The study on the preparation and characterization of gene-loaded immunomagnetic albumin nanospheres and their anti-cell proliferative effect combined with magnetic fluid hyperthermia on GLC-82 cells

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Abstract: As one of the most common malignant tumors, the clinical and socio-economic consequences of lung cancer are significant. Currently, surgery is the main treatment strategy for this disease, but the survival rates of lung cancer patients are not ideal due to the high recurrence rate of the disease. Therefore, many researchers are exploring new specific therapeutic methods that are highly curative and minimally cytotoxic to healthy tissues. To this end, albumin nanospheres simultaneously were loaded with super-paramagnetic iron oxide nanoparticles (as gene vector and anticancer gene), and plasmid pDONR223-IFNG, and modified with anti-EGFR monoclonal antibody cetuximab as therapy. Targeting agents, namely gene-loaded immunomagnetic albumin nanospheres (cetuximab [C225]-IFNG-IMANS), were prepared for targeted lung carcinoma cells (GLC-82 cell lines). Transmission electron microscopy images showed that the C225-IFNG-IMANS were successfully prepared, and the ability of the nanospheres to target GLC-82 cells in vitro was confirmed by Prussian blue staining, immunofluorescence experiments, and magnetic resonance imaging. Transfection photographs and agarose gel electrophoresis proved that pDONR223-IFNG could be encased in the albumin nanospheres. A Cell Counting Kit-8 assay showed that the combination therapy group had significantly more therapeutic effects on GLC-82 cells than other therapy groups. A flow cytometry assay showed that the apoptotic index of the combined treatment group was 67.68%, whereas the indices of the C225 group, gene therapy group, and magnetic fluid hyperthermia group were 12.2%, 16.34%, and 20.04% respectively. Therefore, the combination of thermal treatment, molecular targeted treatment, and gene treatment synergistically targets GLC-82 cells, and the use of C225-IFNG-IMANS as a gene or drug carrier offers a novel and promising approach for the treatment of lung cancer.

Keywords: targeting gene therapy, nano-drug, magnetic nanoparticles

Introduction

Lung cancer is the most common malignant tumor of the respiratory system. Reports from the World Health Organization (WHO) indicate that the incidence of lung cancer ranks first among various types of malignant tumors. The main treatment strategies for lung cancer, including radiotherapy, chemotherapy, surgery, and biological therapy, have achieved certain curative effects,1 however, the survival rate for lung cancer patients remains low. Therefore, identifying a new method or combination of different methods that have mild side effects and accurately target lung cancer have...
attracted the attention of researchers. In this study, the novel treatment method of combining gene therapy with thermo-
therapy and monoclonal antibody therapy was used to treat lung cancer in vitro.

In recent years, many new gene therapy methods (including immune gene therapy, suicide gene, RNA interference, etc) for tumor treatment have been researched. In the immune gene therapy, IFNγ, IL-2, and TNFα have been applied widely and have a more definite curative effect. The local expression of these genes in tumors can improve the immunogenicity of the tumor, and stimulate and enhance the body’s anti-tumor immune responses. After the tumor cells are killed, the initial trigger action is finished. Immune gene therapy, which can be combined with the body’s immune response, has become an attractive new prospect for lung cancer gene therapy.

However, the problem of gene transfer must be solved to successfully implement comprehensive tumor gene therapy. A viral vector system is the most effective method of gene transfer. However, due to its limitations, such as a lack of an efficient and directional vector system, the problem of controlling genes inside the body and, especially the safety issue, its clinical use is strictly controlled. A non-viral vector system can avoid the significant safety risks, but its transfection efficiency is less than that of the viral vector system. Therefore, the bottleneck of gene transfection has become a pressing matter of gene therapy.

Encouragingly, recently developed nanotechnology has offered a new method for solving the problem of the gene transfection vector. Gene transfer vectors based on nanoparticles have attracted extensive attention due to their large surface area and low toxicity. The use of nano-vectors for gene transfer has many advantages compared with traditional carriers, including the possibility of repeated injections, the slow release of the genes to effectively extend response time, the ability to maintain an effective concentration of the products, and the potential to improve the transfection efficiency and bioavailability of the products. Magnetic nanoparticles (MNPs) of Fe₃O₄ are the most commonly employed MNPs. They have interesting properties including biocompatibility, non-immunogenicity, superparamagnetism, and so on. Moreover, Fe₃O₄ MNPs can absorb energy from the high-frequency alternating magnetic field (AMF) and thus, are useful for therapies based on tumor hyperthermia. This treatment approach is referred to as magnetic fluid hyperthermia (MFH); it specifically targets tumors and is thus more effective for deeply situated tumors.

Molecular targeted therapy (MTT) refers to a new therapy designed to interfere with a specific molecular target or signaling pathway that may play a key role in tumor growth or progression. MTT can inhibit cell proliferation and induce apoptosis by specifically interfering with the signaling pathways of tumor cells, which may constitute an effective treatment strategy for cancer. This approach can more effectively kill cancer cells and reduce the damage to normal tissues. In recent years, the use of MTT as a clinical cancer treatment has gradually increased and shown many advantages over traditional methods. EGFR is expressed in various malignant tumor cells of epidermis origin, such as non-small cell lung cancer (NSCLC), breast cancer, colorectal cancer, gastric cancer, and prostate cancer, etc. EGFR and its ligand can form a dimer to activate tyrosine kinase (TK) and promote tumor cells’ proliferation. Researchers have shown that 40%–50% of NSCLC overexpress EGFR. EGFR-targeted therapies, such as gefitinib, erlotinib, cetuximab (C225), and EMD7200, have been widely employed to treat lung cancer.

The affinity of the EGFR monoclonal antibodies for EGFR is higher than that of natural EGFR ligands. This antibody can be used to specifically interrupt cell pathways mediated by EGFR. Healthy cells express much less EGFR than malignant tissue, so monoclonal antibodies’ role in this type of normal cells are very small. Monoclonal antibodies competitively bind EGFR to inhibit the activation of TK and block the signal transduction pathways associated with tumor cell proliferation. As a result, cell proliferation is inhibited and apoptosis is induced. C225, which binds to the extracellular domain of human EGFR, is a chimeric human/mouse immunoglobulin G1 antibody, which is marketed as Erbitux. C225 has been shown to synergistically affect tumors when combined with other chemotherapy drugs. However, the effectiveness rates of most targeted drugs are approximately 10%. Thus, combination therapy has become an important development direction.

In this study, we prepared the gene-loaded magnetic albumin nanospheres (MANS) and the gene-loaded immunomagnetic albumin nanospheres (IMANS), coupled with C225 monoclonal antibodies. We then evaluated the targeting effect of the gene-loaded IMANS and explored the effectiveness of the combination of this thermal treatment, molecularly targeted treatment, and gene treatment as a new combination therapy for GLC-82 cells.

Materials and methods
Main materials
Agarose was purchased from Sigma-Aldrich Co. (St Louis, MO, USA); 1640 medium and fetal bovine serum were obtained from Gibco (Thermo Fisher Scientific, Waltham, MA, USA); Cell Counting Kit-8 (CCK8) was obtained from

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Preparation and characterization of the nanosized Fe₃O₄ and polyetherimide (PEI)-Fe₃O₄

Fe₃O₄ nanoparticles were prepared by chemical coprecipitation. The procedure steps were as follows.

The solutions of FeCl₃·6H₂O (0.1 mol/L, 100 mL) and FeCl₂·4H₂O (0.1 mol/L, 60 mL) were mixed together under nitrogen purging and stirring (the molar proportion of iron(II) and iron(III) was 0.6). Ammonia (1.5 mol/L) was then dropped into the mixture until its pH reached 9. The dark precipitate appeared in the mixed solution rapidly. The solution was stirred for another 30 minutes. After 30 minutes ripening at 90°C, the dark precipitates were isolated using a permanent magnet and dried under a vacuum. They were then examined by transmission electron microscopy (TEM) (JEM-2100, High Resolution). The diameter and zeta potential of Fe₃O₄ were measured by a dynamic light scattering (DLS) instrument. The Fe₃O₄ nanoparticles were subsequently modified by PEI, which was referred to in a previously described method. The detailed steps are as follows.

First, some Fe₃O₄ nanoparticles were dissolved in deionized water to prepare a 4% magnetic fluid; the supernatant was discarded after ultrasonic dispersion and high-speed centrifugation. Second, the precipitate was resuspended in phosphate-buffered saline (PBS) and ultrasonically dispersed. Third, PEI was added and the mixture was blended in a shaker at a constant temperature for 24 hours. The magnetic particles were separated from the solution using a magnetic method and washed repeatedly with distilled water and methanol. MNPs were dried under a vacuum (namely PEI-Fe₃O₄).

The shape of the PEI-Fe₃O₄ particles was observed by TEM, and the adsorption efficiency of PEI was demonstrated by Fourier transform infrared spectroscopy (560; Thermo Nicolet, Madison, WI, USA). The zeta potential of PEI-Fe₃O₄ was measured by a DLS instrument.

Assessment of the capability of PEI-Fe₃O₄ nanoparticles to bind to plasmid

The concentration of plasmid DNA in each experimental group was 0.01 μg/μL. The nanoparticles and pEGFP plasmid DNA were mixed at the ratios of 0:1, 10:1, 30:1, 50:1, and 100:1 (w/w), and then incubated for 30 minutes at room temperature (RT). The binding efficiency was measured by agarose gel electrophoresis (1% w/v agar, voltage: 90 V, time: 30 minutes). The samples were analyzed using the GeneScan Analysis software.

Preparation and characteristics of gene-loaded IMANS

Gene-loaded MANS were prepared by the desolvation-crosslinking method

The PEI-Fe₃O₄ MNPs and plasmid pDONR223-IFNG/pEGFP suspension of the indicated mass ratios were incubated for 30 minutes, bovine serum albumin (BSA) (100 mg) was then added, and the mixture was stirred. The pH of the mixture was adjusted to 9; ethanol was dropped into the solution at a rate of 1 mL/min, followed by the slow addition of 50 μL of 2.5% glutaraldehyde. After stirring for 24 hours at RT, the solution was washed three times and the organic solvent added during the preparation was removed by high-speed centrifugation to yield the gene-loaded MANS (IFNG-MANS/EGFP-MANS).

Gene-loaded IMANS were prepared by using the heterobifunctional cross-linker N-succinimidyl-3-(2-pyridyldithio)propionate (SPDP) to couple nanospheres and cetuximab monoclonal antibody (C225)

The specific steps were as follows.

An adequate amount of C225 was dissolved in 0.01 M PBS (pH = 7.4), and 20 mM SPDP. Ethanol was added to the solution at a molar ratio of 1:15. After 60 minutes, the mixture was dialyzed with acetate buffer at RT to remove excess SPDP. A sufficient amount of DL-dithiothreitol was added to the pyridyl-dithiopropionate-activated C225 (C225-PDP) solution obtained in the previous step, and the mixture was gently stirred for 30 minutes at RT. The reactant was then dialyzed with PBS to remove redundant DL-dithiothreitol and obtain a sulphydryl-activated C225 solution (C225-PDP-SH).

Second, an appropriate amount of gene-loaded MANS was diluted with acetate buffer, and 50 μL of 20 mM SPDP ethanol solution was added while stirring to remove excess SPDP.

Third, the activated gene-loaded MANS and C225-PDP-SH mixture was incubated and shaken at 4°C for 15 hours. After washing several times by centrifugation,
gene-loaded IMANS (C225-IFNG-IMANS/C225-EGFP-IMANS) were obtained. TEM and DLS were used to characterize the nanospheres.

Cell transfection experiment
Prior to transfection, 4 µg of pEGFP plasmid DNA and PEI-Fe$_3$O$_4$ nanoparticles or C225-EGFP-IMANS were (the mass ratio of PEI-Fe$_3$O$_4$ to DNA was 50:1) each diluted in 250 µL of serum-free media. After incubation at RT for 5 minutes, plasmid DNA was added to the PEI-Fe$_3$O$_4$ nanoparticles or C225-EGFP-IMANS suspensions, and the complexes were incubated for 30 minutes. As a control, commercially available transfection reagents, such as Lipofectamine™ 2000 (Invitrogen; Thermo Fisher Scientific) were used in accordance with the manufacturer’s instructions.

GLC-82 cells and HEK293 cells (control cell line) were seeded in six-well plates, at an initial density of $3 \times 10^5$ cells/well in 2 mL of growth medium. After incubation for 24 hours and the cells had reached 80% confluence, the medium was replaced with 1.5 mL serum-free media and 500 µL of PEI-Fe$_3$O$_4$/pEGFP or C225-EGFP-IMANS complexes, and the cells were then incubated for another 4 hours. The medium was replaced with fresh medium containing serum, and the cells were incubated for another 48 hours. The cells were then observed under inverted fluorescence microscopy (TE2000; Nikon Corporation, Tokyo, Japan). The cells were then collected and examined by a flow cytometry instrument to determine the transfection efficiency. All cell studies were approved by the ethics committee of Southeast University, Nanjing, People’s Republic of China.

Targeting evaluation of gene-loaded MANS in vitro
GLC-82 cells were incubated with C225-targeting groups (C225-IFNG-IMANS), and non-C225-targeting groups (IFNG-MANS) for 24 hours. Prussian blue staining, immunofluorescence experiments, and magnetic resonance imaging (MRI) were used to observe the specific binding ability of the immune nanospheres with human lung adenocarcinoma cells in vitro. The HL7702 cell line does not express EGFR and was used as a control.

Prussian blue staining
Aliquots (1 mL) of GLC-82/HL7702 cells ($3 \times 10^5$/mL) in complete 1640 medium supplemented with 10% fetal calf serum (Gibco) were seeded into 12-well plates and cultured at 37°C in 5% CO$_2$ for 24 hours. C225-IFNG-IMANS and IFNG-MANS were added to five wells each, and the cells were incubated for 24 hours. The cells were then washed with PBS and stained with Prussian blue to observe intracellular Fe$_3$O$_4$ nanoparticles. The combination of nanospheres and cells can be observed by light microscopy.

Immunofluorescence experiments
The C225-IFNG-IMANS (immuno-nanospheres) were dissolved in PBS buffer containing serum, and incubated with rabbit anti-mouse IgG labeled with Alexa Fluor 488 for 30 minutes (4°C). The control group was the IFNG-MANS incubated with the IgG. The nanospheres labeled with Alexa Fluor 488 were then incubated with the lung cancer GLC-82 cell lines, which were fixed with cold acetone. After 24 hours, the incubated cells were washed three times with PBS, and the nucleus were stained with 4′,6-diamidino-2-phenylindole. A fluorescence microscope was used to observe the combination of nanospheres and cells. HL7702 cells served as a control for the above groups.

MRI in vitro
GLC-82/HL7702 cells ($3 \times 10^5$/mL) in complete 1640 medium supplemented with 10% fetal calf serum were seeded into six-well plates at 37°C and cultured in 5% CO$_2$ for 24 hours. C225-IFNG-IMANS or IFNG-MANS were added to the cells, which were then incubated for 24 hours. The cells were digested with 0.25% trypsin and collected and then resuspended in Eppendorf (EP) tubes with 0.5 mL 1% agarose after centrifugation. An EP tube with agarose was used as a control. The EP tubes were scanned with 7.0 Tesla small animals MRI. The T2 values were measured in multi-slice, multi-echo-T2-map sequences. The specific parameters were as follows: 3.0 cm diameter body coil, 5×5 cm field of view, 1 mm thick, 256×256 matrix, TR: 2,000 ms, and TE: 30 ms.

Heating test of Fe$_3$O$_4$ MNPs and C225-IFNG-IMANS
Various doses of Fe$_3$O$_4$ nanoparticles were dispersed in 5 mL 0.9% NaCl, at Fe$_3$O$_4$ concentrations of 0.5, 1.0, 1.5, 2.0, 2.5, and 3.0 mg/mL, and 2 mL aliquots of magnetic fluids were added to flat-bottomed cuvettes. These cuvettes were placed at the center of the hyperthermia coil of high-frequency electromagnetic field (SP-04C, Shenzhen, People’s Republic of China). The output frequency was 230 kHz and the output current was 25 A. The temperature was measured every 5 minutes during 1 hour of incubation.

C225-IFNG-IMANS containing different amounts of iron were tested according to the above steps. The concentrations...
of iron in the C225-IFNG-IMANS were 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875, and 1.0 mg/mL.

The therapeutic effects of C225-IFNG-IMANS combined with MFH on cultured lung carcinoma GLC-82

CCK8 assay

GLC-82 cells (4×10⁴) were cultured in three 96-well plates. After 24 hours incubation, the cells were subjected to eight different conditions: negative control group (1640 containing 10% fetal calf serum), C225 group (C225 solution), gene therapy group (gene-loaded albumin nanospheres [ANS], IFNG-ANS), C225 targeted gene therapy group (C225-IFNG-IANS), MFH group (MANS), C225 targeted MFH group (C225-IMANS), gene therapy and MFH combination group (IFNG-MANS), C225 targeted gene therapy and MFH combination group (C225-IFNG-IMANS). After incubation for 24 hours, the 96-well plates of magnetic targeting and double targeting groups were placed on a 96-well strong magnet (Magneto FACTOR plate, Chemicell, Berlin, Germany). The 96-well culture plates of the MFH groups were placed on a high-frequency AMF (f=230 kHz, I=30A) to be heated for 1 hour. All groups continued to be incubated for 24 hours, 48 hours, and 72 hours, and 10 µL of CCK8 solution was then added to each well, and the cells were further incubated at 37°C for 1–4 hours. The optical density (OD) was read at 450 nm using a spectrophotometer (Infinite 50, TECAN, Männedorf, Switzerland). The following formula was used to calculate the cell relative growth rate (RGR):

\[
RGR = \frac{\text{The experimental group OD} - \text{The blank control group OD}}{\text{The negative control group OD} - \text{The blank control group}} \times 100\% \tag{1}
\]

Flow cytometry assay

The GLC-82 cells in logarithmic growth phase were adjusted to the suspension with 3×10⁶ cells/mL. The cells were grouped and handled as described in the CCK8 assay section. After incubation for 48 hours, the cells from all eight groups were collected and rinsed three times with cold PBS, and resuspended in 1× binding buffer at a concentration of 1×10⁶ cells/mL. Subsequently, 100 µL of the solution was transferred into eight 5 mL culture tubes, and 5 µL of fluorescein isothiocyanate Annexin V and 5 µL of propidium iodide (PI) was added to each tube. After gentle blending and incubation for 15 minutes at RT (25°C) in the dark, 400 µL of 1× binding buffer was added to each tube. The tubes were then analyzed by flow cytometry within 1 hour.

Statistical analysis

Values were indicated as mean ± standard deviation. The data were analyzed with the SPSS 16.0 program. A P-value of <0.05 was considered to indicate significant differences.

Results and discussion

Preparation and characterization of the nanosized Fe₃O₄ and PEI-Fe₃O₄

Figure 1A shows the TEM photos of Fe₃O₄ nanoparticles. Under TEM, the Fe₃O₄ nanoparticles showed high electron density. Their diameter was approximately 20 nm. From the DLS results, we determined that the hydrodynamic size of Fe₃O₄ was 64.87 nm and the polydispersity index was 0.152 (Figure 1B). A problem we found was that the size of MNPs detected by DLS was larger than that observed under TEM. The reason is that the DLS measurement shows the hydrodynamic size of the nanoparticles. MNP aggregation is very easy, so the hydrodynamic size of the MNPs is larger than their actual size.

MNPs have recently emerged as a fundamental tool for biomedical technology.24 These advanced nanomaterials are used in key areas of biomedicine due to advantages of their large-scale synthesis25 and complex surface functionalization, including drug/gene therapy,26 magnetic targeting,27 and diagnostic imaging (as contrast enhancers).28

However, Fe₃O₄ nanoparticles cannot directly carry plasmid and require surface modification to deliver plasmids to cells. The monomer (-CH-CH₂-NH₂-) PEI, a common powder surface modification agent, can bind to DNA and adhere to cells, and exerts electrostatic repulsion and steric hindrance effects.29-31 In our previous study, MnₓZnᵧFe₃O₄ nanoparticles were successfully surface-modified by PEI, and used to transfect pEgr1-HSV-TK/GCV into cancer cells to treat hepatoma.32

In this study, we prepared PEI-Fe₃O₄ nanoparticles whose shape and size were similar to those of Fe₃O₄ nanoparticles (Figure 2A). Fourier transform infrared spectroscopy indicated specific peaks at 3,043.6 cm⁻¹, 2,950.1 cm⁻¹, 1,464.8 cm⁻¹, which were due to the chemical structure of the PEI polymer coating on the MNPs (Figure 2B). The surface charge of Fe₃O₄ in pH 7 was ±0.5 mV and turned into 36.3±0.6 mV after being modified with PEI (Figure 2C).
Figure 1. Characterization of Fe$_3$O$_4$.

Notes: (A) TEM of the Fe$_3$O$_4$ nanoparticles that were high electron density with a diameter of approximately 20 nm. (B) Hydrodynamic diameter distribution of Fe$_3$O$_4$ nanoparticles in PBS as measured by a particle size analyzer.

Abbreviations: TEM, transmission electron microscopy; PDI, polydispersity index; PBS, phosphate-buffered saline.

Figure 2. Characterization of PEI-Fe$_3$O$_4$ nanoparticles.

Notes: (A) TEM of the PEI-Fe$_3$O$_4$ nanoparticles whose shape and size were similar to those of the Fe$_3$O$_4$ nanoparticles. (B) FTIR spectra of PEI-Fe$_3$O$_4$ magnetic nanoparticles. The black curve is the Fe$_3$O$_4$, and the red curve is the PEI-Fe$_3$O$_4$. Three blue arrows respectively indicate three characteristic peaks of PEI located at 3,043.6 cm$^{-1}$, 2,950.1 cm$^{-1}$, and 1,464.8 cm$^{-1}$. (C) The zeta potential of Fe$_3$O$_4$ and PEI-Fe$_3$O$_4$ in pH 7. The surface charge of Fe$_3$O$_4$ was 0±0.5 mV and turned into 36.3±0.6 mV after being modified with PEI.

Abbreviations: TEM, transmission electron microscopy; PEI, polyetherimide; FTIR, Fourier transform infrared spectroscopy.
Assessment of the capability of PEI-Fe₃O₄ nanoparticles to be combined with plasmid

Agarose gel electrophoresis (Figure 3) showed that DNA could be effectively bound to Fe₃O₄ MNPs modified with PEI. When the ratio of PEI-Fe₃O₄ and plasmids was 50:1, no obvious DNA strips were seen, whereas obvious DNA strips were seen in agarose gel when the ratios were 0:1, 10:1, 30:1, and the band in the lane with 30:1 was significantly weak. These results suggested that PEI-Fe₃O₄ could be combined with all plasmids in the system at the ratio of at least 50:1 and beyond, and 50:1 was the optimal ratio.

Preparation and characteristics detection of gene-loaded IMANS (C225-IFNG-IMANS)

TEM analysis showed that the self-prepared C225-IFNG-IMANS were approximately spherical and uniform in size. The magnetite materials with a high electron density were incorporated well in the cores of ANS (Figure 4A). DLS showed that the hydrodynamic size of the IFNG-IMANS was 189.5±2.4 nm and the polydispersity index was 0.005 (Figure 4B). The hydrodynamic size of C225-IFNG-IMANS was 211.9±3.1 nm and the polydispersity index was 0.201 (Figure 4C). The surface charge of IFNG-MANS was -37.9±0.4 mV and slightly reduced to -40.2±0.7 mV for C225-IFNG-IMANS (Figure 4D).
The surface charge of IFNG-MANS was 37.9±0.4 mV and slightly reduced to 40.2±0.7 mV for C225-IFNG-IMANS. MANS are ANS that are wrapped with MNPs. ANS are endogenous carriers that are free of the hemolytic or immunogenic problems of albumin. Therefore, ANS may be highly efficient and safe for drug and gene delivery. Many researchers have successfully applied albumin as a drug-delivery system. Additionally, albumin paclitaxel nanospheres (Abraxane) was approved by the US Food and Drug Administration in 2005 for the treatment of metastatic breast cancer based on extensive preclinical and clinical investigations, and this drug has recently been approved for the treatment of NSCLC. ANS have also been successfully used as a gene delivery system. Abbasi et al have used ANS to successfully deliver siRNA into MCF-7 breast cancer cells.

**Cell transfection experiment**

Transfection experiments showed that PEI-Fe$_3$O$_4$ nanoparticles and C225-EGFR-IMANS can effectively deliver plasmid pEGFP into GLC-82 cells and HEK293 cells, which express GFP well (Figure 5A). The transfection efficiency of the PEI-Fe$_3$O$_4$ and Lipofectamine of the HEK293 group were

![Figure 5](https://www.dovepress.com/)

*Figure 5 (Continued)*
the same as that of the GLC-82 group, and the transfection efficiency of the C225-IFNG-IMANS of the GLC-82 group was higher than that of the HEK293 group (Figure 5B).

Gene therapy using nanocarriers is a promising new tumor treatment strategy. Tumor gene therapy using viral and non-viral vectors has been intensively studied, but non-viral gene therapy has been largely ignored in the past due to its inefficient gene delivery, toxicity, and short-term gene expression. However, recent advances in nanoparticle technology and in the genetic material to be transferred are dramatically changing the popularity of this gene transfer methodology. MNPs have been recently developed as a new non-viral gene vector and have shown satisfactory biocompatibility and a large specific surface area.\textsuperscript{37,38} Therefore, MNPs can carry foreign DNA into cells. Because nanoparticles are bound to DNA via electrostatic attraction, the surface of Fe\textsubscript{3}O\textsubscript{4}-MNPs was coated with the positively charged PEI in our experiment, which can better absorb the negatively charged DNA molecules and effectively carry the pEGFR expression plasmid to GLC-82 cells.

**Targeting evaluation of C225-IFNG-IMANS in vitro**

**Prussian blue staining**

Prussian blue staining demonstrated that the cellular iron density (blue particles) was higher in GLC-82 cells that were incubated with C225-IFNG-IMANS (Figure 6A) whereas iron particles were not observed obviously within the cells incubated with IFNG-IMANS (Figure 6B). Because HL7702 cells do not express EGFR, neither the non-C225-targeting groups (Figure 6C) nor C225-targeting groups (Figure 6D) displayed obvious iron particles.

**Immunofluorescence experiments**

Immunofluorescence staining showed that the green fluorescence in the GLC-82 cells of the C225-targeting groups, but not in the non-C225-targeting groups (Figure 7). The C225-targeting groups and non-C225-targeting groups (Figure 7) of the HL7702 cells did not show green fluorescence.
MRI in vitro
The MRI T2 signal intensity of GLC-82 cells significantly decreased after incubation with C225-IFNG-IMANS, but did not markedly decrease after incubation with IFNG-IMANS. The signal intensity of HL7702 cells, which do not express EGFR, did not change in either of the groups (Figure 8A). The T2 relaxation time of the C225-targeting groups was shorter than the non-C225-targeting groups for GLC-82 cells, but did not differ for HL7702 cells (Figure 8B) (*P*<0.05).

In this study, we assessed the targeting of C225-IFNG-IMANS to GLC-82 cells by conducting the above experiments in vitro. Retaining the specific immunoreactivity of monoclonal antibodies is very important when conjugating the monoclonal antibodies to the surface of nanoparticles. Lee et al. prepared methotrexate-loaded immuno-microspheres and evaluated their targeting of an antigen-positive human leukemia cell line. BSA-NS coated with monoclonal antibodies were more readily trapped in the tumor tissue of Lewis lung carcinoma-bearing mice than BSA-nanospheres coated with mouse IgG at 24 hours after injection. Therefore, the targeting ability of the monoclonal antibodies was retained after they were conjugated to the MANS.

Heating test of Fe$_3$O$_4$ MNPs and C225-IFNG-IMANS
The thermodynamic test of Fe$_3$O$_4$ MNPs in vitro showed that when placed in AMF (200 kHz, 20 A), the corresponding magnetic fluid with different concentrations could rise to a steady temperature ranging from 39°C to 56°C within 50 minutes, at which point the temperature of the solution remained stable (Figure 9A).

Figure 9B shows the heating curve of the C225-IFNG-IMANS at different concentrations. Exposure to the magnetic field rapidly warmed fluids of all concentrations to a stable temperature. The concentration of the solution positively correlated with the maximum temperature. Among the tested concentrations, the 0.25 mg/mL solution rapidly warmed within 5 minutes, and then gradually stabilized at a temperature of approximately 44°C, which is appropriate for tumor treatment but does not harm normal tissues. Thus, 0.25 mg/mL was selected for subsequent magnetic induction hyperthermia experiments. A comparison of Figure 8A and C225-targeting groups of GLC-82 cells. (B) C225-targeting groups of GLC-82 cells. (C) Non-C225-targeting groups of HL7702 cells. (D) C225-targeting groups of HL7702 cells.

Abbreviation: C225, cetuximab.
and B, showed the albumin. Coating did not affect the heating capacity of Fe$_3$O$_4$ nanoparticles.

**The therapeutic effects on GLC-82 cells CCK8 assay**

The cell RGRs of eight groups after 24 hours, 48 hours, and 72 hours are shown in Figure 10. The experimental and negative control groups significantly differed ($P<0.05$). Compared with the single therapy groups and the other two therapy groups, the nanosized C225-IFNG-IMANS complex combined with MFH could more significantly inhibit the proliferation of GLC-82 cells ($P<0.05$).

**Flow cytometry assay**

GLC-82 cells were stained with Annexin V and PI and subjected to flow cytometry to detect the ability of various treatments to induce apoptosis.

Generally, cells typically have an intact cell membrane that cannot be stained by PI. However, the externalization of phosphatidylserine changes the cell membrane (membrane

**Figure 7** The specific interaction between cells and C225-IFNG-IMANS.

**Notes:** (A) GLC-82 cells immunofluorescence experiments: a, C225 targeting groups; b, non-C225 targeting groups. 1, DAPI fluorescence; 2, Alexa Fluor 488 fluorescence; 3, merged fluorescence (1 and 2). (B) HL7702 cells immunofluorescence experiments: a, C225 targeting groups; b, non-C225 targeting groups. 1, DAPI fluorescence; 2, Alexa Fluor 488 fluorescence; 3, merged fluorescence (1 and 2).

**Abbreviations:** DAPI, 4′,6-diamidino-2-phenylindole; C225, cetuximab.
Figure 8. In vitro Mri.

Notes: (A) In vitro MRI of GLC-82 and HL7702 cells after treatment with different probe groups. (B) T2 relaxation time of GLC-82 and HL7702 cells after treatment with different probe groups (* indicates P<0.05).

Abbreviations: Mri, magnetic resonance imaging; C225, cetuximab.

Figure 9. Heating test of Fe₃O₄ nanoparticles and C225-IFNG-IMANS in vitro.

Notes: (A) Heating test of Fe₃O₄ nanoparticles fluid in AMF in vitro. The concentration of Fe₃O₄ is 3.0, 2.5, 2.0, 1.5, 1.0, and 0.5 mg/mL. (B) Heating test of C225-IFNG-IMANS fluid in AMF in vitro. The concentration of Fe in albumin nanospheres was 1.0, 0.725, 0.5, 0.375, 0.25, and 0.125 mg/mL.

Abbreviations: IMANS, immunomagnetic albumin nanospheres; AMF, alternating magnetic field; C225, cetuximab.
phospholipids) during the early stages of apoptosis, which can be detected by Annexin V, but not PI due to the intact cell membrane. The membrane ruptures during late apoptosis, and PI can infiltrate the cells. Necrotic cells can be differentiated from apoptotic cells because they lose their membrane integrity, and their phosphatidylserine degrades. Cells positive for only PI are necrotic, whereas those positive for Annexin V and PI are in the late stages of apoptosis and those only positive for Annexin V are in the early stages of apoptosis.

Flow cytometry revealed apoptotic GLC-82 cells in all the experimental groups. The apoptotic indices were 12.2% in the free C225 group, 16.34% in the gene therapy group, 33.88% in the C225-targeted gene therapy group, 20.04% in the MFH group, 39.71% in the C225-targeted MFH group, 39.2% in the gene therapy and MFH combination group, 67.68% in the C225-targeted gene therapy and MFH combination group, and 0.5% in the negative control group (Figure 11).

The results of the CCK8 and flow cytometry assay both suggested that the novel approaches for NSCLC in combination with hyperthermia, gene therapy, and molecularly targeted therapy may be a potential method to treat tumors. However future studies should address remaining challenges, such as the targeting ability of C225-IFNG-IMANS in vivo, the molecular analysis of IFNG gene expression and the treatment effects of C225-IFNG-IMANS in vivo.

Studies in recent years have shown that single treatments are limited and not able to cure cancer alone. Multimodal therapy had been used more and more and exhibited more effectiveness in tumor treatment than single modality treatment. Combination treatments usually show synergistic effects which may observably improve therapeutic effects while reducing the side effects of each single treatment modality. In our study, the tumor-eliminating rate of MFH was greatly enhanced when combined with gene therapy and molecular targeting treatment. However, the mechanism of the combined treatment is inconclusive and the possibilities may be as follows. In our study, we found that with the anti-EGFR monoclonal antibody C225, either gene therapy or hyperthermia therapy brought more significant inhibition of growth and induction of apoptosis in GLC-82 cells than those groups without C225. The results showed that C225 coated MANS did not just have selective targeting of the expression of EGFR cells, but could also inhibit proliferation of lung cancer cells. The product of pDONR223-IFNG is INFγ. INFγ exerts direct anti-tumor effects by inhibiting tumor cell proliferation, inducing differentiation, and promoting apoptosis and exerts indirect anti-tumor effects by activating macrophages and natural killer cells, inducing the expression of major histocompatibility complex class I and class II molecules, enhancing the T cell-mediated death of tumor cells and inhibiting tumor angiogenesis. Hyperthermia in combination with other treatments was thought to affect sensitivity of tumors to other treatments mainly through microenvironment factors. In hyperthermia treatments, the tumor tissue is heated above 42°C, the integrity of the cell membrane is damaged, and the penetrability increased, which causes cellular swelling. Hyperthermia also restrains the repair of DNA damage by
Figure 11 The apoptosis and necrosis of GLC-82 cells analyzed by flow cytometry after different treatments.

Notes: (A) Negative control group; (B) free C225 group (C225 solution); (C) gene therapy group (gene-loaded albumin nanospheres, IFNG-ANS); (D) C225-targeted gene therapy group (C225-iFNG-ANS); (E) MFh group (MANS); (F) C225-targeted MFh group (C225-MANS); (G) gene therapy and MFh combination group (iFNG-MANS); (H) C225-targeted gene therapy and MFh combination group (C225-iFNG-MANS).

Abbreviations: ANS, albumin nanospheres; PI, propidium iodide; MANS, magnetic albumin nanospheres; iMANS, immunomagnetic albumin nanospheres; MFh, magnetic fluid hyperthermia; C225, cetuximab; FITC, fluorescein isothiocyanate; IANS, albumin immuno-nanospheres.
gene therapy. Lastly, heating had direct cytotoxic effects or induced tumor cells’ apoptosis, which contributed to the therapeutic effects.

Conclusion
In this study, the PEI-Fe₂O₃ MNPs were used as a novel gene vector and C225-IFNG-IMANS were successfully prepared. The ability of C225-IFNG-IMANS to target GLC-82 cells was evaluated. A CCK8 test and flow cytometry assay showed that the combination therapy was markedly superior to all tested monotherapies. We explored a new multiple therapy approach for lung cancer by combining thermal treatment, molecular targeted treatment, and gene treatments.

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Disclosure
The authors report no conflicts of interest in this work.

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