Effect of human papillomavirus infection on the immune system and its role in the course of cervical cancer (Review)

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Abstract. Human papillomavirus (HPV) is widely known as a cause of cervical intraepithelial neoplasia (CIN) and cervical cancer. The mechanisms involved have been studied by numerous studies. The integration of the virus genome into the host cells results in the abnormal regulation of cell cycle control. HPV can also induce immune evasion of the infected cells, which enable the virus to be undetectable for long periods of time. The induction of immunotolerance of the host's immune system by the persistent infection of HPV is one of the most important mechanisms for cervical lesions. The present review elaborates on the roles of several types of immune cells, such as macrophages and natural killer cells, which are classified as innate immune cells, and dendritic cells (DCs), cluster of differentiation (CD)4+CD8+ T cells and regulatory T cells, which are classified as adaptive immune cells. HPV infection could effect the differentiation of these immune cells in a unique way, resulting in the host's immune tolerance to the infection. The immune system modifications induced by HPV infection include tumor-associated macrophage differentiation, a compromised cellular immune response, an abnormal imbalance between type 1 T-helper cells (Th1) and Th2 cells, regulatory T cell infiltration, and downregulated DC activation and maturation. To date, numerous types of preventative vaccines have been created to slow down carcinogenesis. Immune response activation-based therapeutic vaccine is becoming more and more attractive for the treatment of HPV-associated diseases.

Contents

1. Introduction
2. HPV infection regulating the host immune response
3. Immune cell changes in the immune response to hrHPV
4. Immune-related markers of CIN and cervical cancer
5. Preventive and therapeutic HPV vaccines
6. Conclusion

1. Introduction

Cervical cancer accounts for 15% of cancers in females aged <65 years old in developing countries (1). Long-term high-risk human papillomavirus (hrHPV) infection must be present, but is insufficient to cause cervical cancer by itself according to the International Agency for Research on Cancer (IARC). Certain other risk factors account for the development of cervical cancer, including an active sexual history, weakened immune system function and smoking (2). The induction of cervical carcinogenesis by persistent hrHPV infection is a multistep process, consisting of persistent infection, different stages of cervical intraepithelial neoplasia (CIN) lesions and ultimately, cervical cancer (3). Although persistent hrHPV infection is an important prerequisite for CIN, the vast majority (90%) of viruses have been cleared by the hosts’ immune system, without medical intervention, as observed on the three-year follow-up examination, and only 1-2% of the remaining 10% become chronic resulting in cervical cancer (3).

To date, >180 types of HPV are known to exist (4). HPV16, 18, 31, 33, 35, 45, 51, 52, 56, 58, 59, 68 and 69 are classified as the hrHPVs (5). HPV6, 11, 40, 42, 54, 55, 61, 62, 64, 71, 72, 81, 83 and 84 are classified as the low-risk HPVs (6). HPV16 and 18 are the most common types, accounting for ~70% of cervical cancers around the world (7). HrHPV infection starts with the contact of the virus with the basement membrane, which is often exposed by micro-abrasions on the cervical surface. Suitable receptors increase the probability of hrHPV infection. Studies have found several receptors involved in this process. Interaction between the virus’ capsid protein and the cell receptors promotes virus capsid conformational changes, thus aiding the cell entry process (8). Tissue-specific heparin sulfate proteoglycan (HSPG), as one of the members of the glycosaminoglycan
family, is a receptor of major capsid protein L1 of HPV (9,10). HSPG could aid HPV16 in binding to the extracellular matrix via laminin-332 (11).

HPV DNA is double-stranded, containing 7,900 base pairs, which are arranged in a circle (12). The genome consists of 8 open reading frames, 6 early genes (E1, E2, E4, E5, E6 and E7) encoding the early protein and 2 late genes (L1 and L2) encoding the late protein. E1 protein aids in viral replication utilizing the host replication machinery, E5, E6 and E7 proteins are considered to be associated with virus immune evasion. The product of E5 could downregulate the histocompatibility leucocyte antigen (HLA) expression of the infected cells, facilitating virus immune escape (13,14). E6 and E7 are considered as tumorigenic genes; their products could bind to tumor suppressor protein p53 and arrest cell cycle, impairing the infected cells apoptosis and enhancing their transformation (15). The L1 and L2 proteins are responsible for forming the structural components of the viral capsid. E2 exhibits a regulatory function for E6 and E7 transcription. HPV genome integration into the host genome would disrupt the regulatory function of E2, leading to the uncontrolled expression of E6 and E7, promoting the progression of the disruption of the normal cell cycle and carcinogenesis (16).

Furthermore, the expression of E6 and E7 is fundamental in hrHPV infection and CIN progression. The expression of these oncoproteins inhibits the immune system response to hrHPV and aids in the persistency of the infection. The oncoproteins could downregulate TLR9 expression, a virus DNA sensor, which is necessary to activate antigen-presenting cells (17), but they could also reduce the expression of transporter-associated antigen processing 1, blocking the activation of specific T lymphocytes. It has been confirmed that neoplastic cervical keratinocytes (KC) expressing high levels of E6 and E7 oncoproteins could escape the attack from cytotoxic T cells (CTLs) (18).

2. HPV infection regulating the host immune response

According to a previous study, the incubation period of HPV infection is ~10 years. The immune response of the host plays a crucial role in the progression or regression of hrHPV infection of the uterine cervix. An effective immune response promotes spontaneous clearance of the virus, while a compromised immune response often starts up the pathological process, developing it into the higher grade (19). HrHPV infection causes several events, which are essential for CIN progression and carcinogenesis (Fig. 1).

HrHPV infection promotes immune cell migration to the dermis. In the squamous epidermis, macrophages, Langerhans cells (LC), KCs, T lymphocytes, dendritic cells (DC), natural killer cells (NK) and B lymphocytes play important roles during the immune response to infection. HrHPV infection could cause the immune system to become more tolerant to the infection, thus creating a microenvironment susceptible to further infection and facilitating CIN progression. The mechanisms that have been proposed and proved are as follows: Firstly, hrHPV remains silent for a long time; its duplication and assembly do not cause cytolysis or the cytopathic death of the host cells (20). Secondly, hrHPV inhibits interferon (IFN) synthesis through E6 and E7 oncoproteins interfering with IFN signaling pathways (21).

Thirdly, hrHPV infection induces regulatory T cell (Treg) infiltration and interleukin (IL)-10 or transforming growth factor β (TGF-β) production. Fourthly, the infected cells express low levels of MHC class I, resulting in impaired CTL function (22). Fifthly, they could induce an accumulation of ineffective CD4 and CD8 T lymphocytes in stage II/III CINs (23).

Additionally, KCs are the major cell type in the epidermis, with certain immune functions, such as cytokine secretion. The compromised innate immune defense in KCs is an important reason for the evasion of the hrHPV infection from the immune response, ultimately resulting in persistent virus infection and the development into pre-neoplastic lesions (24).

Innate immunity affected by hrHPV infection. Usually, once the hrHPV contacts the mucosal epithelium, the innate immunity system mediated by the epithelial barrier starts to conflict with it. LCs are immature DCs, and in the transformation zone their numbers are significantly decreased. The mechanism proposed for this is the direct interaction of E7 with CCAT enhancer-binding protein β, a transcription factor of chemokine (C-C motif) ligand 20 (CCL20) (which has a decisive role in the migration of LC precursors into the epidermis), thereby inhibiting the transcription of CCL20 and thus hindering LC recruitment (25).

Toll-like receptors (TLRs) are an important type of pattern recognition receptor located at the endolysosomal compartments, sensing the bacteria/virus and triggering the associated innate immune response. Several of these receptors have been investigated. HrHPV18 E6 and E7 downregulate TLR9 (specific for the nucleic acid) expression at the infection site, an important strategy for its escaping from immunosurveillance (26). In contrast to TLR9, the TLR3/5/8 pathways are activated in hrHPV-infected KCs. High expression levels of TLR8 in cervical cancer cells are associated with the upregulated expression of B-cell lymphoma-2 and vascular endothelial growth factor in these cells (27). A previous study found that there is marked TLR4 expression in hrHPV-positive cells and that its expression is associated with the virus type, as well as the histopathological grade. Higher TLR4 expression is found in HPV16+ cells compared with HPV18+ cells, and in cervical cancer compared with CIN. Additionally, it has been demonstrated that TLR4 overexpression is correlated with the apoptotic resistance of the HPV-infected cells (28).

It is known that tumor-associated macrophages, also termed M2 macrophages, can promote cancer cell proliferation and migration, angiogenesis and the restriction of immune defenses (29). During cervical lesion progression, the number of M2 macrophages is significantly increased (30). The aggregation of M2 is a key event for the pathological process of carcinogenesis.

HrHPV infection compromises NK cell activation. NK cells predominate at the initial stage of the infection and in the low-grade lesions. Levels of NK-activating receptors, such as NKP30, NKP45, NKP46, NKG2D and NKP80, are significantly decreased in HPV16 cervical cancer. These receptors are closely associated with the low cytotoxic activity of NK cells, facilitating lesion progression and carcinogenesis (31). hrHPV-infected tissues or cervical cancer cells could obtain their immune evasion abilities by making NK cells more tolerant to the hrHPV infection and cervical carcinogenesis (32).
3. Immune cell changes in the immune response to hrHPV

Compromised adaptive immunity is the foundation for the progression from hrHPV infection to cervical cancer. There are different immune cell profiles for the different stages of the disease progression in CIN and carcinogenesis. The changes and modifications induced by hrHPV infection are involved in hrHPV infection adapting the immune system to create a suitable microenvironment for persistent infection and lesion progression.

*HPV infection induces compromised T cell activation.* T cell activation is important in hrHPV infection. It has been proven that in patients with HIV infection, compromised
CD4+ T cells function as the inducer for HPV-associated cancer occurrence (20). For these patients, the induction of inefficient CD4+ T cells by hrHPV infection is the factor promoting CIN lesion progression (33). CTL is the main agent in cancer specific immunity, and it recognizes the antigens with the assistance of MHC/HLA class I. HPV16 E5 could suppress CTL through the downregulation of MHC/HLA class I expression, as aforementioned.

Distorted equilibrium between type 1 T-helper cells (Th1) and Th2 cell is another property of cellular immunity during hrHPV infection (34). There is a marked Th2 cytokine profile in intraepithelial and invasive cervical lesions. Increased Th2 cytokine (IL-10) and reduced Th1 cytokine (IFN-γ, IL-12, IL-2 and tumor necrosis factor-α) levels have been detected in cervical exudates of hrHPV+ patients. This indicated that the reduced Th1 response and increased Th2 response lead to cellular immunity suppression and cervical lesion progression (35,36). Another study proved that the production of certain Th2 cytokines is also decreased during this process, indicating that the Th1 and Th2 phenotypes, but particularly the Th1 phenotype, may be suppressed by hrHPV infection. One of the mechanisms by which hrHPV regulates T cell activation is based on E6 and E7 oncprotein expression, which would upregulate the expression of adhesion molecules, such as intercellular adhesion molecule 1, vascular cell adhesion molecule 1 and E-selectin (classical immunosuppressive molecules), in the infected cells (37).

Overall, compromised T cell activation and the distorted equilibrium between Th1 and Th2 presents hrHPV with the ability to escape from the human immune defense.

Compromised DC activation induced by hrHPV infection. DCs, as one of the important adaptive immune cells, recognize the special patterns of pathogens through TLRs and present the antigens to T cells through MHC molecules to initiate the antigen-specific immune response. A high level of stromal DCs is usually associated with a higher frequency of infection regression. Along with hrHPV infection, immature DCs usually prevent the activation of a correct immune response by CTLs and facilitate lesion progression (38). There is an extremely low distribution or complete absence of functional DCs in patients with cervical cancer. hrHPV E6-expressing cells or hrHPV+ cancer cells can inhibit the differentiation of monocytes into fully functional DCs (39). A low percentage of plasmacytoid DCs in the peripheral blood is significantly associated with persistent hrHPV infection (40). Furthermore, it has been demonstrated that in hrHPV+ patients, decreased expression levels of CD80 and CD86 in DCs have a positive correlation with increased CIN grades (41).

One of the mechanisms involved in the induction of compromised DC activation in hrHPV infection was proposed as the stimulation of programmed death 1 (PD-1)/PD-1 ligand (PD-L1) (CD279/CD274) pathway activation by chronic hrHPV infection in DCs. PD-1/PD-L1 activation is associated with impaired DC-mediated immunity to hrHPV infection, thus creating the immune tolerance microenvironment for persistent infection and CIN progression (41).

Interplay between Treg cells and hrHPV infection. Tregs (CD4+/CD25+/Foxp3+ T cells) are the inducer of immune tolerance. There is increased Foxp3+ Treg infiltration following hrHPV infection. Recent studies have highlighted the role of hrHPV-specific Treg generation induced by hrHPV infection and CIN progression (42). Tregs specific for the E6 and E7 antigens have been detected in high-grade squamous intraepithelial lesion (HSIL)-infiltrating lymphocytes (43), and there are several mechanisms that contribute to their generation and recruitment. Firstly, improper activation of the immune response induced by hrHPV infection provides the possibility of the toleration of T cell generation. Secondly, the weakened innate immune function elicited by the infected KCs creates an immunosuppressive microenvironment, promoting HPV-specific Treg expansion (22).

The regulation of hrHPV infection and CIN progression by Treg cells is mainly based on cytokine production. For example, TGF-β is produced by Treg, and its level has a positive correlation with lesion progression (36). IL-10 is another significant Treg-producing cytokine whose presence would decrease CD8+ T cell infiltration and increase the amount of intratumoral Foxp3+ Treg cells, a critical event for CIN lesion progression (44,45). There is also a notable interaction between IL-10 and HPV-induced lesion progression (46). In hrHPV infected cells, E2 protein binds to the regulatory region of the human IL-10 gene (-2054 nt) and induces its expression; IL-10 can create an immune-tolerant microenvironment, providing the possibility of progressive cervical disease in turn (47).

4. Immune-related markers of CIN and cervical cancer

Immune markers (immune cells, cytokines, chemokines and soluble receptors) during persistent hrHPV infection and CIN progression have not been well studied (48). The percentage of Tregs is not only used as a marker of tumor immune status, but is also considered as a possible marker of immune destruction induced by hrHPV infection. Foxp3, as the only definitive marker of Tregs, exhibits gradually increased expression from CIN 1 to CIN III, which indicates that Foxp3 is involved in the development of cervical cancer (49). However, the exact mechanism of such requires further investigation.

High mobility group box 1 protein (HMGB1), a DNA-binding protein, is critical for host immune suppression. There is a direct correlation between HMGB1 expression and the malignant potential of CIN. A previous study also proved that high levels of HMGB1 expression can be used as an early prognostic marker in recurrent cervical cancer patients (50). Patients with greater HMGB1 expression have a higher rate of hrHPV infection recurrence than those with weak HMGB1 expression (51). Mechanisms employed by HMGB1 to induce the immune tolerance to hrHPV infection are proposed to be as follows: HMGB1 suppresses the human immune function by upregulating Tregs and promoting IL-10 production. It also inhibits the function of T cells by downregulating NF-κB signaling and polarizing Th1 cells to Th2 cells. HMGB1 is becoming a useful biomarker for the evaluation of hrHPV infection persistency and CIN progression (52).

p16INK4a expression is induced by the integration of hrHPV DNA into the host genome; it is the cyclin-dependent kinase inhibitor and is overexpressed in CIN (53). p16INK4a
expression has a positive correlation with the CIN grade, and could be used to discriminate between different CIN grades in cervical biopsies.

Generally, hrHPV L1 capsid protein is expressed in every stage of CIN, with a negative association with the CIN grades (54). The aborption of hrHPV cycles in HSIL results in the L1 capsid protein being non-detectable (55). From the immunological perspective, hrHPV L1 is the inducer for the CD8+ and CD4+ T cell response against hrHPV infection (56). Therefore, reduced L1 capsid expression is a significant mechanism for the immune evasion of higher grade lesions.

5. Preventive and therapeutic HPV vaccines

The majority of clinical trials concerning HPV-targeted therapies are based on eliciting cell-mediated immune responses with vaccines. Finding a target to generate a specific immune response is the strategy for the treatment of hrHPV infection, as well as cervical cancer (57). The preventive vaccination against HPV16 and 18 has now become widely used (58). Antigen-specific immunotherapy is one of the effective methods to elicit immune responses to hrHPV. *Listeria monocytogenes* (LM) has been used to eliminate palpable, vascularized tumors in several mouse models due to its ability to generate CD8+ tumor-infiltrating lymphocytes. Two vaccines, LM-LLO-E7 and LM-ActA-E7, have been created by a truncated listeriolysin O (LLO) fused to E7 and a fragment of the ActA protein fused to E7, respectively. These vaccines overcome central tolerance by expanding low avidity CD8+ T cells specific for E7 (59). A novel fusion protein, HPV16 E7-HBcAg-Hsp65 (VR111), which could elicit an E7-specific CD8+ T cell response, is a novel potential preventive vaccine (60).

Therapeutic vaccines aim to clear hrHPV infections and the hrHPV-related cervical lesions, and are mainly targeted to oncoproteins E6 and E7, the only viral proteins expressed in cervical cancer and precursor lesions. Previously, E6/E7 oncoproteins were used as the target of the therapeutic vaccine, which could increase the lesion-infiltrating CD4+ and CD8+ T cells in the intraepithelial neoplasia lesions (61). In a study, a novel HPV16 E6 and E7 gene plasmid containing oligomannose liposomes (OML-HPV) was generated for the immunotheraphy of cervical cancer. With the stimulation of OML-HPV, HPV16 E6-specific CTLs could be generated from peripheral blood mononuclear cells for HPV16+ cervical carcinoma patients (62).

During the analysis of the HPV vaccines, it has been found that the DC-based HPV vaccine is a promising tool to prevent and treat hrHPV infection, as well as cervical cancer. DCs generated from peripheral blood monocytes with IL-4 and granulocyte-macrophage colony-stimulating factor were treated with HPV16 mE7, which showed the significantly increased expression of co-stimulatory molecules CD80 and CD40, and the marked production of IL-12p70 and IFN-γ. This subset of DCs could upregulate E7-specific CD8+ T cell responses in patients with hrHPV-associated cervical cancer (63). HPV16 E7 polypeptide is another DC-based HPV vaccine. DC loaded with hrHPV16 E7 polypeptide in combination with CpG-ODN2006 also showed distinguished immunotherapeutic activity. When SCID mice were inoculated with such antigen-loaded DCs, they exhibit decreased tumor size, increased IgG and IFN-γ levels and increased CTL activity (64). Another way to activate the DC immune response is via genetic modification. A genetically-modified DC vaccine expressing HPV16 E6/E7 fusion oncoproteins was able elicit the action of E6/E7-specific CTLs against cervical carcinoma CaSki cells and induce their apoptosis. These elicited specific protective immunity functions by the HPV-16 E6/E7 fusion protein produced an effective approach against cervical cancer cell growth (65).

Different types of vaccine have been investigated; one of the most important mechanisms for their utilization is based on their initiating the damaged immune response by hrHPV. Schematic representation of this has been presented in Fig. 2.

6. Conclusion

The present review investigated the association between hrHPV and cervical cancer. The immune system plays an important role from HPV infection to CIN and from CIN to cervical cancer. A compromised immune response is the prerequisite for disease progression. One unique feature of HPV infection is that it can affect the immune system in such as way that it presents a much more tolerant state, which facilitates persistent hrHPV infection and cervical lesion progression. The counteraction of hrHPV infection by vaccines is an important tool to stop the course of the infection or oncogenesis. To date, numerous preventive and therapeutic vaccines have been discovered, with the aim of enhancing the damaged immune response to clear the virus and the tumor cells.

References


