RESEARCH ARTICLE

Preparation of Lysine-Coated Magnetic Fe₂O₃ Nanoparticles and Influence on Viability of A549 Lung Cancer Cells

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Abstract

Objective: To explore the effect of lysine-coated oxide magnetic nanoparticles (Lys@MNPs) on viability and apoptosis of A549 lung cancer cells. Methods: Transmission electron microscopy (TEM), vibrating sample magnetometer (VSM) and Zeta potentiometric analyzer were employed to characterize Lys@MNPs. Then Lys@MNPs and lung cancer A549 cells were co-cultured to study the effect of Lys@MNPs on cell viability and apoptosis. The pathway of Lys@MNPs entering A549 cells was detected by TEM and cell imaging by 1.5 T MRI. Results: Lys@MNPs were 10.2 nm in grain diameter, characterized by small size, positive charge, and superparamagnetism. Under low-dose concentration of Lys@MNPs (< 40 μg/mL), the survival rate of A549 cells was decreased but remained higher than 95% while under high-dose concentration (100 μg/mL), the survival rate was still higher than 80%, which suggested Lys@MNPs had limited influence on the viability of A549 cells, with good biocompatibility and no induction of apoptosis. Moreover, high affinity for cytomembranes, was demonstrated presenting good imaging effects. Conclusion: Lys@MNPs can be regarded as a good MRI negative contrast agents, with promising prospects in biomedicine.

Keywords: Oxide magnetic nanoparticle - lysine - lung cancer cells - magnetic resonance imaging - cell viability

Introduction

Magnetic nanoparticles (MNPs), which have unique properties of superparamagnetism, alternating current magnetic heating, magnetic quantum tunnel effect and good biocompatibility, is widely applied in the field of biomedical domain such as magnetic resonance imaging (MRI) contrast agent, immunoassay, magnetic heat treatment, drug release and cell sorting (Fang et al., 2012; Sterenczak et al., 2012; Rosen et al., 2012; Barick et al., 2014; Bogart et al., 2014; Singh et al., 2014; Wabler et al., 2014; Weis et al., 2014). In order to achieve better diagnosis and treatment effect, MNPs are modified and constructed to deeply study its biological effect according to different application requirements (Basuki et al., 2013; Lee et al., 2014; Riaz et al., 2014; Sun et al., 2014). MNPs shows great potential in MRI contrast agent, however, there is a limitation in application of MNPs achieving cell imaging in the current study. For example, efficient uptake nanoparticles, effective gathering in the lesions, imaging signal amplification and the improvement of imaging quality are needed to be further studied (Bakhru et al., 2012; Laesen et al., 2012; Huang et al., 2013; Huang et al., 2013; Weis et al., 2014).

Small molecule lysine is one of the important components of protein, which can’t synthesize itself, but it is one of eight very necessary amino acids in human body, and selected as coated material widely used in food and pharmaceutical industries. Small molecule lysine has amino and hydroxyl groups, characterized by no toxicity, no antigenicity, good biocompatibility, biodegradability, promotion of the growth and development, improvement of drug efficacy and so on. This study employed lysine to modify oxide magnetic nanoparticles to explore its application in high quality imaging and its influence on cell viability and apoptosis of A549 lung cancer cells.

Materials and Methods

Preparation and surface characterization of lysine coated magnetic nanoparticles

Lysine coated magnetic iron oxide nanoparticles (γ-Fe₂O₃) was prepared using coprecipitation method according to the literature (Ma et al., 2003) and expressed as Lys@MNPs. The specific modification methods are as follows: Proper amount of lysine was dissolved in a small amount of distilled water and after fully dissolved, rapidly added with a certain amount of magnetic liquid (molar ratio of lysine/iron element is 1: 20) through full ultrasonic dispersion. The total reaction system (100 mL) was mechanically stirred for 4 h under the protection of nitrogen, washed by distilled water repeatedly for removing free lysine molecules after magnetic separation.

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and preserved at 4 °C after constant volume.

Transmission electron microscopy (TEM, JEM-200CX, JEOL, Japan) was used for describing morphological characterization of Lys@MNPs. The method was shown below: Magnetic nanoparticles liquid was dripped into 300-mesh copper net and determined after natural drying. The magnetic properties of the sample were described using vibrating sample magnetometer (VSM 7407, Lakeshore, USA). Properties of the surface charge under physiological conditions (pH 7.4) were described using Zeta potentiometric analyzer.

Cell culture
A549 lung cancer cells were purchased from Shanghai Cell Institute, Chinese Academy of Sciences (CAS). RPMI 1640 culture containing 10% FCS, which was used for cell culture in 5% CO₂ incubator at 37 °C, was changed every two days. Cells were inoculated into 6-well or 96-well plate after being fusion to about 90% and cell growth was in a good state and was used when fusion to about 80%.

Detection of cell viability by MTT
A549 cells with concentration of 1×10⁴ cell/mL were inoculated into 96-well plate, 100 μL/well for 24 h incubation, then added with Lys@MNPs culture solution with different final concentrations of 0, 10, 20, 40, 60, 80 and 100 μg/mL for 24 h co-incubation and washed by PBS after removing culture solution. After MTT solution (20 μL) was added into each well for 4 h incubation in the incubator, 200 μL Dimethyl sulfoxide (DMSO) was added and shaken for 30 min. Biv-tek instruments inc was used to select 590nm in wavelength for calculating optional density (OD). Blank control was designed for color comparisons. Cell survival rate (%) = OD measured value / OD blank control * 100.

Detection of cell apoptosis by FCM
Blank cells and marked Lys@MNPs cells were collected, washed two times with PBS and then suspended by 50 μL buffer solution. The suspension was added with 5 μL Annexin V-FITC at room temperature away from light for 10 min, mixed with 10 μL prodium iodide (PI) at room temperature, kept in dark place for 5 min, and after washed by PBS, added with 300 μL buffer solution for resuspended cells which was detected using flow cytometry (FCM).

Detection of cell section by TEM
After A549 cells and Lys@MNPs were co-incubated for 6 hours and digested by trypsin, the collected cells were centrifuged to remove the supernatant, and resuspended by adding with PBS, then centrifuged twice, fixed by 2.5% glutaraldehyde solution at 4 ℃ for 1 h and then by 1% osmium tetroxide at room temperature for 1 h, and washed by PBS three times, 5 min each time. Ethanol dehydration, epoxy resin embedding and routine ultrathin sections were done (50 nm). Sections were observed by TED (HITACHIIH-600, Japan) after was transferred into 300-mesh cooper net and dyed by uranyl acetate.

Analysis of magnetic resonance imaging (MRI)
After 0.25% trypsin digestion, the collected Lys@MNPs-labeled and unlabeled cells were counted, centrifuged and suspended in Eppendof containing 0.5 mL 1% agarose. MRI of cell suspension was analyzed by 1.5 T MR instrument (Philip Company, Eclipse type) and the parameters included inner diameter of surface loop (12.7 cm), field of vision (FOV, 14 cm×14 cm), number of excitation (NEX, 2~4), layer thickness 2 mm, scan matrix 384×256 and signal strength region of interest (ROI, 20 mm²). Imaging sequence: (1) Spin echo (SE) impulse sequence: T1WI, repetition time (TR) 500.0 ms and echo time (TE) 17.9 ms. (2) Fast spin echo (FSE) impulse sequence: T2WI, TR 4 000 ms or 4 063 ms, TE 108 ms, echo train length 16 or 30. (3) Gradient echo (GrE) impulse sequence: T2WI, TR 620 ms, TE 15.7 ms, flip angle 35°. The known concentration of Lys@MNPs was diluted at equal ratio and suspended in Eppendof tube containing 0.5 mL 1% agarose and 0.5 mL distilled water was set as control group.

Statistical data analysis
The result in the experiment was expressed by x±s and SPSS software package was used for data analysis.

Results
Property characterization of Lys@MNPs
Morphology of Lys@MNPs was characterized by approximate sphere, evenly distributed particle, average grain diameter 10.2 nm (Figure 1a and 1b). Figure 1c was Electron diffraction image, from which nano-particles is
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Table 1. The Viability of A549 Cells at the Different Concentrations and Incubation Time (X±s)

<table>
<thead>
<tr>
<th>Lys@MNPs (μg/mL)</th>
<th>A549 cells viability (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>12 h</td>
</tr>
<tr>
<td>0</td>
<td>100.0±3.1</td>
</tr>
<tr>
<td>10</td>
<td>98.1±8.4</td>
</tr>
<tr>
<td>20</td>
<td>97.9±3.4</td>
</tr>
<tr>
<td>40</td>
<td>97.3±6.2</td>
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<tr>
<td>60</td>
<td>93.7±4.7</td>
</tr>
<tr>
<td>80</td>
<td>88.1±4.3</td>
</tr>
<tr>
<td>100</td>
<td>84.4±3.1</td>
</tr>
</tbody>
</table>

Figure 2. Magnetic Hysteresis Chart of Lys@MNPs

Figure 3. The viability of A549 Cells at the Different Concentrations and Incubation Time

γ-Fe₂O₃ nanoparticles of cubic spinel structure.

Saturation magnetization (Ms) and coercive force (H) are the important technical parameters of characterizing magnetism. From Figure 2, saturation magnetization of bare MNPs was 40.58 emu/g while Lys@MNPs reduced to 36.72 emu/g because nonmagnetic substance on nanoparticle surface increased after lysine modification. Coercive force and residual magnetism tended to be zero, showing they were superparamagnetism. When saturation magnetization was up to 3 000 Oe, it tended to be saturated, paralleled with abscissa axis, showing Lys@MNPs had superparamagnetism, connecting with lysine molecules on nanoparticle surface. Further exploration of surface charge characteristic under physiological condition of pH 7.4 revealed bare MNPs and Lys@MNPs were different in electrical property, -316 mV and 4.8 mV, respectively, because there were positively charged amidogen on MNPs surface after lysine modification.

Cell viability and apoptosis of A549 in vitro

After Lys@MNPs and A549 cells co-cultured for 24 h, the viability of A549 cells at the different concentrations and incubation time was as shown in Table 1, showing the survival rate of A549 cells and Lys@MNPs were related in a dose-depended manner. Under low-dose concentration of Lys@MNPs (<40 μg/mL), the survival rate decreased little but higher than 95% while under high-dose concentration (100 μg/mL), the survival rate decreased but still over 80%, which suggested Lys@MNPs had less influence on the viability of A549 cells, with good biocompatibility (Figure 3).

Meanwhile, Annexin V-FITC/PI was used to detect the effect of Lys@MNPs on the apoptosis of cells. From the results of FCM, Lys@MNPs couldn’t cause the apoptosis of A549 cells when the concentration of Lys@MNPs was less than 0.1 mg/mL after 12 h incubation. Viable non-apoptotic cells in the area of Q3 took up over 90% of the total cell number, viable apoptotic cells in the area of Q4 and non-viable apoptotic and necrotic cells in the area of Q2 accounted for 8% and 0.5%, respectively, which showed Lys@MNPs had good biocompatibility and didn’t cause the apoptosis of A549 cells (Figure 4).

Observation of cell section

TEM proved that the large amount of Lys@MNPs entered A549 cells, with high labeling rate. As shown in Figure 5, Lys@MNPs partially entered late endosomes or lysosomes, marked by the formation of cluster embedded in the cytoplasm but didn’t appear in endoplasmic reticulum, mitochondria, golgi apparatus and nucleus.

Cell imaging

As shown in Figure 6, T2WI signal intensity was
In this study, viable non-apoptotic cells in the area of Q3 took up over 90% of the total cell number, viable apoptotic cells in the area of Q4 and non-viable apoptotic and necrotic cells in the area of Q2 accounted for 8% and 0.5%, respectively, which showed Lys@MNPs had good biocompatibility and didn’t cause the apoptosis of A549 cells.

Lysine has a large number of amidogens which can interact with negative charge on the surface of cytomembrane (Ge et al., 2009; Ding et al., 2014). The uptakes of nanoparticles in the non-macrophage cells was mainly through absorption endocytosis, partly through clathrin-mediated endocytosis instead of passive diffusion and fluid-phase endocytosis (Gupta et al., 2005). Therefore, Lys@MNPs connected with cytomembrane through nonspecific absorption and entered cells through endocytosis.

Lys@MNPs is a negative contrast agent, because the more MNPs enter A549 cells, the stronger T2WI signal intensity and the darker the MRI. It has high affinity to cytomembrane, presenting good imaging effect. After 2 h co-incubation of Lys@MNPs and A549 cells, T2WI signal intensity is obviously enhanced with the increase of cell number, especially in cell number of 1×10^4 and significantly in cell number of 1×10^5. The result shows that Lys@MNPs is considered as a good MRI contrast agent, with the further potential application.

**References**


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