Review Article

Regulation of Inflammatory Pathways in Cancer and Infectious Disease of the Cervix

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Cervical cancer is one of the leading gynaecological malignancies worldwide. It is an infectious disease of the cervix, associated with human papillomavirus infection (HPV), infection with bacterial agents such as Chlamydia trachomatis and Neisseria gonorrhoea as well as human immunodeficiency virus (HIV). Furthermore, it is an AIDS-defining disease with an accelerated mortality in HIV-infected women with cervical cancer. With the introduction of robust vaccination strategies against HPV in the developed world, it is anticipated that the incidence of cervical cancer will decrease in the coming years. However, vaccination has limited benefit for women already infected with high-risk HPV, and alternative therapeutic intervention strategies are needed for these women. Many pathological disorders, including cervical cancer, are characterised by the exacerbated activation and maintenance of inflammatory pathways which are considered to be regulated by infectious agents. In cervical cancer, hyperactivation of these inflammatory pathways and regulation of immune infiltrate into tissues can potentially play a role not only in tumorigenesis but also in HIV infection. In this paper we will discuss the contribution of inflammatory pathways to cervical cancer progression and HIV infection and the role of HIV in cervical cancer progression.

1. Introduction

Cervical cancer is the most common gynaecological cancer among women in developing countries [1, 2]. Virtually all cases of cervical cancer follow after infection of the cervical epithelium with oncogenic human papilloma virus (HPV) types [3]. Currently, there are over 150 genotypes of HPV [4]. These are species-specific and tissue-tropic and only infect cutaneous or internal squamous mucosal surfaces in humans [4, 5]. Around 40 types are known to infect the anogenital tract, giving rise to genital warts, condylomata or cancers, and their precursor lesions [4]. The majority of anogenital cancers in humans are associated with the high-risk HPV16 and 18 and there is correlation between percentage of HPV 16 and 18 integration and severity of the cervical lesions [6]. Although it is necessary to have infection of the cervix with oncogenic HPV to develop cancer, HPV itself may not be sufficient. Other associated cofactors including compromised immune system or infections with herpes virus II [7], Chlamydia trachomatis [8], Neisseria gonorrhoeae [9], or bacterial vaginosis [10] have been associated with cervical inflammation and increased risk of cervical cancer. In 1993, cervical cancer was classified as an AIDS-defining disease, together with Kaposi Sarcoma and Non-Hodgkin Lymphoma, in women infected with human immunodeficiency virus (HIV) [11]. This highlighted HIV as a potent cofactor for developing invasive carcinoma of the cervix and highlighted cervical cancer as an infectious disease. Although HPV infection is very common among young sexually active women, only a small percentage of women below the age of 25 years actually develop cervical cancer. In fact, the median age recently reported for women presenting with invasive cervical cancer is around 50 years of age [12, 13]. These observations highlight the long latency of the virus and the need for persistent infection of the cervix to promote disease.

HPV enters the body and infects basal keratinocytes, exposed through mild abrasion or microtrauma to the cervico-vaginal epithelium (Figure 1). The main route of HPV transmission is via exposure of the cervix to virus present in saliva or seminal fluid or in the effected partner's
HIV infects cells and the tat protein causes the amplification of the HPV E1/L1 genes leading to increased HPV replication and release of HPV virions. These then infect the same or adjacent cervical epithelial cell. Within the epithelial cells, the HPV E6 oncoprotein binds to p53 protein targeting it for ubiquitin-dependent degradation while the E7 binds to the Rb protein thus disrupting the Rb and E2F complex and can increase nitric oxide (NO) production, DNA damage and the activation of the COX-2/PG/PG receptor inflammatory pathways leading to increased inflammation and tumorigenesis. Inflammatory and tumor cells can then release cytokines, chemokines and PG which act in autocrine/paracrine to regulate the endothelial, stromal, neoplastic epithelial and infiltrating immune cells function to cause an increased tumour angiogenesis, increased tumour growth, decreased apoptosis, and decreased local immune-surveillance. These conditions favour tumorigenesis and viral survival in the tumor microenvironment. Inhibition of the COX-PG cascade with nonsteroidal anti-inflammatory drugs (NSAIDs) like aspirin could reduce inflammation and tumour progression by inhibiting downstream pathways activated by PG and promote proresolusion by elevating expression of resolution lipid mediators such as Resolvins.

Inflammation involves a coordinate effort by the body to maintain homeostasis in the face of insult, by bacterial or viral pathogen, or by injury. The process of inflammation is highly complex involving a host of resident and recruited cell types, which work together to promote the removal of the insult or injury and initiate the repair of the tissue (reviewed in [18]). When successful, this results in the restoration of tissue homeostasis, a process termed resolution (reviewed in [19–21]).

2. Inflammatory Pathways

2.1. Inflammatory Cell Component of Tumours. The inflammatory infiltrate of tumours can comprise a vast population of different immune cells at any one time—each capable of producing cytokines or other factors which can alter the fate of immune populations within the tumour. The inflammatory environment in tumours is characterised by the presence of leukocytes. Leukocytes are recruited into the tissue in response to cytokines and chemokines released by tumour cells and are resident both in the supporting stroma as well as the tumour itself [22]. Polymorphonuclear leukocytes (PMN’s or neutrophils) are the first immune cell types to be recruited to sites of inflammation, followed by monocytes which are derived from haemopoietic progenitor cells [23]. When in the tissue, monocytes then differentiate into either macrophages or dendritic cells (called Langerhans cells in the epidermis). Langerhans cells generally constitute the first defence against pathogens. Recently, tumour associated neutrophils have been described that are capable of polarising the phenotype of other immune cells and altering the cellular composition of the tumour microenvironment [24]. Here, PMNs are thought to exist in either an N1 state, capable of killing tumour cells by producing and releasing cytotoxic compounds, or an N2 state capable of promoting tumour
growth by modulating the cytokine/chemokine environment in the tumour [24]. A significant proportion of the immune infiltrate in tumours is comprised of tumour-associated macrophages (TAMs). These are derived from circulating monocyte precursors via the release of monocyte chemotactic protein (MCP) chemokines [17]. Although TAMs are a heterogeneous cell population, early stage tumours are thought to have type 1 macrophages (M1). These produce proinflammatory cytokines and chemokines, such as CXCL10 and CXCL19 to recruit Th1, Th17, and natural killer (NK) cells [25]. In more advanced tumours, TAMs polarise towards a type 2 (M2) state to encourage Th2 differentiation and the production of potent angiogenic factors such as VEGF to facilitate tissue remodelling and tumorigenesis [26].

2.2. Cyclooxygenase-Prostaglandin Pathway. The inflammatory cyclooxygenase-(COX-) prostaglandin (PG) axis is a central pathway regulating inflammation and cancer (reviewed in detail in [27, 28]). There are two COX enzyme isoforms in humans, namely, COX-1 and COX-2 [29]. The role of COX enzymes in inflammation has been underscored by studies involving the use of nonsteroidal anti-inflammatory drugs (NSAIDs), which function by blocking COX-derived lipid mediators, namely, prostaglandins, prostacyclins and thromboxanes [30]. In vitro and in vivo studies have highlighted a role of COX-derived lipid mediators in driving tissue remodelling events associated with inflammation. More recently, NSAIDs have been shown to induce active proresolution pathways to facilitate healing, by acetylating COX-2 enzyme and altering its enzymatic capacity to produce classes of proresolution lipid mediators such as Resolvin in humans, namely, COX-1 and COX-2 [29]. The result is uncontrolled replication and a buildup of damaged DNA—ideal conditions for cancer development.

Animal studies using transgenic mouse models of cervical cancer have highlighted more specifically the contribution of individual oncogenes to tumorigenesis and have shown that the E7 protein induces high-grade cervical dysplasia and invasive tumours, whereas the E6 protein only induces low-grade cervical dysplasia. Furthermore, they have highlighted that both oncogenes work in synergy to promote cervical cancer progression since coexpression of E6 and E7 proteins produce larger and more extensive tumours compared with E7 alone [39].

3. HPV Oncogenes: Structure and Role in Inflammation and Tumorigenesis

The HPV genome consists of 3 domains, a non-coding upstream regulatory region, an early region containing open reading frames E6, E7, E1, E2, E4, and E5 and a late region encoding the major (L1) and minor (L2) capsid proteins [5]. Once infected, the E6 and E7 early genes are actively transcribed and the E6 and E7 oncoproteins, both of which are involved in neoplastic transformation, are actively expressed [35]. Although E6 and E7 oncogenes appear to be the main HPV genes involved in transformation, recent studies have highlighted a role for E5 oncogene in immune cell modulation and tumorigenesis [36] and regulation of late viral functions together with E4 oncogene. E1 and E2 oncogenes encode replication factors and are thought to play a role in HPV persistence by allowing episomal copies of the virus to be maintained in the nucleus and partitioned into daughter cells during mitosis [37]. Many of the observations on the roles of HPV oncogenes have been derived using in vitro model systems of HPV-containing cell lines where specific oncogenes have been ablated by RNA interference, or HPV-negative cells that have been transfected with cDNA constructs containing specific HPV oncogenes. These studies have shown that both oncoproteins target different molecular pathways in the cell. E6 oncoprotein binds to P53 protein and targets it for ubiquitin-dependent degradation, whereas E7 binds to family members of the retinoblastoma protein (Rb) and disrupts the complex between Rb and the E2F transcription factor family (Figure 1) [38]. This facilitates immortalisation of cells and triggers the early steps in malignant conversion by interfering with cell cycle control [5, 38]. The result is uncontrolled replication and a buildup of damaged DNA—ideal conditions for cancer development.
and PTGER4 (Figure 1) [36, 40, 41]. Many chronic inflammatory diseases, allergy, asthma, atherosclerosis, autoimmunity, transplant rejection, metabolic, and degenerative diseases and cancer [42] are all associated with upregulation in COX enzyme expression. In the past decade, several studies have emerged from in vitro and in vivo model systems employing cell lines and rodents to demonstrate that prostaglandins, produced as a consequence of elevated COX enzyme expression, can promote extensive tissue remodelling within tumours by evoking all the classical hallmarks of cancer, namely, cellular proliferation, angiogenesis, inhibition of apoptosis, and alteration in vascular permeability to allow immune cell extravasation from the vasculature [18, 43]. The studies of Subbaramaiah and Dannenberg and Oh and colleagues, described above, for the first time demonstrate that these hallmarks of cancer can be driven by HPV oncogenes [36, 40, 41] and provide a direct link between HPV oncogenes and activation of potent inflammatory cascades, with known roles in promoting chronic inflammation and cancer.

3.2. Inflammatory Pathways Regulated by HPV Oncogenes and COX Enzymes. Inflammation involves extensive tissue remodelling events which are orchestrated by complex networks of cytokines, chemokines, and bioactive lipids. These work across multiple cellular compartments to elicit their function. The classic hallmarks of inflammation are the recruitment of immune cells into the tissue and alteration of vascular function to allow for immune cell extravasation. This alteration in permeability causes oedema and the redness generally associated with tissue inflammation. Several reports have correlated inflammatory cell infiltrate with HPV-induced high-grade lesions. Infiltrating lymphocytes are thought to contribute to tumour growth and spread as well as immunosuppression, generally associated with malignant diseases [14]. Although the precise mechanism whereby HPV oncogenes regulate tissue remodelling events is unclear, HPV infections have been shown to promote the release of inflammatory mediators and cytokines from keratinocytes to alter the immune response and promote the infiltration of macrophages, lymphocytes and NK cells (reviewed in [44]). Changes in the vasculature to facilitate immune extravasation and angiogenesis require tissue remodelling of the extracellular matrix, a process facilitated by matrix metalloproteinase (MMP) [13]. Several studies have correlated HPV E6 and E7 transcription with MMP transcription [45, 46] and genes in cervical epithelial cells involved in tissue differentiation and remodelling [47]. In addition, transfection studies have shown that E7 oncoprotein forms a complex with and downregulates leukocyte elastase inhibitor [48]. This would facilitate the activation of neutrophils and promote neutrophil influx into the tissue. Furthermore, transgenic mouse models where the early region genes from HPV16 are expressed under the control of the human keratin 14 promoter have shown that macrophage recruitment to HPV-associated lesions occurs via the release of the chemokine CCL2 and interaction with its receptor CCR2 present on macrophages [49].

Once resident in the tissue, inflammatory cells are known to produce vast amounts of reactive oxygen species and nitric oxide which have been shown to induce DNA damage (Figure 1) [50] and to contribute towards the progression of the disease in high-grade cervical lesions [50]. Furthermore, nitric oxide has been shown to induce transcription of E6 and E7 oncogenes in cervical epithelial cells [51], which can further enhance inflammation via an autoamplifying positive feedback loop via activation of COX-2 and other parallel inflammatory pathways. Taken together, these studies provide compelling evidence for a role for HPV oncogenes in regulation of inflammation in cervical cancers. It is tempting to speculate that targeted inhibition of E6 and E7 actions in cervical epithelial cells could be a potential therapeutic alternative to vaccination for women infected with HPV.

4. HIV, Cervical Cancer, and Inflammation

The human immunodeficiency virus (HIV); the only etiological factor attributed to the acquired immunodeficiency syndrome (AIDS) belongs to the genus Lentivirus within the family Retroviridae [52]. About 33 million people harbour this virus worldwide [53] with high epidemic rates in sub-Saharan Africa. Mature HIV virion is spherical (approximate diameter of 100–120 nm), with a genome of two copies of identical (9.2kb) single-stranded RNA [54]. HIV is divided into two main subtypes HIV-1 and HIV-2. HIV-1 is further subtyped into phylogenetically related clades: types A-K, with subtype C being the most common type in Africa [55]. Transmission is via unprotected sexual intercourse, intravenous drug use, blood transfusion, infection with blood-derived products, or mother-to-fetal transmission. To initiate infection, virus attaches to cellular surfaces via an interaction between the gp120 viral envelope protein and a receptor complex present on the host cell consisting of the CD4 receptor and G protein-coupled receptor (GPCR) coreceptor, usually CCR5 or CXCR4 [54]. Most primary HIV-1 variants are restricted to the use of CCR5 and CXCR4 [56, 57], however, they have been shown to use alternative receptors in vitro. HIV-2 variants are capable of infecting a wider range of cells expressing different coreceptors such as GPR15 and CXCR6 in addition to CCR5 and CXCR4 [58]. This leads to fusion between the viral and cellular membranes and ultimate release of the viral core into the cell cytoplasm [54]. Once inside the host cell the virus is reversed transcribed to full-length double-stranded DNA by the reverse transcriptase enzyme and is integrated into the host genome [59]. The hallmark of infection is characterised by progressive depletion of CD4+ T-cells leading to an immunodeficiency state, paving the way for opportunistic infection and ultimately mortality [60].

4.1. The Interplay between HIV and HPV and Their Role in Cervical Cancer. The interplay between HPV and HIV is complex; however, their synergistic role in exacerbating pathology of the cervix has been well documented. For example, epidemiological studies have shown that women that are coinfected by HPV and HIV have an estimated 41 fold increase in the risk of developing neoplastic cervical lesions [61] and HIV infected immune-compromised women have been shown to have a higher prevalence of HPV-induced
lesions [62]. Furthermore, studies in Sub-Saharan women, where 67% of the population are living with HIV/AIDS, have shown that women with HIV develop cervical cancer at an earlier age than women who are HIV negative [63–65].

Although the precise mechanisms predisposing women infected with HPV to HIV infection are unclear, there is evidence that clearance of HPV infection from the female genital tract elicits a cell-mediated immune response characterised by gross infiltration of lymphocytes and macrophages into the epithelium [66], which can enhance the risk of HIV infecting immune cells in the cervix in these women after unprotected sexual contact.

Central to the role of HIV in cervical cancer is its ability to ablate the systemic immune response to infection, including HPV infection. This can facilitate inadequate clearance of HPV in infected individuals, enhancing HPV persistence or reinfection, and it increases the likelihood that precancerous lesions will develop into cancer. To this end, HIV may modify HPV-related carcinogenesis by altering the expression of inflammatory components (cytokines) in the cervix and diminution of local cervical cellular immunity, thus altering HPV regulation [53]. For example, HIV tat gene has been shown to increase the expression of HPV E1 and L1 genes, hence causing upregulation of the HPV replication (Figure 1) [67,68]. Furthermore, HIV-1 tat protein is capable of transactivating HPV16 transcription [69].

4.2. The Role of HIV in Regulating Inflammation and Its Potential Contribution to Cervical Cancer. Systemic expression of several proinflammatory cytokines has been reported to be a major feature in HIV infection. The virus causes immune dysregulation leading to an increase in the production of proinflammatory cytokines such as TNFα, IL-1, and IL-6 which are detected in the plasma and lymph node of infected patient [70]. This link between HIV and production of proinflammatory cytokine was suggested by the observation that the virus and/or its surface glycoprotein gp120 can induce in vitro secretion of TNF, IL-1, and IL-6 by monocytes isolated from uninfected individuals [71–73]. Supplementary studies also detected high levels of IL-1α, IL-1β, IL-6, and TNF in the serum and cerebrospinal fluid of seropositive individuals [74, 75]. Often associated with the production of these cytokines is the elevated secretion of CC-chemokines such as macrophage inflammatory protein (MIP)-1α, MIP-1β, and RANTES [76, 77]. These proinflammatory cytokines expressed as soluble factors or membrane binding molecules and are directly or indirectly involved in HIV entry and T cell apoptosis [60]. These cytokines have been found to be abundant in microenvironment of several tumours including cervical cancer where they are secreted by the tumour cells, endothelial cells, and/or infiltrating activated immune cells where they act as endogenous tumour promoter by stimulating the production of transcription factors (e.g., NF-κB, AP-1), proliferative and angiogenic proteins (e.g., VEGF, MMP), and adhesion molecules (e.g., E-selectin, VCAM), thus enhancing tumour growth and mediating tumour metastasis [78]. TNF, a major mediator of inflammation, can be detected in various human neoplasias where it is implicated in the induction of MCP-1 which can modulate the infiltration of immune cells in to the tumour microenvironment. It is, therefore, feasible that the elevated level of these proinflammatory mediators in HIV infected individuals can drive tumour progression in HIV-positive women with cervical dysplasia.

In addition to regulation of immune cell infiltrate, alteration in the expression profile of HIV receptors on cells within the cervico-vaginal region could impact on HIV acquisition and cervical cancer progression. The epithelial surface of the female reproductive tract expresses all the receptors necessary for HIV infection including CD4, CCR5, and CXCR4 [79]. Maher and colleagues have recently shown that HIV virions can bind the external surface of cervical epithelium and penetrate beneath the epithelial surface [80]. Thus, it is plausible that epithelial cells lining the cervico-vaginal interface could be the first cells to come into contact with HIV and might play a role in the replication of the virus and transmission to leukocytes present in the submucosa.

Expression of some of the HIV receptors on uterine epithelial cells display a temporal variation in expression during the menstrual cycle, indicating that they are hormonally regulated [79]. This could alter susceptibility to infection depending on the phase of the menstrual cycle. Chemokine receptors such as CXCR4 are elevated in cervical cancer and play a role in lymph node metastasis during advanced-stage disease [81]. These receptors can also be hijacked by HIV for entry in such women. CXCR4 expression can be regulated by HPV oncogenes [82] and prostaglandins [83] in the female genital tract. These observations suggest that HPV infection and inflammation can drive expression of HIV coreceptors on epithelial cells. It is unknown whether the inflammatory milieu of cervical cancer can also alter expression of HIV receptors on immune cells in the tumour periphery. It is plausible that in women with HPV infection or localised cervico-vaginal inflammation, alterations in HIV receptor expression could allow more virus to bind the epithelium and elevate the amount of virus present locally in the genital tract following intercourse. In women with cervical cancers or inflammatory cell infiltrate into the cervix, this could enhance susceptibility to infection.

5. Concluding Remarks

It is clear that inflammation plays a critically important role in regulating pathology of cervix, susceptibility to infection by virus, like HPV and HIV and patient outcome. The grand challenge that lies ahead for the developing world, where a significant proportion of women live with either HPV or HIV infection or both, is how to manage inflammation in these women to prevent disease progression, especially progression of cervical cancer. Clinical trials have shown that long-term treatment with low-dose nonsteroidal anti-inflammatory drugs (NSAIDS) like aspirin can be beneficial for reducing the burden of colorectal cancer [84]. In resource poor countries, it is tempting to speculate that low-dose aspirin, which is affordable and widely available, could be of benefit to women with HPV infections and neoplastic lesions, by suppressing the inflammatory COX-PG axis, promoting resolution of inflammation, and preventing progression of cervical cancer.


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