Construction and evaluation of human papillomavirus genotype 18 pseudovirions

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ABSTRACT

Introduction: Cervical cancer is the second most common cancer in women worldwide and the role of human papillomavirus (HPV) has been proved in its etiology. The currently available L1-capid-protein-based vaccine is highly immunogenic and very high titers of serum antibodies can be obtained by its injection, but unfortunately it is restricted to only a few HPV genotypes and is relatively expensive. Therefore, development of the second-generation HPV vaccines has become the focus of the research and L2 capsid protein that is capable of producing broad spectrum antibodies has become one of the main candidates. Evaluation of the vaccine immunogenicity however, requires development of HPV pseudovirions (HPV PvSs) comprised of L1 and L2 virus protein and the present study was an attempt to produce PvSs of HPV18 genotype by an in-house method. Methods: The HPV18 L1/L2 coding plasmid and the reporter plasmid of pEGFP-N1 were amplified in E. coli DH5α and purified using silica oxide method. The plasmids were co-transfected into HEK 293FT cell line and the preliminary analysis of expression was performed using fluorescence microscopy. The PvSs were partially purified using gel filtration chromatography and were used to transduce the 293FT cells to evaluate the infectivity rate of the PvSs. The results were analyzed by fluorescent microscopy, flow cytometry and atomic force microscopy. Results: The results showed that the HPV18 L1/L2 coding plasmid and the pEGFP-N1 reporter plasmid have been successfully co-transfected into HEK 293FT cells and the PvSs were constructed. The 293FT cells were successfully transduced by PvSs that had packaged reporter plasmid. These findings were confirmed by fluorescence microscopy and flow cytometry as well as AFM imaging. Conclusion: In this study, the cotransfection of HPV18 L1/L2 coding plasmid as well as pEGFP-N1 reporter plasmid into the HEK 293FT cell led to the assembly of the pseudovirion harboring the reporter gene. The protocols used in this study were easy to perform and relatively inexpensive and did not rely on the commercial kits.

KEYWORDS: human papillomavirus, cervical cancer, pseudovirion.

INTRODUCTION

Human papillomavirus (HPV), a non-enveloped, double stranded-DNA virus is the most common cause of sexually-transmitted infections with more than 100 known genotypes and genetically infecting HPVs have been divided into high- and low risk genotypes based on their ability to cause cancer [1]. There are at least 13 high risk oncogenic genotypes of which genotypes 16 and 18 have been strongly associated with cervical cancer. A yearly incidence of nearly 500,000 cases of cervical cancer worldwide has been estimated of which approximately 80% occur in developing countries, with a mortality rate of nearly 50% [1-3]. Prevention through vaccination has been the most effective measure in reducing the burden of the infectious diseases, but since the virus is difficult to culture, producing inactivated or attenuated virus as vaccine has not been possible [4]. Promising progress however has been made using spontaneously self assembled virus-like particles (VLP) obtained from purified L1 capsid protein of the virus which lack the viral genome and therefore are non-infectious, but exhibit conformational epitopes similar to the original HPV virions [5]. It has been shown that vaccination with VLPs induces a high titer of virus-neutralizing antibodies [6-8]. Currently two licensed VLP-based prophylactic vaccines are commercially available. The HPV16, HPV18 bivalent and HPV6, 11, 16, 18 tetravalent vaccines were developed by Merck and GlaxoSmithKline (GSK) respectively and are prepared from purified recombinant L1 proteins. Clinical
evidence has shown the efficacy of these vaccines against the genotypes from which the L1 originates, but are ineffective against other high risk genotypes that are shown to be present at significant levels in many countries [9]. Efforts are underway to develop a nine-valent combination vaccine, but the inclusion of many VLP types might lead to antigenic competition resulting in lower protective efficacy while increasing the cost and complicating vaccine development [10]. To address these issues the use of L2, the minor capsid protein of the HPV has been proposed in animal studies has been shown to induce broad-spectrum cross-neutralizing antibodies, thereby offering broader protection [11]. However it has been reported that the immune response to the low immunogenic L2 protein does not increase due to the immunodominance of L1 making incorporation of the L2 in L1 VLPs less effective necessitating the development of new vaccine strategies [12]. One such strategy is construction of HPV pseudovirions (PsV) consisting of the L1 and L2 virus proteins with the ability to attach and enter the cells [13]. Production of such particles requires the simultaneous expression of the L1 and L2 genes and their assembly on the surface of the PsVs. Furthermore, to evaluate the ability of the PsVs to enter the cells, a plasmid coding for a reporter gene is co-transfected with L1/L2 coding plasmid which would be packaged within the PsVs during the capsid assembly. In this study construction and partial purification of HPV 18 PsVs, harboring EGFP reporter gene as a tool for assessment of neutralizing antibodies produced during vaccination was undertaken.

MATERIALS and METHODS

E. coli DH5α and pEGFPN-1 plasmid (Clontech, USA) were from the stock in Virology Department of Pasteur Institute of Iran and the HEK 293FT cell line was purchased from Invitrogen (USA). The HPV18 L1- and L2-coding sequences cloned in an IRES-containing mammalian bicistronic expression vector which was kindly provided by DKFZ (Cancer Research Center, Germany) and named pHPV 18L1h/L2h. All restriction enzymes and molecular biology materials were purchased from Fermentas (Lithuania). Cell culture medium and supplements were obtained from GibCO (USA).

Amplification and purification of plasmids

Separately transformed pEGFPN-1 and HPV18 L1h/L2h plasmids were maintained in E. coli DH5α and purified for further manipulation using Trigon-Silica method [14]. The purified plasmids were kept at -20°C until used.

HEK 293FT cell line transfection

The HEK 293FT cells (2 x 10^5) were grown in DMEM medium supplemented with 10% FBS, 100 IU/ml penicillin, and 100 μg/ml streptomycin (Gibco, USA) in 10 cm plates until the confluence of the cells reached 40-50%. Calcium phosphate method was used for co-transfection of HEK 293FT cells with purified pEGFP-N1 and HPV18 L1h/L2h plasmids [15]. Briefly, 5 μg of each plasmid in 438 μl deionized sterile water was added by 2 μl of 2 M CaCl2 and mixed with 500 μl of 2x HBs buffer (280 mM NaCl, 10 mM KCl, 1.5 mM NaHPO4, 12 mM Dextrose, and 50 mM HEPS pH 7.05) and kept at room temperature for 10-15 minutes. The solution was then added drop wise to the HEK 293FT cells and incubated in 5% CO2 at 37 °C. The growth medium was replaced with fresh medium after 12 h and the cells were further incubated for 60 h. The harvested cells were suspended in PBS buffer (pH 7.4) and analyzed for EGFP expression by flow cytometry (CyFlow SL, Partec, Germany). About 2 x 10^6 cells were analyzed in the FL1 detector. Transfection efficiency was also determined visually by fluorescent microscopy. Visible light and UV/fluorescent images were captured at 10 x magnification using an inverted microscope with fluorescence attachment (Jenius, China) and Panasonic Lumix DMC-LX5 compact digital camera (Panasonic, Japan). The digital images were level adjusted with Adobe Photoshop software to achieve an equivalent background, before the approximate percent of transfected cells was determined.

Pseudovirion maturation and partial purification

The HEK 293FT–transfected cells were trypsinized 60 h post-transfection, and were washed and pelleted by centrifugation. The cell pellet was washed twice and dissolved in 1.5 cell pellet volume of DPBS-Mg buffer (containing plain PBS added by 0.9 mM CaCl2, 0.5 mM MgCl2, 2.1 mM KCl and 350 mM NaCl pH 7.4) and centrifuged at 200 x g for 10 min. The pellet was dissolved in 1.5 cell pellet volume of DPBS-Mg buffer, and Triton X-100 (0.5% v/v) and RNase A (5 U/μl) were added to the suspension for cell lysis and ribosomal RNA removal. After 24 h incubation in 5% CO2 at 37°C for pseudovirion maturation, the suspension was chilled on ice for 5 min and clarified by centrifugation at 5000 x g for 10 minutes at 4°C. The efficiency of the cell disruption was assessed by light microscopy.

For removal of Triton X-100 detergent, an in-house gel filtration chromatography method was used. Briefly, 1 ml of Sephadex G-25 at the concentration of 30 g/liter with a ratio of Sephadex/PBS of 1:1 v/v was poured into combed cotton-stuffed 1 ml-syringe. The syringe was placed on a drilled microcentrifuge tube, and then the system was entirely placed in a 50-ml conical centrifuge, and centrifuged 1 min at 4500 x g. Then, 0.5 ml of pseudovirus suspension was added to the syringe and centrifuged 1 min at 4500 x g.

Ten microliters of the flow-through was diluted 1:100, 1:500 and 1:1000 and spotted on mica slides, air-dried and visualized by atomic force microscopy (NanoWizard II, JPK, Germany).

Transduction of 293FT cells

To evaluate the infectivity of the pseudovirions, the 293FT cells were grown in DMEM medium containing 10% FBS in 6-well plates (SPL, South Korea) in 5% CO2 at 37°C until cell confluency was 40-50%. The cells were transduced by replacing the culture medium with fresh medium containing 1/50 dilution of pseudovirions suspension flow-through from previous step and incubated in a CO2 incubator at 37°C for 48h. The efficiency of transduction of 293FT cells with the pseudovirions and expression of the reporter gene was evaluated by fluorescence microscopy and flow cytometry as described in the cell transfection methodology.

RESULTS

Plasmid verification

Restriction enzyme digestion of the HPV18 L1h/L2h and pEGFP-N1 plasmids extracted from E. coli DH5α, with HindIII resulted in 2100 bp and 6600 bp fragments for HPV18 L1h/L2h and linearization of pEGFP-N1 respectively, verifying the authenticity of the plasmids (Fig. 1).

Transfection of HEK 293FT cells:

Expression of EGFP in HEK 293FT cells was observed after co-transfection of HPV18 L1h/L2h and pEGFP-N1 plasmids after 72 h (Fig. 2A). The calculated rate of transfection was
approximately 80% which was determined by fluorescent microscopy and confirmed by flow cytometry results (data not shown).

**Pseudovirions purification**

The HEK 293FT cells containing pseudovirions were sedimented and the cells were lysed, and the pseudoviral particles were partially purified by Sephadex G-25 before imaging by AFM (Fig. 2B). The size of pseudovirions was estimated as around 45-55 nm.

**Transduction of 293FT cells**

The infectivity of the purified PsVs was evaluated by transducing the newly cultured HEK 293FT cells. Expression of the EGFP reporter gene in the cells indicated that pEGFP-N1 plasmid had been packaged inside the PsVs. The results showed that approximately 30% of transduced cells expressed the reporter gene as determined by fluorescence microscopy (Fig. 2C) and flow cytometry (data not shown).

**DISCUSSION**

Cervical cancer is the second most common cancer among women and neutralizing antibodies are the most effective immune response to prevent the infection with HPV and the in vitro analysis of the magnitude of this response requires production of the HPV PsVs. Pseudovirions express the major and the minor L2 capsid proteins (L1 and L2), resemble the surface of the intact virus. PsVs can be used for vaccine and drug development and have become the standard tools for studies covering various aspects of the virus biology, for gene transfer or to determine the effectiveness of drug therapies [16, 17].

In this study, we prepared HPV18 PsVs with simple and less expensive methods that did not rely on commercial kits. Our data also showed that HPV PsVs prepared by these techniques are capable of infecting the HEK 293FT cell line in vitro. Various methods and cell types have been used for production of PsVs, but it has been suggested that a single 75 cm² flask of 293TT cells, a variant of HEK 293T cells engineered for high level expression of SV40 large and small T antigen required for amplification of the target plasmid containing SV40 origin of replication [18-19] is capable of producing a minimum of 1 mg of transducing PsVs. However, the drawback to the use of this cell line despite its high yield is the possibility that the SV40 T antigen could be incorporated into the PsVs through promiscuous packaging turning the innocuous PsVs into autonomous tumor viruses [19]. It has been reported that the yield of PsVs in 293FT cell line is lower than that of 293TT cells due to either lower expression of SV40 T antigen in these cells or the lack of the SV40 small t antigen [19].

The transfection rate of 80% obtained in this study using calcium phosphate compared favorably with the more expensive commercial lipofectin. The reason for the low rate of infectivity (30%) with PsVs was not ascertained, but the pseudovirions were only partially purified and the contaminating debris might have affected the infectivity rate. In this study we prepared HPV18 PsVs using an IRES-containing bicistronic vector coding for HPV18 L1 and L2 proteins that were packaged with a reporter plasmid that could be tracked to assess the infectivity of the pseudovirus particles. The methods employed were easy to perform and relatively inexpensive and did not rely on commercial kits which may facilitate their application in developing countries.
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CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

REFERENCES