The Influence of Psychological Stress on HPV Infection Manifestations and Carcinogenesis

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Key Words
Psychological stress • HPV infection • Carcinogenesis • Immune response • Psychoneuroimmunology

Abstract
Psychological stress is an important factor involved in disease manifestations of human papillomavirus (HPV) infection, and it can participate in HPV-associated carcinogenesis. The impact or effect which stress can have (exert) depends on a person’s genetic pool, experiences and behaviors. Due to inconsistencies in some study results, this issue remains a subject of research. Concerning the course of HPV manifestations, it has been observed that a higher number of life stressors in at least the previous 6 months, the absence of social support and the types of personal coping mechanisms employed, all influence HPV progression. In women with cervical dysplasia, a connection between greater stress experiences and dysregulation of specific immune responses has been observed. Once HPV enters a cell via the α6 integrin there are three possible sequences: latent infection, subclinical infection, and clinically manifest disease. HPV proliferation in differentiated epithelial cells induces morphologically cytopathic changes (koilocytosis, epidermal thickening, hyperplasia, hyperkeratosis). Oncogenic transformation requires the integration of the virus genome into the host genome. In doing so, DNA in the E1 region of E2 breaks down, leading to transcription disorders of E6 and E7. For the formation of irreversible malignancy, the following sequence is necessary: initial expression of E6 and E7 genes followed by suppression of apoptosis and the stable expression of E6 and E7 proteins that protect transformed cells from apoptosis. A successful immune response is characterized by a strong, local cell-mediated immune response. Several factors are important for the regression of HPV manifestation/infection, among which is psychological stress which can prolong the duration and severity of HPV disease. Stress hormones may reactivate latent...
tumor viruses, stimulate viral oncogene expression, and inhibit antiviral host responses. In the regression of HPV infection, increased activity of Th1 cells was observed. However, during psychosocial stress, a decrease in the Th1 type of immune response is seen, and there is a shift towards a Th2 response. Understanding perceived stress and biological changes in stress, as well as the evaluation of immune parameters, gives researchers a better picture of how stress influences HPV infections and how to improve disease management and outcomes.

Introduction

Psychological stress is an important factor involved in disease manifestations of HPV infection, and it can participate in carcinogenesis associated with HPV. Due to inconsistencies in some study results, this issue remains a subject of research. It is important to determine to what extent stress plays a role in HPV manifestations and carcinogenesis and how much it participates in the onset, development, and progression of infections.

Features of HPV

Human papillomavirus (HPV) is a DNA virus that belongs to the Papillomaviridae family. It is one of the most ubiquitous viral infections in humans, and it usually manifests as skin or genital mucosa lesions, although it can occur in other mucosa as well. It has long been known that most sexually active people will be infected by this virus at some point of their life, and the incidence of HPV infection is highest in the age group for those 20-40 years old [1-4]. Despite its high prevalence, most who get infected will not have a clinically overt infection. Still, the persistence of the HPV infection may cause a higher risk for developing cervical intraepithelial neoplasia (CIN) and invasive carcinoma. In women, a breakout from a high-risk HPV infection typically takes 14 months to clear up for an oncogenic infection and 5-6 months for a non-oncogenic infection [4].

To date, more than 200 types of HPV have been identified [5]. A strain is considered new when the nucleotides of the L1 part of the HPV genome differ from that of known HPV viruses by more than 10% [6, 7]. HPV viruses can be divided into different groups by their affinity for certain tissues, which in some part depends on their genotype [6]. The specific skin lesions for each type of HPV are as follows: common warts – types 2, 7, 22; plantar warts – types 1, 2, 4, 63; flat warts – types 3, 10, 28; and for verrucous cyst type 60 and epidermodysplasia verruciformis - more than 15 different types. There are also specific HPV types for anal/genital manifestations: anogenital warts—types 6, 11, 42, 44 and others; anal dysplasia (lesions) – types 16, 18, 31, 53, 58; and genital cancers— highest risk types 16, 18, 31, 45, other high-risk types— 33, 35, 39, 51, 52, 56, 58, 59, and probably high-risk— types 26, 53, 66, 68, 73, 82 [5]. The types of HPV for oral/oropharyngeal lesions are: focal epithelial hyperplasia (mouth)—types 13 and 32; mouth papillomas—types 6, 7, 11, 16, 32); and oropharyngeal cancer—type 16; laryngeal papillomatosis—types 6 and 11 [5]. Concerning oncogenic risk, the low-risk HPV types are 6, 11, 49, 42, 43, 44, 54, 61, 70, 72 and 81, while the high-risk types are 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 68, 73, and 82. In addition, 60% of all genital warts are caused by HPV types 16 and 11 [8].

HPV is transmitted through direct skin or mucous membrane contact, and infection can be clinical, subclinical or latent. Microimpairments in the skin or mucous membrane enhance transmission, but for infection to develop it is essential for the complete virus to be transmitted, not only its DNA fragments [4]. In the sexually active population, HPV is present in 80% of women of reproductive age, but its manifestations resolve-spontaneously in most cases (cca 80 % of infections) within 12-24 months [9]. Untreated HPV manifestations in men can cause invasive penile carcinoma, which represents 1% of malignancies in men in developed countries and 10-20% in undeveloped countries. In men, incidence of death due to suffering from HPV-associated cancers of different organs is 0.32 per 100,000 [10].
HPV is a double-stranded DNA virus that has an icosahedron shape and a double-layered capsid made of 72 capsomers (HPV's capsid is not covered by a lipid membrane, which makes it resistant to ethanol and solvents.) It has 8,000 base pairs with a molecular mass of 5200 kDa [8, 11]. The genes are divided into two groups: early (E) and late (L). The group of early genes is composed of six genes (E1, E2, E4, E5, E6, E7) that code for the proteins responsible for replication, transcription, and malignant transformation [5, 8, 11]. Late genes L1 and L2 code for the viral envelope (non-lipid membrane) and for the proteins responsible for the capsid's structure. Gene L1 is the oldest, and it is used to identify the different types of HPV. A 10% difference in the nucleotide sequence of the L1 gene signifies a new strain (type), while a difference of 2-10% is considered a subtype, and a difference of less than 2% is defined as a variant.

The oncogenic potential of HPV depends on genes E6 and E7. The product of the E6 gene binds to the p53 oncosuppressor, and the product of E7 binds to the RB protein. In the case of cervical intraepithelial neoplasia, genes of high-risk types are integrated into the DNA of the host. This integration in some cases leads to disruption of the E2 gene, which results in increased replication of the E6 and E7 genes. The E6 protein binds to p53 causing it to degrade. The E7 protein inactivates the RB protein so that E2F proteins detach from the RB, preventing transcription of the gene that regulates cell growth and differentiation [5, 11]. It has been established that E6 and E7 interfere with the immune response by reducing production of interferon (IFN) [5]. Therefore, the E1 and E2 genes of HPVs are involved in viral replication, while the E6 and E7 proteins function as the promoters of proliferation. The major HPV oncogenes are E6 and E7, which disrupt the normal regulation of the cell cycle and cell progression, giving them an important role in the oncogenesis of HPVs with a high risk of causing anogenital and cervical cancer. Immortalization of epithelial cells induced by HPV requires viral DNA integration into the host cell genome, which causes disruption of the E2 gene. The E2 protein is also a transcription factor, which regulates expression of the E6 and E7 oncoproteins. Integration of the virus in the human genome disrupts the E2 gene and increases expression of E6 and E7 genes in vitro [12-21].

Concerning molecular events during the progression of cervical lesions to carcinogenic lesions, persistent high-risk HPV infection leads to integration of HPV into the host genome and to overexpression of oncogenes E6 and E7 [11]. On the molecular level, interaction of E7 with the pRb protein leads to aberrant initiation of the S-phase. The E7 oncoprotein causes release of E2F transcription factor from the pRb protein, which is then active and can initiate transcription of genes involved in cell cycle progression, contributing to cellular immortalization and transformation. Thus, E6 targets p53 for proteasomal degradation, which leads to inhibition of apoptosis and DNA repair (anti-apoptotic effect). It is important to emphasize that only high-risk HPV types can induce degradation of p53, which can then lead to carcinogenesis. E6 activates the PI3K/Akt pathway, interacts with cellular proteins NFX1, and induces human telomerase reverse transcriptase (hTERT) activation, leading to immortalization and transformation. The interaction of both oncoproteins with DNA methyl transferases leads to aberrant methylation, causing silencing of tumor suppressor genes. Also, E7 interaction with histone deacetylases (HDACs) causes chromosome remodelling and genome instability. So, during lesional progression to carcinogenesis, the cross interaction of E6 and E7 with various pathways plays the crucial role [22]. It is also important to mention that viruses like HPV can create virions and become transmissible at any point in their life cycle (the productive virus replication also known as lytic replication). When lytic replication of the virus begins, it is almost irreversible, and successful replication of the virus begins as well as host cell death. But in tumor cells, these infections are mostly latent, allowing the virus to evade the immune response. Thus, lytic replication of the virus is reduced or absent in the tumor. In viral latency there is no production of unnecessary viral proteins that could initiate cell mediated immune recognition. Integration of the viral genome into the host genome eliminates the virus’s ability to replicate as virions, but the virus can replicate using the host's cellular mechanisms and can be divided whenever the host cell divides. By
evading apoptosis, as previously described, oncogenesis begins [23]. All this indicates that a complex network of actions is involved in the pathogenesis process during the occurrence of HPV manifestations.

**Stress, types of stressors, the hypothalamic-pituitary-adrenal (HPA) axis, the sympathetic autonomic nervous system (ANS) and the impact of stress on the body and disease**

Stress is defined as physical or mental exertion caused by factors that change homeostasis, and it can be observed as an objective stimulus, as an organism's response to stimulus, or as a relation between a person and their surrounding environment. Stress is a state of threat to physical, psychological, and social homeostasis. More recent approaches define stress as any stimulus that causes sudden termination of ordinary activities, that is, an event that goes beyond what is normal for the organism. The impact or effect, which stress can have, depends on a person's genetic pool, experiences, and behaviors [24-28]. The biopsychosocial model of disease and health, for example, asserts that biological, psychological and social factors interdependently affect the course and outcome of disease (Fig. 1). Psychological stress has become an increasingly important factor in the course of disease, due especially to the circumstances of modern life [25].

Stressors can be physical, chemical, psychological and biological. Psychological and social strains are the most common stressors [25]. Stress can be categorized by duration (acute or chronic), relevance (avoidable/averted or unavoidable), and intensity (mild, moderate, or severe). Aside from stress caused by common life events, there are also big traumatic events such as war, a natural disaster, death in the family, job loss, etc. Psych trauma is a state of high-level stress that can result in posttraumatic stress disorder (PTSD) and can cause long-lasting health problems. In physical, chemical, and biological stress, the condition of the person is determined by the harm caused by an external stimulus, whereas in psychological stress, one's assessment of environmental dangers, threats or challenges is the important factor. Stress in humans usually manifests through physical symptoms (e.g. palpitations, shortness of breath, perspiration, angina pectoris, frequent infections) or psychological symptoms (e.g. indecisiveness, poor concentration and memory loss, high sensitivity, sleep...
disturbances, negative thoughts). As a result, stress-related illnesses can occur (e.g. gastric ulcer, hypertension, viral infection, myocardial infarction, psoriasis, allergies, asthma, anxiety disorders, tumors, gastrointestinal disorders) [26-28].

When stressed, the organism reacts in a stress-adapting manner that can take place at the cell-, organ- or organ-system level, or at the level of the entire organism [25]. The organism typically reacts in three phases to a threat or injury with the same set of reactions outlined by the General Adaptation Syndrome (GAS): (1.) alarm; (2.) resistance; and (3.) exhaustion, the long-lasting debilitating phase that makes one susceptible to disease onset [29]. When reacting to external and internal demands by modulating functions and adapting to new conditions (referred to as alostasis), system stability can be achieved through constant adaptation. The main adaptive system includes the hypothalamic-pituitary-adrenal (HPA) axis, sympathetic autonomic nervous system, and cytokine production (Fig. 2) [19-22]. Pathogenetically, the body neutralizes stress with a complex network of physiologic and behavioral responses to reestablish optimal body equilibrium (eustasis) [30]. As crucial components of the stress response, the HPA axis and the autonomic nervous system (ANS) interact with other vital centers in the central nervous system (CNS) and peripheral tissues/organs to mobilize an adequate/appropriate adaptive response against stressors. Thus, different stressful events are recognized by the hypothalamic paraventricular nucleus, which participates in a biological circuit that integrates personal experiences, physiologic signalling and the release of corticotropin-releasing hormone (CRH) [31]. CRH acts on the pituitary gland, which then releases adrenocorticotropic hormone (ACTH), followed by ACTH signals to the adrenal cortex to release glucocorticoids [31]. Thus, the body’s adaptive stress response depends on many interconnected neuroendocrine, immune, cellular, and molecular mechanisms.

During stress, the brain and CNS are the main actors; their response includes a variety of crucial neuroendocrine and autonomic reactions in order to achieve homeostasis [25-28, 32-35]. During that process, neurogenic stressors activate processes in the CNS. Signals are then transferred to periventricular nuclei from prefrontal cortex and limbic structures, where stress is compared to experiential events. This processed signal is transferred to the hypothalamus, which in turn activates the HPA axis. Hypothalamic nuclei receive stimuli from limbic and

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**Fig. 2.** Stress-induced reactions and psychoneuroimmune network during HPV onset, manifestations and course
brain stem catecholaminergic signalling pathways. Periventricular nuclei can be activated by locus ceruleus aminergic signals. Central, medial and cortical amygdala nuclei are connected to periventricular and gabaergic neurons forming a closed circle. Activation of glutaminergic neurons stimulates the hypothalamic release of CRH in the eminentia mediana, which then reaches the hypophysis (anterior pituitary) through portal circulation and stimulates the release of ACTH into the peripheral circulation. Activation of the sympathetic nervous system causes terminal nerves and adrenal gland medulla to secrete more catecholamines [36, 37]. The immune system affects the brain as well; thus, there is a bilateral connection. It is important to emphasize that ACTH stimulates the release of glucocorticoids from the adrenal glands, meaning cortisol is a main stress hormone. Sympatho-adrenomedullary axis (SAM) activation stimulates CRH secretion in the hypothalamic periventricular neuron (PVN) area [25-28, 38]. Stress enhances the activities of many systems and releasing of various substances, including catecholamines, opiates and corticosteroids, which have an immunosuppressive effect by decreasing activity of cytokines and inflammation. It has been shown that immune cells have hormone receptors (corticosteroid, prolactin, growth hormone, sex hormones), neuropeptide receptors (endorphins, vasoactive intestinal peptide, substance P, etc.) and neurotransmitter receptors (adrenaline, noradrenaline, acetylcholine, serotonin, etc.). During acute stress, posterior hypothalamic nuclei, the sympathetic nervous system and adrenal gland medulla are activated, while in chronic stress, the anterior hypothalamus, sympathetic system and adrenal gland cortex are activated.

Finally, the release of glucocorticoids, mainly cortisol, is followed by increased lipolysis and gluconeogenesis to supply the body with available energy sources. A negative feedback system regulates production of cortisol via the hypothalamus and pituitary glands. In addition, the sympathetic nervous system (SNS) is activated and stimulates the adrenal medulla to release catecholamines, adrenaline and noradrenaline, allowing the body’s systems to transport energy to the organs more quickly. Consequently, homeostasis is re-established, provided the stressor falls into the adaptive capacity [31]. However, during severe and/ or chronic stress, dysregulation of the stress system (hyperactivation or hypoactivation) can disrupt homeostasis and lead to cacostasis or allostasis with possible various clinical manifestations [30]. Prolonged activation of stress mechanisms with increased glucocorticoid and catecholamine levels causes a condition where the demand on the individual exceeds their personal adaptive capacity (allostatic load). Therefore, according to research results, chronic stress and increased glucocorticoid/catecholamine levels may participate in cancer progression in different diseases, including HPV-related carcinogenesis [31].

It is significant that cortisol, as the main stress hormone, modifies apoptosis and changes the way in which cytokines are secreted. It is assumed that exposure to relevant stress events can result in dysregulation of the sensitivity and numbers of glucocorticoid receptors (GCR), and that cortisol can affect secretion of local cytokines and trigger stronger inflammation [25, 27, 39]. Glucocorticoids exert their effects through two subtypes of intracellular receptors: (type I) GCR with a high affinity for endogenous corticosteroids (that has a regulating function over the circadian rhythm of the HPA axis) and (type II) GCR with a lower affinity (important for acute stress reactions). In the nucleus, glucocorticoid acts as a transcription factor, and it binds to specific DNA sequences named glucocorticoid response elements (GRE). These sequences modulate gene transcription. Glucocorticoids also interact with other transcription factors, such as AP-1 (activator protein 1) and NF-kB (nuclear factor kappa B). During a stress inflammatory response, glucocorticoids decrease secretion of proinflammatory cytokines and increase secretion of anti-inflammatory cytokines. They also affect redistribution of leukocytes and decrease synthesis and expression of cytokine receptors, lymphocyte proliferation and adhesion molecule expression on the cell surface.

Finally, in stress, adaptive reactions involve the short-term activation of the HPA axis, whereas overproduction of stress hormones and disruption of the regulation of the HPA axis triggers a pathological response/reaction. Just as the cognitive perception of stress is important, so is coping, defined as the constant adaptation of cognitive and behavioral efforts to overcome demands that one finds overwhelming. Adaptive coping styles are
usually related to positive personality characteristics, while maladaptive styles are linked to less desirable characteristics. The coping styles a person uses depend primarily upon the situation but also on the disposition of the person themselves.

Results

The most important results and current knowledge on HPV, stress and carcinogenesis will be presented here, and relevant trends and patterns will be described.

Psychological stress and immunological parameters related to infections

Research in the field of acute and chronic stress have shown that many biochemical changes, e.g. changes in the blood, have been observed in people experiencing stressful events. For example, acute stress increases the number of lymphocytes in the peripheral blood, especially natural killer (NK) cells. Also, during short-term stress events (e.g., an exam), cytokine production disturbances have been observed along with a shift from T helper lymphocytes type 1 (Th1) towards T helper lymphocytes type 2 (Th2), with higher levels of immunoglobulins [25-28, 38].

During chronic stress, lower numbers of lymphocytes, decreased NK cell activity, lymphocyte proliferation to mitogen stimulation and phagocytosis, decreased cellular immunity, and late hypersensitivity reactions were observed. According to previous research results, during psychological stress, higher titers of antibodies against viral infection (Epstein-Barr virus antibodies) as well as reactivation of EBV, cytomegalovirus, and herpes simplex virus were recorded [25]. Thereby, latent viruses can be reactivated, and antibody production decreased. In the setting of chronic stress, there is a shift towards a Th2 response (with limited production of IFN and IL-2), which enhances humoral immunity and decreases cellular immunity, possibly affecting the development of infections and autoimmune and malignant diseases. Immune responses to stress depend on duration of the stress factor effect; subsequently, stress of shorter duration causes translation from specific to innate immunity, while in the setting of chronic stress, this translation is seen in a change from a Th1 type reaction to a Th2 type reaction. It is significant that Th1 cells engage in cellular immunity, secrete IL-2 and IFN-γ, TNF-β and activate macrophages, while Th2 cells induce humoral immunity by activating the B cell response and antibody production, as well as secreting IL-3, IL-5, IL-10, and IL-13.

Increased psychological stress can result in a higher occurrence of viral infections, especially those mediated by IL-6. The influence of stress on cell apoptosis can also be observed, where apoptosis disruption can lead to immunosuppression. For instance, in acute stress (e.g. academic stress) the process of apoptosis, important in carcinogenesis and the function of the immune system, is more frequently blocked. In a specific example, a connection has been observed between women with cervical lesions who have a pessimistic view of the world and immune system disorders, lower infection control and a higher risk of neoplastic cervical lesion progression. Chronically high sympathetic system activity decreases lymphocyte proliferation and changes the Th1/Th2 cell ratio in a way that suppresses Th1 cytokines, therefore reaction evolves in the direction of Th2 cells, which can increase the risk for a high virus replication rate.

Also, the relationship between psychological stress and increased sensitivity to respiratory infections can be explained by the impact of stressful life events on immunological functions. It has been shown that psychosocial stress and limited social support can have a big impact by increasing susceptibility to upper respiratory tract viral infections. Patients with colds, for example, had a statistically significant higher average sum of greater stressful life events in the previous year compared to patients without a cold [40]. Results have also concluded that urinary system infections are more frequent in people who suffer greater stress [41]. Concerning HPV, it has been observed that a higher number of life stressors in at least the previous 6 months, the absence of social support and the types of personal coping
mechanisms employed influenced the progression of HPV infection [25, 42]. In women, it has been shown that HPV infection and stress result in the disruption of IL-6 secretion and the progression of cervical lesions to cervical dysplasia. In chronically stressed caregivers who care for relatives suffering from dementia, a fourfold increase in IL-6 concentration was seen (by longitudinal observation). In women with cervical dysplasia, a connection between greater stress experiences and dysregulation of specific immune responses has been observed.

The connection between stress, immune factors, infections and carcinogenesis often includes a link between HPA/SNS functions, immunosuppression and viral oncogenesis, which together work within the immune response - both glucocorticoids and catecholamines regulate different immune functions (antigen presentation, T-cell proliferation, cell-mediated reactions, humoral immunity, etc.) [43-45]. Stress-induced immunosuppression is especially important for infection-related cancer, whereas disturbed cellular immunity may contribute to increased risk of oncogenic viral infection and DNA damage. Since several human tumor viruses including HPVs (HPV 16, HPV 33, etc.) are sensitive to the glucocorticoid/catecholamine-activating signalling pathways, stress hormones may consequently reactivate latent tumor viruses, stimulate viral oncogenes expression, and inhibit antiviral host responses. In response to glucocorticoids, high-risk HPV may activate gene expression, interact with cellular proto-oncogenes and evade the cellular immune response by down-regulating the HLA expression of class. Conversely, glucocorticoid antagonists suppress infection (the HPV activity) [43].

Pathogenesis of HPV infection and HPV-related carcinogenesis and possible association/relation with stress and psychosocial factors

Once HPV enters a cell via the α6 integrin (which functions as a receptor) there are three possible sequences: latent infection (identified only by molecular diagnostic methods), subclinical infection (identified by colposcopy, peniscopy or microscopic examination), and clinically manifest disease. Incubation lasts 1 to 8 months [44]. HPV proliferation in differentiated epithelial cells induces morphologically cytopathic changes in the form of koilocytosis best seen in condyloma (koilocytosis is not specific to HPV) [44, 46]. HPV infection histologically manifests with thickening of the epidermis, spinous layer, hyperplasia, and hyperkeratosis. Another important pathogenic factor for HPV lesion occurrence is ultraviolet light (UV). Normally, UV induces mutations in basal keratinocytes with consequent repair of DNA mutations or apoptosis. In the case of HPV infection, the cellular response to UV-induced damage is inhibited by E6 and E7 expression. This causes proliferation of damaged cells with a potential for malignant alteration [1].

Aside from the skin, HPV DNA can be found in anogenital and oropharyngeal malignancies as well as precancerous conditions. In these tissues' viral oncogenes, E6 and E7 expression is regularly present. Expression of the E6 and E7 genes is also found in cervical cancer cells. All this supports the observation that HPV has a role in the etiology of epithelial cancer [8]. Furthermore, epidemiology data suggest that HPV infection is a major factor in the etiology of cervical cancer. For the formation of irreversible malignancy, the following sequence is necessary: initial expression of E6 and E7 genes followed by suppression of apoptosis and the stable expression of E6 and E7 proteins that protect transformed cells from apoptosis. This is also the reason for tumor resistance against chemotherapy or hypoxia in HPV malignant lesions [8, 46].

Changes in cellular immunity in HPV infection are associated with changes in cytokine levels, induction of cellular adhesion molecules, and changes in molecules of tissue histocompatibility, human leukocyte antigens (HLA). Thus IFN-γ and cytokines act on the JAK signal transduction pathway. In vitro studies indicate that E region of HPV can bind to the signal transducer and activator of transcription (STAT) protein and inhibit IFN genes. E6 also reduces IFN gene expression and binds to Tyk2 kinase, preventing STAT protein phosphorylation and reduced cellular response to infection. It has also been found that up to 60% of HPV-infected patients have a low titer of antibodies that are specific for the viral capsid [47].
Psychosocial factors may affect the development and persistence of HPV, primarily by acting on the immune response. An association between psychosocial stress and CIN development in women has been established. HPV has developed a number of ways to avoid hosts' responses to infection. It hides the expression of viral proteins and reduces the expression of genes for IFN-α in keratinocytes. A successful immune response is characterized by a strong, local cell-mediated immune response. The immune response depends on the duration of stressors—a shorter duration of stress shifts from specific to innate immunity, while chronic stress shifts it from the Th1 type of reaction to the Th2 type of reaction. Significantly, Th1 cells participate in cellular immunity, secrete IL-2, IFN-γ, and TNF-β and activate macrophages, and Th2 cells stimulate humoral immunity, activate the B cell response and antibody production, as well as promote the secretion of IL-3, IL-5, IL-10, and IL-13. The natural course of HPV infection can vary. It may disappear spontaneously, lead to the appearance of intraepithelial neoplastic lesions and, ultimately, to the appearance of cancer. Oncogenic transformation requires the integration of the virus genome into the host genome. In doing so, DNA in the E1 region of E2 breaks down, leading to transcription disorders of E6 and E7. A number of other risk factors are also involved in the onset of malignant transformation: promiscuity, a lower socioeconomic status, a higher number of pregnancies and abortions, use of oral contraceptives, smoking, co-infections, and stress. The pathogenetic life cycle of HPV can be divided into two stages: (stage 1) persistence, i.e. HPV DNA is found in basal cells lasting from a few weeks to several years, with a low level of virus replication over many generations) and (stage 2) the productive phase (occurs in terminally differentiated suprabasal keratinocytes with the appearance of viral DNA amplification, capsid gene expression and virion formation) [44]. Several factors are important for the regression of HPV manifestation/infection, one of which is the influence of psychological stress, because it has been proven that stress can prolong the duration and severity of HPV disease [25]. In the regression of HPV infection, increased activity of Th1 cells was observed. However, during psychosocial stress, a decrease in the Th1 type of immune response is seen, and there is a shift towards the Th2 type of immune response.

The association between psychologic stress and HPV carcinogenesis can be explained by a few possible mechanisms [43]. The stress-activated HPA axis and SNS act through glucocorticoid- and adrenergic-pathways which impact immune regulation, involving stress-induced immunosuppression and oncogenic infection and other potentially stress-triggered neuroendocrine transmitters which influence cancerogenesis (e.g. serotonin promotes tumor cell growth and angiogenesis; dopamine exerts opposite actions) [43]. The adrenergic pathway and SNS activation support carcinogenesis through tumor growth, malignant transformation, macrophage infiltration, angiogenesis, inflammation, dissemination, etc. Thus, in cervical cancer cells, sustained adrenergic signalling results in protein kinase A (PKA) activation, which then causes inhibition of the tumour suppressive pathway (Hippo Yap pathway). According to research data, a strong association between stress/bereavement and a high viral load infection (HPV16) includes the promoting of overexpression and malignant transformation of HPV oncogenes, and a strong association between repeated infections and bereavement/stress may compromise host immunosurveillance and mediate HPV persistence or reactivation [45]. According to Lu, stress/bereavement was associated with an increased risk for cervical cancerogenesis, especially for women with high screening adherence or when multiple losses, loss of a child, sibling or spouse, and loss due to unnatural causes were analyzed separately [45]. Thus, stress hormones (corticosteroids) affect signal pathways involved in the malignant cell transformation processes induced by oncogenic human viruses.
It is known that stress affects the immune system by redistributing immune cells (leukocyte and neutrophil numbers increase, lymphocyte numbers decrease), which is supported by increased activities/values of catecholamines, cortisol, chemokines and adhesion molecules secreted during stress reactions [48-52]. Leukocytes are especially important because they secrete cytokines, affect the HPA, and have cytokine, hormone, neurotransmitter and growth factor receptors, etc. Current studies have observed that stressful life events can influence genes. That is, they exert activity changes in a variety of genes, which then leads to modulation of immunological reactions, including leukocyte numbers and distribution [52]. Furthermore, it has been seen in animal models that the neutrophil to lymphocyte ratio predicts susceptibility to infection [48]. A higher cortisol level, combined with increased neutrophil and decreased lymphocyte numbers, has been observed in psychologically stressed people [25]. There are some observed (biochemical) changes in plasma protein levels, as confirmed by studies of their values during illness and stress. There are various study results on animals and humans which support the association between psychological stress, infections and carcinogenesis (Table 1 and Table 2) [25, 48, 53-70]. Animal model studies show that stress increases values of acute phase proteins [71].

After stress, haptoglobin significantly increases, so it is used as a stress marker during animal transport and for evaluation of the animal's condition [72]. During acute academic stress, elevated total levels of serum proteins, alpha-2 and beta and gamma globulins, have been observed [73]. Similar results are seen in research by Ctvitano, where elevated leukocytes, alpha-2 and beta globulins were observed among HPV patients with significant experienced stress [25]. Immune-compromised patients are more susceptible to developing a variety of infections, including HPV infection.

Research results in the field of hormonal changes in stressed people, especially changes in cortisol levels and hormones of the HPA axis, are significant as well. Hypocortisolism is evidenced in people subjected to severe stress, as well as in chronic fatigue syndrome, fibromyalgia, and rheumatoid arthritis. On the other hand, elevated glucocorticoid levels with elevated leukocytes, decreased numbers of circulating lymphocytes and a decreased neutrophil to lymphocyte ratio. Cole has observed a significant correlation

### Table 1. Prominent animal study results which support the association between psychological stress, inflammation and carcinogenesis (articles are in alphabetical order by author)

<table>
<thead>
<tr>
<th>Author</th>
<th>Animals/Subjects</th>
<th>Methods and type of study</th>
<th>Main results and conclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arntz et al., 2015</td>
<td>Mice</td>
<td>In mice subjected to daily restraints stress, factors responsible for stress-induced changes in immune parameters were identified</td>
<td>Increase in stress-induced changes in immune parameters</td>
</tr>
<tr>
<td>Jensen et al., 1998</td>
<td>Mice</td>
<td>Infected mice (viral infection) were subjected to social or avoidance-learning stress</td>
<td>Stress applied before viral infection did not impact spleen size, while stress applied after infection markedly reduced spleen enlargement. This phenomenon was not seen in inactivated mice. Stress mice showed slightly lower death rates.</td>
</tr>
<tr>
<td>Magnen et al., 2013</td>
<td>Mice (mouse model)</td>
<td>Analysis of prostate gland cancer for densities of tumor, sympathetic and parasympathetic nerves fibers according to clinical outcomes</td>
<td>The formation of autonomic nerve fibers regulates prostate cancer development and dissemination.</td>
</tr>
<tr>
<td>Pascual et al., 2013</td>
<td>Mice model</td>
<td>Review and analysis of different studies (animal, human) on psychological stress impact and immunological processes in cancer biology, including social disruption (SD) stress</td>
<td>Psychosocial stress triggers sympathetic and neuroendocrine responses, which significantly impacts cancer, in part, through regulation of inflammatory mediators.</td>
</tr>
<tr>
<td>Reuter et al., 2004</td>
<td>Mice</td>
<td>The influence of restraint stress (RTS) on cancer cell (MC) activity and its consequences on IL-12/IFN-γ levels</td>
<td>Restraint stress (RTS) suppressed the chemotaxis responsible for MC cell recruitment into the infected tissues and suppressed several immune-mediated cytokines expressions in the effector response of MC cells (important immune response to control viral replication) and also expressed IL-12 (important for MC cell development and homeostasis) and IL-11 (important for MC proliferation).</td>
</tr>
<tr>
<td>Stelzl et al., 2003</td>
<td>Rat (male Fischer 344)</td>
<td>In socially stressed male rats, the effects of corticosterone stress on the normal distribution of T-cells in lymphoid organs and other tissues were examined</td>
<td>24 hours after injection of SS-C grafted blood T cells, infected rats were inoculated with an immunogenic virus, this resulted in a significantly higher number of viable tumor cells than in control rats. The presence of stress increased overall survival, substantially higher localization was observed.</td>
</tr>
<tr>
<td>Tholou et al., 2006</td>
<td>Mice (arthritic mouse model)</td>
<td>Analysis of ovarian cancer cancer (mouse model) for angiogenesis and neovascularization under stress</td>
<td>In stressed mice, ovarian cancer showed markedly increased neovascularization and enhanced expression of VEGF, MMP2 and MMP9, and angiogenic processes mediated the effects of stress on tumor growth. Behavioural stress enhances tumor angiogenesis primarily through both adrenocorticotropic hormone-dependent and -independent mechanisms (secretion of proinflammatory cytokines).</td>
</tr>
</tbody>
</table>
| Verz FA, 2021           | Rat (21-day-old male Wistar rat) | Social isolation stress effects on chemically-induced caval carcinogenics were examined in rats (isolated and grouped groups) after 90 days of age, the rate (both groups) underwent a marked increase in caval carcinogenics (CVA/NGP) for 20 days after treatment with a 5% solution of nitrate powder under stress, which resulted in significantly higher number of carotid artery lesions (CVA/NGP) diagnostic and screening of ocular tumor inflammatory mediators. | Social isolation stress increased the OSCE occurrence by 20%, compared to controls, and induced ratical higher tumor volume and angiogenesis.gompanied by increased focal formation of tumor microvessels, microvascular density (MDV), leucocytosis, and total numbers of tumor cells, which increased both tumor volume and angiogenesis, accompanied by increased focal formation of tumor microvessels.
between circulating leukocyte numbers and cortisol concentration only in the presence of GCR sensitivity [74]. Observations on the possible mechanism connecting chronic stress and infection are considerable. People who are chronically stressed throughout life could be at risk for an HPA disorder, and decreased immune defense reactions to infections, which are mediated by many cytokines and other transmitters. When cortisol regulation and cytokine production are disturbed, a disbalance of the Th1/Th2 immune reaction and the emergence of clinical symptoms are possible. Consequently, in patients exposed to severe chronic stress, a weakening of the HPA axis is observed [74, 75]. There are specific examples of cortisol levels affecting immune responses in humans, such as the relationship between psychological stress and immune responses in vitro [76].

### Table 2. Main study results (human subjects) which support the association between psychological stress, infections and carcinogenesis

<table>
<thead>
<tr>
<th>Author(s)</th>
<th>Year</th>
<th>Analysis and type of study</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chang, W.</td>
<td>2013</td>
<td>Relationship between depression and the risk of developing any kind of cancer and hormone-related cancers was assessed.</td>
<td>After adjusting for potential confounders, overall risk of developing hormone-related cancers was positively related to minor and major depressive disorders, but not to anxiety disorders. In hormone-related cancers, prostate cancer was positively related to major depression, while cervical cancer was inversely related to major depression. This difference (gender</td>
</tr>
<tr>
<td>Chen, Y.</td>
<td>2016</td>
<td>Review and evaluation of longitudinal associations between stress and cancer: a meta-analytic method.</td>
<td>In generally healthy populations, stress-related physiological responses are significantly associated with higher cancer incidence and poorer survival. A stress-prone personality, maladaptive coping styles, elevated emotional response, and poor quality of life were related to higher cancer incidence and poorer cancer survival/metastasis.</td>
</tr>
<tr>
<td>Celar, A.</td>
<td>2018</td>
<td>Effect of lifestyle factors in predicting the continuation of follow-up care: data collected from programs that provided follow-up care for women with abnormal Papanicolaou results and low-grade cervical lesions to determine whether lifestyle changes increase the risk of discontinuing follow-up care.</td>
<td>Among women who began had discontinued follow-up care (72.7%), more stressful life events in the past year were noted. Life stressors are common presenting demerit of how follow-up care among women.</td>
</tr>
<tr>
<td>Car, B.</td>
<td>2021</td>
<td>Review of the connection between cancer and stress and the role of 9-Formestanes.</td>
<td>ADHR agonists and 9-Formestanes target androgenic anti-tumor effects. Also, PBI and 9TM effectively retain the impact of stress on cancer patients.</td>
</tr>
<tr>
<td>Cotulah, M.</td>
<td>2020</td>
<td>Analysis of psychological and immuno parameters (serum leucocyte, soluble globulin, beta globulin, albumin, and protein) and certain cortisol levels in patients with clinical HPV manifestation: cross-sectional study.</td>
<td>Patients who experienced greater stress had significantly higher values of betaglobulin, soluble globulin, and beta globulin, though decreased after clinical HPV manifestations. Adaptation, coping with a chronic or stressor, stress significantly reduced the chances of having metastragraphic manifestations by 2.6 times. A higher perception of stress significantly increased the risk of developing metastases.</td>
</tr>
<tr>
<td>De Francesco, 1998</td>
<td>Relationship between personality, stress and cervical dysplasia was analyzed by comparing healthy women to those with autoimmune disorder (who personality was reversely attributed for the HPV genotypes causing cervical dysplasia with the fight-back/hormonal responses by using serum and urinary cortisol levels).</td>
<td>Significant differences between women with cervical dysplasia and healthy women were found in intellectual capability and handicaps, as well as in cortisol levels.</td>
<td></td>
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<tr>
<td>La, D.</td>
<td>2019</td>
<td>Analysis of whether psychological distress around the times of diagnosis of invasive cervical cancer leads to a higher risk of cancer-specific mortality: a retrospective cohort study</td>
<td>Psychological distress was associated with an increased risk of cancer-specific mortality. In patients with cervical cancer, psychological and lifestyle factors around the time of cancer diagnosis were linked to increased cancer-specific mortality, independently of tumor characteristics and treatment mortality.</td>
</tr>
<tr>
<td>Fang, C.</td>
<td>2020</td>
<td>In women with cervical dysplasia, a possible association between stress and the immune response to HPV was analyzed: serum concentrations of blood cytokines, stress related events and personal stress and a correlation T-cell response to HPV16 (fluorescent, one cervical sample (HPV typing)</td>
<td>Higher perceived stress levels were associated with a non-response to HPV16, without an association between stress levels and T-cell response to HPV16. Cervical dysplasia, higher perceived stress levels were associated with impaired HPV-specific immune response, suggesting a potential mechanism of stress influences cervical disease progression.</td>
</tr>
<tr>
<td>Fang, E.</td>
<td>2020</td>
<td>Analysis of whether severe emotional stress (loss of a child) influences the risk of getting cancers susceptible to immune modulation, such as cervical dysplasia and cancer: a historical cohort study.</td>
<td>Patients who lost a child had a significantly higher risk of getting cancers potentially associated with HPV infection (e.g., cervical cancer) where higher levels for most cancers were observed within 5 years after child loss, increased risk for liver and stomach cancers was also some evidence that a 3-year period. Thus, severe stress levels may increase the risk for several, clearly HPV-related cancers.</td>
</tr>
<tr>
<td>Bond, A.</td>
<td>2019</td>
<td>Prospective examinations of the association between social support and the risk of cancer and mortality: a cohort of 6,652 Japanese men and women.</td>
<td>Low social support was significantly associated with higher risk of both colorectal cancer and mortality in men. Social support may affect colorectal cancer and prostate through many factors, including healthcare variables and adherence to therapeutic regimens.</td>
</tr>
<tr>
<td>La, D.</td>
<td>2016</td>
<td>The association between harassment (i.e., loss of a family member due to suicide and abnormal cervical cytology, and excessive cervical cancer: and increasing adherence, were examined in a Swedish National Cervical Screening Register (women from 2,844,157 women, two times cancer control information were conducted).</td>
<td>Harassment was associated with a 4.8% higher risk of abnormal cytology, in both and excessive cervical cancer. Harassment was associated with a 49% risk of abnormal cytology, in both and excessive cervical cancer. Harassment was associated with an 8.9% higher risk of abnormal cytology, in both and excessive cervical cancer. The association between harassment and increased risk of developing cervical cancer may be attributed to stress-related increased immune response.</td>
</tr>
<tr>
<td>Krönfeld, 2001</td>
<td>Assessment of marital differentials in cervical 12-cancer causes types, based on individual mortality and cancer death for the whole German population.</td>
<td>The study (provisional cancer mortality among cancer patients compared to similar persons without a cancer diagnosis in the whole, more than 15% higher for more-rural cancer, more rural women and women breast mastectomy. This protective effect of marriage is not due to stage of illness, which was controlled for.</td>
<td></td>
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<tr>
<td>Presti, 2013</td>
<td>Association between life stress and progression/progression of cervical cancer were assessed among women co infected with HPV and HIV after 2-year follow-up. HIV infected women underwent a psychological interview, blood draw, colposcopy, and HPV cervical smear at study entry.</td>
<td>The study was designed to assess the risk of developing progression/progression (CIN) over 1 year by assessing HIV after obtaining relevant psychological and behavioral variables. Life stress may counteract an independent risk factor for stage progression/prevalence in the infected women. Stress management interventions may decrease the risk for IL progression/prevalence in women living with HIV.</td>
<td></td>
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<tr>
<td>Wilmanns, 2019</td>
<td>Elimination of the relationship between psychological factors, behavioral risks for abnormal cervical cytology, and abnormal cervical cytology: The focus for self-help intervention was the identification of psychological state of being compared to groups of women with normal/cervical cancer showing a primary care clinic and setting, attending a colposcopy clinic based on an abnormal test score (20 women participated in the study).</td>
<td>Behavioral risks for developing abnormal cervical cytology are associated with high stress levels, primary factors, and perceived discrimination, although there were no significant relationships between psychological factors and cervical cytology rates.</td>
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</table>

**Previous studies on corticotropes in women with various cancers indicated association/relation [31, 43, 55]. For example, in patients with prostate cancer, cortisol stimulates carcinogenesis by activating the androgen receptor in the absence of androgens. In ovarian cancer, glucocorticoids have anti-apoptotic effects on cancer cell lines, by upregulation of the anti-apoptotic proteins. Also, in colon cancer, the stress hormones pathways induce the proliferation of metastatic cancer cells by upregulating CDK1 expression (a dysregulation of CDK is associated with tumour proliferation activities).**
and lung cancers showed that dexamethasone influences cell apoptosis. Dexamethasone downregulated cisplatin-induced expression of cell death receptor pathway components, which indicates its anti-apoptotic role in many cancers, although pro-apoptotic actions are seen in lymphoid carcinomas. Also, glucocorticoids activate the PI3K/AKT pathway which inactivates pro-apoptotic factors/molecules (eg. caspases). This is the same pathway that HPV uses to cause immortality of transformed cells. HPVs E6 protein can activate oncogenic pathways including phosphoinositide 3-kinase (PI3K)/protein B (Akt), Wnt and Notch. The PI3K/Akt pathway controls cell proliferation, cell growth, angiogenesis and cell survival by influencing multiple downstream targets. In addition, dexamethasone increases some cancers' resistance to cytotoxic therapy (hepatocellular and colorectal), and in some cancers dexamethasone enhanced the growth of cell lines treated with 5-fluorouracil or cisplatin (breast, cervical, melanoma, neuroblastoma). For oncology patients, it is important to consider that glucocorticoids may induce chemotherapy resistance by blocking tumor cell death (i.e. anti-apoptotic effect), although cortisol is routinely administered throughout chemotherapy for treatment/prevention of nausea and allergic reactions. Corticosteroids affect signal pathways involved in the malignant cell transformation processes induced by oncogenic viruses. Glucocorticoid signaling participates in solid tumor progression through increased cell proliferation, inhibited apoptosis and DNA repair activity. In addition, in many cancer patients, chronic stress is associated with flattened diurnal cortisol rhythms and increased nocturnal cortisol, especially in those in advanced disease/cancer stages, and the altered cortisol patterns may partially reflect patient distress and/or avoidant coping. In various cancers (breast, lung, renal), the flattened cortisol slope is associated with compromised survival [43].

Not only are there studies that show the influence of psychological stress on immunological parameters and infection and carcinogenesis, but it has also been suggested that psychological stress impacts clinical manifestations of HPV, though study results are not consistent [25, 45, 68, 70, 76-78]. In several studies, the influence of negative life events, specifically, on clinical manifestations of HPV has been researched. Also, Pereira has observed that at higher levels of exposure to stress there is progression and persistence of precancerous HPV lesions [69]. According to Coker, psychosocial stress influences the course of genital HPV infection. Some other recent studies also show a connection between infections and stress [62]. It has been shown that perceived stress is associated with a longer duration of HPV infection in men over 50 years of age, although the association with HPV infection incidence has not been confirmed. There is some evidence that level of perceived stress (but not the number of negative life experiences) is associated with unreactivity to HPV 16, which was explained by the Th1/Th2 balance shift towards a Th2 type cytokine reaction [51, 78]. Coker also demonstrated that in Caucasian women, negative life events caused deterioration of HPV lesions [62]. The same was not true for Afroamerican women, which was explained by socioeconomic differences. Furthermore, Coker has demonstrated that psychological abuse is associated with pathological findings in Papanicolau cervical smears.

Unlike Coker, Wilkerson did not find any correlation between pathological findings of clinical HPV manifestations and perceived stress or number of negative life events [70]. Likewise, Sharp did not find a significant correlation between the influence of negative life events and HPV lesions [76]. Also, according to Massad, there is no statistically significant correlation between perceived stress and clinical HPV manifestations [79]. According to research by Massad and research by Wilkerson, it hasn't been determined whether stress or depression affects the prevalence of changes associated with HPV [70, 79]. Still, most research points to stress being a potential risk factor for development of HPV infection [51, 68, 80]. The influence of stress on patient compliance, and eventually on development of HPV skin and mucosal lesions, should also be considered because elevated stress levels can lead to decreased patient compliance, thus persistence or progression of intraepithelial lesions [70].
There are also some differences by sex in how stress affects HPV manifestations [25]. Thus, it has been observed that men, after being exposed to psychological stress, are more vulnerable to diseases than women. Longer lasting HPV infection in women who are older and have more notable stressful experiences was also observed. There is also a difference between the sexes in regard to localization of HPV manifestations—in women lesions in the genital region are more common, while in men they are more common in the oral region. Also, HPV transmission rates are higher in women, and the immune response, which is stronger in women, depends on anatomic localization. It has been observed that women are better protected from infection/the effects or manifestations of infection. Another difference seen in men is a lower prevalence of HPV 6, 11, 16 and 18. Men and women also perceive the importance of HPV infection differently by the very fact that there is a much higher incidence of carcinoma in women with evident HPV infection. There is an added impact of socioeconomic status on HPV manifestations for women of lower socioeconomic status, for whom, even though they don't have more prevalent HPV infections, carcinoma caused by HPV is more prevalent. This is presumably due to decreased accessibility to health resources, but also due to greater stress, maladaptive coping styles and limited social support.

With HPV infection, the patient’s perception of stress is important [80-84]. According to research by Cvitanović, a higher level of perceived stress means the probability of genital manifestations significantly increases (3.53 times) [25]. McCaffery demonstrated similar results, and Coker also observed that positive HPV testing results lead to increased anxiety [62, 82]. Also, a higher level of perceived stress has been demonstrated in women with cervical dysplasia and HPV. Increased lytic expression of HPV as a source of stress is also an important concern. A woman might become stressed simply by the appearance of changes on the cervix. This could lead to greater anxiety and the possible progression of those changes/lesions. Thus, a patient’s psychological state can lead to a closed cycle of progression of the manifested disease [85].

Thus, adaptive coping mechanisms for dealing with stress, including stress caused by disease, became a crucial part of avoiding a variety of health problems. According to some research on women, stress and the process of coping with disease can decrease immunity and thus increase the risk for clinically relevant HPV manifestations. Antoni observed a higher incidence of cervical dysplasia in women with maladaptive coping styles than in women with adaptive coping styles [80]. Other research has been done on the association between psychological changes and cervical carcinoma, particularly looking at stressors and stressful events that subjects have little control over. That kind of stress is considerably associated with pathological findings of squamous intraepithelial lesion (SIL). Abuse (in childhood or by a partner) is also associated with pathological findings of SIL. Psychological stress and maladaptive coping styles are also associated with a higher risk of SIL and invasive cervical carcinoma development, which is explained by decreased immune function. It has also been observed that changes in the form of SIL are more frequent in persons suffering from alexithymia, in whom decreased numbers of lymphocytes have been demonstrated. A relationship between infection and psychological factors has also been recorded, e.g. in anxious women, cervical cancer was more commonly found than endometrial cancer (which is not etiopathologically associated with infection). Regarding all of this, it should be emphasized that certain methodological issues arise when investigating the association between stress, the immune response and disease. Tests that investigate the influence of stress on immunological parameters are quite variable and depend upon many nonimmunological factors (age, sex, diet, smoking, substance abuse, lifestyle and socioeconomic status).

Research literature has proposed possible psychological treatments for addressing patients with HPV [25, 70, 76, 78, 79]. Antoni asked female patients with clinically manifested HPV infection in the form of a CIN to undergo cognitive-behavioral therapy and observed that those who underwent treatment experienced substantially less frequent occurrences of severe lesions with higher stages of CIN [80]. This suggests that including stress-management in standard dermatological practice could be effective, especially for patients suffering from anogenital HPV lesions [79]. By using adaptive methods of coping with stress and raising
awareness of the effect a patient’s perception of stress can have on disease manifestations, patient quality of life can be significantly improved and allostatic burden can be diminished.

Chronic stress may negatively influence patients with cancers through suppression of protective immunity related with cancer exacerbation and progression or metastasis, which is important for their immuno-protective responses and successful oncology treatment (primarily tumor immunotherapy), especiall for virally-associated cancers (HPV-associated cervical, anal, and oral cancers etc.). While previous research has thoroughly examined the effects of stress on some cancers (breast, prostate, lung, ovarian), more data for other cancers could be collected and examined in greater detail. Limited data on the stress-HPV-carcinogenesis association indicates a need for more research on the different features/mechanisms of stress-induced cancer progression. Since stress hormones and immune disturbances may affect carcinogenesis, there is a need to expand current knowledge on stress signalling in cancers (e.g. HPV carcinogenesis) and for a more precise understanding of stress. Multidisciplinary approaches to research and therapy including diverse sets of participants and factors/effects related to stress would be greatly beneficial. Also, it is necessary to better understand psychological/psychosocial features of cancer patients, e.g. how stress influences underlying molecular mechanisms. According to current knowledge, there are various possible ways to relieve the impact of psychological stress on cancer patients, such as the use of β-adrenergic receptor (ADRB) antagonists and downstream target inhibitors (which have significant anti-tumor effects), psychosomatic behavioral interventions, and traditional Chinese medicine [63]. Since the influence of psychological stress on HPV infection manifestations and carcinogenesis has been proven by research, specific psychological health measures and treatments should be designed and implemented to help patients deal with stress and its adverse effects in a healthy way.

Conclusion

Present findings suggest that stress is an important risk factor for HPV manifestation development and carcinogenesis. Understanding the key factors and processes clears the way for effective prevention and therapeutic intervention. Even though psychological factors have not been considered in many past epidemiological studies, they are now understood to be important in the setting of frequent and sometimes very serious illnesses such as HPV. Understanding perceived stress and biological changes in stress, as well as the evaluation of immune parameters, gives researchers a better picture of how stress influences HPV infections and how to improve disease management and outcomes.

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Author Contributions

LLM contributed conception and design of the article; LLM, HC, MK and AŠ analyzed the data; LLM, HC and ID wrote the manuscript; all authors contributed to manuscript revision, read and approved the submitted version.

Statement of Ethics

The authors have no ethical conflicts to disclose.
Disclosure Statement

The authors declare that no conflicts of interest exist.

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