Prospects for the use of carbon nanotubes in medicine

Accepted 30th April, 2018

ABSTRACT

Newly discovered compounds showing unique properties can have profound effect on development of medicine. As far as carbon is concerned, a discovery of great consequences for medicine was that of a new allotropic form of carbon known as fullerene. Recently, much interest has been paid to the application of carbon nanotubes as carriers of therapeutic drugs and biosensors in gene therapy or in anticancer therapy.

Keywords: Nanotubes, nanotechnology, medicine.

INTRODUCTION

Nanotechnology has become one of the most intensely developing area of research and it combines the achievements from many branches of science. In 1985, Harold Kroto, Robert Curl and Richard Smalley discovered a new allotropic and molecular form of carbon making icosahedral hollow structures, known as fullerene. As this structure is hollow inside, it can host metal atoms or molecules of chemical compounds. It is expected that this form of drug administration may revolutionize medical industry in near future (Grabowska, 2008). Since 1991, the discovery of carbon nanotubes has been studied in a number of research centres of which the pioneering group was headed by Prof. SumioIijima, NEC, Japan.

New properties and new possibilities of application are discovered daily. New composites with carbon nanotubes have been proposed, showing high mechanical strength, high electric conductivity, exceptional mechanical or electrical features. In medical therapy, the use of carbon nanotubes permits application of active substances to exactly defined target which shortens the time in which the drug reaches the target and increases the effectiveness of therapy.

STRUCTURES PREPARATION AND BASIC CHARACTERISATION

An interesting example of carbon nanostructures are carbon nanotubes. They are made of graphene sheets wrapped to make seamless cylinders. The diameter of nanotubes is by about 10 thousand times smaller than that of a human hair (Grabowska, 2008; Paradise and Goswami, 2007). The nanotube obtained by wrapping a single sheet of graphene is called the single-wall nanotube. Depending on the mode of wrapping of the graphene sheet the nanotubes can be chiral and non-chiral. With respect to the shape of the edge, the non-chiral nanotubes are divided into armchair and zigzag ones. Nanotubes can end with the 0 nm. The ratio of the nanotube length to its diameter can be of an order of 10^{2} to 10^{3}. Depending on the number of graphene layers forming the structure, carbon nanotubes can be divided into single-walled carbon nanotubes (SWCNT) and multi-walled carbon nanotubes (MWCNT), composed of many concentric layers of graphene (Grabowska, 2008).

Varying the chiral angle between hexagons and the tube axis, SWCNTs can be either metals or semi-conductors, with relatively large (circa 0.5 eV for typical diameter of 1.5 nm) or small band gaps (circa 10 meV), even if their diameters are nearly identical (Dresselhaus et al., 1996). With diameters of 1 to 2 nm, lengths range from as short as 50 nm up to 1 cm; SWNTs are one-dimensional (1-D) nanomaterials which may behave distinctly from spherical nanoparticles in biological environment (Liu et al, 2009).

A few methods for production of nanotubes have been proposed, all of which are based on slow condensation of hot vapour of carbon atoms. In the process of their production a mixture of different structures is obtained:
single- and multi-walled nanotubes of different wall configurations, toruses, spirals and fullerenes (Terranova et al., 2006; Chlopek et al., 2006). The methods for production of carbon nanotubes include: laser-induced graphite evaporation, electric arc technique and chemical vapour deposition (CVD) (Pettes et al., 2009). One of the specific properties of nanotubes is their large surface area. Moreover, depending on the diameter and degree of twisting they can behave as a metal or as a semi-conductor. There is the existence of very strong bonds between carbon atoms in the graphene layer, while the nanotubes show high mechanical resistance and their Young modulus is very high (10^12 N/m²), hence, their deformations are elastic and are highly resistant to bending or stretching. Nanotubes can conduct current of very high density, of an order of 10^9 A/cm², as they show very low electric resistance. Their thermal conductivity reaches 6000 W/(K × m) at T = 300 K, which is very useful for removal of heat from electronic elements. The property very attractive from the medical point of view is the possibility of regulating the nanotubes biocompatibility by chemical modifications (Ghorannevis et al., 2010; Odom et al., 1998; Kuryliszyn-Kudelska et al., 2011; Bacsa et al., 2004).

FUNCTIONALIZATION FOR MEDICAL USE OF NANOTUBES

Unmodified carbon nanotubes (CNT) are hardly soluble in water, which restricts their medicinal use. To avoid this problem, the graphene sheets are functionalized in different ways (Iijima, 1991; Jianrong et al., 2004; Kang et al., 2007; Khabashesku et al., 2005). In general, the methods of CNT functionalization are divided into endohedral and exohedral methods. In the endohedral methods, the hollow space inside the tube is filled with substances of different polarity, for example, metals or chemical compounds, while in the exohedral methods the external walls of the tubes are modified (Pagona and Tagmatacharis, 2006; Pastorin et al., 2006).

The functionalization of nanotubes is realized through adsorption of proteins, amino acids, enzymes or nucleic acids (Bacsa et al., 2004; Iijima, 1991; Singh et al., 2005). It can also be performed by adsorption of chemical compounds through pyridine rings. Modification with pyrrolidine rings gives the species soluble in certain organic solvents (Khabashesku et al., 2005). Improvement in nanotubes solubility can also be achieved by introduction of certain biochemical compounds, for example, bovine serum albumin into a solution of nanostructures (Bacsa et al., 2004; Terranova et al., 2006).

APPLICATION IN MEDICINE

The application of nanotubes in medicine is related to the possibility of their bio-functionalization and the control of their biocompatibility (Mikhalovsky and Nikoleav, 2006; McLean, 1974).

DRUG DELIVERY

Much promising is the use of nanotubes as carriers transporting biologically active drugs to certain well-defined sites. This is as a result of their specific properties multi-walled nanotubes which have become the basis of a drug delivery system directly to a target site. CNT have been used to facilitate the absorption of amphotericin B, an antifungal antibiotic. At first, CNTs are subjected to carboxyl acid so that the –COOH groups would attach to the outer surface and thereafter, subjected to diaminetriethylene glycol, which permits incorporation of the antibiotics. On the other hand, CNTs modified with fluorescein isothiocyanate can be used for imaging (Tripisciano et al., 2009; Hampel et al., 2008; Cheung et al., 2010).

CANCER THERAPY

Recently, the use of carbon nanotubes in photodynamic therapy has been studied. Mikhalovsky and Nikoleav (2006) injected carbon nanotubes into the cancer tissue in the rabbit liver and then irradiated this spot with radio waves, which resulted in damaging of the cancer cells. CNT can be modified with certain specific antibodies that would capture some well-defined substances, for example, biological growth factors in tumours. This is as a result of the use of CNTs which has made it possible to resign from the traditional radiotherapy with high-energy radiation also destroying healthy tissues (Tripisciano et al., 2009; Hampel et al., 2008). Their strongly hydrophobic character, responsible for the tendency to aggregation and for possible difficulties with their removal from the organism is an issue of concern. However, as a result of surface functionalization of CNT, for example, with silica, their character can change into hydrophilic form, so that they could form a stable dispersion.

The capturing of therapeutic drug inside the nanotubes is just the first step of goal-directed chemotherapy. They should also be endowed with the properties allowing their accumulation in the tumour and ensuring the release of the active drug at this site and this can be achieved in two ways. The nanotubes can be filled with a ferromagnetic core such that their movements would be controlled by a magnetic field or the nanotubes can be equipped with a cap at the end and this cap would respond to different pH values (Hampel et al., 2008; Cheung et al., 2010).

Chemotherapeutics delivery systems

Due to toxic side effects of most chemotherapeutic agents, there are some limitations in their use. Due to this fact, it is
very important to find a method to develop cell-targeting drug formulations with a wide therapeutic index. Carbon nanotubes have shown great promise as conveyance for targeted drug delivery (Cheung et al., 2010). CNT can be applied as carriers of anticancer drugs to deliver them to a target site within the so-called goal-directed therapy (Bianco et al., 2005; Liu et al., 2008). Drugs administered in traditional chemotherapy affect the whole organism and also destroy healthy cells, while the drugs delivered through goal-directed therapy are released only after having reached the tumour. Another advantage of this method is the possibility of using the carefully adjusted dose of the drug in order to be able to destroy the tumour and not to cause undesired effects. A CNT with a diameter of 80 nm can hold up to 5 million drug molecules (Kam and Dai, 2005).

One of the methods of incorporating drug into CNT is steered molecular dynamic simulation, of which the general principle is to apply an external force to particles in a specific direction by use of harmonic restraint in order to create better change of the particle coordinates. Drugs can either attach to the outer surface of the CNT through functional groups through either covalent or non-covalent bonding, including hydrophobic, π-π stacking and electrostatic interactions (endoedral modification) or can be put inside the CNT (endoedral modification) (Wu et al., 2009; Li et al., 2009). Drug-loaded CNT has to recognize its site of action and the routes by which it can be delivered to target cells. One of the major techniques used involves coating the surface of the CNT with a particular antibody having affinity for the target cancer cell.

Another method used in targeted cancer therapy is modification of the CNT with folic acid and with photosensitizer from the group of porphyrin (Zhang et al., 2011). As a result of using the laser irradiation of the appropriate wavelength on presented structure, singlet oxygen evolution process can be observed. Singlet oxygen destroys the tumor cells depending on the grade of cancer lesions of varying efficacy achieved, but it is not less than 60% (Bianco et al., 2005).

**Thermal ablation**

Specific thermal ablation using single-walled carbon nanotubes targeted by covalently-coupled monoclonal antibodies is used to destroy tumor cells (Marches et al., 2009). Exoedral modifications coupled with those antibodies make the system recognize the cancer cells. The ability of CNTs to absorb near-infrared (NIR) radiation (wavelength 700 to 1100 nm) and convert it into heat gives an opportunity to create a new generation of structures for cancer photo-therapy. NIR light can effectively penetrate healthy tissue and ablate any cell to which the CNTs are attached (Marches et al., 2011).

To increase therapeutical effect of thermal ablation, the chemotherapeutic agent and actinium are placed inside the CNT structure. After introducing modified nanostructure into the patient’s body, determined body area is subjected to the laser radiation with near infrared. As a result of CNT overheating the chemotherapeutic agent is released. Its activity is enhanced by actinium radioactivity (Issels, 2001; Wust et al., 2002). In addition to its lethal activity, hyperthermia has been used in the clinical treatment of solid tumors as a result of enhancing the efficiency of chemo- or radiotherapy. The local increase in temperature also increases the permeability of blood vessels, which can enhance the delivery of drugs to tumors (Iancu and Mocan, 2011).

**T-cell therapy**

T cells, called also T lymphocytes are a type of lymphocyte that plays a significant role in cell-mediated immunity. Their name is derived from the process of their maturation that takes place in the thymus (although some also mature in the tonsils). It was recently found to use tumour specific T cells taken from a patient’s own blood and use them against tumour targets. A promising method to reproducibly expand T cells in human body is by attaching the stimuli for T cells onto artificial substrates with high surface area.

Carbon nanotubes polymers composites (CNPs) can be used as an artificial antigen-presenting cell to efficiently expand the number of T lymphocytes. It was proven that tumour growth was significantly delayed for those mice that was adoptively transferred with CNP-cultured T cells in comparison with those without any treatment at day 14 of therapy (Fadel et al., 2014).

**BIOSSENSORS**

Another interesting application of nanotubes is in biosensors that are able to detect specific molecules (Wang, 2004). For this application, CNTs surface must be functionalized with the enzymes sensitive to a given substance. In such a way it would be possible to make a biosensor detecting in a continuous way the level of glucose in blood.

The conductivity of CNT depends on the functionalization so it will change with subsequent molecules of sugar bound to the enzyme. The nanometric size of the device permits placement of such a detector in the organism. In combination with electochromic materials this device permits design of an intelligent lens whose colour would provide information as regards the level of sugar. After appropriate functionalization CNT can be used for observation of cell properties and changes taking place in cells during their development for control of enzymatic reactions, ion transportation and secretion of proteins or
products of chemical transformations. Detection of DNA particles and neoplastic cells in the early stages of growth is possible as a result of the large surface area of CNT and their ability of electron transportation (Snider et al., 2008; Telega and Latocha, 2012).

BIOCOMPATIBILITY

A very important problem related to carbon nanotubes is their biocompatibility. This problem has been studied by many research groups. Particularly interesting results have been reported by Herzog et al. (2007) who tested the influence of CNT in cells, using osteoblasts and fibroblasts. They tested the effect of MWCNT modified with polysulphone on the lifetime of the cells and the amount of secreted collagen. The presence of CNT only to a small degree weakened the cells’ viability, but promoted the amount of secreted collagen. The effect of increased synthesis of collagen can be used for regeneration of bones and soft tissues with CNT stimulating their growth (Herzog et al., 2007).

GENE THERAPY/DNA DELIVERY

Gene therapy is one of such promising methods for the treatment of cancer and genetic disorders. Genes are transported by special virus-based or not virus-based carriers, the latter groups includes liposomes, polymers and nanoparticles. The use of liposomes brings a risk of undesirable effects such as immunological reaction, inflammatory states or oncosign. In general, the non-virus based carriers do not always ensure the sufficient level of gene expression, which has stimulated the search for new carriers (Malmsten, 2006). The large-molecular and cationic character of functionalized carbon nanotubes (f-CNT) permits electrostatic interaction with plasmid DNA. To evaluate the f-CNT abilities to make complexes with nucleic acids and their translocation, Ali-Boucetta et al. (2008) combined f-CNT at different rates and plasmid DNA containing the marker gene of β-galactosidase. TEM images revealed the presence of CNT-DNA complexes. The functionalised SWCNT seen in the form of bundles among them were the plasmids in the form of ring clusters or highly folded structures. The degree of expression of the marker gene of β-galactosidase confirmed the complexes ability to permeate inside cells. The level of expression of the gene studied was found to be from 5 to 10 times higher for the complexes of f-SWCNT and DNA than for the DNA helix alone (Ali-Boucetta et al., 2008).

Gene transportation by carbon nanotubes can be used for silencing certain genes. Shukla et al. (2002) studied the complexes of f-SWCNT and siRNA of the telomerase gene. They reported a fast penetration of the complexes into a certain line of mouse cancer cells, release of siRNA and effective suppression of the telomerase gene.

TOXICITY

The toxicity of carbon nanotubes can be related to the high ratio of tubes lengths to diameters and to the toxicity of the material of which it is made – graphene. The nanotubes show greater toxicity towards the respiratory system than the particles of diameters larger than 100 nm. CNTs are classified as nanoparticles that can participate in the unknown and unpredictable interactions with biological systems (Lacerda et al., 2006). Their toxicity can be limited by subjecting them to appropriate functionalization. According to the in vitro studies by Sayes et al. (2006), SWCNT covalently functionalized by sulphophenyl and carboxyphenyl groups have weaker cytotoxic effect than the suspension of purified SWCNT in water, stabilized with a 1% solution of surfactant (Sayes et al., 2006). As a result of their size, carbon nanotubes can be treated as fibrous material showing usually high toxicity towards the lungs.

Lam et al. (2004) studied the toxicity of SWCNT in mice. They checked the health risk of exposure to purified and non-purified CNT. According to the results, depending on the dose and the content of a catalyst, the use of SWCNT led to the appearance of granulomas and produced interstitial inflammations; further pathological changes could lead to bronchogenic inflammation of the lungs (Lam et al., 2004).

Shvedowa et al. (2005) studied the effect of SWCNT in different doses on the larynx of mice exposed in CNT in the form of aerosol. The results permitted identification of two SWCNT fractions differing in the size of particles and toxic effect. The first fraction made of CNT aggregates was responsible for the appearance of acute inflammation and formation of granulomas at the site of their accumulation. The second fraction, made of thin delicate CNT of diameters smaller than 50 nm, stimulate the process of fibrosis and contribute to increase in the walls of alveoli in the regions which the primary aggregates did reach (Huczko and Lange, 2007).

Administration of CNT through the trachea and aspiration through the throat led to agglomeration of CNT in the upper part of bronchi and to the beginning of fibrosis. The skin exposure to CNT was also studied. Huczko and Lange (2007) performed a dermatological test on 40 volunteers and Draize test, which revealed the irritating effect of CNT on the skin (Lacerda et al., 2006). On the other hand, the study performed on the lines of human keratinocytes undermines these results.

Shvedowa et al. (2005) studied the effect of non-purified SWCNT on the line of immortalized human keratinocytes (HaCaT) and reported on the increase in the oxidation stress with the simultaneous use of antioxidants, loss of viability and morphological changes in the structure of the cells. To some degree, the results were related to a relatively high content of the catalyst (used for the CNT synthesis) residues (~ 30%) (Shvedowa et al., 2005). For this reason, the authors emphasized the risk related to direct contact of the skin to CNTs. As follows from the
hitherto studies, evaluation of the CNT toxicity is not a simple task. Results of such evaluations are often contradictory. It can be concluded that non-purified carbon nanotubes show rather high toxicity related to the presence of the catalysts (Fe, Ni, Co and Zn) residues. Exposure to purified CNT, especially in high concentrations, leads to much weaker toxic effects. The least toxic are the functionalized nanotubes that are to be used for medical applications (Marches et al., 2011).

CONCLUSION

The application of carbon nanotubes can significantly contribute to solving special problems in medical therapy. Nanotubes can be used for in vivo production of tissues and for controlling of their development. Much promising is the use of carbon nanotubes as carriers of therapeutic drugs in goal-directed therapy or DNA in gene therapy.

REFERENCES

Ali-Boucetta H, Al-Jamil KT, McCarthy D, Prato M, Biancoc A, Kostarelos K (2008). Multiwalled carbon nanotube-doxorubicin supramolecular complexes for cancer therapeutics. Chem Commun (Camb). 4: 459–461.

Bacs RR,  Laurent Ch, Peigney A, Puech P, Hubel H, Dunstan D, Bacs WA (2004). Structural and mechanical properties of double wall carbon nanotubes. NSTI-Nanotech. 3: 214–217.

Bianco A, Kostarelos K, Prato M (2005). Applications of carbon nanotubes in drug delivery. Curr. Opin. Chem. Biol. 9(6): 674–79.

Bianco, A, Kostarelos, K Prato M (2005). Applications of carbon nanotubes in drug delivery. Curr. Opin. Chem. Biol. 9(6): 674–679.

Cheung, W, Pontorriro, F, Taratula, O, Chen AM, He H (2010). DNA and carbon nanotubes as medicine. Adv Drug Deliv. Rev. 62(6): 633-649.

Chlopek J, Czajkowska B, Szaraniec B, Frackowiak E, Szostak K, Bégun F (2006). In vitro studies of carbon nanotubes biocompatibility. Carbon. 44(6): 1106–1111.

Dresselhaus MS, Dresselhaus G, Ealding PC (1996). Science of Fullerenes and Carbon Nanotubes. 2nd ed. San Diego: Academic Press.

Fadel TR, Sharp FA, Vudattu N, Bagheb R, Garty J, Kim D, Hong E, Li N, Hailer GL, Pfefferlei LD, Justesen S, Herold KC, Fahmy TM (2014). A carbon nanotube-polymer composite for T-cell therapy. Nature Nanotechnol. 9(8): 639–647.

Falk MH, Issels RD (2001). Hyperthermia in oncology. Int. J. Hyperthermia. 17: 1–18.

Ghorannevis Z, Kato T, Kaneko T, Hatakeyama Y (2010). Narrow-chirality distributed single-walled carbon nanotube growth from nonmagnetic catalysts. J Am Chem Soc. 132(28): 9570–9572.

Grabowska JF (2008). Przyszeńność zastosowań w medycynie i farmacji. Gazeta Farmaceutyczna. Poland. 6: 38-40.

Hampel S, Kunze D, Hasse D, Kramer K, Rauschenbach M, Ritschel M, Burger M (2011). Magnetic Properties of As-Prepared and Chemically Modified Multiwalled Carbon Nanotubes. Acta. Phys. Pol. A. 119: 597-611.

Hampel S, Kunze D, Hasse D, Kramer K, Rauschenbach M, Ritschel M, Dobrowolski W (2011). Magnetic Properties of "As-Prepared" and Chemically Modified Multiwalled Carbon Nanotubes. Acta Phys. Pol. A. 119: 597-611.

Iijina S (1991). Helical microtubules of graphitic carbon. Nature. pp. 56-58.

Jianrong C, Yuqing M, Nongyuewisp H (2004). Nanotechnology and biosensors. Biotechnol. Adv. 22(7): 505-518.

Kam NW, Dai H (2005). Carbon nanotubes as intracellular protein transporters: Generality and biological functionality. J. Am. Chem. Soc. 127(16): 6021–6026.

Kang S, Pinault M, Pfefferlei LD (2007). Single-walled carbon nanotubes exhibit strong antimicrobial activity. Langmuir. 23(17): 8670-3.

Khabashesku VN, Margrave J, Barrera EV (2005). Fucnialized carbon nanotubes and nanodiamonds for engineering and biomedical applications. Diam. Relat. Mater. 14(3-7): 859-864.

Kuryłyszyn-Kudelska I, Małolepszy A, Mazurkiewicz M, Stohinski L, Dobrowolski W (2011). Nanotechnology. Properties of "As-Prepared" and Chemically Modified Multiwalled Carbon Nanotubes. Acta. Phys. Pol. A. 119: 597-611.

Lam CW, James J, McCluskey R (2004). Pulmonary toxicity of single-wall carbon nanotubes in mice 7 and 90 days after intratracheal instillation. Toxicol Sci. 77(1): 126-134.

Li Y, Cousins BG, Ulijn RV, Kinloch IA (2009). A study of the dynamic interaction of surfactants with graphite and carbon nanotubes using Fmoc-amino acids as a model system. Langmuir. 25(19): 11760–11767.

Liu Z, Tabakman S, Welker K, Dai H (2009). Carbon nanotubes in biology and medicine: in vitro and in vivo detection, imaging and drug delivery. Nano. Res. 2(2): 85-120.

Liu ZI, Robinson JT, Sun X, Dai H (2008). PEGylated nanographene oxide for delivery of water-insoluble cancer drugs. J. Am. Chem. Soc. 130(33): 10876-10877.

Lamstein M (2006). Soft drug delivery systems. Soft Mater. 2: 760-769.

Marches, R, Chakravarty, P, Musselman IH, Bajaj P, Azad RN, Pantano, P, Vitetta, ES (2009). Specific thermal ablation of tumor cells using single-walled carbon nanotubes targeted by covalently-coupled monoclonal antibodies. Int. J. Cancer. 125(12): 2970-2977.

Marches, R, Mikoryak C, Wang, RH, Pantano P, Draper RK, Vitetta, ES (2011). The importance of cellular internalization of antibody-targeted carbon nanotubes in the photothermal ablation of breast cancer cells. Nanotechnology. 22(9): 95-101.

Martel R, Schmidt T, Shea, HR., Hertel T, Avouris, P (1998). Single-and multi-wall carbon nanotube field-effect transistors. App. Physics Letters. 73(17): 2447-2449.

McLean AEM (1974). Host factors in hepatotoxicity. Israel J. Med. Sci. 10: 431-435.

Mikhailovsky SL, Nikoleav VG (2006). Activated carbons as medical adsorbents, [in:], Activated Carbon Surfaces in Environmental Remediation. 4th ed. New York: Elsevier.

Odom TW, Huang JL, Kim, Lieber PC (1998). Scanning tunneling microscopy and spectroscopy studies of single wall carbon nanotubes. Nature. 393(6687): 44(6): 1106–1111.

Pagosa G, Tagmatachis N (2006). Carbon nanotubes: materials for medicinal chemistry and biotechnological applications. Curr. Med. Chem. 13(15): 1789-1799.

Paradise M, Goswami T (2007). Carbon nanotubes - Production and industrial applications. Mater. Des. 28(5): 1477-1489.

Pastorin G, Wu W, Wieczowski S (2006). Zastosowanie nanorurek węglowych w medycynie. Chem. Commun. 21: 1182-1186.

Pettes MT, Shi L (2009). Thermal and Structural Characterizations of Individual Single-, Double, and Multi-Walled Carbon Nanotubes. Adv. Funct. Mater. 19(24): 3918-3925.

Sayes CM, Liang F, Hudson JL, Mendez J, Guo W, Beach JM (2006). Functionalization density dependence of single-walled carbon nanotubes cytotoxicity in vitro. Toxicol. Lett. 161(2): 135-142.

Shukla A, Zhang YH, Dubey P, Margrave J, Li Y, Cousins BG, Ulijn RV, Kinloch IA (2009). A study of the dynamic interaction of surfactants with graphite and carbon nanotubes using Fmoc-amino acids as a model system. Langmuir. 25(19): 11760–11767.

Shukla A, Zhang YH, Dubey P, Margrave J, Li Y, Cousins BG, Ulijn RV, Kinloch IA (2009). A study of the dynamic interaction of surfactants with graphite and carbon nanotubes using Fmoc-amino acids as a model system. Langmuir. 25(19): 11760–11767.

Shukla A, Zhang YH, Dubey P, Margrave J, Li Y, Cousins BG, Ulijn RV, Kinloch IA (2009). A study of the dynamic interaction of surfactants with graphite and carbon nanotubes using Fmoc-amino acids as a model system. Langmuir. 25(19): 11760–11767.
construction of nanotube-based gene delivery vectors. J. Am. Chem. Soc. 127(12): 4388-4396.

Snider RM, Gobatu M, Rue AE, Cliffel DE (2008). A multiwalled carbon nanotube/dihydropyran composite film electrode for insulin detection in a microphysiometer chamber. Anal. Chim. Acta. 609(1): 44-52.

Telega K, Latocha M (2012). Nanotechnologia - przyszłość medycyny. Pol. Merk. Lek. 196: 229-232.

Terranova ML, Sessa V Rossi M (2006). The world of carbon nanotubes: an overview of CVD growth methodologies. Chem. Vapor. Depos. 12(6): 315-325.

Terranova ML, Sessa VM, Rossi M (2006). The World of Carbon Nanotubes: An Overview of CVD Growth Methodologies. Chem. Vapor. Depos. 12(6): 315-325.

Tripisciano C, Kraemer K, Taylor A, Borowiak-Palen E, Single-wall carbon nanotubes based anticancer drug delivery system. Chem Phys Lett. 2009; 478: 200-205

Wang J (2004). Carbon-Nanotube Based Electrochemical Biosensors. Electroanalysis. 17: 7-12.

Wu W, Li R, Bian X (2009). Covalently combining carbon nanotubes with anticancer agent: Preparation and antitumor activity. ACS Nano. 3(9): 2740–2750.

Wust P, Hiklebrandt B, Sreenivas G, Rau B, Gellermann J, Riess H, Felix R, Schlag PM (2002). Hyperthermia in combined treatment of cancer. Lancet Oncol. 3(8): 487–497.

Zhang W, Zhang Z, Zhang Y (2011). The application of carbon nanotubes in target drug delivery systems for cancer therapies. Nanoscale Res Lett. 6: 555. doi: 10.1186/1556-276X-6-555.

Cite this article as:
Igielska-Kalwat J (2018). Prospects for the use of carbon nanotubes in medicine. Acad. J. Biotechnol. 6(6): 074-079.
Submit your manuscript at http://www.academiapublishing.org/ajb