Human endostatin gene transfer, either naked or with liposome, has the same inhibitory effect on growth of mouse liver tumor cells in vivo

Chun-Hong Ma, Yan Zhang, Xiao-Yan Wang, Li-Fen Gao, Hua Liu, Chun Guo, Su-Xia Liu, Ying-Lin Cao, Li-Ning Zhang, Wen-Sheng Sun

INTRODUCTION
Tumor neovascularization is necessary to the growth and metastasis of tumors[1]. Anti-angiogenesis is an effective method to treat cancer[2,3]. Endostatin, an endogenous angiogenesis inhibitor, was found in 1997[4]. It can specifically suppress endothelial cell proliferation. Animal experiments have demonstrated that endostatin can strikingly inhibit the growth of many kinds of tumor cells without drug resistance and toxicity. Endostatin is a promising agent to treat cancer[5,6]. However, to produce a large quantity of biologically active proteins has been proven difficult, the treatment requires repeated administration and high doses of recombinant protein, which limits endostatin’s clinical application. Gene therapy can overcome the above disadvantages and become a potential method for cancer therapy. In this article, we constructed the recombinant eukaryotic plasmid using pVAX1 which is the only vector authorized by FDA in clinical trial. It can express secretory endostatin in vivo. Efficient gene transfer and expression are the key to gene therapy. In order to explore an efficient and easy gene transfer method, we also compared the inhibitory effect of liposome and naked DNA transfection on mouse hepatoma cells in vivo.

MATERIALS AND METHODS

Plasmids
Eukaryotic expression vector pVAX1 authorized by FDA in clinical trial was purchased from Invitrogen(USA). Cloning vector pGEM-T-EN containing human endostatin genes was constructed and preserved in our laboratory[7].

Cells and cultures
Mouse hepatoma cell line H22 was purchased from Shandong Medical Science Institute and grown in BALB/c abdominal cavity.

Animals
BALB/c female mice, 5-6 wk of age, were provided by Experimental Animal Center of Shandong University. A total of 1×10⁶ H22 cells were transplanted subcutaneously into mice. Treatment began 1 wk later, when tumor volume ranged between 0.4 to 0.5 cm³.

Primer design
According to the sequences in GeneBank, primers were designed to amplify endostatin genes. In order to get a secretary protein, signal sequence of human IgG γ chain was added to the 5’ end of forward primer, and sequences recognized by KpnI and EcoRI were respectively added to the forward primer and reverse primer: forward(P1): 5’ TTAGGTACCATGGAAGCCCCAGCTCAG CTTCTTTCTCTGCTACTCTGGCTCCCAGATACCACC GAA 3’; reverse(P2): 5’ CGGCAGATTCC TTGGGAGG CAGTC 3’. All of these oligonucleotide primers were synthesized by Shanghai BioAsia Bio-technology Co.,Ltd.

PCR
The endostatin gene containing signal sequence of human

RESULTS: DNA sequencing showed the endostatin gene with the signal peptide was correctly cloned. In situ gene expression assay indicated that both the ratio of DNA and liposome and the dose of DNA could affect the gene transfection efficiency. Interestingly, naked pVAX-sEN had a similar in situ endostatin expression to pVAX-sEN with liposome. Animal experiments showed that pVAX-sEN together with pVAX-sEN-liposome complex could efficiently suppress the growth of mouse hepatoma cells.

CONCLUSION: Naked endostatin plasmid intratumoral injection can get a similar gene transfection efficiency to liposome-DNA complex when used in situ.

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Abstract
AIM: To explore a safe and efficient strategy of tumor therapy using anti-angiogenetic agents.

METHODS: Endostatin gene with a signal sequence of human IgG γ chain was amplified by PCR and cloned into pVAX1 plasmid which was the only vector authorized by FDA in clinical trial to construct a recombinant plasmid named as pVAX-sEN. The recombinant plasmid was detected with EcoRI/KpnI and DNA sequencing. BALB/c mice bearing hepatocarcinoma cell line H22 were treated with naked pVAX-sEN or liposome-DNA complex in which the dose of DNA and the ratio of DNA and liposome were different from each other. To compare the efficiency of gene transfection, expression of endostatin at the treated tumor site was assayed with ELISA. To investigate the effect of pVAX1-sEN on hepatocellular carcinoma, pVAX-sEN either naked or in liposome-DNA complex was injected into BALB/c mice bearing H22, then the diameter of tumors was measured, microvesSEL density was detected by immunohistochemistry, endostatin expression in vivo was assayed at different time points.

RESULTS: DNA sequencing showed the endostatin gene with the signal peptide was correctly cloned. In situ gene expression assay indicated that both the ratio of DNA and liposome and the dose of DNA could affect the gene transfection efficiency. Interestingly, naked pVAX-sEN had a similar in situ endostatin expression to pVAX-sEN with liposome. Animal experiments showed that pVAX-sEN together with pVAX-sEN-liposome complex could efficiently suppress the growth of mouse hepatoma cells.

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PCR
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IgG γ chain was amplified by PCR taking pGEM-T-EN as template and P1 and P2 as primers. PCR reaction was performed in 25 μL volume containing: 1 μL of plasmid DNA, 2.5 μL of 10xbuffer, 0.5 mmol/L Mg++, 5 mmol/L dNTP, 3 units of Taq polymerase, 10 mmol/L primers (forward and reverse). Reaction conditions were at 95 °C for 5 min, then 30 cycles each at 95 °C for 1 min, at 62 °C for 1 min, at 72 °C for 3 min, followed by a final extension at 72 °C for 5 min. PCR products were purified by UNIQUE-10 Kit (Shanghai Shenggong Biological Co.Ltd) after 20 g/L agarose gel electrophoresis.

**Construction of recombinant eukaryotic plasmid expressing secretive endostatin**

PCR products and pVAX1 were digested by EcoRI and KpnI (TaKaRa Biotechnology, Dalian) respectively, then purified and ligased with T4DNA ligase (TaKaRa Biotechnology, Dalian) at room temperature overnight. E.coli DH5α, competent cells were transfected using CaCl2. Recombinant clones were detected by EcoRI and KpnI and then sequenced by Shanghai BioAsia Bio technology Co.Ltd.

**In situ expression of secretory endostatin**

Five to six week old female BALB/c mice were injected subcutaneously with 1×10⁶ logarithmically growing H22 cells to axillary. One week later, when tumor volume was about 0.4 to 0.5 cm³, naked plasmid or liposome-DNA complex (LipofectamineTM 2000 from Invitrogen) was administered intratumorly. The ratio of liposome and recombinant plasmid in liposome-DNA complex was 1:1, 1:3, 1:8 respectively. Animals were sacrificed 3 d after injection and about 0.5 cm³ of tumor was separated and cut into pieces, then cultured in 0.5 mL MEM medium with 100 mL/L of FCS at 37 °C, 50 mL/L CO₂ atmosphere for 24 h. Supernatants were collected and centrifuged at 1 000 r/min for 5 min, then stored at -20 °C for gene expression assay.

**In vivo inhibition assay**

One week after BALB/c mice were injected subcutaneously with 1×10⁶ logarithmically growing H22 cells to axillary, naked recombinant plasmid (20 μg) or the liposome-DNA complex containing 20 μg recombinant plasmid was administered intratumorly once a week for two weeks. The ratio of liposome and recombinant plasmid in liposome-DNA complex was 1:1, pVAX1 and NS were respectively injected into animals as control. The length (a) and width (b) of tumor were measured twice a week. The tumor volumes were then calculated using the formula: Volume=a×b²×0.5².

The experiment was done in triplicate with 5 animals in each group.

**In vivo expression assay of endostatin**

In order to determine the association of anti-tumor effects and the expression level of endostatin, peripheral blood was prepared from tail at different time point and serum was used to detect the expression of endostatin with ELISA.

**ELISA**

Expression of endostatin was detected with human endostatin™ protein accucyte™ ELIA kit (Oncogene) following the manufacturer’s instructions.

**Immunohistochemistry**

Mice were killed 3 d after the fourth administration. Tumors were excised, fixed in formalin and prepared for paraffin-embedded sections. Factor VIII-related antigen was detected with S-P immunohistochemistry kit (Beijing Zhongshan Biocompany) according to the manufacturer’s instructions.
complex was prepared with the ratio of liposome and DNA 1:1 which contained different doses of DNA. Twenty-four hours after transfection endostatins were detected by ELISA (Figure 3). The result indicated that when DNA was less than 20 µg, with the elevation of DNA doses, the expression of endostatin increased. When the DNA dose reached 20 µg, endostatin reached 152 ng/mL, the peak expression. On the contrary, when the DNA reached 50 µg, the expression of endostatin dropped to 80 ng/mL.

**In vivo inhibitory effect of pVAX-sEN on mouse hepatoma cells**

Changes of tumor volumes indicated pVAX-sEN-liposome complex could obviously inhibit the growth of mouse hepatoma cells when administered intratumorly (Figure 4). The tumor volume was only 0.45±0.26 cm³, obviously less than that in NS treated group (1.86±0.62 cm³) (P<0.05). The inhibition rate was about 73%. Interestingly, pVAX-sEN alone could produce an inhibitory effect. The tumor size in pVAX-sEN treated group was 0.4±0.25 cm³, and the inhibitory rate was about 80%, which was not significant from pVAX-sEN-liposome complex treated group (P>0.05).

**Discussion**

Anti-angiogenesis can effectively suppress tumor growth. Endostatin was found to specifically inhibit endothelial cell proliferation but not actively proliferating cells such as carcinoma cells, intestinal cells, epithelial cells, bone marrow cells[1]. So the application of endostatin did not cause bone marrow ablation, stomach and intestinal reaction, and other side effects[5]. Moreover, endostatin could be used for a long time without drug resistance since its target cells, endothelial cells have a relatively stable gene structure and low mutation rate[5,6]. Consequently endostatin is an attractive new strategy in cancer therapy.

Recombinant endostatin protein has been used in clinical trials at present[9]. However poor solubility of recombinant endostatin protein and its high effective dose have limited its wide spread application[4]. Gene therapy by which endostatin can express in vivo is an efficient method for anti-angiogenesis cancer therapy[10-12].

The key point of gene therapy is to establish an effective gene delivering system. Vectors used to deliver genes nowadays mainly include viral and non-viral vectors. Because of limitations of viral vectors such as potent danger, immune response and limitations of the gene length[13-15], more and more researchers have payed their attention to non-viral vectors. In this paper, a highly safe vector pVAX1 authorized by FDA in clinical trial was selected to construct a recombinant vector expressing...
secretive endostatin which was named as pVAX-sEN. DNA sequencing indicated endostatin gene together with a signal peptide was correctly cloned. Animal experiments showed pVAX-sEN could efficiently suppress the growth of mouse hepatoma cells. This study provides an experimental basis for further study on safe and efficient gene therapy with endostatin.

Non-viral gene can be transferred by physical and chemical methods. The cationic liposome-mediated gene transfer system is popular because of its low immunity, high safety, easy preparation, no limitation to gene sequence and potential to be used repeatedly[16,17]. Gene transfer efficiency of liposome can be affected by many factors. Studies indicated that a suitable quantity ratio of DNA and liposome was one of the essential factors[18]. Naked DNA injection is another simple safe gene transfer method. Many researchers have verified that naked DNA administration could induce a high gene expression level[19,20].

In 2003 VEGF administered into cardiac muscle using naked DNA was applied to clinical trial[21]. In our study, cationic liposome was used to transfer endostatin DNA into mouse hepatoma cells in vivo. It showed that when the quantity ratio of liposome and DNA was 1:1, the expression of endostatin was pretty high. When the quantity ratio was 1:8, endostatin reached the highest expression (92 ng/mL). With the elevation of DNA dose, the expression of endostatin increased. When the DNA dose reached 20 µg, the expression of endostatin amounted to 152 ng/mL. However, if DNA dose reached 50 µg, the expression of endostatin dropped to 80 ng/mL. The possible reason was that the high dose of liposome in the complex might be deposited[19]. So the transfection efficiency was lower. Results also indicated that naked DNA could also get a pretty high gene expression (86 ng/mL). Our in vivo inhibition assay even showed that naked DNA could get an inhibition as efficient as DNA-liposome complex. In 2002, Shi et al.[22] compared the transfection rate of naked DNA with liposome-DNA complex and found the expression of IL-12 was equal. Their result is in accordance with ours. However, in vivo gene expression assay indicated that the peripheral expression of endostatin in naked DNA treated mice decreased more quickly than that in DNA-liposome complex treated mice.

In conclusion, we successfully constructed the recombinant vector secrettively expressing endostatin using highly safe vector pVAX1 named pVAX-sEN. Animal experiments showed it could efficiently suppress the growth of hepatoma cells. It is a promising method in cancer therapy. At the same time we compared the transfection efficiency of naked pVAX-sEN DNA with liposome-DNA complex. Results indicated naked DNA could produce both a high endostatin gene expression and a good tumor inhibition effect when used in situ, strongly suggesting that naked DNA administration is a simple, safe and efficient gene transfer method.

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