

ORIGINAL RESEARCH

Egr I/HSP70 Promoter-Driven Activation of Gene Expression for Synergistic Anti-Hepatoma Using PEI-MZF Nanoparticles and Radiation

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Dongsheng Zhang Imaging and Nuclear Medicine, Southeast University, No. 87 Dingjiaqiao Road, Nanjing, 210000, Jiangsu, People's Republic of China Tel +86-25-83272538 Email zdszds1222@163.com **Introduction:** Spatially restricted gene expression circumvents the gene expression and gene vector problem by enabling localized amplification. The objective of this study is to construct a spatially restricted gene expression for liver cancer therapy based upon the MFH-absorbing properties of PEI- $Mn_{0.5}zn_{0.5}Fe_2o_4$, gene therapy and radiation.

Methods: Mn_{0.5}zn_{0.5}Fe₂O₄ (MZF) magnetic nanoparticles were prepared by an improved chemical co-precipitation method, modified by polyethylene imine (PEI), and then the structure, modification characters, biocompatibility, temperature rise and control ability and binding efficiency of the plasmid were characterized. Then, the dual-promoter plasmid PCDNA3.1-EGR1-HSP70-HSVTK was constructed. The recombinant vectors were identified by enzyme digestion analysis and DNA sequencing. The TK gene expression level was detected by realtime-PCR assay in HEK293 cells. Also, the HSV-TK gene expression was detected in SMMC7721 cells with the help of PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄. In vitro anti-tumor experiment, MTT assay and flow cytometry were used to evaluate the therapeutic effects of the cultured SMMC7721 cells treated by different ways. In vivo anti-tumor experiment, the xenografted mice were treated by different ways for three times to detect the antitumor effect.

Results: The Mn_{0.5}Zn_{0.5}Fe₂O₄ magnetic nanoparticles could be successfully prepared through improved co-precipitation process and showed good biocompatibility. And PEI had been coated on MZF complex. The modified PEI-MZF presented favorable dispensability, responsibility to magnetism, good loading capability and transfect capability. Also, pCDNA3.1-Egr1-Hsp70-HSVTK plasmid had been constructed successfully and could be induced by heat and irradiation. It would be used for further target gene therapy research. The antitumor results in vitro showed: The therapeutic effects of nanosized PEI-MZF-HSV-TK complex could significantly inhibit the proliferation of cultured liver cancer cells (SMC7721), induce cell apoptosis and had a prominent cell cycle disturbance in the S phase in vitro. The results in vivo showed: The combined therapy induced by PEI-MZF-HSV-TK could inhabit the growth of hepatocellular carcinoma xenografts by killing and inhabiting the proliferation of the tumor cells.

Conclusion: The novel site-directed heat/radiation-inducible expression system based upon the hyperthermia (by MFH) and radiation possessed superior antitumor effect in vivo and in vitro.

Keywords: radiation, magnet-induced heating, dual sensitive, magnetic nano vector

Introduction

Traditional treatment of cancer with gene therapy suffers lack of selectivity and specificity even causes some toxic side-effects. Several strategies have been developed, which allow control of both spatial and temporal expression of transgenes. Spatial resolution is often achieved by using tissue- or cell-specific

promoters, or exploitation of the inducible response. Many of these promoters have been developed and applied to the research, whose expression is regulated rigorously by the simple and cheap way. Like early growth response-1 (Egr1) promoter, first described more than two decades ago, is one of the most widely investigated radiationinducible promoters. The researchers find radio-genetic therapy is a good way of combination of gene treatment and radiation. It's effective at a lower radiation dose of the tumor and helpful at a less damage of the normal tissue. Meanwhile, the target gene could express according to their needs.^{2,3} But radio-genetic therapy is not so satisfied in the research, for its short and lower expression efficiency. At the same time, the radiation induction activity is not so high because of the hypoxia inside the tumor.^{4,5} Thermotherapy could improve the hypoxia inside the tumor and boots the radiation therapeutic effects.^{6,7}

Heat shock has been suggested as a non-invasive and non-toxic alternative to induce gene expression locally using a heat-inducible promoter. For example, Tang et al have demonstrated that a hetero gene with hsp70 promoter could be elevated to 10 to 500-fold over background by magnetic heat (43 °C). Isomoto et al reported hsp 70B gene promoter-oriented HSV-tk (HSP-tk)/ganciclovir (GCV) system directed by heat shock significantly inhibited the growth of subcutaneous tumors and prolonged survival of mice with peritoneal carcinomatosis. Thermo resolution may be controlled by external cues, such as frequency magnetic field. Induction of heat shock promoter (HSP)-mediated gene expression by frequency magnetic field is a promising approach for achieving temporal and spatial control of gene expression. 10,12,13

Although many inducible promoters have been developed, there are limitations. Hence, more effective and continual gene expression should be established, preferably with the advantages of different induced methods. The dual-promoter regulation system offering a new concept for gene expression with high efficiency. While there are a lot of evidence that the same gene expression regulated by double promoters in the nature, ¹⁴ but few people used it for the expression of exogenous gene in the research. Some scholars have found that increased the number of the promoters could improve the expression level of target gene. The most common approach for solving this problem is two single promoter in series to form a dual promoter system. ^{15–19} Finn et al studies showed that the dual promoter can effectively increase the gene

expression level with non-viral vector.²⁰ Thus it is more important in clinical application.

Based on the above considerations, we design an Hsp70 and Egr1 promoters-driven eukaryotic expression vector pCDNA3.1-Egr1-Hsp70-HSVTK to increase and control the gene expression mediated by the self-made PEI-Mn_{0.5}Zn_{0.5}Fe₃O₄ magnetic gene delivery system. The magnetic nanoparticles can absorb energy from the high frequency alternating magnetic field (AMF) to warm up. When reaching their curie temperature, they will stop absorbing the energy and the temperature will keep. Once the alternating magnetic field is removed, the temperature will drop. The curie temperature of the self-made nanoparticles is 42-44°C, which is the temperature required for hyperthermia. Thus, we intend to use the temperaturesensitive PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ nanoparticles as the vector of pEgr-1-Hsp70-HSV-TK gene and as a link to construct an anti-tumor gene/magnetic composite nano-system (pEgr1-Hsp70-HSV-TK/GCV/PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄), organically combines gene therapy, hyperthermia, and nuclide therapy. It not only solves the problem of gene transfer vector, automatic heating, temperature control and targeted hyperthermia but also uses Egr1 and Hsp70 dual promoters to solve the expression of target genes in tissues and timing. Furthermore, synergistic therapy of two or more therapies may significantly improve efficacy and reduce adverse reactions through synergistic enhancement of the interactions.²¹

Materials and Methods

Materials

pCDNA3.1-EGFP-C1 (4.7 kb), encoding a green fluorescent protein, driven by an immediate early promoter of CMV, was purchased from Clontech Laboratories (Mountain View, California, USA); pD3SX-HSP70-HSVTK, mammalian expression vector, encoding HSVTK, driven by human Hsp70 promoter was purchased from Stressgen bioreagents corps (Canada); Polyethylenimine (PEI; averageMW25 kDa, average degree of polymerisation 580), and most of the reagents were purchased from Sigma Aldrich (St. Louis, MD, USA);HEK293 cells and SMC7721 cells were purchased from Shanghai Institute of Biochemistry and Cell Biology, Chinese Academy of Sciences (Shanghai, China). Magneto FACTOR plate was purchased from Chemicell (Berlin, Germany). The plasmids were propagated in Escherichia coli, extracted by the alkaline lysis technique, and purified by a QIAGENR kit (Hilden, German). The purity of the

plasmids consisting of supercoiled and open circular forms was checked by electro-phoresis on a 1% agarose gel, and the concentration of DNA was determined by measuring UV absorbance at 260 nm. The purity (LPS) of plasmid is 0.1EU/g DNA and the amount of supercoiled DNA is 90%.

Preparation and Characterization of PEl- $Mn_{0.5}Zn_{0.5}Fe_2O_4$ Nanoparticles

Mn-Zn ferrite of composition $Mn_{0.5}Zn_{0.5}Fe_2O_4$ was prepared by the precipitation method (for details see 7), and PEI-coated $Mn_{0.5}Zn_{0.5}Fe_2O_4$ (PEI-MZF) was prepared by the method described before. They were characterized by HREM (High Resolution Electron Microscopy), TEM (Transmission Electron Microscope), FTIR (Fourier Transform Infrared Spectroscopy) and XRD (X-Ray Diffraction).

Toxicity of PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ Nanoparticles

PEI Cytotoxicity Test (MTT)

PEI-MZF-NPs magnetic nanoparticles were sterilized by low-temperature batch sterilization (60°C10h, 37°C12h, repeated 3 times). L929 cells were seeded in 96-well plates, and they were co-incubated with PEI-MZF-NPs/ DNA complexes or PEI-DNA complexes with different N/ P ratios (1, 3, 5, 7, 9) and (11) (each well contains (0.5)µg) plasmid DNA). The negative group was only 1640 culture medium, and the positive control was added with 0.7% polyacrylamide monomer solution. There were 8 wells in each group. Forty-eight hours later, 20µL MTT was added onto the cells, and were incubated for 4 hours. Then the liquid was discarded. 150µL/well DMSO was added in, then the mixture were shook for 10 minutes to measure the absorbance value at 493nm on the immunoassay reader (Note: There were different reports in the literature about the test wavelengths required for the maximum absorbent value. The purple solution formed by the solution of MTT crystallization with DMSO as a solvent, the maximum light absorption wavelength was about 490nm-515nm, and 493nm was taken in this experiment). Cell viability (%)=average OD of experimental group/average OD of negative group × 100%.

Hemolysis Test

Ten milliliters of New Zealand rabbit blood was extracted, then 0.5mL of 20g/L potassium oxalate was added in, and 10mL of saline per 8mL was added in to obtain diluted anti-coagulated rabbit blood. 0.2mL rabbit blood diluent was added into 10mL distilled water and measured by

spectrophotometer. The wavelength was 540nm and the OD value was 0.685, which met the test requirements. PEI-Mn0.5Zn0.5Fe2O4 nanoparticles were washed twice with distilled water, dried, and suspended with normal saline to 0.1g/mL; Normal saline was used as negative control and distilled water as positive control. After preheating in 37°C water bath, 0.2ml of diluted rabbit blood was added to each group, and then gently mixed. The supernatant of each tube was taken by centrifugation (2500g/min, 5min) after 37°C water bath for 60min. Spectrophotometer to measure the OD value at 545nm, the absorbance value of positive control group should be 0.8±0.3, the absorbance value of negative control group should not be greater than 0.03, hemolysis rate (%) = (mean OD of the tested sample - mean OD of negative control)/(positive control OD - mean OD of negative control) ×100%, If the hemolysis rate <5%, indicating that the material has no hemolysis effect and meets the requirements of hemolysis test for medical materials. Three parallel samples were set for each group.

The Median Lethal Dose (LD50) Test of Mice

Sixty Kunming mice were randomly divided into 6 groups with 10 mice in each group. After fasting overnight, mice in each group were intraperitoneally injected with PEI-MZF nanoparticles suspension at dosage of 1.568g/kg, 2.8g/kg, 5g/kg, 8.92g/kg and 15.94g/kg, respectively. 0.9%NaCl equivalent to the maximum dose was injected as the negative control group. The animals in each group were recorded for 15 days after injection. At the end of the experiment, Karber method was used to calculate the LD50 of the nanoparticles in mice, LD50 = LG-1 (XK-I(\sum P-0.5)). XK is the logarithm of the maximum dose group, P is the response rate of each group, expressed as a decimal, I is the logarithm of the ratio of the two adjacent dose groups (LG (high/low)), and n is the number of animals in each group.

Mice Bone Marrow Micronucleus Test

Thirty-six Kunming mice were divided into 6 groups. Six in each group. PEI-MZF nanoparticle suspension was intraperitoneously injected at dosage of 5g/kg, 2.5g/kg, 1.25g/kg and 0.625g/kg, respectively. 40mg/kg cyclophosphamide was injected in as the positive control, and the maximum dose of 0.9%NaCL was as the negative control. The experiment method was adopted at 30h, that is, the interval between the two administrations of the test substance was 24 hours. Six hours after the second administration, the animals were sacrificed. The femoral bone marrow was taken, fixed with methanol for 5 minutes, dried and stained with Giemsa dye for 15

minutes, and then observed. One thousand polychromatic erythrocytes (PCE) with MN were counted in each mouse. The results were expressed in ‰. Poisson distribution was used to detect whether there were statistical differences between the groups

Determination of PEI Adsorption on Nanoparticles

According to the literature, ²² the Bradford method was used to detect the PEI-MZF magnetic fluids prepared under different pH conditions. The mixture was centrifuged in a high-speed centrifuge for 1 hour. Ten microliters of the supernatant was aspirated from the mixture, and diluted to 100µL with PBS buffer, then 1000µL of Coomassie Brilliant Blue solution was added in. Two minutes later, UV/vis spectrometer (U-2001, HITACHI Japan) was used to measure the optical density of the new mixture at 595nm wavelength. Comparing with the standard curve (drawn under the same experimental conditions), the amount of free PEI in the supernatant was obtained. According to the formula [N] = (C total- C free)/[NP], the amount of PEI adsorbed on nanoparticles (mg/g) under different modification conditions was calculated. Where [N] represents the amount of PEI adsorbed on the nanoparticles, C total and C free respectively represent the concentration of added and remaining PEI (mg/mL), and [NP] represents the concentration of nanoparticles (g/mL).

Heating Characteristics of PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ Nanoparticles in High Frequency Electromagnetic Field

Different doses of PEI-Mn $_{0.5}$ Zn $_{0.5}$ Fe $_2$ O $_4$ nanoparticles were dispersed in 5 mL 0.9% NaCl, to the concentrations of 8, 10, 12.5, 15g/L, respectively. Then, the nanoparticle fluid was placed in a flat-bottomed cuvette, and in turn heated under the high frequency electromagnetic field (SP-04C, Shenzhen, China) machine. The output frequency was 230 KHz and the output current was 30 ampere. Then, the temperature was measured every 5 minutes in 1 hour.

DNA Binding Assay of PEI-MZF-NPs

pEGFP-C1 DNA solution (0.01 mg/mL) was mixed with various amounts of PEI-MZF-NPs (10 mg/mL), to a final volume of 1 mL. The nanoparticle–DNA complexes were prepared at ratios of 0:1, 5:1, 10:1, 20:1, 30:1 (weight/weight), respectively. The complex formation study was performed at room temperature and allowed to stand for 30 minutes to attain

complexes. Then, the PEI-MZF-DNA complexes were electrophoresed by 1% agarose gel electrophoresis at 100~V for 45~minutes.

Construction and Identification of the Eukaryotic Expression Plasmids: EgrI-Hsp70- HSVTK

Construction of Egr I - HSVTK

The primer sequences of HSVTK were designed according to pD3SX-HSP70-HSVTK template (Stressgen), and amplified by PCR. In order to facilitate subsequent experiment, several restriction sites were added on the two sides of HSVTK. Then, the HSVTK segment was subcloned into pCDNA3.1-Egr1-EGFP vector (Constructed and kept in our lab [11]) on EcoRI and XhoI sites. Synthetic products were identified by PCR amplification and gene sequence analysis (sequence primer: Egr1-f, BGH-r).

HSVTK-f:5'-G GAATTC GCCACC ATGGCC TCG TAC CCGGCCAT-3' (EcoRI)

HSVTK-r:5'-CCG CTCGAG TCAGTTAGCCTCCCC CATCT-3' (XhoI)

Product: Length=1131+13+9=1153bp, GC%=65.4, Ta=59.7

PCR Amplification of Hsp70 Promoter Segments

The primer sequences of Hsp70 promoter was designed according to pD3SX template, and amplified by PCR. In order to facilitate subsequent experiment, several restriction sites were also added on the two sides of Hsp70. Synthetic products were identified by enzyme cleavage and gene sequence analysis (gene name: Hsp70; gene length: 434 bp; carrier name: pUC57; cloning site: EcoRI, HindIII).

Hsp70-f:5'-CCC AAGCTT CTCGAGGCGCG TCCT CAGA-3' (HindIII).

Hsp70-r:5'-G GAATTC GGTCGACTAGAGAGCT TCTT-3' (EcoRI).

Product:Length=418+9+7=434bp, GC%=70.6, Ta=58.

Hsp70p Was Sub-Cloned into Egr1-EGFP to Get Hsp70-Egr1-EGFP

The amplified Hsp70p fragments were collected and subcloned into Egr1-EGFP on HindIII and EcoRI sites. After transfection, the plasmids were purified and identified by PCR.

Hsp70-f:5'-CCC AAGCTT TCGAGGCGCGTCCTCA GA-3' (HindIII).

Hsp70-r:5'-G GAATTC GGTCGACTAGAGAGCTT CTT-3' (EcoRI).

Product:Length=418+9+7=434bp, GC%=70.6, Ta=58. According to the PCR electrophoresis results, the correct Hsp70-Egr1-EGFP sequences were selected to be sequenced, EGFP-N-3 as the sequence primer.

Construction of Hsp70-Egr1p-HSVTK

Hsp70- and Egr1p-HSVTK were connected after Egr1-Hsp70-EGFP digested by HindIII and EcoRI. Then, the connected products were transfected into DH5α. At last, the plasmids were isolated and purified after enlargement culturing and tested by HindIII &EcoRI enzyme cleavage. For further, the right clone was verified by MluI enzyme cleavage. (The flow chart of pHsp70-Egr1-HSVTK construction see Figure 1.)

In vitro Optimization of Thermo and Radiation Driven Expression

Cell Transfection and Optimization of Expression Efficiency Assay Using PEI-Mn $_{0.5}$ Zn $_{0.5}$ Fe $_2$ O $_4$

HEK293 cells were transfected with Egr1-Hsp70-HSVTK (or Egr1-HSVTK or Hsp70-HSVTK) using PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ (self-made) at a ratio of 1:20 (DNA: PEI-MZF, w/w), and placed on a Magneto FACTOR plate (magnet-transfection) for 30 minutes. Then, the medium was replaced with fresh medium containing serum,

and continued to incubate for 24 hours. As a control, blank plasmid (pCDNA3.1) were transfected into the cells. To investigate the effect of the radiation and heat on the induction of transgene expression, the transfected cells were exposed to varying conditions.

- Twenty-four hours later, cells were exposed to 2Gy (6 Me V) X-ray radiation, using Siemens medical linear accelerator and harvested at time point of 1h, 6h, 12h, 24h, 48h, respectively. Then, the HSVTK expression level in the cells were tested by real-time PCR analysis.
- The transfected cells were exposed to varying dose of X-ray (1, 2, 4, 8, 16Gy), following the ways as described above. Six hours after treatment, HSVTK expression of the cells were analyzed by real-time PCR.
- 3. The cells transfected with Hsp70-Egr1-HSVTK or Egr1-HSVTK or Hsp70-HSVTK were treated by 2Gy X-ray, thermotherapy mediated by MZF, both of radiation and thermotherapy, and none of the treatment, respectively. Six hours later, the expression of HSVTK in the cells was also tested via realtime PCR.

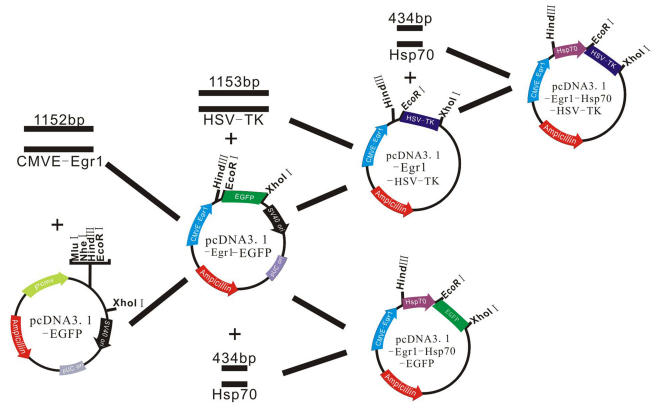


Figure I The flow chart of pHsp70-EgrI-HSVTK construction.

HSVTK Expression Tested by Real-Time PCR

- Total RNA from each cell population was extracted using a commercial kit according to the instructions provided by the supplier (Takara). Total RNA was diluted to 5 ng/mL in water immediately to perform RT-PCRs. GAPDH was used as internal reference.
- 2. The real-time PCRs required the following primers:

HSVTK-f: CCC ACG CTA CTG CGG GTT TAT (153–174).

HSVTK-r: TGT TGG TGC CGG GCA AGG TC (621–602).

Product Length: 469bp Ta=51°C.

GAPDH-f: 5'-GCCACATCGCTCAGACAC- 3'.
GAPDH-r: 5'-CATCACGCCACAGTTTCC- 3'.

Product Length: 614bp Ta=53°C.

1. Reaction and Analysis

Regents:

MgCl ₂	ΙμL
5×PCR Buffer	5 μL
dNTPs	0.25 µL
Primers	f/r 0.25∼I µL
cDNA	2 μL
ddH2O	14.5~16µL
Taq Enzyme	0.25 μL

Mix them on the ice, then centrifuge (4000 rpm, 10s), PCR

1. Data analysis of QPCR

94°C	2min	
94°C	30sec	
51°C	30 sec	35cycles
72°C	90 sec	
72°C	I 0min	
End		
End		

The expression of gene F value= $2^{-\Delta\Delta ct}$

 $\Delta\Delta$ ct= (Average ct value of target gene in experiment group – Average ct value of housekeeping gene in experiment group) – (Average ct value of target gene in control group – Average ct value of housekeeping gene in control group).

The higher F value means the higher gene expression level.

Verification of Transfection and Expression of Egr I/ Hsp70-HSVTK in SMC7721 Cells

To investigate the transfection ability of PEI-MZF NPs in vitro, the SMC7721 cells (a kind of HCC cells, purchased from Shanghai Institute of Biochemistry and Cell Biology) were transfected with Egr1-Hsp70-HSVTK (or Egr1-HSVTK or Hsp70-HSVTK) using PEI-Mn_{0.5}Zn_{0.5}-Fe₂O₄ (self-made) at a ratio of 1:20 (DNA: PEI-MZF), and placed on a Magneto FACTOR plate (magnetofection) for 30 minutes. Then, the medium was replaced with fresh medium containing serum, and continued to incubate for 24 hours. As a control, blank plasmid (pCDNA3.1) was transfected into the cells. To investigate the effect of the radiation and heat on the induction of transgene expression, the transfected cells were exposed to 2Gy (6 Me V) X ray radiation, and harvested at time point of 12h. And HSVTK expression of the transfected cells were analyzed by RT-PCR.

The in vitro Anti-Hepatoma Therapy of Hsp70-Egr1-HSVTK Using PEI-MZF Nanoparticles and Radiation

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MTT Assay for Cell Proliferation

The cytotoxicity of the combination therapy on SMC7721 cells was estimated by MTT assay. Twelve hours before transfection, cells were plated in a 96-well microtiter plate at a density of 4×10^3 cells per well (day 0) and 6 holes each group. The Hsp70-HSVTK or Egr1-HSVTK or Egr1-Hsp70-HSVTK plasmid DNAs were introduced to cells using the PEI-MZF nano-particles and Magneto FACTOR plate (day 1). They were treated with different ways: (1) negative control group (RPMI1640 culture medium), (2) PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ hyperthermia group, (3) 2Gy radiation therapy group, (4) Egr1-HSVTK combined with 2Gy radiation therapy group, (5) Hsp70-HSVTK+ PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ combined with hyperthermia group (MZF-mediated magnetic hyperthermia), (6) Egr1-Hsp70-HSVTK + PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ combined with hyperthermia and 2Gy radiation therapy group. The concentration of PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ magnetic fluid was maintained at 10 mg/mL in the experimental group and hyperthermia (43°C for 1h) was performed by PEI-Mn_{0.5}Zn_{0.5}-Fe₂O₄ fluid under AMF (200kHz, output current 30A, SPG-06A) after overnight transfection (day 2). The radiation therapy group was given 2Gy X-ray irradiation (linear accelerator) after overnight transfection (day 2). The radiation combined hyperthermia group was given heat treatment immediately after receiving the

irradiation (day 2). Cells in groups 4–6 were also treated with GCV at the concentration of 100mg/mL. The medium in each well was replaced on days 3 and 4. On day 5, 20μ L of MTT were added onto the treated cells, and continue to be incubated for 4 hours. Then, the supernatant was replaced with 150 μ L of DMSO, and shook for 10 minutes, then was placed on the microplate reader to read the absorbance (A) value at 492nm. The cell proliferation rate (%) = OD value (experimental group)/OD value (negative group) × 100%.

Flow Cytometry for Detection Cell Cycles

Twenty-four hours before transfection, the cells were plated in a 50mL culture flask at a density of 6×10^5 cells per flask (day 0) and 3 flasks each group. Then, the cells were divided into 6 groups and treated with the ways as described above. After 48 hours, they were harvested to prepare a single cell suspension, washed with PBS, then fixed with 70% cold ethanol for 24 hours. Before detection, the cells were washed twice with PBS, incubated with 50mg/L RNase at 37°C for 30 minutes, stained with 0.5mL PI solution (20mg/mL, 0.25mg/mL RNase A) at room temperature (avoid light) for 30 minutes, and immediately filtered with 300 mesh screen for machine analysis (FACS Vantage SE).

In vivo Anti-Hepatoma Therapy

The 5-week-old male and female BALB/c nu/nu mice were injected subcutaneously in the right forelimb with 2.5×10^6 SMC7721 cells (day 0). When the xenografts tumor reached 80-120mm³, the Hsp70-HSVTK or Egr1-HSVTK or Egr1-Hsp70-HSVTK plasmid DNAs were subcutaneously introduced in using the PEI-Mn_{0.5}Zn_{0.5}-Fe₂O₄ method on days 21, 23 and 25. They were divided into 6 groups: (1) negative control group (0.9% saline), (2) 2Gy radiation therapy group, (3) hyperthermia group (4mg PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ fluid), (4) Egr1-HSVTK combined with 2Gy radiation therapy group, (5) Hsp70-HSVTK+ Mn_{0.5}Zn_{0.5}Fe₂O₄ combined with hyperthermia group, (6) Egr1-Hsp70-HSVTK + PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ combined with hyperthermia and 2Gy radiation therapy group (MZF-mediated magnetic hyperthermia). Negative group was injected in 0.2mL saline on days 22, 24 and 26. The radiation therapy group was given 2Gy/min X-ray irradiation (linear accelerator) on days 22, 24 and 26 and the distance from the skin was 1.0m. The tumor was exposed to the radioactive source, and the rest of the body was protected with a lead block in a suitable plastic mold. Hyperthermia (43°C for 1h) was performed by PEI-Mn_{0.5}Zn_{0.5}-Fe₂O₄ under AMF on days 22, 24 and 26. The radiation combined

hyperthermia group was given heat treatment immediately after receiving the irradiation on days 22, 24 and 26. GCV (30 mg/kg per day) was injected intraperitoneally until day 14 in groups 4-6. During the treatment, the animal's behavior change, weight, daily food intake, daily water intake, tumor surface change were observed. The tumor diameter was measured with calipers, and tumor volume (V) was calculated using the formula: V = abc/2 (where a is the longer diameter, b is the shorter one and c is the height). The results were expressed as percentages relative to tumor size of the negative group. Tumor inhibition rate=(Vave of negative group - Vave of treatment group)/ Vave of negative group×100% (Vave is the average volume of the tumor in the group). After the treatment, the tumor tissue was stripped, weighed, fixed with 4% formaldehyde, sectioned, and HE stained to observe the changes in cell structure. Also, the other tissues were stripped and HE stained to observe the changes in the treatment group. Care, treatment and euthanasia of the animals were performed in accordance with the Institutional Animal Care and Use Committee (IACUC) guidelines and approved by the IACUC (approval number:1,802,135, Southeast University, Nanjing, China). No mice showed signs of toxicity in response to the therapeutic approaches.

Statistical Analysis

Values were expressed as mean \pm SD. The data were analyzed with the SPSS 20.0 program, with P < 0.05 taken as the criterion for statistical significance.

Results and Discussion

Characteristics of MZF-NPs

The self-prepared Mn_{0.5}Zn_{0.5}Fe₂O₄ nanoparticles are high electron-dense and uniform in size with average diameter 30-40 nm observed by HREM (Figure 2A). EDS confirms that the prepared nanoparticles only contain four elements of manganese, zinc, iron and oxygen, and PDF (Powder diffraction File) cards confirm their XRD patterns (Figure 2B and C). Before the modification, the nanoparticles are aggregated, due to the strong interactions between the particles. This aggregation effect is unfavorable for its application. After modification, PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ nanoparticles show little change in appearance and size. They are approximated spherical shape, improved dispersibility, and less agglomeration, indicating that the interaction between magnetic particles is smaller due to the existence of the PEI shell (Figure 2D). Nanomaterials of this size could avoid

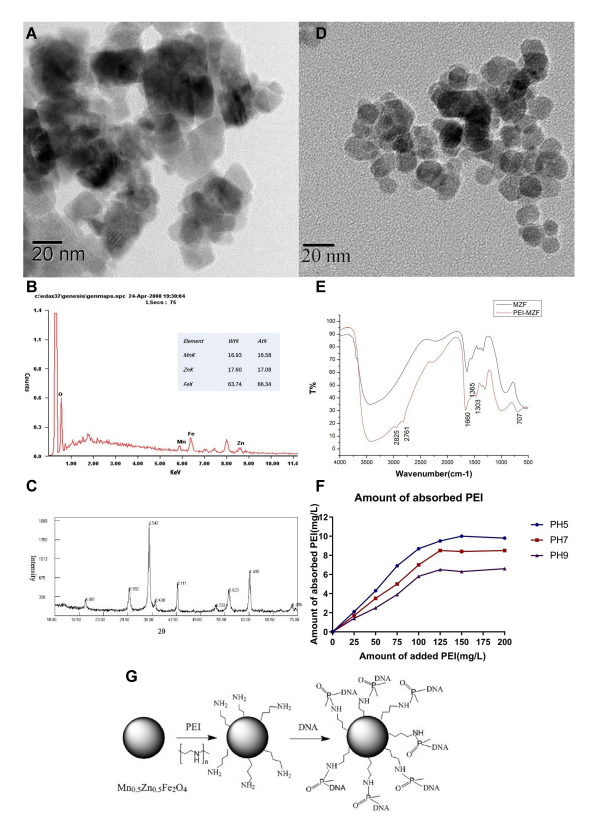


Figure 2 (**A**) HREM image of Mn_{0.5} Zn_{0.5}Fe₂O₄. (**B**) SEM-EDS analysis of Mn_{0.5} Zn_{0.5}Fe₂O₄. (**C**) XRD analysis of Mn_{0.5} Zn_{0.5}Fe₂O₄. (**D**) TEM image of the PEI-Mn_{0.5} Zn_{0.5}Fe₂O₄. (**E**) FTIR of PEI- Mn_{0.5} Zn_{0.5}Fe₂O₄ and Mn_{0.5} Zn_{0.5}Fe₂O₄. (**F**) Adsorption of PEI on the surface of nanoparticles under different pH. (**G**) Scheme of PEI/MZF interaction with DNA.

excretion by the kidneys and phagocytosis by phagocytic cells of the spleen, increasing circulation time and allowing nanoparticles to accumulate in the tumor.²³

Not only the size but also surface modifications, shape, charge, and flexibility can affect the treatment results.²⁴ The FTIR spectra of MZF particles with and without PEI is presented in Figure 2E. Compared the two curves, the dispersant-treated powder presents specific peaks in 713.5 cm-1,1486.8 cm-1, 1577.5 cm-1, 2844 cm-1 and 3013 cm-1 dot, which corresponding to the peaks of -NH2- and -CH2-. -NH2- and -CH2peaks exist in PEI but not in MZF, which proves that PEI is successfully adsorbed on the surface of MZF particles. Figure 2F shows the PEI adsorption amount on the surface of the nanoparticles, under different PH. When PEI is less than 20% (% of the mass), as more PEI adding, adsorption quantity of PEI also gradually increase. The adsorption capacity reached the highest value of 9.81mg/g when PEI was added 15% at pH 5.0. When pH was 7.0 and 9.0, PEI added 12.5%, the adsorption capacity reached the highest values of 8.53mg/g and 6.49mg/g, respectively. Thus, we chose 20% PEI (w/w) and pH5 as the preparation conditions for the modifications.

Why did we chose PEI? PEI was an effective cationic polymer gene carrier, with an active nitrogen atom (-NH2) every two carbon atoms in the molecule. These nitrogen atoms could be protonated to get a positive charge (-NH3+) under physiological conditions, which could help form complexes with negatively charged phosphate groups on DNA by electrostatic adsorption (the scheme see Figure 2G). In addition, the surface charge of PEI-DNA complex was significantly lower than that of PEI, which was conducive to the cell membrane fusion and reduce the toxicity of the vector. The ability of PEI binds to DNA depended on the ratio of the two (N/P) and the particle size of the complex. Nano-sized particles helped to achieve carrier targeting and cell penetration effectiveness.²⁴ The self-prepared PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ nanoparticles were suitable for the carrier with average diameter 30-40 nm.

Toxicity of PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ Nanoparticles

PEI Cytotoxicity Test (MTT)

It is reported that as the N/P ratio increased, so did the toxicity of PEI/DNA complexes to the cells, often due to

the increased adsorption of free cations to the cell membranes on the surface of PEI/DNA complexes. In this study, for the two compounds, PEI-MZF/DNA and PEI/ DNA, as the N/P ratio increases from 1 to 9, the toxicity to the cells increases, as evidenced by a gradual decline in cell viability (Tables 1 and 2). However, for MZF-NPs, although the toxicity to cells increases with the increase of the N/P ratio, the cell survival rate remains above 54.8% in all tested N/P ratios, whereas the toxicity of PEI/DNA to cells increases significantly with the increase of the N/P ratio, and the cell survival rate decreases significantly, to a minimum of only 27%. A comparison of the two groups is shown in Figure 3. In the following study, we used PEI/ MZF at a N/P ratio less than 5. Below this ratio, the cell proliferation rate is >75%, cytotoxicity classification is level 1, and is a qualified biological material.

Hemolysis Test

The absorbance values of negative control group (normal saline), positive control group (distilled water) and experimental group (PEI-MZF normal saline suspension) at 545nm are listed in Table 3, and the hemolysis rate is 0.608%, far less than 5%.

Table I PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ Nanoparticle Toxicity Test MTT $(\overline{X}\pm S, n=8)$

Group	OD Value	Cell Viability %	Cytotoxicity Classification
Negative Group	1.06±0.04	100	0
N/P=I	0.96±0.03	90.8	1
N/P=3	0.87±0.03	85.4	1
N/P=5	0.81±0.02	76.8	Ì
N/P=7	0.71±0.04	66.6	2
N/P=9	0.69±0.01	65.0	2
N/P=II	0.61±0.02	54.8	2
Positive Group	0.16±0.03	15.2	4

Table 2 PEI/DNA Toxicity Test MTT ($\overline{X}\pm S$, n = 8)

Group	OD Value	Cell Viability %	Cytotoxicity Classification
Negative Group	1.32±0.06	100	0
N/P=I	1.15±0.03	87.1	1
N/P=3	1.08±0.02	80.2	1
N/P=5	0.85±0.03	64.4	2
N/P=7	0.62±0.02	46.8	3
N/P=9	0.53±0.01	40.7	3
N/P=II	0.36±0.02	27.1	3
Positive Group	0.18±0.03	13.6	4

Comparison of PEI-MZF and PEI cytotoxicity under different N/P ratio

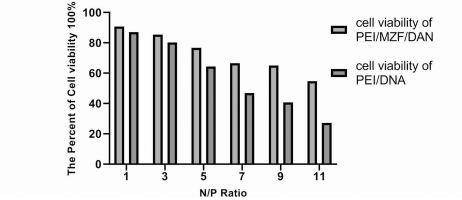


Figure 3 Comparison of PEI-MZF and PEI cytotoxicity under different N/P ratio The N/P ratio of PEI/MZF/DNA and PEI/DNA was 1, 3, 5, 7, 9, 11.

The Median Lethal Dose (LD50) Test of Mice

Mice in experimental groups were injected with different doses of PEI-MZF nanoparticles suspension, and some mice showed symptoms of lethargy, loss of appetite, diarrhea and so on. Some died in the following days. Fifteen days after the administration, the deaths of the animals in each group were recorded, as listed in Table 4. The LD50 of the mice was 3.392g/kg calculated by Karter method, and the 95% confidence interval was 1.245~9.243g/kg, indicating that the nanoparticles have a wide range of safety values.

Mice Bone Marrow Micronucleus Test

Micronucleus (MN) in polychromatic erythrocytes in each group was detected, and no significant difference is found

Table 3 Hemolysis Test Results

Group	Absorbance Value (OD)		Average OD	Hemolysis Rate %	
	I 2 3				
Negative	0.017	0.021	0.025	0.021	0
Experiment	0.024	0.027	0.028	0.026	0.608
Positive	0.834	0.842	0.853	0.843	100

between the experimental group and the negative control group, as shown in Table 5. It can be considered that the material has no teratogenic or mutagenic effect.

The experimental results show that there is no hemolysis effect contacting with blood, and the hemolysis rate is 0.608%, far less than 5%, which meets the requirements of hemolysis test for medical materials. In this experiment, micronucleus formation rate is 2\% when mice were given the highest concentration of 5g/ Kg by micronucleus test. Micronucleus test show that there is no significant difference in the micronucleus formation rate of bone marrow in experimental group compared with that in the negative control group (P > 0.05), but there's significant difference compared with that in the positive control group (P < 0.05), so it could be considered that the material has no teratogenic or mutagenic effect. The LD50 is 3.392g/kg, and the 95% confidence interval is 1.245~9.243g/kg. LD50 in 0.5-5g/ kg body weight is a low toxicity category, but such a large dose will not be used in the actual gene carrier. From the results of biocompatibility of PEI- MZF, PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ is non-toxic in vivo and in vitro, and has good biocompatibility.

Table 4 Results of Acute Toxicity Test of Nanoparticles

Group	Dosage (g/kg)	Dose Logarithm	n	Mortality	P	Q	p×q
I	1.568	3.195346	10	0	0	I	0
2	2.8	0.447158	10	1	0.1	0.9	0.09
3	5	0.69897	10	3	0.3	0.7	0.21
4	8.92	0.950365	10	4	0.4	0.6	0.24
5	15.94	1.202488	10	9	0.9	0.1	0.09
		I = 0.56		$\sum_{p} = 1.7$			

Notes: p is death rate, q is survival rate, n is the number of mice.

Table 5 Micronucleus Test Results

Group	n	Number of PEC	Number of PEC with MN	Micronucleus Formation Rate (‰)
Negative	6	6000	12	2
5.00g/kg	6	6000	П	1.8
2.5g/kg	6	6000	10	1.67
1.25g/kg	6	6000	12	2
0.625g/kg	6	6000	П	1.8
Positive	6	6000	143	23.8

Abbreviation: n, number of mice in each group.

Heating Test of PEI- Mn_{0.5}Zn_{0.5}Fe₂O₄ in vitro

Under the high frequency alternating magnetic field, the heating capacity of the magnetic fluid is positively related to the concentration. As the MZF concentration increases, the maximum temperature rises, but they have common rules: The temperature rises rapidly within 30 minutes, and rises slowly from 30 to 40 minutes. After 40 minutes, the temperature hardly rises and remains constant at a certain level (Figure 4). With different concentrations of magnetic fluid, the maximum temperature can rise to 39°C, 43.5°C, 47°C and 50°C respectively, and it can be kept for 20 minutes in a magnetic field. The temperature of 10g/L MZF solution can rise to 43.5°C, which is the ideal temperature for the treatment of tumors while not harming normal tissues. Later on, 10 g /L was selected for magnet hyperthermia.

DNA Binding Assay of PEI-MZF-NPs

The binding capacity was evaluated based on the relative intensity of the free DNA band in each well with respect to the well with naked DNA. Figure 5 shows the retarded DNA due to nanoparticles binding at varying w/w ratios. As a result, the synthesized PEI-MZF-NPs could bind

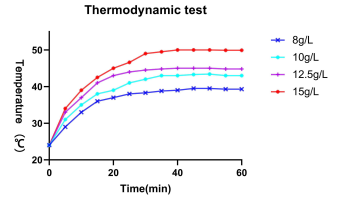


Figure 4 Thermodynamic test of different concentration of PEI- $Mn_{0.5}$ $Zn_{0.5}Fe_2O_4$ in vitro.

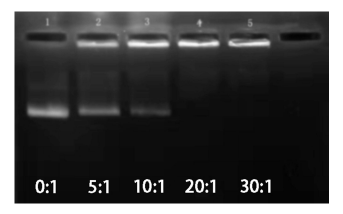


Figure 5 Image of agarose gel electro-phoresis of plasmid DNA and complexed with PEI- $Mn_{0.5}$ $Zn_{0.5}$ Fe₂O₄ (Lane I - 5(PEI-MZF/DNA w/w): 0:1; 5:1; 10:1; 20:1; 30:1).

DNA effectively, resulting in complete retention of the DNA by the agarose gel. When the ratio of 20, DNA migration is completely retarded by PEI-MZF-NPs. Later on, the ratio of 20:1 (nanoparticle/DNA) was selected for gene transfection.

Because of its high stability, chemical simplicity, and high transfection activity, PEI has been widely used to function as gene delivery carrier, either in branched (b-PEI) or linear (l-PEI) form.²⁵ Several studies have attributed the transfection efficiency of PEI to the so-called proton sponge effect.^{26,27}

Construction and Identification of Egrl-HSVTK

Recombinant plasmid of Egr1-HSVTK was confirmed by PCR amplification and gene sequence analysis, which can be seen from Figure 6A with 1153bp HSVTK specific band in lane 1 and Figure 6B with the same sequence as that of HSVTK template.

Construction and Identification of Hsp70-Egr1- EGFG and Hsp70- Egr1- HSVTK

Recombinant plasmid of Hsp70-Egr1-EGFP was also confirmed by PCR amplification and gene sequence analysis, which can be seen from Figure 7A with 434 bp Hsp70 band in lane 1–4 and Figure 7B with the same sequence as that of Hsp70 template.

Recombinant plasmid of Hsp70-Egr1- HSVTK was confirmed by restriction enzyme digestion analysis, which can be seen from Figure 8A (dual enzyme digestion) and Figure 8B (single enzyme digestion). If Hsp70 fragment was inserted correctly into Hsp70-Egr1 -HSVTK, the fragment length between EcoRI

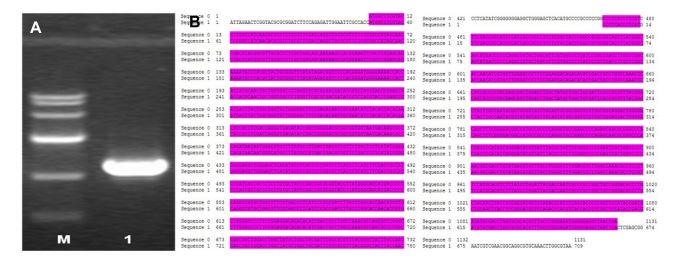


Figure 6 (A) PCR gel electrophoresis of EgrI-HSVTK transfected (lane I: Marker IV (each band in turn from top to bottom is 7 K, 5.5 K, 3.5 K, 2 K, I K, 500 bp); lane I: EgrI-HSVTK). (B) Sequencing and alignment of EgrI-HSVTK (sequence 0: HSVTK; sequence 1:EgrI-HSVTK-EgrI-f.abI; software: Dnassit 2.0).

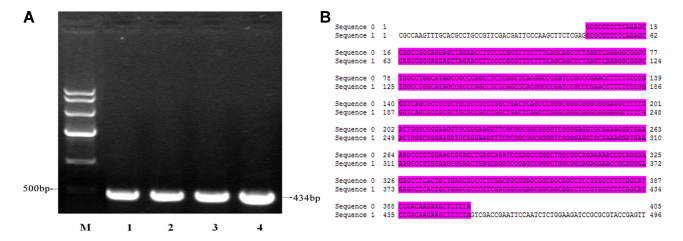


Figure 7 (A) PCR gel electrophoresis of Hsp70-Egr1-EGFP transfected (lane 1: Marker IV each band in turn from top to bottom is 7 K, 5.5 K, 3.5 K, 2 K, 1 K, 500 bp); lane I-4: Hsp70-Egr1-EGFP()-(4). (B) Sequencing and alignment of Hsp70-Egr1-EGFP (sequence 0: HSP70; sequence 1: Hsp70-Egr1-EGFP-f.ab1; software: Dnassit 2.0).

and Hind III should be about 434 bp, because the length of Hsp70 is about 434 bp. As shown in Figure 8A of restriction enzyme digestion of Hsp70-Egr1-HSVTK, Hsp70 promoter was cloned correctly.

Theoretically, there was only one MluI restriction site in the original pCDNA3.1-EGFP, if Egr1 was successfully inserted into the plasmid, another MluI restriction site would be brought in, and the fragment length between the two sites of MluI should be about 1600 bp. Seen from Figure 8B, Egr1 has been successfully inserted. So

it can be seen that the eukaryotic recombinant plasmid Hsp70-Egr1-HSVTK was successfully constructed.

In vitro Optimization of Thermo and Radiation-Driven Expression

Real time-PCR assay was used to detect TK gene expression in HEK293 cells. Amplification was monitored by the uptake of SYBR Green with automated estimation of Ct values throughout each reaction. The gene expression relative quantity was calculated as described above.

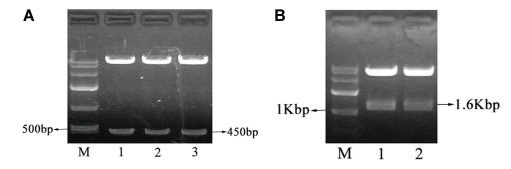


Figure 8 (A) Egr1-Hsp70- HSVTK was identified by EcoRI and HindIII dual restriction enzyme digestion (lane 1: Marker IV each band in turn from top to bottom is 7 K, 5.5 K, 3.5 K, 2 K, 1 K, 500 bp;lane 1–3: pCDNA3.1-Egr1- Hsp70-HSVTK①-③digested by EcoRI and HindIII.). (B) Egr1- Hsp70-HSVTK was identified by Mlul restriction enzyme digestion (lane 1: Marker IV each band in turn from top to bottom is 7 K, 5.5 K, 3.5 K, 2 K, 1 K, 500 bp;lane 1–2: Hsp70-Egr1- HSVTK ①-②digested by Mlul.).

In order to study the radio-sensitivity of the promoter at different time point, TK gene expression in HEK293 cells after 2Gy X ray irradiation at 1 h, 6 h, 12 h, 24 h and 48 h was tested. As Figure 9 shown, the fluorescence intensity began to increase 1 hour later and peak at 6 hour time point (Figure 9). Before 6 hour, Egr1 promoter driven-TK expression increased with the extension of time, then gradually reduced. Compared the highest level with the other one, the expression of TK at 6 hour (4.53±0.29 mean fluorescence intensity) is 5.8 times of that at 48 h (0.33 \pm 0.15), 2.35 times of that at 1 h ((1.86 ± 0.21) , and 1.19 times of that at 12 h (3.72 \pm 0.30). And then the expression level at 48 h reduced to 2/5 of that at 1 h. After induced by radiation, the expression of TK all increased compared to that of none induction. The one-way analysis of variance shows the expression of TK has statistically significant between 6h, 12h groups and 1h, 24h, 48h groups (P<0.05).

In order to study the gene expression induced by different doses of radiation, the cells transfected with Egr1-Hsp70-HSVTK/PEI-MZF complex were given 0 Gy, 1 Gy, 2 Gy, 4 Gy, 8 Gy, and 16 Gy X-ray, respectively. Six hours later, mean fluorescence intensity of TK in cells after radiation is higher than that without radiation, especially the group treated by 2 Gy radiation (Figure 10). As the radiation dose increases, the expression of TK first rises then slightly reduces. Compared the highest level with the other one, the expression level of TK after 2 Gy radiation (2.78 ± 0.12 mean fluorescence intensity) is 4.96 times of that after 1Gy (0.56±0.02). The expression level of TK in cells treated for 8Gy and 16Gy is only 0.7 times of that after 2Gy. The One-way analysis of variance shows the expression of TK has statistically significant between each group (P < 0.05).

TK expression induced by radiation at different timepoint

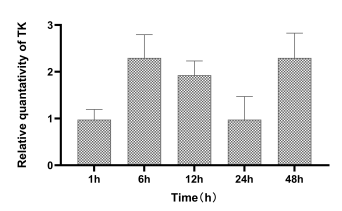


Figure 9 TK expression induced by radiation at different time point. Compared the TK expression at 48h with that in other groups, P < 0.05. Compared the TK expression at 6h with that at other time point, P < 0.05.

TK expression induced by different dose of radiation

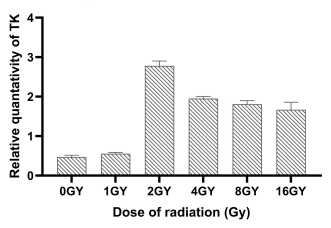


Figure 10 TK expression induced by different dose of radiation. Compared the TK expression with 0Gy radiation with that in other groups P < 0.05. Compared the TK expression after 2Gy radiation with that in other groups P < 0.05.

In order to study the regulation efficiency between the single promoter and double promoter, cells transfected with Egr1-HSVTK, Hsp70-HSVTK and Egr1-Hsp70-HSVTK were treated with different methods: 2 Gy X-ray irradiation (radiation group), magnetic heat treatment under the alternating magnetic field (thermo group) and the 2 Gy radiation combination with thermo (comprehensive group) or without any treatment (control group), and then the TK gene expression was observed. Tested by the real-time PCR, TK gene expression level in the comprehensive group (1.19 ± 0.03) is the highest, which is 2.4 times of that in the control group (0.49±0.03), 1.68 times of that in radiation group (0.71±0.04), and 1.4 times of that in the thermal group (0.84±0.04) (Figure 11). The gene level in thermal group is a little higher than that in radiation group, which is about 1.18 times. The one-way analysis of variance shows the expression level of TK has statistically significant between the experiment group and the control group (P < 0.05).

Verification of Transfection and Expression of PEI-MZF/ Egr1/Hsp70-HSVTK in SMC7721cells

The transfected cells were harvested at time point of 12h. And HSVTK expression of the transfected cells were analyzed by RT-PCR. HSV-TK gene was used as a template to design primers, and the product size was 469bp. Figure 11 shows that TK gene fragments appear in all transfected SMMC7721 cells after RNA extraction and reverse transcription amplification. But the fragments of the TK gene were not found in the un-transfected cells and only GAPDH (the internal control fragment) was found (Figure 12). The

TK expression induced by radiation with different methods

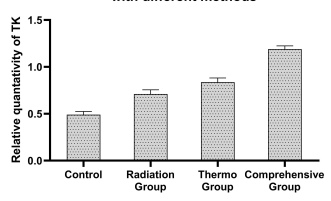


Figure 11 TK expression induced by different methods. Compared the TK expression without treatment, with that in other groups P < 0.05. Compared the TK expression with radiation and thermo, with that in other groups P < 0.05.

results show that pEgr1-Hsp70-HSVTK can be successfully transfected into SMMC-7721 cells (purchased from Shanghai Institute of Biochemistry and Cell Biology, Chinese Academy of Sciences) by PEI-MZF and expressed in them through magnetic heat shock and radiotherapy. GAPDH (614bp) is the internal reference in PCR.

The in vitro Anti-Hepatoma Effect of Hsp70-Egr1-HSVTK Using PEI-MZF Nanoparticles and Radiation MTT Assay for Cell Proliferation

The in vitro antitumor effect of Mn_{0.5}Zn_{0.5}Fe₂O₄ hyperthermia, radiation therapy, Egr1-HSVTK combined with radiation therapy, Hsp70-HSVTK+PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ combined with hyperthermia, and Egr1-Hsp70-HSVTK + Mn_{0.5}Zn_{0.5}Fe₂O₄ combined with hyperthermia and radiation therapy to SMC7721 cells were determined by the MTT assay. The cell viability of each group was calculated according to the measured OD value and the formula, cell viability (%)= Experimental group A/Control group A×100%. The MTT results show that a large number of SMC7721 cells are killed by PEI/MZF/Egr1-Hsp70-HSVTK nano-complexes. As illustrated in Table 6 and Figure 13, low-dose radiation (2Gy) alone could not inhibit tumor growth well, but under the combined effect of heat, radiotherapy and gene therapy, tumor cells show significant growth inhibition. The cell proliferation rate of the triple therapy group is 17.12%, which is much lower than that of radiation alone (86.31%) and hyperthermia alone (58.33%), and also significantly lower than that of radiation combined genome (79.24%) and that of hyperthermia combined genome (44.96%). Compared with

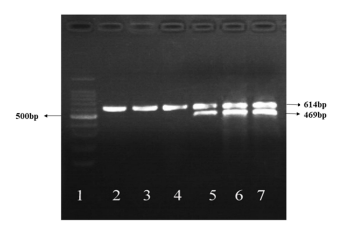


Figure 12 Expression of HSV-TK in SMC7721cells induced by different methods. I 100bp Marker. 2 Negative control (not transfected). 3 Radiation group (not transfected). 4 Hyperthermia group (not transfected). 5 TK transfection by radiation induction group. 6 TK transfection by heat induction group. 7 TK transfection by radiation and heat shock induction group.

Table 6 The Cell Viability of SMMC-7721 cells After Treatment $(\overline{X}\pm S,\ n$ = 6)

Group	A Value	Cell Viability (%)	
Negative group	1.19±0.012	100	
Radiation group	1.027±0.015*	86.31	
Radiation-gene group	0.946±0.015 [△]	79.24	
Heat group	0.696±0.011 [△]	58.33	
Heat-gene group	0.537±0.023 [△]	44.96	
Combined treatment group	0.204±0.029 [△]	17.12	

Notes: *Compared with negative group P > 0.05, $^{\triangle}$ Compared with negative group P < 0.05.

other groups, the difference is statistically significant (P < 0.05). In this study, the radiation and heat-induced dual promoter combined with heat and radiation, which can enhance the therapeutic efficacy, reduce the radiation dose, and reduce systemic side effects.

Flow Cytometry for Detection Cell Cycles

The results of Annexin-PI double staining flow cytometry are listed in Table 7, which is the percentages of apoptotic SMC77212 cells disposed with Egr1-HSVTK, Hsp70-HSVTK and Egr1-Hsp70-HSVTK after 48 hours. It is indicated that the ratio of apoptosis cells and necrosis cells are very low in the negative control group. With different treatment ways, Annexin v-FITC staining cells increase. As well, the ratio of apoptosis cells increase, which is in a treatment-dependent manner. Annexin v staining cells in Egr1-Hsp70-HSVTK combined with radiation hyperthermia group (34.87%) rise significantly compared with hyperthermia alone group (20.22%), radiation genome group (15.16%) and hyperthermia genome group (27.94%), which is consistent with the cell inhibition rate of MTT assay. Moreover, the cell cycle of

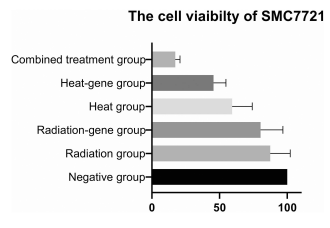


Figure 13 The cell viability of SMC7721 cells after treatment.

each treatment group is blocked in S or G2/M phase to varying degrees (Figure 14A and B).

The HSV-TK/GCV suicide gene system could transform GCV into toxic substances by gene expression products, and plays targeted killing effect on tumor cells. The low-dose radiation and hyperthermia mainly play a role by directly killing tumor cells and inducing tumor cell apoptosis. Therefore, in the mechanism research, the detection and analysis of cell cycle and apoptosis are essential. In this study, the cell cycle analysis of SMMC-7721 dealt with different treatments show that part of cells in the radiation (G2 phase 45.77% and S phase 31.34%), heat (G2 41.16% and S 37.72%), and/or combined gene therapy group (G2 15.91% and S 45.37%) are blocked in the G2 phase and the S phase, especially the S phase increased significantly, indicating that most of cells are blocked in S phase after treatment.

In vivo Anti-Tumor Therapy

Results of Mass and Volume Inhibition Rate Treated with Different Way

Since we previously demonstrated that PEI/MZF-Egr1-Hsp70-HSVTK complex suppressed the growth of HCC cells in vitro, the anti-tumor activity of the nanoparticles containing Egr1-Hsp70-HSVTK was evaluated in SMC7721-bearing mice. As seen in Figure 15, PEI/MZF-Egr1-Hsp70-HSVTK nanoparticles combined with heat and radiation (mass inhibition rate (Im) = 80.10% and volume inhibition rate (Iv) = 83.77%) significantly suppress tumor growth compared to saline and heat (Im = 52.96% and Iv = 59.40%), radiation (Im = 23% and Iv = 22.42%) group at 21st day after first treatment (Figure 15A). It is obvious that the tumor growth of the radiation group is not stunted, but the results need to be compared to the dose of clinical use, which seems to have an effect on tumor growth at much higher dose.

Morphological Changes

During the treatment, the nude mice tumor in groups 3, 5, and 6 involved in hyperthermia had bleeding, necrosis, and eschar formation on the surface. Some eschars could fall off, while such changes are rare in control group (Group 1) and the radiation group (Group 2). After treatment, the tumor tissues of each group were stripped and found that the tumors in the hyperthermia group were dark brown and gray-white in appearance (Figures 15B and C). The pathological sections confirmed that PEI-MZF magnetic nanoparticles were injected into the tissues, while the tumors in the control group were gray-white and more uniform in color. The cells in the control group are tightly arranged, the cytoplasm is rich, the nucleus is

Table 7 Flow Cytometry of Cell Cycles and Apoptosis

Group	Apoptosis (%)	GI Phase	G2 Phase	S Phase
Negative group	0.12%	54.22%	33.78%	11.99%
Radiation group	8.68%	22.89%	45.77%	31.34%
Radiation-gene group	15.16%	30.89%	44.79%	24.31%
Heat group	20.22%	41.16%	37.72%	21.13%
Heat-gene group	27.94%	45.19%	33.66%	21.14%
Combined treatment group	34.87%	38.72%	15.91%	45.37%

large and deeply stained, and the proliferation is active. While the cells in the combined therapy group are significant necrosis, and magnetic material deposition can be seen between the tissues (black). During the treatment time, the tumor in the control group gradually increased over time, the surface was smooth, and no hemorrhagic necrosis was seen. But the tumor in the hyperthermia group gradually decreased during the treatment, part of the tumor surface was uneven, sometimes hemorrhage and necrosis, and the changes are more obvious in Egr1-Hsp70-TK radiation and heating groups. Compared with the control group, the tumor volume and mass of each hyperthermia group are significantly different (p < 0.05), while the tumor volume and mass of the hyperthermia genome (group 5) and radiation hyperthermia genome (group 6) are significantly different (P < 0.05) (Figure 15C) compared with other treatment groups.

Histo-Pathological Changes

Pathological examination reveals that the tumor tissue is poor differentiated cancer; the cells of the negative control group have tightly arranged tumor cells, rich cytoplasm, large and deep stained nuclei, active cell proliferation, and no obvious necrosis. In the Egr1-Hsp70-TK radiation and

PEI-MZF mediated hyperthermia group, the tumor tissues have obvious hemorrhage and necrosis, and the deposition of brown nano-magnetic material is seen (Figure 15C) between the cells. The tumor cells in group 6 are irregular in shape and loosely arranged, and a large number of necrotic areas can be seen. Similar findings are found in groups 3, 4, and 5. The main organs in group 6 were also taken for pathological section observation. There are no obvious pathological changes are found in the heart, liver, lung, spleen and kidney (see Figure 15D). The cells are arranged in an orderly manner, with abundant cytoplasm, proper nucleoplasm ratio, and no degeneration and necrosis. This indicated that the composite material has no significant effect on the tissues.

In conclusion, the tumor transplantation experiment also confirm that the tumor inhibition rate (I_m = 80.10% and 83.54%) of the triple therapy group is much higher than that of the other treatment groups, and the tumor necrosis of the triple therapy group is more obvious than that of the other groups. The combined treatment has achieved a relatively ideal therapeutic effect and almost no effect on normal organs.

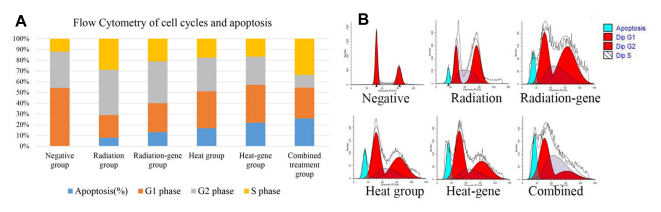


Figure 14 (A) Cell cycle results of SMMC7721 cells after treatment. (B) Flow cytometry results of SMMC7721 cells after treatment.

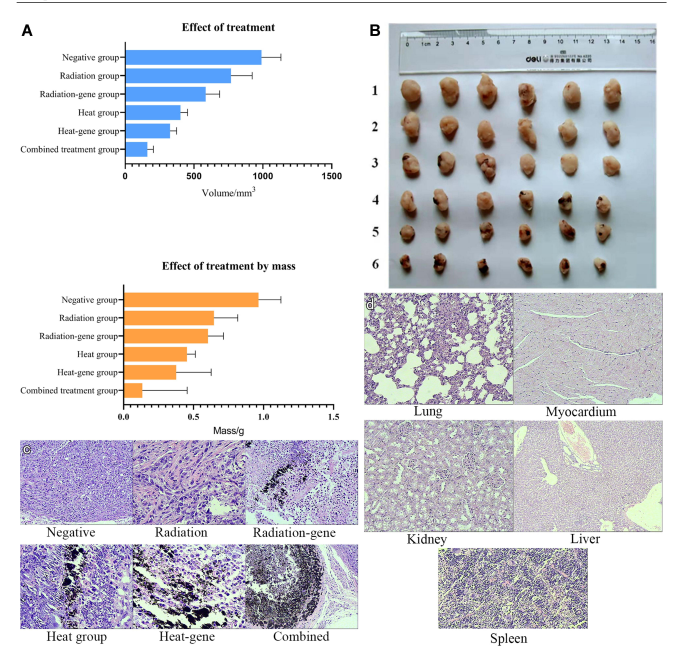


Figure 15 (A) Inhibitory rate of weight and mass in xenograft tumor after treatment. (B) Tumor morphological appearance after treatment. I Negative group. 2 Radiation group. 3 Radiation-gene group. 4 $Mn_{0.5}Zn_{0.5}Fe_2O_4$ hyperthermia group. 5 Heat-gene group. 6 The combined group. (C) The histo-morphology of tumors after treatment (100×). Negative group: The cells are tightly arranged, with rich cytoplasm, large and deeply stained nucleus, and active proliferation. Combined group: The cells are obviously necrotic, and magnetic material deposition can be seen between the tissue. (D) The histo-morphology of organs after treatment (100×). A lung B heart C kidney D liver E spleen.

Conclusions

In this study, we developed an efficient, convenient, dual sensitive and low toxicity system of PEI-MZF/pEgr1-Hsp70-HSVTK nano-complex to active and improve HSVTK gene expression efficiency by combining radiation and thermoradiation based on PEI-MZF. The results showed that the

expression level of HSVTK gene induced by thermo and radiation was significantly higher than that of any single induction in cells. And PEI-MZF-NPs can be used as a novel non-viral gene vector and a hyperthermia strategy in the future. The self-made PEI-MZF, PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ were non-toxic in vivo and in vitro, and had good biocompatibility. In vitro

and in vivo experiments, we found radiation alone or hyperthermia mediated by PEI-Mn_{0.5}Zn_{0.5}Fe₂O₄ or gene therapy or combination of hyperthermia, radiotherapy and gene therapy could inhibit the tumor growth and induce cell apoptosis, and cell cycle arrest. But the radiation and heat shock dual-promoter anti-tumor gene complex group had better tumor suppression effect, and its suppression rate was the highest. In summary, a radiation and heat-inducible gene expression system combination of radiation, hyperthermia and gene therapy described in this study provided a promising therapy approach that may have applications in the treatment of cancer by local or interventional way.

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Disclosure

The authors report no conflicts of interest in this work.

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