Upregulation of Coxsackie Adenovirus Receptor Sensitizes Cisplatin-Resistant Lung Cancer Cells to CRAd-Induced Inhibition

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Abstract

Objective. Conditionally replicating adenoviruses (CRAds) have been proven potent oncolytic viruses in previous studies. They selectively replicate in the tumor cells because of incorporated survivin promoter and ultimately lead to their killing with minimal side effects on normal tissue. Chemotherapy with cisplatin is commonly employed for treating tumors, but its cytotoxic effects and development of resistance remained major concerns to be dealt with. The aim of this study was to explore the anticancer potential of survivin regulated CRAd alone or in combination with cisplatin in the A549 lung cancer cell line and cisplatin-resistant lung cancer cell line, A549-DDPR.

Methods. CRAd was genetically engineered in our laboratory by removing its E1B region and adding survivin promoter to control its replication. A549, H292, and H661 lung cancer cell lines were procured from the CAS-China. The anti-tumor effectiveness of combined treatment (cisplatin plus CRAd) was evaluated in vitro through MTS assays and in vivo through mouse model experimentation. RT-PCR was used to assess MDR gene and mRNA expression of coxsackie adenoviral receptor (CAR).

Results. Results of in vitro studies established that A549 lung cancer cells were highly sensitive to cisplatin showing dose-dependent inhibition. The resistant cells of A549-DDPR exhibited very less sensitivity to cisplatin but were infected with CRAd more efficiently as compared to A549. A549-DDPR cells exhibited higher expression of MDR gene and CAR in the RT-PCR analysis. The nearly similar rise in the CAR expression was seen when lung cancer cell lines received cisplatin in combined treatment (cisplatin plus CRAd). Combined anti-cancer therapy (cisplatin plus oncolytic virus) proved more efficient than monotherapy in the killing of cancer cells. Results of in vivo experiments recapitulated nearly similar tumor inhibition activities.

Conclusion. This study highlighted the significant role of survivin in gene therapy as it has the potential to render CRAd more tumor specific. It also establishes that higher CAR expression plays a vital role in the success of adenovirus-based therapies. Furthermore, a careful combination of chemotherapy drugs and oncolytic viruses can culminate in significant therapeutic achievements against cancer.

Key words: Lung cancer; Resistance; Conditional replication Adenovirus; Chemotherapy; Cisplatin; Survivin; CAR.

Introduction

Administration of chemotherapeutic agents coupled with exposure to radiation and surgical removal of tumor mass are treatment strategies widely used to combat cancer. A chemotherapeutic agent cisplatin or DDP is administered either alone or in combination with other chemo drugs to treat
cancer, but its clinical applications are affected by severe chemotoxic side effects and increasing drug resistance issues [1, 2]. Lung cancer is ranked second amongst all malignant tumors with the highest cancer-related mortality rate [3]. Small cell lung carcinoma is less common as compared to non-small cell lung carcinoma but more severe [4]. Beside achievements have been made in the treatment of lung cancer, recurrence has remained the issue nearly in all patients after initial treatment. This issue is may be due to drug resistance and metastasis of cancer [5, 6]. Research had been conducted to devise a combined therapeutic strategy by using cisplatin and adenoviruses together to obtain better results and to overcome the drawbacks of standard treatments [7, 8].

Oncolytic adenoviruses have been proven clinically as potent anticancer agents owing important advantages like broad cell targets, fewer side effects [9], inordinate gene carrying ability, and relatively lower capacity to alter host genes [10]. ONYX-015 was the first reported adenoviral vector that showed cancer-specific activity [11]. Although the monotherapy of adenovirus (Adv) was discouraged [12], it has been extensively engineered for combined therapies against cancer. Recent studies explained that CAR has a significant role in the entry of adenoviral vector into tumor cells. Investigations have established that the success of an Adv based gene therapy is dependent on higher expression of coxsackievirus/adenovirus receptor (CAR) on cancer cell surface because it is necessary for Adv internalization [13]. On the cell surface, following recognition, Adv is attached to CAR receptor. Interaction of viral pentone base motif with integrins, αvβ3, and αvβ5 facilitates its internalization into the cell [14]. Then through nuclear pore complex, the viral genome is translocated to nucleoplasm where its replication takes place [15].

Ads are genetically engineered to generate CRAds, which specifically replicate in and lyse the tumor cells [16] CRAds are generated either through gene deletion, as in Onyx-015 [17] or induction of promoter sequences like survivin, hTERT, and prostate specific antigen [18]. CRAds internalization into the cells depends on CAR expression which is usually quite low on tumor cell surfaces [19, 20]. Alteration in Adv tropism [21] and introduction of polylysine and heparin sulfate sequences in fiber knot domain has increased the internalization of CRAds [17]

p53, a tumor suppressor gene, is found to be altered in many types of human cancer [22]. In normal cells, its expression is low due to the absence of oncogenic activation [23]. World’s first commercially available gene therapy product of China in 2003, gendicine, also employed a wt p53 gene and delivered it via Adv 5 to treat head and neck squamous cell carcinoma, HNSCC [24]. The working principle of such gene-based therapies has been criticized by some studies which showed that the functional p53 genes might be present in some tumors [25] and Adv replication may be independent of p53 status [26].

Survivin, which is a protein of inhibitor of apoptosis (IAP) family, exhibited many advantages like high expression specifically in tumor cells, broad spectrum anti-tumor effects, and very little expression in normal cells. Owing to these qualities, the use of survivin in gene therapy is encouraged to acquire high tumor specificity [27, 28]. Many studies have reported that survivin expression is regulated at the transcriptional level [29, 30]. It has placed survivin among top five transcriptomes which are highly expressed in cancerous cells with minimal expression in normal cells of same tissue [31].

Currently, one of the major issues faced by the clinical oncologists which impede effective cancer treatment is rapidly developing resistance in tumor cells against chemotherapy agents. The synergistic inhibitory action of combination therapies, involving an oncolytic virus and a chemotherapy agent, has already been established by many investigations [32, 33]. We developed an oncolytic Adv, genetically engineered with a promoter (Sur-P) of IAP member, survivin. In this study, we also developed a cisplatin-resistant lung cancer cell line, A549-DDPR, to examine the impact of our treatment on the cancer cells resistant to chemotherapy. We performed in vitro and in vivo experiments to investigate the therapeutic effect and mechanism of Sur-P controlled CRAd alone or in combination with cisplatin in lung cancer cell lines, A549 and A549-DDPR.

Materials and Methods

Generation of cisplatin resistant cell line (A549-DDPR) and adenovirus

Cisplatin resistant lung cancer cell line, A549-DDPR, was developed from lung cancer cells of A549. Clinically relevant doses of cisplatin were given to A549 cells in a serum free medium for 60 minutes, and the dose was raised stepwise (0-76 μM), afterward, cells were cloned. A549-DDPR cell clone was selected, further treated with cisplatin (76 μM) and propagated for 40 passages with same cisplatin sensitivity. The ability of CRAds to replicate specifically in cancer cells has proven their use as effective anticancer agents. In this study, survivin responsive CRAds were prepared in which the survivin promoter controls the regulation of adenoviral E1A region while the adenoviral E1B
region was deleted. This virus showed efficient cancer selective phenotypes absent reduced anticancer activity. Firefly luciferase expressing Ad-Luc virus was used as a control.

Cell culture

Lung cancer cell lines A549, H661, and H292 used in this study were obtained from the Cell Collection Center, Shanghai (CAS-China). CRAd was multiplied by using a cell line which contains E1A region (HEK-293). RPMI-1640 medium (Gibco-BRL, HyClone) and DMEM containing 10% FBS were used for culturing and propagation of lung cancer cell lines and HEK-293 cell line.

In vitro analysis of tumor inhibition

In 24 wells plate, cells were placed at a concentration of 1×10^5 cells in each well. After 24 hours, different concentrations of CRAds viruses and DDP were given to cells in each well. By the addition of both cisplatin and CRAds serially, two treated cell groups were prepared. Serially treated cells were shifted to a 96 well plate with a concentration of 5×10^3 cells in each well. Then MTS/PMS reagents were added in the treated cells and cells were incubated at standard conditions (37°C, 5% CO2 for 180 minutes). Cells were monitored for five days then spectrophotometrically the absorbance was taken at 490 nm.

Semiquantitative reverse transcription-PCR

Total RNA was extracted by using TRIzol reagent (Ambion, life technologies) RNA samples were subjected to RT-PCR to obtain cDNA by using an RT-reagent kit (TakaRa). The primers employed in this study to amplify CAR (PCR product size 218bp), MDR (417bp) and GADPH control (371bp) are given below:

GAPDH: S: 5’GATTGTTGCCATCAACGACC3’
AS: 5’GTGCAGGATGCATTGCTGAC3’
CAR: S: 5’CCACCTCCAAAAGAGCGGTAC3’
AS: 5’ATCACAGGAATCGCACCC3’
MDR: S: 5’TCGTAGAAGTCTCCGTGGAT3’
AS: 5’CATTGGCGAGCCTGGTAG3’

Amplified PCR products were confirmed by performing 1% agarose gel electrophoresis. After electrophoresis, the gel was stained with ethidium bromide and visualized under UV light.

Tumor model

Six to eight-week-old female BALB/C nude mice were procured from the Chinese Academy of Sciences. Experimentations on animals were conducted by following the ethical guidelines from the NIH. A549 and A549-DDPR cells (4×10^6) were inoculated into the right flanks of the mice subcutaneously (SC). Tumors were visible on the 15th day after inoculation in both A549 and A549-DDPR cells.

In vivo analysis of tumor inhibition

Nude Mice with tumors were divided into four groups a, b, c, d; each group consisted of six mice. Each group was treated differently. Group ‘a’ was treated with only DDP, group ‘b’ was treated with only CRAds, and ‘c’ group was treated with both DDP and CRAds (DDP followed by CRAds order), and group ‘d’ received PBS. Ad-Luc control virus was added into each treatment group. After the one-month volume of tumor in each treatment group was measured to evaluate the effects of treatment.

Statistical analysis

Obtained data was subjected to Student’s t-test to find out the statistical significance, and data values are presented as with standard deviation. Data value at p < 0.05 was considered statistically significant.

Results

Sensitivity of A549 and A549-DDPR to cisplatin

Both cell lines, A549 and A549-DDPR, were treated with different concentrations of cisplatin and for cytotoxicity assessment, MTS/PMS assay was performed after 72 hr of the treatment. Dose-dependent inhibition of cancer cells was observed in both cell lines (Fig. 1) with maximum inhibition in A549 cells (75%) at 64 ug/ml of cisplatin dose. Figure 1 shows that inhibitory rate in A549-DDPR cells is very low even at higher cisplatin doses owing to their reduced sensitivity and high resistance to cisplatin. Cisplatin resistant cell line (A549-DDPR) remained stable throughout the period of observation. Severe cytotoxicity towards normal cells at higher chemo doses and increasing incidence of chemoresistance are the major concerns of oncologists worldwide [34, 35]. This study of combined treatment approach was aimed to get higher tumor inhibition rates with minimum damage to healthy cells.

Enhanced sensitivity of A549-DDPR cells to adenoviral infection

Likewise, to assess susceptibility to CRAds infection and to find optimum MOI, both cancer cell lines (A549 and A549-DDPR) were injected with different MOIs of adenovirus. The results of MTS/PMS assay showed that cisplatin resistant A549-DDPR cells exhibited much higher sensitivity to adenoviral infection as compared to A549 cells. Figure 2 indicates that at 100 MOI nearly
22 % tumor inhibition is achieved in A549 cell line while in cisplatin resistant cell line A549-DDPR, the inhibition rate was almost 2.5 folds higher (55 %). A similar trend was observed at higher doses of CRAd. These observations established that A549-DDPR cells were infected with CRAd more efficiently as compared to non-resistant cells of A549. This significant difference in CRAd transducing ability among two lung cancer cell lines is may be the result of variable CAR expression on their surfaces [2, 36, 37].

Figure 1. Sensitivity of A549 and A549-DDPR to cisplatin. Both cell lines, A549 and A549-DDPR, showed a large difference in response to variable concentrations of cisplatin when MTS/PMS assay was performed. A549-DDPR exhibit very less sensitivity to cisplatin due to resistance. The data in the line graph and error bars represent the mean of triplicate experiments and SD values respectively.

Figure 2. In vitro study to assess the sensitivity of A549-DDPR cells to CRAd. A549 and A549-DDPR cells were given different CRAd concentrations. A549-DDPR cells show comparatively very high inhibition rate which is most probably due to higher CAR expression. The data in the line graph and error bars represent the mean of triplicate experiments and SD values respectively.

Cisplatin increases the cancer cell-killing potential of CRAd in combined treatment

To evaluate that whether cisplatin could augment the tumor suppression capacity of Sur-p regulated CRAd in vitro, both lung cancer cell lines, A549 and A549-DDPR were infected with CRAd at various concentrations of cisplatin (0.25 ug – 64 ug/ml). All treatment groups in in vitro studies were infected with CRAd at 100 MOI. In combined treatment strategy following two sequence approaches were adopted: (a) infecting for 4 hours with CRAd at 100 MOI followed by cisplatin injections for 3 hours; (b) injecting for 3 hours various concentrations of cisplatin followed by infecting for 4 hours with CRAd at 100 MOI. MTS/PMS assay revealed that tumor suppressing effect of the combination treatment was more powerful as compared to monotherapy of DDP and CRAd. Figure 3 indicated that the inhibitory effect on A549 cell proliferation obtained by infecting with 100 MOI of CRAd plus cisplatin (1 ug/ml) was higher than that achieved with CRAd (800 MOI) alone. Synergistic therapeutic activity in cancer cells suppression was observed in combined treatment (Fig. 3, 4). Figures 3 and 4 showed that sequence-based approach b (DDP+CRAd) was more effective in tumor suppression as compared to approach a (CRAd+DDP). The difference was more prominent in non-resistant lung cancer cells of A549. The observations of in vitro studies established that cisplatin augmented the tumor inhibitory effect of CRAd-mediated biotherapy.

Figure 3. Enhanced tumor retarding efficacy of combined treatment in A549 cells. Sequence approach b (DDP+CRAd) appeared more successful as compared to sequence approach a. The data in the line graph and error bars represent the mean of triplicate experiments and SD respectively. p < 0.05 value was set as statistically significant for data.

Raised expression of CAR and MDR

We performed RT-PCR analysis to evaluate mRNA levels of MDR and CAR. The aim of RT-PCR experiments was to confirm the cisplatin resistance in A549-DDPR cells and furthermore to investigate the molecular mechanism which raised the anti-cancer activity of combined treatment (DDP plus CRAd). Specific primers for MDR, CAR, and GADPH
(internal control) were employed for PCR amplifications. This experiment pointed out that CAR expression was much higher in cisplatin-resistant A549-DDPR cells as compared to non-resistant lung cancer cells of A549. A similar trend was observed in lung cancer cell lines which received cisplatin treatment in combined therapy (DDP+CRAd) (Fig. 5). It can be hypothesized from these outcomes that lung cancer cell lines may be sensitized to adenoviral transduction by cisplatin. These results which linked enhanced transduction efficacy of adv with raised CAR expression were also previously confirmed by some other studies [36, 37]. Lung cancer cell line, A549-DDPR, showed higher expression of MDR gene which confirmed that it had developed resistance to cisplatin (Fig. 5).

**Synergistic therapeutic activity of combined therapy**

Our *in vitro* experiments showed that combined treatment has synergistic inhibitory action on cancer cell proliferation, and the significance of sequence in which combined treatment is given was also established. To discover whether the similar trends can be recapitulated *in vivo*, we employed 6-8 week old mice, and $8\times10^6$ A549 and A549-DDPR cells were injected in their right flank subcutaneously. When tumor volume reached 100-150 mm$^3$, mice were divided into four groups (n=6). We used 4 mg/kg of DDP intratumorally and 100 MOI of CRAd intraperitoneally in our *in vivo* studies. Three groups were treated with monotherapies of PBS, DDP, CRAd and one group received combined therapy of DDP plus CRAd. A line graph obtained from the observations of *in vivo* experiments evidenced an antitumor efficacy nearly similar to that was shown by *in vitro* studies. Figures 6A-D indicated noticeable differences in cancer size reduction efficacy among four treatment groups. The combined treatment group (DDP+CRAd) remained most powerful tumor inhibitor exhibiting synergistic therapeutic activity.

Lung cancer is a group of highly heterogenetic diseases. To further strengthen the results of our study we verified the tumor-inhibiting efficacy of CRAd and cisplatin *in vitro* in other lung cancer lines also. Two lung cancer cell lines, H292 and H661, were infected with CRAd at different MOIs, alone or with cisplatin. The inhibitory pattern observed in these cell lines was nearly similar to that was seen in A549 cells. The inhibitory effect of combined treatment (CRAd plus cisplatin) was found more potent than the effect of CRAd or cisplatin alone (Fig. 7A, B). Moreover, the decrease in the number of viable cells in the lung cancer cell lines obviously was time and dose-dependent.

**Discussion**

Gene therapy, manipulating adenoviral vectors to deliver therapeutic gene, has attracted many researchers and oncologists. Genetically engineered adenoviral vector was also employed to transport tumor suppressor gene p53 in world’s first commercially available gene therapy product, gendicine. Cancer therapies involving p53 gene has been criticized for being less potent and tumor protective [25, 38-40]. Chemotherapy, despite its severe damage to normal cells and resistance issues, has been the most efficient warhead against cancer so far. Cisplatin is a potent antineoplastic drug. It is well known for the formation of intrastrand crosslink
adducts to damage DNA and the activation of apoptosis which culminate in cancer cell killing [2]. Now, the trend has shifted towards combined treatment strategies involving gene therapy, virotherapy, and chemotherapy.

In the present study, instead of the p53 gene, we used survivin promoter (Sur-P) to modify adenoviral vector for specifically targeting cancer cells. Replication of ONYX-015 may not depend on the status of p53 [41], but its anti-cancer activity is beyond doubt. Our Sur-P incorporated adenoviral vector, CRAd, was nearly similar to ONYX-015 but was more tumor cell specific because Sur-P is overexpressed in almost all cancers with limited expression in normal cells [42, 43]. Results of this study showed that CRAd was successful in achieving high cancer cell specificity with very little toxicity towards normal cells. It is in agreement with many recent studies which supported Sur-P role in cancer gene therapy [29, 44, 45].

The results of our experiments established that a combined anti-cancer therapy (cisplatin plus oncolytic virus) was more efficient in the killing of cancer cells.
than the monotherapies of respective treatments. Many previous experiments have published the similar results [46, 47]. One of the major obstacles in the use of adenoviruses in gene therapy is their interaction and dependency on a transmembrane protein CAR for cell internalization. Many studies have pointed out that CAR has a very low expression in cancer cells [37, 48, 49]. In our study, RT-PCR analysis confirmed that cisplatin has the potential to enhance CAR expression on cancer cells. Cisplatin resistant lung cancer cell line, A549-DDPR, also exhibited high expression of CAR as well as MDR genes. Many late studies showed similar results [36, 51].

The objective of our study was to enhance tumor-specific killing with minimum toxicity towards normal cells and moreover to unveil the molecular mechanisms involved. Our strategy of combined treatment (DDP+CRAd) to combat lung cancer appeared synergistic in action to suppress tumor growth; this finding is in agreement with many recent studies [46, 47]. The viable number of mice was raised, and high rate of tumor inhibition was attained. It was because of the highly tumor-targeted oncolytic virus, CRAd, and reduced cisplatin dose.

Against lung cancer, it was the first study of this kind which used Sur-P regulated oncolytic virus in combination with cisplatin. A cisplatin-resistant lung cancer cell line, A549-DDPR, was also produced and employed in this study to examine the impact of treatment on resistant lung cancer cell line. Approximately 2.5 folds higher tumor inhibition was achieved in cisplatin resistant cell line as compared to non-resistant cell line, A549. These results suggested that adenoviral transgenic expression was probably enhanced in A549-DDPR because of drug resistance. We also witnessed in RT-PCR data that levels of CAR messenger RNA were comparatively higher in cisplatin-resistant lung cancer cell line. Recent studies reported similar results for cisplatin-resistant cells of other organs [36, 51]. This study has highlighted the significant role of the survivin promoter in gene therapy which is not restricted to lung cancer only. It further showed that the dose adjustment is very critical in combined therapy as the cells which are pre-exposed to cisplatin before adenoviral-mediated gene therapy and cisplatin-resistant cells showed enhanced adenoviral transduction.

It is concluded from this study that an anti-cancer therapy can be designed by carefully combining cisplatin and genetically engineered oncolytic virus to make significant therapeutic achievements in the war against cancer. Furthermore, adenovirus transduction efficiency should be carefully evaluated before performing gene therapy to adjust doses of a chemotherapy agent and adenovirus, because additive or synergistic action of combined therapy may culminate in the incidence of cytotoxicity or immune response activation.

Drug resistance has emerged as an unavoidable issue with chemotherapy. This study not only indicated the anticancer potential of combined treatment but also suggested a novel treatment approach of using CRAd as monotherapy in chemotherapy-resistant cancer patients. This research offered a highly tumor targeting oncolytic virus, CRAd, for further investigation as lethal warhead against cancer.

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Competing Interests

The authors have declared that no competing interest exists.

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