Protective Sliding Carbon-Based Nanolayers Prepared by Argon or Nitrogen Ion-Beam Assisted Deposition on Ti6Al4V Alloy

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The microstructure and the surface properties of samples coated by carbon-based nanolayer were investigated in an effort to increase the surface hardness and reduce the coefficient of friction of the Ti6Al4V alloy. Protective carbon-based nanolayers were fabricated by argon or nitrogen ion-beam assisted deposition at ion energy of 700 eV on Ti6Al4V substrates. The Raman spectra indicated that nanolayers had a diamond-like carbon character with sp² rich bonds. The TiC and TiN compounds formed in the surface area were detected by X-ray diffraction. Nanoscratch tests showed increased adhesion of a carbon-based nanolayer deposited with ion assistance in comparison with a carbon nanolayer deposited without ion assistance. The results showed that argon ion assistance leads to greater nanohardness than a sample coated by a carbon-based nanolayer with nitrogen ion assistance. A more than twofold increase in nanohardness and a more than fivefold decrease in the coefficient of friction were obtained for samples coated by a carbon-based nanolayer with ion assistance, in comparison with the reference sample.

1. Introduction

Carbon-based layers attract much attention due to their excellent sliding properties [1, 2] and their good wear resistance [3]. They could potentially be used as a protective film or as a functional layer in many industrial applications, for example, as protective thin films for recording media, sensors, and medical implants [4–6]. The deposition technique and the processing conditions have a direct influence on the chemical composition, the structure, and the physical, mechanical, and surface properties [7]. Various techniques have been used for depositing a carbon-based layer, including ion-beam assisted deposition (IBAD), chemical vapor deposition (CVD), plasma-enhanced chemical vapor deposition (PECVD), sputtering, and filtered cathodic vacuum arc deposition (FCVA) [8–12]. The IBAD method is very promising for the deposition of carbon-based layers. Ion bombardment significantly affects the performance of the layers. Preferred surface properties are produced by carbon-based coating surfaces with sp² and sp³ hybridizations [13]. In order to optimize the carbon-based layer deposition method, it is necessary to understand the mechanisms involved in the growing process and in the degradation process [13, 14]. In his review paper on amorphous carbon, Robertson presents his theory of the growth of amorphous carbon layers and introduces a model of subplantation [15]. Dopants such as nitrogen and hydrogen also play a critical role in respect to properties such as hardness, friction, and internal stress. Incorporating nitrogen in amorphous carbon increases the fraction of sp² hybridized carbon atoms [16]. Thus carbon-based layers with nitrogen incorporation can provide low-friction behavior, high wear resistance, and reduced internal stress [16–18]. In biomedical engineering, titanium alloys are used especially for their good biocompatibility and their low elastic modulus [19, 20]. However, the surface properties of titanium-based implants, such as friction and wear resistance, are not fully satisfactory for use in biomedicine, and they are often modified.

In this work, the surface properties of Ti6Al4V titanium alloy were modified by ion-beam assisted deposition of
carbon-based nanolayers. Protective sliding carbon-based nanolayers were deposited with the assistance of argon ions or nitrogen ions. The nanolayers were characterized by X-ray diffraction (XRD) and by Raman spectroscopy. The elemental chemical composition was measured by glow discharge optical emission spectroscopy (GD-OES). Nanohardness was measured by nanoindentation, adhesion was investigated by the nanoscratch test, and the sliding properties were tested on a ball-on-disc tribometer. The aim of our work was to investigate the effect of the deposition conditions on the microstructure and on the properties of carbon-based nanolayers, in order to improve the surface properties of biomedical Ti6Al4V alloy.

2. Materials and Methods

2.1. Preparation Process. Ti6Al4V titanium alloy with the α + β phase microstructure was used as the substrate material. The disc samples (20 mm in diameter and 6 mm in thickness) were cut from a titanium alloy bar and were then ground with a series of waterproof abrasive papers up to 2000 grit. Final polishing was performed with 5 μm and 1 μm diamond paste. Before the deposition process, the samples were ultrasonically degreased in acetone and in isopropyl alcohol, both of them for 20 minutes, respectively.

The samples were placed on a sample holder mounted on a rotary manipulator in a vacuum chamber and were sputtered with 700 eV argon ions to etch and activate the surface. A carbon nanolayer approximately 100 nm in thickness with a tungsten interlayer approximately 20 nm in thickness was deposited by electron beam evaporation of high purity carbon and tungsten targets. The deposition process was carried out with simultaneous assistance of argon or nitrogen ion bombardment. The deposition rate of the carbon and tungsten nanolayers was about 0.10 nm s⁻¹, and the work pressure was about 5 × 10⁻³ Pa. The energy of the accelerated argon ions and nitrogen ions was 700 eV, and the beam current density was about 80 μA cm⁻².

2.2. Characterization Methods. The elemental chemical composition was measured by glow discharge optical emission spectroscopy (GD-OES). A LECO GDS850A optical emission spectrometer was used. GDS850A is an optical emission spectrometer with a Paschen-Runge polychromator with a focal length of 750 mm, and it has a glow discharge spectral source with a flat cathode (Grimm-type). The sample itself serves as a cathode in this arrangement. Atomization of the sample surface is performed by the mechanism of cathode sputtering under an anomalous discharge. The process gas was argon. The spectrometer was calibrated using reference materials with a known composition and known sputter rates, and the “standard” calibration model was used.

The Raman spectra were measured using a Renishaw RM 1000 Raman microscope with Ar laser excitation at 514.5 nm. The data were acquired from 3 μm diameter spots with neutral filters limiting the laser power to 10%, that is, around 0.1 mW μm⁻². This level of power was chosen after a set of measurements in which the power was slowly stepped up in a series of spectra acquisitions.

The phase composition was investigated by the X-ray diffraction (XRD) method. The measurements were performed on a PANalytical X’Pert PRO powder diffractometer. We chose a geometry with the parallel beam with an incident angle of 0.5° with respect to the intensity of the diffraction pattern. The diffraction was recorded for the 2θ range 10°–80°. Cobalt radiation with a wavelength of 0.1789 nm was used.

The nanohardness of the modified samples was investigated by nanoindentation testing. The dependence of the nanohardness on depth was registered. The measurements were performed on a Hysitron TribolIndenter™ with a Berkovich diamond tip, using the CMX dynamic mode (continuous measurement of X) with a maximum indentation force of 3000 μN and a prescribed load/unload frequency of 85 Hz. Nanoindentation testing proceeded at a temperature of 23.2°C. Grids of 4 × 4 indents with 5 μm separation were applied for each sample; see Figure 6. The tip area function was calibrated on the fused quartz sample according to a standard calibration procedure [21, 22].

The adhesion of carbon-based nanolayers was investigated by nanoscratch testing on the Hysitron TribolIndenter with scratch load resolution of 3 × 10⁻⁴ μN, scratch force resolution of 0.5 μN, and displacement resolution of 3 nm. The measurements were performed with a 60° conical tip <1 μm in radius. The maximum load was up to 10000 μN on a scratch length of 4500 nm. The scratch scan recorded the penetration depth with increasing scratch distance, increasing scratch load, and increasing scratch force during nanoscratching.

A sliding test was performed in order to find out the coefficient of friction. A ball-on-disc tribometer was used. A constant normal load of 1 and 2 N was set during the test. The velocity was 8 cm s⁻¹. The second part of the sliding couple was a 100Cr6 steel ball 6 mm in diameter. The sliding tests proceeded at a laboratory ambient temperature of 25°C, without lubrication and with relative humidity of 54%. The measurements were performed without sucking off the abrasive wear products.

3. Results and Discussion

3.1. Elemental Chemical Composition. Typical GD-OES depth profiles of elements of the modified surface area are shown in Figure 1. The concentration profiles of the elements indicate that there is no sharp interface between the titanium alloy substrate, the tungsten interlayer, and the carbon-based nanolayer. The carbon-based nanolayer mixed with substrate atoms can be observed in both types of samples in Figures 1(a) and 1(b). The sample modified by a carbon-based nanolayer with argon ion assistance shows a larger area of atomic mixing. This indicates particularly the concentration profile of titanium, which was detected near the surface. Carbon and other elements detected at a deeper level can also indicate atomic mixing, but the concentration of these elements deep inside the substrate is influenced by the crater effect. A crater is formed during the GD-OES measurement, and sputtering from the crater wall influences the depth profiles. Diffusion may play some role, but the dominant influence is from sputtering from the crater wall. In the sample modified by a carbon-based nanolayer with
nitrogen ion assistance, the average nitrogen concentration was found to be approximately 40 at% at a depth down to approximately 75 nm. Nitrogen was also detected in the sample modified by a carbon-based nanolayer with argon ion assistance. The nitrogen on the surface of the sample and a shallow layer up to a few nanometers originated from contamination due to air exposure. Argon was used as the process gas, so the argon content cannot be evaluated from the GD-OES spectra.

3.2. Microstructure. During the IBAD process a continuous carbon-based monolayer was deposited on the Ti6Al4V alloy surface. Figure 2 presents SEM micrograph of the transversal section in which a continuous carbon-based layer with tungsten interlayer is shown. The continuous carbon-based monolayer was observed in all modified samples. SEM micrograph of the transversal section supplemented by EDS analysis (Figure 3) confirms the chemical composition of the created surface layer.

The microstructure and the phase composition of the modified samples were examined by XRD. This method can provide direct evidence of compounds formed in the modified surface area. An incident angle of 0.5° was chosen with respect to the intensity of the diffraction spectra of a thin surface layer. Figure 4 shows the diffraction spectra of the modified samples. The alpha structural phase of the titanium alloy ($\alpha$-Ti) was observed in all samples, while no minority beta structural phase ($\beta$-Ti) was identified in the diffraction spectra. An additional $\alpha$-Ti phase with a larger lattice parameter appeared in the diffractograms of samples modified by a carbon-based nanolayer with nitrogen ion assistance. The implanted nitrogen ions were placed in the interstitial sites, and a solid solution of nitrogen was formed in the surface area. Some of the interstitial nitrogen atoms...
These findings show that nitrogen ion assistance during the deposition of carbon-based nanolayers leads to the formation of a TiN compound on the interface of a carbon-based nanolayer/titanium alloy substrate and stabilizes the \( \alpha \)-Ti phase in the surface area. Schmidt et al. [23] and Vlack et al. [24] have confirmed that nitrogen ion implantation on Ti-6Al-4V stabilizes the \( \alpha \)-Ti phase and forms TiN compounds.

In both types of samples, XRD analysis showed the presence of the TiC phase; see Figure 4. The results of XRD analysis confirm that ion assistance during deposition of carbon-based nanolayers led to a mixed surface area in which TiN and TiC compounds were found. The broad structure of the carbon was investigated by Raman spectroscopy. Figure 5 compares the Raman spectra of a carbon-based nanolayer deposited with argon or nitrogen ion assistance and the spectrum of a carbon nanolayer deposited without ion assistance. The Raman spectra of samples coated by a carbon-based nanolayer with argon ion assistance (C:Ar nanolayer) and with nitrogen ion assistance (C:N nanolayer) were compared with the spectrum of a C nanolayer. The Raman spectra show that the ratio of the integrated areas under peaks 1, 2, and 3 decreases and the position of peak 3 (the G peak) moves slightly to lower frequencies due to ion assistance during the deposition process. The observed changes in the Raman spectra characterize the transformation of graphitic carbon into a DLC (diamond-like carbon) character, which explains the increase in hardness shown in Figure 7. Similar trends indicating the DLC character of the coatings were observed by TiN phases, as shown in Figure 3. The SEM image of the polished transverse section of a sample modified by a carbon-based nanolayer with nitrogen ion assistance and corresponding EDS line scan analysis.
### Table 1: Peak parameters of the fitted Raman spectra.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Peak 3 (cm$^{-1}$)</th>
<th>Peak 2 (cm$^{-1}$)</th>
<th>Peak 1 (cm$^{-1}$)</th>
<th>Integrated areas ratio $(1+2)/3$</th>
</tr>
</thead>
<tbody>
<tr>
<td>C:Ar</td>
<td>1563</td>
<td>1402</td>
<td>1178</td>
<td>2.43</td>
</tr>
<tr>
<td>C:N</td>
<td>1565</td>
<td>1407</td>
<td>1198</td>
<td>2.67</td>
</tr>
<tr>
<td>C</td>
<td>1567</td>
<td>1394</td>
<td>1123</td>
<td>3.09</td>
</tr>
</tbody>
</table>

Figure 6: Representative 3D post scans of indents after nanoindentation testing of a sample modified by a carbon-based nanolayer with argon ion assistance (a) and a sample modified by a carbon-based nanolayer with nitrogen ion assistance (b).

observed by Morshed et al. [26] and by Ziebert et al. [27]. The correlation with the nanohardness results shows that the nanohardness increases as the ratio of the integrated areas decreases. It can be estimated from the Raman spectra that the carbon-based nanolayer consists predominantly of sp$^2$ bonds with a small sp$^3$ content [15].

### 3.3. Nanohardness

The nanohardness of the modified samples was investigated by nanoindentation testing. Representative post scans of the indents are shown in Figure 6. A small plastic deformation region can be seen around the indents. The resulting nanohardness depth profiles are shown in Figure 7. It is seen that the nanohardness at shallow depths up to few nanometers is small, and it increases rapidly with contact depth to the maximum value. The nanohardness values in this region are influenced by the error originating from the standard calibration of the tip area function on fused quartz [21, 22]. As the contact depth increases further, the nanohardness decreases from the maximum value and converges to the hardness of the reference sample (see Figure 7). The reduction in nanohardness is a consequence of the influence of the softer titanium alloy substrate. The maximum indentation hardness values of the modified samples are shown in Table 2. The nanohardness of samples coated by a carbon-based nanolayer with argon ion assistance was increased by a factor of approximately 2.31, and the nanohardness of a sample coated by a carbon-based nanolayer with nitrogen ion assistance was increased by a factor of approximately 2.22, compared to the maximum hardness value of the reference sample. The increase in nanohardness can be attributed mainly to the bond structure of carbon. It can be assumed that the mixing area may also have an impact on the increase in nanohardness. The hard TiC and TiN compounds detected by XRD in the interface of the carbon-based nanolayer/titanium alloy substrate also hardened the surface area. The nanohardness depth profiles in Figure 7 contain the depth profile of the nanohardness of the sample coated by a carbon nanolayer without ion assistance. The maximum hardness value of the sample coated with a 100 nm thick carbon nanolayer without ion assistance is $8.74 \pm 0.64$ GPa at a depth of 30 nm. The nanohardness of the reference sample decreases with increasing contact depth from a maximum value of $5.33 \pm 0.41$ GPa to almost constant value of 4.6 GPa. The nonconstant course of the nanohardness of the reference sample may be caused by mechanical strengthening of the surface due to cutting and polishing.

### 3.4. Adhesion

The adhesion of nanolayers to a titanium alloy substrate was investigated by nanoscratch testing over a continuous range of scratch loads. The trend of the penetration depth according to the scratch load for samples coated by carbon-based nanolayers with ion assistance and for the sample coated by a carbon nanolayer without ion assistance is presented in Figure 8. The penetration depth increases with increasing scratch load. A comparison of the curves in Figure 8 shows that the penetration depth for the same scratch load value is greater for the sample coated by a carbon nanolayer without ion assistance. This shows that the resistance to penetration of the scratch tip increases due to
Table 2: Maximum nanohardness values of the modified samples and the reference sample.

<table>
<thead>
<tr>
<th>Sample</th>
<th>C:Ar</th>
<th>C:N</th>
<th>Ref. sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum indentation hardness (GPa)</td>
<td>12.34 ± 0.64</td>
<td>11.84 ± 0.34</td>
<td>5.33 ± 0.41</td>
</tr>
<tr>
<td>Contact depth (nm)</td>
<td>11</td>
<td>14</td>
<td>6</td>
</tr>
</tbody>
</table>

Figure 7: Resulting nanohardness depth profiles.

Figure 8: The trend of the penetration depth according to the scratch load for samples coated by carbon-based nanolayers with ion assistance and the sample coated by a carbon nanolayer without ion assistance.

Figure 9: Scratch force versus scratch distance during nanoscratch experiments.

ion assistance during the nanolayer deposition process. This finding is in agreement with the nanohardness results. The sample coated by a carbon-based nanolayer with argon ion assistance and sample coated by a carbon-based nanolayer with nitrogen ion assistance show similar trends. A small difference can be observed for a greater scratch load from approximately 6000 μN. Figure 8 shows a smooth course of the penetration depth along the scratch load for samples coated by a carbon-based nanolayer with ion assistance, while the course of the penetration depth of the sample coated by a carbon nanolayer without ion assistance shows step increases in the penetration depth at a constant scratch load. Similar trends can be observed in Figure 9, which presents the scratch force versus the scratch distance. The visible jump of the scratch force curve for the sample coated by a carbon nanolayer without ion assistance represents the increase in the scratch distance without an increase in the scratch force. The jump indicates delamination of the carbon nanolayer, which is confirmed by the slight decrease in the scratch force in Figure 9 and is accompanied by the step increase in the penetration depth shown in Figure 8. The carbon-based nanolayers deposited with ion assistance show good adhesion with no signs of delamination of the nanolayers throughout the course of the nanoscratch tests. It can be assumed that the mixing area detected by GDOS and XRD improved the adhesion of the nanolayers.

3.5. Sliding Behaviour. Figure 10 shows the evolution of the coefficient of friction at a normal load of 1N for samples coated by carbon-based nanolayers with ion assistance, without ion assistance, and for the reference sample as a function of the number of cycles. The friction curve of the sample coated by a carbon nanolayer without ion assistance can be divided into two parts. The first part represents the low-friction regime, while the second part represents the increased friction regime. The friction coefficient in the low-friction part is low, having a value of about 0.13. The
Figure 10: A comparison of the friction coefficient as a function of the number of cycles for samples coated by a carbon-based nanolayer with ion assistance, without ion assistance, and for the reference Ti6Al4V sample (ball-on-disc test, 1 N normal load, 100Cr6 ball, 8.00 cm s⁻¹ linear speed).

Figure 11: Coefficient of friction of samples coated by a carbon-based nanolayer with ion assistance and the reference Ti6Al4V sample (ball-on-disc test, 2 N normal load, 100Cr6 ball, 8.00 cm s⁻¹ linear speed).

4. Conclusion

Protective carbon-based nanolayers were fabricated by argon or nitrogen ion-beam assisted deposition at ion energy of 700 eV on Ti6Al4V substrates. The carbon-based nanolayers were found to have a diamond-like carbon character with sp² rich bonds. TiC and TiN compounds were found in the surface area. Increased adhesion of carbon-based nanolayers deposited with ion assistance was observed, in comparison
with carbon nanolayers deposited without ion assistance. The nanohardness of a sample coated by a carbon-based nanolayer with argon ion assistance was increased by a factor of 2.31, and the nanohardness of a sample coated by a carbon-based nanolayer with nitrogen ion assistance was increased by a factor of 2.22, in comparison with the reference sample. The very stable course and the same coefficient of friction value of 0.11 of the samples coated by a carbon-based nanolayer with argon ion assistance or with nitrogen ion assistance were maintained at a normal load of 2 N. In comparison with the reference sample, a greater than fivefold decrease in the coefficient of friction at a normal load of 2 N for samples coated by carbon-based nanolayers with ion assistance was obtained.

**Conflict of Interests**

The authors declare that there is no conflict of interests regarding the publication of this paper.

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