rho$ are 1.57E-7 Ωm (on glass) and 1.74E-7 Ωm (on PET) after 56min Ag ELP which are 9.8× and 10.9× the bulk Ag resistivity, respectively.

In conclusions, fine lines of Ag with low electrical resistivity were fabricated by sequential inkjet printing of mussel-inspired poly(dopamine) nanoparticle ink and Ag electroless plating. The process yields Ag lines with thickness linearly dependent on plating time and electrical resistivity 10× and 11× that of bulk silver, on glass and PET substrates, respectively. This process demonstrates a simple, low temperature and low-cost method of fabricating fine conductive patterns for flexible electronics applications. The combined approach of inkjet printing organic
nano-materials and ELP was reported for the first time.

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**Notes and references**

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