Antitumor activities of dFv-LDP-AE: An enediyne-energized fusion protein targeting tumor-associated antigen gelatinases

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Abstract. Gelatinases play an important role in tumor growth and metastasis, and overexpression of these molecules is strongly correlated with poor prognosis in a variety of malignant tumors. Lidamycin is an enediyne antitumor antibiotic with potent cytotoxicity. We previously reported that a tandem scFv format (dFv-LDP-AE) showed enhanced binding ability with gelatinases compared with the scFv-lidamycin conjugate (Fv-LDP-AE). In this study, the antitumor activities of dFv-LDP-AE on hepatocellular carcinoma (HCC) were evaluated in vitro and in vivo. By SDS-PAGE analysis, it was found that partial fusion protein dFv-LDP existed as dimer; the results of ELISA and immunofluorescence demonstrated that the fusion protein dFv-LDP could efficiently bind to hepatoma cells in vitro. The apparent arrest of cell cycle at G2/M phase and induction of apoptosis at nanomole levels indicated that the dFv-LDP-AE was very potent against HCC. In in vivo experiments, dFv-LDP-AE shown enhanced cytotoxic effects compared to those of LDM. Administration at mouse tolerable dosage level, the inhibition rate of tumor growth was 89.5% of dFv-LDP-AE vs. 73.6% of LDM on hepatoma Bel-7402 in mice (P<0.05) and, 87.3% of dFv-LDP-AE vs. 63.4% of LDM on hepatoma Bel-7402 in mice (P<0.01). Small animal optical imaging showed that the FITC-labeled dFv-LDP preferentially localized in the tumor site in less than 30 min, which demonstrated remarkable tumor-targeting properties. Taken together with the above findings, the enediyne-energized fusion protein dFv-LDP-AE showed potential application as a new agent for therapeutic applications in HCC.

Introduction

Hepatocellular carcinoma (HCC) is one of the most common malignancies in the whole world. Each year more than 360,000 new cases are diagnosed with HCC in China and the incidence is continuing to rise (1). It is known that surgical resection is the primary curative treatment for HCC in clinic, however, the major treatment of patients with unresectable or metastatic hepatocellular carcinoma (HCC) is chemotherapy (2). Though numerous patients with HCC could be successfully induced into first remission by chemotherapy, most of them ultimately experienced a relapse within 3-5 years. In addition, chemotherapy is often accompanied with substantial side effects for the reason of non-specific cytotoxicity. Therefore, it is imperative to develop novel or other therapy approaches or agents for clinical HCC therapy. Targeted therapy by antibody-mediated therapy has emerged as a novel approach for the effective and innovative treatment of HCC (3,4).

Gelatinase (including matrix metalloproteinase (MMP)-2 and MMP-9), a member of MMPs, plays an important role in tumor growth and metastasis, and overexpression of these molecules is strongly correlated with poor prognosis in a variety of malignant tumors (5,6). It is known that the gelatinases are abundantly presented in certain hepatocellular carcinoma (7,8), therefore, gelatinases is a possible target for cancer therapy or cancer interference. Antibody targeting therapy of hepatoma is a new approach. To augment the antitumor efficiency, a potent cytotoxic drug conjugated with antibody has been attempted and already obtained distinct clinical outcome (9).

The enediyne family of antibiotics is among the most toxic antitumor compounds described to date (10). Lidamycin (LDM, also called C-1027) is an enediyne antibiotic with potent cytotoxicity causing DNA strand breaks at very low concentration (11). Therefore, lidamycin is a potential moiety or ‘warhead’ to conjugate with the antibody. It has been documented that antibody-lidamycin conjugate showed very good antitumor efficacy in vitro and in vivo (12-14). Our previous results demonstrated that a tandem scFv format conjugated with lidamycin (dFv-LDP-AE) improved the antitumor efficacy compared with the scFv-lidamycin conjugate, and showed excellent tumor targeting capability to certain...
tumor cell lines, which indicated that the dFv-LDP-AE is a promising anticancer agent for development (15). As HCC is a leading cause of cancer death in China, developing new therapeutic strategies is warranted. In this study, we observed the potential application of the tandem scFv-LDM conjugate (dFv-LDP-AE) in HCC. To evaluate the antitumor efficacy of dFv-LDP-AE, ELISA, immunofluorescence, cell cycle arrest and apoptosis experiments were performed, the inhibition of tumor growth and the targeting capability in vivo was also investigated.

Materials and methods

Materials. The preparation of the dFv-LDP fusion protein and its enediyne-energized product dFv-LDP-AE were described in our previous report (15). MTT and FITC were purchased from Sigma Chemical Co. (USA). Other chemical agents used were of analytical grade.

Cell culture. Human hepatoma Bel-7402 cells, and HepG2 cells were stored in our lab and cultured at 37°C in DMEM medium supplemented with 10% fetal bovine serum, penicillin G (100 U/ml) and streptomycin (100 µg/ml). All cell lines were passaged every 3 days and maintained in exponential growth to ~80% confluence for experiments.

Binding specificity of dFv-LDP with cancer cells by immunofluorescence. The human hepatoma Bel-7402 cells were grown on slides and fixed with ice-cold 70% methanol for 30 min at 4°C. Non-specific binding was blocked with 3% BSA-PBS. After washing with PBS, the cells were incubated with dFv-LDP. The cells were then overlaid with mouse anti-His-Tag monoclonal antibodies after being washed with PBS. Then, the slides were mounted with fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse antibody and then re-stained with propidium iodide (PI). Following a final washing step with PBS, the fluorescence images were captured by a fluorescence microscope.

Binding affinity assay. To quantitatively determine the binding affinity of fusion protein dFv-LDP to hepatoma cells, the cell-based ELISA was used. Briefly, serial dilutions of re-folded dFv-LDP or Fv-LDP in 1% BSA-PBS were added into Bel-7402 and HepG2 tumor cells in pre-coated plates, incubated and washed. Then, the plate was incubated with anti-His-tag HRP-conjugate and washed, 3,3',5,5'-tetramethylbenzidine was used as the chromogen for the color development, absorbance values at 450 nm were measured on microplate reader (Bio-Rad Laboratories).

MTT assay. Cells were detached by trypsinization, seeded at 3,000 cells/well in a 96-well plate (Costar, Cambridge, MA) overnight. Then different concentrations of LDM were added and incubated for an additional 48 h. The effect on cell growth was examined by MTT assay. Briefly, 20 µl of MTT solution (5 mg/ml) was added to each well and incubated at 37°C for 4 h. The supernatant was removed, and the MTT formazan formed by metabolically viable cells was dissolved in 150 µl of DMSO, and then monitored with a microplate reader (Bio-Rad Laboratories) at a wavelength of 570 nm. Survival ratio was calculated according to the following formula: Survival ratio = (A_{test}/A_{blank})/(A_{control}/A_{blank}) x 100%.

Cell cycle analysis. Cells were treated with dFv-LDP-AE (0.01 and 0.1 nM) up to 48 h. Floating and adherent cells were harvested and centrifuged at 500 x g for 5 min. Then cells were fixed in ice-cold 70% ethanol and stored at -20°C for 24 h before analysis. For cell cycle analysis, cells were washed twice in PBS and stained with 50 mg/ml propidium iodide and 200 µg/ml RNase A for 30 min. The samples were analyzed on a fluorescence-activated cell sorter (Becton-Dickinson).

FITC-Annexin V/PI apoptosis assay. Cells were collected and resuspended in 200 µl binding buffer. Then 10 µl FITC-labeled enhanced Annexin V and 100 ng propidium iodide were added. Upon incubation in the dark (15 min, room temperature or 30 min at 4°C), the samples were diluted with 500 µl binding buffer. Flow cytometry was carried out on a FACScan instrument (Becton-Dickinson) and data were processed by WinMDI/PC-software.

In vivo antitumor experiments. The antitumor experiments were carried out using two hepatoma cancer cell lines, the mouse hepatoma 22 (H22) and human hepatoma Bel-7402. Kunming (KM) mice and the male athymic nude mice (Balb/c nu/nu) were purchased from the institute of animal research, Chinese academy of medical science, and allowed to acclimatize in the institutional animal house for >5 days before use. The study protocols were in accordance with the regulations of Good Laboratory Practice for non-clinical laboratory studies of drugs issued by the National Scientific and Technologic Committee of People's Republic of China.

The mouse H22 animal experiment was performed with 7-week-old female Kunming mice. H22 cells suspended in saline were inoculated subcutaneously (Day 0) at the right axilla of the mouse with 1.5x10^6 cells/0.2 ml/mouse. The mice were randomized into seven groups with 10 mice in each group. On Day 2, dFv-LDP-AE was administered at doses of 0.75, 1.0 and 1.25 mg/kg of body weight. LDM, and dFv-LDP fusion protein was administered at doses of 0.06, and 1.25 mg/kg, respectively. All treatments were administered by intravenous injection into the tail vein in 0.2 ml of sterile PBS.

To further evaluate the antitumor efficacy of dFv-LDP-AE, a second animal experiment was performed with 16-18-week-old nude mice. Exponentially growing human hepatoma Bel-7402 cells were implanted into two male athymic nude mice by the subcutaneous injection of 5x10^6 cells on the right flank. Tumors resulting after 2 weeks in donor animals were aseptically dissected and mechanically minced. Pieces of tumor tissue (2 mm^3 in size) were transplanted (subcutaneously) by a trocar needle into nude mice. When tumors reached about 100 mg in size, the mice were randomized into treatment groups (n=6 per group). Then the treatments were started (intravenously injection twice approximately on Day 7 and 14 after tumor implantation), the nude mice were injected in the tail vein with dFv-LDP-AE at different doses (0.4, 0.6 and 0.8 mg/kg), and with LDM (0.05 mg/kg), dFv-LDP (0.8 mg/kg), respectively. Control nude mice were injected with saline. Tumor growth was measured with a caliper, and tumor volumes were calculated.
with the following formula: \( V = 0.5a \times b^2 \), where \( a \) and \( b \) are the long and the perpendicular short diameters of the tumor, respectively (15). The data are presented as the mean ± SD. Tumor growth curves were plotted of time against the mean tumor volume ± SD.

The inhibitory rates of tumor growth were calculated according to the weight of excised tumor. The Student’s t-test was used to determine statistically significant differences. \( P<0.05 \) was considered significant.

Fluorescein isothiocyanate (FITC) labeling of the dFv-LDP and the small animal optical imaging. FITC were purchased from Sigma and was dissolved in dimethylsulfoxide. The protein dFv-LDP was dialyzed against 0.1 M sodium bicarbonate (pH 9.6) three times. Then the protein and FITC solutions were mixed in a final volume of 2.5 ml and incubated for 8 h at 4°C in the dark with constant shaking, after that 25 µl ammonium chloride (5 mol/l) was added to terminate the reaction for another 2 h. The FITC-labeled dFv-LDP were separated from free FITC on a Sephadex G-25 column equilibrated with phosphate-buffered saline (PBS). The protein-FITC ratio was measured with the formula \( 3.1 \times A_{495}/(A_{280} - 0.31 \times A_{495}) \) and the concentration of labeled protein was determined by the formula \( (A_{280} - 0.31 \times A_{495})/1.4 \) (16). The small animal optical imaging was performed with an IVIS Imaging System (Xenogen) comprised of a highly sensitive, cooled CCD camera mounted in a light-tight specimen box. Images and measurements of fluorescence signals were acquired and analyzed using Living Image software (Xenogen). Ten to 15 min prior to \textit{in vivo} imaging, animals received the FITC-labeled dFv-LDP at 40 mg/kg by tail vein and were anesthetized using 1-3% isoflurane. Animals were placed onto the warmed stage inside the camera box and received continuous exposure to 1-2% isoflurane to sustain sedation during imaging. The imaging time was 20 sec and two to three mice were imaged at a time. The light emitted from the FITC was detected \textit{in vivo} by the IVIS Imaging System, digitized and electronically displayed as a pseudocolor overlay onto a gray scale animal image.

\textbf{Results}

\textit{The preparation of dFv-LDP fusion protein.} The fusion protein dFv-LDP was produced by \textit{E. coli} and expressed mainly as inclusion body. After purification and refolding as reported previously (15). In this study, the PAGE was used to analyze the fusion protein dFv-LDP under reduced and non-reduced conditions. As shown in Fig. 1, there is a tiny band under the non-reduced condition, which indicates that a small percent of dFv-LDP exists as a dimer.

\textit{Binding capability with tumor cells.} To investigate the binding of fusion protein dFv-LDP with HCC, enzyme-linked immunosorbent assay (ELISA) was used to determine the affinity of dFv-LDP with Bel-7402 and HepG 2 cell lines. As shown in Fig. 2, fusion protein dFv-LDP could bind with tumor cells in a dose-dependent and saturable manner (Fig. 2), and the affinity constant is about 2- or 3-fold increased compared to the Fv-LDP, which contained just a single scFv. The binding of dFv-LDP with tumor cells was also displayed by immunofluorescence, as shown in Fig. 3, which indicated that the
gelatinases are abundantly expressed around the tumor cells, and the fusion protein dFv-LDP could efficiently bind them.

**Cytotoxicity of dFv-LDP-AE in vitro.** As examined by MTT assay, the enediyne-energized fusion protein dFv-LDP-AE displayed potent cytotoxicity to two HCC cell lines. As shown in Fig. 4, the IC\textsubscript{50} of dFv-LDP-AE to Bel-7402 and HepG2 tumor cells were ~4.5x10^{-10} M and 7.7x10^{-10} M, respectively.

**Cell cycle analysis.** HepG2 and Bel-7402 tumor cells were PI stained and the cell cycle progression was evaluated using flow cytometry. As shown in Fig. 5, an apparent increase of cells in the G2 phase by 3.06-68.71% with a corresponding decrease of cells in the G1 phases, was observed in Bel-7402 cells upon treatment with 0.01 nM and 0.1 nM dFv-LDP-AE. Similarly, an increase of cells arrest in G2 phase by 7.4-49.26% with a corresponding decrease of cells in the G1 phase was detected in HepG2 cells after exposure to dFv-LDP-AE at indicated doses, thus, indicating that dFv-LDP-AE had potent inhibition on the progression of HCC tumor cells.

**Apoptosis analysis.** FITC-Annexin V/PI analysis showed that the energized fusion protein dFv-LDP-AE induced apoptosis in Bel-7402 and HepG2 cells. As shown in Fig. 6, 0.01 and 0.1 µM dFv-LDP-AE induced apoptosis in 15.48 and 34.7% of Bel-7402 cells, respectively. Similarly, 0.01 and 0.1 µM dFv-LDP-AE resulted in apoptosis in 9.9 and 35.16% of HepG2 cells, respectively.

**The antitumor effect of dFv-LDP-AE in vivo.** The in vivo antitumor effects of dFv-LDP-AE were investigated on the inhibition of subcutaneously transplanted murine H22 in Kunming mice. The tumor-bearing mice were treated by tail vein injection once at 48 h after transplantation. The results showed that dFv-LDP-AE significantly inhibited the growth of hepatoma 22 tumors (Table I). As evaluated on Day 12, dFv-LDP-AE at 1.25, 1.0 and 0.75 mg/kg suppressed the tumor growth by 89.5% (P<0.05 vs. LDM), 85.2% (P<0.01 vs. control) and 77.4% (P<0.01 vs. control), respectively. 

![Figure 3. The immunofluorescence analysis of fusion protein dFv-LDP with Bel-7402 cancer cells.](image)

![Figure 4. The cytotoxicity of energized fusion protein dFv-LDP-AE to Bel-7402 and HepG2 cells using MTT assay.](image)
Figure 5. Cell cycle analysis of cells treated with dFv-LDP-AE. Bel-7402 cells and HepG 2 cells were stained with PI after a 48-h exposure to 0.01 and 0.1 nM dFv-LDP-AE. Percentages of the total cell population in the different phases of the cell cycle were determined. Representative results of experiment are shown.

Figure 6. Apoptosis induced by dFv-LDP-AE. Cells were cultured for 48 h with control medium, or with 0.01 and 0.1 nM dFv-LDP-AE, respectively. Cells were then harvested and labeled with a combination of Annexin V-FITC and propidium iodide. The percentage of early apoptotic cells (lower right quadrant) and late apoptotic/necrotic cells (upper right quadrant) is indicated.

Table I. Tumor growth inhibition effects of dFv-LDP-AE on hepatoma 22 in mice.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Dose (mg/kg)</th>
<th>Mouse no. (begin/end)</th>
<th>BWC (mean ± SD, g)</th>
<th>Tumor weight (mean ± SD, g)</th>
<th>Inhibition ratio (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td></td>
<td>10/10</td>
<td>11.63±0.95</td>
<td>3.1±0.36</td>
<td>-</td>
</tr>
<tr>
<td>LDM</td>
<td>0.06</td>
<td>10/10</td>
<td>3.57±2.48</td>
<td>0.82±0.61</td>
<td>73.6</td>
</tr>
<tr>
<td>dFv-LDP-AE</td>
<td>0.75</td>
<td>10/10</td>
<td>3.28±1.46</td>
<td>0.7±0.44</td>
<td>77.4</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>10/10</td>
<td>3.36±1.24</td>
<td>0.46±0.32</td>
<td>85.2</td>
</tr>
<tr>
<td></td>
<td>1.25</td>
<td>10/10</td>
<td>2.79±2.47</td>
<td>0.33±0.21</td>
<td>89.5</td>
</tr>
<tr>
<td>dFv-LDP</td>
<td>1.25</td>
<td>10/10</td>
<td>10.56±2.17</td>
<td>1.93±0.70</td>
<td>37.6</td>
</tr>
</tbody>
</table>

*aCompared to the control, P<0.05; bcompared to LDM, P<0.05.
in nude mice; and the therapeutic efficacy of dFv-LDP-AE was apparently stronger than that of LDM (P<0.01), as compared under the tolerable dosage, i.e. the targeting delivery of LDM by an anti-gelatinase antibody fragment enhanced the anti-tumor effects significantly. It was also noteworthy that the enediyne-energized fusion protein dFv-LDP-AE was shown to be significantly more effective than dFv-LDP or LDM used alone in prolonging the survival of mice (P<0.05, Fig. 7B).

The small animal optical imaging using FITC-labeled fusion protein. We previously reported the tumor targeting capability of dFv-LDP to certain tumor cells and the injection dose was 20 mg/kg. As the FITC is not a very good fluorescence in imaging in vivo (17), its wavelength is relatively short and it could not penetrate thick tissue, in order to make the in vivo imaging much more clear, the higher dose of 40 mg/kg FITC-labeled fusion protein dFv-LDP was injected via tail vein. As
shown in Fig. 8, we observed significant tumor retention of FITC-labeled dFv-LDP in the tumor foci, the FITC-labeled dFv-LDP saturated the tumor and its surrounding sites within 30 min after injection. The signal of fluorescence was still visible at the tumor site and maintaining the signal at 12 h after administration. As we expected, the dFv-LDP demonstrated very good tumor targeting potential, which was mainly located at tumor sites. However, the question regarding the dFv-LDP with high liver and kidney uptake remained unsolved, therefore causing a problem in the systemic toxicity to normal organs.

Discussion

In this study, we demonstrate that the tandem anti-gelatinase scFv format improved the biodistribution and therefore enhanced the antitumor effects of LDM. By covalent association of a tandem anti-gelatinase scFv to the N' terminal of LDM, the accumulation of LDM in the peritumor tissue could be increased, and the tumor killing capability, the G2/M phase arrest or the induction of apoptosis would be greatly enhanced. Therefore, the inhibition rate of LDM and dFv-LDP-AE on tumor growth is apparently improved from 73.6 to 89.5% in hepatoma 22 xenograft tumor model and from 63.4 to 87.3% in hepatoma Bel-7402 xenograft tumor model. The explanation for this difference is the tumor targeting ability of the tandem scFv format. From the results of the optical imaging, the degree and specificity of tumor localization were due to dFv-LDP-AE, which was presumably as a result of its affinity with gelatinase overexpression in tumor sites.

Gelatinases are abundantly overexpressed in HCC and its stromal cells, and play an important role in tumor microenvironment (18,19). In this study, we found that the ‘naked’ fusion protein dFv-LDP had an inhibition rate of 38.8% at a dosage of 0.8 mg/kg in Bel-7402 xenograft, which indicated that the fusion protein dFv-LDP not only played as a carrier of AE, but also possessed certain antitumor efficacy in vivo. This might be attributed to the dual effects of dFv-LDP on tumor growth inhibition and the disturbed balance of tumor microenvironment.

In the small animal optical imaging experiment, the fast tumor location of dFv-LDP at 30 min after administration was observed. When accumulation of dFv-LDP-AE in the target tissue is achieved, therapeutic efficacy is displayed by a potent tumor-killing power of AE. The rapid accumulation of dFv-LDP-AE in tumor site indicated that the dFv-LDP-AE might not decompose or decay before being delivered to the target site. Thus, the dFv-LDP-AE effects could be internalization or display of a ‘bystander’ effect, killing nearby tumor cells in addition to the directly targeted cells.

In summary, in this study we demonstrated that the tandem format dFv-LDP-AE showed improved antitumor effect and biodistribution compared to the LDM by increasing the targeting delivery of LDM indicating that the gelatinases (MMP-2 and MMP-9) are a valid target for antibody-directed therapy. Although, more studies are warranted to further explore targeting ability and decreasing the immunogenicity for the murine origin of the scFv used, as well as lowering the kidney and liver adsorption.

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References