Adenoviral vector-based strategies against infectious disease and cancer

Chao Zhang and Dongming Zhou

Vaccine Research Center, Key Laboratory of Molecular Virology & Immunology, Institute Pasteur of Shanghai, Chinese Academy of Sciences, Shanghai, China

ABSTRACT
Adenoviral vectors are widely employed against infectious diseases or cancers, as they can elicit specific antibody responses and T cell responses when they are armed with foreign genes as vaccine carriers, and induce apoptosis of the cancer cells when they are genetically modified for cancer therapy. In this review, we summarize the biological characteristics of adenovirus (Ad) and the latest development of Ad vector-based strategies for the prevention and control of emerging infectious diseases or cancers. Strategies to circumvent the pre-existing neutralizing antibodies which dampen the immunogenicity of Ad-based vaccines are also discussed.

Introduction
Ad is non-enveloped, double stranded DNA virus with icosahedral capsids. It was first discovered by Rowe and his colleagues when they tried to culture the adenoid tissue in the laboratory in 1953. Ad infection is usually mild to human beings, but sometimes could be life threatening, especially to the immunocompromised individuals. In the 1970s, the United States army developed live Ad vaccines to prevent acute respiratory disease caused by AdHu4 and AdHu7. In 1991, Rosenfeld et al developed the first in vivo gene transfer using an Ad vector, and demonstrated that human α 1-antitrypsin gene delivered by the E1-E3-deleted Ad could be detected in the lung of a cotton rat. In 1993, the first human gene therapy study based on Ads was performed, a 23-year-old man with cystic fibrosis homologous received the first in vivo gene therapy with administration of an E1-E3-deleted rAd vector expressing the normal human CFTR, and the subsequent clinical studies were then initiated. In recent two decades, Ads have been widely applied as vaccine carriers since they are capable of eliciting T and B cell responses. Furthermore, Ads can be genetically modified to induce the apoptosis of the cancer cells, which are known as the oncolytic Ads. Ads are not only generally safe and can replicate in almost all the living cells, but also can be expanded easily in HEK293 cells and purified by CsCl gradient ultracentrifuge, and administered through oral, intranasal or intramuscular routes without adjuvants. Here, we review the Ads’ potential in vaccine development against infectious pathogens or in cancer treatment, and address the latest advances in the field.

Genome and structure of the Ad
Ads are DNA virus with icosahedral capsids of approximately 90 nm in diameter. Several studies have illustrated the structure of the Ads by cryo-electron microscopy. The genomic DNA of Ads is about 26–45 kb, with two inverted terminal repeats of 100–140 bp flanking at both ends. The genes that express during the life cycle of Ads are generally divided into two types: the early genes and the late genes. The early genes include E1A, E1B, E2, E3 and E4, and they are mainly responsible for facilitating the replication of Ads by changing the expression levels of related host genes. The early genes can be further classified into two types: the immediate early genes (E1A) and the delayed early genes (E1B, E2, E3 and E4). E1A promotes the expression of the delayed early genes. The E1B protein generally suppresses the apoptosis of the host cells by binding to p53, Bak and Bax proteins. The late genes are mainly responsible for the lysis of the host cells, assembly and release of the virions.

Biological characterization of Ad

Classification of the Ad
Ads are isolated from different mammalian species, such as human being, bovine and simian, among which the human Ads and chimpanzee Ads are widely used in the laboratory research or clinical study. Human Ads include more than 50 serotypes classified into subtype A to G, and chimpanzee Ads have more than 6 serotypes. Human Ads are distributed widespread in the nature and most people have been infected, thus high neutralizing antibody titers were detected among the population. Human subtype C Ads are the most common Ads which usually infect the children and cause upper respiratory tract infections or urinary tract infections. Human subtype B Ads sometimes cause severe eye or urinary tract infections. Some other serotypes, such as AdHu4 from subtype E, cause acute respiratory diseases. However, most of the Ad infections are mild, which promotes Ad vectors into a new era as vaccine carriers.

CONTACT Dongming Zhou dmzhou@sibs.ac.cn No. 320 Yueyang Road, Shanghai, 200031 China.
© 2016 Chao Zhang and Dongming Zhou. Published with license by Taylor & Francis. This is an Open Access article distributed under the terms of the Creative Commons Attribution-Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0/), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited. The moral rights of the named author(s) have been asserted.
DNA-associated proteins. The V protein mainly contacts with the nucleoli of the host cells and are involved in the viral assembly process.14,15 The VII protein plays a key role in the manipulation of DNA, such as the DNA binding, the initiation of DNA replication and the viral genome’s protection, et al.16-18 The X protein is responsible for the viral chromosome condensation.19 The capsid proteins comprise Hexon, Penton, fiber, IIIa, VIII and IX. There’re 240 trimers of hexons on the surface of the Ad virions, and the hexons are the major structural protein on the capsids.20 On the hexons, there’re several hypervariable regions which are the major neutralization sites for the Ads, and the hypervariable regions can be replaced with other foreign antigens as potential vaccine carriers.21-23 There have 12 pentamers of pentons on the top of the 12 icosahedral vertices, and they serve as the receptors for Ad internalization into the host cells,24 and each vertex has 12 trimers of fibers protruding from the capsid surface which are mainly responsible for the interaction with the cellular receptors to initiate the viral entry.10,25 The IIIa proteins are on the inner capsid surface, and are mainly responsible for the correct viral assembly, stabilization of the vertex region and the assembly of the packaged genome.11,26 The VI proteins are inside the capsids, linking the core to the icosahedral shell, and are critical lytic factors of Ads during the endosome disruption.27 The VIII proteins provide bonds between the peripental hexons and are involved in the stability of the capsids.28 The IX proteins function to dampen the innate immune response, and affect the viral tropism and stability of the capsids.29-30

Many Ads can be engineered for laboratory or clinical use. The recombinant Ads can be replication-incompetent or replication-competent. E1 gene is essential for the replication of the Ads, total or partial E1-deletion results that the vector can infect most of the living cells but cannot be expanded as it being replication-incompetent in normal cells. However, E1-deleted Ad can be propagated on dedicated helper cells, specialized cells that provide the E1 functions in trans, such as HEK 293 and PER.C6,31 in which E1-deleted Ad is replication-competent.

**Cellular receptors of the Ad**

The entry of the Ads into host cells is initiated by the binding of fiber knob to the cell receptors. The CAR functions as the receptor for the fiber protein in subtype A and C-F Ads,32-34 but in some cells, such as cancer cells and mature skeletal muscle cells, CAR is expressed in low levels.35-37 In CAR-deficient lymphocytes, subtype C Ads can cause latent infection,38 and the mutated CAR does not affect the tropism when Ads are administered systematically.39 These results suggest that there might be other receptors for Ads except CAR. In fact, many other receptors for the entry of Ad have been found, for examples, CD46 or DSG2 for subtype B Ads,40,41 and sialic acid and integrins, et al.42-45

After the binding of the fiber knob to receptors, the virion internalization starts through endocytosis. Generally, the endocytosis is initiated by the binding of penton bases to the integrins,46-48 but some reports revealed that the uptake of Ad virions could use lipid rafts or caveolae as entry route.49 A review article suggested that when the Ad virions were coated with charged polymers, the entry routine might be changed accordingly.50

**Ad vector-based vaccine candidates for infectious diseases**

Ad vectors are one of the most effective carriers for delivery of foreign antigens into the host cells. Compared with other viral vectors such as lentivirus, retrovirus and adeno-associated virus, etc, Ads are highly immunogenic and can induce both robust innate and adaptive immune responses in mammalian hosts. Ads have a large genome size, making the manipulation of the genetic DNA much more convenient. Unlike lentivirus or retrovirus, Ads do not integrate the viral genomic DNA into the hosts’ genome, which reduces the risk of insertion mutagenesis. Adeno-associated virus is less pathogenic than Ads, but it is not yet suitable for mass production.50 All above features make Ad a good vaccine carrier for the infectious diseases.

**Table 1** and **Table 2** show a list of vaccine candidates based on Ad vectors against some certain infectious diseases.

In the early stage, some Ads were modified as replicating-competent vectors with only E3 deletion which is not indispensable for the replication.89 Nowadays, most of Ad vectors are replication-deficient with the deletion of E1 or both E1 and E3. E1-deficient vectors can only be rescued and expanded in the E1-trans-compensating cell lines. E3 deletion increases the packaging capacity of the Ad vectors, such E1-E3-deleted Ads can be incorporated with up to 7–8 kb foreign genes.90 E4 can also be deleted, but the E4-deleted vectors only propagate in the E4-compensating cell lines.91 The fully gutted Ad vectors were developed with containing the replication origins and packaging signals while most of the viral coding sequences were deleted. The fully gutted Ads can only be amplified with the appearance of helper virus.92 Compared with the traditional Ad vectors, the fully gutted Ads have less toxicity caused by T cell responses, and the transgene products can be stably expressed.92,93

Different Ad vectors can elicit different immune responses in various laboratory animals or species.94 A study of SIV Gag-
specific CD8(+) T cell responses in mice vaccinated with AdHu5, AdHu26 and AdHu35, respectively demonstrated that AdHu5 was more immunogenic than AdHu26 and AdHu35, but AdHu26 and AdHu35 generated long-lived memory T cells, whereas AdHu5 elicited more terminally differentiated phenotypes of T cells. In another study, AdHu35, AdHu26 and AdHu48 were found to substantially produce higher levels of IFN-γ, IL-6 and 10-kDa gamma interferon-induced protein than AdHu5 in rhesus monkeys. Based on their different immunogenicity profiles, certain serotype of Ads can be selected alternatively for specific researches. Ad vectors can be administered by injection and oral immunization, both of which elicited well immune responses, whereas oral administration can elicit mucosa immune response compared to injection, and greatly circumvent the pre-existing anti-vector immunity.

HIV vaccine based on Ad vector

Safe and efficient HIV vaccine is urgently needed since HIV still remains a severe public health threat. Several strategies have been developed for HIV vaccine design, of which Ad vectors are widely tested. One of the most well-known clinical trials is AdHu5 based HIV vaccine which was developed by Merck, Inc. In the clinical trial, the replication-incompetent AdHu5 vectors encoding gag, pol and nef genes were administered to 1494 participants at a dose of 3×10^10 vp, while placebo administered to 1506 participants. This vaccine induced CD8+ T cell responses in homosexual men, but failed to prevent the HIV infection or reduce the early viral load. Further research revealed that the vaccine appeared to increase the risk of HIV infections in the AdHu5 serotype positive individuals. To explain the phenomenon, several studies have been performed and suggested that one possible mechanism was that the immune complexes of AdHu5 and anti-AdHu5 antibodies could activate the dendritic cells and CD4+ T cells which might serve as the targets for HIV infection.

After the failure of the Merck HIV vaccine trial, other improved strategies have been tested, such as the regimen of AdHu5 with AdHu5 boost. As reported by Churchyard GJ and Koup RA, et al, a DNA plasmid encoding multiple HIV genes from multiple clades for priming at 0,1 and 2 month respectively at a dose of 4 mg, and AdHu5 expressing multiple HIV genes for boosting at 6 month induced polyfunctional CD4+ and CD8+ T cells as well as the anti-envelop binding antibodies, which revealed the heterologous prime-boost regimen was a potent immunization strategy for inducing both antibody and T cell responses. However, similar strategy used in another clinical trial with the regimen of priming 4 mg DNA encoding multiple HIV genes at week 0, 4 and 8, respectively, and boosting with 10^10 pu rAdHu5 at week 24 reduced neither the rate of HIV-1 acquisition nor the viral-load set points in the participants.

As the AdHu5 based vaccine carriers are not suitable for HIV prevention, other serotype Ad based vaccines, such as AdHu26, AdHu35 or chimpanzee Ad vectors have been developed. In a study of AdHu26 expressing HIV-1 envelope as a new vaccine candidate, both the AdHu26-serotype positive and negative participants received a single intramuscular immunization with 5×10^10 vp rAdHu26. The result revealed that rAdHu26 elicited both systemic and mucosal envelop-specific humoral and cellular immune responses, but interestingly the individuals with pre-existing AdHu26-specific neutralizing antibodies had comparable immune responses to the AdHu26-serotype negative ones. HIV vaccines based on the rAdHu35 expressing the HIV-1 envelope antigen have been studied. In this phase Ib study, 192 healthy, HIV-uninfected participants were recruited and divided into one of following groups: rAdHu35/rAdHu5, DNA/rAdHu5, and DNA/rAdHu35 in AdHu5-seronegative persons, and DNA/rAdHu35 in AdHu5-seropositive persons, and a placebo group. The participants received three doses of 4 mg DNA or just one dose of 10^10 pu rAd at the first 2 month, then were boosted with 10^10 pu rAd at month 6. 4 weeks post boost, the immune responses were detected. The results indicated that all regimens were generally well tolerated and similarly immunogenic, and elicit cross-clade antibody responses including envelope V1/V2-specific IgG responses.

Presently, more novel Ad vectors are being discovered and developed for the HIV vaccine design, but none of them is ready for the market, thus more improvement is needed for the Ad vector-based HIV vaccine.

Influenza vaccine based on Ad vector

Ad vectors have been applied in the development of influenza vaccine. In most of Ad-based influenza vaccines, the influenza protein, such as HA, NP or M2 is expressed by the Ad vectors to induce neutralizing antibodies and T cell responses in the host. For example, HA protein of PR8 strain (H1N1) expressed by Ads can elicit HA-specific antibodies and cellular responses against the PR8 virus. Besides Ad vectored vaccine against particular strain of influenza virus, the universal influenza vaccines based on Ads have been explored. In the multivalent influenza vaccines based on the replication-incompetent AdHu5, HAs from different subtypes and NP from one subtype were expressed on the Ads. The mice were immunized intramuscularly with 10^10 PU of rAds twice at 4-week interval. 4 weeks post the boost, high levels of humoral and cellular immune responses were well induced and the mice were protected from lethal challenge with H5, H7 and H9 avian influenza virus subtypes. In another multivalent influenza vaccine based on AdHu4 and AdHu5, HA genes from the H1, H3, H5 subtypes of influenza virus were expressed by the Ad vectors, and then immunized mice by rAdHu4-prime/rAdHu5-boost regimen at a doses of ranging from 10^7 to 10^10 vp with a 4 week interval. The

---

**Table 2. List of rAd vaccine candidates in pre-clinical trials.**

| Pathogens | Ad vectors | Antigens | Study models | Clinical trials | References |
|-----------|------------|----------|--------------|----------------|------------|
| Rabies virus | AdHu5, Canine Ad type 2,AdC68 | GP | Mice, fox, dog, sheep, nonhuman primates | No | 63-66 |
| Dengue virus | AdHu5 | E, prM | Mice, nonhuman primates | No | 67-69 |
| MERS | AdHu5, AdHu41 | S | Mice | No | 70-71 |
vaccination results revealed that the highest dose vaccine groups were 100% protected from the heterologous lethal challenge of different subtypes of influenza virus, indicating that Ad-based multivalent influenza vaccines had great potential in the prevention and control of the influenza virus.

Ad-based influenza vaccines have been tested in clinical trials. A non-replicating AdHu5 vector expressing HA from avian influenza and a TLR3 ligand were tested in humans.60 Most of the participants received only one dose of Ads by capsule, with titers ranging from 10^8 to 10^10 IU, but some were boosted with another dose of 10^9 IU Ad at 4 weeks post prime. The vaccination results revealed that the antigen specific cytokotoxic and IFN-γ responses were induced in a dose dependent manner and cytotoxic responses increased after boost, which demonstrated that Ad-based vaccine administered orally could induce antigen specific immune responses and was safe as a vaccine candidate.60 AdHu4 based avian influenza vaccine was developed and tested in a clinical trial. AdHu4 expressing HA of H5N1 was orally administered 3 times at a dose of 10^7 to 10^10 vp within 56 days, and then boosted with 90 μg inactivated H5N1 viruses. The results demonstrated that cellular immune responses were well induced and oral administration of Ad might enhance the efficacy of poorly immunogenic vaccines such as H5N1, but the limitation of this study was that the HI titers were hardly measured.61 By orally administered replication-competent AdHu4 vaccine, another improved strategy was reported.111 In the study, the individuals were primed with AdHu4-H5-Vtn three times at dose of 10^7 to 10^10 vp in 56 days, and then boosted with the 90 μg inactivated H5N1 subunit vaccine at 3.5 to 12 months. The new regimen induced high HI titers compared with unprimed individuals, which compensated for the disadvantages in the previous vaccination routine.

With the development of RNAi technology, Ad based RNAi strategies have been applied in prevention and control of influenza infection. For example, a novel chimpanzee Ad termed as AdC68 was used as the microRNAs expression vector and tested in mice. AdC68-expressing amiRNAs targeting M1, M2 or NP genes of influenza virus could efficiently suppress the viral replication and confer complete protection from the lethal challenge of H9N2 and H5N1.112

**Ebola vaccine based on Ad vector**

Ebola virus was first discovered in 1976 with the outbreaks in Democratic Republic of Congo and Sudan, but the outbreak in Africa in 2013 made it a public concern again. There have been several strategies for the Ebola vaccine development, of which the Ad vectors are selected as a priority.

The NIH vaccine research center firstly developed a vaccine based on heterologous prime-boost regimen in 2000. In the research, the primates were firstly immunized with a DNA vaccine three times at 0, 4, 8 weeks at a dose of 4 mg, and then boosted with 10^10 PFU AdHu5 which expressed GP of Ebola virus 2 weeks post prime. The results showed high antibody titers and CD4+T cell responses were induced in vaccinated animals, and the vaccinated groups had a higher survival rates than the control groups after challenged with Ebola virus. Since this experiment took more than six months to complete the immunization flow chart, researchers developed an accelerated immunization method for the vaccination.117 In the accelerated experiments, the animals were given AdHu5 expressing GP and NP of Ebola virus twice at doses of 2×10^12 vp with a 9-week interval. The vaccinated animals were challenged with lethal Ebola virus and the protection was highly effective since the Ebola-specific CD8+ T cell and antibody responses were well induced. In the subsequent study, the animals were only primed with rAd-GP/NP and challenged 28 days later, but they still had high survival rates with either low or high doses of challenged virus. As the AdHu5-based Ebola vaccine showed a good prospect in the non-human primates, the first clinical trial based on AdHu5 was performed, with a recombinant vaccine encoding the envelope GP from the Zaire and Sudan Ebola virus tested in a randomized, placebo-controlled, double-blinded, phase I human study. Thirty-one healthy adults received a single dose of the rAdHu5 at 2×10^9 vp (n = 12), or 2×10^10 vp (n = 11) or placebo (n = 8). The results indicated the antibody responses to the two GPs in subjects were not well balanced. In the low dose group, antibody responses to Zaire GP were 50%, and 58% to Sudan GP. However, the antibody responses in the high dose group were 55% and 100%, respectively. In this study, the pre-existing neutralizing antibodies to AdHu5 was also noted, but it didn’t appear to affect the T cell response to Ebola GP since 32% to 82% subjects responded with more CD4+ than CD8+ T cells.

Recently, Ebola vaccines based on the other serotypes are being developed, one of which is the chimpanzee Ad serotype 3 (cAd3). In a clinical trial of cAd3 based Ebola vaccine, 20 volunteers were recruited and divided into 2 groups. The volunteers in each group received the rAds expressing the GP protein from both Zaire and Sudan Ebola virus. One group received high titers of Ads with 10^11 vp while another received 10^10 vp, and two individuals in the high dose group experienced transient fever. The final results demonstrated that the rAds induced well specific antibody responses and T-cell responses, with higher levels of responses in the high dose group.118 In another trial based on cAd3, 60 volunteers who were divided into 3 groups received a single dose of Ads ranging from 1×10^10 vp to 5×10^10 vp, and only 2 volunteers developed transient fever. After vaccination, specific antibody responses and T-cell responses were successfully elicited, but the levels were lower than those detected in the non-human primates.119 In a recent phase I clinical trial which was performed between Oct 8, 2014, and Feb 16, 2015, 91 participants in Mali and 20 in the USA were recruited to receive cAd3 expressing GP of Ebola with a single dose ranging from 10^10 to 10^11 pu. After the prime of Ads, some Malians were boosted with vaccinia Ankara expressing GP of Zaire Ebola virus and filovirus antigens (MVA-BN-Phyllo). The vaccination results showed that 1×10^11 pu single-dose rAds could suffice for an efficacy trials and the regimen of MVA-BN-Phyllo boosting could confer long-lived protection which might be needed for the health-care workers.77

**Other vaccines against infectious diseases based on Ad vector**

Ad vectored vaccines have been developed for some other infectious diseases besides influenza virus, HIV and Ebola virus. A tetavalent dengue virus vaccine based on the rAds was tested in non-
human primates. In the study, the prM and E gene from different subtypes of dengue virus were expressed by Ads. The vaccination included two doses of $10^9$ IU rAds administration with a 57-day interval, and 85 days or 253 days post prime, the animals were challenged with dengue virus. The vaccination results revealed that the animals produced high-titer antibodies that could neutralize all four serotypes of dengue viruses in vitro. The challenge studies showed that significant protection from viremia was observed against all four dengue virus serotypes, but the protection efficacy was better in dengue-1 and dengue-3 challenges than in dengue-2 and dengue-4 challenges.67

In addition to the Ad-based virus vaccine, Ads have been developed for the bacteria vaccine or protozoan vaccine. Mycobacterium tuberculosis causes serious bacterial infections in humans, and a vaccine based on AdHu5 expressing Ag85A has been tested in a phase 1 clinical trial. The results showed the polyfunctional CD4+ and CD8+ T cell responses were well stimulated, and the pre-existing neutralizing antibodies to AdHu5 had little influence on the potency of the vaccine.62 Malaria, which is caused by Plasmodium falciparum, poses a serious threat to public health. An AdHu5 vector encoding the apical membrane antigen 1 and circumsporozoite protein of P. falciparum was evaluated in a clinical trial.85 In the study, the DNA prime with Ad boost regimen was proved to be effective in eliciting specific T cell responses. Furthermore, some other serotypes of Ad, such as AdHu35 were developed for the malaria vaccines which listed in Table 1.

Human Ad serotypes such as AdHu5 have been extensively used for vaccine development mainly due to their excellent immunogenicity and safety. As the effect of pre-existing immunity on AdHu5-based vaccines, the clinical use of AdHu5 is greatly limited, while the rare human serotypes of Ads or non-human-originated Ads such as chimpanzee Ads have been extensively tested in both preclinical research and clinical trials.

**Ad vector-based cancer therapy**

Oncolytic Ads have shown great promise in cancer treatment since they exhibit distinct anti-cancer characteristics. During the life cycle of Ads, the Ad-infected cancer cells can be lysed in the end, and after the release of the Ad virions, they infect other cancer cells to initiate the next life cycle. Generally, two strategies are widely adopted for the modification of oncolytic Ads. The first one includes Ads expressing the therapeutic genes or combining RNAi technology to degrade the tumor promotion proteins. The second one mainly focuses on the capsids modification of the Ads, making Ads have specific tropism for the tumor cells or replicate to higher titers in the tumor tissues than in normal ones. These two strategies might be integrated to generate better anti-cancer effect.

One of the most well-known Ad-based anti-cancer drugs is the Advexin.120 It is an E1-E3-deleted AdHu5 vector expressing p53 under the drive of CMV promoter in the E1 region. The Advexin has been applied in multiple cancer treatments, such as head and neck cancer, breast cancer and colon cancer, et al.121-123 In the hand and neck squamous cell carcinoma, the Advexin was tested in a phase III clinical trial, patients were randomly treated with either Ad-p53 intratumorally on days 1 and 3 of each week at a daily dose of $2 \times 10^{12}$ vp or methotrexate once weekly at a starting dose of 40 mg/m², and each treatment cycle include 21 days. The results revealed that the vector was well tolerated and the anti-tumor activities were significant.124 Another well-known drug against glioma is Sitimagene Ceradenovec. The drug is an E1-E3-deleted, AdHu5-based vaccine that expresses the herpes simplex virus’ thymidine kinase at downstream of CMV promoter in E1 cassette. In the phase III trial, 250 patients were recruited with 124 in the AdHu5-treated group while 126 in the standard care group. Different groups received standard care plus injection of $1 \times 10^{12}$ vp rAd or just standard care alone. Almost all the individuals experienced adverse events in the trial. The clinical results suggested that use of Sitimagene Ceradenovec increased the survival time or re-intervention in patients with newly diagnosed supratentorial glioblastoma multiforme.125

Transduction of specific tumor antigen into dendritic cells is one of the most effective strategies against cancer. In an in vitro study, the DCs transduced with rAds expressing livin α induced strong specific cytotoxic T lymphocytes against different cancer cells.126 Ads can be armed with immune modulator such as GMCSF or REIC to induced cytotoxic T lymphocytes against cancers. In a clinical trial, the patients received the combination treatment of Ad-GMCSF and alkylation agents had higher survival rates than the ones only treated with alkylation agents, possibly due to the activation of anti-tumor T cells.127 In a E-G7 tumor-xenograft mouse model, Ad-REIC induced tumor-associated antigen specific cytotoxic T-lymphocytes, and the secreted REIC protein in the tumor generated a proper microenvironment for inducing of activated dendritic cells, resulting in decreased tumor size in the tumor-bearing mice receiving the Ad-REIC compared to the control groups.128

RNAi technology is widely used in the downregulation of the specific gene’s expression by sequence-specific degradation of the RISC complex.129 The application of RNAi technology based on Ads may be extremely effective since the small RNA molecules can be steadily expressed, thus the targeted protein remains at low levels for long. One of the firstly used RNAi technologies based on Ads was the vascular endothelial growth factor (VEGF)-specific targeting small RNAs. To induce and maintain the long-lasting silencing of VEGF, the study constructed E1A-mutated, E1B-deleted Ads with shVEGF expressing at the E3 region under the drive of U6 promoter. After the vaccination of the Ad-shVEGF in tumor-bearing mice, potent anti-angiogenesis was induced and resulted in tumor suppression and survival benefits.130 Recently, a study revealed that the amiRNAs based on the AdC68 vector could downregulate the survivin which was highly expressed in tumors, and the rAdC68 caused blockade of mitosis and cell cycle arrest at the G2/M phase. In the tumor-xenograft nude mice models, survivin-targeting amiRNAs expressed by rAdC68 effectively delayed growth of hepatic and cervical carcinomas.131

Ads have great potential as anti-cancer vectors. However, the clinical use of Ads is limited due to their limited infectivity in some cancer cells. Modifying the tropism of the Ads is an alternative way to generate better anti-cancer effect. In a recent study, the epidermal growth factor-like domain of the human heregulin-α (HRG) was inserted into the HI loop of AdHu5 fiber without adverse effect on the Ad growth or yields. The fiber-modified Ad virions showed enhanced infection of cells expressing the cognate receptors HER3/ErbB3 and HER4/
Capsid-incorporation of foreign antigen into ad virions as vaccine candidate

The most common method for Ad vaccine development is the expression of the foreign antigens in the E1 or E3 region of Ad vector as previously described in Table 1 and Table 2. However, pre-existing antibodies to the vectors may result in the failure of the vaccine. The “antigen capsid-incorporation” strategy has been developed to compensate for the drawbacks associated with the conventional antigen-expression system by the Ad vector to evade the pre-existing immunity. The Ad capsid proteins such as hexon, penton base, and pIX have variable sites for the antigen incorporation.13,14

Hexon is the most abundant structural protein on the capsid and has several hypervariable regions which can be modified to display the foreign antigens without affecting the Ad’s rescue and infectivity. For example, the AdC68 vectors were modified to express a linear B-cell epitope of the ectodomain of matrix 2 (M2e) of influenza virus within hypervariable regions 1 (HVR1) or HVR4 of the Ad hexon. Additional vectors with wild-type or M2e-modified hexon with influenza A virus NP as a transgene product in the E1-deleted region were also tested in the study. The vaccination regimen included priming with $10^{10}$ vp rAd and some mice boosted 2 months later. The pre-clinical study demonstrated that Ads expressing M2e within HVR1 of hexon induced higher magnitude and avidity of M2e-specific antibody responses than those carrying M2e within HVR4 or vectors expressing the M2e as part of a transgene product, and the M2e-specific antibody responses could be boosted by a second dose of the HVR1 hexon-modified vector but not by repeated immunization with the HVR4 hexon-modified vector.25 Besides influenza virus vaccine, other studies reported that the insertion of the neutralizing epitopes of HFMD virus into hexon could elicit neutralizing antibodies against HFMD virus lethal challenge in the mice models.22,137 As the HVRs of hexon contain the neutralizing epitopes of Ads,20 the hexon-modified Ads might change the immunogenicity compared to the wild type Ads, and the anti-sera from the hexon-modified-Ad vaccinated animals cannot well neutralize the wild type Ads, which provides a good platform for the prime-boost regimens for Ad based vaccines.22

Besides hexon, fiber can be engineered as an antigen-display system. Fiber modification makes Ads have specific tropism for cancer cells as above described, and can be incorporated with foreign antigens as vaccines against infectious diseases as well. A vector of AdHu5 expressing the 14-mer Pseudomonas aeruginosa immune-dominant outer membrane protein F (OprF) epitope 8 (Ep8) in five distinct sites of fiber was immunized in the mice. The results demonstrated that the FG-loop and HI-loop inserted sites were better than the other insertion sites in fiber since higher levels of protective immunity against P. aeruginosa were induced by FG-loop or HI-loop modified vectors.138 The penton base and pIX were tested for antigen-incorporation in some studies.139-142 However, compared with the other three proteins on the capsids, the penton base was rarely incorporated with foreign antigens perhaps due to the structural constraints.

Outlook and conclusions

Generally, Ad vectors are easy to be manipulated for genetic modification and capable of inducing potent antigen-specific immune responses. Most of the Ad species are rarely pathogenic to humans. Compared to the conventional vaccines, Ad vector-based vaccines can express a wide range of antigens from virus, bacteria or protozoan, and elicit long-term immune responses against infectious diseases. Despite the pre-existing neutralizing antibodies to the human Ads, the rare serotypes of Ads from different species have been developed to circumvent the disadvantages. All above advantages make Ads very attractive and potential vaccine candidates. Furthermore, Ad vectors show priority in anti-cancer research since they can be armed with therapeutic genes or modified to expand to higher titer in tumors than in the normal tissues. Many Ad vectors have been studied in animals against either infectious diseases or cancers, and revealed a good prospect of the further development.

Despite the incomplete success of Ad-based vaccines, Ad vectors still show great potential and are being extensively tested in the clinical trials recently. With more information obtained from Ad-related clinical trials, our understanding of the Ad vectors will be greatly enlarged, which will further promote the use of Ad vectors in the prevention and control of infectious diseases and cancer.

Abbreviations

| Abbreviation | Definition |
|--------------|------------|
| Ad           | adenovirus |
| CAR          | coxsackie and adenovirus receptor |
| AdHu         | human adenovirus |
| rAd          | recombinant Ad |
| DSG2         | desmoglein 2 |
| HIV          | human immunodeficiency virus; |
| SIV          | simian immunodeficiency virus |
| MERS         | Middle East respiratory syndrome coronavirus |
| HA           | hemagglutinin |
| NP           | nucleoprotein |
| HI           | hemagglutination-inhibiting |
| GP           | glycoprotein |
| amiRNA       | artificial microRNA |

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

Funding

This work was supported by grants from the “Knowledge Innovation Program” (No: Y014P31503) and the “100 Talent Program” (No: Y316P11503) from the Chinese Academy of Sciences and the Shanghai Pasteur Foundation.

References

[1] Takaifuji ET, Gaydos JC, Allen RG, Top FH, Jr. Simultaneous administration of live, enteric-coated adenovirus types 4, 7 and 21
vaccines: safety and immunogenicity. J Infect Dis 1979; 140:48-53; PMID:458200; http://dx.doi.org/10.1093/infdis/i40.1.48

[2] Rosenfeld MA, Siegfried W, Yoshimura K, Yoneyama K, Fukayama M, Stier LE, Pääkkö PK, Giliardi P, Stratford-Perricaudet LD, Perricaudet M. Adenovirus-mediated transfer of a recombinant α1-antitrypsin gene to the lung epithelium in vivo. Science 1991; 252:431-4; PMID:1907689; http://dx.doi.org/10.1126/science.1907689

[3] Khuri FR, Nemunaitis J, Gany JI, Arseneau J, Tannock IF, Romet L, Gore M, Ironside J, MacDougall RH, Heise C, et al. A controlled trial of intratumoral ONX-015, a selectively-replicating adenovirus, in combination with cisplatin and 5-fluorouracil in patients with recurrent head and neck cancer. Nat Med 2000; 6:879-85; PMID:10932224; http://dx.doi.org/10.1038/78638

[4] Bischoff JR, Kirn DH, Williams A, Heise C, Horn S, Muna M, Ng I, Nye JA, Sampson-Johannes A, Fattaey A, et al. An adenovirus mutant that replicates selectively in p53-deficient human tumor cells. Science 1996; 274:373-6; PMID:8832876; http://dx.doi.org/10.1126/science.274.3286.373

[5] Lasaro MO, Erlc HC. New insights on adenovirus as vaccine vectors. Mol Therapy 2009; 17:1333-9; http://dx.doi.org/10.1038/mtna.2009.130

[6] Tatsis N, Erlc HC. Adenoviruses as vaccine vectors. Mol Therapy 2004; 10:616-29; http://dx.doi.org/10.1038/jmt.2004.07.013

[7] Walsh MP, Seto J, Liu EB, Dehghan N, Hasan V, Lukashev AN. New insights on adenovirus as vaccine vector. J Infect Dis 1979; 140:48-53; PMID:458200; http://dx.doi.org/10.1093/infdis/i40.1.48

[8] Walsh MP, Seto J, Liu EB, Dehghan N, Hasan V, Lukashev AN. New insights on adenovirus as vaccine vector. J Infect Dis 1979; 140:48-53; PMID:458200; http://dx.doi.org/10.1093/infdis/i40.1.48

[9] Hendrix RM, Linde AE, Benton FR, Monteith SC, Tuchscherer MA, Gray GC, Gaydos JC. Large, persistent epidemic of adenovirus type 5 infections in healthy adults in China. Emerg Microbes Infect 2014; 3:e30; PMID:26038738; http://dx.doi.org/10.1038/jm.2014.30

[10] Wang X, Xing M, Zhang C, Yang Y, Chi Y, Tang X, Zhang H, Xiong S, Yu L, Zhou D. Neutralizing antibody responses to enterovirus and adenovirus in healthy adults in China. Emerg Microbes Infect 2014; 3:e30; PMID:26038738; http://dx.doi.org/10.1038/jm.2014.30

[11] Xia D, Henry LJ, Deisenhofer J. Crystal structure of the receptor-binding domain of adenovirus type 5 fiber protein at 1.7 Å resolution. Structure 1994; 2:1259-70; PMID:7704534; http://dx.doi.org/10.1016/S0969-2126(94)00126-X

[12] Anderson CW, Young ME, Flint SJ. Characterization of the adenovirus type 2 virion protein, m1. Virology 1989; 172:506-12; PMID:2800334; http://dx.doi.org/10.1016/0042-6822(89)90119-1

[13] Crawford-Miksza L, Schnurr DP. Analysis of 15 adenovirus hexon proteins reveals the location and structure of seven hypervariable regions containing serotype-specific residues. J Virol 1996; 70:1836-44; PMID:8627708

[14] Hendrix RM, Lindner JL, Benton FR, Monteith SC, Tuchscherer MA, Gray GC, Gaydos JC. Large, persistent epidemic of adenovirus type 5 infections in healthy adults in China. Emerg Microbes Infect 2014; 3:e30; PMID:26038738; http://dx.doi.org/10.1038/jm.2014.30

[15] Loskutoff DJ, Miescher P, Thompson JA, Beaudet AL. Adenovirus type 5 capsid. EMBO J 2005; 24:1645-54; PMID:15861131; http://dx.doi.org/10.1038/sj.emboj.7600653

[16] Wickham TJ, Cartron ME, Kovesi I. Targeting of adenovirus penton base to new receptors through replacement of its RGDF motif with other receptor-specific peptide motifs. Gene Therapy 1995; 2:750-6; PMID:8750015

[17] Hendrix RM, Linde AE, Benton FR, Monteith SC, Tuchscherer MA, Gray GC, Gaydos JC. Large, persistent epidemic of adenovirus type 5 infections in healthy adults in China. Emerg Microbes Infect 2014; 3:e30; PMID:26038738; http://dx.doi.org/10.1038/jm.2014.30

[18] Karen KA, Hearing P. Adenovirus core protein VII protects the viral genome from a DNA damage response at early times after infection. J Virol 2011; 85:4135-42; PMID:21345950; http://dx.doi.org/10.1128/JVI.02540-10

[19] Rosenfeld MA, Siegfried W, Yoshimura K, Yoneyama K, Fukayama M, Stier LE, Pääkkö PK, Giliardi P, Stratford-Perricaudet LD, Perricaudet M. Adenovirus-mediated transfer of a recombinant α1-antitrypsin gene to the lung epithelium in vivo. Science 1991; 252:431-4; PMID:1907689; http://dx.doi.org/10.1126/science.1907689
Roelvink PW, Lizonova A, Lee JG, Li Y, Bergelson JM, Finberg RW, Brough DE, Kovesdi I, Wickham TJ. The coxsackievirus-adenovirus receptor protein can function as a cellular attachment protein for adenovirus serotypes from subgroups A, C, D, E, and F. J Virol 1998; 72:7909-15; PMID:9733828

Sachs MD, Rauen KA, Ramamurthy M, Dodson JL, De Marzo AM, Puttali MJ, Schonthal MP, Rodriguez R. Integrin α(+) and coxsackie adenovirus receptor expression in clinical bladder cancer. Urology 2002; 60:531-6; PMID:12350512; http://dx.doi.org/10.1006/s090-4295(02)01748-X

Fux J, Liu L, Malin S, Philipson LV, Peetersson RF. Expression of the coxsackie and adenovirus receptor in human astrocytic tumors and xenografts. Int J Cancer Int Du Cancer 2003; 103:723-9; http://dx.doi.org/10.1002/ijc.10891

Nalbantoglu J, Pari G, Karpati G, Holland PC. Expression of the primary coxsackie and adenovirus receptor is downregulated during skeletal muscle maturation and limits the efficacy of adenovirus-mediated gene delivery to muscle cells. Hum Gene Therapy 1999; 10:1009-19; http://dx.doi.org/10.1089/1043039950841409

Garnett CT, Talekar G, Mahr JA, Huang W, Zhang Y, Ornelles DA, Gooding LR. Latent species C adenoviruses in human tonsil tissues. J Virol 2009; 83:2417-28; PMID:19109894; http://dx.doi.org/10.1128/JVI.02392-08

Smith TA, Idamakanti N, Marshall-Neff J, Rollence ML, Wright P, Kaloss M, King L, Mech C, Dinges L, Iverson WV, et al. Receptor interactions involved in adenoviral-mediated gene delivery after systemic administration in non-human primates. Hum Gene Therapy 2003; 14:1595-604; PMID:14633402; http://dx.doi.org/10.1089/10430340322542248

Gaggar A, Shayakhmetov DM, Lieber A. CD46 is a cellular receptor for group B adenoviruses. Nat Med 2003; 9:408-12; PMID:14566335; http://dx.doi.org/10.1038/nm952

Wang H, Li ZY, Liu Y, Persson J, Beyer I, Moller T, Koyuncu D, Drescher MR, Strauss R, Zhang XB, et al. Desmoglein 2 is a receptor for adenovirus serotypes 3, 7, 11 and 14. Nat Med 2011; 17:96-104; PMID:21151137; http://dx.doi.org/10.1038/nm.2270

Nilsson EC, Storm BJ, Bauer J, Johansson SM, Lookene A, Angstrom J, Hedenstrom M, Eriksson TL, Frangsmyr L, Rinadli S, et al. The G1α1a glycan is a cellular receptor for adenoviruses causing epidemic keratoconjunctivitis. Nat Med 2011; 17:105-9; PMID:21151139; http://dx.doi.org/10.1038/nm.2267

Huang S, Kamata T, Takada Y, Ruggeri ZM, Nemerow GR. Adenovirus interaction with distinct integrins mediates separate events in cell entry and gene delivery to hematopoietic cells. J Virol 1996; 70:4502-8; PMID:8676475

Roelvink PW, Kovesdi I, Wickham TJ. Comparative analysis of adenovirus fiber-cell interaction: adenovirus type 2 (Ad2) and Ad9 utilization of the same cellular fiber receptor but use different binding strategies for attachment. J Virol 1996; 70:7614-21; PMID:8929881

Arnberg N. Adenovirus receptors: implications for targeting of viral vectors. Trends Pharmacological Sci 2012; 33:442-8; http://dx.doi.org/10.1016/j.tips.2012.04.005

Davidson E, Diaz RM, Hart JR, Santis G, Marshall JE. Integrin alpha5beta1-mediated adenovirus infection is enhanced by the integrin-activating antibody TS2/16. J Virol 1996; 71:6204-7; PMID:9223518

Davidson E, Kirby I, Whitehouse J, Hart I, Marshall JF, Santis G. Adenovirus type 5 uptake by lung adenocarcinoma cells in culture correlates with Ad5 fibre binding is mediated by α(v)β3 integrin and can be modulated by changes in β1 integrin function. J Gene Med 2001; 3:550-9; PMID:11778901; http://dx.doi.org/10.1002/jgm.223

Colin M, Mailly L, Rogge S, D’Halluin JC. Efficient species C HAAdV infectivity on placental cell lines using a clathrin-dependent lipid raft/caveola endocytic route. Mol Therapy 2005; 11:224-36; http://dx.doi.org/10.1038/j.ymthe.2004.10.007

Choi JW, Lee JS, Kim SW, Yun CO. Evolution of oncolytic adenoviruses for cancer treatment. Adv Drug Delivery Rev 2012; 64:720-9; http://dx.doi.org/10.1016/j.addr.2012.11.011
Prevec L, Campbell JB, Christie BS, Belbeck L, Graham FL. A recombinant human adenovirus vaccine against rabies. J Infect Dis 1990; 161:27-30; PMID:2295855; http://dx.doi.org/10.1093/infdis/161.1.27

[64] Voy A, Neubert A, Tommereng E, Muller T, Dohner L, Neubert L, Hughes K. Immunogenicity of an E1-deleted recombinant human adenovirus against rabies by different routes of administration. J General Virol 2001; 82:2191-7; http://dx.doi.org/10.1099/0022-1317-82-9-2191

[65] Bouet-Cararo C, Contreras V, Fournier A, Jallet C, Guibert JM, Prevec L, Campbell JB, Christie BS, Belbeck L, Graham FL. A recombinant adenovirus vaccine against rabies by different routes of administration. J Infect Dis 2015; 212 Suppl 2:S359-83; PMID:25957963; http://dx.doi.org/10.1093/infdis/jiv102

[66] Tapia MD, Sow SO, Lyke KE, Haidara FC, Dallal F, Dombia M, et al. Use of ChAd3-EOB-Z. Ebola virus vaccine in Malian and US adults, and boosting of Malian adults with MVA-BN-Filo: a phase 1, single-blind, randomised controlled trial, a phase 1b, open-label and double-blind, dose-escalation trial, and a nested, randomised, double-blind, placebo-controlled trial. Lancet Infect Dis 2015; 16:31-42; PMID:26546548; http://dx.doi.org/10.1016/S1473-3099(15)00362-X

[67] Choi JH, Schaefer SC, Zhang L, Kobinger GP, Jueelich T, Freiberg AN, Croyle MA. A single sublingual dose of an adenovirus-based vaccine protects against lethal Ebola challenge in mice and guinea pigs. Mol Microbiol 2013; 9:156-67; http://dx.doi.org/10.1111/j.1365-2958.2013.13586.x

[68] Li W, Li M, Deng G, Zhao L, Liu X, Wang Y. Prime-booster vaccination with Bacillus Calmette Guerin and a recombinant adenovirus co-expressing CFP10, ESAT6, Ag85A and Ag85B of Mycobacterium tuberculosis induces robust antigen-specific immune responses in mice. Mol Med Reports 2015; 12:5088-98; PMID:22926955; http://dx.doi.org/10.3892/mmr.2015.4512

[69] Smill F, Jeyanathan M, Smieja M, Medina MF, Thanthri-Don N, Zganiacz A, Yin C, Heriazen A, Damjanovic D, Puri L, et al. A human type 5 adenovirus-based tuberculosis vaccine induces robust T cell responses in humans despite preexisting anti-adenovirus immunity. Sci Translational Med 2013; 5:205ra134; http://dx.doi.org/10.1126/scitranslmed.3006843

[70] Chuang I, Sedegah M, Maiolatesi S, Fedders C, Reyes S, Reyes A, Vaquez C, Alcorta Y, Chuang I, Spring M, et al. Human adenovirus 5-vectored Plasmodium falciparum NMRC-M3V-Ad-PfCA vaccine expressing CSP and AMA1 is safe, well-tolerated and immunogenic but does not protect against controlled human malaria infection. Hum Vaccines Immunotherapeutics 2013; 9:2165-77; http://dx.doi.org/10.4161/hv.24941

[71] Schwenk R, Banaria G, Epstein J, Kim Y, Peters B, Belmonte M, Ganesan H, Huang J, Reyes S, Stryhn A, et al. Ex vivo tetramer staining and cell surface phenotyping for early activation markers CD38 and HLA-DR to enumerate and characterize malaria antigen-specific CD8+ T-cells induced in human volunteers immunized with a Plasmodium falciparum adenovirus-vectored malaria vaccine expressing AMA1. Malaria J 2013; 12:376; http://dx.doi.org/10.1186/1475-2875-12-376

[72] Chuang I, Sedegah M, Cicatelli S, Spring M, Polhemus M, Tamminga C, Patel A, Soule G, Ennis J, Javaher A, Ketter G, et al. A replication-competent adenovirus capsid display recombinant elicits antibodies against Plasmodium falciparum sporozoites in Aotus nancymaae monkeys. Infect Immun 2015; 83:268-75; http://dx.doi.org/10.1128/IAI.02626-14
Perreau M, Pantaleo G, Kremer EJ. Activation of a dendritic cell-T
Koup RA, Roederer M, Lamoreaux L, Fischer J, Novik L, Nason
MSP1 and AMA1: assessment of efficacy against mosquito bite
challenge in humans. Mol Therapy 2012; 20:2355-68; http://dx.doi.
or/10.1038/mn.2012.223

Ginberg HS. The life and times of adenoviruses. Adv Virus Res
1999; 54:1-13; PMID:10547672; http://dx.doi.org/10.1016/S0065-
3527(08)60363-2

Kochanek S. High-capacity adenoviral vectors for gene transfer and
somatic gene therapy. Hum Gene Therapy 1999; 10:2451-9; http://
dx.doi.org/10.1089/10430349950016807

Yeh P, Dedieu JF, Orsini C, Vigne E, Denelle P, Perricaudet M. Effi-
cient dual transcomplementation of adenovirus E1 and E4 regions
from a 293-derived cell line expressing a minimal E4 functional
unit. J Virol 1996; 70:559-65; PMID:8523570

Toietta G, Mane VP, Norona WS, Finegold MJ, Ng P, McDonagh
AF, Beaudet AL, Lee B. Lifelong elimination of hyperbilirubinemia
in the Gunn rat with a single injection of helper-dependent aden-
viral vector. Proc Natl Acad Sci U S A 2005; 102:3930-5; PMID:
15753322; http://dx.doi.org/10.1073/pnas.0500930102

Gilbert R, Dudley RW, Liu AB, Petroff BJ, Nalbantoglu J, Karpati G.
Prolonged dystrophin expression and functional correction of mdx
mouse muscle following gene transfer with a helper-dependent (gutted)
adeno virus-encoding murine dystrophin. Hum Mol Genet-
ics 2003; 12:1287-99; http://dx.doi.org/10.1093/hmg/ddg141

Dicks MD, Guzman E, Spencer AJ, Gilbert SC, Charleston B, Hill
AV, Cottingham MG. The relative magnitude of transgene-specific
adaptive immune responses induced by human and chimpanzee
adenovirus vectors differs between laboratory animals and a target
species. Vaccine 2015; 33:1121-8; PMID:25629523; http://dx.doi.
or/10.1016/j.vaccine.2015.01.042

Tan WG, Jin HT, West EE, Penaloza-MacMaster P, Wieland A, Zill-
xioi MJ, McElrath MJ, Barouch DH, Ahmed R. Comparative analy-
sis of simian immunodeficiency virus gag-specific effector and
memory CD8+ T cells induced by different adenovirus vectors. J
Virol 2013; 87:1359-72; PMID:23175355; http://dx.doi.org/10.1128/
JV1.2012055-12

Teigler JE, Lampierto MJ, Barouch DH. Vaccination with adenovirus
serotypes 35, 26, and 46 elicits higher levels of innate cytokine responses
than adenovirus serotype 5 in rhesus monkeys. J Virol 2012; 86:9590-8;
PMID:22787208; http://dx.doi.org/10.1128/JVI.00740-12

Xiang ZQ, Gao GP, Reyes-Sandoval A, Li Y, Wilson JM, Ertl HC. Oral
vaccination of mice with adenovirus vectors is not impaired by preexist-
ing immunity to the vaccine carrier. J Virol 2003; 77:10780-9;
PMID:14512528; http://dx.doi.org/10.1128/JVI.77.20.10780-10789.2003

Liebowitz D, Lindblom JD, Brandl JR, Garg SJ, Tucker SN. High titre
neutralising antibodies to influenza after oral tablet immunisation:
a phase 1, randomised, placebo-controlled trial. Lancet Infect Dis
2015; 15:1041-8; PMID:26333337; http://dx.doi.org/10.1016/S1473-
3099(15)00266-2

Richert L, Lhomme E, Fagard C, Levy Y, Chene G, Thiebaut R.
Recent developments in clinical trial designs for HIV vaccine
development. PloS One 2013; 8; http://dx.doi.org/10.1371/jour-
nal.pone.0062496

Webby RJ, Weaver EA. Centralized Consensus Hemagglutinin
Genes Induce Protective Immunity against H1, H3 and H5 Influen-
za Viruses. PloS One 2015; 10:e0140702; PMID:26469190; http://dx.
doi.org/10.1371/journal.pone.0140702

Khrura S, Coyle EM, Manischewitz J, King LR, Ishioka G, Alexan-
der J, Smith J, Gurwith M, Golding H. Oral priming with replicating
adenovirus serotype 4 followed by subunit H5N1 vaccine boost pro-
 motes antibody affinity maturation and expands H5N1 cross-clade
neutralization. PloS One 2015; 10:e0115476; PMID:25629161;
http://dx.doi.org/10.1371/journal.pone.0115476

Zhang H, Tang X, Zhu C, Song Y, Yin J, Xu J, Ertl HC, Zhou D.
Adenovirus-mediated Artificial MicroRNAs targeting matrix or
nucleoprotein genes protect mice against lethal influenza virus chal-
lenge. Gene therapy 2015; 22:653-62; PMID:25835311; http://dx.
doi.org/10.1038/gt.2015.31

Feldmann H, Jones S, Klenk HD, Schnittler HJ. Ebola virus: from
discovery to vaccine. Nat Rev Immunol 2003; 3:677-85;
PMID:12974482; http://dx.doi.org/10.1038/nri1154

Feldmann H, Geisbert TW. Ebola haemorrhagic fever. Lancet 2011;
377:849-62; PMID:21084112; http://dx.doi.org/10.1016/S0140-6736
(10)60667-8

Sacks JA, Zehe E, Redick C, Bah A, Cowger K, Camara M, Dia llo A,
Gigo AN, Dhillon RS, Liu A. Introduction of Mobile Health Tools
to Support Ebola Surveillance and Contact Tracing in Guinea.
Global Health Sci Pract 2015; 3:646-59; http://dx.doi.org/10.107-
45/GHSP-D-15-00207

Sullivan NJ, Sanchez A, Rollin PE, Yang ZY, Nabel GJ. Develop-
ment of a preventive vaccine for Ebola virus infection in primates.
Nature 2000; 408:605-9; PMID:11117750; http://dx.doi.org/10.1038/35046108
C. ZHANG AND D. ZHOU

[117] Sullivan NJ, Geisbert TW, Geisbert JB, Xu L, Yang ZY, Roederer M, Koup RA, Jahrling PB, Nabel GJ. Accelerated vaccination for Ebola virus haemorrhagic fever in non-human primates. Nature 2003; 424:681-4; PMID:12904795; http://dx.doi.org/10.1038/nature01876

[118] Ledgerwood JE, DeZure AD, Stanley DA, Novik L, Enama ME, Berkowitz NM, Hu Z, Joshi G, Ploquin A, Sitar S, et al. Chimpanzee Adenovirus Vector Ebola Vaccine - Preliminary Report. N Eng J Med 2014; 374:1635-46.

[119] Rampleting T, Ewer K, Bowyer G, Wright D, Imoukhuede EB, Payne R, Hartnell F, Gibani M, Bliss C, Minhinnick A, et al. A Monovalent Chimpanzee Adenovirus Ebola Vaccine - Preliminary Report. N Eng J Med 2016; 374:1635-46.

[120] Wolf JK, Bodurka DC, Gano JB, Deavers M, Ramondetta L, Ramirez PT, Levenback C, Gershenson DM. A phase 1 study of Ad5p3 (INGN 201; ADVEXIN) for patients with platinum- and paclitaxel-resistant epithelial ovarian cancer. Gynecologic Oncol 2004; 94:442-8; http://dx.doi.org/10.1016/j.ygyno.2004.05.041

[121] Wu J, Zhu Y, Xu C, Xu H, Zhou X, Yang J, Xie Y, Tao M. Adenovirus-mediated p53 and ING4 gene cotransfer elicits synergistic antitumor effects through enhancement of p53 acetylation in breast cancer. Oncol Reports 2016; 35:243-52.

[122] Nemunaitis J. Head and neck cancer: response to p53-based therapy. Hum Gene Therapy 2005; 16:1133-40; http://dx.doi.org/10.1089/hum.2004.15.1167

[123] Rux JJ, Burnett RM. Adenovirus structure. Hum Gene Therapy 2004; 15:1167-76; http://dx.doi.org/10.1089/hum.2004.15.1167

[124] Nemerow GR, Pache L, Reddy V, Stewart PL. Insights into adenovirus immunity. J Virol 2006; 80:5523-30; PMID:16699033; http://dx.doi.org/10.1128/JVI.02667-05

[125] Westphal M, Yla-Herttuala S, Martin J, Warnke P, Menei P, Eckerd R, Kinley J, Kay R, Ram Z. Adenovirus-mediated gene therapy with sitimagene ceradenovec followed by intravenous ganciclovir for patients with operable high-grade glioma (ASPECT): a randomized, placebo-controlled, phase 3 trial. Lancet Oncol 2013; 14:823-33; PMID:23996201; http://dx.doi.org/10.1016/1535-7163.CCTR-09-1044

[126] Xie J, Guo X, Liu F, Luo J, Duan F, Tao X. In vitro antitumor immune response induced by dendritic cells transduced with human livin within adenovirus α recombinant adenovirus. Cell Immunol 2015; 297:46-52; PMID:26100880; http://dx.doi.org/10.1016/j.ccellimm.2015.06.003

[127] Liikanen I, Ahtiainen L, Hirvinen ML, Bramante S, Cerullo V, Nemunaitis J, Clayman G, Sitar S, et al. Oncolytic adenovirus with temozolomide induces autophagy and oncolytic adenovirus immune responses in cancer patients. Mol Therapy 2013; 21:1212-23; http://dx.doi.org/10.1038/m.2013.51

[128] Arioishi Y, Watanabe M, Iikawa S, Yamazaki C, Sakahira T, Hiran T, Araki M, Ebara S, Nasu Y, Udono H, et al. The induction of antigen-specific CTL by in situ Ad-REIC gene therapy. Gene Therapy 2016; PMID:26836118; http://dx.doi.org/10.1038/2016.7

[129] Yao C, Sasaki HM, Ueda T, Tomari Y, Tadakuma H. Single-Molecule Analysis of the Target Cleavage Reaction by the Drosophila RNAi Enzyme Complex. Mol Cell 2015; 59:125-32; PMID:26140368; http://dx.doi.org/10.1016/j.molcel.2015.05.015

[130] Yoo JY, Kim JH, Kwon YG, Kim EC, Kim NK, Choi HJ, Yun CO. VEGF-specific short hairpin RNA-expressing oncolytic adenovirus elicits potent inhibition of angiogenesis and tumor growth. Mol Therapy 2007; 15:295-302; http://dx.doi.org/10.1038/sj.mt.6300023

[131] Chi Y, Wang X, Yang Y, Zhang C, Erli HC, Zhou D. Survivin-targeting Artificial MicroRNAs Mediated by Adenovirus Suppress Tumor Activity in Cancer Cells and Xenograft Models. Mol Therapy Nucleic Acids 2014; 3:e208; PMID:25368912; http://dx.doi.org/10.1038/mt.2014.59

[132] MacLeod SH, Elgadi MM, Bossi G, Sankar U, Pisio A, Agopsowicz K, Sharon D, Graham FL, Hitt MM. HER3 targeting of adenovirus by fiber modification increases infection of breast cancer cells in vitro, but not following intratumoral injection in mice. Cancer Gene Therapy 2012; 19:888-98; PMID:22099884; http://dx.doi.org/10.1038/cgt.2012.79

[133] Zhan Y, Yu B, Wang Z, Zhang Y, Zhang HH, Wu H, Feng X, Geng RS, Kong W, Yu XH. A fiber-modified adenovirus co-expressing HSV-TK and Coli.NTR enhances antitumor activities in breast cancer cells. Int J Clin Exp Pathol 2014; 7:2850-60; PMID:25031704

[134] Takagi-Kimura M, Yamano T, Tamamoto A, Okamura N, Oka-mura H, Hashimoto-Tamaoki T, Tagawa M, Kasahara N, Kubo S. Enhanced antitumor efficacy of fiber-modified, midkine promoter-regulated oncolytic adenovirus in human malignant mesothelioma. Cancer Sci 2013; 104:1433-9; PMID:23962292; http://dx.doi.org/10.1111/cas.12267

[135] Rux JJ, Burnett RM. Adenovirus structure. Hum Gene Therapy 2004; 15:1167-76; http://dx.doi.org/10.1089/hum.2004.15.1167

[136] Nemerow GR, Pache L, Reddy V, Stewart PL. Insights into adenovirus host cell interactions from structural studies. Virology 2009; 384:380-8; PMID:19019405; http://dx.doi.org/10.1016/j.virol.2008.10.016

[137] Tian X, Su X, Li X, Li H, Li T, Zhou Z, Zhong T, Zhou R. Protection against enterovirus 71 with neutralizing epitope incorporation within adenovirus type 3 hexon. Plos One 2012; 7:e41381; PMID:22848478; http://dx.doi.org/10.1371/journal.pone.0041381

[138] Sharma A, Krause A, Xu Y, Sung B, Wu W, Worqall S. Adenovirus-based vaccine with epitopes incorporated in novel fiber sites to induce protective immunity against Pseudomonas aeruginosa. Plos One 2013; 8:e56996; PMID:23437292; http://dx.doi.org/10.1371/journal.pone.0056996

[139] Krause A, Joh JH, Hackett NR, Roelvink PW, Bruder JT, Wickham TJ, Kovesdi I, Crystal RG, Worqall S. Epitopes expressed in different adenovirus capsid proteins induce different levels of epitope-specific immunity. J Virol 2006; 80:5523-30; PMID:16699033; http://dx.doi.org/10.1128/JVI.02667-05

[140] Vigne E, Mahfouz I, Dedieu JF, Brie A, Perricaudet M, Yeh P. RGD inclusion in the hexon monomer provides adenovirus type 5-based vectors with a fiber knob-independent pathway for infection. J Virol 1999; 73:5156-61; PMID:10233980

[141] Dmitriev IP, Kashentseva EA, Curiel DT. Engineering of adenovirus vectors containing heterologous peptide sequences in the C terminus of capsid protein IX. J Virol 2002; 76:6893-9; PMID:12072490; http://dx.doi.org/10.1128/JVI.02667-05

[142] Parks RJ. Adenovirus protein IX: a new look at an old protein. Mol Ther 2005; 11:19-25; http://dx.doi.org/10.1038/mt.2005.09.018