INTRODUCTION.

Cancer targeting gene-viro-therapy (CTGVT) approach has become a hotspot and a trend in the field of cancer biotherapy and oncolytic adenovirus is an ideal vector to carry the targeting genes. In this study, we used human telomerase reverse transcriptase (hTERT) promoter to control the adenovirus early region 1a (E1A) and the human α-fetoprotein (AFP) promoter integrated with hypoxia response element (HRE) to control the adenovirus early region 1b (E1B). Then the novel double-regulated adenovirus Ad-hTERT-HREAF (named SG505) was engineered. The short-hairpin RNA against focal adhesion kinase (FAK) was inserted into SG505 and thus forming Ad-hTERT-HREAF-shRNA (called SG505-siFAK). Then various oncolytic adenoviruses were examined to verify whether they could suppress liver cancer cells selectively and efficiently, therefore SG505-siFAK could be a potential agent for future clinical trials of hepatocellular carcinoma.

Abstract. Cancer targeting gene-viro-therapy (CTGVT) approach has become a hotspot and a trend in the field of cancer biotherapy and oncolytic adenovirus is an ideal vector to carry the targeting genes. In this study, we used human telomerase reverse transcriptase (hTERT) promoter to control the adenovirus early region 1a (E1A) and the human α-fetoprotein (AFP) promoter integrated with hypoxia response element (HRE) to control the adenovirus early region 1b (E1B). Then the novel double-regulated adenovirus Ad-hTERT-HREAF (named SG505) was engineered. The short-hairpin RNA against focal adhesion kinase (FAK) was inserted into SG505 and thus forming Ad-hTERT-HREAF-shRNA (called SG505-siFAK). Then various oncolytic adenoviruses were examined to verify whether they could suppress liver cancer cells selectively and efficiently both in vitro and in vivo. Both replicative and replication-defective adenoviruses carrying FAK-shRNA significantly inhibited the expression of FAK in Hep3B and SMMC-7721 cell lines and efficiently suppressed the growth of liver cancer cell lines with minor effect to normal cells. Furthermore, the recombined oncolytic adenoviruses, SG505-siFAK, SG505-EGFP and SG505 were able to selectively propagate in AFP-positive liver cancer cells in vitro and the SG505-siFAK efficiently suppressed the expression of FAK. SG505-siFAK showed the most potent tumor inhibition capability among the three recombined adenovirus with IC50 levels of 0.092±0.009 and 0.424±0.414 pfu/cell in the Hep3B and HepG2 cell line, respectively. Animal experiment further confirmed that SG505-siFAK achieved the most significant tumor inhibition of Hep3B liver cancer xenograft growth by intratumoral injection comparing to the intravenous injection among the three recombined viruses. Immunohistochemical results indicated that FAK expression was downregulated significantly in the tumors treated with SG505-siFAK. The dual-regulated oncolytic adenovirus SG505-siFAK was proven to inhibit the growth of liver cancer cells selectively and efficiently, therefore SG505-siFAK could be a potential candidate for future clinical trials of hepatocellular carcinoma.

Introduction.

Hepatocellular carcinoma (HCC) is one of the most frequently diagnosed cancers and is also one of the major causes of tumor-related deaths worldwide (1). With the improvement of the surgical techniques and the application of transarterial chemoembolization (TACE), the radiofrequency ablation (RFA), and liver transplantation, the overall survival and the prognosis have made great progress and the 5-year survival rate of the early diagnosed HCC with curative treatments ranges from 50 to 70%, but the recurrence of early-stage HCC after resection occur in ~20, 50 and 75% of patients at 1, 3 and 5 years, respectively (2,3). Therefore, the recurrence and the metastasis after resections are intractable problems for achieving total control in HCC treatment, while there are few drugs to resolve the crucial problem. At present, the only approved systematic therapy for the advanced-stage HCC patients is the multikinase tyrosine kinase inhibitor sorafenib which used alone improves the median overall survival by ~3 months (4,5). Thus, it is urgent to pursue new agents to improve the prognosis of HCC.

Cancer targeting gene-viro-therapy (CTGVT) is a promising approach to conquer the malignant tumor as it is endowed with the ability to selectively infect and damage the tumor cells with minimum harm to the normal tissue (6). The new strategy takes advantage of gene therapy and virotherapy by utilizing the oncolytic adenovirus containing the anticancer gene, which produces more effective antitumor effects than either gene therapy or the viral therapy alone (7,8). The adenovirus vectors are the most common used gene delivery system, especially the adenovirus serotype 5 (Wad5) due to its excellent characteristics. Adenovirus can replicate in both dividing and non-dividing cells and it hardly integrates into the host genome posing low risk of mutagenesis. The easy manipulation and the capability to produce high titers of virus are favorable. Moreover, the adenovirus activates the humoral and cellular immune response which might activate immune system for the recognition of tumor antigens. The side effects of adenovirus therapy are mild and rare. So adenovirus is an ideal candidate...
vector for the CTGVT (9). The oncolytic adenovirus, also called conditionally replicating adenovirus, could theoretically selectively propagate in and lyse the tumor cells and the released progeny virus will evade the neighboring tumor cells and the virus replication will stop in the normal cells (9,10). To achieve the targeting therapy of HCC and safety for the normal cells, a dual-regulated oncolytic vector by using liver cancer-specific promoters was then constructed.

Human telomerase reverse transcriptase (htERT) is highly active in >80% of human tumor cells but not in most normal cells and the hTERT promoter has been cloned by several scientists and was utilized to drive the exogenous gene expression (11,12). The adenovirus early region 1a (E1A) plays a central role in the virus replication and cell cycle and the E1A promoter can be replaced by the hTERT promoter (13). However, hTERT is also expressed in the hematopoietic stem cells and generative cells, so the system has a potential detriment to the normal cells and will produce side effects in humans. To minimize the potentially adverse effects to normal cells, more delicate regulatory systems should be engineered. Human α-fetoprotein (AFP) is re-expressed in ~70% of the HCC, so the AFP promoter is widely used to drive the adenovirus E1B gene as an excellent tool for HCC targeting therapy (14,15). The system can be used to target to the AFP-positive HCC specially, but does not work in the low-AFP-generating cells or AFP-negative cells so we created a fused AFP promoter, HREA, by connecting the hypoxia reactive element (HRE) enhancer with the AFP and the system is able to enhance the replication capability of oncolytic adenviruses in both AFP-positive and AFP-negative liver cancer cells (16,17). However, there exists a disadvantage in this system that it can also kill the hepatic stem cells which also express AFP, thus weakening the hepatic reserve and worsening the prognosis of the HCC patients. It has been reported that the HREA promoter enhanced the translational strength and selectivity of the oncolytic virus to achieve HCC-specific and tumor environment-selective viral replication and tumor killing (17,18). In order to achieve the hepatoma-restricted cytoxity and enhanced replication, we integrated the hTERT promoter and HREA promoter into our oncolytic adenovirus where the hTERT promoter drove the E1A expression and the HREAf drove the E1B expression, then the Ad-hTERT-HREAf was constructed and we named it SG505.

Previous studies revealed that the focal adhesion kinase (FAK) expression was upregulated in HCC and the high expression was associated with a poor prognosis (19,20). FAK besides contributing to the invasion and metastasis of the HCC, also proved to be an important mediator of cell adhesion, growth, proliferation, survival, angiogenesis and migration which were often disturbed in the cancer cells, so it was a promising therapeutic target for the HCC (21). Herein, we created the short-hairpin RNA against FAK and inserted the gene in the recombined oncolytic adenovirus to confirm the antitumor effect and the new system was Ad-hTERT-HREAf-shRNA, or SG505-shFAK for short. We also created other recombinant adenoviruses, namely, Ad-hTERT-HREAf-EGFP-shRNA (SG505-EGFP), Ad-PPE3-siFAK, and Ad-DC311-siFAK. In this study, we explored the selective and efficient antitumor effect of the recombined oncolytic adenovirus to offer better treatments for HCC.

Materials and methods

Cell lines and culture conditions. HEK293 cells (embryonic kidney cell line containing the E1A region of serotype adenovirus 5) were purchased from Canadian Microbiex Biosystem Inc. Hep3B and HepG2 (AFP-positive human liver cancer cell), SMMC-7721 (AFP-negative human liver cancer cell), L02 (human normal liver cell line), BJ (skin fibroblasts), PANCl-1 (poorly differentiated pancreatic carcinoma cell line) and H460 (human large cell lung carcinoma cell line) were obtained from Cell Bank of Shanghai Institutes for Biological Sciences of the Chinese Academy of Sciences (Shanghai, China). Cells were cultured in DMEM or RPMI-1640 supplemented with 10% fetal bovine serum, 4 mmol/l glutamine, 50 U/ml penicillin and 50 µg/ml streptomycin under an atmosphere of 95% air and 5% CO₂.

Recombinant adenviruses. Short hairpin RNA targeting FAK was constructed in our lab. The hybrid HREAf promoter and hTERT were generated referring to the previous studies (16,22). Therefore, the shuttle plasmids Ad-DC311-siFAK, Ad-PPE3-siFAK and the dual-specific antitumor oncolytic adenvirus SG505, SG505-EGFP, SG505-shFAK were constructed. Briefly, shuttle plasmids were constructed to generate the recombinant. Packing and production of the recombinant adenviruses were performed in the HEK-293 cells using Lipofectamine 2000 (Gibco BRL, Grand Island, NY, USA) according to the manufacturer’s protocol. Viral plaques appeared 9–14 days after cotransfection and were sublimated three times. The recombinant adenviruses were identified separately by PCR. The viral titers were determined by the tissue culture infectious dose 50 (TCID₅₀) assay in HEK293 cells.

Western blot analysis. Cells were infected with the recombinant adenviruses at a MOI of 3 pfu/cell. The cells were trypsinized, harvested and resuspended in RIPA lysis buffer 3 days later. The protein concentration was confirmed by the avidin-biotin complex (ABC) technique as described by the manufacturer. Then, protein samples were separated by 10-15% SDS-polyacrylamide gel electrophoresis and transferred to polyvinylidene fluoride (PVDF) membranes. Membranes were blocked in 5% bovine serum albumin (BSA) and incubated with primary antibodies at 4°C overnight and then were detected by the appropriate secondary antibodies marked with horseradish peroxidase at 37°C for 2 h. Signals were visualized via the enhanced chemiluminescence (ECL) Western blotting substrate kit. Extracts of uninfected cells were used as the negative control and β-actin was used as the internal control.

MTT assay and BrdU incorporation assay. The MTT colorimetric assay was carried out to detect cell viability (23,24). To assess the cytotoxic effect of the virus, cells on logarithmic phase were seeded onto 96-well plates at 1x10³-2x10³ cells per well and infected with oncolytic adenovirus. The viruses were diluted by serum-free culture solution. The cells were then infected with various concentrations (at MOI of 0.1, 0.2, 0.5, 1, 2, 5, 10 and 100 pfu/cell) of recombinant adenovirus 24 h later. The cells without virus infection were used as controls
and 4 duplications were set corresponding to a certain MOI value. The media was replaced by 100 µl phosphate-buffered saline (PBS) per well. The cells were treated with 5 µl MTT (5 mg/ml) and incubating for 4 h at 37˚C to allow MTT metabolization. Then the culture medium was removed and the crystals formed were dissolved in 150 µl MTT lysate solution (70% dimethylsulfoxide and 0.02 mol/l hydrochloric acid) for 4 h at 37˚C. Cell viability was determined by measuring absorbance at 595 nm, using a microplate reader. The controls at each day were set as 100% viability. The percentage of cell survival rate treated with the recombinant adenovirus was expressed using the following formula: [100% x (absorbance value of experimental cells) / (absorbance value of control cells)]. All measurements were performed in triplicate. Bromodeoxyuridine (BrdU) incorporation was measured using BrdU Cell Proliferation Assay Chemicon cat. no. 2750. Transfected cells were incubated with BrdU for the last 12 h of the 72-h treatment with FAK siRNA and BrdU incorporation was assessed as described in the manufacturer's protocol. The concentration resulting in 50% of cell growth inhibition (IC50) was calculated using SPSS version 17.0.

**Real-time RT-PCR analysis.** Real-time PCR was carried out on the Real-Time PCR system (Applied Biosystems) with the following conditions: 95˚C, 15 sec for 1 cycle; then 95˚C, 5 sec; and 60˚C, 30 sec for 45 cycles. The mRNA expression of FAK was measured on ABI 7,500 Real-Time PCR system with β-actin used as a control. Each experiment was repeated three times. RNA levels were calculated from the mean relative quantity (RQ) (RQ=2^(-∆∆Ct)).

**Tumor xenograft experiment.** Eighty female BALB/c nude mice (4-week-old) were purchased from the Shanghai Experimental Animal Center (Shanghai, China). To establish xenograft tumors, 5x10^6 Hep3B cells in 500 µl normal saline were subcutaneously injected into the right flank of each mouse. When the largest diameter of tumors reached 5 mm, 40 mice were divided randomly into four groups (10 mice per group). PBS or 2x10^8 pfu of Wad5, SG505, SG505-siFAK were administrated to each mouse by intravenous injection every other day into the tumors five times. All injections were given every 2 days during the first week (days 3, 5 and 7 after grouping) and once weekly for 2 more weeks (days 14 and 21 so on after grouping). The remaining 40 nude mice received the treatments and measurements as described above by intratumoral injection. Tumor size was measured using calipers twice a week and calculated with a formula of [(0.52 x (smallest diameter)^2 x (largest diameter)] (25), tumor inhibition rates was calculated using the formula (1 - tumor weight of experimental group/average tumor weight of control group) x 100%. During the animal experiment, all animals were monitored daily and sacrificed at the end of the experiment. Ethics approval was granted by the Ethics Committee of the Affiliated Sixth People's Hospital of Shanghai Jiao Tong University.

**Immunohistochemical analysis.** Tumors were harvested and fixed in 4% paraformaldehyde, embedded in paraffin and cut into 4-µm sections for immunohistochemical analysis after sacrifice. These sections were stained with monoclonal anti-FAK antibodies at a 1:500 dilution. The slides were then washed with PBS and incubated with the avidin-biotin-peroxidase complex reagent. Then the differential expression of FAK in various tissues were studied. The criteria applied to evaluate the FAK staining intensity were described previously (19).

**Statistical analysis.** Statistical analysis was carried out by SPSS Statistical analysis. All continuous data were presented as mean ± standard deviation (SD). Categorical variables were compared by the χ^2 test or Fisher's exact test. The independent sample t-test or ANOVA was used to compare the mean values of different groups. P-value <0.05 was considered statistically significant.

**Results**

**Construction and characterization of the recombinant oncolytic adenovirus.** SG505-siFAK was a dual-regulated oncolytic adenovirus with E1A driven by the hTERT promoter and E1B driven by the HREAF promoter and it was armed with the therapeutic gene siFAK. The SG505 was blank vector and SG505-EGFP was armed by the reporter gene enhanced green fluorescent protein (EGFP). Replicative Ad-PDC311-siFAK virus and replication-defective Ad-PPE3-siFAK virus were initially introduced by the siFAK to confirm the function of RNA interference targeting to the FAK (Fig. 1A). The sequences of the inserted parts in all of the recombinant vectors were analyzed and proved to be correct.

It has been reported that the FAK was overexpressed in the HCC tumor but not in the normal tissue, so we constructed a targeting siFAK which will inhibit the FAK expression by the RAN interference technology (19,20,26). The synthetic gene was then inserted into the replicative adenovirus Ad-PPE3 and replication-defective adenovirus AD-DC311. To evaluate FAK expression in tumor cells, we employed the real-time PCR to analyze the levels of the FAK mRNA in vitro. All of the samples were analyzed and the expression of β-actin mRNA transcripts were used as internal control. We found that the relative expression level of FAK mRNA in Hep3B cells infected by the Ad-PPE3-siFAK was significantly lower than that infected by the Ad-PPE3 (0.13±0.015 vs. 1.00±0.06; P<0.01) and the relative expression level of FAK mRNA in Hep3B cells infected by the Ad-DC311-siFAK was also significantly lower than that infected by the Ad-DC311 (0.39±0.021 vs. 1.00±0.044; P<0.01), so we can confirm that the FAK-siRNA has potential to be an effective therapeutic gene for HCC (Fig. 1C).

As FAK is a potential therapy targeting the tumor and knockdown of the FAK suppressed the growth of the HCC, as established herein, to evaluate the antitumor ability of the oncolytic adenovirus expressing the FAK shRNA in vitro, we measured the cell viability by MTT assay. We found that the liver cancer cell lines (Hep3B and SMCC7721) were sensitive to the Ad-PPE3 and Ad-DC311 infection, whereas the normal cell lines (BJ and L02) showed much lower sensitivity, demonstrating safety of the vectors expressing siFAK to the normal cells (Fig. 1D). Similar results were also confirmed by BrdU incorporation assay (Fig. 1E). Collectively, these data indicated that RNA interference against FAK functioned as a suppressive factor of FAK. The primary studies paved the way for further investigations for HCC treatment.
Selective replicative ability of the recombined oncolytic adenovirus to AFP-positive liver cancer cells. To confirm the virus infective ability, the virus yields of different viruses were assayed using the standard tissue culture infectious dose 50 (TCID$_{50}$) assay. Virus replication rates were determined at 96 h after the cells were infected with Wad5, SG505, SG505-EGFP and SG505-siFAK separately at a MOI of 5 pfu/cell. Previous studies have already assayed the AFP secretion ability of various liver cell line, which indicated that Hep3B and HepG2 cell lines secreted AFP at level of 1,174±17.33 and 38.20±1.29 ng/ml and the secretion of AFP by L02 and SMMC-7721 was undetectable (7). The recombined oncolytic adenoviruses showed more potent replicative abilities in the Hep3B and HepG2 cell lines with the replication increasing by ~10$^8$-fold, compared to the AFP-negative cell lines, including the SMMC-7721, L02, BJ, PANC-1 and H460 cell lines (P<0.05) (Fig. 2A). Notably, Wad5 displayed the strongest replicative ability among various kinds of viruses in the cell lines. Although the recombined adenoviruses were slightly low in virus yield compared to Wad5, it was potent enough to drive the expression of the exogenous gene. The FAK expression was significantly downregulated in the Hep3B and HepG2 cell lines infected by the SG505-siFAK (Fig. 2B). Therefore, it also confirmed that SG505 was a promising vector to deliver the siFAK which could effectively suppress the FAK expression. In addition, the virus titers of SG505, SG505-EGFP and SG505-siFAK in the normal BJ and L02 cell were remarkably weaker than that of Wad5, which indicated less damage to the normal cells. The results demonstrated that recombined adenoviruses had significantly higher replication rates in the AFP-positive liver cancer Hep3B, and HepG2 cells and the safety concerning the recombined adenoviruses was remarkably balanced.

To further investigate the selective invasion of the dual-regulated oncolytic adenovirus, we applied western blot analysis to analyze the E1A and E1B expression in different cell lines infected by various viruses. E1A and E1B gene expression in the infected cells provided an independent verification of virus replication because they were essential for the virus replication (27-29). In BJ cells, it was clear that detectable levels of E1A and E1B were only produced by the Wad5-infected cells while they were not detected in the other
four oncolytic adenoviruses (Fig. 3A). Besides, in the normal liver cell line (L02), the E1A and E1B were weakly expressed in the cells infected by SG505-siFAK, SG505 and SG505-EGFP compared to Wad5, which confirmed the safety to the normal viable cells (Fig 3A). The expression of E1A and E1B in HepG2 and Hep3B infected by the oncolytic adenovirus was comparable to the ones by Wad5, which implied that the oncolytic adenoviruses produced large amounts of progeny viruses (Fig. 4B).

The cytotoxic effects of dual-regulated oncolytic adenovirus in AFP-positive liver cancer cell lines. Next, we investigated the antitumor effect of the viruses on cell viability. The AFP-positive liver cancer cell lines (Hep3B and HepG2), AFP-negative liver cancer cell line (SMMC-7721), human normal liver cell line (L02), PANC-1 and H460 were infected with Wad5, SG505, SG505-siFAK and SG505-EGFP at consistent MOIs ranging from 0.1 to 100 pfu/cell. Overall, the cell viability declined gradually with the increasing
Figure 2. The replication ability of the oncolytic adenoviruses in various cell lines. (A) Replicative ability of Wad5, SG505, SG505-EGFP and SG505-siFAK in various cell lines are shown. Cells were infected with 5x10^5 pfu of the indicated viruses (MOI=5) and viruses were harvested 96 h later, virus titers were qualified by the TCID_{50} assay. Amplification fold (yield virus/input virus) is shown. The bars represent the mean amplification fold of each adenovirus infecting different cells; the data are given as the mean ± SD of triplicate samples. (B) Analysis of FAK expression in the liver cancer cells. Western blotting was applied to analyze the FAK expression in AFP-positive liver cancer cell lines Hep3B and HepG2. β-actin expression was used as internal control.

Figure 3. Western blot analysis of E1A and E1B gene expression in BJ, L02 cells (A) and Hep3B and HepG2 cells (B). Cells were infected by Wad5, SG505, SG505-siFAK and SG505-EGFP at a MOI of 3 pfu/cell, and were harvested after 3 days, the lysates were analyzed by western blot analysis. The β-actin expression served as a loading control.
MOI values in various cell lines. SG505-siFAK and Wad5 showed most evident cytotoxic effects in the Hep3B and HepG2 cell lines. To further study the cytotoxic effects of the viruses, we compared the IC₅₀ values in the cell lines infected with Wad5, SG505, SG505-EGFP and SG505-siFAK. The IC₅₀ values of SG505-siFAK were 0.092±0.009, 0.424±0.414, 14.796±2.520, 48.709±0.927, 5.970±0.945 and 2.710±0.244 pfu/cell in Hep3B, HepG2, SMMC-7721, l02, PANC-1 and H460, respectively. The IC₅₀ values further confirmed that the SG505-siFAK showed strong cytotoxic effects to the AFP-positive tumor cell line, and was weakly cytotoxic in the pancreatic cancer line PANC-1 and large cell lung cancer cell line H460. Although Wad5 showed similar cytotoxic effects in Hep3B and HepG2 comparing to the SG505-siFAK, it had more potent cytotoxic effect to the normal liver cell l02 (P<0.01), therefore SG505-siFAK was mild to the normal liver cell comparing to Wad5.

Antitumor effect of dual-regulated oncolytic adenovirus in the nude mouse xenografts. To evaluate and compare the antitumor activity of the recombined oncolytic adenovirus in vivo, a liver cancer cell tumor xenograft model was established by inoculating AFP-positive Hep3B cells into the right back of nude mice. Mice were monitored for tumor volume and body weight for 28 days after inoculation. The antitumor effect was measured by the growth delay and body weight loss. The results showed that SG505-siFAK had the strongest antitumor activity, followed by Wad5, SG505, SG505-EGFP, and control group. The tumor volume and body weight loss were significantly reduced in mice treated with SG505-siFAK compared to those in control group. The results indicated that the dual-regulated oncolytic adenovirus had a great potential for treating liver cancer in vivo.
the nude mice. When the long diameter reached 5 mm, PBS, SG505-EGFP and SG505-siFAK (2×10⁸ pfu/mice per time) were administered to the mice by either intravenous injection or intratumoral injection in order to determine which kind of administration would cause more potent antitumor effect. Each nude mouse in the experimental group received a total of 1×10⁹ pfu of virus and all of the mice were sacrificed after 8 weeks. The volumes of tumors in the mice treated with PBS

Figure 5. SG505-siFAK inhibited the growth of xenograft tumors. (A) The tumor volumes of different groups treated with PBS, Wad5, SG505 and SG505-siFAK by intravenous or intratumoral injection. The tumor volumes treated intravenously with PBS, Wad5, SG505 and SG505-siFAK at day 56 were 3,237.5±223.2, 645.6±107.6, 1,168.3±38.0 and 659.2±82.4 mm³, respectively. The tumor volumes treated intratumorally with PBS, Wad5, SG505 and SG505-siFAK at day 56 were 2,952.1±435.3, 456.5±155.7, 1,339.2±147.6 and 379.6±95.6 mm³. Tumor volumes were recorded as mm³ ± SD, points indicated the mean value (n=10) and bars indicated SD; *P<0.01; †P<0.05; NS, no significant difference. (C) Data collected 56 days after the initial virus treatment were analyzed to compare the tumor inhibition rates between the intravenous group and intratumoral injection. In the intravenous injection group, tumor inhibition rates of Wad5, SG505 and SG505-siFAK were 80.06, 54.14 and 79.63%, respectively; in the intratumoral injection group, tumor inhibition rates of Wad5, SG505 and SG505-siFAK were 84.54, 56.64 and 87.14%, respectively. (D) Representative micrographs of immunohistochemical staining for FAK in the groups. The specimens from various tumors injected with different adenoviruses were assayed by IHC and they revealed evidence of antitumor effects of SG505-siFAK in vivo. The FAK in cytoplasm of tumor cell was yellow or brown. FAK in the cells treated by the SG505-siFAK was stained negatively and FAK in the cells treated by the PBS, Wad5 and SG505 was stained positively, implying the successful gene knock-down by the shRNA. Arrow indicated cells that stained positively for FAK.
in both group grew rapidly and reached ~1,300 mm^3 within 4 weeks, suggesting the malignancy of the Hep3B in the study. SG505 and SG505-siFAK began to show remarkable inhibition effect by day 14 compared to the blank control group. Nude mice treated with SG505-siFAK at a total dosage of 1x10^7 pfu per mouse showed similar antitumor ability when compared to the Wad5 in both groups (P>0.05) and stronger antitumor efficiency than SG505-EGFP in each group (P<0.01). It is worth noting that SG505-siFAK resulted in potent tumor growth inhibition without tumor regression, which resembled survival with tumor in the clinical scenario (Fig. 5A and B). The discrepancy of antitumor efficiency between two groups was accomplished by comparing the tumor inhibition rates, which demonstrated that SG505-siFAK and Wad5 had more robust antitumor capability in the intratumoral injection group than that in the intravenous injection (Fig. 5C). The main reason for the phenomenon was that the oncolytic adenovirus would be feasibly trapped and eliminated in the liver and the intratumoral injection could directly produce faster antitumor effects.

To confirm that the antitumor efficiency was mediated by the FAK siRNA engineered in the dual-regulated adenovirus, expression of FAK was assayed by immunohistochemistry in the tumor specimen when the mice were sacrificed. The FAK expression levels in the tumors treated with SG505-siFAK intratumorally or intravenously were markedly lower than those treated with PBS or SG505. Although SG505-siFAK and Wad5 showed similar antitumor capability, the FAK expression in the tumor tissue infected by the former was positive, and weakly positively infected by the latter (Fig. 5D). We thus deduced that SG505-siFAK might exert antitumor efficiency in vivo by combining the selective suppression effects of double-regulated oncolytic adenovirus to the liver cancer cells and the downregulation of FAK expression.

Discussion

In this study, we constructed double-regulated oncolytic adenovirus which proved to be a safe and efficient approach to suppress the development of AFP-positive liver cancer both in vitro and in vivo. Several promoters have been applied to achieve the selectivity for the tumor cells in the oncolytic systems, such as HRE, the prostate-specific antigen, osteocalcin, MUC1, midkine, Lplastin, E2F-1, UPII and Survivin genes (30-32). The promoters of these reconstructed oncolytic adenoviruses can be only activated in certain tumor cells which produced the special transcription factor and finally resulted in massive virus replication. We were inspired by the previous meaningful studies and decided to utilize the hTERT promoter to drive the E1A expression and hybrid HREAF promoter to drive the E1B expression, thus, SG505 was engineered based on the framework of Wad5 and the targeting therapeutic gene was introduced into the vector. The reason why AFP-promoter was chosen was that AFP is a widely-used clinical marker and it was feasible to choose suitable objects to receive the treatments. The recombined virus replication was potent in the AFP-positive liver cell lines (Hep3B and HepG2), exceeding that in the AFP-negative liver cell line (SMMC-7721), large cell lung cancer line (H460) and pancreatic cancer line (PANC-1) in spite of the enhanced HREAP promoter. Therefore, the recombined adenovirus showed specific infection capability in the AFP-positive liver cancer cells. SG505-siFAK was designed to selectively invade the AFP-positive liver cancer cell lines (Hep3B and HepG2) and produced more progeny viruses compared with the normal cell lines (BJ and L02). SG505-siFAK and SG505-EGFP derived from the SG505 shown similar selective replication (Fig. 2A). The western blotting also demonstrated the replication abilities of SG505, SG505-EGFP and SG505-siFAK were attenuated in the normal cells but were weaker than the wild adenovirus (Fig. 3). Thus, our data demonstrated the SG505 was a safe and efficient vector in the CTGVT.

The current strategy of CTGVT is to insert the targeting gene in the oncolytic adenovirus (33), therefore an appropriate antitumor gene is crucial for the success of CTGVT. FAK has been reported in the liver cancer tissue and proved to be a potential therapeutic target for HCC. The present reports clarified that FAK was required for c-Met/β-catenin-driven hepatocarcinogenesis and activation of the FAK- Src signaling pathways contributes to HCC growth and metastasis, so gene knockdown of FAK provided a promising strategy to treat HCC (34,35). Therefore, we decided to introduce a robust tool to knock down the FAK expression by RNA interference (36,37). RNA interference has rapidly become a powerful tool for target validation and the most widely used gene-silencing technique in functional genomics due to its extremely high inhibitory activity and the fact that the inhibition is very specific (38). Our results demonstrated that a significant suppression of SG505-siFAK was achieved in AFP-positive Hep3B and HepG2 cell lines without causing too much damage to the normal liver cells (Fig. 4), which rendered the double-regulated oncolytic adenovirus remarkable safety. Regarding to the antitumor efficiency, The IC_{50} value of SG505 in Hep3B and HepG2 was 1.199±0.073 and 15.026±0.926 pfu/cell, respectively, which was larger than that of SG505-siFAK (P<0.05). The results meant that the CTGVT therapy combined the benefits of virotherapy and gene therapy, producing comprehensive and synergetic effect (39).

We further confirmed the strong antitumor ability of SG505-siFAK in nude mouse xenograft model (Fig. 5). The SG505-siFAK induced more significant tumor regression than the SG505-EGFP in both intratumoral injection group and intravenous group. Besides, the SG505 was able to produce a considerable tumor growth inhibitory effect compared with the blank control group. It indicated that oncolytic adenovirus could cause tumor inhibition by itself and with the oncolytic adenovirus armed with antitumor gene. The potential side effects of the oncolytic adenovirus is hardly under the manipulation in nude mice because mice are known to be non-permissive to the human adenovirus, therefore the antitumor efficiency discrepancy among the Wad5 and SG505-siFAK was not remarkable.

Although RNAi was more potent than other gene silencing methods, several inherent limitations in RNAi technology need to be overcome before carrying out clinical trials, including transfection, incomplete silencing of tumor cells, low specificity, immunological reactions, undesired gene insertions (40). Above all, the RNA interference technology only knocked down gene expression but generally didn't eliminate it (41).
Consistent with the previous studies, our data also indicated that the tumors were not eradicated totally (Fig. 5) (42). To accomplish the total tumor inhibition, we postulated that two therapeutic genes could be used in the dual-regulated adenovirus vector, which will completely eliminate the hepatoma xenograft (39). The liver cancer suppressor gene included tumor necrosis factor-related apoptosis inducing ligand (TRAIL), hepatocellular carcinoma suppressor 1 (HCCCS1), and suppressor of cytokine signaling 3 (SOCS3) (43–45). The previous studies on the HCC and colorectal tumor xenografts have proved the viability and feasibility (7). Besides, the CTGVAT could also be combined with chemotherapeutic drugs, thus reducing the side effects and drug resistance (46).

In conclusion, the present data demonstrated that SG505-siFAK, a novel recombined dual-regulated oncolytic adenovirus, could infect and kill liver cancer cells specifically and effectively. In addition, our study provided evidence that siRNA directly delivered against FAK by the SG505 possessed the ability to downregulate the targeting gene and suppressed the progression of human liver cancer cells in the xenografts in the mouse model. Hence, the CTGVAT approach provides a promising method for the effective treatment of HCC and more studies are needed to verify its potential therapeutic effects in clinical trials.

Acknowledgements

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