I. Introduction

Carbon is the sixth most abundant element in the universe. In addition, carbon is a very special element because carbon is your body, food you eat, clothes you wear, cosmetics you use and gasoline that fuels your car. Carbon, discovered in prehistory was known to the ancients, who manufactured it by burning organic material for charcoal production. In Fig. 1 there are four popular allotropes of carbon: amorphous carbon, graphite, diamond and fullerene (nanotubes) [1].

Diamond and DLC have many excellent electrical, chemical, mechanical and biological properties [2]. Diamond is the hardest material known, it is extremely chemically inert and biologically compatible.

Electrical properties of diamond are: the highest electrical breakdown voltage, the highest thermal conductivity and the widest electromagnetic radiation transparency range of any material, as well as wide bandgap and high carrier mobilities. As an illustration of the excellent properties of diamond, Fig. 2 compares various wide-gap materials in terms of carrier mobility and thermal management properties, represented by surface area of the plotted circles. Recent progress in chemical vapor deposition (CVD) diamond technology has enabled the preparation of high-quality n-type CVD diamond layers using phosphorus as an n-type dopant and boron, nitrogen, hydrogen as p-type dopants. CVD diamond can therefore be considered as a new interesting conventional wide-gap semiconducting material having both n-type and p-type conductivity, which makes it attractive for numerous applications in high-temperature, high-voltage and high-frequency devices [3].

Fig. 1 Carbon Allotropes: Diamond, Buckminster fullerene, Carbon nanotube, and Graphite

Fig. 2 Comparison of mobility materials together with thermal conductivity as represented by the surface area of the circles. The inset shows predicted performances, expressed as the square root of the output power, for high-frequency applications by using various semiconducting materials.
The electrochemical approach is attractive because of sensitivity and dynamic range, portability and ease of application. Diamond and DLC is an attractive alternative electrode for electroanalysis on account of the reproducibility and chemical inertness, low background currents, and wide working potential window which this material exhibits in electrochemistry. One of the applications of diamond and DLC, carbon nitride CNx, is for fast detection and determination of trace metals in solutions. Major importance lies in electrochemical analysis, where single or multiple heavy metals-containing systems are invariably encountered. Electrochemical stripping voltammetry methods offer a simple, quick and cheap way of detecting trace metals such as (Pb, Cu, Cd, Mn, Ag) in water [4, 5].

Biomaterials are an important aspect in the development of biomedical devices and implants and this area of research has been expanding rapidly over the last 50 years. It is the surface of a biomaterial which first contacts with the living tissue when biomaterial is placed in the body. Titanium alloy (Ti-6Al-4V) is well established as a primary metallic biomaterial for orthopedic implants. However, the host response to Ti-6Al-4V is not always favorable, whereby a fibrous layer may form at the skeletal tissue-device interface, resulting in implant failure. Therefore, there is a need to develop novel micro-engineered surfaces to provide better biological outcomes. DLC and carbon nitride CNx are excellent candidates for use as biocompatible coatings on biomedical implants, which are due to not only their excellent properties but also their chemical composition containing only carbon, hydrogen and nitrogen, which are biologically compatible elements [6].

2. Experimental

Diamond deposition
Deposition of polycrystalline diamond (PCD) over surface was carried out in the double bias enhanced HF CVD reactor described previously [7]. The gas phase was a mixture of 2% CH4 in H2, the total pressure in the reactor was 3 000 Pa and flow rates were 6:300 sccm. Gases were activated by 5 tungsten filaments above-situated grid (positive grid bias of 100 V with respect to filament). Plasma formed in the region between heated filaments and the stage of plasma formation depended on account of the reproducibility and chemical inertness, low background currents, and wide working potential window which this material exhibits in electrochemistry. One of the applications of diamond and DLC, carbon nitride CNx, is for fast detection and determination of trace metals in solutions. Major importance lies in electrochemical analysis, where single or multiple heavy metals-containing systems are invariably encountered. Electrochemical stripping voltammetry methods offer a simple, quick and cheap way of detecting trace metals such as (Pb, Cu, Cd, Mn, Ag) in water [4, 5].

DLC deposition
DLCs were deposited in a UVNIPA-I-001 vacuum system with three sources (gas ion source for cleaning, electric arc source for non-magnetic metal sputtering and pulse arc carbon source for DLC deposition). The pulse sputtering of graphite target is a possible setup in the range of (/f = 1, 2, 3, 5, 10, 15, 20 and 30 Hz).

Samples entering were sputtered in one vacuum cycle. All of substrates (microelectrode arrays and medical hips) were cleaned for 10 min with Ar ions. After cleaning the hip was covered with 50 nm Ti interlayer for better adhesion of DLCs and for heated up bulk substrate to temperature about 200 °C. For deposition of MEA (the technological process is described later) is this step missing, but a Ti interlayer is not needed. Consequently the DLC layer (~ 150 °C) was deposited at low deposition temperature. Nitrogen with 30 sccm flow rate was added during the DLC deposition into a working chamber. The substrates were planetary rotating through all the deposition steps for homogeneous deposition.

MicroElectrode Array
Fig. 3 shows all the technological steps of manufacturing a 1 × 1 mm microelectrode array (MEA) structure consisting of 50 625 microdiscs with 3 μm in diameter and distance of 20 μm between microdiscs on Si with low resistivity (0.008 – 0.024 Ωcm).

The 1-st step was to clean the substrate in ultrasonic acetone bath. Then, the 2-nd step, about 300 nm thick SiO2 layer was created during 6 hours with the substrate at 1200 °C in dry O2 atmosphere in a furnace. During the standard photolithographic process, as steps 3 to 6, MEA structure in SiO2 layer was produced through a lithographic mask. After MEA preparation there are two ways for deposition of carbon layers (diamond or DLC). The 7-th step.

After DLC deposition the lift off technique is potentially used for SiO2 removal from the substrate. The overgrown diamond or DLC layer is removed if necessary, the 8-th step. At this stage of the overall process we have a completely prepared MEA for electrochemical measurement, the 9-th step.

Hip replacement joints
Standard biocompatible alloy of Ti-6Al-4V supplied by prof. Zitánsky from the Faculty of Mechanical Engineering of the Slovak University of Technology in Bratislava was used as the substrate. The substrate face corresponds to special human hip replacement joints with the surface roughness after rough machining of about Rq = 10 μm, were 10 min. rinsed in acetone in an ultrasonic bath and than washed in deionized water.
3. Results and discussion

We can see in Fig. 4 SEM images of a deposited layer surface after the complete technological process. In Fig. 4 a) MEA polycrystalline diamond structure, in Fig. 4 b) CNx layer structure are displayed, both deposited on Si. In case of polycrystalline diamond, the diamond layer was deposited on a freshly etched silicon surface. The areas which act as working electrodes (microdiscs) were defined photolithographically. Finally, the insulating resist mask layer was hardbaked to improve its adhesion. The behavior of a bare electrode array was characterized by cyclic voltammetry at the scan rate of 50 mV/s in 0.1 mol/l KNO₃ (acidified by HNO₃ to pH 2.5). Differential pulse anodic stripping voltammetry was used for determination of Pb²⁺ ions in solution on bare CNx and polycrystalline electrodes. A preconcentration step was carried out at −700 mV vs. Ag/AgCl/Cl⁻ for 180 s. Lead ions were co-deposited with Hg²⁺ ions (1 × 10⁻³ mol/l) in 0.1 mol/l KNO₃ (pH 2.5). All the experimental solutions were prepared from chemicals of analytical grade in 18 × 10⁴ Ωm water.

CNx microelectrode array for in-situ determination of Pb²⁺ ions was prepared by co-deposition of Pb²⁺ at −700 mV vs. Ag/AgCl/Cl⁻ in 0.1 mol/l KNO₃ with different content of Pb²⁺ for 180 s preconcentration time under constant magnetic striping. Stripping currents at −566 mV vs. Ag/AgCl/Cl⁻ from reoxidation of Pb (Pb⁰ → Pb²⁺ + 2e⁻) into the same solution were recorded by DP ASV. The minimum measurable current response to Pb²⁺ concentration on PCD was 1.10⁻⁴ mol/l, and on CNx 5 × 10⁻⁶ mol/l.

The homogeneity of CNx layer was very high, it can be seen in Fig 5. DLC is in a dense, metastable form of amorphous carbon a–C, or hydrogenated amorphous carbon a–C:H containing a significant number of sp³ bonds. In other words, DLC consists of amorphous carbon and disordered graphite, and its atomic structure contains mixture of sp² and sp³ sites. The π state of sp² controls the electrical properties and sp³ controls the mechanical properties. Experimentally obtained CNx films are composed of two phases that have sp³ and sp² bonded structure [8]. Therefore, the concentration of nitrogen in the structure of CNx, coatings is a key factor affecting the coating characteristics, up to now not yet studied.

4. Conclusion

We have demonstrated a very wide range of applications for polycrystalline diamond prepared by the unique HF CVD processing method and for CNx layer prepared by a pulse arc system.

In an electronic application such as microelectrode array for measurement of heavy metals in water there is a possibility to use an arc system for CNx deposition on a large area substrate with high homogeneity at very low temperature of about 150 °C opposite to 700 °C in HF CVD.

Low deposition temperature opens a new window for many cheaper applications in medicine such as cover layers of hip replacement joints, nails, or cover microscopic glass for research of DNA or any biological tissue growth.

5. Acknowledgement

The presented work was supported by the Scientific Grant Agency of the Ministry of Education of the Slovak Republic and Slovak Grant Agency, No. 1/2061/05, 1/0170/03 and by Science and Technology Assistance Agency under the contract, No. APVT-20-034404 and APVT-20050702/04.
References


