Enhanced shRNA delivery by the combination of polyethylenimine, ultrasound, and nanobubbles in liver cancer

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Abstract.

\textbf{BACKGROUND:} Traditional cancer treatments such as surgery, radiation, and chemotherapy destroy both cancer and normal cells, which limit their clinical application. It is difficult to achieve the best results for any liver cancer patients using any single treatment method. Gene therapy for HCC demands non-invasive, efficient, targeted and safe gene transfection strategies.

\textbf{OBJECTIVE:} In this study, a nonviral shRNA gene delivery system utilizing a combination of PEI, US, and NBs was developed for targeting survivin in liver Cancer.

\textbf{METHODS AND RESULTS:} The PEI-shRNA-NBs cumulated in the tumor tissue because of the EPR effect. By exposure to the US, micelles shRNA may be released from PEI-shRNA-NBs in tumor tissues and the shRNA then transmitted efficiently to cancer cells. Considerably enhanced therapeutic outcome was obtained with the gene silencing effect enhanced.

\textbf{CONCLUSIONS:} PEI-shRNA-NBs possess the potential to become promising tools intended for shRNA delivery.

Keywords: Nanobubbles, transfection, shRNA, gene carrier, ultrasound

1. Introduction

Hepatocellular carcinoma (HCC) has a poor prognosis and the therapeutic options currently available are largely ineffective [1]. RNA interference (RNAi) is a biological pathway of gene silencing. Many studies have demonstrated the great potential associated with RNAi in HCC [2]. Despite the great potentials, significant medical effect hasn’t been demonstrated. One major challenge is to deliver shRNA into cancer cells [3]. On account of the poor half-life of uncoated shRNA in the bloodstream and the widely existing RNAase, uncoated shRNA may be degraded quickly. Furthermore, large size shRNA and negatively charged backbone have low accumulation and are unable to enter the cell membrane [4].

Ultrasound (US)-mediated gene delivery (USMD) exhibits possibilities for increasing the efficiency of gene into tissues and cells, a technique generally known as sonoporation (the transient permeabilization of cell membrane) [5]. Nanobubbles (NBs) as cavitation nuclei oscillate and collapse under US
stimulation. Cell permeability can be altered and pores in the endothelial layer produced, which allow macromolecules to enter the target tissue [6]. As a physical transfer methodology, USMD methods have much preferred quality, include reduced toxicity, low immunogenicity, low invasiveness, organs targeted with high specificity, and broad usefulness [7].

NBs have been recommended as a prospective device for gene delivery [8]. Due to the large size, microbubbles cannot pass through the tumor vascular endothelium. NBs offer several advantages in targeted gene transfection. Due to the enhanced permeability and retention (EPR) effect, NB can accumulate in tumor tissue, which can increase cell permeability during US exposure. The sonoporation effects of NBs can also be detected under low frequency US exposure [9]. NBs has become a desirable tool for shRNA delivery.

Extracellular nucleases can degrade the released DNA at the target site and the cells are not efficient at taking DNA. NB therefore sends DNA with only a small amount of DNA uptake and expression. USMD combined with nonviral polycationic polymer carriers has also been used to enhance gene transfer efficiency. PEI is one of the most efficient non-targeted polycations utilized for gene delivery [10]. Acid-catalyzed polymerization of aziridine produces a highly branched network, which ensnares DNA. On account of the near neighborhood of numerous linker amino groups, PEI maintains a substantial buffering ability at almost any pH. Sonoporation can improve the efficiency of PEI [11].

Survival targeting strategies have become a promising strategy for HCC [12]. Steady survivin knockdown restricted HCC cell proliferation, increases sensitivity to apoptosis, imprisoned cell cycle in the G1 phase and also resulted in significant mitotic catastrophe. Additionally, cells stably expressing survivin shRNA in vivo study indicated that tumor growth is inhibited by spontaneous apoptosis of cancer cells and animal survival was significantly prolonged [13].

In this study, we evaluated the therapeutic potential of Polyethylenimine (PEI) conjugated shRNA with the use of nanobubbles (NBs) as carriers and ultrasound triggered release for enhanced delivery of complex to the cells for targeting anti-apoptosis gene survivin. Herein, we expect PEI-shRNA-NBs with effective shRNA protection and high transfection efficiency under US exposure, possess the potential to become promising tools intended for shRNA delivery.

2. Methods

2.1. Materials

Phospholipids including DSPC (1, 2-distearoyl-sn-glycero-3-phosphocholine), DSPE-PEG2000 (Distearoyl phosphoethanolamine-PEG2000) and DPPA (1,2-dipalmitoyl-sn-glycero-3-phosphate) were used in powdery form (Lipoid, Germany). Pluronic F-68 and bPEI (Branched polyethylenimine) from Sigma Aldrich. AnnexinV-FITC and PI from Key Gen Biotech (Nanjing, China). Dimethyl sulfoxide (DMSO) from Sigma (USA); Sulfur hexafluoride (SF₆) gas from the Specialty Gas plant.

2.2. Preparation of NBs

Chloroform: dimethyl carbinol (9-mL volume, 2:1, v/v) solution in a round-bottom flask were used to dissolve phospholipids (DSPC, DPPA and DSPE-PEG2000, molar ratio 80:12:8) dried by evaporation under reduced pressure at 50°C water bath until phospholipid membrane was formed. The membrane was then dissolved with 5 mL of liquid (10% glycerol and Pluronic F-68).
The air above the obtained liposomal suspension was substituted with replaced with SF$_6$. The NBs were finally produced by intense shaking using a shaker. After low speed centrifugation, prepared bubbles were divided into two parts. After removing the bottom liquid layer, the NBs in the lower fraction were collected. The NBs concentration was determined using a hemacytometer.

2.3. Preparation of PEI-shRNA-NBs

The survivin-sh plasmid, encoding green fluorescence protein, was designed to target survivin gene. Survivin shRNA plasmid (the sequence 5′-GCCGCAGTTTCTTCTGCT -3′) were synthesized in GenePharma Co. Ltd. Zero point five mg of shRNA and a designed amount of PEI was dissolved separately in PBS. Gel electrophoresis was performed to evaluate the shRNA condensation capability connected with PEI. PEI and shRNA were mixed together at various N/P ratios of PEI to shRNA (1: 4, 1: 2, 1:2:1, 3:1, and 6:1). After the development of the polyplex micelle, the binding ability of PEI and shRNA was checked. PEI-shRNA-NBs complexes were prepared by surface conjugation. Morphological characteristics of the NBs and PEI-shRNA-NBs were characterized using a TEM (H600, Hitachi, Japan). The average diameters and zeta potential of the NBs, PEI-shRNA, and PEI-shRNA-NBs were analyzed by Dynamic Light Scattering and zeta potential analyzer.

2.4. Investigation on cell sonoporation

To see the effects of US exposure upon cell membranes, SMMC-7721 cell was observed with the scanning electron microscope (JEOL JSM-6360LV, Japan). The ultrasound was generated by US system with a single-element 1.0 MHZ frequency transducer. The incubated MMC-7721 cells with PEI-shRNA-NBs solution were put under US exposure. Different intensity (0, 0.3, 0.5 and 1.1 W/cm$^2$) at a frequency of 1 MHz and pulse repetition frequency of 1 kHz with 50% of a duty cycle, was applied to the transducer for 60 s respectively. After US exposure, each sample was fixed, dehydrated and observed with SEM.

2.5. In vitro cell inhibition and apoptosis evaluation

There were five groups (n = 6 for all groups): (1) DMEM medium containing 10% FBS; (2) PEI-shRNA; (3) PEI-shRNA-NBs; (4) PEI-shRNA with US exposure; (5) PEI-shRNA-NBs with US exposure. The medium of SMMC-7721 cells was changed to 0.2 mL of the above-mentioned transfer solution and incubated for 10 min. The parameters of US exposure (groups 4 and 5) were as follows: intensity 1.1 W/ cm$^2$ and exposure time 1 min. Later on, the medium was altered. After incubation for 48 h, 120 µL of MTT solution was added and incubated for 4 h. Next, DMSO (1 mL) was added. The solution was blended till the precipitate was dissolved. The absorbance was measured at 492 nm using a universal microplate reader and the cell growth inhibition rate (IR) was calculated. The apoptosis rate was identified by flow cytometry. Grouping and processing methods were the same as previously mentioned. Annexin V-FITC (5 µL) and propidium iodide (PI, 5 µL) were added and the samples were analyzed in an hour on a FACSCalibur flow cytometer.

2.6. In vivo antitumor effect

Xenograft tumors were induced by the subcutaneous injection of SMMC-7721 cells. There were five groups of 6 each: (1) a negative control group [physiological saline solution]; (2) PEI-shRNA group; (3) PEI-shRNA-NBs group; (4) PEI-shRNA with US exposure group; (5) PEI-shRNA-NBs with US exposure group.
Table 1

<table>
<thead>
<tr>
<th>Preparation</th>
<th>Diameter (nm)</th>
<th>PI</th>
<th>ZP (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NBs</td>
<td>395.6 ± 9.6</td>
<td>0.16</td>
<td>−19.35 ± 1.1</td>
</tr>
<tr>
<td>PEI-shRNA</td>
<td>60.2 ± 1.8</td>
<td>0.13</td>
<td>+35.3 ± 1.9</td>
</tr>
<tr>
<td>PEI-shRNA-NBs</td>
<td>382.3 ± 8.9</td>
<td>0.12</td>
<td>+20.3 ± 1.6</td>
</tr>
</tbody>
</table>

Abb: PI, polydispersity index; ZP, zeta potential.

Fig. 1. Electrophoretic mobility of shRNA in agarose gel at various N/P ratios of PEI to shRNA (1:4, 1:2, 1:1, 2:1, 3:1, and 6:1).

exposure group. The PEI-shRNA-NBs containing 6 µg of plasmid complexes were prepared and injected through tail veins. The parameters of US exposure (groups 4 and 5) were as follows: intensity of 2.06 W/cm² and exposure to low-frequency US for 10 min at the designated points. We used a commercial clinical ultrasound instrument (Technos MPX; Esaote, Italy) with a linear array transducer to monitor the treatment processing. The xenograft tumor areas were localized before the nanobubbles were injected. With low mechanical index (MI 0.08), the arrival of the nanobubbles into the xenograft tumor areas and the entire treatment process can be visualized. The treatment was repeated every 5 days (20 days in total). Tumor sizes were measured every five days after treatment. All of the mice were euthanized, weighed and then sectioned to hematoxylin-eosin (HE) staining. The rate of inhibition of tumor volume and weight was assessed.

2.7. Immunohistochemistry

Tumor sections were used for immunohistochemical studies after undergoing a series of process. Primary antibody against survivin (1:200 dilution in 5% BSA), bcl-2 (1:100 dilution in 5% BSA) and bax (1:30 dilution in 5% BSA) were used. Immunohistochemical processes include dewaxing, washing, incubation, and blocking inactivation of endogenous peroxidase. The tissues were then incubated with primary antibody, washed with PBS, and further incubated with appropriate biotinylated secondary antibody.

2.8. Statistical analysis

The data was statistically analyzed with SPSS software (version 18.0, SPSS Inc.).

3. Results

3.1. Preparation of PEI-shRNA-NBs

Figure 1 represents the results of agarose gel electrophoresis. A clear DNA band could be seen in each lane with 1:4, 1:2, 1:1, and 2:1 of PEI and shRNA. The band (3:1) was considerably weakening. There
Table 2
Cell growth inhibition and apoptosis

<table>
<thead>
<tr>
<th>Group</th>
<th>Optical density</th>
<th>Inhibition rate (%)</th>
<th>Apoptotic cells (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) negative control</td>
<td>0.982 ± 0.029</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>b) PEI-shRNA</td>
<td>0.833 ± 0.027</td>
<td>15.1 ± 2.75</td>
<td>16.46 ± 1.08%</td>
</tr>
<tr>
<td>c) PEI-shRNA-NBs</td>
<td>0.773 ± 0.009</td>
<td>21.16 ± 2.82</td>
<td>19.49 ± 1.01%</td>
</tr>
<tr>
<td>d) PEI-shRNA with US</td>
<td>0.716 ± 0.011</td>
<td>27.07 ± 2.26</td>
<td>22.01 ± 1.52%</td>
</tr>
<tr>
<td>e) PEI-shRNA-NBs with US</td>
<td>0.541 ± 0.032</td>
<td>44.92 ± 3.48</td>
<td>37.1 ± 2.63%</td>
</tr>
</tbody>
</table>

Notes: *Comparison of experimental groups with the negative control group, P < 0.05, and #comparison of the PEI-shRNA-NBs with US group with other groups, P < 0.05. Abb: PEI, polyethylenimine; NBs, nanobubbles; US, ultrasound; PEI-shRNA-NBs, NBs bearing PEI-shRNA.

Fig. 2. TEM of (A) unloaded and (B) shRNA-loaded NBs.

was no band to be observed in the lane (6:1). These outcomes recommended that the ratio of 6:1 was the optimal proportion of all the plasmids combined with PEI. Both NBs and PEI-shRNA-NBs were spherical with particle size (300 to 500 nm). TEM showed the PEI-shRNA-NBs with a well-defined core-shell structure and the shell thickness was about 60 nm (Fig. 2). Unlike NBs with a fairly smooth surface, the PEI-shRNA-NBs had uneven surface. The average diameter, PI, and zeta potentials of NBs, PEI-shRNA, and PEI-shRNA-NBs were reported in Table 1. Weak positive charge prefers the adhesion of PEI-shRNA-NBs to cancer cells.

3.2. Investigation on cell sonoporation

Figure 3 shows SEM images of SMMC 7721 cells under different US exposure energy. Normally, the morphology of the cells was spherical and their surfaces were smooth (Fig. 3A). Under the 0.3 and 0.5 W/cm² ultrasound energy, conspicuous pits or pores on the cell membranes were not identified. However, when the acoustic-pressure amplitude was increased to 1.1 W/cm², rough regions and small pits began to appear on the cell membrane (Fig. 3D). Therefore, proper ultrasonic energy can improve the uptake of the nanocarrier while retaining a good morphology.

3.3. In vitro cell inhibition and apoptosis evaluation

The combination of PEI-shRNA-NBs with the US showed the best inhibition compared to the other groups (Table 2). The IRs of the PEI-shRNA and PEI-shRNA-NBs groups were 14.75% and 20.97%, respectively. After using ultrasound with the same plasmid concentration, the IR increased to 26.85% and
Table 3

<table>
<thead>
<tr>
<th>Group</th>
<th>Tumor volume (cm³)</th>
<th>Tumor mass (g)</th>
<th>IR of volume (%)</th>
<th>IR of weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) negative control</td>
<td>4.845 ± 1.944*</td>
<td>5.713 ± 1.365*</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>b) PEI-shRNA</td>
<td>4.458 ± 0.764*</td>
<td>4.223 ± 0.261*</td>
<td>8.0</td>
<td>17.5</td>
</tr>
<tr>
<td>c) PEI-shRNA-NBs</td>
<td>3.548 ± 1.053*</td>
<td>3.485 ± 0.444*</td>
<td>26.8</td>
<td>38.0</td>
</tr>
<tr>
<td>d) PEI-shRNA with US</td>
<td>3.440 ± 1.014*</td>
<td>3.062 ± 0.376*</td>
<td>29.0</td>
<td>46.4</td>
</tr>
<tr>
<td>e) PEI-shRNA-NBs with US</td>
<td>1.584 ± 0.513*#</td>
<td>1.977 ± 0.247*#</td>
<td>67.3</td>
<td>65.4</td>
</tr>
</tbody>
</table>

Notes: *Comparison of experimental groups with the negative control group, P < 0.05, and #comparison of the PEI-shRNA-NBs with US group with other groups, P < 0.05.

Fig. 3. SEM images of SMMC 7721 cells irradiated at an intensity of 0, 0.3, 0.5 and 1.1 W/cm² ultrasound.

44.78%. These data indicate that the PEI-shRNA-NBs combined with US were much more effective in inhibiting the cell proliferation. The number of apoptotic SMMC-7721 cells was observed. PEI-shRNA-NBs with US group achieved the highest apoptosis level of 37.41%, indicating that a combination of PEI, US, and NBs enhanced shRNA delivery. The percentages of apoptotic cells after treatment with PEI-shRNA-NBs with US, PEI-shRNA with US, PEI-shRNA-NBs, and PEI-shRNA were 37.1 ± 2.63%, 22.01 ± 1.52%, 19.49 ± 1.01%, and 16.46 ± 1.08% respectively. These results are similar to the MTT data.

3.4. Antitumor effect in vivo

After repeated treatments by PEI-shRNA-NBs with US exposure, tumor volume was much smaller than other groups from day 10 and tumor growth was efficiently slowed down (Fig. 4). Consistent with
In our previous in vitro findings, PEI-shRNA-NBs with US group yielded significantly higher volume and weight inhibition rates than the other groups (Table 3). In the control group, HE staining showed that the tumor cells actively proliferated and had low differentiation whereas in the treatment group, some tumor tissues showed obvious necrosis and hemorrhage, especially in the PEI-shRNA-NBs plus US group. Immunohistochemical assays of hepatoma sections from mice after the treatments provided further evidence of the expression of the surviving, bcl-2 and bax proteins. Tumor tissues from the PEI-shRNA-NBs with US group had the lowest level of surviving and bcl-2 protein and the highest level of bcl-2 protein whereas in the saline groups expression of surviving, bcl-2 and bax were reversed (Fig. 5). These outcomes exhibited that treatment method utilizing the PEI-shRNA-NBs by making use of US exposure could reduce the surviving and bcl-2 expression and to trigger the bax gene very efficiently.

4. Discussion

US is a common non-invasive and well-controlled visual tool that is inexpensive and safe for clinical applications. It has sufficient penetration depth to reach the remote target anatomy. Some scholars have studied MBs with US-mediated drug therapy for deep tumors. The study found that the combination of ultrasound, microbubbles and chemotherapy drugs can prolong the quality of life and extend survival in patients with inoperable pancreatic ductal adenocarcinoma [14]. The intimal barrier is a major factor in limiting gene therapy. With the recent developments in the study of micro- as well as nanoparticles, drug delivery has shown great potential for systemic treatment [15]. The cavitation effect caused by USMD enhances the permeability of the cell membrane and cause micro vascular rupture, widening the extracellular space, and the gene can pass through the ruptured microvessels and the extracellular space to the tissue cells for targeted delivery.

Gene therapy for HCC demands non-invasive, efficient, targeted and safe gene transfection strategies. MB combined with US has been used to enhance the efficiency of gene transfection and the ability to target cationic polymer micelles [16]. Nevertheless, MB with larger particle size is not able to penetrate
Fig. 5. Immunohistochemical assays of hepatoma sections from mice after the treatments. Brown staining indicated the expression of surviving, bcl-2 or bax protein, and the blue staining showed the nuclei, respectively (400×).

PEI as an efficient non-viral vector has been utilized successfully for in vivo [21]. Especially, Deshpande and Mrausinitz [22] documented that the combination of PEI and US can together improve the DNA transfection. In this work PEI-shRNA micelle were used instead of naked shRNA. We also
examined if ultrasound and PEI possess a synergistic effect to improve shRNA transfection in liver cancer. Vitro and vivo outcomes demonstrated PEI-shRNA-NBs combined with US therapy could reduce the number of viable liver cancer cells, induce apoptosis in cancer cells and inhibit tumor growth. Histopathological results showed that tumor structures in the PEI-shRNA-NBs with US were destroyed more completely than that of the other groups. The mechanisms involved may be as follows: First, when ultrasound irradiated the tumor area, due to the existence of NB, cavitations appear in the capillaries and tumor tissues, which improved permeability of the vascular endothelium and cancer cell membranes [23]. On the other hand, PEI-shRNA cationic polymers could protect the plasmid from degradation, and the positively charged surface also facilitates their delivery to cancer cells. In addition, gene could be focused on the target site by US regional irradiation.

Survivin, the smallest molecular weight member of apoptosis protein IAP (inhibitor) family, has been demonstrated to play a key role in tumorigenesis [24]. It is highly expressed in liver cancer cell lines and liver cancer tissues, and promotes the rapid deterioration of HCC through its anti-apoptosis, promoting mitosis, proliferation promotion and angiogenesis [25]. These biological functions make Survivin an attractive target for cancer treatment, including HCC treatment [26]. Several independent studies have shown the introduction of survivin siRNA oligonucleotides into the body for the treatment of liver cancer [27]. The bcl-2 proteins adjust a variety of steps in apoptosis as well as obstruct cell death, while pro-apoptotic proteins for instance bax encourage apoptosis [28]. In the study, immunohistochemical method was used to detect apoptosis-related proteins in tumor tissues. The results showed that the inhibition of survivin by management of PEI-shRNA-NBs led to apoptosis induction by down regulating bcl-2 and up regulating of bax.

5. Conclusion

In conclusion, we developed a nonviral shRNA gene delivery system utilizing a combination of PEI, US, and NBs for targeting survivin in liver cancer. The PEI-shRNA-NBs cumulated in the tumor tissue because of the EPR effect. By exposure to the US, micelles shRNA may be released from PEI-shRNA-NBs in tumor tissues and the shRNA then transmitted efficiently to cancer cells. According to vitro and vivo outcomes, PEI-shRNA-NBs possess great potential to become promising tools of shRNA delivery of targeted cancer therapy.

Acknowledgments

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Conflict of interest

None to report.
References


