Evaluation of Pre-Clinical Efficacy to HPV16 L2E6E7 Vaccine and HPV16 E6E7 Adenovirus-5 Vector Vaccine with Different Dosages and Prime-Booster Regimens in Mouse Model

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Abstract

Purpose: This study evaluates the dose-response and immunization procedure in mouse model and the efficacy of prime-booster regimens with HPV16 L2E6E7 vaccine and HPV16 E6E7 Ad5 vector vaccine. Methods: Experimental animals were C57 BL/6 mice. Each group included 10 or 20 C57 BL/6 mice. The tumor model used TC-1 tumor cells. The HPV16 L2E6E7 vaccine groups were treated using the following dosage: 15 μg/ml, 30 μg/ml, 60 μg/ml, 120 μg/ml, 240 μg/ml, and then 120 μg/ml was used for the following regimens: 0-7 days, 0-15 days, 0-7 days, 0-15 days. The HPV16 E6E7 Adenovirus-5 vector vaccine groups were treated using the following dosage: 3.00×107 IU/ml, 3.00×108 IU/ml, 3.00×109 IU/ml, and then 3.00×108 IU/ml was used for the following regimens: 0-7 days, 0-15 days, 0-7-15 days, and control group. Prime-booster combined regimens with HPV16 L2E6E7 vaccine (P, 120 μg/ml) and HPV16 E6E7 ad5 vector vaccine (V, 3.00×107 IU/ml) were set as follows: 0P-7P days, 0P-7V days, and 0P-7P-15V days, 0P-7V-15V days, and 0P-7P-15V-21V days. Results: Upon challenge with 104 TC-1 tumor cells, mice developed palpable, rapidly growing tumors within 7–14 days. These tumors became lethal to the mice within 21–28 days. HPV16 L2E6E7 vaccine (120 μg/ml, 0-7-15 day’s procedure) protective efficacy was 85% and the HPV16 E6E7 Ad5 vector vaccine (3.00×107 IU/ml, 0 day procedure) that was 80%. Prime-booster regimens showed a protective efficacy of 80–90% for the 0P-7V days and 0P-7V-15V day’s schedules. Conclusion: HPV16 L2E6E7 vaccine and HPV16 E6E7 Ad5 vector vaccine are proved the candidate vaccine for therapeutic intervention against HPV16-induced tumor.

Keywords: Human papillomavirus-16; Therapeutic vaccine; Pre-clinical; Efficacy

Introduction

Vulval intraepithelial neoplasia (VIN) and cervical cancer are commonly associated with human papillomavirus (HPV) type 16 infection [1,2]. This discovery has provided the opportunity for an immunotherapeutic approach to the treatment of these conditions. HPV infection by one or more “high-risk” or “oncogenic” types, most commonly 16 and 18, are present in 90-100% of invasive and grade 3 intraepithelial cervical lesions and in the majority of noncervical anogenital neoplasias. HPV oncoproteins E6 and E7 play a crucial role in lower genital tract carcinogenesis and are required for maintenance of the transformed state and malignant phenotype [3,4]. This makes the HPV16 E6 and E7 oncoproteins attractive targets for immunotherapy because they are exclusively expressed in virally infected and neoplastic cells, never in normal or healthy cells.

An alternative approach is to induce systemic cell-mediated immune responses with the aim of effecting local cytotoxicity against the virus infected cells. Nonreplicating recombinant adenoviruses have been shown to induce cell-mediated and humoral immune responses and, in some cases, protection against a variety of pathogens [5,6]. A recombinant HPV 16-L2E6E7 fusion protein (HPV16 P-vaccine) and adenovirus 5 vector delivered HPV E6E7 protein (HPV16 Ad5-vaccine) were built using pilot plant technology. In order to develop the therapeutic vaccine, we need to evaluate the dose-response and immunization procedure in mouse model and the efficacy of prime-booster regimens. Only then can we seek permission from the China’s FDA to perform clinical trials.

Methodology

Antigens and vaccine formulations: HPV16 P-vaccine, a recombinant HPV 16-L2E6E7 fusion protein, was isolated from solubilized E. coli inclusion bodies under reducing conditions and purified by chromatography. The 80 kD L2E6E7 monomer comprises 725 amino acids, as predicted by the nucleotide sequence of the L2E6E7 gene. The protein was formulated using 5 mM glycine buffer (pH 8.0) containing 0.9 mM cysteine and finally freeze-dried and routinely stored at 2-8°C until use. Prior to administration of the vaccine protein, 0.5 ml sterile water was added for injection. The final vaccine consisted of a white homogeneous liquid.

HPV16 Ad5-vaccine, adenovirus 5-vector-delivered HPV E6E7 protein, consisted of the fused E6 and E7 open reading frames of HPV16, each under the control of an adenovirus promoter. The fused E6 and E7 genes had previously been found to show no transforming activity. The virus was prepared at a concentration of 1×109 infective units (IU)/ml. HPV16 Ad5-vaccine is routinely stored at -70°C until use.

Experimental animal and tumor model establishment: C57 BL/6
mice (female, 6-8 weeks, 20-25 g) were obtained from Shanghai SLAC Laboratory Animal Center and held under specific pathogen-free conditions. All experiments in C57 BL/6 mice were performed in compliance with "Experimental Animal Management Regulations in Zhejiang Province, China".

TC-1 tumor cells derived from primary epithelial cells of C57 BL/6 mice were co-transformed with HPV16 E6 and E7 and c-Ha-ras oncogenes. These cells were cultured in IMDM +10% FCS.

C57 BL/6 mice were injected subcutaneously with 1×10^6 TC-1 cells in the left leg, and were vaccinated with HPV 16 P-vaccine next day (recorded as day 0, 24 hours later after injection TC-1 cells) according to the dosage strategies selected for that mouse’s treatment group. Control group mice were injected only with the freeze-dried formulizer.

HPV-specific cellular immunity and tumor development were monitored from day 7 through day 40, until the tumor growth remained stable through three consecutive observations. At that point mice were killed.

Different groups of mice were immunized at different dosages and on different regimens: HPV16 p-vaccine for dose-response groups was set as follows: 15 μg/ml, 30 μg/ml, 60 μg/ml, 120 μg/ml, 240 μg/ml, and a control group (amino-salt formulizer). Based on the dose-responses results, different immunization procedures were established: 120 μg/ml on days 0 and 7 (0-7), on days 0 and 15 (0-15), on days 7 and 15 (7-15), and on days 0, 7, and 15 (0-7-15), and a control group (also at 0-7-15). Each group included 10 or 20 C57 BL/6 mice.

HPV16 Ad5-vaccine for dose-response groups was set as follows: 3.00×10^6 IU/ml, 3.00×10^7 IU/ml, 3.00×10^8 IU/ml, and 3.00×10^9 IU/ml, with control group (adv5+PBS). Based on the dose-response results, different immunization procedures were established: 3.00×10^7 IU/ml at 0-7, 0-15, and 0-7-15, and a control group (Ad5, 0-7-15).

Combined prime-booster regimens with HPV p-vaccine (marked P) and HPV ad5-vaccine (marked V) were set as follows: 0P-7V, 0P-7P-15V, 0P-7V-15V, and 0P-7P-15V-21V with 120 μg/ml HPV p-vaccine and 3.00×10^7 IU/ml HPV ad5-vaccine.

Observation index: onset of the tumor, size of the tumor; tumor outgrowth (%) in vaccine group.

Results

1. HPV p-vaccine. Upon challenge with 10^5 TC-1 cells, mice developed palpable, rapidly growing tumors within 7-14 days. These became lethal to the mice within 21-28 days. As such, newly challenged mice were regarded as a proper model for immune-intervention against minimal residual disease. Mice challenged with 10^5 TC-1 cells were vaccinated 24 hours after challenge (on day 0) and monitored for the development of the tumors. All control mice quickly developed tumors. Therapeutic vaccination with 120 μg/ml HPV p-vaccine on a 0-7-15 schedule protected the majority of mice against tumor outgrowth (19/20). The onset of tumor growth in one tumor-positive mouse was delayed (21 days after challenge). The protective effect of 15 μg/ml and 30 μg/ml was considerable weaker. Dosages of HPV16 p-vaccine at 60 μg/ml, 120 μg/ml, and 240 μg/ml showed an efficacy of 40%, 60%, and 60% respectively. The most effective immunization regimen was the 0-7-15 schedule at a dosage of 120 μg/ml (Figures 1 and 2). Taken together, these results show that HPV p-vaccine administered at a dose of 120 μg/ml on a 0-7-15 schedule is highly effective 85% in therapeutic settings.

2. HPV16 Ad5-vaccine. All control mice quickly developed tumors within 7-14 days. The onset of tumor growth in the HPV ad5-vaccine group was delayed for 14 days after challenge (Figure 3). The dose of 3.00×10^6 IU/ml showed 80% efficacy whereas both 3.00×10^7 IU/ml and 3.00×10^8 IU/ml showed 100% efficacy. Single doses of 3.00×10^6 IU/ml or more were so highly effective, there was no significant difference between the different immunization schedules. Therapeutic vaccination with HPV ad5-vaccine at single doses of 3.00×10^6 IU/ml or more was found to protect the majority of mice from tumor outgrowth.

![Figure 1](https://example.com/fig1.png)

**Figure 1:** Each group consists of 10 C57 BL/6 mice. HPV-p vaccine with different dosage groups was designed as 15 μg/ml, 30 μg/ml, 60 μg/ml, 120 μg/ml and 240 μg/ml; control group was injected with amino-salt formulizer.

![Figure 2](https://example.com/fig2.png)

**Figure 2:** Each group consists of 20 C57 BL/6 mice. HPV-p vaccine in different immunization procedure was designed as 0-7 days, 0-15 days, 0-7-15 days injection the vaccine with dosage of 60 μg/ml and 120 μg/ml respectively; control group was injected with amino-salt formulizer.

![Figure 3](https://example.com/fig3.png)

**Figure 3:** Each group consists of 10 C57 BL/6 mice. HPV ad5-vaccine in different dosage were designed as 3×10^6 IU/ml, 3×10^7 IU/ml, 3×10^8 IU/ml, 3×10^9 IU/ml, 3×10^10 IU/ml and 3×10^11 IU/ml; Control group was injected with adv5+PBS.
3. Monitoring of vaccine induced immunoresponses. The number of T-cells per 300,000 splenocytes that spontaneously appeared following stimulation with E749–57 peptide produced IFN-γ upon stimulation was not found to be significant. No such cells were detected by ELISPOT, but ELISA showed HPV16-specific IgG antibodies to be present at a high titer in the sera of the mice (Figure 4). In the different regimens, the results showed that only HPV ad5-vaccine failed to significantly increase IgG levels. The HPV p-vaccine was significantly associated with increases in IgG. The combined regimens with p-vaccine boosted with p-vaccine or with ad5-vaccine showed highly increased IgG levels.

4. Prime-boost regimens with HPV p-vaccine and HPV ad5-vaccine. In control groups, the greatest frequency of tumor onset was observed from day 7 through day 14. Our results showed that injection of HPV ad5-vaccine at day 7 was more efficacious at suppressing tumor outgrowth than injections of HPV p-vaccine. Heterologous prime-boost immunization regimens, in which two different types of vaccine sharing the antigen of choice are used, have been proven more effective in stimulating the T-cell responses than homologous immunization regimens. Our experiment demonstrated that heterologous regimen works as well or better when HPV ad5-vaccine is administered in a single dose of $3.00 \times 10^7$ IU/ml. Regimens 0P-7V and 0P-7V-15V showed strong protective efficacy against tumor outgrowth, with an efficacy of 80–90% (Figure 5).

**Discussion**

Over the past decade, substantial progress has been made in the understanding of host immune responses to HPV infection and in the development of candidate vaccines [7]. The rationale behind immunotherapy for intraepithelial neoplasia is based upon the expression of HPV E6 and E7 proteins, which distinguish neoplastic from normal epithelia, thereby acting as tumor-specific antigens [8]. Animal model experiments have demonstrated that the induction of E6- and E7-specific T cells can effectively control established tumors, and several candidate vaccines are being tested in early-phase clinical trials [9,10].

We here evaluated therapeutic HPV p-vaccine, and HPV ad5-vaccine in a pre-clinical mouse model. Our data show that the majority of mice were protected against tumor outgrowth when HPV p-vaccine and HPV ad5-vaccine were therapeutically administered in a setting of minimal residual disease. For the HPV p-vaccine, INF-γ was not significantly induced, but the levels of the specific antibody IgG were found to be highly increased after vaccination. This differs from the findings of some other research teams [11].

The viral vector in therapeutic HPV vaccine as reported was a vaccinia virus-based vaccine TA-HPV, which has been tested in cervical cancer patients [12,13]. In our research, we used the adenovirus-5 vector carrying the HPV 16 E6/E7 fusion proteins. The results indicated that HPV ad5-vaccine at a dose of $3.00 \times 10^7$ IU/ml or more, even when only a single dose was administered protected 80% of the mice from tumor outgrowth. This suggests that it is possible for viral vector vaccines to use vaccinia virus and adenovirus-5 as vectors. Both of these showed the same efficacy in the mouse model.

The HPV ad5-vaccine, developed through pre-clinical mouse studies, was found to be capable of inducing substantial numbers of specific CTL [14,15]. In this study we did not find any significantly increased levels of INF-γ of CTL. In the prime-booster regimens HPV p-vaccine combined with HPV ad5-vaccine demonstrated greater efficacy than all other single or homologous strategies.

The prime-boost regimens with HPV p-vaccine and HPV ad5-vaccine showed the greatest ability to protect mice from tumor outgrowth, at about 80–90%. These results provide a scientific basis for the evaluation of HPV p-vaccine and HPVAd5-vaccine in human trials for therapeutic intervention against HPV16-induced disease.

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**References**


