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Deposition of hard thin films from HMDSO in atmospheric pressure dielectric barrier discharge

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Abstract. The atmospheric pressure dielectric barrier discharge burning in nitrogen with small admixture of hexamethyldisiloxane (HMDSO) was used for the deposition of thin organosilicon films. The thin films were deposited on glass, silicon and polycarbonate substrates, the substrate temperature during the deposition process was elevated up to values within the range 25°C – 150°C in order to obtain hard SiOₓ-like thin films. The properties of the discharge were studied by means of optical emission spectroscopy and electrical measurements. The deposited films were characterised by Rutherford backscattering and elastic recoil detection methods, x-ray photoelectron spectroscopy, infrared spectroscopy measurements, ellipsometry and depth sensing indentation technique. It was found that the films properties depend significantly on substrate temperature at deposition. An increase of substrate temperature from 25°C to 150°C leads to an increase of film hardness from 0.4 GPa to 7 GPa and the film chemical composition changes from CHₓSiₓO₂ to SiOₓHᵧ. The films were transparent in visible range.


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1. Introduction

Dielectric barrier discharges (DBDs) are widely used for industrial purposes such as the modification of polymer surface properties, e.g. wettability or adhesion, owing to the possibility of the atmospheric pressure processing and the on-line treatment. Although this method is very useful, the main disadvantage of DBDs is lack of uniformity. This is due to the fact that at atmospheric pressure DBD is normally a filamentary discharge. Due to of this drawback an effort to homogenise DBDs has been made and it was found that under certain conditions homogeneous DBD can be obtained. A homogeneous DBD can be obtained in helium or neon and it is called atmospheric pressure glow discharge (APG discharge, APGD) [1, 2]. Another type of homogeneous DBD can be obtained in nitrogen and this discharge type is called atmospheric pressure Townsend-like discharge (APTD) [3]. The uniformity of plasma favours these discharge types for thin film deposition techniques such as plasma enhanced chemical vapour deposition (PECVD). PECVD methods enable the preparation of coatings with a wide range of properties, e.g. wetting and sticking behaviour, mechanical, optical or electrical properties. The disadvantage of the low pressure PECVD techniques is the necessity to use expensive vacuum pumping systems. Moreover, the difficulties to arrange the deposition system for large area depositions have to be overcome. Recently plasma deposition at atmospheric pressure has become a promising technology due to its economical and ecological advantages. Sawada et al [4] reported the organosilicon thin films deposition in APGD in helium with the admixture of tetraethoxysilane (TEOS) or HMDSO and oxygen, Prat et al [5] reported fluoro-polymer film deposition in helium APGD with admixture of hexafluoropropylene or tetrafluoroethylene, Gherardi et al [6] used N₂-SiH₄-N₂O APTD for SiO₂ deposition, Foest et al [7] used APGD in helium with an admixture of HMDSO for organosilicon thin film deposition. Starostine et al [8] reported the silica-like coating deposition in APGD in argon, nitrogen, oxygen and HMDSO gas mixture. Recent reviews of theoretical understanding of homogeneous DBD can be found in [9, 10].

In the present paper APTD in nitrogen was used with a small admixture of hexamethyldisiloxane (C₆H₁₈Si₂O - HMDSO) for the deposition of thin films with desired mechanical and optical properties and surface energy. HMDSO is already widely used for SiOₓ thin film deposition in different discharge types [11, 12, 13, 14]. In a previous study it was possible to deposit thin films from HMDSO in nitrogen APTD [15]. However, the deposited films were polymer-like, their maximum hardness was 0.6 GPa and contained about 40% of carbon. Because such films cannot be used as hard protective layers, procedures for obtaining harder films were needed. The first possibility is an annealing of deposited films. The annealing of deposited films increases their hardness up to 5 GPa and the content of Si–O–Si groups increases also [16]. The second possibility is to add an oxidizer gas (O₂ or N₂O) to HMDSO. Massines et al [17] reported the deposition of SiOₓ thin film in N₂ - HMDSO - N₂O APTD. In this case, N₂O acts as oxidizer gas and it was possible to obtain SiO₂-like material. However, this
research found that the addition of N$_2$O achieves an insufficient increase of film hardness. In the present paper we studied another possibility to increase the film hardness, namely the deposition at higher substrate temperatures.

2. Experimental

The experiments were carried out in a metallic discharge reactor with the dimensions 500 mm × 500 mm × 500 mm. The discharge burned between two planar metal electrodes, the upper covered with Simax glass, 1.5 mm in thickness. The bottom electrode was rectangular with dimensions 150 mm × 60 mm, the upper electrode was circular with a diameter of 36 mm. The bottom electrode could be heated using a heating spiral and the electrode temperature was measured with a thermocouple. The films were deposited on thin glass substrates with dimensions 160 mm × 65 mm × 1 mm, polycarbonate 160 mm × 65 mm × 0.8 mm substrates and 500 µm silicon wafers, 76 or 102 mm in diameter. The glass and polycarbonate substrates were placed directly on the bottom electrode. In the case of silicon (Si) substrate, a 1 mm thick glass plate was placed between the bottom electrode and Si. The discharge gap between the substrate and the upper electrode was set to 0.5 mm.

The films were deposited in homogeneous mode of DBD (APTD) from HMDSO vapours mixed with synthetic air and pure nitrogen. The 6 or 16 sccm of synthetic air bubbled through liquid HMDSO in a glass bottle container. It was then mixed with the main nitrogen flow of 6 slm. The HMDSO flow rate was determined by weighting the liquid before and after the deposition. The concentration of HMDSO in nitrogen was 70 ppm and 173 ppm whereas the concentration of oxygen was 200 ppm and 532 ppm for the air flow rates of 6 sccm and 16 sccm, respectively. The film homogeneity depended on the type of gas flow [18]. Configuration of gas supply was optimized in preliminary deposition experiments. The final set-up, used for the depositions discussed in this paper, was as follows. The working gas mixture was supplied through an inlet in the upper corner of the discharge reactor. The reactor was pumped out from the opposite bottom corner. The deposition uniformity was further improved by gas exhaustion of 50 sccm through an opening in the centre of the upper electrode.

Before starting the experiments the discharge chamber was pumped down to a pressure of 20 Pa and then filled with nitrogen to a pressure of 101 kPa. Atmospheric pressure during the deposition was maintained by slight pumping. High voltage with a frequency of 6 kHz was used for discharge generation. The type of discharge, as concerns filamentary or homogeneous mode, was determined from the current-voltage measurements recorded by the digital oscilloscope HP 54820A Infinium (500 MHz, 2 GS/s). The input power to the 6 kHz high voltage generator was 26 W for the homogeneous mode deposition on glass substrates. Maintenance of the homogeneous discharge mode restricted the input power as well as applied voltage in a small range, especially for Si and polycarbonate substrates. In this case, the input power ensuring the homogeneous mode was only 15 W and the flow of air bubbled through liquid HMDSO
was only 6 sccm. The power consumed in the discharge was several watts depending on input power and used substrate [19]. Besides electrical measurements, the discharge was studied by means of the optical emission spectroscopy. The emission spectra were recorded with the Jobin-Yvon TRIAX 550 spectrometer equipped with a CCD detector.

The chemical structure of the films was studied by Fourier Transform Infrared Spectroscopy (FTIR) using a Bruker Vertex 80v spectrometer. The films deposited on Si were measured in transmittance mode taking bare Si substrate as a reference. The films on glass substrates were measured in attenuation total reflection (ATR) mode using diamond crystal (one ATR reflection, angle of incidence 45°). The sample compartment was evacuated down to 2.51 hPa in both measurement modes. The resolution of the spectrometer was set to 4 cm$^{-1}$.

The surface composition of the deposited films was studied by X-ray induced photoelectron spectroscopy (XPS) by means of an ADES-400 photoelectron spectrometer (VG Scientific, UK) using Mg Kα excitation (1253.6 eV) at an incidence angle of 70° and emission angle 0° measured from the surface normal. Since the sampling depth of XPS is about 5 nm, a surface contamination layer was removed by a short argon ion cleaning (5 keV, 5 min). Surface composition was determined from C 1s, O 1s, Si 2p, and N 1s spectra for both, as-deposited and sputter-cleaned films.

The complete atomic composition was determined by ion beam methods combining Rutherford backscattering (RBS) and elastic recoil detection analysis (ERDA). The atomic fractions of C, O and Si were measured by RBS using 2.4 MeV protons perpendicularly bombarding the surface. The sensitivity to carbon content was enhanced using proton resonance effect at the energy of 1.74 MeV. ERDA and RBS with an incident beam of 2.75 MeV α-particles at 75° to the surface normal were simultaneously used to determine the percentage of H. The measured spectra were evaluated by GISA [20] and SIMNRA [21] procedures.

The optical properties of the films on Si were investigated by ellipsometry combined with reflectometry. The associated ellipsometric parameters $I_s$, $I_{II}$ and $I_{III}$, representing three coordinates on Poincaré sphere defined as $I_s = \sin 2\Psi \sin \Delta$, $I_{II} = \sin 2\Psi \cos \Delta$ and $I_{III} = \cos 2\Psi$, were measured with a UVISEL Jobin Yvon ellipsometer at five angles of incidence (55°–75°) in the spectral range 190–2000 nm. The reflectance $R$ was measured with a Perkin Elmer Lambda 45 spectrometer within the spectral range 190–1000 nm at an angle of incidence of 6°. The fitting procedure utilized the least-squares method, i.e. minimizing squares of differences between measured optical quantities $R$, $I_s$, $I_{II}$ and $I_{III}$ and their theoretical values calculated by 2 × 2 matrix formalism [22]. The models used for the calculation of theoretical values are briefly described in the next section.

The hardness and elastic modulus values were assessed from depth sensing indentation tests using a Fischerscope H100 tester with Vickers indenter. The load and the corresponding indentation depth were recorded as a function of time for both loading and unloading processes. The desired material parameters were obtained from analysis of loading and unloading curves. The indentation tests were carried out for
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several different indentation depths (i.e. several different applied loads) in order to map the mechanical properties of the film/substrate system from near surface up to the film–substrate interface. The applied load $L$ ranged from 1 to 100 mN with the force resolution of 0.04 mN. The accuracy of the indentation depth measurement was $\pm 1$ nm. Each indentation test was repeated at least 16 times.

3. Results and discussion

3.1. Discharge diagnostics

The discharge diagnostics was done by electrical measurement and optical emission spectroscopy. The spectra were recorded in the range of 300–800 nm, but above 450 nm only the second spectral order was registered. The influence of electrode temperature on the discharge conditions, manifesting e.g. in the deposition rate and film hardness, was investigated with optical emission spectroscopy. The radiation coming from the discharge was recorded with a spectrometer during controlled continual decrease of electrode temperature. Other parameters were kept similar as during depositions (nitrogen flow 6 slm, synthetic air flow 6 sccm, applied voltage 17 kV$_{pp}$, frequency of applied voltage 6.7 kHz). The spectra consisted of the nitrogen second positive system ($N_2, C^3\Pi_u \rightarrow B^3\Pi_g$), the first negative system ($N_2^+, B^3\Sigma_u^+ \rightarrow X^3\Sigma_g^+$) and of CN violet system ($^3\Pi \rightarrow ^3\Sigma$). The integrated intensities of 0-2 band of the nitrogen second positive system at 380 nm, 0-0 band of the nitrogen first negative system at 391 nm and 1-1 band of CN are plotted against electrode temperature in figure 1. The CN 1-1 band was chosen in order to avoid the inclusion of overlapping nitrogen bands.

The intensities of the bands strongly depended on electrode temperature, especially in the low temperature region. While the intensity of $N_2$ band increases linearly with the electrode temperature, CN band intensity increases at first quickly and then it becomes saturated at high electrode temperatures. Thus maximum at approx. 60°C is observed in their intensity ratio CN/$N_2$.

The bands of the second positive system of nitrogen $N_2$ 0-2, $N_2$ 1-3 and $N_2$ 2-4 were used to calculate vibrational temperature. Determined vibrational temperature was at about 1720 ± 20 K and it this not depend substantially on electrode temperature.

3.2. Chemical structure and composition of films

The chemical structure and composition of the deposited films were assessed by comparison with FTIR spectroscopy, XPS or RBS combined with ERDA. The films deposited on Si substrates using 6 sccm of air bubbling through liquid HMDSO were studied by FTIR transmittance measurements and RBS/ERDA methods. Since the glass substrates are not transparent in IR range, the films on glass deposited with the air flow rate of 16 sccm were investigated by FTIR-ATR spectroscopy. Additionally, these films were studied by XPS. The chemical structure of the films and its evolution
with increased deposition temperature was similar for both substrates and air flow rates. The details are discussed below.

3.2.1. Films on silicon substrates - air flow rate of 6 sccm

The IR absorbances for the films on Si deposited at 25, 50, 75 and 150°C are compared in figure 2. The spectra show three characteristic peaks of silica at 440–450, 802–810 and 1043–1072 cm\(^{-1}\) corresponding to the rocking, bending and antisymmetric stretching vibrations of the Si-O-Si groups, respectively [23]. The position of the stretching peak shifts significantly to higher wavenumbers with increased deposition temperature. This can be caused by several effects such as geometrical reasons, i.e., different film thickness, oxygen deficiency, material densification or carbon admixture. The first reason can be disregarded because the film thicknesses were similar (about 450 nm). Oxygen deficiency causes the shift of this peak to lower wavenumbers [24, 25]. Studies of plastically densified SiO\(_2\) showed the shift of Si-O-Si stretching peak to higher wavenumbers with increased film density [26]. Some authors relate the absorption in the range 1000–1100 cm\(^{-1}\) to Si-O-C groups [27, 28, 29] and Aumaille et al. observed downshift of this peak with increased organic content in the HMDSO/O\(_2\) deposited films [30]. The results of RBS/ERDA analyses showed that the carbon content decreased with temperature from 23 to 2.5 % (figure 3) and the density increased from 1.0 to 2.0 g cm\(^{-3}\) (figure 4). Additionally, there is an oxygen deficiency at 25°C because the O/Si ratio is only 1.4.

![Graph](image.png)

**Figure 1.** The dependence of different band intensities (derived from OES) and their ratios on electrode temperature.
All these results are, therefore, consistent with observed shift of the peak from 1043 to 1072 cm\(^{-1}\). It should be noted that its position for the 400 nm thick thermally grown film was observed by Martinet et al [23] at 1089 cm\(^{-1}\).

Hydrocarbon groups can be responsible for part of the absorption in the region around 800 cm\(^{-1}\) in the case of low deposition temperature at which the carbon and hydrogen percentages were quite high (figure 3). A single peak at 810 cm\(^{-1}\) detected for 150°C film was obviously composed of minimally three peaks at about 782, 802 and 840 cm\(^{-1}\) for the temperatures 25–75°C. Their positions corresponded to CH\(_3\) rocking or Si-C stretching in Si-(CH\(_3\))\(_x\) groups (\(x = 1 \ldots 3\)) [27, 31, 32]. Further evidence of methylsilyl groups was revealed by a narrow peak at 1273–1281 cm\(^{-1}\) and weak peak at 1410 cm\(^{-1}\) associated with symmetric and asymmetric bending of Si-(CH\(_3\))\(_x\), respectively [31, 33]. The former peak was quite strong for all the low temperature films but almost diminished at 150°C. The latter weak peak and weak peak at 2970 cm\(^{-1}\) associated with asymmetric stretching of CH\(_3\) [31, 34] were observable only for 25–75°C. A weak broad peak at about 1360 cm\(^{-1}\) detected for 25°C was assigned to bending of CH\(_2\) in Si-CH\(_2\)-Si [31, 33].

![Figure 2. Infrared absorbances of thin films deposited on Si substrates (air flow rate of 6 sccm through liquid HMDSO) for different substrate temperatures (given in figure legend).](image-url)
The O/Si ratio determined from RBS measurements as 1.4 for 25°C increased to 2.0 for 50°C and even 2.2 and 2.3 for 75 and 150°C, respectively. Obviously, not all oxygen atoms were bonded in the Si-O-Si network. The presence of OH groups was evidenced by a broad peak ranging from 3000 to 3750 cm$^{-1}$. It composed of two peaks centered approximately at 3350 and 3600 cm$^{-1}$ associated with the stretching of hydrogen-bonded and free O-H groups, respectively [35]. Further evidence of OH groups comes from the peak at 930 cm$^{-1}$ assigned to Si-OH stretching [32]. The shift of this peak to 920 cm$^{-1}$ for 25°C could be caused by the presence of the peak at 890 cm$^{-1}$ corresponding to CH$_3$ rocking or Si-C stretching in Si-(CH$_3$)$_2$ [31, 33]. Additional peaks in the region 1490–1780 cm$^{-1}$ observed mainly for low temperatures were also associated with non-silica oxygen. The peaks were relatively well resolved for 25°C at 1585 cm$^{-1}$, 1680 cm$^{-1}$ and 1728 cm$^{-1}$. These positions were assigned to carboxylate ion, trapped water and carbonyl group, respectively [36, 37, 38]. Presence of CO and OH groups can be explained by incorporation of by-products formed during oxidation of HMDSO. Existence of these by-products was proven for low pressure deposition. An increase of optical emission of OH and CO was observed when oxygen was added to HMDSO [30] and mass spectrometer revealed the presence of CO and CO$_2$ in exhaust gas [39].

As already mentioned, the mass and molar densities determined from RBS using
known film thicknesses exhibited an increase when the deposition temperature was raised from 25 to 50°C and a further temperature increase led to a continuous increase of the mass density (figure 4). The latter effect was rather related to the reduction of hydrogen than to the closer packing of atoms as seen from the constant value of the molar density and rapid decrease of hydrogen content illustrated in figure 3.

### 3.2.2. Films on glass substrates - air flow rate of 16 sccm

The apparent differences between the films on glass and Si substrates were caused by different instrumentation used for the investigation of their chemical composition. The striking differences between RBS/ERDA (figure 3) and XPS results (figure 5), i.e. differing amount of Si, O and C as well as progressive changes of the composition with the deposition temperature, were caused by the fact that the amount of hydrogen could not be determined by XPS. If the RBS/ERDA results were recalculated without taking into account the hydrogen, the atomic composition of the films deposited on Si at temperatures 50, 75 and 150°C would be very similar to the XPS results obtained after the film sputtering. The sputtering was necessary because it removed carbon contamination of the film surface. This fact is supported by RBS analyses of the films on Si in which the carbon content also rapidly decreased to 2.5 at.% for the temperature of 150°C. In agreement with RBS the incorporation of nitrogen in the films was below 2 at.%. 

![Figure 4. Mass and molar densities of thin films deposited on Si substrates (air flow rate of 6 sccm through liquid HMDSO) obtained from RBS measurements and known film thickness determined by optical methods in UV/VIS/NIR.](image-url)
The position of Si 2p XPS atomic signal, corrected for a sample charging, corresponded to the binding energy (BE) of silicon oxides, 103.0–103.5 eV [40, 41, 42]. The corrections were performed by shifting the whole spectra to 285.0 eV of C 1s signal. More detailed analyses of chemical bonding were obtained by peak fitting of high resolution measurements of the C 1s signals for the as-deposited samples. The peaks were clearly asymmetric towards higher BE because of oxygen bonded to carbon. A small peak related to these bonds was fitted at about 286.3 eV.

The FTIR-ATR spectra of the films deposited on glass at the same temperatures are shown in figure 6. Since ATR measurements were restricted to the range above 530 cm\(^{-1}\), only two of three characteristic absorption bands of silicon oxides could be observed. Moreover, the ATR spectra are strongly influenced by the changes of refractive index around the position of strong absorption peaks and, therefore, these peaks are shifted to lower wavenumbers and any close peaks at lower wavenumbers increase in intensity. These effects caused distortion of the FTIR-ATR spectra, when compared to FTIR transmittances, in the region of strong absorption bands of silicon oxides. The peak of Si-O-Si rocking vibrations was observed at 790 cm\(^{-1}\), i.e. shifted to lower wavenumbers.
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by 10–20 cm\(^{-1}\) with respect to the position observed in transmittances of the films on Si. Similarly, the antisymmetric stretching peak moved to lower wavenumbers by approximately 20 cm\(^{-1}\). Its shift from 1020 to 1050 cm\(^{-1}\) was again related to the increased deposition temperature and the possible reasons have already been discussed in details for the films on Si.

The stretching peak of the glass substrate was observed at very low wavenumber, 910 cm\(^{-1}\), and could be seen also in the spectra of the films as a weak shoulder at about 900 cm\(^{-1}\). The relatively strong peak observed at 920–940 cm\(^{-1}\) was associated to Si-OH groups. Its high intensity, compared to transmittances of the films on Si, was caused by the adjacent strong Si-O-Si stretching peak. Other absorption features and their behaviour were similar as in the case of the films on Si. The absorption related to OH groups was also detected in the region 3000–3700 cm\(^{-1}\). At temperatures above 25°C, some OH groups were not hydrogen-bonded as evidenced by the appearance of second peak at higher frequencies, about 3600 cm\(^{-1}\). Absorption in the range 3300–3500 cm\(^{-1}\) could be influenced also by a presence of N-H stretching [35], but since the concentration of nitrogen determined by XPS was below 2 %, its contribution was disregarded.

The presence of CH\(_3\) groups, reminiscent of original HMDSO reactant, was
significant especially at lower temperatures. As discussed for the films on Si, their fingerprints were the peaks at 1265–1277 cm\(^{-1}\) and 1412 cm\(^{-1}\). The C-H stretching absorption was also detected in the region 2830–3000 cm\(^{-1}\) as three sharp peaks. Presence of additional groups, that do not have their origin in HMDSO, was revealed by a multipeak absorption in the range 1520–1750°C. It corresponded to several chemical groups such as carboxylate ion, trapped water and carbonyl groups (see above).

3.3. Film optical properties in UV/VIS/NIR range and deposition rate

The UV/VIS/NIR optical properties and thickness of the films deposited on Si using 6 sccm of air bubbling through liquid HMDSO were obtained from combined ellipsometry and reflectometry. Measured data were fitted using a similar dispersion model of the optical constants as in [13]. The model took into account an absorption in the UV range resulting from interband transitions, existence of localized states inside the band gap and a phonon absorption in IR. The interband transitions were described using three parameters (i) band gap \(E_g\), (ii) maximum energy limit of interband transitions \(E_h\) and (iii) parameter proportional to the total density of states \(A\). The transitions corresponding to the localized states as well as the phonon absorption were both parameterized by Gaussian broadened peaks. For details see [43]. Sample imperfections such as non-uniformity and roughness were also taken into account.

The average film thickness obtained from the fitting was used for the calculation of deposition rate plotted in figure 7. The deposition rate on Si decreased quadratically with the deposition temperature. It was quite low as compared to the values reported e.g. by Massines et al [17] because 4 ppm of HMDSO in their gas mixture resulted in three times higher deposition rate. However, deposition rate is a function of several parameters including a local supply of depleted precursor molecules that needs to be optimized in our experimental set-up.

The optical properties, refractive index and extinction coefficient, are given in figure 8 and figure 9. The refractive index did not exhibit any monotonic dependence on the temperature. Its value at 632 nm ranged from 1.45 to 1.47. The film absorption in UV range evidently decreased with the temperature. The extinction coefficient at 200 nm was in the range 0.014–0.030 for the films deposited at 23–75°C. It decreased down to 0.007 for 100°C and even more to 0.001 for 120 and 150°C. Therefore, the material deposited at higher temperatures approaches the structure of SiO\(_2\) as already suggested by the results of the previous section.

3.4. Film mechanical properties

In case of the indentation tests on thin films, the measured properties depend on the indentation depth due to the combined response of the coating and the substrate (here glass, silicon or polycarbonate). Therefore, the measured elastic modulus and hardness values should be corrected for the substrate influence [44]. In the case of glass and silicon substrates, the elastic modulus and the plastic hardness of the films were lower
than those of the substrate and their values could be calculated using the approach of
Oliver and Pharr [45], if the indentation depth does not exceed approx. 10% of the
film thickness. The resulting film elastic modulus and plastic hardness increased with
increasing deposition temperature approaching the mechanical properties of SiO$_2$ in the
temperature range 120–150$^\circ$C. The film microhardness in dependence on temperature
is depicted in figure 10.

In the case of films on polycarbonates, the hardness and elastic modulus of film
were always substantially higher than those of the substrate. Therefore, it represented
the system of a thin hard coating deposited on a soft viscoelastic-plastic substrate
(polycarbonate) and the conditions of the Oliver and Pharr method were not fulfilled.
The Martens hardness (HM) that measures both elastic and plastic resistances against
indentation was more suitable for the characterisation of the film/substrate systems.
The dependencies of HM on indentation depth for the films deposited at several different
deposition temperatures are shown in figure 11. According to this figure, the surface
hardness of the polycarbonate after nitrogen plasma treatment slightly decreased. The
films deposited at elevated temperatures, on the other hand, substantially increased its
surface hardness. The trend in the mechanical properties with increasing deposition
temperature was similar as for the glass and Si substrates. The films deposited at
the maximum temperature suitable for polycarbonate substrates, i.e. 120$^\circ$C, exhibited

![Deposition rates for different substrate temperatures.](image)

**Figure 7.** Deposition rates for different substrate temperatures.
mechanical properties very close to SiO$_2$. However, crack creations were observed at higher indentation depths as shown in figure 11. The crack creation may be suppressed by deposition of film with graded structure increasing the temperature during the deposition (i.e. with gradual change of film properties from polymer-like to SiO$_2$) [46, 47].

4. Conclusion

The plasma enhanced CVD of silicon oxide thin films was investigated in atmospheric pressure Townsend-like discharge burning in nitrogen. The films were deposited from the mixture of hexamethyldisiloxane (HMDSO) with oxygen in nitrogen working gas on glass, silicon and polycarbonate substrates. The substrate temperature, varied from 25°C to 150°C, had significant influence on the film’s chemical structure and functional properties. The increasing deposition temperature resulted in a decreasing content of carbon and hydrogen and an increasing oxygen-to-silicon ratio in the films. This ratio reached 2.3 for 150°C but not all the oxygen atoms were bonded in the Si-O-Si network as revealed by the presence of OH groups in the FTIR spectra. The hardness of the films on the glass and silicon substrates increased with increasing substrate temperature from 0.4 GPa to 7 GPa. Increasing temperature caused increased hardness also for the
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Figure 9. Extinction coefficient of the films deposited on silicon substrates.

films on polycarbonate substrates. The Martens hardness of the film/polycarbonate system for the deposition temperature of 120°C was several times higher than this of bare polycarbonate at small indentation depths. The deposition rate decreased with increasing temperature from 5 nm min$^{-1}$ at 25°C to 2.8 nm min$^{-1}$ at 150°C. However, the chemical structure, optical and mechanical properties of the films deposited at 150°C approached those of SiO$_2$ films deposited in low pressure discharges [12, 48].

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![Figure 10. The film microhardness for different deposition conditions. The numbers at curves indicate air flow in sccm / nitrogen flow in slm during the deposition.](image)

Figure 11. The Martens hardness of films on polycarbonate at different deposition
temperatures using air flow rate of 16 sccm compared with the Martens hardness of
polycarbonate treated in nitrogen plasma.

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