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Abstract. The research and development for biomedical applications are recently focused on multifunctional nanoparticles. To integrate various functionalities, different methods of modifying the particle’s physical properties are developed. Among the considered, nanodiamond (ND) is a promising candidate for the development of multifunctional complex due to its variable features in size, structure, surface chemistry, physical properties, and biocompatibility. In addition to its well-studied structural, surface, electrochemical and photonic properties, strong magnetism of ND can be observed. In the present work, magnetically modified ND is introduced in terms of its bioapplications. Along with the soft ferromagnetism of ND, the increased fluorescence at one- and two-photon excitation is realized. Utilizing the combined magnetic and fluorescence properties of the magnetically modified ND, fluorescence imaging, fluorescence lifetime imaging and manipulation of cells by magnetic field are demonstrated. The perspectives to use the magnetic ND for drug delivery, cells magnetic separation and filtration, in bioengineering to control the cell distribution combined with imaging and treatment are discussed. © 2018 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JBO.23.9.091404]

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1 Introduction

The recent intense developed nanoparticle (NP) applications in biology and medicine are essentially determined by their multifunctional uses. Particularly, for example, theranostic application generally includes combining functionalities of drug delivery or other medical treatments with multimodal imaging. Therefore, the particles that are able to combine the desired physical/chemical properties attract close attention. Among all NPs, nanodiamond (ND) appears to be a promising candidate for the multifunctional application due to its variable features in size, structure, surface chemistry, physical properties, and availability in various size and surface structure, and, most importantly, the demonstrated biocompatibility in spite of some limitations, a number of facilities for ND theranostic applications have been demonstrated. These include stimulus-responsible ND drug delivery and release, multimodal imaging utilizing luminescence properties of ND, allowing fluorescence imaging at one- and two-photon excitation, fluorescence lifetime imaging (FLIM), Raman spectroscopic analysis and mapping, and optical sensing on the base of energy transferred and magnetic sensing utilizing spin properties of structural defects. One possibility to get multifunctional nanoparticles is to modify the nanoparticle to enhance the desired properties for certain application, for example, methods to enhance ND fluorescence in different spectral ranges.

In this work, we discuss magnetically modified ND and its biomedical applications. Magnetic ND (MND) can combine properties and advantages of the ND and magnetic nanoparticle. The most developed and clear way to prepare magnetic ND is doping with magnetic atoms. However, in some cases, such ND can show some level of cytotoxicity and be not biocompatible.

Here we report, the magnetic ND was synthesized and treated without the use of metals and metal oxides using the method proposed by ray techniques. The origin of its magnetism is unclear. As for magnetic properties of diamond in general, pure diamond structure combines diamagnetic and paramagnetic components, but in an intermediate graphite–diamond structure the conditions for spin ordering and magnetic interactions can be created. Such structure can be realized in ND. Ferromagnetic properties of carbon nanostructures are discussed for theoretically predicted structures and are found experimentally; the methods to obtain such properties without embedding magnetic atoms are developed using mechanical, electromagnetic treatments, hydrogenation, creation defects and doping (with nonmagnetic atoms) and surface state modification. Additionally, spin properties of structural defects of ND are studied and the applications of ND are discussed, such as magnetic resonance imaging (MRI) agent for MRI and for quantum sensing.

Bioapplications of magnetic nanoparticles have also developed into a topical issue. The magnetic nanoparticles of various composition, size, shape, and surface properties are developed...
for different purposes. For instance, they can serve as probes in living tissue and be controlled noninvasively from the outside by an external magnetic field, wherein the magnetic field action on the diamagnetic biotissues is negligible. The most developed applications of magnetic nanoparticles are used as a MRI marker at local magnetic hyperthermia and its applications; controlled drug delivery, guided by magnetic field; cell separation and filtration. With the aforementioned possibilities, in the present work, the magnetic-modified ND is characterized, analyzed, and demonstrated from the point of view of bioapplications.

2 Methods and Materials

2.1 Magnetic Fluorescent Nanodiamond

The ND, called further RayND, was synthesized by laser treat-
ment of specially prepared hydrocarbon targets in liquid me-
dium without the use of metals or metal oxides. These NDs produced by controlled technology of light hydrodynamic pulse (LHDP) have average size of 4.3 nm and are rather sim-
ilar to ND obtained by detonation synthesis (DND), just purer and more uniform due to the use of metal-free raw materials and the controlled method of the synthesis. After isolation from hydrocarbon blend, synthesized RayND particles underwent a two-step chemical–thermal modification using metal-free chem-
icals. First, RayND was treated thermally for ordering carbon radicals on the ND surface resulting in the unusual ferromagnet-
ism of unknown nature. Then to provide the colloidal stability, to enhance the ND dispersion in aqueous solutions and increase in the sedimentation stability, of diamond nanofluids, the obtained magnetic ND powder was further chemically modified in metal-free solvents, nitrogenized, which for unclear reason changed its optical properties (e.g., PL). Note, that the process of RayND surface modification is not disclosed in this paper. Magnetically and optically modified ND (RayND-M) were ana-
yzed and compared with DND received with nominal average size of crystallites 3 to 10 nm (Microdiamant AG, Switzerland).

2.2 RayND-M Characterization

The diamond structure of ND (DND, RayND, and RayND-M) was confirmed by x-ray diffraction analysis (XRD) using D8 Advance diffractometer (Bruker AXS, Karlsruhe, Germany), performed as described in Ref. The obtained XRD patterns are identical and indicate that all samples have almost the same crystalline structure close to 100% diamond with the average size of primary particles automatically defined by Scherrer’s formula of 4.3 to 4.5 nm.

Magnetic properties of the RayND-M, and the comparison with DND, were measured using LakeShore 7407 vibrating sample magnetometer (VSM, Lake Shore Cryotronics) at ambient conditions. The ND samples (about 15 mg) were packed in the plastic diamagnetic bag with a size of 5 × 5 mm and a thickness of 0.5 mm. The free bag was also investigated for magnetic moment field dependence. The sample magnetic moment was determined as a difference of the sample and free bag signals at the same magnetic field.

The composition of the investigated samples was estimated via energy-dispersive x-ray analysis (EDX), with scanning electron microscopy imaging using FE-SEM JEOL JSM 7000F (JEOL, Japan) to check the presence of magnetic atoms in the RayND-M.

For EDX-SEM analysis as well as for measurements of Raman, FTIR, fluorescence spectra, and fluorescence lifetime measurements, ND water suspension was dropped on the Si substrate (1 cm × 1 cm) and dried in a desiccator at room temperature.

The particles and their aggregates size distribution as well as c-potential in water were measured using the dynamic light scattering (DLS) method in a Zetasizer nano ZS (Malvern Instruments, United Kingdom), with a 4-mW, 633-nm wave-
length He–Ne laser, detection angle 173 deg at pH ~ 6.8.

To estimate predominant structure of crystallites, Raman spectra were measured using a Renishaw 1000B (Renishaw, United Kingdom) spectrometer equipped with a CW laser (DPGL-2100F; Photop Suwtech, China) with 532-nm wave-
length; and a confocal Raman α-300S (Witec, Germany) with a 488-nm wavelength argon laser (Melles Griot). Surface functional groups were analyzed using a Fourier transform infrared spectroscopy (Bomem MB154 FTIR spectrometer, ABB Bomem, Switzerland) in situ using a mercury cadmium telluride liquid nitrogen cooled detector under vacuum with chamber base pressure ~3 × 10^{-5} Torr.

To study photonic properties of the RayND-M, absorption spectra were measured for ND water suspensions using a UV–visible JASCO V-550 spectrometer (JASCO, Japan) at room temperature. For the fluorescence spectra, the Renishaw 1000B and α-300S confocal Raman spectrometers were used. Analysis of the RayND-M fluorescence lifetime at two-photon excitation and FLIM at their interaction with target biological systems were performed using a system on the base of a femtosecond tunable Ti-Sapphine laser Chameleon Ultra-II (Coherent), the used excitation wavelength was in the 760- to 800-nm range, repetition rate 80 MHz, pulse duration 140 fs, and laser power at about 3 mW. For FLIM, a 2-D scanner (EINST Technology, Singapore) was used. The registration was recorded in the 450- to 650-nm spectral range with the single photon counting system PicoHarp 300 (PicoQuant, Germany) and cooled PMT connected with an Olympus IX 71 optical microscope, 40x objective.

2.3 RayND-M Interaction with Cells

The biointeraction of RayND-M was studied using baby hamster kidney cells (BHK) and human lung alveolar carcinoma cells (A549) cell cultures obtained from Biosource Collection and Research Center, Taiwan (BCRC, Taiwan). The A549 cells were cultured in RPMI1640 medium (Gibco, Invitrogen, United Kingdom), BHK cells were cultured in Dulbecco’s modified eagle medium (DMEM) (Gibco, Invitrogen, United Kingdom), all are supplemented with 2-mM L-glutamine (Invitrogen), 10% fetal bovine serum (FBS) (Gibco/Life Technologies). Cells were maintained under standard cell culture conditions in an incubator (Galaxy 1705, Eppendorf) at 95% air and 5% CO_{2} and 37°C. Culture medium was replaced with fresh medium every 48 or 72 h. Cells were detached by treatment with 0.5% trypsin and 2.6 mM ethyl-enediaminetetraacetic acid (Gibco/Life Technologies); cultures were subcultured routinely at ~80% confluence.

Human serum albumin (HSA, Sigma, Ronkonkoma) was physically adsorbed on RayND-M. The albumin solution with a concentration of 40 mg/ml and an aqueous suspension of the ND with concentration of 4 mg/ml were mixed in a 1:1 ratio. After thorough mixing for 2 h, the mixture was centrifuged.
to separate ND from the protein adsorbed on the surface. The separated sediment was washed twice in distilled water. This method provides the formation of a stable ND-protein complex.

To confirm the adsorption, the albumin content in the solution was determined using a UV–Visible spectrometer JASCO V-550 before its mixing with ND and after the isolation of ND–protein complexes.

To study the interaction with the BHK cells, the cells were treated with 20 μg/ml of RayND-M added to the cell growth medium for 4 h, whereas nonreacted ND has been washed out. Then the cells were fixed with 3.7% formaldehyde (FA) on the coverslips. The cells with ND were investigated with a confocal microscope and FLIM, to visualize the ND penetration into the cell. For investigations with the confocal microscope, cell cytosol was dyed with Alexa Fluor 674 (Sigma); the nuclei were stained with Hoechst33342 (Sigma). A TCS SP5 Laser Scanning Fluorescence Confocal Microscope (Leica, Germany) was used. To define the ND location in the cell structures, the scan along vertical direction (Z-scan) was performed.

For ND intercellular distribution visualization, the RayND-M fluorescence was excited by 488 nm and collected in the range from 560 to 600 nm; Hoechst fluorescence was excited with 405 nm and detected at 450 to 490 nm; cytoplasm stained with Alexa Fluor 647 was excited with 633 nm with detection at 640 to 690 nm.

For FLIM investigation, the cells were not dyed. The FLIM investigation at two-photon excitation was performed as described above.

The cytotoxicity was estimated by MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay using ND-HSA complex and BHK cell line. BHK cells were seeded in a 96-well microtiter plate in DMEM supplemented with 2-mM l-glutamine (Invitrogen) and 10% of FBS and incubated at 37°C and 5% CO2 for 24 h. After incubation, the medium was removed. Then the cells were treated with a fresh medium containing ND-HSA of different RayND-M concentrations (5 to 200 μg/ml) for 24 and 48 h. After this procedure, the medium was removed and the cells were treated for 4 h with MTT reagent (with the concentration of 2.5 mg/ml). The surviving cells converted MTT-agent to formazan, a blue-purple color dissolved in dimethyl sulfoxide (DMSO). The solution was removed and 200 μl of DMSO was added to dissolve the formazan.

The optical absorption of the treated cells and control (nontreated) cells was measured at 570 nm (OD 570) with ELISA reader (MRX revelation Microplate Reader, DYNEX). Relative percentages of surviving cells were calculated by dividing the absorbance of the treated cells with that of the control measured in each experiment.

For the experiment with magnetic guidance of the cells containing RayND-M by a magnetic field, the cells were detached from the bottom of the culture dish, suspended in the cellular medium, and investigated with microscopy and FLIM.

2.4 Utilizing RayND-M Magnetic Properties

2.4.1 Magnetic field guidance of RayND-M/cell complexes

The possibility of magnetic guidance of ND and ND-containing cells was demonstrated for particles in water and for RayND-M-containing living cells. To create a nonuniform magnetic field, a thin steel needle (with diameter of 0.18 mm) was magnetized with neodymium magnet (NdFeB, disk of diameter of 5 mm; National Imports). For a preliminary testing, RayND-M was suspended in the distilled water with a concentration of 20 μg/ml. To demonstrate a magnetic guidance and control by the magnetic field of the spatial distribution of the ND-containing cells, the cells were detached from the bottom of the culture dish and suspended in the cellular medium. The guidance of particles and their aggregates and the cells with embedded RayND-M by the magnetic field were observed with a conventional microscope (Olympus IX 71, objective with magnification 40×) and with FLIM.

2.4.2 Magnetic field-controlled RayND-M/cell complex distribution and positioning

To demonstrate the possibility to control the spatial distribution of the cells during culturing by the magnetic field, suspension of RayND-M-containing cells was mixed with the cells without the ND and recultured together for 12 to 36 h in the presence of the magnetic field. The magnetic field was applied locally to the selected area of the cultural dish, using the permanent magnets. After removing the magnets, the areas treated and nontreated with the magnetic field were analyzed with a confocal microscope and FLIM to see the cells distribution.

3 Results and Discussion

3.1 Characterization of the Nanodiamond

In Fig. the magnetization curve of RayND-M is shown and compared with the magnetization curve of diamagnetic DND. For the RayND-M, the strong magnetization with magnetic hysteresis was observed revealing soft ferromagnetism of this ND. In comparison with nanomagnetite particles, the magnetization of RayND-M is almost 100 times less, so the interaction (and aggregation also) of the particles caused by magnetic dipole–dipole interaction should be weaker and this fact could be considered as the advantage. Figure shows the elemental analysis of the RayND-M; the sample’s SEM image is shown in Fig. Characteristic EDX spectrum of RayND-M demonstrates a very low content of magnetic atoms (Fe, Co, and Ni). Numerically estimated for several measurements the highest content was observed for Fe, being from 0 to 0.2 wt. %. Note that DND powders from diverse suppliers usually have incombustible impurities of 0.5 to 3.2 wt. % with Fe content defined as 0.10 to 0.55 wt. %. however, they do not exhibit tangible ferromagnetic properties. Therefore the origin of ferromagnetism in RayND-M could be considered due to their surface structural modification rather than the elemental contents. However, the mechanism is still unclear.

The magnetic properties of carbon are recently discussed in some theoretical and experimental works. The models that in principle can be realized in ND are suggested. For example, it has been shown that if some carbon sp3-hybridized atoms in a graphite two-dimensional network are substituted by three-valence atoms such as sp3-hybridized carbon atoms, the resulting magnetic moment can be high. The system becomes nonplanar as the tetrahedral-coordinated carbon atoms link to atoms of the graphite-like layers. Thus, a stable all-carbon structure with the ferromagnetic interaction can be created. However, the magnetism nature of nanocarbon is still unclear and is not fully understood.
recently, especially imparting paramagnetism and ferromagnetism to ND significantly increases the ND attractiveness for its use in biomedical researches and applications. Figure 1(a) shows large aggregates of the studied nanoparticles in the state of dry powder. The DLS analysis of the nanoparticles in the water suspension showed an average size of 90 ± 10 nm. These results also correspond to the particles aggregation. In contrast to the magnetization curve, the Raman spectrum of the RayND-M [Fig. 2(a)] is similar in principle with aggregation. In contrast to the magnetization curve, the Raman characteristic spectra of fine ND (DND). It reveals a peak of sp3-hybridized carbon, confirming the diamond structure of the particles. The peak is broadened and shifted, relative to the peak of bulk diamond at 1332 cm$^{-1}$, to 1322 to 1325 cm$^{-1}$. This shift is attributed to the phonon confinement effect at decreasing the crystallites size to 3 to 10 nm. The shift of the diamond band to lower wavenumbers at the decrease of crystallite size is often accompanied by shifting the characteristic band for sp2-bonded carbon or G-band from 1590 cm$^{-1}$ to higher wavenumbers (1630 cm$^{-1}$ in presented case). It is ascribed as a superposition of G-band with a vibration of some surface groups or to interstitial defect, which consists of an isolated sp2-bonded carbon pair in the position of one carbon atom in a diamond lattice. Note that the cross section of Raman scattering of diamond is low, so relatively intense diamond and D-band reveals significant diamond (sp$^3$) fraction.

The FTIR spectrum [Fig. 2(b)] of RayND-M shows the surface carboxyl group characterized by resolving the C=O stretching in the 1720 to 1780 cm$^{-1}$ range and O–H bending 1630 to 1640 cm$^{-1}$. The lower frequency range from 1000 to 1300 cm$^{-1}$ is usually assigned for the C–O functional group in ether/ester structure present on RayND-M. A corresponding OH stretching feature is observed at 3000 to 3600 cm$^{-1}$, which could be related to the carboxyl group and water adsorption. Atmospheric CO$_2$ is in wavenumber 2300 to 2350 cm$^{-1}$ and C–H related to hydrocarbon or surface adsorbed hydrogen on sp$^3$-dangling carbon is found in the range of 2800 to 3000 cm$^{-1}$. As compared with RayND-M, there is no substantial difference with nonmodified DND. The $z$-potential was measured in bidistilled water suspension to be $-32.8$ mV at pH 6.8, which is consistent with carboxylated ND. Therefore, the crystal structure of RayND-M, observed with Raman, and the surface properties characterized with FTIR are similar to ND and RayND; this suggests some well-developed methods can be used to attach molecules of interest to the RayND-M surface for drug delivery, targeted imaging, etc.

In contrast to most NDs, the photonic properties of RayND-M would be quite promising for multifunctional applications. It was found well detectable at confocal imaging at excitation in UV–visible range and two-photon excitation also gives a strong signal, which allow lifetime analysis and FLIM. Figure 3(a) shows the UV–visible absorption and photoluminescence spectra measured using 488- and 532-nm wavelengths CW lasers excitations. Lifetime decay curve and FLIM of RayND-M powder at two-photon excitation using a 790-nm wavelength femtosecond laser, pulse duration 140 fs, and repetition rate 80 MHz are shown in Fig. 3(b). Fluorescence lifetime is short and estimated to be $\sim 0.2$ to 0.5 ns, being close to instrument response function (IRF). The signal is intense and the lifetime differs significantly from lifetimes of the most endogenous fluorophores, so RayND-M is well detectable both at two-photon excitation of fluorescence lifetime and at one-photon fluorescence measurements. Note that the origin of enhanced
fluorescence emission from magnetic-modified ND is also not clear yet and can be discussed together with the origin of magnetic properties, but it is subject to other consideration.

3.2 Interaction with Cell

RayND-M penetration into the cell is demonstrated using confocal microscopic imaging. In Fig. 4(a), the confocal fluorescence image of the BHK cell is presented after the cell incubation with RayND-M for 4 h, with the concentration of ND in the culture medium (DMEM) set at 20 μg/ml. In the image, ND is shown in green, the cell cytoplasm in red, and nuclei is shown in blue. Yellow reveals the ND entered the cytoplasm. The cross-sectional z-scan images on one cell at changing positions in the z-axis with 1-μm step were performed for observation of the ND inside the cell. It clearly reveals ND engulfment by the cell and their localization in the cytoplasm, like other ND, the penetration of which into cells of various types was observed and the clathrin-dependent endocytosis was studied and confirmed.

Fluorescence lifetime analysis [Fig. 4(b)] and imaging [Figs. 4(c) and 4(d)] at two-photon excitation also allow visualizing ND interaction with the cell. Since the RayND-M fluorescence lifetime is much shorter than that of the cell autofluorescence, RayND-M is well observed in FLIM, which enables to define the ND location inside the cell, to monitor, and to quantify the metabolic state of the living cell and its viability can be used as lifetime analyzing, for example, endogenous fluorophores, such as NADH and FAD.

Low cytotoxicity of the RayND-M was observed using the MTT assay with the viability of the BHK cell line (Fig. 5). No concentration dependence or time dependence (in the times and concentrations range used) are observed. The encapsulation of RayND-M with protein albumin (HSA) can additionally reduce the low cytotoxicity of RayND-M to a negligible level. Albumin adsorption shown previously as a factor increasing the ND biocompatibility, not only provides the safe interaction with a biological target, but also helps to control the ND deaggregation and dispersibility. In addition, the ND encapsulation with albumin enables controlling the desired interaction with surface-attached drugs, which is highly important for the development of new reliable nanoparticles-based drug delivery systems.

With these observations, magnetic ND was shown very promising for bioapplications due to its low toxicity and to fluorescence properties that provide possibilities for multimodal imaging combining excitations in very wide range, including near-IR, and different methods of signal detection. Note that observed fluorescence properties of RayND-M are significantly more convenient for bioimaging as compared with other nonmodified NDs (such as DND). And additional new facilities for the RayND-M multifunctional use are opened due to the significant magnetism of this ND. In the following, we consider some options and demonstrate with preliminary experiments and discuss the possibilities of the applications development.

3.3 RayND-M Magnetic Properties Applications

The possibility of the RayND-M guided with magnetic field was demonstrated using a nonuniform magnetic field from a magnetized steel needle of diameter 0.18 mm, magnetized with a strong permanent magnet. The RayND-M particles were suspended in the bidistilled water. When the needle was immersed in the suspension, the ferromagnetic particles were attracted to
the region of the magnetic field gradient as a result of magnetophoresis.\textsuperscript{30,31} Figure 6(a) shows the scheme of this simple experiment. The moving of particles and their aggregates in the needle’s magnetic field were observed with an optical microscope and with lifetime imaging, respectively. The optical image in Fig. 6(b-1) was obtained at the beginning of the experiment; Fig. 6(b-2) is the FLIM of the particles attracted to the needle; and Fig. 6(b-3) shows the microscopic image of the needle with attracted particles just after FLIM measurement. This series of images show that particles in water or in a medium can be moved by magnetic field and in principle can be used for developing methods of drug delivery.

The motion of the water- or medium-suspended particles driven by a magnetic field can be used in controlled dispersion of therapeutic agents within liquid- and gel-filled cavities of an organism. Nonspecific delivery and distribution in such a system are driven by diffusion and are limited by the molecule (or NP carrier) concentration and size, surrounding medium properties, such as viscosity. Application of magnetic NP controlled by the external magnetic field provides the possibility to optimize...
the controlled delivery. This scheme [Fig. 6(a)] can also be used to demonstrate the possibility of the magnetic field’s guided cells labeled with RayND-M.

Magnetic labeling of cells provides new possibilities for the remote cells manipulations for various purposes, as well as for investigations including MRI and treatments, such as a local magnetic hyperthermia. The methods of cells labeling with magnetic nanoparticles are developed, among these particles, ND with superbiocompatibility and well-studied pathways of cells penetration also become a very promising candidate when being magnetically modified. The magnetically labeled cells manipulations with magnetic field and cells magnetic separation have been realized in applications such as detection of rare cancer cells in circulating blood, detection of fetal cells in maternal blood, and modification of cellular content of the graft for cellular therapies.

To validate this idea, A549 cells was incubated with RayND-M for 4 h, then detached from the cultivation dish bottom to be suspended in the medium (PBS). Their moving in the magnetic field created by the magnetized needle was observed. The effect of the applied magnetic field on these cells was observed with microscopic imaging and FLIM. The cell attraction evidenced from the cells moving toward the needle is shown in Fig. 6(c). Importantly, the attracted cells [Fig. 6(c)-left] contain RayND-M, as the FLIM shows [Fig. 6(c)-right].

Using magnetic nanoparticles can also be considered in the context of bioengineering. It should be noted that the combining of bioengineering and magnetic fabrication for the purpose of graft engineering was first tried about 30 years ago and references therein. Recently it is continuing to be developed, so the problem of the nanoparticles optimization remains important to overcome some obstacles and limitations. On the other hand, among the NP, the use of ND in bioengineering has been presented. For example, NDs are considered for bone engineering due to their mechanical properties and biocompatibility. Thus, it is sensible to discuss using RayND-M for bioengineering combined with magnetic nanofabrication, MND allows also controlling and visualization of the tissue forming.

For simple estimation of the MND applicability in this capacity, monolayer cell culturing was used. To show the possibility of MND used for the control of cell distribution during the growth of cell culture, the mixture of MND-labeled and nonlabeled A549 cells was recultured in the presence of the local magnetic field. The magnetic field from the neodymium permanent magnet was applied to selected area of the culturing dish with the labeled and nonlabeled cells mixture in a cellular medium. The cells were cultivated together for 12 to 36 h. Then the cell culture was observed with confocal microscopy and FLIM to see how the cells with and without MND distribute in the presence of the local magnetic field. The schematic of the experiment and result is shown in Fig. 7(a). Both confocal...
fluorescence imaging and FLIM show cells with MND localized close to the magnet. Note also that the distribution of RayND-M-labeled cells in the magnetic field could be controlled by the field geometric configuration and parameters of cell cultivation (ND concentration, cocultivation time). This demonstrates the principle to use the magnetic ND to control the cell distribution combined with multimodal imaging; and opens new prospects for its applications in bioengineering allowing simultaneous controlled growth of the tissue, tracking of particles during the process and visualization of obtained structures.

4 Conclusion

In the present work, magnetically modified ND is introduced in terms of its multifunctional bioapplications. Fluorescent ferromagnetic metal-free ND powder (RayND-M) has been obtained by surface modification of ND produced by the LHDP method (the process of the surface modification is not disclosed in this paper). The diamond structure and purity of ND were defined by EDX, Raman, SEM, and the surface was characterized with FTIR; soft ferromagnetism of the ND was detected by a vibrating sample magnetometer. Together with the ferromagnetism of the nanoparticles, the increased fluorescence at one- and two-photon excitation is demonstrated. Note, however, that the origins of the ferromagnetism as well as of enhancement of fluorescence emission in the result of magnetic modification of the ND are still unclear and can be considered together. High biocompatibility of RayND-M was confirmed by MTT analysis using the BHK cell line. RayND-M uptakes and its localization in the cytoplasm similar to other NDs have been observed. The possibility of magnetic guidance of ND in cells and magnetic manipulation of ND containing cells in tissue has been demonstrated by the example of A549 cells cocultured with RayND-M. Strong magnetism, fluorescence, and non-cytotoxicity of the modified magnetic ND enables significant breakthroughs in controlled bioengineering, tissues regeneration, early diagnostics, and targeted drug delivery.

Disclosures

The authors have no relevant financial interests in this article and no potential conflicts of interest to disclose.

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