Biocompatibility of Carbon Nanoparticles in HeLa Cells is Dictated by Synthesis and Sterilization Procedures

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Abstract: Due to their unique chemical and physical properties, carbon-based NanoMaterials (C-NMs) are largely exploited in biomedicine, i.e., cell and tissue imaging, drug delivery and tissue engineering scaffold, even if reports regarding their toxicity are still conflicting. In fact, biological effects strictly depend on the dynamic physicochemical characteristics of C-NMs, which in turn are strongly influenced by the procedures of their synthesis, and nanometrological techniques, e.g., Electron Microscopy (EM)-based analysis, are becoming the main tool for researchers to characterize nanoparticles. The aim of the present work is the study of the influence of synthesis and sterilization protocols on the size, shape, stability and biocompatibility of carbon NanoParticles (C-NPs). C-NPs were synthesized by using graphite as bulk material through an electrochemical method applying a constant voltage of 30 V and different times of synthesis. The C-NPs solution was sterilized by adopting different sterilization protocols during and/or after the synthesis. Size, shape and stability were studied by TEM and spectroscopy, while biocompatibility was tested in HeLa cells. Synthesis and sterilization procedures did not influence size, shape and stability of C-NPs, but interfered with C-NPs biocompatibility. In fact, irrespective of time of electrolysis process, the NPs show spherical shape with an average diameter of 7 nm. UV-visible spectra show typical peak of carbonaceous materials that falls at 236 nm without aggregation and sedimentation. However, when NPs obtained at 90 min of synthesis were twice autoclaved the peak shifted to 257 nm. HeLa cells were incubated with different C-NPs solutions administered at different concentrations, ranging from 8×10^5 to 1.6×10^7 C-NPs/cell, for different times (4, 24 and 48h). Cell viability was C-NPs concentration- and time of culture-dependent; interestingly, also the time of electrolysis process used during particles synthesis and procedures adopted to sterilize C-NPs solutions largely influenced cells response.

Keywords: Carbon Nanoparticles, Biocompatibility, Synthesis Parameters, Sterilization Procedures

1. Introduction

Carbon-based NanoMaterials (C-NMs), such as single or multi-walled nanotubes, fullerenes, nanodiamonds, and graphenes, are becoming key materials for various industrial applications due to their unique combinations of chemical and physical properties (i.e., thermal and electrical conductivity, high mechanical strength, and optical properties) [1]. They have promising applications in nanoelectronics [2], microelectrical devices [3], electrochemistry [4,5], sensors [6], catalysis [7] and ultracapacitors [8-10]. In recent years, C-NMs are receiving many attention also in biomedical research [11], mostly in bioimaging [12,13] and drug delivery [14,15]. Engineered carbon NanoParticles (C-NPs) can be synthesized by numerous methods such as the candle soot method [16], laser induced pyrolysis of hydrocarbons [17], low temperature solution synthesis [18], electrochemical oxidation of graphite [19], microwave pyrolysis of sucrose [20], proton-beam irradiation of nanodiamonds [21], thermal decomposition of organic compounds [22,23], using mesoporous silica NPs as template [24], and using polyacrylonitrile as a NPs precursor [25].

Due to the widespread application of nanosized materials, a precise characterization is need for their safe and responsible exploitation. This has opened new nanometrology challenges for an accurate morphological characterization, near atomic resolution, to ascertain NPs elemental composition or to exclude the presence of contaminants introduced during the synthesis procedures. Thus, the precise knowledge of the NPs characteristics,
monitored by nanometrological techniques, is pivotal to ensure their biocompatibility and define their biological effects [26].

Unfortunately, determining biocompatibility or toxicity of C-NPs is quite complicated. The surface coating, the presence of metal catalysts and/or graphite, the dispersion properties, and the tendency to deposit as aggregates, because of high Van der Waals’s forces, influence greatly the behavior of these NPs. Epidemiological and experimental studies, for example, have confirmed the role of C-NPs in aggravating pulmonary disorders such as asthma as well as lung cancer, pulmonary fibrosis and systemic cardiovascular disorders [27]. C-NPs have also a negative impact on the correct development of sea urchin, thus opening environmental risk concerns [28-30]. At the same time, C-nanotubes are largely exploited in bioimaging and in drug delivery [31].

Taking into account the fact that no data are available in the literature about the effect of the synthesis methods of C-NPs on the biocompatibility, in this study we evaluated the cytotoxicity of electrochemical synthesized C-NPs by using graphite as bulk material. In particular, we investigated the influence of synthesis time and sterilization procedures on biocompatibility of C-NPs solution obtained. C-NPs were characterized by TEM and spectroscopy and biocompatibility was evaluated by using HeLa cells in *in vitro* experiments.

2. **Materials and Methods**

2.1. **Chemicals**

All chemicals were of analytical grade and were purchased from Sigma-Aldrich (Sigma, St. Louis, MO, USA) unless otherwise indicated.

2.2. **Synthesis of C-NPs**

High purity graphite rod (99.99%, SPI Supplies Division Structure Probe Inc., West Chester, PA, USA) was used as an anode (5 mm diameter), and a stainless steel rod AISI 1016, 5 mm diameter (Swagelok, Solon, OH, USA), was used as a cathode. The electrodes were immersed in a beaker containing Milli-Q water at a distance of 1 mm. A constant voltage of 30 V was applied during the electrolysis process. Simultaneously, the colloidal solution was forcedly dispersed by an ultrasonicator (1000 W power output; 100 ± 5 kHz frequency) (Flexonomic-1200-35/72/100G, Mirae Ultrasonic Tech., Co., Ltd., Guro gu Seoul, South Korea) to prevent the aggregation of NPs. Different times of the synthesizing reaction and sterility conditions have been tested as reported in Table 1.

20 mL of NPs solution was diluted with 30 mL of distilled water. The average size, size distribution, morphology of NPs were studied by high resolution TEM and UV-visible spectroscopy. The stability of different C-NPs concentrations over the time was assayed in complete Eagle’s Minimum Essential Medium (EMEM).

2.3. **Physical Analyses of C-NPs**

UV-visible spectra were recorded in the range between 200 and 1000 nm by using a T80 spectrophotometer (PG Instruments Ltd., Wibtoft, UK). Optical spectra were obtained from the prepared solution by measuring the absorption of the solution in a quartz cuvette with a 1 cm optical path. The optical properties were monitored for a period of 6 days to test the solution stability. Ultrastructural observations were performed by a Hitachi 7700 TEM, operating at 100 kV (Hitachi High Technologies America Inc., Dallas, TX, USA). For TEM observations specimens were placed as small droplets of NPs solutions onto standard carbon supported 600-mesh copper grids and let to slowly air dry. To determine the NPs distribution sizes, 50 fields (150×100 nm) were randomly chosen and analyzed by using the image software Image J (US NIH, Bethesda, MD, USA).

2.4. **Biocompatibility Test**

Human epithelioid cervix carcinoma cells (HeLa) were routinely cultured in EMEM (Cambrex, Verviers, Belgium) supplemented with 10% fetal calf serum, 2 mM L-glutamine (Cambrex, Verviers, Belgium), 100 IU/ml penicillin and streptomycin solution and 10000 U/ml nystatin (antimycotic solution) (Cambrex, Verviers, Belgium), in a 5% CO2 humidified atmosphere at 37°C.

For all experiments, HeLa cells were harvested from sub-confluent cultures by trypsinization, resuspended in fresh complete medium and plated into 6-well plates at a density of 1.5×10^5 cells/mL.

### Table 1. Scheme of the experiments.

| Parameters for electrochemical synthesis of C-NPs | Sterile protocols (before and/or after C-NPs administration to cells) | Investigations performed on C-NPs | on C-NPs treated cells |
|-----------------------------------------------|---------------------------------------------------------------|----------------------------------|------------------------|
| Electric potential difference ΔV (volt)        | Time (min)                                                   | C-NPs solution autoclaved twice at 121 °C, 2 atm for 20 min | HeLa cells incubated with: |
|                                               |                                                              | C-NPs synthesized under sterile conditions (materials and workplaces) & TEM characterization of freshly synthesized and 6 days old C-NPs | i) 8×10^5 C-NPs/cell |
| 30                                            | 90                                                          | C-NPs solution microfiltered with filters (pores sized 0.2 µm) & UV-visible spectra from 200 to 1000 nm) | ii) 1.6×10^6 C-NPs/cell |
| *                                             | 150                                                         |                                                                | iii) 3.2×10^6 C-NPs/cell |
| *                                             | 190                                                         |                                                                | iv) 8×10^6 C-NPs/cell |
|                                               |                                                              |                                                                | v) 1.6×10^7 C-NPs/cell |
|                                               |                                                              |                                                                | at 4, 24 and 48h of incubation: MTLM (phase contrast) |
After 24h incubation, the culture medium was replaced with a fresh one containing variable concentrations of C-NPs (8×10^6, 1.6×10^6, 3.2×10^6, 8×10^6, 1.6×10^7 C-NPs/cell). Cells viability was assayed after 4, 24 and 48h of incubation with C-NPs. In each experiment, untreated cells served as negative control. At least six independent experiments and two replicates for each C-NPs concentration were done. At fixed times after C-NPs exposure, the culture medium was discharged, the cells were washed twice with Phosphate-Buffered Saline (PBS) and fresh culture medium containing 1 mg/mL of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide salt (MTT) was added to each well. After 2h incubation at 37°C in a 5% CO2 humidified atmosphere, the MTT, reduced by the mitochondrial reductase of vital cells, produced a dark insoluble product, formazan salt. Then, the formazan salts were dissolved in DimethylSulfoxide (DMSO), leading to a violet solution whose absorbance was measured with Ultrospec 4000 UV-visible spectrophotometer (Pharmacia Biotech, Stockholm, Sweden) at 570 nm. Viability was expressed in percentage as the Relative Growth Rate (RGR) by the equation:

\[
\text{RGR} = \left( \frac{D_{\text{sample}}}{D_{\text{control}}} \right) \times 100
\]

where \(D_{\text{sample}}\) and \(D_{\text{control}}\) are respectively the absorbance of the test samples and the negative controls.

2.5. Statistical Analysis

Data were analyzed by performing one-way ANalysis Of VAriance (ANOVA) at the 95% confidence level. P values less than 0.05 were considered significant.

3. Results

3.1. C-NPs Characterization

A basic step in the study of NPs cytotoxicity is the detailed description of their characteristics that are strictly dependent on synthesis procedures. The shape, size and stability of the C-NPs used in this work were carefully characterized using TEM and spectroscopy. TEM images (Fig. 1) of C-NPs show that NPs have spherical shape and are polydisperse in the solution. Morphometry of the TEM images provides C-NPs size and size distribution, that ranges from 4 to 10 nm with an average diameter of 7 nm as reported in Table 2. The time of the electrochemical synthesis (i.e., 90, 150 and 190 min) as well as the sterility protocols did not affect NPs size, size distribution and dispersion (Table 2).

UV-visible spectra are recorded soon after synthesis and after 6 days in complete EMEM culture medium in the range between 200 and 1000 nm. In Fig. 1 we report the data about stability of C-NPs solutions obtained applying 90 min of synthesis. In fact, the time of the electrochemical process does not influence the stability of the C-NPs solutions. The peak at 236 nm indicates the presence of C-NMs in the solution (Figs 1a,c,d). The treatment at 121°C and 2 atm for 20 min affects the spectroscopic absorbance of C-NPs as showed by the shift of the peak from 236 nm to 257 nm (Fig. 1b). The well-defined peaks at 236 nm or at 257 nm show stability of C-NPs over time without aggregation and sedimentation.

### Table 2. Size distribution of freshly synthesized C-NPs.

| Application time of AV (30 V) | Sterile protocols (before and/or after C-NPs administration to cells) | Average diameter ± SD (nm) |
|-------------------------------|---------------------------------------------------------------------|---------------------------|
| 90 min                        | None                                                                | 7± 3                      |
|                               | Autoclaved twice                                                    | 7± 2                      |
| 150 min                       | Synthesis under sterile conditions                                  | 7± 2                      |
| 190 min                       | Microfiltration                                                     | 6± 2                      |

*To determine the C-NPs distribution sizes 50 randomly fields of 150×100 nm were analysed by using the image software Image J (US NIH, Bethesda, MD, USA); ΔV = electric potential difference.

3.2. Biocompatibility of C-NPs

To investigate whether the different parameters, i.e., time of synthesis and sterilization procedures, used for obtaining C-NPs solution can interfere with biocompatibility and/or toxicity, HeLa cells were exposed to increasing C-NPs concentrations (8×10^5, 1.6×10^6, 3.2×10^6, 8×10^6, 1.6×10^7 NPs/cell) for different times (4, 24 and 48h). As expected, our results indicate that cell viability is C-NPs concentration- and time of culture-dependent. Interestingly, for the first time is here reported that also the time of electrolysis process used during particles synthesis and procedures adopted to sterilize C-NPs solutions largely influenced cells response. In Fig. 2 we report representative data about cell viability of C-NPs solutions. In general, the highest toxicity was always measured at 24 and 48h, irrespective of the modality of synthesis of C-NPs and concentration used. Conversely, at the shortest time of incubation examined, 3.2×10^6 NPs/cell prepared during 90 min of electrolysis was the most toxic NPs preparation, inducing more than 20% of viability reduction. The greater reduction in cell viability (ranging between 30 and 50%) was observed in the presence of the highest C-NPs concentrations (8×10^5 and 1.6×10^7 NPs/cell) at 24 and 48h of culture (Fig. 2a). In the presence of C-NPs prepared during 90 min of electrolysis process, a generalized cell viability reduction was found after 4h of culture; differences were related to NPs concentrations. Conversely, C-NPs prepared during 90 min of electrolytic process and autoclaved for two cycles at 121°C and 2 atm for 20 min were not toxic for HeLa cells even at 48h of treatment (Fig. 2b). Noteworthy, they induced a significant cell viability increase (120% vs 100% of the untreated cells) (Fig. 2b). By increasing the electrolytic synthesis times, i.e., 150 and 190 min, C-NPs treatment caused a significant inhibition of cell growth, reaching values of around 40% survival in the presence of the 1.6×10^7 NPs/cell (Figs 2c,d). Shorter treatments, i.e., 4h with C-NPs, were not toxic independently on C-NPs concentrations; in addition, 8×10^6 and 1.6×10^7 C-NPs/cell induced HeLa cells proliferation respectively of 130% and 140% (Figs 2c,d). A similar trend was observed when C-NPs were obtained at the longest electrolytic synthesis time, i.e., 190 min; indeed, the peak of proliferation was significant.
higher (Fig. 2d).

The major morphological changes of HeLa cells treated with C-NPs, monitored at inverted Light Microscope (LM), are elongation of the cellular body and substrate detachment (Fig. 2).

Figure 1. Spectroscopy and TEM of C-NPs freshly prepared and 6 days old in EMEM after different sterilization procedures. UV-visible absorbance spectra and TEM pictures are referring to C-NPs synthesized via electrochemical process at 30 V electric potential difference for 90 min in different condition of sterility: C-NPs (a) synthesized without sterility, (b) autoclaved twice, (c) synthesized under sterile conditions or (d) microfiltered. Each curve represents the mean of six independent experiments, each done in duplicate. The SDs are under the limit to be plotted. TEM micrographs of C-NPs synthesized at 30 V for 90 min with different sterilization protocols as above reported; bar = 5 nm.
Figure 2. Cell viability of HeLa cells up to 48h of incubation with different amounts of C-NPs synthesized at 30 V for different times with or without different sterilization protocols. HeLa cells viability was assayed in the presence of different amounts of C-NPs by MTT assay: C-NPs (a) synthesized for 90 min in non sterile condition, (b) autoclaved twice after 90 min of synthesis, (c) synthesized for 150 min under sterile conditions or (d) synthesized for 190 min and microfiltered. The values are reported as percentage of the control untreated cells considered as 100%. Each value represents the mean ± SD of six independent experiments, each done in duplicate. Star indicates significant value with respect to the corresponding control value at the same time point (p<0.05). Ctrl (control) = untreated cells; h = hours. LM micrographs show HeLa cells in culture and the treatment refers to the histogram above. The number on the picture refers to the column with the corresponding number. The changes of cell viability are indicated as increased (+) or decreased (-) percentages with respect to the control untreated cells taken as 100%; bar = 10 µm.

4. Discussion

In this work, we used an electrolysis process to obtain C-NPs from graphite. In particular, we evaluated different times and sterile conditions during and/or after electrochemical synthesis on features of C-NPs, in terms of size, shape and stability. Moreover, we evaluated the biocompatibility of the different C-NPs solutions in HeLa cells by using different NPs amounts and different times of incubation.

Here we demonstrated that synthesis and sterilization protocols have no effect on the average diameter (7 nm) and shape (spherical) of the C-NPs neither on the stability over 6 days of storage. Indeed, NPs show typical peak of absorbance at 236 nm and do not aggregate and sediment. Conversely, high temperature (121°C) and pressure (2 atm) caused a shift of the peak at 257 nm, suggesting putative
chemical features change as demonstrated by preliminary Raman characterizations (data not shown), without affecting the stability over time.

C-NMs are the most exploited in biomedical engineering as cell and tissue labeling and imaging [31], drug delivery systems [32] and reinforcing tissue engineering scaffold [33]. Despite many successful applications in biomedical engineering, there is a growing concern for safety with C-NPs and biocompatibility, which are essential requisites for biomedical applications. Conflicting reports exist regarding the toxicity of C-NMs, that largely stem from the variability of material properties (i.e., size and surface) and model used (i.e., in vivo vs in vitro, cell types, tissues considered and animals types) [34]. In vitro studies suggest that cytotoxicity of C-NMs depends on their cellular uptake, agglomeration and induced oxidative stress; conversely, most studies in vivo have shown that C-NMs do not cause significant toxicity [35-36]. Our previous studies have demonstrated the toxicity induced of C-NPs on P. lividus development [28-30]. In particular, C-NPs interfere with biomineralization process during skeletogenesis. In addition C-NPs ingested by P. lividus sea urchin pluteus induce a defense response by the animals [30].

The results here reported suggest that the time of synthesis is pivotal in cellular response. The toxicity of the C-NPs solution obtained at 90 min of synthesis process without any sterilization procedure is the more toxic, probably due to the presence of contaminants in the solution. On the other hand, by sterilizing the solution, the viability of HeLa cells significantly increased. In particular, the change in chemical characteristics of C-NPs observed in solution sterilized by twice autoclaving positively influence the interaction with HeLa cells, as suggested by the absence of cell toxicity. Conversely, the C-NPs synthesized during 150 min of electrolysis process and microfiltered after synthesis affect cell viability in a concentration and time dependent manner.

5. Conclusion

In conclusion, our results suggest that the synthesis procedure is crucial in inducing cellular response in terms of cytotoxicity. In fact, synthesis affects size, morphology, stability and properties (chemical and physical) of the NMs. Thus, very important is the effort of researchers to develop procedures for producing reproducible, stable and biocompatible NMs. Also, more systematic biological evaluations of C-NMs induced effects are important in order to determine their precise pharmacokinetics, cytotoxicity and optimal dosages. In fact, deeper investigation, such as uptake by the cells, intracellular localization, and cellular metabolism, are need to be better evaluated in relation to synthesis and sterilization protocols of C-NPs biocompatible.

These concerns highlight the pivotal role that nanometrological techniques play in different scientific and industrial fields, since they allow to yield information of surface features, shape, size and structure of NMs as well as of their intracellular uptake and localization.

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