HPV16E7d Candidate Vaccine Modified Genes Expression in Tumor Environment: Flic as Adjuvant Changed TGF-B Gene Expression

Abstract

Cervical cancer is one of the most important venereal disease and leads to mortality in women. Human papilloma virus causes this disease. This virus has three protein regions that include early, late and non-coding areas. The early region consists of E1 to E7 that E6 and E7 are two early oncoproteins of HPV which are responsible for disordering in cell cycle and tumor induction. In this study, the efficiency of Flic-E7d-HPV16 and HPV16-E7d-Flic were investigated in tumor bearing Mice models. Tumor was induced in mice with transplantation method. After tumor growth, the mice were divided into five groups. Two groups of C57BL/6 mice were injected with 20 µg of the vaccine Flic-E7d-HPV16 and E7d-HPV16-Flic subcutaneously three times over a one week interval. The third group only E7d was injected and for fourth group only Flic was injected and for fifth group PBS was injected. One week after final immunization, tumor was assessed for gene expression and pathologic smear. According to pathological studies on the tumor samples of mice which indicated the tumor was kind of Sarcoma, Necrosis, Apoptosis, Mitosis and vessel generation were evaluated in each group. Apoptosis and angiogenesis in A, B group were not observed. According to investigation of TGF-β and VEGF genes and HPRT control gene using PCR-real-time, there was a decrease of VEGF and TGF-β genes in group A and B. So the vaccine of group A, B were potent in the reduction of angiogenesis. As a result, Flic as an adjuvant improved anti-tumor effect of HPV16 E7d vaccine.

Keywords: HPV16E7d Flic; Gene expression; Pathology

Introduction

Cervical cancer is one of the most important disease leads to mortality in women the entire world. HPV is directly related to cervical cancer [1]. This virus has three protein regions that include early, late and non-coding areas. The early region consist of E1 to E7 that E6 and E7 are two main oncoproteins of HPV, which are responsible for disturbance in cell cycle and tumor progression of infection [1]. E7 protein with binding to retinoblastoma protein result in uncontrolled cell mitosis [2-5]. So that, Because of the risk of oncogenic, it was utilized as a type of mutant protein with three substitution in amino acids. Therefore called E7 detox (E7d). While changes have been prevented from binding proteins to Retinoblastoma protein (pRb), it has been maintained in immunogenic property. Carcinogenesis potency of high risk Papilloma virus is associated with E6 and E7 protein that is expressed in cancerous cells constantly. Although these proteins are small, they have high capacity in binding to regulatory proteins of host cell to create malignancy in target tissues [6-9]. Todays, it is believed that a bare protein is not enough immunogenic and should be formulated in adjuvants. Actually adjuvants are some substances that increase vaccine immunogenicity and thereby vaccine potency. Adjuvants used in combination or mixture with vaccines by various mechanisms improve immune responses [10,11]. Such adjuvants are microbial product which bind to the Toll like receptors and induce inflammation. The microbial products that bind to the TLR5 is flagellin (Flc). In fact, Flc as adjuvant is able to bind to TLR5 on dendritic cells and macrophages [12,13]. So that, some signals are generated to activate adaptive immune system [13,14]. Some studies showed that VEGF could induce angiogenesis through effect on endothelial cells directly and also facilitate access oxygen and nutrients to tumor growth. VEGF plays a central role as a mediator in tumor growth and angiogenesis [15,16]. Another factor that should be considered as tumor progression is TGF-β. TGF-β suppresses immune responses in favor of tumor growth. On the other hand, studies show that tumor cells with secretion of TGF-β could escape from immuno surveillance mechanisms [17]. Until now, different procedures were utilized to create therapeutic vaccines against of human Papilloma virus such as DNA, peptide, protein, whole cell vaccines. Todays, utilization of TLRs agonists is a method to increase vaccine efficacy [13,14]. Herein, type B pseudomonas aeruginosa recombinant FliC was used as adjuvant. In this study, Flic-E7d-HPV16, HPV16-E7d-Flic protein efficacy was investigated as vaccines in tumor bearing mice models. Also, tumor environment events were investigated, furthermore, after vaccination, TGF-β, VEGF genes expression level and apoptosis, necrosis, mitosis and angiogenesis was appraised in pathologic slides.
Material and Methods

Mice

Female six-to-eight week old inbred C57BL/6 mice were purchased from Pasteur Institute of Iran (Karaj) and kept them for one week before experiment. Mice were kept in animal room with suitable ventilation at 20-22°C. Given free access to water and food and maintained in a 12hr light/12hr dark cycles. Handling of the mice was carried out with a professional technician and in accompanied by the Animal Care and Use Protocol of Pasteur Institute of Iran.

Tumor modeling

Tumor was induced by cell line injection manner using TC1 cell line. Briefly, after cell suspension preparation of TC1 cell line, the cells washed and PBS was added and suspension adjusted to 1×10⁶ cells/ml. Finally, one million cells was injected at the flank of C57BL mice. After two weeks, tumor developed and used as stock for induction of tumor induction in native mice via transplantation method. Briefly tumor dissected to 3-5 mm pieces and transplanted to the flask of anesthetized mice.

Vaccine preparation

Flic-E7d, E7d-flic, Flic and E7d recombination protein was produced and purified by Meysam Gachpazan at Department of Virology from Pasteur Institute of Iran and gifted for this study. Flic protein was utilized as a recombination flagellin of pseudomonas aeruginosa which was produced previously by Dr. Sobhan Faezi [18] from Pasteur Institute of Iran.

Study groups and immunization

Tumor Bearing mice were divided into five groups. Two groups of tumoral C57BL/6 mice were injected with 20µg of vaccine Flic-E7d-HPV16, and E7d-HPV16-Flic subcutaneously, for third group only E7d was injected and fourth group only Flic was injected and for fifth group control PBS was injected. Adjuvant (Flic) and PBS as control groups. Each group was marked by punctuation on the ear. Immunization was done three times with one week interval. After last inoculation, tumor samples of experimental mice removed and analyzed for gene expression and pathology parameters.

Primer design

Primers (VEGF1 forward 5’-TGT ACC TCC ACC ATG CCA AGT-3’; VEGF1 reverse 5’-TGG TAG ACC TTC ACC ATG AAC TTG-3’), (VEGF2 forward 5’-GGA GAT CCT CCG AGG AGC ACT-3’; VEGF2 reverse 5’-GCC GAT TTA GCA GCA GAT ATA AGA-3’) and (TGFβ forward 5’-CAC CTG GAA GAC CAT CGA CAT-3’; TGFβ reverse 5’-GAG CCT TAG TTT GGA CAG GAT CTG-3’) were designed to amplify 145, 94, 130 and 85 base pairs fragments respectively. These small primers (less than 250 bp) were attached to mRNA on the border between two exons. All of the primers were apprised with primer express 3.0 software and adopted among papers. Selected primers sequences were used by blast software.

RT-PCR assay

The method was performed according to the optimum set up of our laboratory. Briefly, RNA of tumor samples was extracted with RNX Plus (Qiagen) and cDNA was synthesized. Gene expression was assessed by Real Q-PCR 2x Master Mix Kit. Along with, SYBR green was used to identify amplified PCR production. Also, 5 µL of SYBR Taq II, 1µL of PCR forward primer, 1µl of PCR reverse primer, 1µl of Template, 2µl of dH2O in reaction mixture was used. To carry out Real time PCR reaction, Rotor Gene 6000 thermal cycler device (Corbett Company) was used. Temperature plane includes the following steps: initial denaturation (95°C, 15 min) denaturation (95°C, 10 s), annealing and extension (62°C, 20 s) and the steps consist of 40 cycles. In other to properties evaluation of PCR, Melting curve was used at 43°C. RT-PCR method was optimized with primer binding temperature and primer concentration optimization.

Smear pathology preparation

Experimental tumor samples were placed in Tissue-Process for 12h. After that, the fixed samples are ready to section and were stained by H&E. eventually; prepared slides were investigated at Pathology Department of Firoozgar Hospital (Tehran, Iran).

Statistical analysis

All analysis was carried out by Graph pad prism V6.01 software and ANOVA test for statistical analysis and P value < 0.05 was considered as significant difference.

Results

Real time PCR

Results of VEGF gene expression showed significant decrease in Flic-E7d , E7d-Flic and E7d vaccine candidate groups as in comparison to PBS and Flic control groups (p<0.0005) (Figure 1). Also, the VEGF gene expression level in the groups immunized with Flic-E7d, E7d-Flic and E7d did not show significant differences (p>0.05). Results of TGF-β gene expression showed significant decrease in Flic-E7d vaccine candidate group as compared with PBS control group (p=0.004). Insignificant decrease was observed in Flic-E7d vaccine candidate group in compared to E7d-Flic group (p=0.44). TGF-β gene expression at group received vaccine formulated with Flic-E7d does not show significant differences versus Flic control group (p>0.05). Evaluation of TGF-β gene expression results at group immunized with E7d-Flic versus E7d vaccine candidate group did not show significant difference (p=0.05). Also, immunization with E7d-Flic showed decrease versus PBS control group, however that difference was not significant (p>0.05). Group received E7d didn’t show significant differences as compared with Flic and PBS control groups (p>0.05) (Figure 2).

Pathologic results

Apoptosis: Evaluation of apoptosis Results shows that immunization with Flic-E7d cause significant increase apoptosis versus control groups (Flic vaccine and PBS) (P=0.0012, P=0.0185 respectively). Also, immunization with E7d-Flic vaccine candidate shows significant increase of apoptosis level versus Flic control group (P=0.0186). Apoptosis level at group received E7d vaccine has increased as compared with PBS and Flic control groups (P=0.0437, P=0.003 respectively). The apoptosis level at Flic-E7d, E7d and E7d-Flic groups shows no significant differences versus each other (p>0.05) (Figure 3).
Necrosis: Necrosis results showed that immunization with E7d-Flic significantly increases necrosis percent versus other experimental groups (Flic-E7d, Flic, E7d, PBS) (P<0.05). Necrosis percent at group received Flic-E7d didn’t show significant differences as compared with Flic, E7d and PBS groups (p=0.05). Immunization with Flic-E7d does not show significant differences when compared with E7d, Flic and PBS control groups (P>0.05). (Figure 4).

Mitosis: Immunization with Flic-E7d significantly decreases mitosis percent versus E7d-Flic and E7d groups (P<0.0004) but
does not show significant differences versus PBS and FliC control groups (P>0.05). Immunization with E7d-Flic shows significant increase versus PBS and FliC control groups (P<0.0001), but not versus E7d group (P>0.05) (Figure 5).

**Angiogenesis:** Immunization with Flic-E7d and E7d-Flic show a significant decrease angiogenesis percentage versus E7d vaccine group (P<0.0042). But, immunization with Flic-E7d and E7d-Flic does not show significant differences as compared with Flic and PBS control groups (P>0.05). Immunization with E7d significantly increase angiogenesis percentage versus Flic and PBS control groups (P=0.0042) (Figure 6).

**Figure 5:** Assessment of mitosis levels in experimental groups. Immunization with Flic-E7d significantly decrease mitosis percent versus E7d-Flic and E7d groups (P<0.0004).

**Figure 6:** Results of angiogenesis percent in experimental groups. Immunization with Flic-E7d and E7d-Flic show a significant decrease angiogenesis percentage versus E7d vaccine group (P<0.0042).

**Discussion**

In the present study, HPV vaccine with E7d mutated protein of type 16 papilloma virus accompanied with Flic adjuvant was used as a model to describe of immunization versus HPV vaccine candidate in tumor bearing mice. E7d protein is expressed in tumor cells constantly and is considered to cancer treatment followed by HPVs infection [7]. Herein, we evaluated TGF-β cytokine gene expression as key immuno-suppressor cytokine in tumor milieu and also apprised pathology markers that reflect immune responses against tumor tissue. Our results show that administration with Flic-E7d vaccine candidate significantly decreased level of VEGF and TGF-β genes expression. Several studies show that VEGF is up-regulated by hypoxia and some inflammatory cytokines and also through effects on endothelial cells could induce angiogenesis and promote tumor growth. Along with, studies show that TGF-β as a potent regulator factor is considered to suppress immune system and important promoter of cancerous cell growth and progression [19]. Elevated TGF-β converted naïve T cells to regulatory T cells that in turn provide immune response escape and tumor progression [20]. Also, Branca et al. [21] showed that VEGF increased in cervical cancer, and related to high risk HPV in cervical lesions closely [21]. Boil et al. [20] showed that VEGF was elevated in patient with cervical cancer. Yang et al. showed that TGF-β promote tumor growth through changing microenvironment and establishment of suppress condition in immune system [20]. So that, VEGF and TGF-β which are really vital to tumor survival and growth are compatible with other study in this respect. Therefore, our results showed that Flic-E7d could inhibit immune response suppression and tumor growth by decrease of VEGF and TGF-β even than E7d vaccine that in turn reflects positive effect of Flic as adjuvant that modified immune response in the tumor microenvironment. Results of pathology experiments showed that Flic-E7d, E7d-Flic, and E7d candidate vaccine groups significantly increased level of apoptosis as compared with control groups. Moreover, E7d-Flic could significantly decreased angiogenesis versus E7d candidate vaccine group and also Flic-E7d significantly decreased anaplasia percent as compared with two other vaccinated groups. Along with, E7d-Flic significantly increased necrosis percent versus all experimental groups. The apoptosis and necrosis mechanism implicated in tumor elimination [22] as here Flic as adjuvant in E7d vaccine increased these parameters.

Also, studies showed that blood vessels formed by adequate nutrients and oxygen are called angiogenesis by which leads to metastasis of tumor cells. In addition, some studies demonstrated that, in cancer condition mitosis was forced to P53 arrest and followed by this situation uncontrolled proliferation occurred [23,24]. Herein, E7d-Flic and Flic-E7d could decrease significantly angiogenesis percent and efficacy of angiogenesis suppression in these candidate vaccine groups was higher than E7d vaccine. Inhibition of tumor angiogenesis is an important mechanism in tumor treatment via immunotoxins [25]. From this regard, we have shown that E7d as vaccine candidate and Flic as a benefit adjuvant be able to increase of apoptosis, necrosis, and decrease in mitosis and angiogenesis in the tumor tissue samples, also Flic as adjuvant could affects on VEGF and TGF-B genes expression in tumor environment and suppress angiogenic and
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References

Conflict of Interest

Authors declare no conflict for this work.

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