

## Adhesion, Growth and Differentiation of Osteoblasts on Surface-Modified Materials Developed for Bone Implants

M. VANDROVCOVÁ<sup>1</sup>, L. BAČÁKOVÁ<sup>1</sup>

<sup>1</sup>Department of Growth and Differentiation of Cell Populations, Institute of Physiology, Academy of Sciences of the Czech Republic, Prague, Czech Republic

Received June 10, 2010

Accepted September 17, 2010

On-line March 14, 2011

---

### Summary

This review briefly outlines the history and possibilities of bone reconstruction using various types of artificial materials, which allow interaction with cells only on the surface of the implant or enable ingrowth of cells inside the material. Information is also provided on the most important properties of bone cells taking part in bone tissue development, and on diseases and regeneration. The most common cell types used for testing cell-material interaction *in vitro* are listed, and the most commonly used approaches to this testing are also mentioned. A considerable part of this review is dedicated to the physical and chemical properties of the material surface, which are decisive for the cell-material interaction, and also to modifications to the surface of the material aimed at integrating it better with the surrounding bone tissue. Special attention is paid to the effects of nanoscale and microscale surface roughness on cell behaviour, to material surface patterning, which allows regionally-selective adhesion and growth of cells, and also to the surface chemistry. In addition, coating the materials with bioactive layers is examined, particularly those created by deposition of fullerenes, hybrid metal-fullerene composites, carbon nanotubes, nanocrystalline diamond films, diamond-like carbon, and nanocomposite hydrocarbon plasma polymer films enriched with metals.

### Key words

Surface roughness and topography • Surface wettability • Surface coating • Surface patterning • Osteogenic cells • Bone tissue engineering

### Corresponding author

L. Bačáková, Dept. of Growth and Differentiation of Cell Populations, Institute of Physiology, Academy of Sciences of the Czech Republic, Videnska 1083, 142 20 Prague 4 – Krc, Czech Republic. Fax: +420 2 4106 2488. E-mail: lucy@biomed.cas.cz

### Introduction

The lifestyle and attitudes of people nowadays are responsible for the more and more frequent manifestation of so-called “diseases of civilization”. The most common diseases include heart and vascular diseases, cancer, obesity, diabetes, inflammatory rheumatic joint diseases, premature birth or miscarriage, depression, etc. The causes of these diseases are various and can be combined. They comprise mainly the intake of high-calorie foods, high consumption of animal products, decreased physical exertion, excessive alcohol drinking and cigarette smoking, and mental stress. Humans usually have difficulty in adapting to the excessive amount of conveniences brought to us by civilization, in the best sense of this word. The food industry produces fatty, salty meals and sugary drinks. The development of transport has caused a decline in physical movement, and also contributes to injuries due to accidents. The lifestyle of “modern” humans paradoxically reduces their quality of life. Nevertheless, the human life span is constantly lengthening, and thus there is a need to extend people’s productive life. Modern medicine has come on to the scene, and has been developing new medicaments and new therapeutic procedures, including the creation of

tissue and organ replacements by tissue engineering methods. At the same time, however, disease prevention remains of great importance.

This review concentrates on ways to replace irreversibly damaged bone and joint tissue, especially by surface-modified materials that are highly attractive for adhesion, growth and osteogenic differentiation of cells, and thus supportive for integration of the implant with the bone tissue and its secondary stability. Prior to the construction of such implants, it is necessary to know about the physiology and pathophysiology of the bone tissue in health and disease.

Natural bone is composed of several types of cells of mesenchymal origin, such as osteoblasts, osteoclasts, stem cells and vascular cells. Osteoblasts are responsible for bone growth, while osteoclasts are responsible for bone resorption. Bone is resorbed and replaced in a physiological process referred to as bone remodelling (Barrett *et al.* 2010). Osteoblasts are very similar to fibroblasts; all the genes expressed in fibroblasts are also expressed in osteoblasts. The only difference is in the expression of two osteoblast-specific transcripts: one encoding *Cbfa1*, and the other encoding osteocalcin. *Cbfa1* is an osteoblast-specific transcription factor, considered as the earliest and most specific marker of osteogenesis. Osteocalcin is the most abundant non-collagenous protein of the bone matrix. This molecule inhibits bone growth by inhibiting the activity of transglutaminase, binds calcium and is also an important marker of osteoblast differentiation (Kaartinen *et al.* 1997, Ducy *et al.* 2000). During embryonic development, osteoblast differentiation can occur through two distinct pathways, i.e. intramembranous and endochondral ossification. Intramembranous ossification mainly occurs during formation of the flat bones of the skull. In this case, the mesenchymal progenitor cells differentiate directly into osteoblasts. Endochondral ossification, on the other hand, occurs in long bones, such as limbs. Long bones are formed from cartilage. Upon vascular invasion into the cartilage template, the chondrocytes die through apoptosis, are replaced by osteoblasts, and the template is transformed into bone by the ossification process (Barrett *et al.* 2010).

In healthy bone, there is a delicate balance between the synthetic activity of osteoblasts and the resorptive activity of osteoclasts. For example, during ageing and in some kinds of diseases, the balance shifts in favour of osteoclasts, which can lead to increased bone resorption. This process affects the bone structure, and

makes the bone brittle and susceptible to fractures. One of the most common bone diseases is osteoporosis. It is mostly caused by a decrease in estrogen production in women during menopause (Rodan and Martin 2000). Estrogen loss is associated with elevated bone resorption caused by a rise in osteoclast number, which is driven by an increase in the cytokine production that regulates osteoclast generation by the following cascade: receptor for the activator of nuclear factor- $\kappa$ B (RANK) ligand; tumor necrosis factor- $\alpha$  (TNF- $\alpha$ ); interleukin-1, interleukin-6, interleukin-11; macrophage-colony stimulating factor (M-CSF); and prostaglandin E (Pacifici 1998).

Degeneration of the articular cartilage (by aging, wear, etc.), i.e. primary osteoarthritis, can be considered as the main cause of joint diseases. However, secondary osteoarthritis (caused e.g. by trauma or by disability) is also an important factor. Both primary and secondary osteoarthritis are manifested by pain during walking, which can be partially suppressed by palliative remedies, e.g. painkillers. Movement itself, however, may be a problem, even an impossibility. The only way to restore mobility is often by implanting an artificial joint (Sedlák and Piška 2008). The worldwide investment in bone implants was about 23 billion dollars in 2005, and this figure has been increasing year by year. Metal implants form the main category, though these materials are less suitable for bone tissue engineering.

## Tissue engineering and artificial materials

Tissue engineering has been defined as “an interdisciplinary field that applies the principles of engineering and life sciences toward the development of biological substitutes that restore, maintain, or improve tissue function or a whole organ” (Langer and Vacanti 1993). In order to aid tissue repair or regeneration, tissue engineering can use purely biological building blocks, e.g. living cells and extracellular matrix (ECM) molecules, or cells on carriers made of artificial materials. Thus, interactions between biomaterials and cells play an important role in this field. It has been repeatedly shown that cell behaviour is strongly dependent on the physical and chemical properties of the surface of the material (Linez-Bataillon *et al.* 2002, Eisenbarth *et al.* 2002, Webster and Smith 2005).

Artificial materials have been used for hundreds of years. The Mayans, for example, created teeth from sea shells in ~600 A.D. Similarly, an iron dental implant was

found in Europe in a corpse dated from 200 A.D. However, biomaterials as we think of them today did not exist just 50 years ago. There were no manufacturers of medical devices (except glass eyes, dental devices, some external prostheses, etc.), no understanding of biocompatibility, no legal regulations, etc. (Ratner *et al.* 2004).

Materials used nowadays can be divided into several basic groups: synthetic polymers, natural polymers, ceramics, metals, and composites which combine together the properties of two or more materials of different classes.

Each of the above-mentioned groups of materials can be used in bone tissue engineering. Generally, synthetic polymers are used for fabricating some orthopaedic and dental devices, and also for artificial tissue replacements. Poly(methyl methacrylate) is, for example, a main component of bone cement for orthopaedic surgery. Poly(dimethyl siloxane), well-known from its application in constructing artificial heart valves, is used in finger joint prostheses. Medical fibres, made e.g., of polytetrafluoroethylene (PTFE) or poly(lactide-co-glycolide) (PLGA), can be used as ligament and tendon substitutes (e.g. to fix separated shoulder joints or to repair knee joints). However, the application of polymers is limited due to relatively frequent problems with abrasion and wear, poor bone attachment, excessively high flexibility or low hardness. High hardness is characteristic for ceramic materials, but these materials are usually excessively brittle. Ceramic materials, e.g. hydroxyapatite (HAp) or tricalciumphosphate, can be used advantageously in combination with polymers, particularly in the form of bioactive and biodegradable micro- and nanoparticles (for a review, see Abramson *et al.* 2004, Vagaská *et al.* 2010). Biodegradable materials are important for temporary mechanically supportive healing appliances, such as sutures and bone fixation devices. These materials also have great potential for bone tissue engineering, because temporary biodegradable scaffolds are expected gradually to be replaced by newly formed, fully regenerated and functional bone tissue. Natural polymers are similar to and often identical with native macromolecules. These molecules are non-cytotoxic, because they are of the body origin, and are also better available for cells, e.g. for binding with cell adhesion receptors. However, natural polymers are frequently immunogenic, and are also associated with a risk of pathogen transmission. Moreover, their structural conformation is more complex than in synthetic molecules, which can lead to worse technological manipulation. One of the most interesting molecules for

bone tissue engineering is collagen. Collagen is a major protein of the ECM of various tissues; especially collagen I is predominant in the bone, tendon and skin. It has been found that, in combination with chondroitin 6-sulfate, collagen modifies the kinetics and the mechanism for healing full-thickness skin wounds in rodents. A positive influence of this combination on the bone tissue is proposed.

A final group of materials – metals – are the dominant materials used in present-day orthopaedics (for a review, see Abramson *et al.* 2004). Materials that are currently widely used for surgical implants include 316L stainless steel, cobalt-chromium alloys, and titanium and its alloys. Amongst the metallic materials available for implant applications, titanium is considered to be the best for bone implants, and is extensively used in biomedical applications. There are several reasons for choosing titanium and its alloys, namely mechanical properties, chemical stability, and last but not least the well-known biocompatibility of Ti. In comparison with other alloys, titanium-based materials have a low modulus of elasticity, varying from 110 to 55GPa, and approach the elasticity modulus of natural bone (30GPa). In comparison, 316L stainless steel and chromium-cobalt alloys have a much higher modulus of elasticity (210GPa, 240GPa, respectively), and this can lead to prosthesis failure through loosening or fracture. Thus, the modulus of elasticity of an artificial bone implant should be as similar as possible to that of the bone (Geetha *et al.* 2009). As for the chemical properties of metallic implants, local adverse tissue reactions or allergic reactions caused by these implants originate from the release of metal ions from the implant (Long and Rack 1998). Titanium is a reactive metal. This means that in air, water or any other electrolyte, an oxide is spontaneously formed on the surface of the metal. This oxide is about 4 nm in thickness. The specific mode of oxide growth on titanium has the positive effect that no metal ion will reach the surface. Titanium oxide is a good insulator and can occlude anionic impurities such as chlorine and fluorine of phosphates. By contrast, conductible materials imply redox processes which lead to denaturation of macromolecules (Steinmann 1998).

### **Cell models and culture conditions for studying cell-material interaction**

Materials designed for bone implantation are tested *in vitro* on cell cultures and then *in vivo* on

laboratory animals. Material tests under *in vitro* conditions usually start with the use of cell lines. These lines represent homogeneous, well-defined, relatively easily available and cultivable cell populations, which enable screening of multiple material samples and are capable of giving reproducible results. Animal cell lines often used for biomaterial testing include osteoblast-like UMR-106 cells, mouse bone marrow MBA-15 cells and mouse progenitor MC3T3-E1 cells. The last-mentioned are cells originally with a fibroblast-like phenotype, capable of differentiating into osteoblasts in appropriate culture environments (Kanazawa *et al.* 2007). Human bone-derived cell lines include osteoblast-like cells CPC-2, TE-85, MG-63, SaOS-2, U-2OS. Although these cell lines were derived from osteosarcoma, they generally retained the most important markers of osteogenic cell differentiation, such as the activity of alkaline phosphatase and production of osteocalcin (Zhao *et al.* 2007, Rudnik *et al.* 2008, Grausová *et al.* 2009a, Kalbáčová *et al.* 2009).

Selected interesting and promising results obtained on cell lines are then usually verified on primocultured and low-passaged cells derived from animal and human bones. These cells included differentiated osteoblasts, such as osteoblasts from neonatal rat calvaria (Webster *et al.* 2000a,b, de Oliveira and Nanci 2004, de Oliveira *et al.* 2007), human osteoblasts (hOB) obtained from surgical bone specimens (Anselme and Bigerelle 2005), and also non-differentiated stem cells, such as mesenchymal stem cells derived from the bone marrow (MSCs; Filová *et al.* 2009a) or embryonic stem cells (Bedi *et al.* 2009). Primocultured and low-passaged animal and human cells can be purchased from specialized companies (Webster and Ejiófor 2004), which makes their availability and use in biomaterial testing easier.

For cell cultivation on the tested materials, classical static and more advanced dynamic cell culture systems can be used. In this system, the material samples are inserted into cultivation vessels (usually dishes). They are immersed in the cell culture medium and seeded with cells suspended in the medium. For cell seeding, a drop in the cell suspension in the culture medium can also be applied selectively to the surface of the material. The classical static system is suitable mainly for cells on planar ("two-dimensional") material samples, though these cells can also be exposed to dynamic conditions, such as fluid shear stress generated by a flow of cell culture media (Scaglione *et al.* 2008, Kokkinos *et al.*

2009, Tan *et al.* 2010), hydraulic pressure (Gardinier *et al.* 2009) or stretch strain if the cells are cultured on elastic deformable materials, such as silicone rubber (Kaspar *et al.* 2000). These types of mechanical stimulation of osteoblasts or other osteogenic cells (e.g. bone marrow stromal cells, osteoprogenitor MC3T3-E1 cells, human osteoblast-like MG 63 cells) led to reorganization of the focal adhesion plaques and cytoskeleton, increased cell stiffness and particularly led to more pronounced osteogenic cell differentiation, manifested by a higher expression of the main osteogenic transcription factors Cbfa1 and Osterix, extracellular matrix molecules collagen I, osteopontin, osteocalcin and bone sialoprotein, and higher alkaline phosphatase activity.

Dynamic cultivation is essential for cells grown on three-dimensional polymeric, ceramic or composite porous or fibrous scaffolds that are often used in bone tissue engineering. Dynamic cultivation, especially if coupled with dynamic cell seeding, facilitates the penetration of cells into the pores of the material and colonization of the deep parts of the scaffolds with cells. This is almost impossible in the conventional static cell culture system. In addition, dynamic systems usually improve the perfusion of the scaffolds with cell culture media, and thus they improve the delivery of oxygen and nutrients to the cells and the removal of waste products of the cell metabolism and of degradation of the scaffold (for a review, see Pamula *et al.* 2008, 2009). The most frequently used dynamic systems are perfusion bioreactors (Olivier *et al.* 2007), horizontally or vertically rotating bioreactors (Belfiore *et al.* 2009, Meretoja *et al.* 2009) and bioreactors generating pressure (Mauney *et al.* 2004). Various types of mechanical stimulation can also be combined in a single bioreactor, for example cyclic hydraulic pressure and fluid shear stress (Gardinier *et al.* 2009), centrifugal-force-induced fluid pressure in a "rotating-cup" bioreactor and compression pressure (Belfiore *et al.* 2009) or perfusion-generated fluid shear stress and compression stress (Bölgen *et al.* 2008). Similarly as on planar surfaces, the cells in 3D materials cultured under dynamic conditions also exhibited enhanced proliferation, viability and osteogenic differentiation in comparison with cells in the conventional static cell culture system (Mauney *et al.* 2004, Olivier *et al.* 2007, Meretoja *et al.* 2009). Osteogenic cell differentiation can also be significantly enhanced by the use of special osteogenic media, i.e. supplemented with dexamethasone, ascorbic acid,

vitamin D<sub>3</sub> and beta-glycerolphosphate (Mauney *et al.* 2004, Kokkinos *et al.* 2009).

### Physicochemical properties of the material surface

When constructing artificial materials, it is desirable to create materials promoting the attachment, migration, proliferation, differentiation, long-term viability and proper functioning of the cells. The adhesion of cells to artificial materials and their further performance are mediated by ECM molecules, such as vitronectin, fibronectin, collagen, laminin and also fibrin, a molecule of provisional ECM taking part in wound healing and often used for modifying the surface of a biomaterial (Brynda *et al.* 2005, Filová *et al.* 2009b). These molecules are spontaneously adsorbed on the surface from the biological fluids, such as cell culture media, blood or intercellular fluid, and specific active sites on these molecules are then recognized and bound by adhesion receptors on cells, mainly integrins. Thus, cell adhesion-mediating ECM molecules have to be adsorbed in an appropriate amount, spectrum, and particularly spatial conformation and flexibility in order to be accessible for cell adhesion receptors. The adsorption behaviour of proteins is markedly influenced by the physical and chemical properties of the material surface, such as its polarity, wettability, electrical charge and conductivity, roughness and topography, compliance and others (Engler *et al.* 2004, Bačáková *et al.* 2004, Bačáková and Švorčík 2008). Modification of the tissue response by the roughness and topography of the material is one of the most important criteria for the production of biomaterials, because these criteria are decisive for the cell-material interaction and for integration of the material with the surrounding tissue (Clark 1994, Ito 1999, He *et al.* 2008).

#### Surface roughness

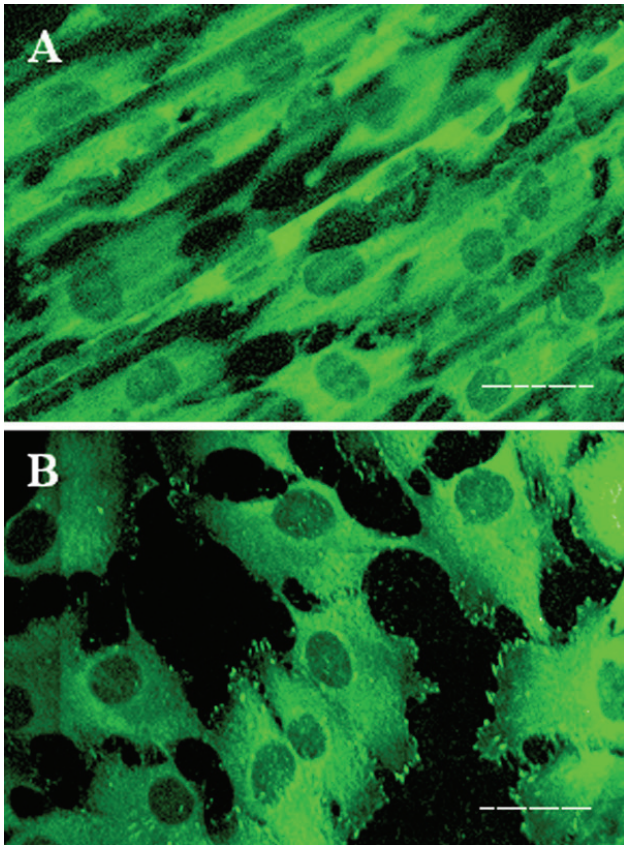
Intensive research is being carried out on modifications of the material surface aimed at enhancing the attractiveness of the material for cells and thus optimizing the integration of the material with the surrounding tissue. One of these modifications is induction of the nanoscale roughness and topography of the material surface. Nanostructured surfaces, i.e. those containing irregularities less than 100 nm, are believed to mimic the nanoarchitecture of natural tissues, e.g. the size of some ECM molecules or irregularities on these

molecules, such as folding, branching, etc., and also the size of the extracellular parts of cell adhesion receptors. On nanostructured surfaces, cell adhesion-mediating ECM molecules are adsorbed in advantageous, almost physiological geometrical conformations, and thus the specific bioactive sites in these molecules, such as amino acid sequences like RGD or osteoblast-binding sequence KRSR, are well accessible to cell adhesion receptors. In addition, among all ECM proteins, nanostructured surfaces preferentially adsorb vitronectin (due to its relatively small, linear and non-complicated molecule), and this protein is preferentially recognized by osteoblasts over other cell types (Webster *et al.* 2000a,b, Price *et al.* 2004).

A wide range of studies have compared the effects of micro- and nanoscale material surface topography on various types of cells (Lincks *et al.* 1998, Bačáková *et al.* 2001, Zhao *et al.* 2007, Khang *et al.* 2008, Liu *et al.* 2008, Mendonça *et al.* 2008). Most of these researchers consider that nanoscale topography provides greater support for the proliferation of bone cells. This result has been explained by the advantageous interaction between nanosize irregularities on the material surface and adsorbed cell adhesion-mediating molecules, as mentioned above, and thus improved osteoblast adhesion and spreading, which is a prerequisite for their good subsequent proliferation activity (Webster *et al.* 2000a,b, Christenson *et al.* 2007, Khang *et al.* 2008, Liu *et al.* 2008). Microscale irregularities can hamper cell spreading (Fig. 1), and thus they may slow down cell proliferation (Bačáková *et al.* 2001, Starý *et al.* 2003a,b). In accordance with this, Rosa and Beloti (2003), who compared the influence of various submicron- and microscale roughnesses of titanium substrates on human bone marrow cell growth, found that cell adhesion and proliferation decreased with increasing material surface roughness. In this study, surface roughness was measured by the  $R_a$  parameter, defined as the departure of the roughness profile from the mean line, which was 0.24  $\mu\text{m}$ , 0.69  $\mu\text{m}$ , 0.80  $\mu\text{m}$  and 1.90  $\mu\text{m}$ . Similarly, human osteoblasts in cultures on titanium substrates of the mean roughness amplitude (parameter  $S_a$ ) from 0.53  $\mu\text{m}$  to 2.52  $\mu\text{m}$  spread more intimately on surfaces with low roughness amplitude than on rougher surfaces (Anselme and Biggerelle 2005). On the other hand, cells on rougher Ti samples ( $R_a$  0.80  $\mu\text{m}$  and 1.90  $\mu\text{m}$ ) contained higher amounts of total protein and showed higher activity of alkaline phosphatase, i.e. an enzyme participating in bone tissue mineralization and an



important marker of osteogenic cell differentiation (Rosa and Beloti 2003). Therefore, microscale roughness can support osteogenic cell differentiation, although the initial cell responses, monitored by cell adhesion and proliferation, were not ideal.



**Fig. 1.** Immunofluorescence of vinculin, an integrin-associated focal adhesion protein, in human osteoblast-like MG 63 cells in 4-day-old cultures on carbon fibre-reinforced carbon composites (CFRC) with an unmodified surface (**A**), and with the surface ground using metallographic paper, coated with pyrolytic graphite, again ground using metallographic paper and polished with diamond 3/2 (**B**). **A:** Less spread cells on pristine composites of microscale surface roughness ( $R_a = 6.5 \mu\text{m}$ ,  $S = 42 \mu\text{m}$ ), elongated along the carbon fibres bulging on the material surface, and without visible vinculin-containing focal adhesion plaques. **B:** Well-spread polygonal cells with clearly developed vinculin-containing focal adhesion plaques in cultures on modified composites with submicron-scale surface roughness ( $R_a = 0.67 \mu\text{m}$ ,  $S = 81 \mu\text{m}$ ). Bio-Rad MRC600 confocal laser scanning microscope, oil immersion 60 $\times$  objective, (numerical aperture = 1.4), excitation wavelength 488 nm. Scale bar = 50  $\mu\text{m}$ .  $R_a$ : departures of the roughness profile from the mean line (i.e., size of the irregularities),  $S$ : mean spacing of the adjacent local peaks (i.e., distances among the irregularities).

Similar cell behaviour was observed on surfaces with various nanoscale roughnesses. In a study by Webster *et al.* (1999), performed on primary rat calvarial osteoblasts in cultures on nanophase titania (grain sizes

from 20 to 56 nm) and alumina (grain sizes from 20 to 67 nm), the highest numbers of initially adhering cells were found on  $\text{TiO}_2$  with grain size of 20-32 nm and  $\text{Al}_2\text{O}_3$  with grains of 20-49 nm. These numbers were significantly higher than on conventional titania and alumina, which were markedly rougher (grain sizes of 2120 nm and 177 nm, respectively). These favourable results on nanostructured surfaces were explained by the relatively large material surface area that was available for cell adhesion on ceramics with optimal surface nanoroughness, and also the improved adsorption, configuration and bioactivity of proteins that mediate osteoblast adhesion (Webster *et al.* 1999).

In our studies performed on human osteoblast-like MG 63 cells in cultures on  $\text{TiO}_2$  or glass, nanoroughness of  $R_a=40$  nm (i.e., a value close to the optimum surface roughness suggested by Webster *et al.* 1999) also induced a larger cell spreading area and significantly higher cell numbers on day 7 after seeding than was induced by higher surface roughness of 100 and 170 nm (Vandrovcová *et al.* 2010). Similar cell behaviour was observed on nanocrystalline diamond films (NCD). Among these films of root mean square (RMS) roughness of 20, 270 and 500 nm, surfaces with RMS of 20 nm provided the best support for the initial adhesion of human osteoblast-like SaOS cells (measured by cell number 1 hour after seeding), for metabolic activity of these cells (measured by the activity of cellular dehydrogenases 48 hours after seeding), and also for their osteogenic differentiation (measured by alkaline phosphatase activity and mineral deposition by cells 11 days after seeding). These beneficial effects of a nano-rough coating with RMS of 20 nm were explained by its similarity to the topography of a real bone surface (Kalbáčová *et al.* 2009).

The supportive effect of nano-rough surfaces on osteogenic cell differentiation was also demonstrated when flat and nano-rough surfaces were compared. Although the osteoblast colony occupancy (i.e., the total surface area occupied by each cell colony) was larger on conventional flat borosilicate glass surfaces than on nanophase ceramics (represented by alumina, titania and HAp with grain sizes of 24-67 nm), the synthesis of alkaline phosphatase and the deposition of calcium-containing mineral were significantly higher in osteoblasts on nanophase ceramics after 21 and 28 days of cultivation (Webster *et al.* 2000b).

Similarly, de Oliveira and Nanci (2004) compared the osteogenic differentiation of neonatal rat

calvarial osteoblasts in cultures on conventional unmodified Ti and TiAlV discs and discs with nanoscale irregularities induced chemically by etching in a solution of  $\text{H}_2\text{SO}_4$  and  $\text{H}_2\text{O}_2$ . Osteoblasts grown on the nanotextured surfaces, which contained nanopits with a honeycomb-like appearance and were in the 10 nm range in diameter, increased their secretion of bone sialoprotein and osteopontin, i.e. non-collagenous proteins typical for the bone matrix. The activity of alkaline phosphatase and the formation of mineralized bone-like nodules were also higher in cells on etched surfaces than on non-etched surfaces (de Oliveira *et al.* 2007).

Therefore, it can be concluded that surfaces of roughness in tens of nanometers are preferred by bone cells for adhesion, growth, differentiation and phenotypic maturation, rather than flat surfaces and surfaces with submicron- or micro-scale roughness.

#### *Surface topography and patterning*

Not only the size of the surface irregularities, but also their shape and distribution on the surface of the material are important for the cell-material interaction. For example, structures with different diameter (0.5 and 2  $\mu\text{m}$ ) and  $R_a$  parameter (0.2 and 0.4  $\mu\text{m}$ ) but with the same porous shape, which were created on titanium surfaces by anodic oxidation, increased the number of initially attached cells to similar values and induced similar morphological features in cells such as formation of filopodia, induced similar shape and distribution of vinculin-containing focal adhesion plaques, and induced a similar spatial arrangement of the actin cytoskeleton (Zhu *et al.* 2004).

The density of the irregularities on the surface of the material was crucial for the cell-material interaction. For example, Rice *et al.* (2003) deposited white-sulphate latex particles (shaped as hemispherical protrusions 110 nm in height) on titanium oxide surfaces in densities covering 3 %, 19 %, 30 %, or 43 % of the material surface. Increasing densities of nanoparticles decreased the spreading and proliferation of rat calvarial osteoblasts but increased the expression of osteocalcin in these cells. Similar results were obtained in experiments by Kunzler *et al.* (2007) performed on poly(ethylene imine)-coated silicon wafers adsorbed with silica nanoparticles (diameter 73 nm) in a gradient ranging from 0 % to 21 % coverage of the material surface. Rat calvarial osteoblasts in cultures on these surfaces exhibited a significant decrease in spreading, proliferation and formation of actin cytoskeleton, which was explained by more

pronounced bending of the cell membrane, cell stress and reduced cell-material contact area. Thus, it can be concluded that increased density of surface irregularities had similar effects to increased size of these irregularities, as described above (paragraph 4.1.).

#### *Surface chemical state of the material*

Several studies have investigated whether surface roughness and topography, on the one hand, or surface chemistry, on the other, has a higher influence on cell behaviour on a material surface. Webster *et al.* (1999) stated that enhanced adhesion of rat calvarial osteoblasts in cultures on nanophase alumina and titania was independent of the chemistry of the material surface, and dependent only on the optimal surface topography of nanophase ceramics. This conclusion was also drawn in another study by Webster and Ejiófor (2004), performed on low-passaged human osteoblasts in cultures on nanophase metals (Ti, Ti6Al4V and CoCrMo alloys).

However, this conclusion was not confirmed in our experiments performed on MG 63 cells in cultures on  $\text{TiO}_2$  films (in anatase form) and on microscopic glass slides, where the cells spread and grew better on a  $\text{TiO}_2$  surface than on glass slides of the same roughness (Vandrovcová *et al.* 2010). Similar results were obtained in rat calvarial osteoblasts in primary cultures on silicon substrates coated with  $\text{TiO}_2$  in anatase, rutile and amorphous crystal phases (He *et al.* 2008). Although all three films were of similar roughness and topography, the cell spreading area, measured 8 hours after seeding, was largest on  $\text{TiO}_2$  in anatase form. On anatase films, the cells also reached the highest cell numbers 36 and 72 hours after seeding. In addition, on day 7 and 14 after seeding, they displayed the greatest activity of alkaline phosphatase, i.e. an enzyme participating in bone tissue mineralization, and thus an important marker of osteogenic cell differentiation (He *et al.* 2008). These beneficial effects of anatase on cell colonization and cell phenotypic maturation were attributed to the highest surface hydrophilia of anatase (water drop contact angle about  $60^\circ$ ) compared with rutile and amorphous  $\text{TiO}_2$  (contact angle  $\sim 90^\circ$ , He *et al.* 2008). This explanation was further supported by Sawase *et al.* (2008), who described further enlargement of cell spreading area *in vitro* and also enhanced bone apposition *in vivo*, when the hydrophilicity of anatase was increased by being irradiated with ultraviolet light prior to seeding the cell or implanting it into the rabbit tibia. Thus, the positive effect of the material surface nanostructure on cell performance

can be further enhanced by surface hydrophilia, which (in synergy with the nanostructure) induces the adsorption of cell adhesion-mediating ECM molecules in advantageous spatial conformations, i.e. with specific amino acid sequences, serving as ligands for cell adhesion receptors exposed to these receptors (Bačáková *et al.* 2004, Bačáková and Švorčík 2008).

#### *Novel bioactive films on the material surface*

The physical and chemical properties of the surface of a material can also be modified by deposition of other materials (atoms, molecules, particles) in the form of continuous or micropatterned layers. A wide range of organic and inorganic materials can be used for modifying the material surface, such as entire ECM molecules and ECM-derived adhesion oligopeptides (i.e., ligands for cell adhesion receptors), fibrin, ceramics represented mainly by hydroxyapatite or tricalcium phosphate (for a review, see Bačáková *et al.* 2007a, Bačáková and Švorčík 2008, Filová *et al.* 2009a, Vagaská *et al.* 2010). Recently, we carried out an intensive study of the influence of layers made of carbon nanoparticles, namely fullerenes, nanotubes, nanocrystalline diamond, on adhesion, growth, viability, metabolic activity and osteogenic cell differentiation, and we found that these layers are promising for surface modifications of bone implants (Bačáková *et al.* 2007b, Grausová *et al.*, 2008a,b, 2009a,b, Vandrovcová *et al.* 2008; for a review see Bačáková *et al.* 2008).

#### *Fullerenes*

Fullerenes are spherical molecules made of carbon atoms, which are of a hollow cage-like shape similar to clathrin-coated vesicles in cells. This suggested the idea of using them for drug and gene delivery, or even for constructing artificial cellular organelles (for a review, see Bačáková *et al.* 2008). These carbon allotropes were discovered by Kroto *et al.* (1985), and have been considered as promising for various biomedical applications. However, it should be pointed out that fullerenes can be cytotoxic. For example, after irradiation with visible or ultraviolet light, they convert oxygen molecules into highly reactive radicals, which can damage the cell membrane, various intracellular molecules, enzymes, and also DNA. This property of fullerenes can be utilized in photodynamic therapy against tumours, viruses and bacteria resistant to multiple drugs. On the other hand, fullerenes can also act as potent radical scavengers (for a review, see Bačáková *et al.* 2008). An interesting issue is

the promoting action of fullerenes C<sub>60</sub> on chondrogenesis, probably due to stimulatory effects of these compounds on proteoglycan synthesis (Tsuchiya *et al.* 1995). Some derivatives of fullerenes also have affinity to the bone tissue. For example, Gonzalez *et al.* (2002) found that some fullerene derivatives can affect bone tissue mineralization. These authors investigated the interaction between a bisphosphonate fullerene C<sub>60</sub>(OH)<sub>16</sub>AMBP, i.e. (4,4-bisphosphono-2-(polyhydroxyl-1,2-dihydro-1,2-methanfullerene(60)-61-carboxamido)butyric acid) and HAp, i.e., an important inorganic component of the bone tissue. C<sub>60</sub>(OH)<sub>16</sub>AMBP has strong binding sites for HAp, which was manifested by capturing HAp from a solution and by its lower availability for bone tissue mineralization. On the other hand, this new knowledge on bone-vectored compounds can be useful in bone radiation therapy, where the radionuclide-containing fullerene can be bound to the diseased site in the bone, enabling a low dose to be used to avoid damage to the non-diseased tissue.

Fullerenes can be deposited on the material surface as nanostructured layers of various thicknesses, which can be controlled by the temperature and time of deposition. These layers can also be deposited through a metallic mask in micropatterned forms, which are useful in applications where regionally-selective cell adhesion and directed growth is desirable (Fig. 2), e.g. in tissue engineering, microarrays for advanced proteomics and genomics, and constructing biosensors (Grausová *et al.* 2008a, 2009b, Vandrovcová *et al.* 2008). In addition, fullerene molecules can be combined with other materials, particularly atoms of Ti, Co or Ni, and form binary fullerene-metal composites, also promising for biomedical applications (Vandrovcová *et al.* 2008, Vacík *et al.* 2010; for a review, see Bačáková *et al.* 2008).

#### *Carbon nanotubes*

Carbon nanotubes are cylindrical molecules made of carbon atoms. They exist in two basic forms: single-walled nanotubes (SWNT; formed by one cylindrical graphene sheet) and multi-walled nanotubes (MWNT; containing two or more concentrically arranged graphene sheets) (Iijima 1991, Dresselhaus *et al.* 1996). Carbon nanotubes can also be cytotoxic under some circumstances. In the same manner as fullerenes, carbon nanotubes can cleave molecular oxygen into free radicals, which can induce oxidative stress, inflammation, changes in the structure of proteins such as enzymes, extracellular matrix molecules and cell membranes. They can also disrupt DNA (Cui *et al.* 2005, Davoren *et al.* 2007, Kisin



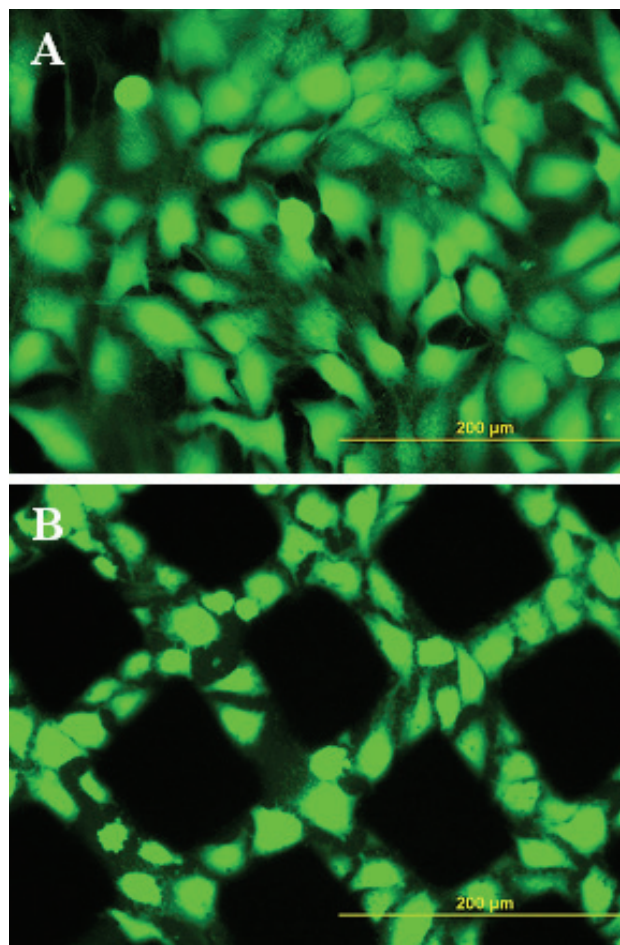
*et al.* 2007, Zhang *et al.* 2007). However, several studies have cast doubt on the cytotoxic action of nanotubes (Chen *et al.* 2006, Dumortier *et al.* 2006, Yehia *et al.* 2007, Zhu *et al.* 2006). These controversial results may be explained by the use of nanotubes of different size, purity, functionalization, water solubility and different tendency to form agglomerates (for a review, see Bačáková *et al.* 2008). Many studies have focused on applications of carbon nanotubes for tissue engineering in neural and also vascular systems, mainly as components of scaffolds for cell colonization and functioning (Matsumoto *et al.* 2007, Mattson *et al.* 2000, MacDonald *et al.* 2005). However, carbon nanotubes are also promising for bone tissue engineering. Abarategi *et al.* (2008), for example, tested MWNT on composites with chitosan loaded by human bone morphogenetic protein-2, and these authors found osteoinductive effects of nanotubes on mouse myoblast cell line C2C12. A positive effect of nanotubes in combination with HAp on spreading and phenotypic maturation of human osteoblasts has also been described (Balani *et al.* 2007).

#### Nanocrystalline diamond

Nanocrystalline diamond has become widely investigated for its excellent properties, such as low friction coefficient, chemical stability and particularly high biocompatibility (Tjong and Chen 2004, Amaral *et al.* 2008, Bačáková *et al.* 2007b, Grausová *et al.* 2008a,b, 2009a,b, Kovalchenko *et al.* 2011). Unlike the two groups of carbon allotropes mentioned above, nanocrystalline diamond is non-cytotoxic (Schrand *et al.* 2007, Aspenberg *et al.* 1996) and non-immunogenic (Tang *et al.* 1995, Nordsletten *et al.* 1996). These properties of nanocrystalline diamond enable its application not only in electronics and optics but also in biology and medicine. Due to their hardness, nanodiamond films proved to be suitable for coating the heads and cups of artificial joints (Papo *et al.* 2004). These films also supported adhesion, growth and differentiation of bone cells (Bajaj *et al.* 2007, Amaral *et al.* 2008). Thus, nanocrystalline diamond could be applied for coating the parts of joint prostheses or dental implants that anchor the bone with the surrounding bone tissue.

#### Diamond-like carbon

Another important carbon-based material is diamond-like carbon (DLC), also referred to as amorphous carbon. There are seven different forms of



**Fig. 2.** Human osteoblast-like MG 63 cells in 7-day-old cultures on fullerene  $C_{60}$  layers micropatterned with prominences  $128 \pm 8$  nm in height (**A**) or  $1043 \pm 57$  nm in height (**B**). **A**: homogeneous distribution of cells on the surfaces with relatively low prominences; **B**: preferential localization of cells in grooves among relatively high prominences. Cells stained with LIVE/DEAD viability/cytotoxicity kit (Invitrogen). Olympus IX 51 microscope, DP 70 digital camera, obj. 20x, bar = 200  $\mu$ m

amorphous carbon materials, differing in the content of hydrogen (hydrogen-free or hydrogenated with various hydrogen concentrations), hybridization ( $sp^2$  or  $sp^3$ ) or presence of additional non-carbon atoms, such as metals (W, Ti) or non-metallic elements (Si, O, N, F, B) (Fraunhofer Institute, Name Index of Carbon Coatings).

DLC displays some of the special properties of diamond, such as high hardness, chemical inertness, and good tribological characteristics. Further, DLC elicited no inflammatory response *in vitro* and induced no histopathological changes *in vivo* (Schroeder *et al.* 2000, Bruinink *et al.* 2005). Similarly as nanocrystalline diamond films, DLC can be deposited on orthopedic implants in order to prevent the release of metal ions (Dearnaley 1993, Dowling *et al.* 1997). Most DLC-based materials, particularly those containing only C and/or H

atoms, act as bioinert, i.e. not supporting cell adhesion, and thus they have been utilized for coating blood contacting devices, such as heart valves or coronary stents, in order to prevent thrombus formation (Grill 2003). However, in some applications, it is necessary to promote cell adhesion (e.g. for better integration of bone implants with the surrounding bone tissue). In such cases, DLC and related materials (e.g. hydrocarbon plasma polymers) have been rendered bioactive by adding titanium atoms (Bačáková *et al.* 2001, Grinevich *et al.* 2009).

## Conclusion and further remarks

A wide variety of materials have been used for constructing bone implants and replacements, such as synthetic and natural polymers, ceramics, metals and their composites. Bone replacements can be constructed in two forms: “two-dimensional”, i.e. interacting with cells only at the implant surface, and “three-dimensional”, allowing ingrowth of the bone tissue inside the implant, which is usually combined with degradability of the material and its gradual replacement by regenerated bone tissue. Nevertheless, for both forms of implants, the physicochemical properties of the material at the cell-material interface are decisive for the cell-material interaction. There are many ways to modify the surface of a material in order to enhance its attractiveness for cell colonization and osteogenic cell differentiation. Creating the nanostructure of the surface, especially in combination with surface hydrophilia, seems to be the most efficient of these approaches.

## References

- ABARRATEGI A, GUTIÉRREZ MC, MORENO-VICENTE C, HORTIGÜELA MJ, RAMOS V, LÓPEZ-LACOMBA JL, FERRER ML, DEL MONTE F: Multiwall carbon nanotube scaffolds for tissue engineering purposes. *Biomaterials* **29**: 94-102, 2008.
- ABRAMSON S, ALEXANDER H, BEST S, BOKROS JC, BRUNSKI JB, COLAS A, COOPER SL, CURTIS J, HAUBOLD A, HENSCH LL, HERGENROTHER RW, HOFFMAN AS, HUBBELL JE, JANSEN JA, KING MW, KOHN J, LAMBA NMK, LANGER R, MIGLIARESI C, MORE RB, PEPPAS NA, RATNER BD, VISSER SA, VON RECUM A, WEINBERG S, YANNAS IV: Classes of material used in medicine. In: *Biomaterials Science. An Introduction to Materials in Medicine*. BD RATNER, AS HOFFMAN, FJ SCHOEN, JE LEMONS (eds), Elsevier Academic Press, London, 2004, pp 67-137.
- AMARAL M, DIAS AG, GOMES PS, LOPES MA, SILVA RF, SANTOS JD, FERNANDES MH: Nanocrystalline diamond: In vitro biocompatibility assessment by MG 63 and human bone marrow cells cultures. *J Biomater Res A* **87**: 91-99, 2008.
- ANSELME K, BIGERELLE M: Topography effects of pure titanium substrates on human osteoblast long-term adhesion. *Acta Biomaterialia* **1**: 211-222, 2005.

The research in the field of bone tissue replacement and regeneration is endless. What usually happens is that new materials and new modifications to them are found, which are at least slightly better than the systems of the previous generation. Despite all the efforts of scientists, innovations are delayed by the occurrence of new complications, such as new diseases, allergies or problems related to long-term exposure of the organism to artificial tissue replacements. Everybody has to accept that death is inseparable from our destiny, and that it will occur in exactly 100 % of cases. Thus, any effort to avert death in all circumstances seems to be a completely unscientific activity, because death comes to all of us mortals (Komárek and Djakow 2010). However, biomaterial science and tissue engineering are capable of improving the quality of the life that remains between the onset of a disease and death. This is a desirable contribution, and is of great importance for an advanced human society.

## Conflict of Interest

There is no conflict of interest.

## Acknowledgements

This study was supported by the Academy of Sciences of the Czech Republic (grant No. KAN101120701 and KAN400480701) and by the Grant Agency of the Czech Republic (grant No. P108/10/1858 and 106/09/1000). Mr. Robin Healey (Czech Technical University, Prague) is gratefully acknowledged for his language revision of the manuscript.

- ASPENBERG P, ANTTILA A, KONTTINEN YT, LAPPALAINEN R, GOODMAN SB, NORDSLETTEN L, SANTAVIRTA S: Benign response to particles of diamond and SiC: bone chamber studies of new joint replacement coating materials in rabbits. *Biomaterials* **17**: 807-812, 1996.
- BAČÁKOVÁ L, STARÝ V, KOFROŇOVÁ O, LISÁ V: Polishing and coating carbon fiber-reinforced carbon composites with a carbon-titanium layer enhances adhesion and growth of osteoblast-like MG63 cells and vascular smooth muscle cells in vitro. *J Biomed Mater Res* **54**: 567-578, 2001.
- BAČÁKOVÁ L, FILOVÁ E, RYPÁČEK F, ŠVORČÍK V, STARÝ V: Cell adhesion on artificial materials for tissue engineering. *Physiol Res* **53** (Suppl 1): S35-S45, 2004.
- BAČÁKOVÁ L, FILOVÁ E, KUBIES D, MACHOVÁ L, PROKS V, MALINOVÁ V, LISÁ V, RYPÁČEK F: Adhesion and growth of vascular smooth muscle cells in cultures on bioactive RGD peptide-carrying polyactides. *J Mater Sci Mater Med* **18**: 1317-1323, 2007a.
- BAČÁKOVÁ L, GRAUSOVÁ L, VACÍK J, FRACZEK A, BLAZEWCZ S, KROMKA A, VANĚČEK M, ŠVORČÍK V: Improved adhesion and growth of human osteoblast-like MG63 cells on biomaterials modified with carbon nanoparticles. *Diamond Relat Mater* **16**: 2133-2140, 2007b.
- BAČÁKOVÁ L, ŠVORČÍK V: Cell colonization control by physical and chemical modification of materials. In: *Cell Growth Processes: New Research*. D. KIMURA (ed), Nova Science Publishers, New York, 2008, pp 5-56.
- BAČÁKOVÁ L, GRAUSOVÁ L, VANDROVCOVÁ M, VACÍK J, FRACZEK A, BLAZEWCZ S, KROMKA A, VANĚČEK M, NESLÁDEK M, KOPEČEK M: Carbon nanoparticles as substrates for cell adhesion and growth. In: *Nanoparticles: New Research*. F. COLUMBUS (ed), Nova Science Publishers, New York, 2008, pp 39-107.
- BAJAJ P, AKIN D, GUPTA A, SHERMAN D, SHI B, AUCIELLO O, BASHIR R: Ultrananocrystalline diamond film as an optimal cell interface for biomedical applications. *Biomed Microdevices* **9**: 787-794, 2007.
- BALANI K, ANDERSON R, LAHA T, ANDARA M, TERCERO J, CRUMPLER E, AGARWAL A: Plasma-sprayed carbon nanotube reinforced hydroxyapatite coatings and their interaction with human osteoblasts in vitro. *Biomaterials* **28**: 616-624, 2007.
- BARRETT KE, BARMAN SM, BOITANO S, BROOKS H: Hormonal control of calcium & phosphate metabolism & the physiology of bone. Chapter 23, *Ganong's Review of Medical Physiology*, Electronic access at <http://www.accessmedicine.com/content.aspx?aid=5244785>
- BEDI RS, BEVING DE, ZANELLO LP, YAN Y: Biocompatibility of corrosion-resistant zeolite coatings for titanium alloy biomedical implants. *Acta Biomater* **5**: 3265-3271, 2009.
- BELFIORE LA, BONANI W, LEONI M, BELFIORE CJ: Pressure-sensitive nutrient consumption via dynamic normal stress in rotational bioreactors. *Biophys Chem* **140**: 99-107, 2009.
- BÖLGEN N, YANG Y, KORKUSUZ P, GÜZEL E, EL HAJ AJ, PIŞKIN E: Three-dimensional ingrowth of bone cells within biodegradable cryogel scaffolds in bioreactors at different regimes. *Tissue Eng Part A* **14**: 1743-1750, 2008.
- BRUININK A, SCHROEDER A, FRAN CZ G, HAUERT R: In vitro studies on the effect of delaminated a-C:H film fragments on bone marrow cell cultures. *Biomaterials* **26**: 3487-3494, 2005.
- BRYNDA E, PACHERNÍK J, HOUSKA M, PIENKA Z, DVOŘÁK P: Surface immobilized protein multilayers for cell seeding. *Langmuir* **21**: 7877-7883, 2005.
- CHEN X, TAM UC, CZLAPINSKY JL, LEE GS, RABUKA D, ZETTL A, BERTTOZZI CR: Interfacing carbon nanotubes with living cells. *J Am Chem Soc* **128**: 6292-6293, 2006.
- CHRISTENSON EM, ANSETH KS, VAN DEN BEUCKEN JJP, CHAN CK, ERCAN B, JANSEN JA, LAURENCIN CT, LI W-J, MURUGAN R, NAIR LS, RAMAKRISHNA S, TUAN RS, WEBSTER TJ, MIKOS AG: Nanobiomaterial applications in orthopedics. *J Orthop Res* **25**: 11-22, 2007.
- CLARK P: Cell behaviour on micropatterned surfaces. *Biosensors and Bioelectronics* **9**: 657-661, 1994.
- CUI D, TIAN F, OZKAN CS, WANG M, GAO H: Effect of single wall carbon nanotubes on human HEK293 cells. *Toxicol Lett* **155**: 73-85, 2005.
- DAVOREN M, HERZOG E, CASEY A, COTTINEAU B, CHAMBERS G, BYRNE HJ, LYNG FM: In vitro toxicity evaluation of single walled carbon nanotubes on human A549 lung cells. *Toxicol In Vitro* **21**: 438-448, 2007.

- DEARNALEY G: Diamond-like carbon: a potential means of reducing wear in total joint replacements. *Clin Mater* **12**: 237-244, 1993.
- DE OLIVEIRA PT, NANJI A: Nanotexturing of titanium-based surfaces upregulates expression of bone sialoprotein and osteopontin by cultured osteogenic cells. *Biomaterials* **25**: 403-413, 2004.
- DE OLIVEIRA PT, ZALZAL SF, BELOTI MM, ROSA AL, NANJI A: Enhancement of in vitro osteogenesis on titanium by chemically produced nanotopography. *J Biomed Mater Res A* **80**: 554-564, 2007.
- DOWLING DP, KOLA PV, DONNELLY K, KELLY TC, BRUMITT K, LLOYD L, ELOY R, THERIN M, WEILL N: Evaluation of diamond-like carbon-coated orthopaedic implants. *Diamond Relat Mater* **6**: 390-393, 1997.
- DRESSELHAUS MS, DRESSELHAUS G, EKLUND PC: *Science of Fullerenes and Carbon Nanotubes*. Academic Press Inc., New York, 1996.
- DUCY P, SCHINKE T, KARSENTY G: The osteoblast: a sophisticated fibroblast under central surveillance. *Science* **289**: 1501-1504, 2000.
- DUMORTIER H, LACOTTE S, PASTORIN G, MAREGA R, WU W, BONIFAZI D, BRIAND JP, PRATO M, MULLER S, BIANCO A: Functionalized carbon nanotubes are non-toxic and preserve the functionality of primary immune cells. *Nano Lett* **6**: 1522-1528, 2006.
- EISENBARTH E, LINEZ P, BIEHL V, BREME J, HILDEBRAND HF: Cell orientation and cytoskeleton organisation on ground titanium surfaces. *Biomol Eng* **19**: 233-238, 2002.
- ENGLER A, BAČÁKOVÁ L, NEWMAN C, HATEGAN A, GRIFFIN M, DISCHER D: Substrate compliance versus ligand density in cell on gel responses. *Biophys J* **86**: 617-628, 2004.
- FILOVÁ E, BULLETT NA, BAČÁKOVÁ L, GRAUSOVÁ L, HAYCOCK JW, HLUČILOVÁ J, KLÍMA J, SHARD A: Regionally-selective cell colonization of micropatterned surfaces prepared by plasma polymerisation of acrylic acid and 1,7- octadiene. *Physiol Res* **58**: 669-684, 2009a.
- FILOVÁ E, BRYNDA E, RIEDEL T, BAČÁKOVÁ L, CHLUPÁČ J, LISÁ V, HOUSKA M, DYR JE: Vascular endothelial cells on two- and three-dimensional fibrin assemblies for biomaterial coatings. *J Biomed Mater Res A* **90**: 55-69, 2009b.
- FRAUNHOFER INSTITUTE: *Name Index of Carbon Coatings*. Electronic table at <http://www.ist.fraunhofer.de/english/c-products/tab/complete.html>
- GARDINIER JD, MAJUMDAR S, DUNCAN RL, WANG L: Cyclic hydraulic pressure and fluid flow differentially modulate cytoskeleton re-organization in MC3T3 osteoblasts. *Cell Mol Bioeng* **2**: 133-143, 2009.
- GEETHA M, SINGH AK, ASOKAMANI R: Ti based biomaterials, the ultimate choice for orthopaedic implants – a review. *Materials Science* **54**: 397-425, 2009.
- GONZALES KA, WILSON LJ, WU W, NANCOLLAS GH: Synthesis and in vitro characterization of tissue-selective fullerene: vectoring C60(OH)16MBP to mineralized bone. *Bioorg Med Chem* **10**: 1991-1997, 2002.
- GRAUSOVÁ L, VACÍK J, BÍLKOVÁ P, VORLÍČEK V, ŠVORČÍK V, SOUKUP D, BAČÁKOVÁ M, LISÁ V, BAČÁKOVÁ L: Regionally-selective adhesion and growth of human osteoblast-like MG 63 cells on micropatterned fullerene C60 layers. *J Optoelectron Adv Mater* **10**: 2071-2076, 2008a.
- GRAUSOVÁ L, KROMKA A, BAČÁKOVÁ L, POTOCKÝ S, VANĚČEK M, LISÁ V: Bone and vascular endothelial cells in cultures on nanocrystalline diamond films. *Diamond Relat Mater* **17**: 1405-1409, 2008b.
- GRAUSOVÁ L, BAČÁKOVÁ L, KROMKA A, VANĚČEK M, LISÁ V: Molecular markers of adhesion, maturation and immune activation of human osteoblast-like MG 63 cells on nanocrystalline diamond films. *Diamond Relat Mater* **18**: 258-263, 2009a.
- GRAUSOVÁ L, VACÍK J, VORLÍČEK V, ŠVORČÍK V, SLEPIČKA P, BÍLKOVÁ P, VANDROVCOVÁ M, LISÁ V, BAČÁKOVÁ L: Fullerene C60 films of continuous and micropatterned morphology as substrates for adhesion and growth of bone cells. *Diamond Relat Mater* **18**: 578-586, 2009b.
- GRILL A: Diamond-like carbon coatings as biocompatible materials - an overview. *Diamond Relat Mater* **12**: 166-170, 2003.
- GRINEVICH A, BAČÁKOVÁ L, CHOUKOUROV A, BOLDYRYEVA H, PIHOSH Y, SLAVINSKÁ D, NOSKOVÁ L, ŠKUCIOVÁ M, LISÁ V, BIEDERMAN H: Nanocomposite Ti/hydrocarbon plasma polymer films from reactive magnetron sputtering as growth supports for osteoblast-like and endothelial cells. *J Biomed Mater Res* **88A**: 952-966, 2009.

- HE J, ZHOU W, ZHOU X, ZHONG X, ZHANG X, WAN P, ZHU B, CHEN W: The anatase phase of nanotopography titania plays an important role on osteoblast cell morphology and proliferation. *J Mater Sci Mater Med* **19**: 3465-3472, 2008.
- IJIMA S: Helical microtubules of graphitic carbon. *Nature* **354**: 56-58, 1991.
- ITO Y: Surface micropatterning to regulate cell functions. *Biomaterials* **20**: 2333-2342, 1999.
- KAARTINEN MT, PIRHONEN A, LINNALA-KANKKUNEN A, MÄENPÄÄ PH: Transglutaminase-catalyzed cross-linking of osteopontin is inhibited by osteocalcin. *J Biol Chem* **272**: 22736-22741, 1997.
- KALBÁČOVÁ M, REZEK B, BAREŠOVÁ V, WOLF-BRANDSTETTER C, KROMKA A.: Nanoscale topography of nanocrystalline diamonds promotes differentiation of osteoblasts. *Acta Biomater* **5**: 3076-3085, 2009.
- KANAZAWA I, YAMAGUCHI T, YANO S, YAMAUCHI M, YAMAMOTO M, SUGIMOTO T: Adiponectin and AMP kinase activator stimulate proliferation, differentiation, and mineralization of osteoblastic MC3T3-E1 cells. *BMC Cell Biology* **8**: 51, 2007.
- KASPAR D, SEIDL W, NEIDLINGER-WILKE C, CLAES L: In vitro effects of dynamic strain on the proliferative and metabolic activity of human osteoblasts. *J Musculoskelet Neuronal Interact* **1**: 161-164, 2000.
- KHANG D, LU J, YAO CH, HABERSTROH KM, WEBSTER TJ: The role of nanometer and sub-micron surface features on vascular and bone cell adhesion on titanium. *Biomaterials* **29**: 970-983, 2008.
- KISIN ER, MURRAY AR, KEANE MJ, SHI XC, SCHWEGLER-BERRY D, GORELIK O, AREPALLI S, CASTRANOVA V, WALLACE WE, KAGAN VE, SHVEDOVA AA: Single-walled carbon nanotubes: genotoxic and cytotoxic effects in lung fibroblast V79 cells. *J Toxicol Environ Health A* **70**: 2071-2079, 2007.
- KOKKINOS PA, ZARKADIS IK, PANIDIS TT, DELIGIANNI DD: Estimation of hydrodynamic shear stresses developed on human osteoblasts cultured on Ti-6Al-4V and strained by four point bending. Effects of mechanical loading to specific gene expression. *J Mater Sci Mater Med* **20**: 655-665, 2009.
- KOMÁREK S, DJAKOW J: With Professor Stanislav Komárek not only about a scientific conference, medicine, death and dying of man. (in Czech) Electronic article at <http://www.lf2.cuni.cz/Informace/2010/komarek.htm>
- KOVALCHENKO AM, ELAM JW, ERDEMIR A, CARLISLE JA, AUCIELLO O, LIBERA JA, PELLIN MJ, GRUEN DM, HRYN JN: Development of ultrananocrystalline diamond (UNCD) coatings for multipurpose mechanical pump seals. *Wear* **270**: 325-331, 2011.
- KROTO HW, HEATH JR, O'BRIEN SC, CURL RF, SMALLEY RE: C60: Buckminsterfullerene. *Nature* **318**: 162-163, 1985.
- KUNZLER TP, HUWILER CH, DROBEK T, VÖRÖS J, SPENCER ND: Systematic study of osteoblast response to nanotopography by means of nanoparticle-density gradient. *Biomaterials* **28**: 5000-5006, 2007.
- LANGER R, VACANTI JP: Tissue engineering. *Science* **260**: 920-926, 1993.
- LINCKS J, BOYAN BD, BLANCHARD CR, LOHMANN CH, LIU Y, COCHRAN DL, DEAN DD, SCHWARTZ Z: Response of MG63 osteoblast-like cells to titanium and titanium alloy is dependent on surface roughness and composition. *Biomaterials* **19**: 2219-2232, 1998.
- LINEZ-BATAILLON P, MONCHAU F, BIGERELLE M, HILDEBRAND HF: In vitro MC3T3 osteoblast adhesion with respect to surface roughness of Ti6Al4V substrates. *Biomol Eng* **19**: 133-141, 2002.
- LIU H, YAZICI H, ERGUN C, WEBSTER TJ, BERMEK H: An in vitro evaluation of the Ca/P ratio for the cytocompatibility of nano-to-micron particulate calcium phosphates for bone regeneration. *Acta Biomaterialia* **4**: 1472-1479, 2008.
- LONG M, RACK HJ: Titanium alloys in total joint replacement – a materials science perspective. *Biomaterials* **19**: 1621-1639, 1998.
- MACDONALD RA, LAURENZI BF, VISWANATHAN G, AJAYAN PM, STEGEMANN JP: Collagen-carbon nanotube composite materials as scaffolds in tissue engineering. *J Biomed Mater Res* **74A**: 489-496, 2005.
- MATSUMOTO K, SATO C, NAKA Y, KITAZAWA A, WHITBY RLD, SHIMIZU N: Neurite outgrowths with neurotrophin-coated carbon nanotubes. *J Biosci Bioeng* **103**: 216-220, 2007.
- MATTSON MP, HADDON RC, RAO AM: Molecular functionalization of carbon nanotubes and use as substrates for neuronal growth. *J Mol Neurosci* **14**: 175-182, 2000.



- MAUNEY JR, SJOSTORM S, BLUMBERG J, HORAN R, O'LEARY JP, VUNJAK-NOVAKOVIC G, VOLLOCH V, KAPLAN DL: Mechanical stimulation promotes osteogenic differentiation of human bone marrow stromal cells on 3-D partially demineralized bone scaffolds in vitro. *Calcif Tissue Int* **74**: 458-468, 2004.
- MENDONÇA G, MENDONÇA DBS, ARAGÃO FJL, COOPER LF: Advancing dental implant surface technology – from micron- to nanotopography. *Biomaterials* **29**: 3822-3835, 2008.
- MERETOJA VV, MALIN M, SEPPÄLÄ JV, NÄRHI TO: Osteoblast response to continuous phase macroporous scaffolds under static and dynamic culture conditions. *J Biomed Mater Res A* **89**: 317-325, 2009.
- NORDSLETTEN L, HØGÅSEN AKM, KONTTINEN YT, SANTAVIRTA S, ASPENBERG P, AASEN AO: Human monocytes stimulation by particles of hydroxyapatite, silicon carbide and diamond: in vitro studies of new prosthesis coatings. *Biomaterials* **17**: 1521-1527, 1996.
- OLIVIER V, HIVART P, DESCAMPS M, HARDOUIN P: In vitro culture of large bone substitutes in a new bioreactor: importance of the flow direction. *Biomed Mater* **2**: 174-180, 2007.
- PACIFICI R: Cytokines, estrogen, and postmenopausal, osteoporosis – the second decade. *Endocrinology* **139**: 2659-2661, 1998.
- PAMULA E, BAČÁKOVÁ L, FILOVÁ E, BUCZYNSKA J, DOBRZYNSKI P, NOSKOVÁ L, GRAUSOVÁ L: The influence of pore size on colonization of poly(L-lactide-glycolide) scaffolds with human osteoblast-like MG 63 cells in vitro. *J Mater Sci Mater Med* **19**: 425-435, 2008.
- PAMULA E, FILOVÁ E, BAČÁKOVÁ L, LISÁ V, ADAMCZYK D: Resorbable polymeric scaffolds for bone tissue engineering: the influence of their microstructure on the growth of human osteoblast-like MG 63 cells. *J Biomed Mater Res A* **89**: 432-443, 2009.
- PAPO MJ, CATLEDGE SA, VOHRA YK, MACHADO C: Mechanical wear behavior of nanocrystalline and multilayer diamond coatings on temporomandibular joint implants. *J Mater Sci Mater Med* **15**: 773-777, 2004.
- PRICE RL, ELLISON K, HABERSTROH KM, WEBSTER TJ: Nanometer surface roughness increases select osteoblast adhesion on carbon nanofiber compacts. *J Biomed Mater Res A* **70**: 129-138, 2004.
- RATNER BD: A history of biomaterials. In: *Biomaterials Science. An Introduction to Materials in Medicine*. BD RATNER, AS HOFFMAN, FJ SCHOEN, JE LEMONS (eds), Elsevier Academic Press, London, 2004, pp 10-19.
- RICE JM, HUNT JA, GALLAGHER JA, HANARP P, SUTHERLAND DS, GOLD J: Quantitative assessment of the response of primary derived human osteoblast and macrophages to a range of nanotopography surfaces in a single culture model in vitro. *Biomaterials* **24**: 4799-4818, 2003.
- RODAN GA, MARTIN TJ: Therapeutic approaches to bone diseases. *Science* **289**: 1508-1514, 2000.
- ROSA AL, BELOTI MM: Effect of cpTi surface roughness on human bone marrow cell attachment, proliferation, and differentiation. *Braz Dent J* **14**: 16-21, 2003.
- RUDNIK V, SANYAL A, SYED FA, MONROE DG, SPELSBERG TC, OURSLER MJ, KHOSLA S: Loss of ERE binding activity by estrogen receptor-alpha alters basal and estrogen-stimulated bone-related gene expression by osteoblastic cells. *J Cell Biochem* **103**: 896-907, 2008.
- SAWASE T, JIMBO R, BABA K, SHIBATA Y, IKEDA T, ATSUTA M: Photo-induced hydrophilicity enhances initial cell behavior and early bone apposition. *Clin Oral Implants Res* **19**: 491-496, 2008.
- SCAGLIONE S, WENDT D, MIGGINO S, PAPADIMITROPOULOS A, FATO M, QUARTO R, MARTIN I: Effects of fluid flow and calcium phosphate coating on human bone marrow stromal cells cultured in a defined 2D model system. *J Biomed Mater Res A* **86**: 411-419, 2008.
- SCHRAND AM, HUANG H, CARLSON C, SCHLAGER JJ, OSAWA E, HUSSAIN SM, DAI L: Are diamond nanoparticles cytotoxic? *J Phys Chem B* **111**: 2-7, 2007.
- SCHROEDER A, FRAN CZ G, BRUININK A, HAUERT R, MAYER J, WINTERMANTEL E: Titanium containing amorphous hydrogenated carbon films (a-C:H/Ti): surface analysis and evaluation of cellular reactions using bone marrow cell cultures in vitro. *Biomaterials* **21**: 449-456, 2000.
- SEDLÁK J, PÍŠKA M: Production of implants. (in Czech) *MM Průmyslové spektrum* **3**: 74, 2008; electronic article at <http://www.mmspektrum.com/clanek/vyroba-implantatu>
- STARÝ V, BAČÁKOVÁ L, HORNÍK J, CHMELÍK V: Bio-compatibility of the surface layer of pyrolytic graphite. *Thin Solid Films* **433**: 191-198, 2003a.

- STARÝ V, GLOGAR P, BAČÁKOVÁ L, HNILICA F, CHMELÍK V, KOŘÍNEK Z, GREGOR J, MAREŠ V, LISÁ V: A study of surface properties of composite materials and their influence on the biocompatibility. *Acta Montana AB* **11**: 19-36, 2003b.
- STEINEMANN SG: Titanium – the material of choice? *Periodontology* **17**: 7-21, 1998.
- TAN L, MEYER T, PFAU B, HOFMANN T, TAN TW, JONES D: Rapid vinculin exchange dynamics at focal adhesions in primary osteoblasts following shear flow stimulation. *J Musculoskelet Neuronal Interact* **10**: 92-99, 2010.
- TANG L, TSAI C, GERBERICH WW, KRUCKEBERG L, KANIA DR: Biocompatibility of chemical-vapour-deposited diamond. *Biomaterials* **16**: 483-488, 1995.
- TJONG SS, CHEN H: Nanocrystalline materials and coatings. *Mater Sci Eng* **R45**: 1-88, 2004.
- TSUCHIYA T, YAMAKOSHI YN, MIYATA NA: A novel promoting action of fullerene C<sub>60</sub> on the chondrogenesis in rat embryonic limb bud cell culture system. *Biochem Biophys Res Commun* **206**: 885-894, 1995.
- VACÍK J, LAVRENTIEV V, NOVOTNÁ K, BAČÁKOVÁ L, LISÁ V, VORLÍČEK V, FAJGAR R: Fullerene (C<sub>60</sub>)–transitional metal (Ti) composites: Structural and biological properties of the thin films. *Diamond Relat Mater* **19**: 242-246, 2010.
- VAGASKÁ B, BAČÁKOVÁ L, FILOVÁ E, BALÍK K: Osteogenic cells on bio-inspired materials for bone tissue engineering. *Physiol Res* **59**: 309-322, 2010.
- VANDROVCOVÁ M, HANUŠ J, KYLIÁN O, BIEDERMAN H, LISÁ V, BAČÁKOVÁ L: Effect of different surface nanoroughness of TiO<sub>2</sub> films on the growth of MG63 cells. Abstracts 2<sup>nd</sup> International Conference on Cellular and Molecular Bioengineering (ICCMB2), August 2-4, 2010, Singapore.
- VANDROVCOVÁ M, VACÍK J, ŠVORČÍK V, SLEPIČKA P, KASÁLKOVÁ N, VORLÍČEK V, LAVRENTIEV V, VOSEČEK V, GRAUSOVÁ L, LISÁ V, BAČÁKOVÁ L: Fullerene C<sub>60</sub> and hybrid C<sub>60</sub>/Ti films as substrates for adhesion and growth of bone cells. *Phys Stat Sol (a)* **205**: 2252-2261, 2008.
- WARD BC, WEBSTER TJ: The effect of nanotopography on calcium and phosphorus deposition on metallic materials in vitro. *Biomaterials* **27**: 3064-3074, 2006.
- WEBSTER TJ, SIEGEL RW, BIZIOS R: Osteoblast adhesion on nanophase ceramics. *Biomaterials* **20**: 1221-1227, 1999.
- WEBSTER TJ, ERGUN C, DOREMUS RH, SIEGEL RW, BIZIOS R: Specific proteins mediate enhanced osteoblast adhesion on nanophase ceramics. *J Biomed Mater Res A* **51**: 475-483, 2000a.
- WEBSTER TJ, ERGUN C, DOREMUS RH, SIEGEL RW, BIZIOS R: Enhanced functions of osteoblasts on nanophase ceramics. *Biomaterials* **21**: 1803-1810, 2000b.
- WEBSTER TJ, EJIOFOR JU: Increased osteoblast adhesion on nanophase metals: Ti, Ti6Al4V, and CoCrMo. *Biomaterials* **25**: 4731-4739, 2004.
- WEBSTER TJ, SMITH TA: Increased osteoblast function on PLGA composites containing nanophase titania. *J Biomed Mater Res A* **74**: 677-686, 2005.
- YEHIA HN, DRAPER RK, MIKORYAK C, WALKER EK, BAJAJ P, MUSSELMAN IH, DAIGREPONT MC, DIECKMANN GR, PANTANO P: Single-walled carbon nanotube interactions with HeLa cells. *J Nanobiotechnol* **5**: 8-24, 2007.
- ZHANG LW, ZENG L, BARRON AR, MONTEIRO-RIVIERE NA: Biological interactions of functionalized single-wall carbon nanotubes in human epidermal keratinocytes. *Int J Toxicol* **26**: 103-113, 2007.
- ZHAO G, RAINES AL, WIELAND M, SCHWARTZ Z, BOYAN BD: Requirement for both micron- and submicron scale structure for synergistic responses of osteoblasts to substrate surface energy and topography. *Biomaterials* **28**: 2821-2829, 2007.
- ZHU X, CHEN J, SCHEIDERLER L, ALTEBAEUMER T, GEIS-GERSTORFER J, KERN D: Cellular reactions of osteoblasts to micron- and submicron-scale porous structures of titanium surfaces. *Cells Tissues Organs* **178**: 13-22, 2004.
- ZHU Y, RAN T, LI Y, GUO J, LI W: Dependence of the cytotoxicity of MWCNTs on the culture medium. *Nanotechnology* **17**: 4668-4674, 2006.