Genetic susceptibility to oral cancer by human papillomavirus infection

Martha Rebolledo-Cobos¹* and Zuleima Yáñez-Torregroza²
¹Universidad Metropolitana, Dentistry Program, Atlántico, Colombia; ²Universidad Simón Bolívar, Faculty of Medicine, Basic and Biomedical Sciences, Atlántico, Colombia

Abstract

Environmental carcinogens and human papillomavirus (HPV) are the main responsible factors for oral cancer. Susceptibility factors in the human genome play a risk-modulating role; however not all individuals exposed to these carcinogens suffer from cancer. The purpose of the present review is to describe the main factors of genetic susceptibility to oral cancer due to HPV infection. A systematic search was carried out in three databases in English, with only 7 articles meeting the selection criteria. Genetic polymorphisms are shown in three categories, which are related to HPV and participate in oncogenesis. Three articles related to deregulation of cell cycle control mechanisms were identified, as well as one referring to mutations in the apoptosis pathway and three about polymorphisms in inflammatory and immune response genes. The association of polymorphisms for the development of oral cancer by HPV is evident, although it remains under study. Oral neoplasms’ oncogenesis pattern is not always associated with HPV, but with other environmental or epigenetic factors.


Introduction

The description of the variants in human DNA sequences and protein sequences started with two independent publications by Beaudet et al. and Beutler et al. in 1993. Those authors proposed that any abnormal change in the nucleotide sequence is usually a mutation,¹,² which may or may not cause phenotypic changes in affected individuals and not necessarily corresponds to a disease and can be inherited from the parents (germline mutation) or acquired and manifest itself throughout an individual’s life (somatic or multifactorial mutation).

In the latter mutation type, there is specifically a variation in the DNA sequence that can occur in a population with a frequency of 1 % or higher, and that is known as polymorphism.³ Polymorphisms occur naturally, with a neutral effect, beneficial or promoter of disease. They can also involve one or more nucleotide changes, as well as other mutations. Single nucleotide polymorphism is the most common, and it arises from every 1000 base pairs (bp) in the human genome and is usually found in areas that flank protein encoding genes or in regions recognized as critical for microRNA binding and gene/protein expression regulation.³ However, single nucleotide polymorphisms can also occur in coding sequences, introns or intergenic regions.⁴

In the context of hereditary and multifactorial diseases, a mutation has been defined as an alteration...
in the gene sequence that nullifies its biological function; it is also possible to find it in malignant tumors of the oral cavity when the individual is exposed to certain viruses or epigenetic risk factors. In general, there are few types of cancer where one or more genes whose mutations confer a high risk for developing them have been able to be identified. As for oral cancer (OC), especially squamous cell carcinoma and other head and neck cancers, although it is true that they have multiple etiologies and acquired risk factors such as alcohol consumption or smoking, sexually transmitted diseases or infections with human immunodeficiency, herpes simplex or human papilloma virus (HPV) play an important role. In the latter case, mutations of different kinds are identified, which confer susceptibility to neoplastic diseases originating in a primary viral infection at some point in life. HPV is the cause of other neoplastic lesions that sometimes persist, but not necessarily evolve into cancer, which is related to its genotypes: high risk (hrHPV) and low risk (lrVHP); the most commonly associated with OC are genotypes 16, 18, 31 and 45, considered to be of high risk.

There are investigations that associate the somatic component with the appearance of hrHPV-attributed OC due to changes in the genes involved with the cell cycle (which play a role in the modulation of cell DNA repair), control of the cell cycle, cell growth, apoptosis and inflammatory and immune response, which in physiological conditions usually provide stability to the human genome. This systematic review has as its goal to describe the main factors of genetic susceptibility for the development of OC or squamous cell carcinoma in association with HPV infection.

Method

- Population: Patients infected with HPV and who developed OC.
- Interventions: Molecular diagnosis of mutations and HPV genotyping with regard to the presence of OC.
- Objective: To identify susceptibility factors for OC associated with HPV infection.
- Included studies: case-control studies, systematic reviews or meta-analyses.

Inclusion criteria

- Full-text articles written in English.
- Studies assessing genetic susceptibility to OC due to HPV infection.
- Studies provided information on cell cycle control polymorphisms in OC associated with HPV.
- Studies that provided information on polymorphisms by variations in apoptosis.
- Studies that would provide information on polymorphisms of inflammatory and immune response genes.

Exclusion criteria

- Studies that provided inadequate information.
- Narrative reviews.
- In vitro studies.
- Animal studies.
- Studies involving potentially malignant alterations.
- Studies that included other infections, such as human immunodeficiency virus or herpes simplex.
- Studies where no molecular diagnosis and genotyping had been performed.

Sources or information and search

An extensive systematic literature search was manually conducted in the PubMed, Elsevier and Clinical Key databases, using the combination of the following descriptors: HPV and oral cancer, human susceptibility, HPV infection, oral cavity, polymorphisms of a single nucleotide, polymorphisms of cell cycle control, variations in apoptosis and polymorphisms of inflammatory and immune response genes. The articles involved only investigations in humans, with no gender or age distinction (Figure 1).  

Study selection

Initially, 115 articles were recorded in an electronic spreadsheet. After eliminating duplicate articles, only 64 were considered relevant due to their titles and abstracts; 18 were identified to assessed susceptibility to HPV infection in the oral cavity, out of which 11 were excluded because they failed to meet the selection criteria. Only seven articles were included in the final analysis (Figure 1).  

Data collection

A standard pilot-type instrument was provided for data extraction by the expert; only the appropriate elements were taken into account for this review. The
Results

For hrHPV infection to persist and extrapolate to cancer development, there are host genetic factors that contribute to the variability these malignant neoplasms develop with after viral infection. HPV viral oncoproteins E6 and E7 deactivate tumor suppressor genes p53 and pRb, which allows the cell to escape the normal cell cycle checkpoints, with subsequent cell transformation, immortalization and oncogenesis. Several authors point at three simultaneous pathways for genetic polymorphisms analysis that can help understand the different mechanisms involved in the appearance of HPV-associated OC and squamous cell carcinoma: cell cycle control polymorphisms, apoptosis variations and inflammatory and immune response genes polymorphisms.

Cell cycle control polymorphisms

The first type of polymorphism corresponds to the interactions between the human genome and the hrHPV genome; the mechanism of action lies in the relationship between tumor suppressor gene p53 and HPV oncoprotein E6. This non-structural protein binds to the p53 gene and starts its degradation, which causes uncontrolled cell proliferation. Gene p53 is located on the short arm of chromosome 17, band 13 (17p13.1), has approximately 20 kb and consists of 11 exons, the first one non-coding and placed between 8 and 10 kb of exons 2 to 11. Ninety percent of this type of mutations have been located in exons 5 to 8 of the gene; approximately 20 to 30% of mutations occur in all five hot-spot codons found in these exons.

Other p53 alterations are deletions, insertions, mutations in splicing sites and heterozygosity losses. The most common mechanism of p53 functionality loss is point mutation of one of the alleles and deletion of the other. In this sense, it should be noted that some p53 mutations are dominant, which is an
exception to the rule that suppressor genes manifest their oncogenic action only if there is an alteration of both gene copies;9,10 a single defective protein monomer stemming from the mutation of one of the two alleles of the p53 gene is enough to cause total inactivation of the tetrameric protein. When mutated monomers form complexes with normal monomers, a mutant p53 protein with noticeably lengthened half-life is formed.7,8

**Apoptosis variations**

During carcinogenesis, tumors have to develop multiple mechanisms to overcome host immune surveillance and intrinsic apoptosis or cell cycle arrest. The individual difference in resistance to apoptosis through the FAS pathway might allow many cancers to escape the immune system or counterattacking it. The germ line variants in extrinsic and intrinsic pathways might affect apoptotic efficacy and resistance to apoptosis and, consequently, influence on HPV infection.7,11 This can be particularly relevant to a viral mechanism that works through the cell cycle and apoptotic mechanisms. Gene polymorphisms of the FAS and FASL promoters have been suggested to contribute to the risk of HPV-associated cancer by inducing immune cells differential apoptosis in response to micro-environmental signals after HPV infection. The polymorphism at position 670 of the FAS promoter has been found to suppress the binding site for the transcription nuclear element and to alter the expression of the FAS gene; pro-72 homozygote p53 polymorphism in codon 72 appears to be an important apoptosis regulator through the FAS/FASL pathway in head and neck cancer.11

**Inflammatory and immune response gene polymorphisms**

While hrHPV are well understood as risk factors for OC and head and neck cancer, research on host genetic factors in inflammatory and immune responses to HPV infection might help understanding the association between HPV infection and OC. It is precisely these responses or immune surveillance efficacy that modifies HPV elimination or persistent infection.7,12 Inflammation is part of the host response to internal or external environmental stimuli, which is promoted by the action of pro-inflammatory cytokines, including interleukins (IL) 1 and 6, tumor necrosis factor and interferon-γ, and is commonly resolved with anti-inflammatory agents. Cytokines IL-4, IL-10, IL-13, as well as interferon-α and transforming growth factor-β play a role in the control of HPV-infected cell growth. Viral persistence, disease progression or transformation into cancer involve escaping from these mechanisms. Therefore, germine variants of these cytokines might modify the effectiveness of the defense against HPV and, consequently, infection rates.12

The success of HPV in decreasing immune responses may be important in the pathogenesis of HPV-associated cancer, since the etiology of HPV-induced cancer triggers a persistent viral infection that can be minimized by an effective immune response. The polymorphisms of several cytokine genes have been implicated in an induction of susceptibility or resistance to cancers caused by HPV infection due to their role in determining the host immune response. Therefore, genetic variants of cytokine genes in promoter or coding regions that are thought to influence on expression levels or functional efficacy may be involved in susceptibility to the HPV status of patients with squamous cell carcinoma and OC.13,14

Some studies have associated polymorphisms in pro-inflammatory and anti-inflammatory cytokine genes, particularly in regulatory regions, with intra-individual variations in cytokine production and cancer risk. The association of cytokines interferon-γ and IL-10 polymorphisms with cancer risk has been well documented: interferon-γ decreased transcription and IL-10 increased transcription has been observed in patients with HPV-positive cervical cancer.12,14 Interferon-γ plays an essential role in the defense against intracellular viruses and pathogens by inducing immune-mediated inflammatory responses. Single nucleotide polymorphism T + 874A, located at interferon-γ gene translation start site, which coincides with a binding site for the putative nuclear factor-kappa B transcription factor, could be fundamental in the induction of a constitutively high interferon-γ production. The +874 T to A alleles with low (AA), medium (AT) and high (TT), have also been significantly associated with cytokine production. IL-10 has a suppressive effect on cell-mediated immunity, which can be critical in eliminating QTs that host HPVg.13 There are different polymorphisms in the IL-10 gene, among which that of a single nucleotide in position 1082 of the promoter region plays an important role in determining high, medium and low production of IL-10.

The association of G/A single nucleotide polymorphism in position 1082 has been associated with low
Table 1. Cell cycle control polymorphisms related to human papillomavirus in oral cancer and squamous cell carcinoma

<table>
<thead>
<tr>
<th>Polymorphism</th>
<th>Mutation</th>
<th>Genotype frequency</th>
<th>n</th>
<th>p</th>
<th>Oncologic diagnosis</th>
<th>Country</th>
<th>Author</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell cycle control pathway</td>
<td>p53 codon 72</td>
<td>Arg/Arg = 23 % (8) Arg/Pro = 57 % (20) Pro/Pro = 20 % (7)</td>
<td>35</td>
<td>&lt; 0.809</td>
<td>SCC</td>
<td>India</td>
<td>Tandon et al.8</td>
</tr>
<tr>
<td></td>
<td>p53 codon 72</td>
<td>Arg/Arg = 8.47 % (22) Arg/Pro = 43.46 % (113) Pro/Pro = 4807 % (125)</td>
<td>260</td>
<td>&lt; 0.001</td>
<td>SCC</td>
<td>Pakistan</td>
<td>Saleem et al.4</td>
</tr>
<tr>
<td></td>
<td>p27</td>
<td>p27 (rs34329; 3.05 ratio, 95% CI = 2.12-4.40), cyclin E (rs1406), cyclin H (rs3093816), cyclin D1-1 (rs47451), cyclin D2 (rs3217901) and Rb 1-2 (rs3092904)</td>
<td>6/9</td>
<td>&lt; 0.0001</td>
<td>OC</td>
<td>India</td>
<td>Murali et al.10</td>
</tr>
</tbody>
</table>


Table 2. Apoptosis pathway polymorphisms11

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Genotype</th>
<th>n</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>FAS (mutación)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FAS 670 A &gt; G</td>
<td>AA</td>
<td>7</td>
<td>16.7</td>
</tr>
<tr>
<td></td>
<td>AG + GG</td>
<td>25</td>
<td>83.3</td>
</tr>
<tr>
<td></td>
<td>GG</td>
<td>34</td>
<td>80.9</td>
</tr>
<tr>
<td></td>
<td>AG+AA</td>
<td>8</td>
<td>19.1</td>
</tr>
<tr>
<td>FAS 1377 G &gt; A</td>
<td>AA</td>
<td>31</td>
<td>73.8</td>
</tr>
<tr>
<td></td>
<td>AG + GG</td>
<td>11</td>
<td>26.2</td>
</tr>
<tr>
<td></td>
<td>CC</td>
<td>11</td>
<td>26.2</td>
</tr>
<tr>
<td></td>
<td>CT + TT</td>
<td>31</td>
<td>73.8</td>
</tr>
<tr>
<td>FASL (mutación)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FASL 124 A &gt; G</td>
<td>AA</td>
<td>31</td>
<td>73.8</td>
</tr>
<tr>
<td></td>
<td>AG + GG</td>
<td>11</td>
<td>26.2</td>
</tr>
<tr>
<td></td>
<td>CC</td>
<td>11</td>
<td>26.2</td>
</tr>
<tr>
<td></td>
<td>CT + TT</td>
<td>31</td>
<td>73.8</td>
</tr>
<tr>
<td>FASL 844 C &gt; T</td>
<td>AA</td>
<td>31</td>
<td>73.8</td>
</tr>
<tr>
<td></td>
<td>AG + GG</td>
<td>11</td>
<td>26.2</td>
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<td></td>
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<tr>
<td></td>
<td>CT + TT</td>
<td>31</td>
<td>73.8</td>
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</tbody>
</table>

AA, medium AG and high GG cytokines production. Polymorphisms in human IL-1β and tumor necrosis factor-α genes have also been reported to influence on cytokine and transforming growth factor expression, which regulates epithelial cell proliferation and apoptosis.7,13,14 Tumor necrosis factor-α can directly control HPV infection by apoptosis induction in HPV-infected cells, such as cervical cancer and OC cells. Disturbance of the balance between pro and anti-inflammatory cytokine levels may be caused by inherited genetic mutations, from which common genetic variants can also modify key genes expression or function, disrupting cytokine balance and affecting cancer risk and outcome.13,14

The investigations examined for the synthesis of this review show biotransformation, detoxification, elimination or immune control of carcinogens such as HPV, together with cell regulatory mechanisms, DNA repair and apoptotic pathways, as intrinsic strategies to evade cancer.7 Tandon et al. and Saleem et al. describe polymorphisms in the p53 gene located on the short arm of chromosome 17, band 13 (17p13.1). In turn, Murali et al. associate them with the p27 gene and consider that they contribute to OC pathogenesis. Only three articles related to this type of polymorphism were found, two from India and one from Pakistan10 (Table 1).

Only one article showed evidence that there are mechanisms to overcome host immune surveillance and intrinsic apoptosis. Sun et al. consider that the FAS/FASL promoter variants in the apoptosis pathways alter the transcriptional activity of those genes and cell death regulation. However, they state that no study has investigated if tumor sites contribute to the association between the FAS/FASL polymorphisms and the risk of tumor recurrence.13 They also identified that patients with genotype index for oropharyngeal cancer and FASL 844 C/T had a significantly higher risk of cancer recurrence (cHR = 2.5, 95 % CI = 1.1-5.8, p = 0.043; HR = 2.7, 95 % CI = 1.2-6.0, p = 0.032) in comparison with patients with the FASL 844 CC genotype as the reference group, while patients without oropharyngeal cancer with the FAS 670 AG/GG and FASL 844 C/T genotypes had a significantly higher risk than those patients at risk of tumor recurrence (cHR = 2.2 and 1.7, 95 % CI = 1.1-5.1 and 1.0-3.0, p = 0.043 and 0.049, respectively) in comparison with their corresponding AA and CC genotypes11 (Table 2).

In the examined investigations, hrHPVs were widely described as risk factors for head and neck cancer and OC. The host produces inefficacious immune surveillance responses against HPV infections, which causes for viral presence and malignant transformation to perpetuate; three articles were found in this regard.

Jin et al. found that HPV 16 seropositivity was only associated with an increased risk of OC (OR = 3.1, 95 % CI = 2.1-4.6) and that the risk of oropharyngeal cancer associated with HPV 16 was modified for each
In addition, similar results were observed for combined-risk genotypes of four variants, and all these significant associations were more pronounced in several subgroups, particularly in patients with oropharyngeal cancer that never smoked.12

In turn, Hsu et al. stated that there are two cytokines involved in carcinogenesis.13 They observed an association between the polymorphism in TGF-b1 (G to C polymorphism at codon 25 < þ915> and IL-10 (1082 G/A), -819 C/T and 592 C/A) and OC risk in 162 patients. They found that the TGF-b1 genotype 25 GC codon is significantly more common in patients with OC in comparison with a healthy control group (p < 0.0001). Patients with genotype GC at codon 25 had an 11.09-fold higher risk of OC (OR = 11.09, 95% CI = 6.16-113.23). They demonstrated that IL-10 polymorphisms in positions 819 and 592 were correlated with OC risk (p < 0.0001). The genotypes that exhibited IL-10-592 C alleles had a higher risk of OC (OR = 3.1, 95% CI = 2.1-4.6). HNC Jin et al.12

Conclusions
The studies examined in this review show ample evidence of the relationship between HPV infection and the presence of OC, apart from epigenetic risk factors, such as tobacco and alcohol consumption, age and modified sexual intercourse. Viral integration in the cell genome of the oral mucosa makes the human being genetically susceptible to the development of malignant tumors. The presence of three categories of genetic polymorphisms and mechanisms of simultaneous participation has been observed in the oncogenesis process, as well as the identification of transforming genes or oncoproteins that induce deregulation of the cell cycle control mechanisms, thus producing instability in the genome, apoptosis alteration and instability in inflammatory and immune response genes.

The study of the association of these polymorphisms with the development of OC in association with HPV infection is still under development; in addition, the oncogenesis pattern of all of head and neck cancers or OC is not always exclusively related to the presence of HPV, but with different environmental or hereditary risk factors. In the Colombian context, no exhaustive studies were found emphasizing the susceptibility of human genetics to OC, squamous cell carcinoma or head and neck cancer associated with HPV infection.

References