A review of the research progress in T-lymphocyte immunity and cervical cancer

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Abstract: Cervical cancer develops as a result of T-cell immune evasion by human papillomavirus (HPV). T-cell immunity requires the participation of many factors, such as antigen-presenting cells (APCs), cytokines, co-stimulatory molecules, etc. HPV vaccines are promising treatments to prevent HPV infection and cervical cancer. This article mainly provides a summary of the number and function changes of T cells during HPV infection and cervical cancer development. Studies on T-cell immunotherapy, which is expected to become a new treatment for cervical cancer after surgery, radiotherapy, and chemotherapy, are also reviewed in this article.

Keywords: T-cell; cervical cancer; human papillomavirus (HPV); immunotherapy

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Introduction

Cervical cancer is one of the malignant tumours that seriously threaten the life and health of women. Nearly 500,000 women are diagnosed with cervical cancer each year (1), and nearly 288,000 of them are dying from the disease. It is widely acknowledged that high-risk human papillomavirus (HR-HPV) is a major risk factor for the development of cervical cancer, while persistent HR-HPV infection is a high-risk factor for the recurrence of cervical intraepithelial neoplasia (CIN). Even though nearly 80% of HPV infections are transient (2), persistent infections are not necessary to develop cervical cancer. When HPV invades the human body, it triggers immune reactions, and cervical cancer results when HPV overcomes the immune system's responses (3). Immunity is responsible for the recognition and exclusion of foreign material and for maintaining the balance of the body—functions which are closely related to the occurrence and development of tumours (4). Recent studies have shown that persistent infection of HPV has links with the body's immunity, particularly cervical T-cell immunity. Thus far, the majority of related studies have focused on tumour immunity therapy, but T-cell immunotherapy after surgical, radiological, or chemical therapy, is expected to become a new treatment method for cervical cancer. We require a more comprehensive understanding of the participating factors of T-cell immunity in order to clarify the changes in T-cell immunity in cervical tumour tissues and peripheral blood. It is only working from this level of comprehension that novel diagnosis and treatment methods for cervical cancer immunotherapy can be developed. This review provides a brief description of the relationship between T-cell immunity and cervical cancer.
Overview of T cells

Thymus-dependent lymphocytes (abbreviated as T cells) are the main components of lymphocytes, comprising 65–75% of the total number of lymphocytes in peripheral blood and >95% of lymphocytes in the thoracic duct. T cells act as “fighters” that resist diseases and tumours through varieties of biological effects, including the direct killing of target cells, regulating B cell antibody production, responding to specific antigens, and producing cytokines. T-cell immunity is called “cell immunity” because the T cells kill the targets directly instead of producing antibodies. T cells can be divided into different subgroups according to different classification methods. Based on surface markers, they are divided by clusters of differentiation into CD4+ and CD8+ T cells. Based on function, they are divided into cytotoxic T cells/lymphocytes (Tc, or CTLs), helper T (Th) cells, regulatory/suppressor T (T reg) cells, and memory T cells.

T-cell subsets and cervical cancer

Helper T cells

Th1 cells can be classified into Th1, Th2, and Th17 subtypes. Th1 play crucial roles in cellular immunity, during which they secrete interferon-gamma (IFN-γ), interleukin (IL)-2, and tumour necrosis factor-beta (TNF-β) (5). These lymphokines further enhance and expand the immune response process and help stimulate other anti-tumour immunity processes. More specifically, IFN-γ can inhibit tumour growth and enhance the anti-tumour activity of natural killer (NK) cells. Th2 play a role in humoral immunity, which bolsters the production of antibodies from B cells, and Th2 may itself secrete IL-3, IL-4, IL-5, IL-6, etc., of which IL-4 and IL-3 are known tumour growth factors which promote the rapid growth of tumour cells. IL-4 exerts an anti-tumour response by inhibiting the activation of NK cells and reducing tumour cell antigen expression. The Th1/Th2 ratio has proven to be a good indicator of the dynamic changes of the anti-tumour response, as the transfer of Th1 to Th2-type cells in the peripheral blood of cancer patients, especially those with cervical cancer, is responsible for the growth of malignant tumours and their evasion from the host’s immunity response (5).

Th17 is the third independent Th cell subtype. In 2005, a study of the immunological pathogenesis of the central nervous system was first to identify Th17 (6). IL-17A is secreted by Th17 and exerts a strong immune effect in the early stages of many inflammatory diseases. As a consequence, the Th17/Th1 ratio was used to assess the changes in immunity (7). Another cytokine secreted by Th17, IL-23, can block certain extracellular pathogens and has a crucial role in inflammation response and autoimmune diseases (2). A study that focused on the transcriptional regulation of Th17 expression evidenced that autoimmune diseases were pathogenically driven by IL-17/IFN-γ-double-producing cells (8).

In recent years, the relationship between Th17 cells and cervical cancer has become an area of intense research focus. Studies have revealed that Th17 cells are elevated in patients with cervical cancer and in those with CIN. The higher levels of Th22, Th17, and Th1 cells in the circulation system may responsible for the pathogenesis of cervical cancer (9). Clinical stage, lymph node metastasis, and invasion were proven to be associated with the increased Th17 cell level in cervical cancer patients, whereas increased T reg cell frequency was associated with tumour differentiation. Furthermore, the Th17/T reg ratio was found to be significantly higher in patients with uterine cervical cancer with lymph node metastasis or invasion than in patients without cancer (10). A similar study showed that the Th17/T reg balance was broken, and the related cytokines were significantly higher in the blood of Uygur patients with cervical cancer or CIN (11). The imbalance of T cells that expressed Th17/fork head box protein 3 (FOXP3) is responsible for the development of uterine cervical cancer by fostering angiogenesis (12). Though the roles of Th17 cells in promoting autoimmune disorders and inflammation responses have been extensively demonstrated, whether and how Th17 cells affect tumour immunity response are still controversial. On one hand, Th17 cells can promote tumour development by inducing angiogenesis and suppressing immune functions. On the other hand, Th17 cells can activate anti-tumour responses by recruiting immune cells to the tumours and activating CD8+ T cells, or by activating the phenotype converting Th1 and producing IFN-γ. This characteristic of Th17 cells might be used for developing it as a target in future therapeutic strategies (13).

Regulatory/suppressor T cells

T reg cells, which are a group of cells with special phenotypes and functions, can be divided into native T reg cells and induced T reg cells depending on their source. Native T reg cells mainly maintain immune tolerance and
immune homeostasis (14). Induced T reg cells can inhibit the anti-tumour response and consequently promote the development of tumours. For instance, CD4+ T reg cells are involved in the anti-tumour response when it is an important component of specific immunity. Since being discovered in previous studies, the transcription factor FOXP3 has become the most commonly and widely used marker for T reg cells, whose expression is closely related with the survival of invasive carcinoma patients (15).

A related study of cervical cancer found that large numbers of CD4 and CD25 cells were recruited by T reg cells in the blood of patients with high-grade squamous intraepithelial lesions (16). The T reg cell numbers in the blood of the cervical cancer and CIN patients were significantly higher than that in the control patients (11). Similar studies reported that T reg cell levels in cervical cancer patients were correlated with the stages of the disease (staged by the Federation International of Gynecology and Obstetrics staging system) and the degree of tumour differentiation, which indicates that T reg cells are closely related to HPV infection (17). A large number of studies have confirmed that T reg cells are associated with cervical immune regulation. As a marker of T reg cells, FOXP3 increment is accompanied by an increased number of cervical lesions and lower tumour cell differentiation (18). In cervical cancer, T reg cells dominate the microenvironment of tumour-positive lymph nodes in high frequencies, thereby increasing the deterioration risk of patients (19). FOXP3+ Treg cells accumulate in the stroma of CIN3, micro-infiltration, and invasive cancers, while FOXP3+ Treg cells gather in the lymph nodes of metastatic invasive cancer, thus possibly creating a network for immune escape (20).

Th17 and T reg cells are unlikely allies, and have broad prospects for the treatment of autoimmune diseases and cancers in humans. T reg cells were shown to promote Th17 cell differentiation in vitro and in vivo, and enhance the function of Th17 cells, which included a protective effect in the host’s defence. In contrast, they exerted detrimental effects in inflammation and in the support of tumour growth. An in vivo study suggested that Th17 cells are the most effective Th subtype in stimulating and supporting the phenotypic stability and expansion of T reg cells. These observations suggest that two kinds of Th cells may reciprocally stimulate each other, the effects of which should be considered when future Th17 and T reg cell-targeted therapies are conducted (21).

Changes in CD4 and CD8

CD4 and CD8 are both glycoproteins and differentiated by their morphology, which is expressed on the surface of T cells as cell receptors. CD4 is expressed mainly on Th and T reg cells, whereas CD8 is expressed mainly on CTLs. Most studies have provided evidence that T lymphocyte subsets are abnormal and have imbalanced proportions in patients with tumours of various types. One study showed that the ratio of CD4+ to CD8+ subtypes in cervical cancer tissue was significantly lower than that in the peripheral blood, and this alteration was more obvious in higher stages of the disease (22). The decreased proportions of tumour-infiltrated CD4+ T cells, as well as the reversed CD4+/CD8+ ratio, are closely related with rapid tumour growth and lymph node metastasis of cervical cancer patients. Regional immune escape is of great importance to cancer progression (23).

In regressing CIN1, CD4+ cells predominate within both the stroma and epithelium, presenting the highest CD4+/CD8+ ratio compared with the progressing CIN1, CIN3, and invasive carcinoma (24). The CD4+/CD8+ ratio is also associated with the 5-year survival rate of cervical cancer, patients with a lower CD4+ T reg cell level with the up-regulated CD4+/CD8+ ratio usually obtaining better clinical outcomes (25). Another study showed that the expression of CD8+ and CD4+ was inhibited or even deleted in CIN and cervical cancer (26). It was also found that in the early stage of cervical cancer, CD8+ cells and CD8+/CD4+ ratio were significantly elevated in patients without lymph nodes metastasis. When the CD8+/CD4+ ratio decreased below 2, both the total survival rate and disease-free survival rate decreased, a finding which was also reported in other research studies (11,27).

Unlike CD4 and CD8 antigens, CD3 is expressed on all subsets of T cells. The density of peritumoral CD3+ cells was found to have the strongest ability in relapse prediction, and the hazard ratio reduced to 0.27 when the CD3+ cell density increased from 795 to 2,043 cells/mm² (28).

T-cell immunity and cervical cancer

HR-HPV, a major risk factor for cervical cancer development, can produce two kinds of oncoproteins (viz. E6 and E7), which affect cell cycle and inhibit cell apoptosis. E6 and E7 are closely related to the genesis and development of tumours. When these two oncoproteins are recognized, and their expression is then inhibited by T-cell-specific immunity, this is known as an anti-tumour immune
response. At present, a large number of studies are being performed to investigate HPV-specific immunity and the immunosuppression of cervical cancer.

**Antigen-presenting cells (APCs)**

APCs exert critical roles for the antigen transportation from the periphery to locally organized lymphoid tissue (29). Dendritic cells (DCs) are a group of APCs that possess unique abilities to activate primary immune responses. DCs capture antigen and transfer the antigen information to activate adaptive immune response. They are not only crucial for primary immune responses, but also important for immune tolerance and T cell-mediated immune regulation (30). DCs can be precisely and effectively delivered in vivo by optimally adjusting the net charge after intravenous administration of RNA-lipoplexes. This thus mediates the efficient uptake and expression of antigens including major histocompatibility complex (MHC), costimulatory molecules, and adhesion molecules, which are required for T-cell activation by the DC populations (31). As the most professional APCs, they induce strong effector and memory T-cell response and mediate IFN-dependent rejection in progressive melanoma tumours (31). Furthermore, DCs produce IL-12, a cytokine crucial for anti-tumour responses, after strong activation (32). CD83+ DCs can be found in cancer biopsies, and phenotypically mature CD83+ DCs in cancer cells express matrix metalloproteinase 9 (MMP-9), a kind of protein known to have vasculogenic and tumor-genesis-promoting properties (33). Studies reported that a massive amount of MHC class II+ CD83+ DCs were observed in the intra-tumoral region of HPV-induced head and neck cancer, where the maturation of DCs (assessed by the CD83/DC-SIGN ratio) was significantly lower than that in the peritumoral stroma (P<0.05), indicating a distinct immune regulatory mechanism between the intra- and peritumoral sites (34). DC-derived exosomes (Dexo) were demonstrated to join ex vivo and in vivo anti-tumour immune responses. Dexo loading the E720-57 peptide can efficiently activate CD8+ T cells against tissue culture-1 (TC-1) tumour cells, via promoting their proliferation and excretion of IFN-γ ex vivo (35). NK cells were also shown to enhance DC maturation and secretion of CD86, human leukocyte antigen-DR isotype (HLA-DR), and IL-12p70 in the presence of HPV virus (36). The DCs further activated NK cells by up-regulating cell surface markers (CD69 and HLA-DR), secreting IFN-γ and activating cytotoxicity toward an HPV-positive cell line. Another study showed that monocyte-derived DCs that pulsed with recombinant Mycobacterium tuberculosis heat shock protein 70-formyl peptide receptor 1 (MtHSP70-FPR1) expressed high levels of HLA-DR, CCR7, CD80, CD83, and CD86, and secreted more TNF-α, IL-12p70, and IL-1β to induce stronger CTL activation and proliferation, thus resulting an effective killing effect of cervical cancer cells (37).

**Cytokines**

Cytokines are soluble low molecular weight proteins which are produced by many immune-related cells under the stimulation of antigens and exhibit immune regulatory functions, including ILs, IFNs, TNFs, etc. Research indicates that, with the progression of CIN lesions in the uterine cervical increased, the expression of IL-1β and IL-8 is enhanced, meaning that the Th2 inflammatory response is strong. Meanwhile, a decrease in the macrophage inflammatory protein-1beta (MIP-1β) level may be helpful for the immune escape of HPV (38). The levels of IL-10, IL-17, IL-23, and transforming growth factor-beta (TGF-β) were more highly expressed in the peripheral blood of patients infected with high-risk cervical HPV than in those infected with low-risk cervical HPV (P<0.05) (39). The logistic regression analysis indicated that, after adjusting for other variables (age, cervical cancer history, and smoking history), the single nucleotide polymorphism of three Th2 cytokines (IL-4, IL-6, and IL-10) and one Th3 cytokine (TGF-β1) were found to have a significant association with cervical cancer (40). Tumour necrosis factor receptor 2 (TNFR2)+ T reg cells and relevant cytokines have been confirmed to be associated with cervical cancer development. Both the peripheral and tumour-infiltrated TNFR2+ T reg cells were found to be significantly higher in patients with CIN or cervical cancer, and the percentage of peripheral TNFR2+ T reg cells was negatively correlated with the clinical stages of cervical cancer (41). In HPV-positive head and neck squamous cell carcinoma, the defection of TNF receptor-associated factor 3 (TRAF3) activated an alternative nuclear factor-kappa B signalling, which suppressed the expression of anti-viral cytokine and promoted cancer cell survival and resistance (42). A higher serum TNF-α level was also shown to contribute to a lower clearance of oral infected HPV in men than in women (43). HPV possibly interferes with the trafficking of leukocyte by synthesizing and releasing specific pro-inflammatory cytokines and chemokines to change the
tumour micro-environment, which may be helpful for
tumour development (44).

Co-stimulatory molecules

Co-stimulatory signals are important for T-cell activation.
Although CD8+ T cells induced by APCs lacking co-
stimulatory molecules were found to retain their cytotoxic
activity, the lack of cloning and proliferation ability was
observed and their abilities on adaptive immune response
and IL-2 secretion were inhibited (45). Previous studies
showed that HPV could escape from the host immunity
system. The HPV-mediated suppression of co-stimulatory
signals by Langerhans cells (LCs) may be responsible
for persistent HPV infection and immune escape (46).
When the proper stimulus was given to women with HPV
persistent infection history, then the LCs presented HPV
antigens and induced an adaptive T-cell immune response.
For instance, stabilized polyinosinic: polycytidylic acid
[s-Poly(I:C)] compounds are indicated to be a promising
immune-modulator for the clearance of persistent HPV
infection (47).

The B7 family consists of the following important co-
stimulatory molecules: B7-H1, B7-H2 (ICOSL), B7-1
(CD80), B7-2 (CD86), B7-H3 (CD276), programmed
death-ligand 1 (PD-L1), B7-H4 (B7X), and B7-DC (PD-
L2). The interactions between B7 molecules and CD28
family receptors are crucial for adaptive cellular immunity
regulation. In cancer, the aberrant expression of co-
inhibitory B7 was demonstrated to reduce anti-tumour
response and was attributed to cancer immune escape.
In addition, cytotoxic T-lymphocyte-associated protein-4
(CTLA-4) could reduce the level of B7 ligands and inhibit
T-cell response, mainly via inhibiting IL-2 and blocking cell
cycle progression (48).

Co-stimulatory receptors on the T cell surface (e.g.,
CD28) could bind with the co-stimulatory molecules
on the APC surface, including CD80 (B7-1) and CD86
(B7-2). The combination could then trigger the survival
and proliferation of activated T cells (signal 2), which is
important for immune tolerance and T-cell responsiveness
regulation (49). B7-H3 co-stimulates the proliferation of
both CD4+ and CD8+ T cells, which enhances the induction
of CTLs and stimulates IFN-γ production via T-cell
receptor signalling (50). In addition, B7 molecules may
promote the proliferation, invasion and migration of cancer
cells (51). The expression level of B7-H3 is significantly
higher in cervical cancer tissues than in normal tissues,
indicating a worse prognosis for patients (52). The HPV
E6 and E7 oncoproteins have strong antigenicity. The
introduction of B7 molecules into cervical cancer cells could
enhance the immunogenicity of the cells via synergistic
stimulation and the down-regulation of immuno-
suppression, thus inhibiting their proliferation and invasion.
They may therefore be a potential target for cervical cancer
therapy.

Cervical cancer immunotherapy

At present, cervical cancer can be prevented by HPV
vaccine injection and regular cervical cancer screening,
but protective vaccines cannot eliminate an existing HPV
infection and cannot prevent the development of cervical
precancerous lesions or cervical cancer. Therefore, it is
urgent to find effective counter measures for existing HPV
infections and related diseases.

Adoptive T-cell therapy

The adoptive transfer of tumour-specific T cells, which
are obtained from tumour-infiltrating lymphocytes or
peripheral blood, is a promising immunotherapeutic
approach for cancer. In a clinical trial, 9 patients, who
were diagnosed with metastatic cervical cancer, were
given a signal infusion of tumour-infiltrated T cells that
were HPV E6 and E7 reactive after finishing platinum-
based chemotherapy or chemoradiotherapy; the tumour
completely regressed in 2 patients (53). After treatment,
2 complete responses were ongoing at 22 and 15 months,
respectively, and 1 partial response was received at 3 months
into the trial (53).

The tumour-draining lymph nodes of patients with HPV-
induced cervical cancer can be used as a source for adoptive
cell transfer. One study demonstrated that the stimulation
of lymph node mononuclear cells from the tumour-draining
lymph nodes resulted in a 36-fold increment of polyclonal
HPV-specific T cells under a clinical grade condition for
adoptive immunotherapy in patients with cervical cancer (54).

Inhibitors of immune checkpoints

In normal conditions, immune checkpoints are crucial
for the maintenance of self-tolerance, which prevents
autoimmunity and normal cells from being attacked by the
immune system. Unfortunately, cancer cells can utilize this
function to escape from immunization surveillance. Drugs
that block the immune checkpoints (viz. the programmed death protein-1 (PD-1)/PD-L1 and CTLA-4 pathway) have shown promising results for solid and hematological malignancies by significantly improving the survival of newly diagnosed and heavily pre-treated patients (55).

In 2014, the US Food and Drug Administration (FDA) approved a humanized antibody against PD-1 (Pembrolizumab, Merck & Co. Inc., White house Station, NJ, USA) and a human anti-PD-1 antibody (Nivolumab, Bristol-Myers Squibb, New York, NY, USA) to treat advanced melanoma (56,57). Recently, a complex method consisting of anti-PD-1 antibody (CT-011), low-dose cyclophosphamide depletion of T reg cell, and the HPV 16 E7 peptide vaccine, was shown to activate antigen-specific immune responses and induce the complete regression of tumours to prolong the survival time of a significant percentage of treated animals (58). Another study showed that tumours may form an “immune-privileged” region for adaptive immunity resistance via the PD-1/PD-L1 pathway, which may reasonably explain the therapeutic hypo-responsiveness of this pathway in HPV-positive oropharyngeal squamous cell carcinoma patients (59).

New methods are emerging in cancer immunotherapy beyond CTLA-4 and PD-1. PD-L1 and lymphocyte-activation gene-3 (LAG-3) inhibitors have made progress in clinical trials. B7 family members and the TNFR superfamily members have also shown promise or variable success in preclinical models and clinical studies, respectively (60).

**Therapeutic HPV vaccines**

Recently, various therapeutic HPV vaccines are being studied with the aim to activate an HPV-associated T-cell immune response and eliminate HPV-related diseases and cancer. The HPV E6 and E7 tumour antigens are expressed only on cervical cancer cells surface, but not in normal tissues (61), which make them ideal targets for therapeutic vaccines that do not damage normal tissue.

**Live vaccines**

Live vaccines can activate strong cellular and humoral immune response (62), and can be classified into bacterial- and viral-based vectors, for which *Lactobacillus casei* and *Listeria monocytogenes* have been widely used to develop vaccines for HPV infection therapy. In a study which was the first to report a correlation between cell-mediated immune response specific to E7 in the cervix after immunotherapy for human mucosal neoplasia, oral administration of a *Lactobacillus*-based vaccine expressing E7 activated a specific mucosal immunity to E7 in uterine cervical lesions. About 70% of the patients experienced a pathological downgrade from CIN3 to CIN2 after taking the optimized dose (63).

The immunotherapy ADXS11-001 is a live attenuated *Listeria monocytogenes* cancer vaccine that secretes a fusion product of the HPV16E7 antigen and listeriolysin O protein-targeting HPV-transformed cells. A phase I clinical trial showed that patients with persistent/recurrent metastatic cervical cancer who received a high ADXS11-001 tolerated it well, and the most common treatment-related adverse events (TRAEs) that occurred in ≥3 patients were chills, vomiting, hypotension, tachycardia, fever, and nausea (64). Only one grade 3 TRAE (hypotension) and no grade 4 or grade 5 TRAEs were reported in the study (64). In a phase II clinical trial, 69% of patients (18/26) who received all three per-protocol doses of ADXS11-001 had a median survival duration of >1 year [12.1 (95% CI: 6.8–NR) months], and the 12-month overall survival rate was 55.6%. The Stage 2 trail is currently recruiting patients and has been amended to allow for continuous cycles of therapy (65).

Viral vectors enter host cells with high efficiency and can activate the T-cell responses. For safety reasons, viral vectors are limited to use in patients who are immunocompromised. In a first stage clinical trial, a recombinant vaccinia virus was administrated to 8 patients with advanced stage of cervical cancer (66). The result showed that this kind of vaccination generated only a mild and tolerable response in patients with single-dose inoculation. Another clinical study that was performed in 29 patients with stage IB or IIA cancer with 2-time injection of TA-HPV vaccine showed that >28% of the patients exhibited specific serological responses, but the CTL response was transient (67).

**Peptide vaccines**

Compared to live virus vaccines, peptide vaccines are easier to produce, more stable, and safer. However, single epitope-based peptide vaccines tend to generate a narrow and weak immune response. Researchers proposed using overlapping peptides or mixed overlapping long peptides as vaccines to remedy this weakness (68). One study showed that the HPV16 E-and-E7-overlapping long peptide-based vaccine was well-tolerated and capable of inducing a broad IFN-γ-associated T-cell response even in patients with advanced cervical cancer (69). Other clinical research has demonstrated that a mixed overlapping long peptide vaccine
could generate a complete or partial response in 47% and 80% of the participants, respectively (70). A recent clinical trial study showed that the combination of an overlapping long peptide vaccine with immune-check point inhibitor was more efficient than the use of the check point inhibitor alone (71).

### Protein vaccines

TA-CIN, a 725-amino-acid-based protein vaccine, has been shown to generate HPV16-specific antibodies and CTL responses in TC-1 tumour-bearing mice. In a phase I clinical trial, though no adverse events were observed, the HPV16 E6- and E7- specific responses were low (72). In another clinical study, a group of CIN 3 patients received a loop electrosurgical excision procedure or cone biopsy, with 78% of them presenting complete or partial responses, and two patients (3.5%) progressing to microinvasive cancer after they were subcutaneously injected with SGN-00101. No other drug-related adverse effects were reported (73).

### Recombinant overlapping peptide vaccines

Mixed synthetic overlapping peptides or long peptides vaccines have many advantages, but challenges in manufacture quality control and the drug registration have arisen. Although protein-based vaccines have less of the above-mentioned disadvantages, they only stimulate CD4 T and B cell responses and not the CD8 T cell response.

Considering the disadvantages of synthetic overlapping peptides and protein-based vaccines, the recombinant overlapping peptide was proposed as a new category of vaccine. To put briefly, the overlapping peptide sequences of a target protein are linked to an intracellular protease. The linked sequences are then expressed as a recombinant protein (also called recombinant overlapping peptides, ROP). ROPs have been tested as vaccines successfully in proof-of-concept in vitro and in vivo studies (74). The ROP vaccine can be produced as one entity to avoid the multiple disadvantages in quality control and registration. Moreover, ROP vaccines can be cleaved into overlapping peptides to stimulate both CD4 and CD8 T cell responses once they are ingested by APCs (74).

### DNA vaccines

The DNA vaccine has emerged as a promising immunotherapeutic agent against cancers by virtue of its stability, simplicity and safety. A number of clinical trials have demonstrated that the DNA vaccines is well-tolerated by patients and does not trigger significant adverse effects (75).

More recently, sequential pNGVLa-Sig/E7(detox)-HSP70, with a recombinant vaccine virus encoding an immunomodulatory E6/E7 fusion protein, was tested in 12 patients with CIN3 for the phase I trial, resulting in 5 of the subjects showing complete tumour regression (76). A phase I–II test of the VGX-3100 vaccine (a mixture of two plasmids encoding HPV16/18 E6/E7 antigens) showed that 49.5% (53/107) of the patients treated with CIN2/3 who were administered by electroporation showed complete tumour regression. VGX-3100, by inducing both T-cell and B-cell responses to E6 and E7, was proven to be the first effective therapeutic vaccine against HPV 16- and HPV18-associated CIN2/3 (77).

Despite these advantages, DNA vaccines for cancers still entail a few disadvantages, including the relatively low immunogenicity, which have impeded the expected success of clinical application. Naked DNA is not easily spread between cells in vivo, and APCs do readily take up and present antigens to start effective immune responses. Thus, effective strategies need to be explored to enhance the potency of DNA vaccines.

### DC-based vaccines

DCs possess strong abilities in activating and controlling the T-cell response, which makes them ideal candidates for immunotherapy strategies. A previous study showed that DCs stimulated with E6/E7 RNA could activate antigen-specific CTL to recognise and lyse DCs that were transfected with E6 or E7 RNA and cervical carcinoma cells expressing E6/E7 proteins. The efficiency was similar to that which the researchers achieved in DCs that were activated with E6 and E7 peptides. Moreover, DCs co-transfected with both E6/E7 RNA showed more efficiency in stimulating CTLs to lyse human cervical cancer cells (78).

A modified HPV16 E7-pulsed DC vaccine enhanced by an adenovirus-mediated suppressor of cytokine signalling 1 (SOCS1) reduced the transformation activity and improved its antigenicity, thus receiving a better immunotherapeutic effect in allografted tumour mouse models (79).

Collectively, Dexo loaded with the E749–57 peptide could effectively induce CD8+ T cell proliferation and IFN-γ excretion to exert cytotoxic activity toward TC-1 tumour cells ex vivo. Moreover, the Dexo vaccine promoted the re-stimulation responses of E7 antigen vaccinated mouse splenocytes in vitro. Notably, Poly (I:C) dramatically increased the potent anti-tumour ability of antigen-pulsed Dexo in cervical cancer amelioration (35). Although various
cellular immunotherapies have made substantial progress in the treatment of HPV-related diseases, further research needs to be performed and large-scale clinical data need to be collected to confirm their efficacy.

Conclusions

Cell immunity is closely related to the development of cervical cancer. APCs, co-stimulatory molecules, cytokines, and HPV subtypes are also involved in the immune response of cervical cancer. CD8^+ T cells and the ratio of CD4^+/CD8^+ may influence the immune function of patients. A review of the relevant studies was provided above with the of aim creating a platform for future research studies to derive effective treatments from the various specific T-cell immune responses, and to use the effective immune response to control tumour development and improve therapy.

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Footnote

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