Influence of Cu impurities and surface temperature to the formation of thin a-C:H(Me) film

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The aim of this experimental work is identify most important physical and chemical processes on the surface during amorphous carbon films formation on the silicon surface with copper nanoclusters (Cu). Duration of films deposition on n type silicon (100) surface by using different substrate temperatures (25 °C, 100 °C and 250 °C) was 45s. Velocity of film growth from C$_2$H$_2$ gas plasma depended on surface temperature and was variable (0.2-0.5 nm/s). Data of null ellipsometry, Raman spectroscopy and element analysis showed that formation of amorphous carbon film is in the early stage and mixture of Si-C, Si-COH and GLC fragments is dominant on the surface. The experimental RS curves were fitted by few Gaussian-shape lines in the spectral range from 500 cm$^{-1}$ to 700 cm$^{-1}$, from 700 cm$^{-1}$ to 900 cm$^{-1}$, from 900 cm$^{-1}$ to 1100 cm$^{-1}$ and from 1100 cm$^{-1}$ to 1900 cm$^{-1}$ and analysis of the additional peaks in all ranges confirmed complexes of different carbons structures in the film. Carbon films with more ordered C-C bonds grows during film formation at higher temperatures (>100°C) on the silicon surface with Cu particles, because atoms of copper penetrate into the deeper layers of silicon. The substrate temperature influenced the surface amorphisation because of active oxygen and hydrogen diffusion into deeper layers and formation of hydrogenated silicon carbon with fragments of amorphous carbon films becomes not significant.

Key words: amorphous carbon films, metal impurities, Raman spectroscopy.
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Introduction

The insertion of metallic phase into the a-C: H matrix improves its mechanical performance and also gives the rise to new material properties [1]. Carbon films with metal impurities are interesting because of applicable in medicine like toxins of microorganisms [2] or in microelectronics as single electron devices [3]. Early stage of amorphous carbon film with or without metal impurities formation is interestingly because single theory of this process not is in nowadays. The few theoretical models is dominant: complete condensation, growth with re-evaporation, nucleation on defects, and total or null cluster-cluster coalescence [4]. It is known, that particles of metals on the surface catalyzed growth of nanoformation during amorphous carbon films deposition. One of the earliest mechanism for CNT growth proposed A. These and others, and it is the so-called “scooter mechanism” [5], where metal atoms with high electronegativity chemisorbs to the open edge of a curving graphene sheet. It avoids formation of an endohedral fullerene and rearrange of carbon rings. Other mechanisms explaining the CNT growth using metal or non-metal catalyst exist also [6-8], but two of them are most acceptable: tip-growth model and base growth model. Interaction between catalyst particle and surface determined which of them would be dominant. Type of nanoformation (single or multy wall carbon nanotube, fullerene et. all) depends on the catalyst size [9]. Surface temperature is other important factor for nanostructure growth. High temperature is the reason for nanoclustering [10] because graphite like carbon films will forms on the surface during deposition in these conditions. Type of nanoformation (single or multy wall carbon nanotube, fullerene and other) depends on the catalyst size [9]. High temperature is the reason for nano-
clustering [10, 11] because graphite like carbon films will forms on the surface during deposition in these conditions [12]. Formation of graphite like carbon film at comparative low temperatures is possible using PECVD methods, so addition of metals particles can be reason of nanostructures growth. Influence of metal additions and temperature to the properties of thin amorphous carbon films and possibilities of nanostructures formation are analyzed in this experimental work. The aim of it is to identify of most important physical and chemical processes on the surface during thin amorphous carbon films formation on the silicon surface with metals clusters (Cu) by PECVD method.

**Experimental setup**

a-C: H films were formed on silicon surface with copper clusters by PECVD method using C\textsubscript{2}H\textsubscript{2} gas plasma. The temperature of silicon substrate was 25 °C, 100 °C and, 250 °C. Duration of deposition was 45 s bombarding ion energy was 200 eV. The copper was deposited before carbon film formation during bombardment the copper net by argon ions with ion energy 500 eV, deposition duration 20 min. The deposition and film formation took place in a single two steps technological process. The films optical properties were determined by null-ellipsometry (Gaertner L117 with a He–Ne laser (632.8 nm)), Raman spectroscopy (Ivon Jobin spectrometer with a Spectra Physics YAG: Nd laser (532.3 nm, 50 mW, spot size 0.32 mm)). The experimental RS curves were fitted by few Gaussian-shape lines in the spectral range from 500 cm\textsuperscript{-1} to 700 cm\textsuperscript{-1}, from 700 cm\textsuperscript{-1} to 900 cm\textsuperscript{-1}, from 900 cm\textsuperscript{-1} to 1100 cm\textsuperscript{-1} and from 1100 cm\textsuperscript{-1} to 1900 cm\textsuperscript{-1}. The relative element concentrations were measured by Bruker AXS Microanalysis GmbH.cm\textsuperscript{-1}.

**Results**

The main properties of films and peaks values of Raman spectra are shown in the table 1 and the table 2. Deposition rate depended on the nature of metal clusters and substrate temperature (0.2 -0.5 nm/s). Differences in the Raman spectra of silicon substrate (shift 517 cm\textsuperscript{-1}, cm\textsuperscript{-1}, 561 cm\textsuperscript{-1}, 617 cm\textsuperscript{-1} and 669 cm\textsuperscript{-1} were obtained in the Raman spectra of amorphous carbon films deposited on the silicon surface with Cu clusters at the room temperature. The peak at 517 cm\textsuperscript{-1} associated with silicon substrate but it shifts to the lower value when surface temperature increased (516 cm\textsuperscript{-1} at 100 °C and 509 cm\textsuperscript{-1} at 250 °C). Shifting of the silicon peak to lower values corresponds with the process of amorphization and intensity of it depends on substrate temperature [13]. Existing peak at 532 cm\textsuperscript{-1}-535 cm\textsuperscript{-1} correlate with the strain in the silicon [14] and at the higher temperature it become more intensive. The next two peaks in this range (561 cm\textsuperscript{-1} and 617 cm\textsuperscript{-1}) are related with Cu\textsubscript{2}O because this compound have three characteristic vibrations: at 570 cm\textsuperscript{-1}, 618 cm\textsuperscript{-1} and 624 cm\textsuperscript{-1} [15]. Intensities of that are variable and depend on the temperature: the third peak is weak and is not fixed in the spectra when substrate temperature is 25 °C and 250 °C, but becomes dominant at 100 °C. Intensity of the second peak (617 cm\textsuperscript{-1}) decreasing at 100 °C, but it is dominant and shifted to the lower values (614 cm\textsuperscript{-1}) at 250 °C when other two peaks disappear. Variations of these peaks show dependence of structure and concentration of metal in the film.
cooper bonds with oxygen in the film and interlayer on the surface temperature. Wide peak at 669 cm\(^{-1}\) is related to the Si-O-Si stretching [16] and its shift to the lower values with the temperature increasing is conditioned by the bond deformation or deficiency of oxygen.

Table 2 a – C:H(Me) film peaks values of Raman spectra

| Vibration modes                        | Peaks n RS spectra of the films, cm\(^{-1}\) | References |
|----------------------------------------|---------------------------------------------|-------------|
| Si-Si TO mode                          | 509-518; 520;                               | [13]        |
| Strain in Silicon                      | 533; 535; 537;                              |             |
| Cu\(_2\)O                              | 570; 614; 617; 623; 570; 618; 624;          | [15]        |
| SiOH                                   | 586-610; 606;                               |             |
| Symmetric Si-O-Si stretching           | 664-669;                                    | [16]        |
| Si-CH\(_3\) rocking                    | 690; 687;                                   | [17]        |
| Si-C                                   | 682; 732; 784; 763-765; 809-813; 841, 874; 855; | [18]        |
| C-H                                    | 1036-1037; 1079;                            | [21]        |
| Oxyethylene ring deformation           | 1122;                                       | [22]        |
| D4                                     | 1181-1197; ~1190;                           | [23]        |
| C-O-C stretching                       | 1247; 1252;                                | [24]        |
| C-C inter-ring stretching              | 1288; 1286;                                | [25]        |
| D peak                                 | 1313-1396; 1350;                           | [26]        |
| Transpolyacetylene \(v_c\) C=C stretching mode | 1410-1454; ~1450;                          | [25]        |
| Semicircle stretch vibrations in benzene or benzene clusters | 1484; 1486;                               | [26]        |
| G peak \(E_g\) symmetry                | 1510-1553; 1550;                           | [23]        |
| D\(^{\prime}\) (G2 mode)               | 1611-1637; 1620;                           | [26, 27]    |
| C=O stretching                         | 1713-1788; 1680-1820;                       | [28]        |

The characteristics peaks for hydrogenated amorphous silicon (SiOH) (586 cm\(^{-1}\) and 639 cm\(^{-1}\)) and Si-C (682 cm\(^{-1}\)) becomes more intensive in the spectra when the substrate temperature is 100 °C, but peak of Si-O-Si is less intensive. It means that competition of processes hydrogenation and oxidation of silicon is very important for interlayer formation of the early stage film deposition when substrate temperature increase.

The peak at 732 cm\(^{-1}\) associated with O-C-O bonds in plane bending or CH\(_2\) in rocking mode is obtained in the spectra range of 700 cm\(^{-1}\) – 900 cm\(^{-1}\) for samples with Cu impurities. This range is typical for silicon carbide. Most likely that this peak is related with amorphous SiC because other peaks characteristic for it at 763 cm\(^{-1}\), 784 cm\(^{-1}\), 809 cm\(^{-1}\), 841 cm\(^{-1}\) and 874 cm\(^{-1}\) were obtained also. Their intensity decreased with increasing temperature and it show that structure of silicon carbide becomes more orderly because weak peak of crystalline SiC at 855 cm\(^{-1}\) is defined in the spectra. It is important that peaks related with C-H bonds deformation at 750 cm\(^{-1}\) (also peaks at 1037 cm\(^{-1}\) and 1079 cm\(^{-1}\) in the RS range of 1000 cm\(^{-1}\) – 1800 cm\(^{-1}\)) becomes more intensive with increasing temperature. And it confirms that the oxygenation becomes less intensive than the hydrogenation of silicon or carbon when temperature of surface increases.

Typical asymmetric curve for amorphous carbon films was not obtained in the Raman spectra range at
1000 cm\(^{-1}\) to 1800 cm\(^{-1}\). The separate peaks at 1197 cm\(^{-1}\), 1313 cm\(^{-1}\), 1409 cm\(^{-1}\), 1510 cm\(^{-1}\), 1552 cm\(^{-1}\), 1611 cm\(^{-1}\) and 1738 cm\(^{-1}\) are identified in the spectra of samples deposited in the 25 °C. First of them together with peak at 1409 cm\(^{-1}\) can be related to the vibration of transpolyacetylene carbon groups, but confirmation of it is complicated because other authors [22] demonstrate that intensity of these peaks increase with increasing of hydrogen concentration in the plasma and it is not typical for transpolyacetylene. So, the peak at 1197 cm\(^{-1}\) can be attributed to any known sp\(^2\) – related feature also, so in agreeing with the other authors it was remarked that this peak shows highly defective graphitic structure relaxes through the formation of occasional four-coordinated atoms between the planes [29, 30].

So, the shift to the lower values with increasing of the surface temperature is related with increasing of order in the graphitic structure, the relaxing of internal stress and the decreasing of defect. Some authors peak at 1450 cm\(^{-1}\) ascribe to fullerene-like structures [31]. Confirmation of it is formation of these structures during longer time (300 s) deposition at the same experimental conditions [32]. It was obtained that this peak shifted to higher values with increasing temperature of samples and becomes equal to 1453 cm\(^{-1}\) at 250 °C. Other peaks at 1510 cm\(^{-1}\), 1552 cm\(^{-1}\), 1611 cm\(^{-1}\) can be attributed to variation of G band.

Peak at 1611 cm\(^{-1}\) is dominated when temperature of surface is 25 °C and it can be related with D’ or G2 mode because the tangential G-band (at ~1580 cm\(^{-1}\)), which derived from the graphite-like in-plane mode, can split into several modes where two are the most distinct: G1 (1577 cm\(^{-1}\)) and G2 (1610 cm\(^{-1}\)) [33]. This splitting was also observed in the other papers [33-35].

The peak at 1738 cm\(^{-1}\) comes from C=O stretching vibration and is intensive at low temperature. It confirmed that SiCOH groups formed on the surface. This peak becomes weak at 100 °C substrate temperature, but it arises becomes more intensive in the spectra at 250 °C.

Discussion

EDS analysis shows, that initial concentration of Cu is same (1.2%) for all samples but films thickness and growth velocity decreased with increasing of sample temperature. Theoretically, only relative concentrations of carbon and silicon will depend on the film thickness, but cooper concentration will remain same because cooper clusters are in the interlayer and depths of EDS measurement are constant. But experimental results show that cooper concentration also depends on surface temperature and it increased with temperature increasing. So, it is reason for assumption that cooper can be removed from the surface because of desorption or ion bombardment during film deposition at 200 eV ion energy and 25 °C temperature. For that reason concentration of cooper become,
lower (0.8%) and traces of cooper oxides were obtained in the RS spectra. From another side, cooper diffused into the deeper layers using silicon defects when substrate temperature is 100 °C. Concentration of it become lower in the film, but not in the film/interlayer/substrate structure and EDS measurements shows that total Cu concentration is higher (1.16%) but intensity of cooper oxides peaks becomes lower in the Raman spectra. Peaks of Cu₂O disappears, but relative concentration of Cu remain similar (1.18%) when temperature increases till 250 °C and it show that diffusion process becomes more intensive at higher temperature. Amorphisation of silicon confirms these assumptions (shift of the silicon peaks into the lower values at higher temperatures, table 2). Single peak at 614 cm⁻¹ remains in the RS spectrum at highest temperature case, but it not related with cooper oxide because other additional peaks not exist there. Also single similar peaks at 613 cm⁻¹ – 617 cm⁻¹ were obtained in the samples with Ag and Au [32], so it was decided that this peak is attributed to the C–C–C deformation in-plane vibration [36].

Analysis of the RS spectra in the other ranges shows that hydrogen remains in the forming film while Cu atoms bonds with oxygen and only part of it penetrate into the silicon substrate at low temperature. Slow film formation and more intensive penetration of hydrogen and oxygen into the silicon at higher temperatures create condition for hydrogenated silicon carbide formation. More intensive peaks related with Si-H, Si-C and Si-O and obtained in the spectra confirm this assumption. It is important also, that typical for the amorphous hydrogenated carbon films D and G modes in the spectra have low intensities and it shows that a-C:H films is in early stage formation at low temperature. C-C bond become more order when temperature increased and it conditioned that thickness of this ordered film decreased. Measurement of refracting indexes also confirms these assumptions. Low refractive index (1.4) of films deposited at 25 °C is more characteristic for SiCOH films formed by PECVD method [37] than amorphous carbon film. Refractive index increased with temperature increases and becomes ~1.6 at 250 °C. Featureless visible Raman spectra with no clear Raman peaks shows that films becomes polymer like (PLC) with low Young modulus (≤20 GPa) and low density (≤1.2 g/cm³) [38].

Conclusions

Formation of hydrogenated silicon carbon with fragments of amorphous carbon films on the silicon surface was obtained during early stage of a: C-H films deposition by PECVD methods. Carbon films with more ordered C-C bond grows during film formation at higher temperatures (>100° C) on the silicon surface with Cu particles, because atoms of cooper penetrate into the deeper layers of silicon. Amorphisation of substrate increases also. Lower refractive index than typical amorphous carbon films confirmed that the films deposited at 25 °C is more characteristic for SiCOH films. The typical for the amorphous hydrogenated carbon films D and G modes in the spectra have low intensities when films were formed at low temperature. More polymer like films forms at higher temperatures of the substrate.

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