NANOSTRUCTURED FILMS OF PLASMA POLYMERIZED HEXAMETHYLDISILAZANE AND LACTIC ACID DEPOSITED BY ATMOSPHERIC PLASMA JET

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Abstract

In this work, composite layers of hexamethyldisilazane (HMDSN) and lactic acid (LA) were prepared using chemical vapor deposition process by a nitrogen atmospheric pressure plasma jet source. The HMDSN vapours were introduced into the plasma jet. Varying LA admixture was added into the jet in the form of aerosol. The chemical composition, morphology and wettability of resulting films were studied. Infrared spectra and XPS analysis show both Si-O like and hydrocarbon structures. With increasing amount of LA admixture, the oxidation of the layers increases. The morphology of the films was measured with SEM. The films prepared with no LA have a foam-like structure. With increasing amount of LA the structure changes into more fractal-like. The X-ray diffraction analysis suggests the presence of amorphous SiO₂ domains in the films. The films exhibited superhydrophobic properties that can be modified by the post-treatment of the samples in DBD discharge.

Keywords: Lactic acid, hexamethyldisilazane, atmospheric plasma jet

1. INTRODUCTION

(Bio)degradable polymers, like polyethylene oxide and polylactic acid, are used in biomedical field [1], food industry, packaging or medicine [2]. For such applications, preparation of these materials in the form of thin films with controlled properties via plasma-based methods was found viable [3][4], including deposition in atmospheric pressure plasma [5].

Atmospheric pressure plasma jets have been used to produce thin films based on siloxanes [6][7] or silazanes [8][9]. While the monomer is more typically introduced into the discharge in the form of vapours, deposition from aerosols has also been demonstrated [10][11]. In this work, a combined deposition from HMDSN vapours and lactic acid aerosol in atmospheric pressure plasma jet has been studied.

2. EXPERIMENTAL

2.1. Preparation of the films

The films were prepared using atmospheric pressure DC plasma jet fed with nitrogen (Figure 1) [12][13]. The total N₂ flow used was 7 l/minstp. Of this, 0.1 l/minstp (100 sccm) was fed into the bubbler with HMDSN, effectively then carrying about 3 sccm (0.025 ml/min) of HMDSN. Lactic acid was supplied by a syringe pump into an ultrasonic sprayer (nebulizer). The resulting aerosol was carried with the rest of the N₂ flow. The resulting mixture of N₂, HMDSN vapours and LA aerosol was fed into the plasma jet. The plasma jet was
operated at 10.2 kV and 11.3 mA. The deposition time was 5 minutes. The films were deposited on single-side polished silicon and gold-coated silicon.

![Figure 1 Scheme of deposition setup](image)

2.2. Characterization of composition and morphology of the films

The chemical structure of the films was characterized using infrared spectroscopy (FTIR-RAS, Bruker Equinox 55) and X-ray photoelectron spectroscopy (XPS, Specs Phoibos 100). Morphology of the films was characterized by scanning electron microscopy (SEM, Tescan Mira III) and X-ray diffraction (XRD, Rigaku SmartLab). Water contact angle measurement and the dielectric barrier discharge (DBD, Ceplant) setup for post-treatment [14] were custom built.

3. RESULTS AND DISCUSSION

3.1. Infrared spectra

The infrared spectra of the films (Figure 2) generally resemble more the HMDSN monomer than LA, despite the relatively high LA input into the nebulizer. That can be probably explained by a worse transmission of the LA aerosol through the gas conduit in comparison with the HMDSN vapours. The most prominent features of the films can be attributed to bonds for silazanes (Si-N-Si, N-H), siloxanes (Si-O-Si) and alkyl-substituted silanes (Si-H, Si-(CH$_3$)$_n$, CH$_x$) with the presence of OH groups and a very small amount of C=O bonds.

With increasing LA admixture most visible changes can be seen in the peak around 1100 cm$^{-1}$. The shape of the peak shifts in favour of Si-O-Si bonds compared to nitrogen and carbon containing bonds, but the overall intensity of the peak decreases in comparison to the signal of Si-bound CH$_3$ group.

3.2. X-ray photoelectron spectroscopy

The elemental composition of the films (Figure 3) was established using XPS. The amount of silicon in the films remains roughly constant. However, with increasing LA admixture, the amount of oxygen in the films increases and the amount of carbon and nitrogen decreases.

The shape of the high-resolution peaks was found to be too symmetrical to allow reliable assignment of all the various bonds possibly present in the films. The elemental composition of films on gold-coated substrates was measured as a double-check to avoid the influence of the silicon substrate on the apparent film composition. No significant difference to the composition of the films on pure silicon was found.
3.3. Chemical composition - summary

The amount of silicon in the films seems to be almost independent of the concentration of LA added into the working gas. The concentration of nitrogen is very low (in HMDSN monomer the Si/N ratio is 2, in the films it is 8 or more) and for layers prepared with some LA, the nitrogen nearly disappears. We can also see that there is a lot of oxygen in all layers, including those without LA admixture. The plasma jet column itself carries in the ideal case only nitrogen with precursors, so we suppose that oxidation from the ambient air around the jet takes place in the process. An increase of the oxidation with increasing LA admixture shows a fragmentation of LA molecule, bringing more oxygen into the process. The plasma polymerization of lactic acid takes place in a very small amount if at all - if it were the case, the C/O ratio would not drop with LA admixture.
3.4. Morphology of the films

The SEM images of the films are shown in (Figure 4). The films show a foam-like or fractal-like structure for lower or higher LA admixture, respectively. The size of the structures seems also to slightly increase with increasing LA admixture.

The film thickness increases with increasing LA amount input (Figure 5). However, the films then also appear to be visually less dense. The effective deposition rate of the films is 1-2 μm/min.

Figure 4 SEM images of the film for varying LA admixture a) top view b) close-up top view c) side view

Figure 5 Thickness of the films in dependence on the LA admixture
3.5. X-ray diffraction

The X-ray diffraction data were obtained for the samples prepared at 0.1 ml/min and 0.2 ml/min of LA admixture. A weak peak around 2Φ=23° was detected. The corresponding lattice constant can match SiO$_2$ domains that can be present in the structure of the films. The width of the peak (~10° FWHM) suggests that these domains are amorphous.

3.6. Wettability properties of the films and DBD treatment of the films

The films were found to be superhydrophobic and slippery (water contact angle ~180°, no hysteresis in advancing/receding contact angle). Films prepared with 0 and 0.2 ml/min LA admixture were treated with dielectric barrier discharge in air. After treatment with the discharge with a power density of 3 W/cm$^2$ for 10 s these films were found to be superhydrophilic. This switching between the extremes of wetting behaviour of nanostructured films due to chemical changes on their surface have previously been demonstrated on other nanostructured films [15][16].

4. CONCLUSIONS

Thin films from hexamethyldisilazane and lactic acid have been prepared using atmospheric pressure plasma jet. The LA is found to be strongly fragmented in the discharge. The resulting nanostructured films have a general composition of oxidized organosilicones with domains of SiO$_2$. The films exhibit superhydrophobic properties.

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372
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