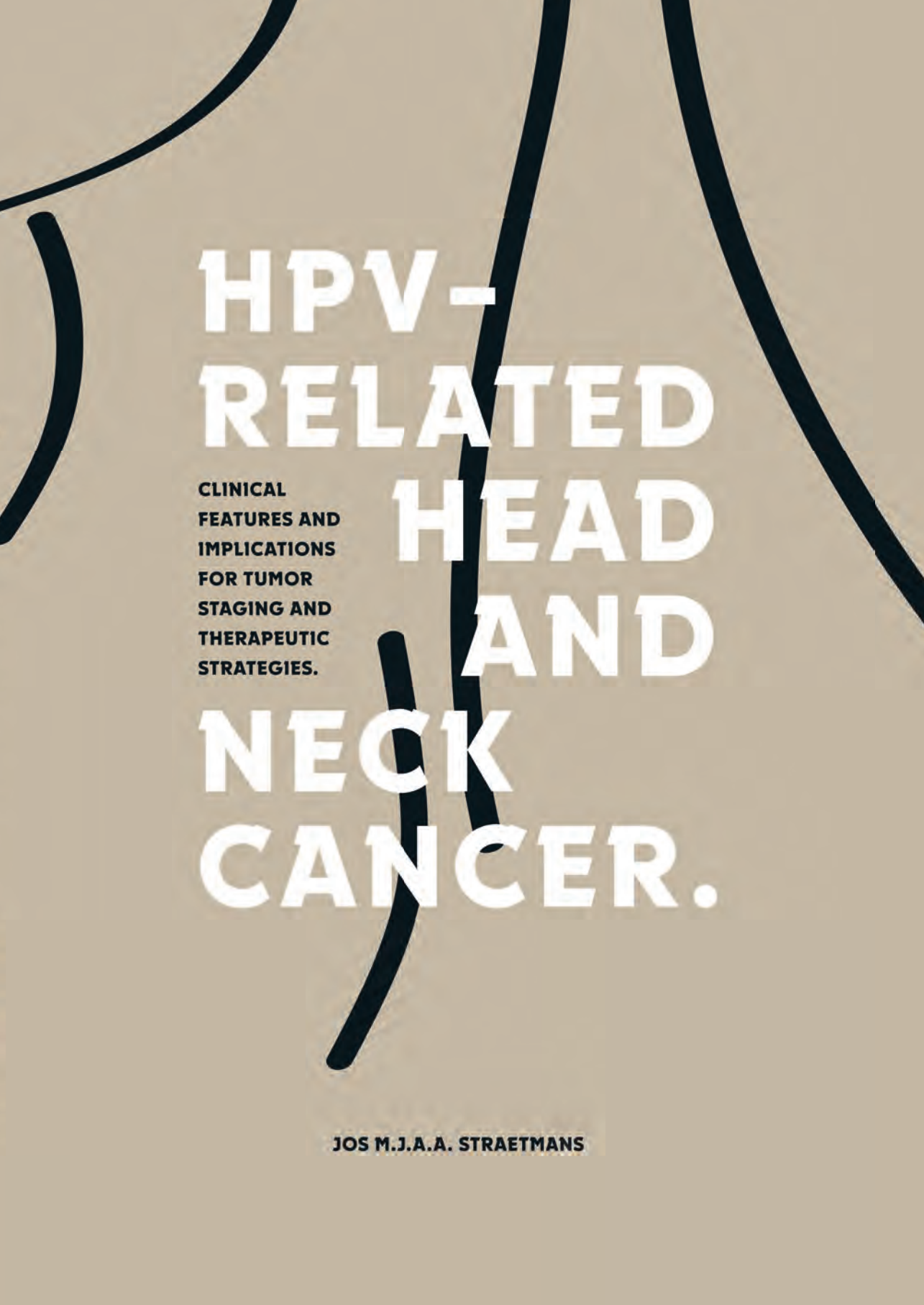




HPV- RELATED HEAD AND NECK CANCER.

CLINICAL
FEATURES AND
IMPLICATIONS
FOR TUMOR
STAGING AND
THERAPEUTIC
STRATEGIES.

JOS M.J.A.A. STRAETMANS

HPV-related head and neck cancer: clinical features and implications for tumor staging and therapeutic strategies

10 november 2020

Jos M.J.A.A. Straetmans, MD

Colofon

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HPV-related head and neck cancer: clinical features and implications for tumor staging and
therapeutic strategies

Proefschrift

Ter verkrijging van de graad Doctor aan de Universiteit Maastricht, op gezag van de Rector
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Door

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General introduction

1.1. HUMAN PAPILLOMAVIRUS (HPV) AND HEAD AND NECK SQUAMOUS CELL CARCINOMAS (HNSCCS)

Head and neck squamous cell carcinomas (HNSCCs) affect approximately 600000 patients per year and are associated with a mortality of 40-50%.¹ Abuse of tobacco and alcohol are well-established risk factors for the development of HNSCCs. The incidence of head and neck cancer as a consequence of tobacco and alcohol has decreased during the past decades.² However, the incidence of oropharyngeal squamous cell carcinomas (OPSCCs) has not decreased implying the role of another agent in its carcinogenesis, id est human papillomavirus (HPV). In fact, epidemiological evidence has revealed a rapid increase in the prevalence rates of HPV-associated OPSCCs.^{3,4} In the USA and Northern Europe more than 70% of OPSCCs are estimated to be HPV-associated, as compared with only 17% in Southern Europe.³⁻⁶

Regarding the role of HPV in human lesions, mucocutaneous warts already were described in the classic Greek era. The viral etiology of HPV causing warts in humans was demonstrated more than a century ago in 1907 by Cuffio et al.⁷ In 1983, the first evidence was published for HPV involvement in benign and malignant squamous cell tumors of the oral mucosa by Syrjänen et al., and it was demonstrated that HPV may be the etiological agent of a subgroup of oral squamous cell carcinomas.⁸ It was not until 1989 that the first report on the presence of HPV16 DNA in tonsillar squamous cell carcinomas was published (TSCCs).⁹

Nowadays, HPV - and HPV16 as the predominant type - has been well established as additional risk factor for OPSCCs, as has been demonstrated in several meta-analyses.¹⁰ The oropharyngeal subsites of the palatine tonsils and base of the tongue (BOT) are particularly involved.¹¹

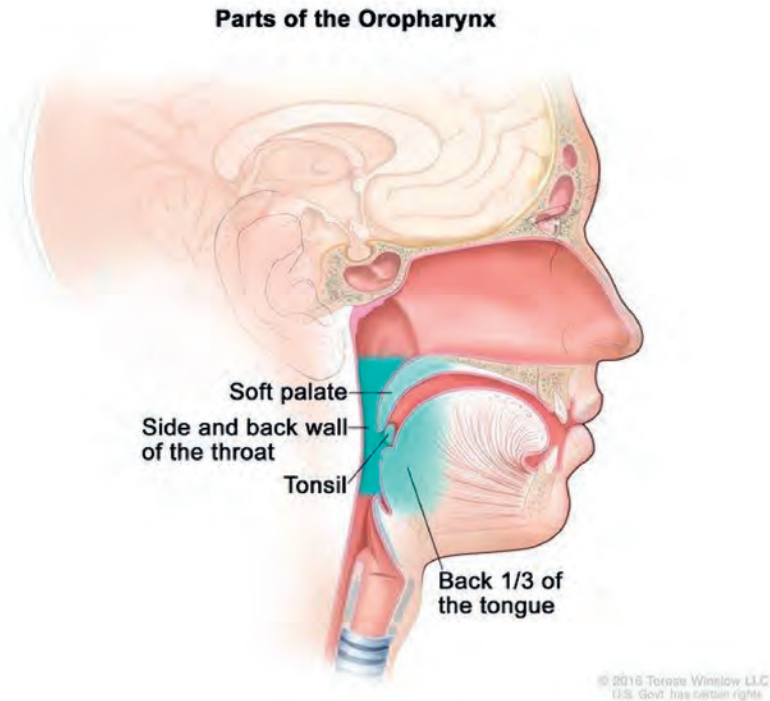
1.2. ANATOMY OF THE OROPHARYNX

The oropharynx is situated posterior of the oral cavity and cranially and caudally connected to the nasopharynx and the larynx/hypopharynx, respectively.

The oropharynx contains: 1) BOT, 2) the palatine tonsils and tonsillar pillars, 3) the soft palate, and 4) the pharyngeal wall (Figure 1).

Anteriorly, the border of the oropharynx is demarcated by the circumvallate papillae indicating the beginning of the BOT. The BOT is mainly composed of lymphoid tissue and extends laterally to the glossopalatine sulcus. The BOT ends inferiorly in the vallecula. The inferior border of the oropharynx is situated at this point, at the top of the hyoid bone.

Figure 1. Anatomy of the oropharynx



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The palatine tonsils are located on the lateral wall of the oropharynx and are -just as the BOT- mainly composed of lymphoid tissue in a fibrous capsule. They are situated posterior to the retromolar trigone and are surrounded by the anterior and posterior tonsillar pillars. The anterior tonsillar pillar is formed by the palatoglossal muscle, whereas the posterior pillar is formed by the palatopharyngeal muscle.

Together with the adenoid located in the nasopharynx, the lymphoid tissue of the tonsils and the BOT form the ring of Waldeyer.

The soft palate forms the cranial border of the oropharynx and caudal border of the nasopharynx. It connects anteriorly with the hard palate demarcating the border with the oral cavity.

Posterior to the tonsillar pillars, the oropharynx is surrounded by pharyngeal wall, which is composed of the pharyngeal constrictor muscles, and forms as such the posterior wall and lateral walls of the oropharynx.

1.3. HPV: CARCINOGENESIS AND DEREGLATION OF THE CELL CYCLE

The tonsillar crypts may serve as the probable site for HPV to enter the tonsillar mucosa as it is considered to function as a storage location for HPV during and/or after an HPV-infection. The crypt also lacks the highly structured barrier function of the squamous epithelium and has a high number of epithelial reserve cells/stem cells.^{12, 13} The possible occurrence of microlesions in the tonsillar squamous epithelium subsequently allows the virus to penetrate the mucosa and infect the basal cell layer of stratified epithelium. It has to be noted that primary premalignant lesions that can be attributed to the presence of high-risk HPV (HR-HPV) are only rarely observed.¹⁴ Therefore, studies on uterine cervical carcinogenesis provide most current knowledge on the initiation of HPV-associated mucosal disease and the dysregulation of the cell cycle by HR-HPV.^{6, 15} After infection by HR-HPV, early HPV genes *E1* and *E2* are expressed and the viral DNA replicates from episomal DNA. The infected cells replicate and move into the parabasal layers. Then *E6* and *E7* are expressed, which result in suppression of differentiation and re-entering of the cell cycle. Infected cells move forward to the superficial layers in the epithelium, where the viral genome is replicated and late genes *L1* and *L2* and *E4* are expressed. *L1* and *L2* proteins enable encapsulation of the viral genomes in the nucleus to form progeny virions (infectious virus particles), which are shed from the squamous epithelium to initiate new infections. Known uterine cervical lesions in which these non-neoplastic productive HPV infections occur are cervical intraepithelial neoplasia 1 (CIN1) or low-grade intra-epithelial lesions (LSIL). Regression of these lesions by an adequate immune response is still possible.^{6, 15}

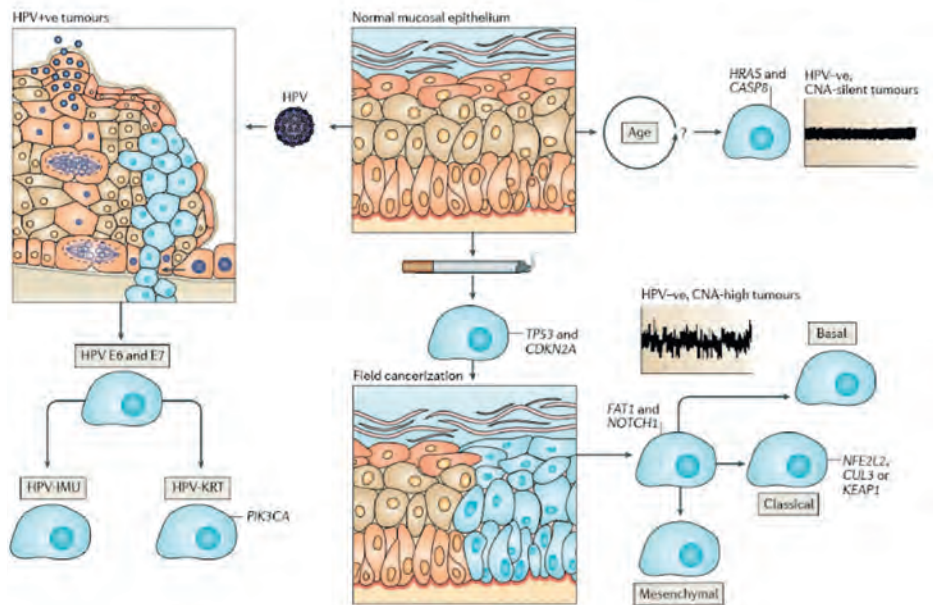
Approximately 5% of HPV infections are persistent, leading to local immune suppression, accumulation of chromosomal changes in the infected host cells, deregulated expression of HPV early genes, and reduced viral production. Approximately 0.3-1.2% of initial infections progress to invasive cancer.¹² In this transition, viral DNA often integrates into the host genome, leading to disruption or loss of *E2* and upregulation of *E6* and *E7* oncogene expression. However, expression of viral oncoproteins *E6* and *E7* can also occur independently of HPV DNA integration in the host genome.^{12, 16} Although much research have been carried out to unravel the physical status and copy number of HPV in HNSCCs, the involvement of these parameters in deregulating human gene expression and their value in predicting prognosis has not been sufficiently clarified yet.^{12, 17}

The ubiquitous expression of viral oncoproteins *E6* and *E7* is essential for tumor development and results in inactivation of p53 and the retinoblastoma protein (pRb), respectively. This leads amongst others to cell cycle deregulation and increased cellular proliferation, and inhibition of apoptosis. As a consequence, cyclin-dependent kinase (CDK) inhibitors, including p16^{Ink4a} and p21Cip1/Waf1, and the Mdm2 inhibitor p14Arf, are upregulated, which subsequently leads to CDK4/6 inhibition and downregulation of Cyclin D1.¹⁸⁻²¹ Zhang et al. recently identified two possible HPV-positive OPSCC subgroups based on molecular expression profiles: HPV-KRT (HPV-keratinocyte differentiation and

oxidative reduction process) or HPV-IMU (HPV-immune response and mesenchymal cell differentiation) tumors (Figure 2).²² Further study is required to confirm these findings.²³

In contrast, HPV-negative carcinomas, induced by smoking and alcohol consumption, are generally characterized by near universal loss of function *TP53* mutations and *CDKN2A* inactivation (p16^{Ink4a} loss of function) or *CCND1* amplification (overexpression Cyclin D1), also resulting in deregulation of the cell cycle and apoptosis.^{14, 23} Within HPV-negative HNSCCs, two distinct subgroups have been identified based on the presence of numerous or absent copy number alterations (CNA), resp. CNA-high and CNA-silent.^{23, 24}

Figure 2. Genomic carcinogenesis models of head and neck squamous cell carcinoma.



A schematic overview of squamous cell carcinogenesis in the head and neck is shown. The main distinction among head and neck squamous cell carcinomas (HNSCCs) is the presence of three genetic subgroups: tumors that contain transcriptionally active human papillomavirus (HPV+ve), tumors that are HPV-negative (HPV-ve) and have numerous copy number alterations (CNA-high), and tumors that are HPV-ve but CNA-silent.

HPV infection in oral squamous epithelium mostly leads to productive infections, while infection particularly in specific oropharyngeal crypt cells (light blue) might lead to an oncogenic event resulting in either HPV-KRT (HPV-keratinocyte differentiation and oxidative reduction process) or HPV-IMU (HPV-immune response and mesenchymal cell differentiation)²² tumors. These subgroups have been identified by expression profiling but have not been definitively verified and are still under investigation.

The P53 and RB pathways that play a key role in cell cycle control are frequently abrogated in HPV–ve tumors, except they seem to remain active in CNA-silent tumors. In addition, the etiology of this latter subgroup remains unclear, and ageing is hypothesized to be the risk factor. Many cancer genes and pathways seem to be involved in the progression of the HPV–ve, CNA-high tumors, but FAT1 and NOTCH1, which might act in the WNT– β -catenin pathway, are worth mentioning, and smoking is a known risk factor. At least three subgroups of tumors can be identified based on expression profiling, indicated as classical, basal and mesenchymal, but more may exist. The classical HPV–ve, CNA-high subgroup is characterized by nuclear factor erythroid 2-related factor 2 (NFE2L2) pathway mutations. HPV–ve tumors typically develop from mucosal precursor changes that can present as leukoplakias. Cells in these ‘fields’ progress to cancer by an accumulation of mutations. The current lack of data on precursor changes hampers the precise timing of events, but it is likely that the accumulation of events is the most important factor. Specific details and references are indicated in the main text. CASP8, caspase 8; CDKN2A, cyclin-dependent kinase inhibitor 2A; CUL3, cullin 3; KEAP1, kelch-like ECH-associated protein 1. (source: Leemans CR, et al. *The molecular landscape of head and neck cancer*. *Nat Rev Cancer*. 2018;18:269-282. Approval for reproduction of the figure was obtained by Nature Cancer Reviews)²³

1.4. PREVALENCE OF HPV: INFLUENCE OF DETECTION METHOD

In contrast to HNSCCs caused by tobacco and alcohol, the incidence of HPV in TSCCs has increased over the past decades and ranges from 20% to 90% in different studies.⁴ Incidence numbers of HPV involvement in HNSCCs depend on the detection approach used and also a wide geographical variation in HPV distribution has been described.^{5, 18, 25-27} Furthermore, the presence and detection of HPV-DNA in the tumor as only criterion for HPV-positivity might overestimate the true role of HPV in head and neck carcinogenesis, as it may reflect, for example, a transient infection unrelated to the carcinogenic process occurring for example after oral sexual intercourse, or DNA contamination during tissue processing.^{28, 29} Currently there is no single test that is considered the ‘gold standard’ for HPV-positivity in formalin fixed, paraffin-embedded (FFPE) HNSCC tissue. In order to detect a true biological HPV infection, it therefore has been proposed to combine HPV DNA detection by PCR with a second test that demonstrates viral activity. One option is reverse transcriptase PCR for HPV E6/E7 transcripts, which is easy to carry out on fresh frozen tissue, but may be cumbersome on FFPE tissue.²⁹ As an alternative, p16^{ink4a} overexpression has been reported as the most reliable surrogate marker for the presence of HR-HPV in OPSCC.^{18, 30-32} Many studies have shown, that the combination of p16^{ink4a} immunohistochemistry (p16^{ink4a}-IHC) followed by detection of HR-HPV DNA by PCR is a reliable, and accurate algorithm to distinguish HPV-positive from -negative HNSCCs.^{18, 19, 33-36} Nevertheless, false-positive rates for p16^{ink4a}-IHC has been reported (up to 7%).³² Results on HPV-prevalence and consequently all results of clinical data associated with HPV in OPSCCs, thus, should be taken with care and with special regard to the detection methods used.

1.5. FEATURES DISTINGUISHING HPV-POSITIVE FROM -NEGATIVE HNSCCS: ARE THEY STRAIGHTFORWARD?

There is increasing evidence that the pathogenesis of HPV-positive tumors is different from their HPV-negative counterparts, which is confirmed by molecular and clinical differences between the two subgroups.¹¹ But is it really that simple?

Weinberger et al. (2006) proposed a model for the development of HNSCCs, in which three classes are differentiated, divided into two arms. In the first arm, tumorigenesis is induced by alcohol consumption and/or smoking tobacco (Class I). However, in this group HPV-superinfection of the tumor site may occur, with biological features that do not differ from alcohol/tobacco-related tumors (Class II). In contrast, in the second arm, the inducer of tumorigenesis is HPV (Class III), which is proposed to be independent of smoking tobacco and alcohol consumption.³⁷ In general, patients with HPV-associated HNSCCs tend to be younger, smoke less tobacco and consume less alcohol, and have a more favorable prognosis when compared to patients with HPV-negative carcinomas. The true etiological role of HPV in HNSCCs, however, is challenged by factors such as concomitant tobacco and alcohol use together with the mentioned limitations caused by detection approaches used. This leads to questions like: Is the presence of HPV 16 DNA in the oropharyngeal tumor of a tobacco-smoking patient indicative of a HPV- or smoking-driven carcinogenesis? And why is the prevalence of OPSCCs increased in smoking patients, independent of the presence of HPV in their tumors?³⁸

From a molecular point of view, literature has provided evidence that there are also molecular differences within HPV-positive and HPV-negative HNSCCs (Zhang et al (2016), Leemans et al (2018), (TGCA: 2015), indicating the presence of different subgroups even within the three “Weinberger”-classes: CNA-high and -silent HPV-negative HNSCCs, and the subgroups HPV-KRT and HPV-IMU within the HPV-positive carcinomas. Verification of these subgroups and their association with clinical characteristics remains to be studied.²²⁻²⁴ On top of this, Wichmann et al. highlighted the possible importance of variants within the HPV 16-virusses worldwide, and their possible associated differences in clinical behavior and outcome in head and neck cancer, as is observed in uterine cervical cancer.^{39, 40}

Clinical characteristics and demographics associated with HPV-positive tumors are also not always unequivocal. The numerous studies published are more than once contradictory and are once again often biased by definition and detection of HPV.^{28, 41-44} Identifying the risk profile prone to HPV-induced HNSCC development seems to be complicated by the interaction of the different clinical variables as tobacco and alcohol use, age, gender, and sexual behavior. Moreover these variables are also affected by culture differences worldwide and even within continents (e.g. southern versus northern Europe).⁴¹

Therefore, the identification of the ‘typical’ HPV-positive HNSCC patient may be less straightforward than expected. Therefore, research should be directed to improve the identification of the HPV-positive “persona”, because it may help in the selection of true-HPV-positive tumors as well as in identifying subgroups of HPV-positive tumors associated with different risk levels regarding disease progression and survival.

1.6. PROGNOSIS OF HPV-POSITIVE OPSCC: THE INFLUENCE OF ADEQUATE STAGING AND THERAPEUTIC MANAGEMENT

1.6.1. Lymphatic drainage of the oropharynx

To describe the role of TNM classification in OPSCCs and how classification is affected by HPV, the first part of this section is dedicated to the description of the lymphatic drainage of the oropharynx. This is important for the understanding of possible differences in drainage dependent on the oropharyngeal subsites.

Lymph node drainage from the BOT and the palatine tonsils is typically to levels II to IV and the lateral retropharyngeal nodes.⁴⁵ The most common site of lymph node metastasis is to the jugulodigastric nodes in region II. Occasionally, levels Ib and V become involved. Level Ib is at increased risk of involvement in case of significant invasion of the oral tongue. Midline tumors, such as BOT tumors, are at higher risk for bilateral lymph node metastasis. Small lateralized TSCCs without midline extension have less risk of contralateral lymphadenopathy. Sood et al. confirmed the differences in lymphatic drainage between TSCCs and carcinomas of the BOT.⁴⁶ The first group is found to have more ipsilateral node involvement, whereas bilateral node involvement occurs almost only when the BOT is involved.

1.6.2. Influence of HPV on the UICC staging system: consequences for therapeutic decision-making and prognosis

At the start of our research on HPV in OPSCCs, the most important prognostic marker for all head and neck carcinomas including carcinomas originating from the oropharynx was lymph node involvement (N-status).⁴⁷⁻⁴⁹ Smoking and alcohol consumption were associated with a worse survival. HPV-presence in OPSCCs was reported to be associated with a better survival when compared to HPV-negative tumors. TNM-classification (7th edition UICC staging system) was leading in therapeutic decision making.³⁰ The standard treatment of OPSCCs in the Netherlands has been dependent on tumor location and TNM stage (UICC), independent of HPV-status. Smaller primary tumors can be treated surgically or with radiotherapy, although currently tumors are predominantly treated with radiotherapy. Larger tumors (T3 and T4) are preferably treated with radiochemotherapy or dependent on the overall condition and age of the patient with radiotherapy alone. In residual/recurrent cases, salvage surgery may be performed.⁵⁰

Literature on the prognostic importance of tumor spread to the cervical lymph nodes in TSCCs was not consistent when our research started. A higher N status was not always reported to result in a worse prognosis. It was reported that HPV-associated tumors show another clinical behavior than HPV-negative tumors. They often present with advanced lymph nodes in early primary tumor stages, resulting in advanced staged OPSCCs despite small (T1-T2) primary tumors.^{30, 51, 52} Moreover, HPV-positive OPSCCs respond significantly better to therapy than HPV-negative tumors.⁵³ Therefore, the presence of HPV in OPSCCs alters the consequences of staging for determining patients' prognosis and choice of treatment.^{37, 54-57} As a consequence in the course of the present PhD research the TNM classification for OPSCC has been adapted. In the newly introduced 8th edition UICC staging system, HPV-classification based on P16^{ink4a}-IHC testing alters the staging of lymph node involvement of OPSCCs which is comparable to the staging of nasopharyngeal carcinomas as demonstrated in the ICON-S-study.⁵⁸

1.6.3. Role of HPV in cervical metastases of unknown primary tumors

Knowledge on HPV-positive OPSCCs and its related pattern of lymph node metastases suggests a role for HPV detection in the diagnostic work-up of cervical metastases of unknown primary tumors (CUP). The additional value of HPV-testing in the diagnostic work-up in CUP patients was advocated in order to find the primary tumor, help improve outcomes and investigate de-intensification of treatment protocols in this group of patients because patients are often treated extensively with neck dissection, bilaterally applied radiotherapy and radiotherapy of the pharyngeal axis.^{59, 60} However, studies on the prevalence of HPV in lymph node metastases of which the primary tumor could not be detected after diagnostic workup, so-called "true" CUPs, are scarce and contradictory, and HPV prevalence rates range from 0% to 100% and were tested in relatively small sample numbers (range 1-25).⁶⁰⁻⁶⁵

1.6.4. De-escalation of therapy in HPV-positive TSCCs: *primum non nocere*

HPV-related OPSCC has emerged as a separate entity in terms of etiology, biology and clinical behavior; importantly, it has a more favorable prognosis and may require less intensive therapy. The ultimate goal in the treatment of patients with OPSCCs is to improve the efficacy and minimize the toxic effects of treatment. There are increasing indications that de-escalation could be possible. Different de-escalation trials have been recently published and/or are going on, however, with no clear evidence for adjusting current treatment protocols.⁶⁶⁻⁶⁸ Furthermore, it has to be kept in mind, that despite advances and innovations in multimodality treatment and a better understanding of head and neck carcinogenesis, the survival rates of a subgroup of locally advanced OPSCCs have not improved substantially and that the prognosis for patients with recurrent disease and/or distant metastases remains very poor even when tumors are HPV-positive.⁶⁹

Current de-escalation trials therefore focus on low-risk HPV-positive OPSCC subgroups. To better identify which patient groups are favorable for less intense therapy, adequate understanding of risk-profiles within HPV-positive OPSCCs is necessary.

1.7. AIMS OF THIS THESIS

Within the above mentioned context a couple of questions arose, which are given below. This thesis aims to provide the necessary answers to these questions and to attribute to a better understanding of the role of HPV in the clinical presentation of OPSCCs and CUPs, and its implications for tumor staging and therapeutic strategies.

1.7.1. To what extent are lymph nodes affected in HPV-positive carcinomas and what is the value of N-status in HPV-positive carcinomas?

For this purpose, the prognostic value of N status in TSCCs was examined in a population of 81 patients, while also taking into consideration the HPV status, clinicopathological features (age, gender, TNM classification, tumor differentiation grade, smoking tobacco, and alcohol consumption), and treatment of the tumor.

1.7.2. Is p16^{Ink4a} immunohistochemistry (p16^{Ink4a}-IHC) a reliable surrogate marker for the presence of high-risk- and low-risk-HPV in benign and malignant head and neck lesions?

To address this question, p16Ink4A-IHC immunohistochemistry was performed on a series of benign and dysplastic head and neck lesions and more specifically on paraffin-embedded tissue sections of 162 oropharyngeal squamous cell carcinomas (OPSCC), 14 tonsillar and 23 laryngeal dysplasias, and 20 tonsillar and 27 laryngeal papillomas. PCR, enzyme-immunoassay and FISH analysis were used to assess HPV-presence and type.

1.7.3. Is there an additional value for HPV-testing in the diagnostic work-up of CUP to identify a possible missed primary tumor?

29 true-CUP patients were analyzed for the presence of p16^{Ink4a} and HPV 16 DNA in the histologic specimens of the resected cervical lymph node metastases. Patients with CUP underwent a comprehensive diagnostic-work-up and were considered as true-CUP patients if a primary tumor was not detected through staging examination.

1.7.4. If HPV-positive tumors present more often with metastases to the regional lymph nodes, what is the frequency of HPV in cervical metastases of carcinomas with unknown primary tumor (CUP) in the head and neck area? And independent of HPV-status, is de-escalation of therapy an option?

For this purpose, data of patients with 'true'-CUP were evaluated for the presence of HPV by detection of p16^{ink4a} and HPV DNA, and how this affects outcome. First, the presence of HPV in neck metastases of CUP was analyzed. The data of a Dutch and a German academic cancer center (Maastricht and Cologne) were merged and a retrospective analysis was carried out. Second, the different therapeutic strategies and survival rates of both centers were analyzed. This gave us the possibility to study the additional value of irradiating the pharyngeal axis, as well as that of bilateral versus ipsilateral postsurgical radiation, with or without concomitant chemotherapy.

1.7.5. Does de-escalation of therapy in CUP results in different outcome and is it safe?

To critically evaluate the results of the above mentioned bi-national study, the outcome of patients with cervical CUP (n=124) in relation to the applied treatment in two Dutch academic head and neck clinics was examined. Both centers had a congruent history of de-escalation of therapy over time. Results of unilateral versus bilateral post-operative irradiation and radiotherapy of the pharyngeal axis were compared in terms of disease-free survival (DFS), regional recurrence rate (RCR) and overall survival (OS). In the same series the relation of HPV-positivity (assessed by detection of p16^{ink4a} and HPV DNA) of affected lymph nodes with outcome was investigated retrospectively.

1.7.6. What is the prognostic value of lymph node metastases in HPV-positive TSCCs and to what extent do the 7th and the 8th edition of the UICC classification system take this into account? Which parameters besides TNM enhance reliability in predicting prognosis and should be considered for risk stratification in therapeutic decision-making in the future?

The aim of this study was to contribute to a better tumor classification system for HPV-associated TSCCs. We focused on a large series of 368 TSCCs, which were subjected to HPV-analysis using p16-overexpression, HPV-specific PCR and/or FISH. All cases were staged with the 7th and the 8th edition of OPSCC and the prognostic value of T-, N- and M-status was then examined. Then additional parameters (age, smoking behavior, alcohol consumption, tumor differentiation grade, and treatment) were included in the analysis. It was shown that the addition of some of those parameters could improve the prognostic value of the latest (8th) TNM staging system for TSCCs.

1.7.7. Does HPV-related tumor biology in head and neck offer new targets for therapy?

In a review clinical and molecular characteristics of HPV-positive and – negative HNSCC were studied and possible ways to target specifically the HPV-infected cells were explored to elucidate possible new therapeutic strategies for these tumors.

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Human papillomavirus reduces the prognostic value of nodal involvement in tonsillar squamous cell carcinomas.

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2.1. ABSTRACT

Objective/Hypothesis: Assessment of the prognostic value of nodal status in relation to human papillomavirus (HPV) status and the various treatment modalities in tonsillar squamous cell carcinomas (TSCC).

Study design: Retrospective 5-year survival-analysis

Methods: A 5-year follow-up of disease-free, disease-specific and overall survival in a group of 81 patients with TSCC was conducted. The nodal status and integration of HPV-DNA in the genome (detected with fluorescence in situ hybridization) as prognostic indicators was examined while correcting for other clinical parameters (smoking habits, alcohol consumption, treatment modality, differentiation, TNM-classification).

Results: Of TSCCs, 41% were positive for HPV type 16. In these TSCCs, the primary tumor was significantly smaller when compared to HPV-negative TSCCs ($p=0.04$), whereas the percentage of cases with cervical metastases was identical. In the total tumor population, it was not nodal involvement, but rather HPV manifestation, which was related to patient prognosis. Within the treatment modalities (surgery combined with radiotherapy and radiotherapy alone), neither nodal status nor HPV were prognostic indicators.

Conclusion: Since a substantial percentage of TSCCs is HPV-positive and metastasizes to cervical lymph nodes in less advanced primary tumors, the N-status is an unreliable prognostic indicator in TSCCs. HPV is only prognostically relevant in the total tumor population, but loses its value within patient groups receiving a single treatment modality. The value of HPV for prognosis of patients with TSCC requires further study.

Key words: Human papillomavirus, tonsil, carcinoma, nodal involvement, treatment, survival

2.2. INTRODUCTION

During the past decades numerous investigations have shown an etiologic relationship between oncogenic or high-risk human papillomavirus (HPV) and squamous cell carcinomas of the tonsil (TSCC).¹⁻⁴ The incidence of HPV in TSCC is increasing, possibly related to changes in sexual behavior, and ranges from 20% to 80% in different studies depending on the detection approach used.⁵⁻⁸ There is increasing evidence that the pathogenesis of HPV-positive tumors is different from their HPV-negative counterparts, which is confirmed by molecular and clinical differences between the two subgroups. Therefore, a different clinical approach may be advisable for both TSCC groups.^{2-4,7,9} Weinberger et al. proposed a model for the development of HPV-positive carcinomas, in which three classes are differentiated and divided into two arms.⁹ In the first arm, tumorigenesis is induced by alcohol consumption and/or smoking tobacco (class I). In this group, HPV superinfection of the tumor site may occur, resulting in an HPV-positive tumor with biological features that resemble alcohol/tobacco-related tumors (class II). In the second arm, the inducer of tumorigenesis is HPV (class III), independent of smoking tobacco and alcohol consumption. Many studies suggest a better prognosis for patients with HPV-associated head and neck carcinomas,⁷ as a consequence of an improved response to treatment when compared to HPV-negative tumors.^{10,11} Therefore HPV testing has recently been proposed to be included in the standard diagnostic work-up for oropharyngeal carcinomas.^{8,10}

Another factor influencing prognosis is tumor spread to cervical lymph nodes. For head and neck tumors in general, a positive N-status is considered the most reliable prognostic marker for an unfavorable prognosis.¹²⁻¹⁴ However, TSCC literature is not consistent in confirming the prognostic value of N-status in TSCC.¹⁵⁻¹⁷ The present study aims to examine the prognostic value of N-status in a series of 81 TSCC, while also taking into consideration the HPV-status, clinicopathological features (age, gender, TNM-classification, tumor differentiation grade, smoking tobacco and alcohol consumption), as well as treatment of the tumor.

2.3. MATERIALS AND METHODS

2.3.1. Tumor material and patient data

The study population consists of 81 TSCC patients (mean age 58,9 yrs; range 39-87 yrs; 73% male)², diagnosed between 1992 and 2001 at the Maastricht university Medical Centre. The formaldehyde-fixed, paraffin-embedded archival biopsy and resection materials of these patients had been classified by histopathology at the Department of Pathology, University Hospital Maastricht, The Netherlands and analyzed for the presence of oncogenic HPV16 DNA by means of polymerase chain reaction (PCR), fluorescence in situ hybridisation (FISH) as well as p16 immunostaining.¹⁷ Data on age, gender, TNM classification, tumor differentiation grade, smoking habits, amount of

alcohol consumption, treatment modality, and follow-up were collected from the head and neck tumor database of our institute and from reviewing clinical, pathological, radiological and surgical reports. Tumors of patients treated before 1997 were re-classified according to the UICC classification. Data on tumor differentiation grade were unavailable for three patients.

Patients were classified as smokers (> 1 cigarette, pipe, and/or cigar per day) (n=69) or non-smokers (n=12). The latter group consisted of patients who had never smoked (n=10) and former smokers (n=2), who had stopped smoking more than 10 years before diagnosis of TSCC. Patients were also classified as drinkers (consumption of > 2 whiskey equivalents (~10g alcohol) per day) or non-drinkers (0-2 whiskey equivalents per day). All patients were treated irrespective of their HPV status by multimodal regimens including local resection (LR), combined resection (CR): neck dissection plus local resection, i.e. tonsillectomy or combined mandibular operation (command-procedure), radiotherapy (RT), local resection plus radiotherapy (LR/RT), combined resection plus radiotherapy (CR/RT), chemotherapy (ChT) or chemotherapy in combination with previous other treatment modalities.

Patients with tumors feasible for resection without unacceptable compromising organ functionality were treated with radical resection and elective neck dissection and postoperative radiotherapy if indicated. In patients who were primarily treated with radiotherapy the neck was also treated with radiotherapy.

During multidisciplinary counseling, treatment plans were designed dependent on tumor size, neck staging, presence of distant metastases, feasibility of surgery, histopathology of resection specimens, clinical condition and comorbidities. With the exception of microcarcinomas, elective treatment of the neck was performed routinely, including in the N0 neck, because of the high incidence of occult metastases in TSCC.^{14, 18-20}

For patients treated with radiotherapy alone, histopathological data from surgical neck dissection were consequently unavailable. Therefore clinical staging (including panendoscopy, magnetic resonance imaging of the neck, ultrasound with fine-needle aspiration, and X-thorax) was used in this study. In the CR/RT group, there was one patient with a N0 neck according to diagnostic work-up, where a positive lymph node was observed in histopathological analysis after ipsilateral elective neck dissection. This patient was considered as N0 in this study.

Follow-up data were collected with a minimum of 5 years after treatment for all patients.

The investigation was conducted in accordance with the declaration with the 18th meeting of the World Medical Association in Helsinki 1964 and the subsequent revisions. The study protocol was approved by the institutional ethical committee. Written consent was obtained from all the patients included in this study.

2.3.2. Statistical analysis

The association of N-status and HPV status with other factors associated with prognosis, including age at time of diagnosis, gender, TNM-status, tumor differentiation grade,

smoking habits and amount of alcohol consumption, were analyzed by cross-tabulations using the 2-tailed χ^2 -test or 2-tailed Fisher exact test adhering to a significance level of $p < 0.05$.

Survival curves for disease-free survival (DSF), disease-specific survival (DSS) and overall survival (OS), were calculated using the Kaplan-Meier method.²¹ Five-year survival was calculated from the date of diagnosis until death or until discharge from follow-up. DFS was calculated from the date of diagnosis until the date of recurrence (local, regional or distant). Patients without recurrence were censored at the date of the last follow-up or the date of death. Differences between survival times was determined by the log rank test in univariate analyses (significance level of $p < 0.05$).²¹ Four patients initially presented with distant metastases were excluded from the survival analysis.

Multivariate analyses were performed using the Cox proportional hazards model. Variables included were HPV, smoking tobacco, alcohol consumption and T-classification. The inclusion of treatment modality as a variable depended on the number of patients within each treatment group. Variables remained in the model if their p-values were below 0.10.²

SPSS Base System version 12.0 was used for the statistical analysis.

2.4. RESULTS

2.4.1. N-status and HPV-status in relation to patient characteristics.

A positive N-status, as determined in the diagnostic work-up by the clinical investigation and radiology, was more frequently observed in non- and former-smokers ($p=0.048$). Other clinico-pathological factors (age, alcohol consumption, TNM-stage and tumor differentiation grade) and HPV-status did not appear to be associated with N-status (Table 1).

HPV-status, however, correlated with a poor tumor differentiation grade ($p=0.015$), as did less or no smoking of tobacco ($p=0.011$) and alcohol consumption ($p=0.003$) (Table 1). Primary tumor size at time of presentation was found to be significantly smaller in the HPV-positive group than in the HPV-negative group ($p = 0.041$) in spite of comparable frequencies of nodal involvement in both groups. Moreover, in case of nodal involvement, a swelling in the neck was the main reason for visiting the ENT-outpatient department for 15 of the 24 patients with HPV-positive TSCC, compared to only 6 of the 30 patients with HPV-negative TSCC ($p=0.001$).

2.4.2 N-status and HPV-status in relation to patient treatment.

The different treatment modalities for TSCC are listed in Table 2. The two largest patient groups receiving a single treatment modality included thirty patients treated with combined surgery and radiotherapy (CR/RT) and 30 patients received radiotherapy (RT). The other treatment modalities were not taken into consideration in this analysis because of the limited number of patients in these groups. Functional resectable tumors

were preferably treated with CR/RT, whereas patients with functional irresectable tumors or other contra-indications for surgery were treated with RT alone. T-status was significantly lower in the CR/RT group than in the RT-group ($p=0.001$), whereas nodal involvement did not differ. HPV-positive TSCC were more often treated with CR/RT ($p=0.035$), whereas HPV-negative TSCC were more often treated with RT ($p=0.048$). In the CR/RT group, all HPV-positive patients had a positive N-status whereas in HPV-negative patients significantly less nodal involvement was present ($p=0.003$). Patients treated with CR/RT (resectable tumors) smoked significantly less than patients treated with RT ($p=0.001$). Smoking habits observed in both treatment modality groups were HPV-dependent. Nine of the 16 HPV-positive patients who were treated with CR/RT were nonsmokers, whereas the eight HPV-positive patients treated with RT (irresectable tumors) all smoked ($p=0.007$). In patients with HPV-negative tumors, no significant differences in smoking habits were found between the treatment modality groups. Alcohol consumption did not differ between treatment modality groups. Patients treated with CR/RT had a far more favorable 5-year overall, disease-specific and disease-free survival compared to patients treated with RT (log rank, $p<0.001$) (Figure 3: A-C). This outcome based on patient treatment did not differ between the HPV-positive and HPV-negative patients with TSCC (OS, DSS, log rank $p<0.001$; DFS, log rank $p=0.001$). Moreover, in both treatment modality groups, there were no differences noticed for the development of local and regional recurrence or for the development of distant metastases between HPV-positive and HPV-negative patients with TSS (Fisher exact test).

2.4.3. N-status and HPV-status in relation to outcome.

Nodal involvement did not correlate with survival in either the entire group of TSCC (Fig. 1A–C) or in the treatment subgroups CR/RT and RT. Remarkably, a trend for a favorable 5-year DFS rate was observed in patients with a lymph node metastasis ($P=0.067$) (Fig. 1C).

A statistically significant difference between the prognosis of HPV-positive and HPV-negative TSCCs was not found in the univariate analysis. However, there was a trend for a better DSS in the HPV-positive group (log rank, $p=0.094$) (Figure 2: A-C). Also within the treatment sub-groups CR/RT and RT, HPV-status proved to be an unreliable prognostic indicator.

The influence of N-status on prognosis was also analyzed in patients with HPV-positive and HPV-negative TSCC. In the HPV-negative group, the presence of nodal involvement seemed to be related to an unfavorable 5-year OS, DSS and DFS (Fig. 3). In the HPV-positive group, however, the presence of nodal involvement seemed to be related with a better OS, DSS and DFS.

In multivariate analysis neither N-status, nor gender, age and tumor differentiation grade, appeared to have a statistically significant influence on survival. For HPV-negative TSCCs a 2 times higher risk of cancer death was found (95% confidence interval (CI) = 0.9–4.2) compared to patients with HPV-positive tumors. Patients with a tumor staged T3

or T4 had a 2.6 times increased risk of cancer death (95% CI = 1.4-4.9) compared to patients with tumor staged T1 or T2. However, the strongest prognostic factor was smoking: smokers had a 5.5 fold higher risk (95% CI = 1.3-23.6) of dying from cancer when compared to non-smokers.¹⁷

Multivariate analysis within treatment modality groups CR/RT and RT were not performed, due to the limited number of patients in each of these groups (n=30).

2.5. DISCUSSION

In head and neck squamous cell carcinoma (HNSCC), N-status is known to be an important prognostic factor.²²⁻²⁷ Nodal involvement reduces survival by more than 50% in patients with HNSCCs.²²⁻²⁷ However in recent years, the prognostic value of nodal involvement in TSCC is becoming increasingly controversial.¹⁵⁻¹⁷ A possible explanation for this finding is the heterogeneity in etiological factors underlying tumorigenesis in different head and neck mucosa areas. For example, HPV appears to play a much more prominent etiological role in TSCC than in other head and neck tumors.^{28,29} Moreover, the incidence of HPV in TSCC has increased substantially in the past few decades.⁵⁻⁸ Therefore, investigation into whether the presence of HPV underlies the decreased prognostic value of nodal involvement is warranted.

In our study we noticed that the only clinical parameter associated with a positive neck status in TSCCs was absence of tobacco smoking. This parameter, however, was found to correlate much stronger with the presence of HPV in the tumor (41% of cases). This observation has also been described in other studies.⁷⁻⁹ Additional parameters correlating to HPV included less or no alcohol consumption, a poor tumor differentiation grade and a smaller T-stage. The latter finding suggests a different tumor biology for HPV-positive TSCCs with regional spreading occurring at smaller primary tumor sizes. Possibly, HPV-positive tumors were detected in smaller T-stages because of an earlier clinical presentation of a lymph node metastasis in this group. We observed a relationship between a swelling in the neck as presenting symptom and the presence of HPV in the primary tumor. This has also been described in literature.¹⁷ This pattern of regional tumor spreading in smaller primary tumor sizes may contribute to an earlier detection of a so far “unknown” primary tumor at a smaller size. According to this hypothesis, the tumor biology of HPV-positive tumors would result in the detection of tumors with smaller T-stages.

In this study patients with tumors feasible for resection with respect to organ functionality were treated with CR/RT, and (functionally) inoperable tumors were treated with RT. As a consequence, the CR/RT group showed a significantly better outcome. Because of their smaller primary tumor sizes, HPV-positive TSCCs were more often feasible for resection and subsequently more often treated with CR/RT. We would like to stress that conclusions with respect to the efficacy of these treatment modalities should not be extracted from the data presented here.

To what extent patient-dependent factors as life-style and co-morbidities influence clinical choice and treatment outcome remains to be studied. A favorable performance status appears to be related to HPV-positive tumors.¹¹ The inverse relation between tobacco smoking, alcohol consumption and HPV-status, suggests also that the difference in life-style may result in a decreased prevalence of co-morbidities in the HPV-positive tumor population.

N-status was not found to be of prognostic value in TSCC. Patients with HPV-negative TSCC were found to have a 2 times greater risk of cancer death. Separate analysis of the CR/RT- and RT did not indicate that these two parameters had an effect on prognosis. HPV-positive tumors, thus, show no significant improvement of response to therapy within the different modality groups. Although this could be caused by the relatively limited number of patients in the different treatment groups, a recent study also showed no significantly favorable prognosis of HPV-positive oropharyngeal tumors when treated with combined radio/chemotherapy.¹¹ Multiple hypotheses for a better outcome of HPV-positive tumors have been put forward. They are all based on factors related to therapy-outcome: absence of field cancerization as a consequence of the inverse relation of HPV and tobacco smoking and alcohol consumption, an intact apoptotic tumor response to radiation and an immune surveillance to viral-specific tumor antigens.³⁰⁻³³ However, in most cases multimodal treatment modalities have been used.

In our study, the presence of nodal involvement in HPV-negative TSCC, seemed to expose a negative influence on the prognosis. However in HPV-positive TSCC, nodal involvement even appeared to ameliorate outcome. This suggests that the tumor biology of HPV-positive TSCC is not only different from the HPV-negative TSCC but also has a great influence on the clinical presentation and outcome. As mentioned, nodal involvement in HPV-positive TSCCs is often the presenting symptom and seems to indicate the presence of a smaller primary tumor in HPV-positive TSCCs (squealer node in unknown primary tumors). Subsequently, these HPV-positive TSCCs are more feasible for a radical therapeutic approach. This indicates that the outcome of HPV-positive tumors is not only dependent on a better response to different (multimodal) treatment modalities, but more importantly, the presence of HPV in TSCCs seems to determine the choice of treatment as a result of its biology. As a consequence of the controversial prognostic value of nodal involvement, we advise the implementation of testing HPV diagnostically to stratify in TSCC tumor staging.

2.6. CONCLUSION

HPV-positive tumors, which are associated with less smoking and alcohol, have a different tumor biology. They have smaller primary tumor sizes while regional lymph node involvement is comparable to HPV-negative tumors. Our data indicate that the relatively favorable prognosis of HPV-related TSCC is determined by the choice

of treatment as a result of its biology. The prognostic value of nodal involvement is reduced by the presence of HPV. HPV-testing in the diagnostic work-up is therefore advised in TSCC tumor staging.

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Figure 1. A) N-status-dependent 5-year Overall Survival, B) N-status-dependent 5-year Disease Specific Survival, C) N-status-dependent 5-year Disease Free Survival.

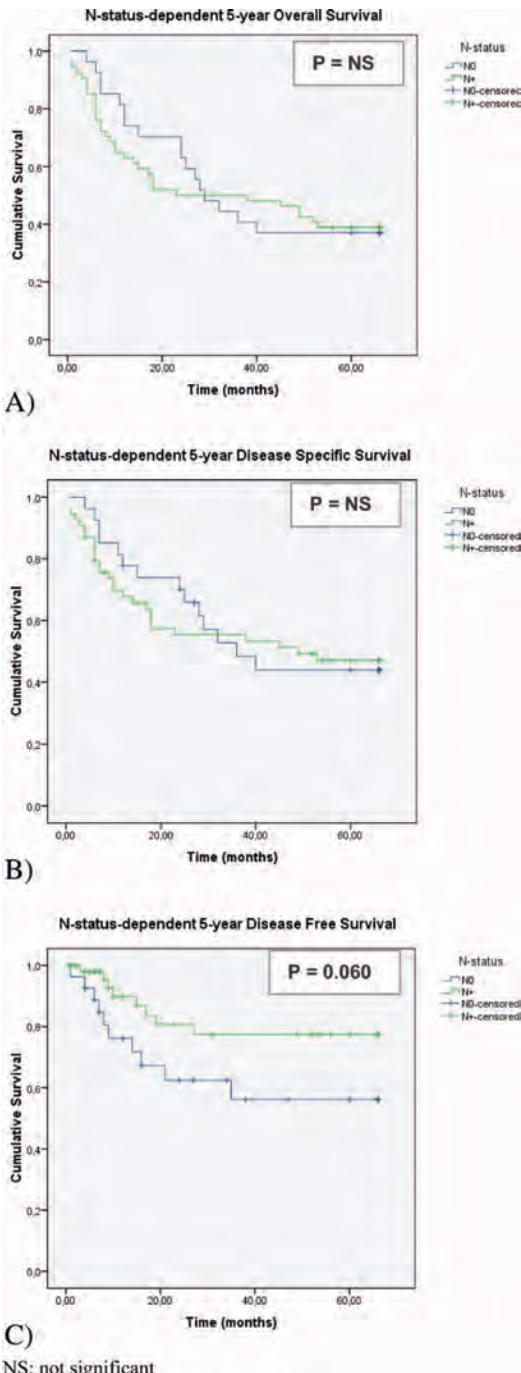
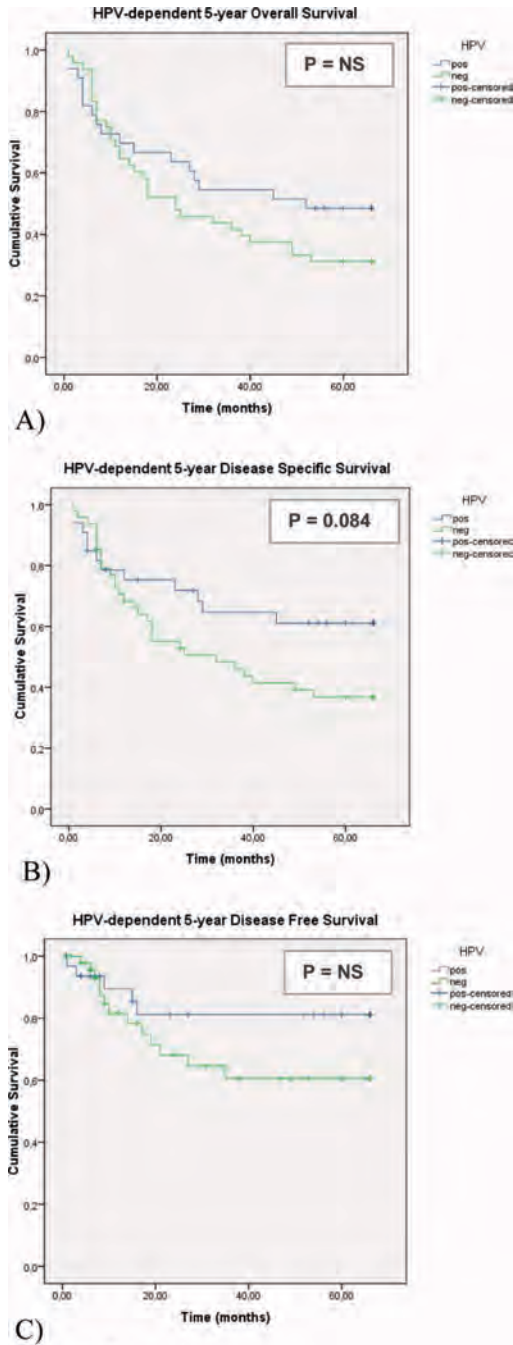


Figure 2. A) HPV-dependent 5-year Overall Survival, B) HPV-dependent 5-year Disease Specific Survival, C) HPV-dependent 5-year Disease Free Survival.



NS: not significant

Figure 3. A) N-status-dependent 5-year Disease Specific Survival in patients with HPV-positive TSCCs and B) N-status-dependent 5-year Disease Specific Survival in patients with HPV-negative TSCCs.

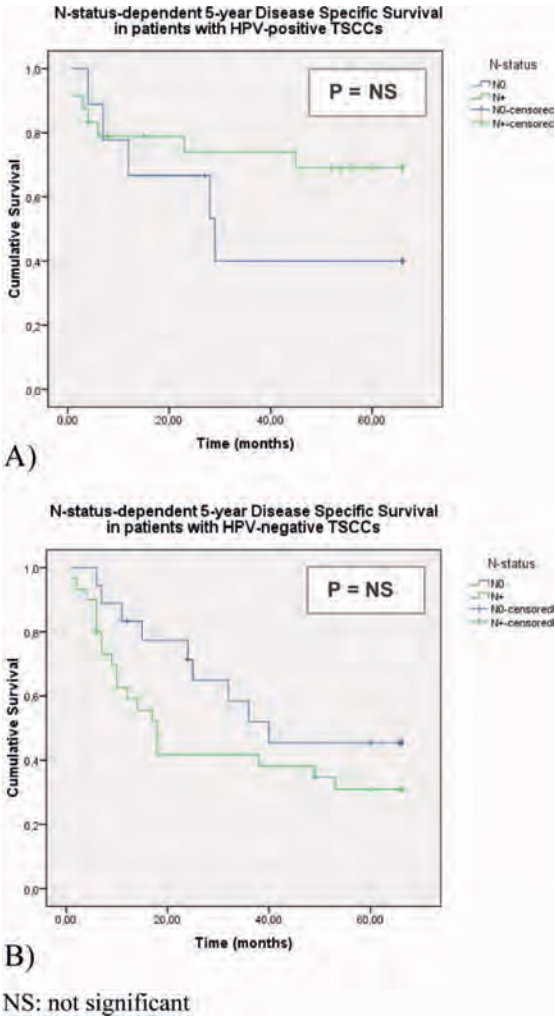


Table 1. N-status, HPV-status and treatment modality in correlation to patient characteristics.

Patient Characteristics	N0	N+	P Value	HPV+	HPV-	P Value	CR/RT	RT	P Value
Male - female	19 - 8	38 - 16	NS	24 - 9	31 - 15	NS	20 - 10	24 - 6	NS
Mean age, yr	56.9	59.1	NS	59.3	57.85	NS	58.53	56.4	
Age range, yr	41-77	39-87		39-79	41-87		39-74	44-74	
Smoking									
Nonsmoker/former smoker >10yrs.	2	14	.048	11	5	.011	11	1	.001
Smoker	25	40		22	43		19	29	
Alcohol									
None/social	12	26	NS	22	16	.003	14	13	NS
>2 U/day	15	28		11	32		16	17	
TNM									
T1, 2 - T3, 4	12 - 15	26 - 28	NS	20 - 13	18 - 30	.041	21 - 9	8 - 22	.001
T1	3	6		3	6		4	2	
T2	9	20		17	12		17	6	
T3	12	15		9	18		8	11	
T4	3	13		4	12		1	11	
N0/N+				9 - 24	18 - 30	NS	6 - 24	11 - 19	NS
N0				9	18		6	11	
N1				6	11		7	7	
N2				16	14		17	7	
N3				2	5		0	5	
M+	0	4	NS	1	3	NS			

Table 1. Continued.

Patient Characteristics	N0	N+	P Value	HPV+	HPV-	P Value	CR/RT	RT	P Value
Tumor differentiation grade									
Poor	7	22	NS	17	12	.015	11	10	NS
Moderate/well	20	29		15	34		18	20	
HPV16 status									
Positive	9	24	NS				16	8	.035
Negative	18	30					14	22	
Local recurrence	8	2	.002*	2	8	NS	0	6	0,024*
Regional recurrence	3	2	NS	1	4	NS	2	2	NS
Distant metastasis	1	3	NS	2	2	NS	1	3	NS
Total: 81 patients	27	54		33	48		30	30	

P-values were obtained using 2-tailed χ^2 -test

*asterisks indicate the use of Fisher exact test

Abbreviations. CR/RT: patients treated with tumor surgery and neck dissection in combination with radiotherapy; RT: patients treated with radiotherapy;

NS: no significance detected (p-value > 0.05).

Table 2. HPV in correlation with treatment.

Treatment	HPV+	HPV–	Total
Local resection (LR)	0	1	1
Combined resection (CR)	3	3	6
LR/RT	3	2	5
CR/RT	16	14	30
Radiotherapy (RT)	8	22	30
Chemotherapy (ChT)	2	1	3
CR/RT/ChT	1	0	1
RT/ChT	0	3	3
None	0	2	2
Total	33	48	81



3

P16^{INK4A} immunostaining is a strong indicator for high-risk-HPV-associated oropharyngeal carcinomas and dysplasias, but is unreliable to predict low-risk-HPV-infection in head and neck papillomas and laryngeal dysplasias.

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3.1. ABSTRACT

Human papillomavirus (HPV) is a risk factor for the development of benign and malignant mucosal head and neck lesions. P16^{INK4A} is often used as a surrogate marker for HPV-infection, although there is still controversy with respect its reliability. Our aim was to determine if p16^{INK4A} overexpression can accurately predict both high-risk and low-risk-HPV-presence in (pre)malignant and benign head and neck lesions. P16^{INK4A} immunohistochemistry was performed on paraffin-embedded tissue sections of 162 oropharyngeal squamous cell carcinomas (OPSCC), 14 tonsillar and 23 laryngeal dysplasias, and 20 tonsillar and 27 laryngeal papillomas. PCR, enzyme-immunoassay and FISH analysis were used to assess HPV-presence and type. Of the 162 OPSCC and 14 tonsillar dysplasias, 51 (31%) and 10 (71%) were HPV16-positive, respectively. All tonsillar papillomas were HPV-negative and four laryngeal dysplasias and 26 laryngeal papillomas were positive for HPV6 or –11. P16^{INK4A} immunohistochemistry revealed a strong nuclear and cytoplasmic staining in 50 out of 51 HPV16-positive and 5 out of 111 HPV-negative OPSCC ($p < 0.0001$) and in all HPV16-positive tonsillar dysplasias, whereas highly variable staining patterns were detected in the papillomas and laryngeal dysplasias, irrespective of the HPV-status. In addition, the latter lesions generally showed a higher nuclear than cytoplasmic p16^{INK4A} immunostaining intensity. In conclusion, our data show that strong nuclear and cytoplasmic p16^{INK4A} overexpression is a reliable surrogate indicator for HPV16 in OPSCC and (adjacent) dysplasias. For HPV6 or –11-positive and HPV-negative benign and premalignant lesions of the tonsil and larynx, however, p16^{INK4A} immunostaining is highly variable and cannot be recommended to predict HPV-presence.

Keywords: human papillomavirus, immunohistochemistry, FISH, PCR

3.2. INTRODUCTION

Human papillomavirus (HPV)-infections may lead to the development of head and neck mucosal lesions. High-risk (HR)-HPV16 is involved in the carcinogenesis of oropharyngeal squamous cell carcinomas (OPSCC) and is an indicator of favorable prognosis, independent of the applied treatment modality. The reported incidence of HPV16 in OPSCC ranges from 25% to 93% in different studies, and appears to have increased during the last decade. Also the incidence of OPSCC and its proportion within the total head and neck squamous cell carcinoma (HNSCC) population are increasing.^{1,2} Low-risk (LR)-HPV6 and -11 especially play a role in the development of benign laryngeal lesions. HPV-positivity may help in these lesions to predict a higher risk for recurrence and a lower risk for malignant progression in relation to HPV-negative, smoking-induced laryngeal lesions.³

In general, two methods are used for the detection of HPV-DNA in clinical diagnosis and biomedical research, i.e., polymerase chain reaction (PCR)-based methods and fluorescence in situ hybridization (FISH). PCR is a highly sensitive method, but may not be able to distinguish biologically and clinically relevant HPV-containing lesions from cases with an extracellular virus contamination. Although FISH is a less sensitive method, it is highly specific and has the advantage to visualise the virus-DNA in situ and allows the distinction between an episomal and integrated status. Because both techniques are relatively costly, technically demanding and requiring sophisticated laboratory facilities as well as experienced personnel,⁴ practical alternatives or complementary procedures for HPV-testing have been explored, including the immunohistochemical detection of p16^{INK4A}.⁵

HPV-positive carcinomas are characterized by overexpression of viral oncoproteins E6 and E7 leading to inactivation of p53 and the retinoblastoma protein (pRb), respectively, thereby inducing cell cycle deregulation and inhibition of apoptosis. As a result, cyclin-dependent kinase (CDK) inhibitors, including p16^{INK4A} and p21Cip1/WAF1, and the MDM2 inhibitor p14ARF are generally upregulated, which subsequently leads to downregulation of cyclin D1 and inhibition of its complex formation with CDK4.6-9 Of these HPV-associated changes in protein expression, p16^{INK4A} overexpression has been reported to be the most reliable surrogate marker for the presence of HR-HPV in OPSCC.^{6, 10-12} In contrast, HPV-negative carcinomas, induced by smoking and alcohol consumption, are generally characterized by inactivation of the p16^{INK4A} gene through loss of 9p21. Therefore, the choice of p16^{INK4A} immunostaining to distinguish HPV-positive and -negative malignancies is most obvious.^{6, 7, 13, 14}

P16^{INK4A} immunostaining has long been identified as an objective biomarker for HR-HPV-positive (pre)malignancies of the uterine cervix, allowing the unambiguous identification of truly dysplastic cells in biopsies, thereby reducing interobserver disagreement and improving diagnostic specificity.^{6, 15, 16} In contrast, normal cervical epithelium and inflammatory or metaplastic lesions are usually p16^{INK4A} negative, while genital lesions containing LR-HPV6 or -11 often show only weak p16^{INK4A} immunostaining.¹⁴

¹⁷⁻¹⁹ Moreover, several studies reported a strong correlation between p16^{INK4A} immunostaining intensity and frequency of positive cells with increasing severity of the cervical lesion.^{18, 20-22} The College of American Pathologists and the American Society for Colposcopy and Cervical Pathology has recently published their recommendations on using p16^{INK4A} immunostaining as a biomarker in lower anogenital squamous lesions, introducing the concept of “block-positivity,” defined as continuous strong nuclear or nuclear plus cytoplasmic p16^{INK4A} staining of the basal cell layer with upward extension. The authors state that this scoring method allows a categorization of precancerous disease when the differential diagnosis is between a premalignancy [-intraepithelial neoplasia (IN) 2 or -IN 3] and a mimic of a premalignancy known to be not related to neoplastic risk. Furthermore, it can be used when -IN 2 is in the differential diagnosis to distinguish between a premalignant disease and a low-grade lesion/a non-HPV-associated pathology. As such, the method is used as an adjudication tool for cases in which there is a professional disagreement in histologic specimen interpretation.²³ Despite the fact that p16^{INK4A} immunohistochemistry is a low-cost test that can easily be implemented in daily diagnostic practice, its reliability as a surrogate marker for the presence of HR-HPV has been subject of debate.^{18, 24} HPV-negative but p16^{INK4A} positive cases, although small in number, have been described in many series. It has been suggested that this p16^{INK4A} positivity results from pRb inactivation by means of mechanisms independent of HPV-infection.^{4-9, 25, 26} Also the lack of standardization of p16^{INK4A} immunohistochemistry and its evaluation criteria may be a source of misinterpretations. The foregoing arguments have been a reason for hesitation to introduce the p16^{INK4A} test into the clinic.¹⁵

Although numerous studies analyzed the expression of p16^{INK4A} in HR- and LR-HPV-associated cervical premalignancies, these data are scarce for premalignant and benign head and neck lesions, and therefore the aim of the present study was to determine the usefulness of this biomarker in neoplasms of the oropharynx and larynx. For this purpose we analysed a series of 162 OPSCC as well as a range of tonsillar and laryngeal dysplasias and papillomas, part of which was expected to harbor HR- and LR-HPV-types.

3.3. MATERIAL AND METHODS

3.3.1. Tumor material and patient data

Biopsy and resection material was obtained from 227 patients resulting in 246 samples, taken at different moments in time. Since our main goal was to evaluate the potential of p16^{INK4A} immunostaining patterns as a surrogate marker for HPV in daily routine diagnostic practice, i.e. , in every single biopsy sample a pathologist receives from the clinic, we present the analyses in the total number of samples in the main text (Table 1 and Figs. 1a and 1b). For comparison we have included the analyses per patient in Supporting Information Table 1 and Supporting Information Figs. 1a and 1b.

Formaldehyde-fixed and paraffin-embedded tissues were selected from the archives of the Departments of Pathology of the University of Cologne, Germany and the Maastricht University Medical Centre, The Netherlands. Patients were diagnosed between 1986 and 2007.

Table 1 summarizes the clinicopathological features of the cases used in this study. Patient age at first diagnosis ranged from 39–87 years (mean 60) in the OPSCC group with no significant difference between the HPV-positive and the HPV-negative cases. In the papilloma group, the age ranged from 1 to 72 years (mean 43) and in the dysplasia group from 36 to 82 years (mean 62). The specimens included 162 OPSCC (from 162 patients) and 84 benign and premalignant samples (from 65 patients), i.e., 14 tonsillar dysplasias from 12 patients, 13 of which were adjacent to an invasive carcinoma, 20 tonsillar papillomas (from 20 patients), 23 laryngeal dysplasias (from 19 patients) and 27 laryngeal papillomas (from 14 patients).

A correlation of the HPV16 status with the disease-free survival (DFS) data with a maximum follow-up time up to 180 months and a mean of 36 months revealed that the after 5 years was 55% for patients with an HPV-negative carcinoma and 74% for patients with an HPV-positive carcinoma (Hazard ratio = 0.4; 95% Confidence Interval = 0.2–0.8), indicating that we used a representative OPSCC patient group.

The OPSCC and dysplasias were histologically classified according to the criteria of the World Health Organization (C.J.P.-K. and H.P.D.).²⁷ Eleven out of 14 tonsillar dysplasias were graded as severe and three as moderate. From the laryngeal dysplasias six were graded as mild, eight as moderate and nine cases as severe (see Table 1).

3.3.2. P16^{INK4A} immunohistochemistry

Immunostaining of p16^{INK4A} was performed on 4-μm-thick formaldehyde fixed, paraffin-embedded tissue sections, which were deparaffinized using xylol. As a primary p16^{INK4A} antibody clone E6H4 (Roche, Almere, The Netherlands) was used and detected using Powervision (DAKO A/S, Glostrup, Denmark) and peroxidase-DAB visualisation, as previously described.⁷

Three independent observers (J.J.M., S.E.G., E.-J.S.) performed evaluations of the immunostained samples and consensus was acquired about the scores.

P16^{INK4A} immunostaining patterns were first scored on a semiquantative scale, using the following evaluation criteria: Score 0 indicates that the majority of the cells are negative and that only up to 5% of the cells show nuclear immunoreactivity with or without cytoplasmic staining; Score 1 indicates that 5–10% of the cells show a nuclear reactivity with or without cytoplasmic positivity; Score 2 indicates a focal staining pattern with 10–25% of the cells showing nuclear reactivity with or without cytoplasmic positivity; Score 3 indicates that >25% cells show a strong nuclear staining reaction with or without cytoplasmic positivity.¹⁴

The proportion of positive cells is the primary parameter, with the intensity as a secondary scoring parameter. All cases scored as 2 and 3 showed a strong immunostaining intensity, while few cases scored as 1 stained less intensely positive.

In addition, we scored the p16^{INK4A} staining patterns according to the “block-type” immunopositivity approach, defined as p16^{INK4A} only being block positive if continuous (>70%) strong nuclear with or without cytoplasmic staining is present (in all head and neck lesions) and staining is observed in the basal cell layer with extension upwards (in the benign and premalignant lesions).²³ Controls included sections of uterine cervical IN grade 2 and 3 (positive controls) and sections from the same cases on which the p16^{INK4A} antibody was substituted by a monoclonal mouse IgG2a antibody to *Aspergillus Niger* or by buffer without a primary antibody (negative controls).

3.3.3. DNA isolation and HPV typing by PCR

For DNA isolation, tissues were processed with the QIAamp Tissue Kit (Qiagen, Hilden, Germany) according to the manufacturers’ instructions. Total cellular DNA was eluted with 250 µl of the AE-buffer (Qiagen) and 5 µl were used in each of the PCR analyses.

To test the quantity and quality of the DNA samples and to demonstrate that the samples were free from inhibitory substances, PCR was performed for the β-Globin gene, resulting in a 268 bp PCO4/GH20 PCR product.²⁸ As negative controls water and as positive controls human placental DNA were included in each PCR run.

HPV sequences were detected by highly sensitive group-specific nested PCR protocols with degenerate primers A5/A10 and A6/A8 for HPV as previously described.²⁹ PCR products (5 µl) were separated on 2% agarose gels and visualized by ethidium bromide staining. For HPV typing internal biotinylated A6/A8-PCR products (270 bp) were hybridized with 37 type-specific digoxigenin-labeled oligonucleotide probes in an enzyme-immunoassay as described earlier.^{30, 31}

3.3.4. Detection of HPV6-, 11-, 16- and 18 by FISH

FISH was performed on 4-µm thick formaldehyde fixed and paraffin-embedded tissue sections tissue as described previously.³²⁻³⁴ Briefly, sections were deparaffinized, pre-treated with 85% formic acid/0.3% H₂O₂, 1 M NaSCN and 4 mg/ml pepsin, post-fixed in 1% formaldehyde in PBS, dehydrated in an ethanol series and hybridized with digoxigenin-labeled HPV6-, 11-, 16- and 18-specific probes (PanPath, Budel, The Netherlands), respectively, depending on PCR results, according to the manufacturer’s instructions. After hybridization the preparations were washed stringently in 50% formamide, 2× SSC at 42°C (two times 5 min). The probes were detected by subsequent application of mouse anti-digoxigenin (Sigma, St. Louis, MO), peroxidase-conjugated rabbit anti-mouse and peroxidase-conjugated swine anti-rabbit (both DAKO), and visualized by a peroxidase based amplification reaction using rhodamine-labeled tyramide.^{35, 36} Preparations were mounted in Vectashield (Vector Laboratories, Burlingame, CA) containing 4,6-diamidino-2-phenyl indole (DAPI; Sigma: 0.2 g/ml). Microscope images were recorded with the Metasystems Image Pro System (black and white CCD camera; Sandhausen, Germany) mounted on top of a Leica DM-RE fluorescence microscope equipped with DAPI and rhodamine filters.

Evaluation of nuclear hybridization signals was performed by two investigators (J.J.M., E.-J.M.S.) according to the criteria described by Mooren et al.,³⁷ i.e., nuclear punctate signals were considered to indicate integrated HPV-DNA and diffuse signals to indicate episomal HPV-DNA. Controls included hybridizations on HPV6-, -11-, -16- and 18-positive formaldehyde-fixed, paraffin-embedded tissues of known human uterine cervical and head and neck (pre)malignancies. Negative controls consisted of HPV-negative tissue sections as determined by PCR and hybridizations omitting the viral probe.

3.3.5. Statistical analysis

HPV-status and p16^{INK4A} immunostaining were correlated using cross-tabulations and the two-tailed Fisher exact test and/or Chi-square test. We regarded a p-value ≤ 0.05 as level of significance. DFS in the OPSCC group was calculated from the date of diagnosis until the date of recurrence (local, regional or distant, whichever occurred first). Patients without recurrence were censored at the date of the last follow-up or the date of death. The statistical significance of differences between survival times was determined by the log rank test in univariate analysis. All statistical analyses were performed by PASW Statistics version 18.0.

3.4. RESULTS

3.4.1. HPV-status of the head and neck lesions examined in this study

The HPV-status of the OPSCC, the premalignant and the benign head and neck lesions is presented in Table 1. Only when the presence of HPV was detected by both PCR and FISH the sample was judged to be HPV-positive.

Of the 162 OPSCC 51 (31%) were found to be positive for HPV16. All 51 cases showed integrated HPV16, as concluded from the FISH-analyses. No other HPV-subtypes were detected in these lesions.

Forty of the 84 benign and premalignant lesions (48%) were HPV-positive: 10 (71%) of the 14 tonsillar dysplasias harbored HPV16; 4 (16%) of the 23 laryngeal dysplasias harbored HPV6, of which two also contained HPV11; of the 27 laryngeal papillomas 17 (63%) were positive for HPV6, and nine (33%) for HPV11. The frequency of HPV-infections in these lesions is in accordance with previous studies.³⁸⁻⁴¹ Remarkably, all 20 samples of tonsillar papillomas were HPV-negative.

In all HPV16-positive tonsillar dysplasias the virus was integrated (Fig. 2b), whereas in one sample additionally an episomal FISH pattern was detected in the cell nuclei surrounding tonsillar crypts (data not shown). In all, except one sample, of the HPV6- or 11-positive laryngeal lesions the virus was episomal (Fig. 2h).

In the Supporting Information Table 1, the HPV-positivity per patient is presented, for comparison, showing only a slightly difference in the frequency of HPV-positivity in the

tonsillar dysplasias (83% per patient vs . 71% per sample) and the laryngeal papillomas (93% vs . 96%).

3.4.2. Strong correlation between p16^{INK4A} overexpression and HR-HPV16 in OPSCC and tonsillar dysplasias

The p16^{INK4A} scores in the HPV-positive and -negative lesions, using the two different scoring methods, are shown per subgroup in Figs. 1a and 1b . The analyses per patient are presented in Supporting Information Figures 1a and 1b.

The presence of HR-HPV16 in the OPSCC was significantly associated ($p < 0.0001$) with a strong p16^{INK4A} immunostaining reaction (staining Score 3) and with a block positive p16^{INK4A} immunostaining: 50 of 51 HPV16-positive OPSCC showed strong nuclear and cytoplasmic p16^{INK4A} immunopositivity in more than 70% of the tumor cells, whereas only five of the 111 HPV16-negative tumors were p16^{INK4A} positive.

Also all 10 HPV16-positive tonsillar dysplasia samples showed strong (Score 3) p16^{INK4A} expression (Fig. 2a). In eight of these an equally strong staining intensity in the nuclei and cytoplasm could be observed in more than 70% of the dysplastic cells and these were scored as block positive. The remaining two HPV16-positive samples showed a strong (Score 3) p16^{INK4A} immunostaining reaction in 40–50% of the lesion, being block negative.

Of the 4 HPV-negative tonsillar dysplasia samples, two showed an intermediate p16^{INK4A} staining pattern (Score 2) and two samples from one patient were negative (Score 0), implying that these were all block negative.

3.4.3. Highly variable p16^{INK4A} immunostaining patterns in the tonsillar papillomas and laryngeal lesions

In the 20 HPV-negative tonsillar papillomas, the p16^{INK4A} immunostaining patterns were highly variable and ranged from a negative score to a strong staining intensity in 30–80% of the cells (Figs. 2d –2f and Fig. 1a). Only one sample was p16^{INK4A} block positive (Fig. 1b). In 12 (86%) out of the 14 p16^{INK4A} positive tonsillar papillomas, the immunostaining intensity of the nucleus was stronger than that of the cytoplasm.

Of the 23 laryngeal dysplasias, four samples (two from one patient) showed a strong, four a low to intermediate and 15 (two from one patient) no p16^{INK4A} expression (Fig. 1a). Three of the four HPV6-positive lesions, of which two samples from the same patient harbored HPV11-DNA, were p16^{INK4A} negative. The remaining HPV6-positive dysplasia sample showed a strong p16^{INK4A} immunostaining (Score 3) and was also block positive (Figs. 2c and 2g and Figs. 1a and 1b). There was no association between p16^{INK4A} score and the degree of dysplasia. Although all five mild dysplasias were also p16^{INK4A} negative, the moderate and severe dysplasias showed negative to strong p16^{INK4A} staining patterns.

P16^{INK4A} staining patterns in the HPV6 and -11-positive laryngeal papillomas varied from negative to strongly positive. Five out of the 26 HPV-positive laryngeal papillomas were p16^{INK4A} block positive, three of these being from the same patient. Similar to most

tonsillar papillomas, all of the p16^{INK4A} positive laryngeal lesions showed a stronger nuclear than cytoplasmic staining intensity.

In the Supporting Information Figures 1a and 1b, the p16^{INK4A}-positivity per patient is presented, showing only minor differences in the frequencies of p16^{INK4A}-immunostaining scores, compared to those presented in Figures 1a and 1b (analyses per sample).

3.5. DISCUSSION

Overexpression of p16^{INK4A} has been put forward as a specific surrogate biomarker for the presence of HR-HPV, in particular in (pre)malignancies of the uterine cervix, as well as those in the head and neck region.^{6, 7, 13, 14} However, there still exists controversy on the reliability of p16^{INK4A} expression as indicator for the presence of HPV, particularly in the latter group. In our study, we analysed the relationship between HR- and LR-HPV-status and p16^{INK4A} immunostaining patterns in 246 samples of benign, premalignant and malignant head and neck lesions. A strong (Score 3) and block positive nuclear and cytoplasmic p16^{INK4A} staining pattern was predominantly found to correlate with HPV16-containing OPSCC and tonsillar dysplasias. In contrast, the tonsillar and laryngeal papillomas, as well as the laryngeal dysplasias showed a highly variable p16^{INK4A} staining pattern, independent of the HPV-status. By using the block-type scoring approach, however, most of these lesions were interpreted as being p16^{INK4A} block negative.

It is now well established that HPV-associated HNSCC represent a separate entity, which is clinically and molecularly distinct from its tobacco- and/or alcohol-induced counterpart.^{2, 7, 37} The upregulation of p16^{INK4A} results from inactivation of pRb by the HPV-E7 oncoprotein, or alternatively through E7-mediated epigenetic induction of KDM6B, subsequently leading to activation of the p16^{INK4A} gene.^{6, 16}

Although several studies have reported a high interobserver conformity in the evaluation of p16^{INK4A} immunostaining patterns,^{4, 14, 18, 19, 42} others have indicated limitations in this procedure. The immunohistochemical p16^{INK4A} staining procedure is subject to variations in the protocol and to difficulties in the interpretation of the staining patterns as a result of different scoring criteria.¹⁷ There is an emerging consensus that only samples showing strong nuclear and cytoplasmic p16^{INK4A} immunostaining, observed in more than 25% of tumor cells, should be considered HPV-positive.^{5, 13, 14, 18, 20-22} Recent studies even suggest to use 70% immunopositivity of the carcinoma tissue as a cut-off point.^{4, 11, 12} In our study, we indeed observed the strong and block positive p16^{INK4A} immunostaining in all except one HPV16-positive OPSCC. Despite the fact that a small fraction of the HPV-negative OPSCC did show p16^{INK4A} positivity, the correlation of this surrogate marker with HPV16-presence is highly significant.

Also in all HPV16-positive tonsillar dysplasias of the present study a strong, and in most cases block positive, p16^{INK4A} immunostaining pattern could be detected, which is in accordance with the results in cervical dysplasias.^{14, 19} Similar to OPSCC, also in these

lesions we identified p16^{INK4A} positive staining in two HPV-negative samples, albeit with an intermediate score and being block negative.

The use of p16^{INK4A} immunostaining as surrogate marker for HPV-presence in laryngeal dysplasias and head and neck papillomas, however, is unreliable. In the HPV6/11-positive laryngeal dysplasia samples the majority of cases was p16^{INK4A} negative, while 1/3 of the HPV-negative samples did show p16^{INK4A} positivity. LR-HPV-integration seldomly occurs and the only laryngeal papilloma sample harboring nuclear punctate HPV6-FISH signals only showed weak p16^{INK4A} immunostaining. In a recent study on juvenile onset laryngeal papillomatosis progressing to carcinoma in one patient, the papilloma with episomal and the carcinoma with integrated HPV6 were all p16^{INK4A} negative.⁴³

All our tonsillar papilloma samples were HPV-negative, whereas 70% of the samples showed variable p16^{INK4A} positivity, indicating that p16^{INK4A} immunostaining appears of no use for predicting HPV-presence in these lesions. This is in accordance with a previous study on tumor-free palatine tonsils, all proven to be negative for HPV, but showing p16^{INK4A} overexpression in a quarter of these normal cases.²⁵ By using the block-type scoring system, however, most tonsillar and laryngeal papillomas and laryngeal dysplasias were interpreted as being p16^{INK4A} block negative. This fits with the fact that these head and neck lesions are usually HPV-negative or contain LR-HPV types. Block-positivity is used to identify HR-HPV-containing precancerous anogenital lesions, which is in accordance with the block-positivity observed in our study in OPSCC and tonsillar dysplasias, being associated with HR-HPV.²³

Also at a subcellular level variability in p16^{INK4A} immunostaining patterns could be observed: in the HPV16-positive OPSCC and tonsillar dysplasias we predominantly found an equally strong p16^{INK4A} staining intensity in nuclei and cytoplasm, whereas in the other head and neck lesions the nuclear intensity was generally stronger than that of the cytoplasm.

The fact that studies on oral papillomas showed an HPV-positivity in 13–60%, predominantly HPV6 or –11,^{44–46} makes the absence of HPV in the tonsillar papillomas in our series remarkable. The lack of HPV in these samples might be explained by elimination of the virus by the immune system with persistence of the lesions, or by the fact that other viruses, yet to be discovered, play an etiological role in the development of the tonsillar papillomas. Also the mechanisms underlying the predilection of different HPV-types for different anatomical head and neck lesions remain to be further explored.

The highly variable p16^{INK4A} immunostaining patterns in the benign and premalignant head and neck lesions may be explained by several other factors. It is well described in the literature that in uterine cervical lesions LR-HPV6 results in a less intense p16^{INK4A} immunostaining pattern as compared to HPV16, which is caused by the fact that the affinity of the HPV6 E7 protein for cellular pRb is 10-fold lower than that of HPV16 E7.^{19, 47, 48} Therefore, it is surprising that our HPV6/11-containing benign and premalignant head and neck samples were often strongly positive, or completely negative for p16^{INK4A}.

Since a subgroup of HPV-negative lesions also shows overexpression of p16^{INK4A}, several factors have been proposed to explain this observation, which remain to be studied. These include infection with other viruses (i.e., cytomegalovirus and adenovirus), which functionally inactivate pRb in a similar fashion as the HPV-oncogene E7,^{49, 50} physiological stress, oncogene-driven senescence by functional overactivation of (proto)oncogenes including Ras, Raf, MEK and E2F or replicative senescence due to DNA-damage or oxidative stress.⁵¹⁻⁵⁴

In summary, our results indicate that a strong nuclear and cytoplasmic p16^{INK4A} immunostaining pattern can accurately predict the presence of HR-HPV16 in OPSCC and tonsillar dysplasias. Our data underscore the proposed cut-off level of 70% p16^{INK4A} positive cells, corresponding to block positive p16^{INK4A} immunopositivity, in these lesions as indicator for HR-HPV16-presence. In the other premalignant and benign head and neck lesions, however, caution is recommended when using this surrogate marker for HPV-infection.

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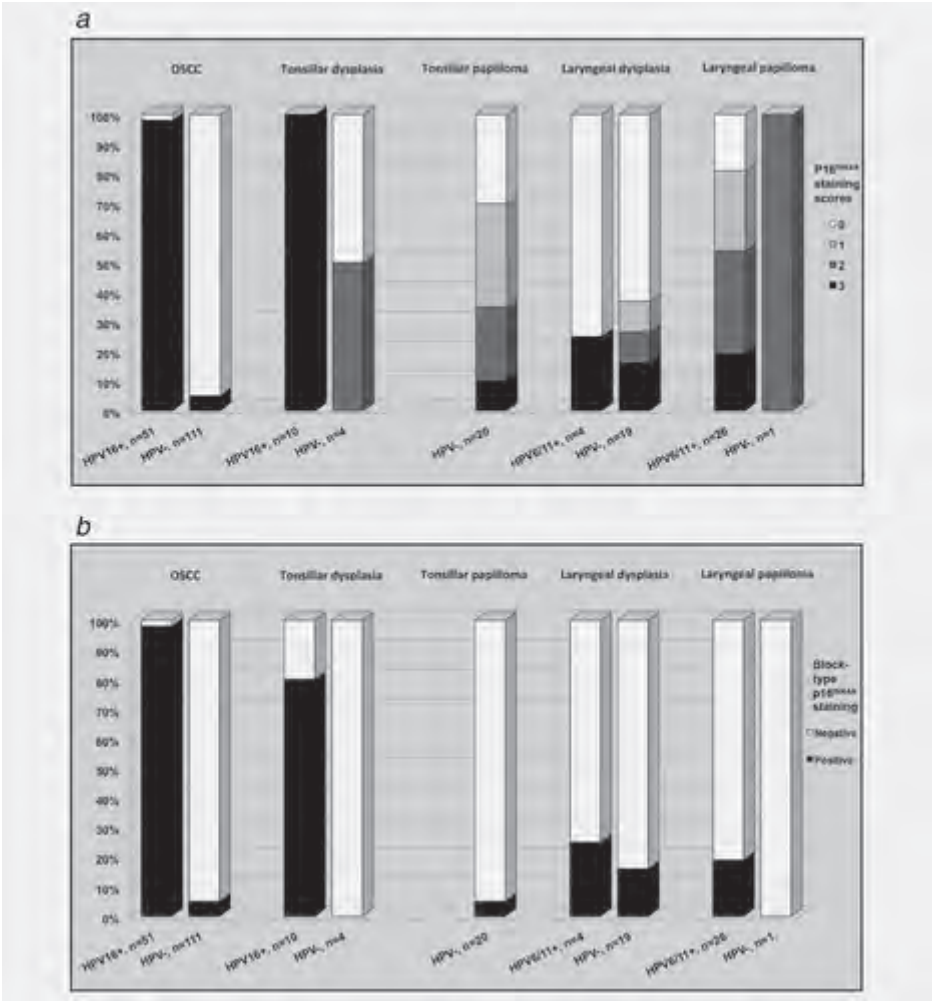
Table 1. HPV-status and clinicopathological features of all head and neck lesion studied

	OPSCC (n = 162)	Tonsillar dysplasia (n = 14)	Tonsillar papilloma (n = 20)	Laryngeal dysplasia (n = 23)	Laryngeal papilloma (n = 27)
Number of different patients	162	12	20	19	14
Gender					
Male	117 (72%)	9 (75%)	12 (60%)	18 (95%)	12 (86%)
Female	45 (28%)	3 (25%)	8 (40%)	1 (5%)	2 (14%)
Mean age at first diagnosis (range)	60 (39–87)	60 (36–79)	42 (5–66)	63 (43–82)	44 (1–72)
TNM status					
T1	23 (14%)				
T2	54 (33%)				
T3	46 (29%)				
T4	39 (24%)				
N0	51 (31%)				
N1	29 (18%)				
N2	64 (39%)				
N3	15 (9%)				
Missing	3 (3%)				
M+	4 (3%)				
Histology grade (all samples)					
Mild		0 (0%)		6 (26%)	
Moderate		3 (21%)		8 (35%)	
Severe		11 (79%)		9 (39%)	
Number of HPV-positive samples	51 (31%)	10 (71%)	0 (0%)	4 (17%)	26 (96%)
HPV-PCR (all samples)					
HPV 6				4a	17
HPV 11				2	9
HPV 16	51	10			
HPV-FISH (all samples)					
Integrated	51	9			1b
Episomal		1		4	26

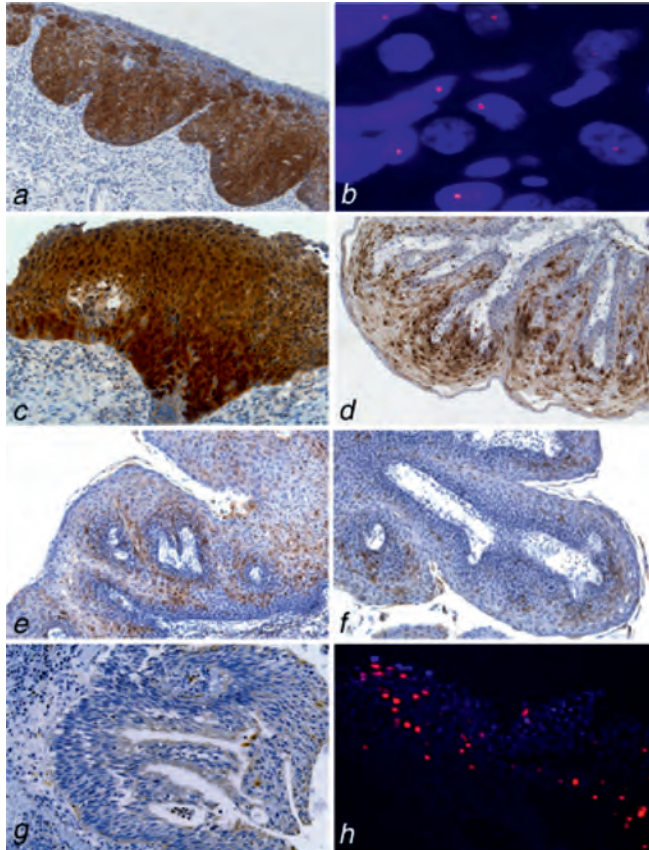
a Two out of these four HPV6-positive samples were also positive for HPV11.

b This sample showed both episomal and integrated HPV6.

Figure 1



(a) P16^{INK4A} immunostaining scores in all (pre)malignant and benign head and neck lesions. Scoring criteria: Score 0 negative; Score 1: 5–10% of the cells show a nuclear reactivity with or without cytoplasmic positivity; Score 2: focal staining pattern with 10–25% of the cells showing nuclear reactivity with or without cytoplasmic positivity; Score 3: >25% cells show a strong nuclear staining reaction with or without cytoplasmic positivity.¹⁴ (b) Immunostaining results for p16^{INK4A} in all (pre)malignant and benign head and neck lesions based on the block-type scoring approach, defined as p16^{INK4A} only being interpreted as positive if continuous (>70%) strong nuclear with or without cytoplasmic staining is present (in all head and neck lesions) and is observed in the basal cell layer with extension upwards (in the benign and premalignant lesions).²³

Figure 2

Representative examples of p16^{INK4A} immunohistochemistry (a , c–g) and HPV-specific FISH analysis (b , h) in head and neck lesions. (a) Strong nuclear and cytoplasmic p16^{INK4A} immunostaining (Score 3) and also block-positive p16^{INK4A} staining pattern in a tonsillar dysplasia; (b) Punctate signal (red) per nucleus (blue DAPI) indicating integrated HPV16 in a tonsillar dysplasia; (c) Strong nuclear and cytoplasmic p16^{INK4A} immunostaining (Score 3) and also block-positive p16^{INK4A} staining pattern in a laryngeal dysplasia sample; (d) Tonsillar papilloma sample showing strong nuclear and weaker cytoplasmic p16^{INK4A} immunostaining (Score 3), but scored as being block-negative. (e , f) Tonsillar papilloma samples showing a focal staining pattern with 10–25% of the cells showing nuclear and cytoplasmic positivity (Score 2; e), and showing 5–10% of the cells with a nuclear and cytoplasmic positivity (Score 1; f). (g) Laryngeal dysplasia sample showing 0–5% of the cells showing nuclear and cytoplasmic p16^{INK4A} immunopositivity (Score 0); (h) Diffusely (red) stained nuclei (blue DAPI) indicating episomal presence of HPV6 in a laryngeal papilloma.





Value of human papillomavirus testing in the diagnostic workup of lymph node metastases from an unknown primary tumor to the neck.

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In their article, Strojan et al¹ promote human papillomavirus (HPV) testing in patients with metastases of unknown primary tumors to the neck (CUP). This advice is merely based on retrospective studies on patients with CUP in which the HPV status of the metastasis was compared with that of the primary tumor identified during diagnostic workup, suggesting that HPV positivity is indicating an origin in the oropharynx.²⁻⁶ If the authors are right, HPV testing could improve the sensitivity of current extensive diagnostic workup protocols applied to patients with true CUP by identifying “missed” (or formerly undetected) oropharyngeal primary tumors.

However, studies on the prevalence of HPV in lymph node metastases of which the primary tumor could not be detected after diagnostic workup so-called “true” CUPs are scarce and contradictory. HPV prevalence rates range from 0% to 100% and were tested in very small sample numbers: Weiss et al⁷ 0/1 (0%), Armas et al⁸ 0/4 (0% in N3 necks; 4/4 Epstein Barr virus–positive), Begum et al⁵ 3/10 (30%), Barwad et al⁹ 9/17 (53%), Hofmann et al¹⁰ 2/3 (66%), Desai et al¹¹ 4/6 (66%), and Goldenberg et al³ 2/2 (100%). Compton et al¹² investigated HPV16 presence by DNA in situ hybridization in a cohort of resected, nonirradiated true CUPs: of 11/25 p16-positive, there were 7/25 HPV-positive (28%) tumors. However, a relationship with survival was not found. So far, no studies have been reported in which additional HPV testing has led to the discovery of the primary tumor in case the regular diagnostic workup revealed a true CUP.

In our institute, 29 true CUPs (all squamous cell carcinomas) were analyzed with no prior history of head and neck carcinoma. Diagnostic workup consisted of MRI or CT of the head and neck, CT of the thorax, and ultrasonography with fine-needle aspiration cytologic study in all patients. In 18 patients, positron emission tomography–CT of the whole body was executed. Panendoscopy was routinely performed with biopsy of the base of the tongue, epipharynx, and ipsilateral tonsillectomy. After treatment with ipsilateral neck dissection (29/29) and ipsilateral radiotherapy of the neck (27/29), without radiation of pharyngeal axis and contralateral neck (21/27), 5-year follow-up revealed no primary tumors. HPV was tested on all (nonirradiated) surgical specimens (paraffin-embedded): 5/29 were p16-positive without association with survival. However, in none of the specimens was HPV-DNA detected by fluorescence in situ hybridization and polymerase chain reaction (HPV16-DNA), or GP5 /6 PCR (0%). Therefore, in our cohort, HPV testing did not add value to the described diagnostic workup and did not influence therapeutic decision-making, for example, additional postsurgical radiotherapy of the oropharynx.

Currently, the prevalence of HPV in true CUPs seems insignificant despite the relationship of oropharyngeal viral carcinogenesis and its pattern of lymph node metastasis. The prognostic and therapeutic implications of HPV in true CUPs are therefore limited until now. Probably, most HPV-positive primary tumors of the oropharynx are detected by tonsillectomy or “blind biopsies” of the oropharyngeal region, resulting in low percentages of HPV-positive true CUPs. A prospective study is warranted to investigate the value of HPV presence in lymph node metastases of true CUPs regarding the detection of the primary oropharyngeal tumor and to explore whether de-escalation of therapy in HPV-positive true CUPs can be considered.

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Management of neck metastases of unknown primary origin united in two European centers.

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5.1. ABSTRACT

Combined analysis of diagnostic and therapeutic management of neck metastases of carcinoma of unknown primary origin ('true CUP') in two European tertiary referral centers (University Medical centers of Maastricht, NL and cologne, D) to contribute to the ongoing discussion on management in CUP. Retrospective analysis of 29 (Maastricht) and 22 (cologne) true cervical CUP syndrome patients (squamous cell carcinoma). The diagnostic and therapeutic approaches were correlated with clinical follow-up data and HPV status. In total, 48 out of 51 true CUP patients received postsurgical adjuvant radio- therapy. In eight patients from cologne, this was combined with concomitant platin-based chemotherapy. neither in cologne nor in Maastricht, radiotherapy of the pharyngeal mucosa was commonly performed (n=6, 12.5%). The percentage of patients who were irradiated ipsilaterally or bilaterally did not differ between both institutes (N= 21/27 in Maastricht vs. 11/21 in Cologne), nor did the 5-year overall survival differ significantly. Oncogenic HPV was only found in 4 out of 51 CUPs (7, 8 %). Therefore, no relation with overall and recurrence-free survival could be detected. no occult primary tumors were revealed during follow-up despite de-escalation of therapy by abandoning irradiation of the pharyngeal mucosa in both institutes. There were no significant differences between ipsilateral and bilaterally irradiated patients regarding overall and recurrence-free survival. The occurrence of distant metastases was more often noticed in ipsilaterally treated patients as compared to bilaterally radiated patients (8 vs. 2, p=0.099). Those patients all had been classified N2b or higher. International guidelines still are not unified and there is an urgent need for a consented therapeutic regimen. Comparison of two international strategies on the management of CUP patients is presented and further research is recommended regarding the role of radiotherapy of the pharyngeal axis, the value of unilateral and bilateral radiotherapy and the role of concomitant or induction chemotherapy in CUP patients, particularly in N2b or higher-staged neck disease. The prevalence and role of HPV in true CUP after thorough diagnostic work-up seem limited in our case series, particularly when compared to the role in oropharyngeal carcinomas.

Keywords cervical carcinoma of unknown primary (CUP) · neck · Diagnosis · Treatment · Outcome

5.2. INTRODUCTION

Carcinoma of unknown primary cancer (CUP) is defined as a metastatic disease with its origin remaining unknown despite extensive clinical, laboratory and imaging examinations.¹ Up to 10 % of all cervical lymph node metastases present without a known primary site.¹ The histopathology is most commonly squamous cell carcinoma, but CUP lymph nodes can also consist of adenocarcinoma, as reported in a review of 223 patients by Lee et al.²

A unified strategy for CUP management is desirable, not only to optimize diagnostics and therapy, but also to minimize side effects and finally to improve the outcome and survival of those patients. However, for CUP syndrome, no joined diagnostic or treatment strategies have yet been determined. In multiple countries, national consensus guidelines (e.g., German Oncologic Society³, Dutch national Guidelines⁴) recommend diagnosis and treatment strategies of cervical CUP. However, they vary,^{5–7} and no unified regimen has yet been established.^{1, 8, 9}

Rodel et al.¹⁰ already in 2009 called for a unified treatment strategy in cervical CUP syndrome, and outlined in a heterogeneously treated group of 58 patients that the treatment needs to be unified and optimized.

In Germany, ultrasonography of the neck, in some centers including fine needle aspiration cytology (USgFNAC) of the suspected lesion, is generally recommended.¹¹ Fluoro-deoxy-glucose positron emission tomography (FDG-PET) may be helpful to identify a primary cancer due to its enhanced glucose metabolism.^{12, 13} However, in Germany, it is not universally conducted – be it for financial and insurance aspects (high costs which are not always covered by insurances) or for availability (not every hospital has a positron tomogram). Thus, variations in diagnostic and therapeutic approaches to patients presenting with CUP syndrome may lead to differences in prognosis and outcome within one country and even between centers.

In the Netherlands, general diagnostic approaches in patients with suspected cervical CUP syndrome include ultrasonography of the neck with guided fine needle aspiration cytology, CT and/or MRI of the head and neck, as well as chest X-ray.^{11, 14, 15} In the last decade, PET-CT scanning has been introduced and has established an important role in detecting occult tumor sites.^{14, 16} Therefore, a combined FGD-PET-CT of the whole body is currently recommended as stated in the Dutch national guidelines.^{8, 14, 15, 17–22}

In case of squamous cell carcinoma (SCC), there is international consensus that the above-mentioned diagnostic approaches should be completed with panendoscopy (pharyngeal and laryngeal endoscopy, combined with cervical esophagoscopy and a bronchoscopy under general anesthesia), to search for a potential primary tumor in the mucosal alignment of the larynx, pharynx, trachea and upper esophagus.¹⁶ Systematic biopsies of suspected regions, biopsies of the nasopharynx and base of the tongue, as well as an ipsilateral tonsillectomy are routinely performed to detect occult primary tumors. If a tonsillectomy was performed in the past, biopsies of the tonsillar fossa should be obtained.

Retrospective studies on patients with CUP syndrome, in which the HPV status of the neck lymph node metastasis was compared to that of the eventually identified primary tumor, showed that HPV positivity is an indicator for a primary cancer originating in the oropharynx.^{23–26} HPV testing could, therefore, improve the sensitivity of current extensive diagnostic work-up protocols applied to true CUP patients by indicating an occult oropharyngeal primary tumor in HPV-positive metastasis. Thus, the addition of HPV testing to the diagnostic work-up in patients with CUP is recommended by some authors^{23–27} and the recommendations particularly concern the investigation of HPV status in true CUP patients to detect potential HPV-positive primary cancers which were missed during the regular diagnostic work-up in CUP.^{28, 29} However, it is questionable whether HPV-positive primary tumors of the tonsil or base of tongue are not already detected by tonsillectomy, biopsies of the base of tongue and/or biopsies during panendoscopy.^{24, 25, 28–32}

After the diagnosis of a ‘true CUP syndrome’ with exclusion of a primary cancer, an appropriate treatment modality should be established. To achieve loco-regional control, a combination of surgery (tonsillectomy as a diagnostic tool to exclude a primary cancer and ipsilateral neck dissection) and adjuvant radiotherapy are most commonly performed in CUP syndrome.^{14, 18, 33} Singular treatment modalities, such as neck dissection or radiotherapy alone, may be appropriate for patients with more favorable clinical N-stages.^{14, 18, 34} Also, adjuvant chemoradiation (with or without a neck dissection) can be applied in patients with CUP, but may be associated with an increased toxicity.^{35, 36}

There is a current debate on most efficient therapies regarding oncologic results and associated toxicity of more aggressive treatment protocols. The extent of radiotherapy, in particular whether administered ipsilaterally or bilaterally in the neck and with or without additional radiation of the pharyngeal mucosa, is subject of controversies; and a general consensus is yet to be established.³⁷

In the literature, depending on the used treatment modalities and patient characteristics, 5-year loco-regional disease-free (LR-DF) survival rates vary from 44.0 to 66.0 %.^{20, 33, 38,}

³⁹ Five-year overall survival rates vary from 27.0 to 66.0 %.^{14, 16, 18–20, 33, 36, 39, 40}

5.2.1. Hypothesis and aim

This present study aimed at analyzing the current therapeutic regimen conducted in two European oncologic centers. The two collaborating head and neck cancer centers, both tertiary referral hospitals in University medical centers (Maastricht, Netherlands, 121,050 inhabitants with a referral area containing about 1,500,000 inhabitants, and Cologne, Germany, 1,000,000 inhabitants) united in this retrospective analysis of patient records to analyze and compare their data.

Patient data were evaluated regarding the role for HPV testing and the influence of HPV prevalence on outcome was explored, as well as the value of irradiating the pharyngeal mucosa to treat occult primary tumor sites. Lastly, regional control rates were described

and compared in patients treated with bilateral versus ipsilateral postsurgical radiation, with or without concomitant chemotherapy.

5.3. MATERIALS AND METHODS

5.3.1. Patient population

The data of patients presenting with CUP syndrome to the Department of Otorhinolaryngology, Head and neck Surgery of the University of Cologne and Maastricht Medical Centre, from 1997 to 2010 were retrospectively assessed. Patients were included if no primary tumor was detected during thorough staging examinations ('true CUP') and if histopathological material was available (Cologne: n = 22; Maastricht: n = 29). Thus, patients who only received radiation without surgical therapy were excluded. This retrospective investigation was approved by the ethics committees of the University of Cologne as well as Maastricht Medical Centre.

5.3.2. Diagnostic work-up in both centers

In Cologne, all patients underwent ultrasound of the neck with ultrasound guided fine needle aspiration cytology (USgFNAC) of the suspected mass, computed tomography (CT) or magnetic resonance imaging (MRI) of the head and neck, CT of the abdomen and thorax, abdominal ultrasonography, skeletal scintigraphy and positron emission tomography (FDG-PET, since 2008).

In Maastricht, patients underwent ultrasonography of the neck and USgFNAC of the suspected lesion, MRI or/CT of head and neck, chest X-ray or CT-thorax and combined PET/CT scanning (whole body, since 2002).

In both institutes, a panendoscopy was performed: In Cologne, patients underwent panendoscopy of the upper aerodigestive tract (pharyngeal and laryngeal endoscopy combined with rigid esophago-(gastroduodeno-)scopy and a rigid bronchoscopy under general anesthesia. If no primary tumor was found in these staging examinations, bilateral tonsillectomy and biopsies of the base of tongue as well as curettage of the nasopharynx for histopathological evaluation were executed.

In Maastricht, patients underwent a panendoscopy similarly, during which biopsies of base of tongue and nasopharynx and an ipsilateral tonsillectomy were performed. The contralateral tonsil was not removed.

5.3.3. Patient collective

In Maastricht, true CUP Syndrome is classified as no known primary cancer found—when a primary cancer can be identified, patients are registered by tumor location. It was, therefore, not possible to investigate retrospectively how many patients initially presented with a neck mass suspicious of CUP. In Cologne, similarly, patients presenting with a neck mass are treated with intensive staging, however, we were able to retrospectively show in how many patients a primary cancer could be found in this

patient collective (in 20 out of 47 patients, 30 %).⁴¹. Only the 27 true CUP patients were analyzed in the presented study.

Of all neck dissections in Cologne, the resection margins were classified by the pathologist as resected radically (R0). In Maastricht, the width of resection margins was specified in mm (Table 3).

5.3.4. Histopathology: additional HPV testing in the diagnostic work-up

In both institutes, HPV testing was assessed as follows: the lymph node metastases obtained via neck dissection were investigated retrospectively for HPV-DNA using PCR, and HPV typing was subsequently performed, as well as p16-immunohistochemistry. Since our two cancer centers collaborate, the HPV-investigations, including HPV-analysis, were performed at the same laboratories to exclude inter-investigator variations. Tissue biopsies from lymph node metastases were analyzed according to the following protocols. HPV-16-FISH (fluorescence in situ hybridization) analyses were performed retrospectively in 2011, as previously described by Hafkamp et al.^{42, 43}

5.3.5. Treatment protocols in both centers

Cologne

In Cologne, patients underwent a neck dissection of the ipsilateral neck: lymph nodes of levels I to V, as classified by Robbins,^{44, 45} were dissected with intended preservation of the jugular vein, the sternocleidomastoid muscle and the accessory nerve. The tissue from each level was placed in a separate pathology jar, submersed in 4 % formaldehyde and then submitted for histopathological investigation, where the lymph nodes were dissected, embedded and sectioned as described by Schroeder et al.⁴⁶ Regarding adjuvant radio- and chemotherapy, no standard protocol was used for determining the choice between ipsi- and bilateral radiotherapy whether or not combined with chemotherapy. Adjuvant radiotherapy was performed with linear accelerators (6 MeV) after three-dimensional treatment planning. The clinical target volume included the involved lymph node area, ipsi- or contralateral cervical and supraclavicular lymph nodes. The pharyngeal mucosa was not irradiated as such. Total dose to this volume was in the mean 50.4 Gy. Macroscopically involved lymph nodes were boosted to 59.4–63 Gy. Chemotherapy consisted of carboplatin administration (20 mg/m² body surface daily in week one and five, n = 8 in Cologne). no standard protocol was used for the choice of administering concomitant chemotherapy. In general, the decision for adjuvant therapy was made in consensus with the patient: after surgery, in an interdisciplinary informative discussion with the patient, all advantages and risks and side effects of chemoradiation were explained. The patient could then decide on the adjuvant therapy he desired, and in case of lymph nodes larger than N1, the general recommendation for adjuvant chemoradiation was given.

Maastricht

Regarding the treatment protocol in Maastricht, a modified radical neck dissection was performed (levels I to V), in which the jugular vein and sternocleidomastoid muscle routinely were resected.^{9,47} The accessory nerve was preserved if feasible. Patients treated before 2002 in general underwent an ipsilateral neck dissection and post-operative radiotherapy of the pharyngeal mucosa and the bilateral neck. The elective radiation dose to the uninvolved neck regions and the pharyngeal mucosa was 46–50 Gy. The regions of the involved nodes were treated up to 60–66 Gy. After 2002, the treatment protocol was changed with abandoning radiation treatment of the pharyngeal mucosa and the contralateral neck, only performing post-operative ipsilateral radiation of the neck following ipsilateral neck dissection.

Also, after 2002, in the N1 (single metastatic lymph node <3 cm) without radiographic signs of extracapsular growth in the staging examinations, single treatment modalities, e.g. ipsilateral surgery or radiotherapy alone, were occasionally performed in Maastricht (in 25 % of CUP patients in Maastricht in the decade). In patients with neck dissection, radiological findings were verified by histopathology. Patients not undergoing surgical therapy were thus excluded because their lymph node state was not pathologically confirmed and HPV status could not be examined. Concurrent chemotherapy during radiation was not used in the Maastricht cohort.

5.3.6. Statistical analysis

Statistical analysis was performed using SPSS® (Statistical Package for the Social Sciences, USA, Versions 18.0 and 19.0). Disease-free survival (DFS) and overall survival (OS) rates were estimated using the Kaplan–Meier algorithm for incomplete observations. The overall survival time was defined as the interval between the date of diagnosis and the last date when the patient was known to be alive (censored) or date of death for any reason (uncensored). The disease-free survival rate was measured as the period of time between the date of diagnosis and the date of the last follow-up examination in which the patient was disease free (censored), or the date of first recurrence independently if it was a local, regional, or distant recurrence (uncensored). The log-rank test was used to test for differences between subgroups. Statistical correlation was performed with the Chi-square test. A p value <0.05 was considered statistically significant in two-sided tests.

5.4. RESULTS**5.4.1. Demography: comparison of patient groups between Maastricht and Cologne**

There were no significant differences concerning age, tobacco smoking behavior, alcohol consumption and tumor differentiation grade between the patients of Maastricht and Cologne. Regarding gender, the Maastricht-group consisted of 22 men and 7

women whereas in the Cologne-group, there were only men ($n = 22$, $p < 0.001$). No differences were found in both institutions comparing the clinical N-status determined after radiology with the pathological N-status determined after neck dissection. Both clinical and pathological N-status were distributed equally in both study groups (Table 1). In the Maastricht-group, however, nine pN3-neck lymph node metastases were included, compared to three pN3-metastases in Cologne [31 % (9/29) and 13.63 % (3/22), $p = 0.147$].

None of the patients presented with contemporary distant metastases at the time of diagnosis.

Prior to panendoscopy and neck dissection, two patients from Maastricht underwent a diagnostic nodal excision at the neck at other hospitals as compared to seven patients from Cologne ($p = 0.021$). The average time from first presentation of disease until neck dissection at the University Medical Centers was 40 days in Maastricht and 26 days in Cologne ($p = 0.005$). In histopathological analysis, extracapsular growth of the metastases was found in 22 patients from Maastricht (75.9 %) as compared to 18 patients from Cologne (81.8 %, p value: NS).

5.4.2. Treatment of CUP patients

Out of the 51 included patients, 48 patients received adjuvant radiotherapy. Concomitant chemotherapy was administered to eight patients in Cologne (carboplatin 20 mg/m² body surface daily in week 1 and 5). Three patients were not irradiated: two patients refused further treatment and in the third patient, a distant metastasis was discovered at the planning PET-CT of the whole body, as it has become routine work-up in Holland in recent years.

Two patients received adjuvant radiotherapy elsewhere and the records were unclear whether treatment was administered ipsi- or bilaterally. Those patients were excluded from the analysis when comparing the extension of adjuvant radiotherapy.

Thirty-two patients received postsurgical radiotherapy in the ipsilateral neck and 14 patients bilaterally. Eight patients received concomitant chemotherapy (ipsilateral RT: $n = 2$; bilateral RT: $n = 6$) and another six patients were treated with additional radiotherapy of the pharyngeal mucosa (bilateral RT: $n = 6$).

No differences were noted between adjuvantly treated groups regarding age, gender, institute of inclusion in Maastricht or Cologne, smoking and alcohol behavior. Patients treated with adjuvant bilateral radiotherapy were all but two (pN2a) staged with pN2b or higher involved lymph nodes.

5.4.3. Five-year overall survival and disease-free survival rates related to adjuvant radiation strategy

The 5-year overall survival (5-year OS) was 54.9 % in our patient population ($n = 51$). For patients treated with ipsilateral adjuvant radiotherapy, the 5-year OS was 67.6 % as compared to 35.7 % for the bilateral RT-group (log rank, $p < 0.001$, Fig. 1). This

difference is caused by the difference in pN-status between both groups, as patients treated with neck dissection and adjuvant bilateral radiotherapy were all —but two (pN2a)— staged with pN2b or higher involved lymph nodes. The 5-year OS for patients with a pathologically staged neck pN1-N2a was 81.8 % as compared to 46.2 % for pN2b or higher. In the latter group, however, 5-year OS also did not differ significantly between both adjuvant radiotherapy-groups.

5.4.4. Local control: the value of radiotherapy of the pharyngeal mucosa in treating possibly missed primary tumors

Only six patients in our patient population of $n = 51$ received radiotherapy of the pharyngeal mucosa. There was only one oropharyngeal tumor found during follow-up presenting 52 months after therapy. This tumor was treated as a second primary cancer considering the period of more than 4 years after the initial CUP syndrome (Table 2; Fig. 1). Table 2 shows tumor characteristics and clinical follow-up data.

5.4.5. Regional control: the value of adjuvant ipsilateral versus bilateral radiotherapy

Nine patients experienced regional recurrence during follow-up and in one patient, there was residual disease after treatment. All recurrences were seen in patients with a maximum nodal size of >3 cm or a conglomerate of lymph nodes.

Regarding regional recurrence rates, there was no significant difference between the ipsilaterally or bilaterally adjuvant RT-group (6 vs. 2 recurrences in resp. 32 and 14 patients). There was also no difference in sidedness of the recurrence between both groups: 50 % of recurrences in both groups were found in the contralateral neck. However, in two patients with recurrent disease in the contralateral neck, remission of disease could be established after dissection of the contralateral lymph nodes.

In one patient, staged pN3, PET/CT scanning after radiotherapy revealed incomplete response to therapy after R2 resection and also a contralateral recurrence and a newly diagnosed distant metastasis to the lung.

5.4.6. Occurrence of distant metastases

In total, nine patients developed distant metastases. They were all initially treated for pN2b or higher-staged lymph node involvement.

There was no significant difference in the occurrence of distant metastases between patients treated with adjuvant ipsilateral radiotherapy (8/32) and patients treated with bilateral radiotherapy (1/14) or with concomitant chemotherapy (1/8). There were no further patient characteristics associated with the occurrence of distant metastases.

5.4.7. Disease-free survival

In total, 13 patients presented with recurrence of disease. All recurrences or distant metastases were found within 2 years after therapy. For ipsilateral radiotherapy, the 2-year disease-free survival was 68.8 % ($n = 32$). When comparing pN1-N2a with pN2b

or higher-staged neck, the 2 year-DFS was 88.9 % compared to 60.9 %. For bilateral radiotherapy, the 2-year disease-free survival was 78.6 % only (n = 14: all but two pN2a necks were pN2b or higher).

One patient experienced residual disease after adjuvant radiotherapy together with a contralateral recurrence and distant metastasis (mainly to the lung, but also bone and liver). Interestingly, six patients experienced recurrent regional disease within 6 months after radiotherapy, five of which simultaneously showed distant metastases.

Table 3 shows the different diagnostic and treatment strategies of CUP patients in Maastricht and Cologne.

5.4.8. The role of HPV testing in CUP syndrome

In none of the 29 tested samples of Maastricht, HPV could be found when tested for p16-/HPV-PCR and HPV-FISH. In the 22 tested neck dissection samples of Cologne, four samples were identified as HPV-positive (18.2 %). In one of the four HPV-positive patients, an oropharyngeal tumor originating from the base of tongue was found after 52 months of follow-up. In Maastricht, no HPV-positive tumor was found.

In this total investigated population, the prevalence of HPV was 7.8 %. The relationship of HPV association with local control, regional control and the 5-year OS was absent as a consequence of the low prevalence. In Maastricht, no HPV-positive tumor was found. Table 2 shows the results of HPV testing in Cologne, related with tumor characteristics and clinical follow-up.

In comparison to oropharyngeal cancers, in which percentages of HPV infection are reported to be as high as 30–70 %, this appears to be a different tumor entity.^{41, 48}

5.5. DISCUSSION

The diagnostic and therapeutic approach of cervical metastases of unknown primary tumors is debated in the current literature.^{16, 17, 19}

International guidelines still are scarce and diverse and there is an urgent need for a unified regime. This current description of treatment strategies in two European tertiary medical centers highlights the international diversity in therapeutic approaches, despite the high medical and scientific standards. Most international guidelines focus on CUP syndrome in general (also inguinal and para-aortal, published by hemato-oncological societies) and are not specific for the management of cervical CUP (e.g. <http://www.NICE.org.uk>, <http://www.dgho-onkopedia.de>).

Comprehensive guidelines for the management of cervical CUP syndrome have been published by the national comprehensive cancer network (NCCN, <http://www.NCC.org> guidelines) and also by the Dutch head and neck society (NWHHT). These guidelines are similar, but allow multiple treatment modalities for the different N-status. For N1-status, monotherapy is suggested (neck dissection or radiotherapy). N2-N3-staged neck statuses are preferably treated with neck dissection of levels I to V. Postsurgical

treatment consists of ipsilateral radiotherapy and its extension is dependent on the size and number of affected lymph nodes, the presence of extracapsular spread and the resection status (R0/R1). The possibility to treat the neck bilaterally is mentioned, but not explicitly recommended.

Also, the addition of concomitant chemoradiation or induction chemotherapy is indicated as optional in guidelines, and needs consideration in the presence of extracapsular spread. Discussion on the role of chemotherapy in the guidelines is based upon findings in studies on patients with neck metastases in known primary head and neck carcinomas rather than upon results in studies on CUP patients.

Regarding the role of radiotherapy of the pharyngeal mucosa, guidelines include this in the discussion for consideration, and mention a possible restriction of radiotherapy only to the oropharyngeal and nasopharyngeal mucosa when HPV or EGFR is detected.

HPV-associated oropharyngeal carcinomas are present with smaller primary tumors and often bulky neck disease.⁴⁹ The HPV prevalence in 'true CUP syndromes' can, therefore, be related with occult and missed primary tumors of the oropharynx. Moreover, in the Dutch guideline, one can consider primary

(chemo-)radiation instead of surgery as a consequence of probable relation of CUP-disease with a primary tumor of Waldeyer's ring.^{50, 51} A positive HPV association could support this change of treatment protocol in the affected CUP patients.

This study combines the treatment strategies in two European tertiary referral centers (University Medical centers of Cologne and Maastricht). To attribute to the voids in the described guidelines, the prevalence of HPV and the postsurgical management of CUP patients in both institutes were retrospectively assessed comparing ipsilateral and bilateral radiotherapy, radiotherapy of the pharyngeal axis and the addition of chemotherapy. Therefore, the presented data included only true CUP patients who were primarily treated surgically and excluded all cases in which a primary cancer was found. In both institutes, patients routinely underwent neck dissection levels I to V and all but three patients were treated with postsurgical radiotherapy. From all patients, histopathologic samples were available for analysis of the prevalence of HPV.

The pitfalls of this retrospective analysis are the low case number and the diverse approaches to cervical CUP syndrome in diagnostics and therapies — the broad variety prohibits any statistical evaluation or conclusion. However, it does indicate the need for a unified regimen and calls for clear guidelines on the background of the ongoing discussions.

The diagnostic work-up in both institutes was similar, however, in Cologne, a bilateral tonsillectomy was performed as compared to a unilateral tonsillectomy in Maastricht. This might have led to selection bias as possible contralateral occult primary tumors can be missed. Nevertheless, no occult primary tumor revealed during follow-up or that HPV prevalence was 0 % in the Maastricht population.

As described, neck dissections were performed through levels I to V in both institutions, however, in contrast to Maastricht, the sternocleidomastoid muscle was routinely preserved in Cologne.

5.5.1. Role of additional HPV testing in CUP patients

Studies on the prevalence of HPV in lymph node metastases, of which the primary tumor could not be detected after diagnostic work-up — so-called true CUPs — are scarce and contradicting. HPV prevalence rates range from 0 until 100 % tested in very small samples, ranging from $n = 2$ to $n = 25$.^{23–25, 27, 52} In none of these studies, a relation between HPV status and survival was found. Moreover, in the current literature, no study reports of HPV testing during diagnostic work-up leading to the discovery of the primary tumor, in case the regular diagnostic work-up revealed no primary cancer, i.e., a true CUP syndrome. In this study, no HPV was detected in 29 true CUPs of Maastricht. In Cologne, only four HPV-positive CUP patients were found. Therefore, HPV positivity in those patients did not result in different outcome, although it has to be acknowledged that the number of patients is very limited for implications of HPV in true CUPs in this group of patients. This stands in great contrast to oropharyngeal tumors, in which HPV prevalence rates range from 20 to 90 %⁴⁸ and they appear to be a different tumor entity.^{41, 48} This seems important, as some institutes treat CUP syndrome akin to oropharyngeal tumors. Probably, most HPV-positive occult primary tumors of the oropharynx are detected by tonsillectomy or biopsies of the base of tongue resulting in low percentages of HPV-positive ‘true CUPs’. However, in the diagnostic work-up of patients referred with the clinical diagnosis ‘CUP syndrome’, HPV detection in USgFNA could show an important role in helping to identify a primary tumor in the oropharynx.⁴¹

A prospective study is suggested, in which the removed tonsils and oropharyngeal biopsies are thoroughly inspected for the presence of small (microscopic) primary tumors in case of HPV presence detected during the diagnostic work-up of CUP. It seems, however, that after a thorough diagnostic work-up, the prevalence, which in this study is only 7.8 %, and consequently the role of HPV in true CUP is very limited.

5.5.2. Therapeutic decision making in CUP

As in recent years, multiple institutes down-regulated the treatment of CUP patients from adjuvant bilateral radiotherapy including the pharyngeal mucosa towards adjuvant ipsilateral radiotherapy, studies offering a clear comparison are lacking in literature. Both collaborating institutes have undergone a similar down regulation of therapy, as it was observed that morbidity in the first group was high and the occurrence of recurrent, mostly contralateral, disease most often was curable and did not affect the mortality rates. This resulted in a comparison between patients adjuvantly treated with ipsilateral radiotherapy ($n = 32$) and contralateral radiotherapy ($n = 14$). There was, however, a prolonged time from initial diagnosis until treatment in Maastricht as compared to Cologne (40 vs. 26 days). Whether this difference affected outcome remains unclear. Moreover, there was no significant difference in distribution between pN1-2a and pN2b-3 staged necks in both institutes. The latter group of patients more frequently developed recurrent disease and distant metastasis during follow-up.

5.5.3. Adjuvant radiotherapy of the pharyngeal mucosa

During follow-up, only one oropharyngeal tumor manifested after 52 months, located in the base of the tongue. Regarding the period after first presentation, this tumor was considered as a second primary cancer. One possible explanation for the fact that no primary cancer manifested during follow-up after therapy might be that small primary tumors of the base of the tongue or other pharyngeal subsides could be combated by the patient's immuno-system.⁵³ Another reason could be that unintended radiation dose to the mucosa, especially when not applying conformal radiotherapy, may result in successful eradication of a microscopic, initially undetected primary tumor in the pharyngeal region, although no radiotherapy of the pharyngeal mucosa is given. A third reason could be that histopathologically undetected micro carcinomas have been resected during diagnostic tonsillectomy and biopsies.

5.5.4. Overall and disease-free survival: postsurgical ipsilateral versus bilateral radiotherapy of the neck with or without chemotherapy

Patients with pN2b or higher-staged necks experience a worse overall and disease-free survival. Bilateral radiotherapy was administered in only pN2b or higher-staged necks. The overall survival rate of bilaterally treated patients was worse as compared to ipsilaterally treated patients. This difference disappeared after stratification for lymph nodes staged pN2b or higher.

Neither were there differences between patients treated with ipsilateral nor bilateral radiotherapy regarding a recurrence-free survival rate, nor regarding the occurrence of distant metastases between patients treated with ipsilateral or bilateral radiotherapy with or/without chemotherapy.⁵⁴

Regarding total disease-free survival, it was noted that all recurred during the first 2 years after treatment. The disease-free survival for ipsilaterally treated patients was almost 90 % in lymph nodes staged lower than pN2b. When comparing lymph nodes staged pN2b or higher, the DFS was just over 60 % for ipsilaterally treated patients compared to almost 80 % for bilaterally treated patients. This was, however, not significant.

Despite the similar recurrence rates in both ipsilaterally and bilaterally treated patients, for higher-staged necks (or maximum nodal size more than 3 cm), additional treatment could be considered. Bilateral radiotherapy showed to be associated with a slightly better disease-specific survival ($p = \text{NS}$), but on the other hand increased radiotherapy-associated morbidity. Moreover, in case of contralateral recurrences, initial ipsilateral treatment offered the potential of salvage surgery in combination with radiotherapy. We, therefore, suggest a large prospective multicenter study in which subgroups can be expanded since in the current literature these data are lacking but are crucial.

5.6. CONCLUSION

This study united patient data from two European institutes to compare the therapeutic strategy for cervical metastasis of unknown primary squamous cell carcinoma. The presented data support the ongoing discussion in a call for a consensus strategy for the management in diagnostics and therapy of cervical CUP syndrome, particularly regarding the role of omission of radiotherapy of the pharyngeal axis, ipsilateral as compared to bilateral postsurgical radiotherapy and the role of concomitant chemotherapy and the prevalence of HPV in true CUPs. To increase evidence and to unify the treatment strategies for patients with CUP syndrome, a systematic (international) multicenter comparative study is needed.

Regarding HPV prevalence in true CUPs, in only four patients presenting with CUP, association with HPV was found (7.8 %) and data, therefore, did not provide insight in the influence of HPV presence on outcome after therapy in this patient population. The value of HPV testing in neck metastases of unknown primary tumors seems, in the presented patient collective, to be of limited value particularly when compared with prevalence rates in oropharyngeal carcinomas. Further investigation is advised.

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Figure 1. Five-year disease-free survival in patients treated with adju- vant ipsilateral (a) and bilateral (only n2b or higher) (b) radiotherapy

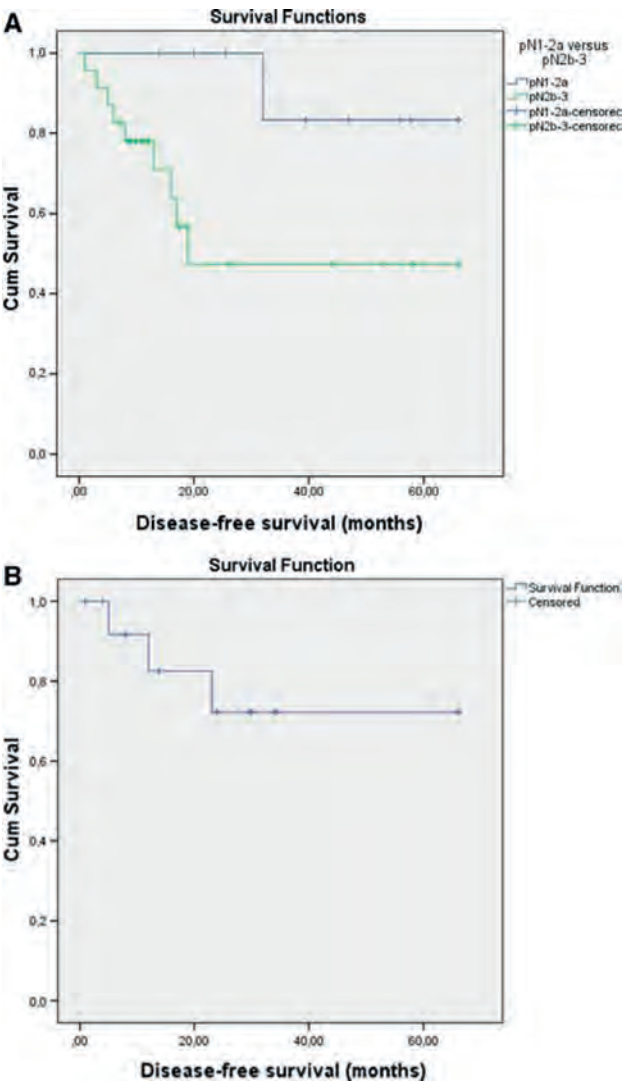


Table 1. Comparison of N-status in the Maastricht- and in the Cologne-group

	Maastricht	Cologne	p value
	n = 29	n = 22	
pN status ^a			
N1	2	2	
N2a	3	4	
N2b	15	13	
N3	9	3	0.147

^aDiagnostic work-up did not reveal any cN2c-staged neck. Therefore, no bilateral neck dissections were performed in both institutes

Table 2. Results of HPV testing in Cologne, related with tumor characteristics and clinical follow-up

	HPV-positive CUPs (n = 4)	HPV-negative CUPs (n = 18)	p value
Primary tumor	1 (base of tongue: 52 month)	0	NS
N-status			
pN1	1	1	
pN2a	0	4	
pN2b ^a	2	11	
pN3	1	2	NS
Regional recurrence	0	2	NS
Distant metastases	0	2	NS
Five-year overall survival	3 (75 %)	12 (66.7 %)	NS
			NS (log rank)

^a n = 1: 24 months

Table 3. Treatment of CUP patients in Maastricht and Cologne

	Maastricht (n = 29)	Cologne (n = 22)	p value
Nodus extirpation	2	7	0.021
Neck dissection	29	22	NS
Extracapsular growth	22	18	NS
Radical resection (R0)		18	
Resection margin < 1 mm	18	Unknown	
Resection margin 1–5 mm	4	Unknown	
Resection margins ≤ 5 mm	7	Unknown	
Angio-invasive growth	9	Unknown	
Perineural growth	4	Unknown	
Adjuvant RT	27	21	NS
Unilateral	21	11	NS
Bilateral	6	8	NS
Pharyngeal axis	6	0	NS
Unknown	0	2	NS
Plus chemotherapy	0	8	0.000
Time until OR (days)			
Average	40	26	0.005





Tumor control of cervical lymph node metastases of unknown primary origin: the impact of the radiotherapy target volume.

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6.1. ABSTRACT

Purpose: Debate on the extent of treatment of neck metastasis of cancer of unknown primary tumors (CUPs) is still ongoing. In two Dutch tertiary referral centers, the post-surgical radiation target volume changed from the bilateral neck including the pharyngeal axis to the unilateral neck only, in the course of the last decade. This study aims to investigate the outcome of patients with CUP before and after de-escalation of post-surgical radiotherapy.

Methods: Data of two Dutch tertiary referral centers were merged. Disease-free survival (DFS), overall survival (OS), and regional control rate (RCR) of 80 patients diagnosed with CUP (squamous cell and undifferentiated carcinomas) between 1990 and 2009 were retrospectively analyzed.

Results: Thirty patients received bilateral neck and pharyngeal axis radiotherapy and 42 patients ipsilateral radiotherapy only. In another eight patients, the postsurgical radiation target volume was expanded to the contralateral neck or to the pharyngeal axis, due to suspicious lesions on imaging. The 5-year DFS, OS and RCR were 60%, 51.2%, and 80% respectively, in the total patient population. RCR did not differ in patients treated with ipsilateral as compared to bilateral radiotherapy, nor did 5-year OS and DFS. No tumors occurred in the pharyngeal axis.

Conclusion: In this study, omitting elective treatment of the contralateral neck and pharyngeal axis did not lead to a decrease in locoregional control or survival rates when treating patients with CUP.

Keywords: [Lymph node, *pathology], [Neoplasms, Unknown primary], [Neck dissection], [Radiotherapy], [Survival Rate]

6.2. INTRODUCTION

Cervical lymph node metastases of carcinomas of unknown primary origin (CUP) represent 2-5% of all malignancies in the head and neck region [1]. Diagnostic approaches in patients with CUP are comprehensive, including a full history, a physical examination, ultrasonography (US) of the neck combined with fine needle aspiration cytology (FNAC), magnetic resonance imaging (MRI) of the head and neck region and/or computed tomography (CT) with (whole body) FDG-PET scan. The latter has been introduced in the last decades. Next, a panendoscopy is performed together with systematic biopsies of suspect regions and blind biopsies of the nasopharynx and base of the tongue, as well as an ipsilateral or bilateral tonsillectomy.^{1, 2}

Human papillomavirus (HPV)-positivity in the lymph node metastasis is an indicator for a primary cancer originating in the oropharynx.³ Sensitivity of diagnostic work-up may therefore be improved by testing on p16^{INK4A}-immunohistochemistry and HPV-16 DNA polymerase chain reaction (PCR).⁴ Correspondingly, detection of Epstein-Barr virus (EBV) nucleic acids in lymph node metastases of unknown origin suggests a nasopharyngeal primary tumor.⁵ Current TNM-classification (Eight Edition) has adopted special staging systems for HPV and EBV-associated lymph node metastases in CUP in which these entities have been classified as oropharyngeal or nasopharyngeal carcinoma respectively (T-stage as T0).⁶

In literature, 5-year loco-regional disease-free survival rates and 5-year overall survival (OS) rates vary from 17-85 and 22-79%, respectively, dependent on the treatment modalities applied and patient characteristics.^{2, 7-10} Generally, treatment consists of primary surgery (with or without postoperative radiotherapy) or primary radiotherapy. Radiotherapy comprises uni- or bilateral neck radiation, with or without radiation of the pharyngeal axis. In the last years, chemoradiotherapy has also been applied as a treatment option for selected cases.¹¹ However, the optimal treatment for CUP and which tumor- and patient characteristics should steer treatment decision-making, is still a matter of debate.¹⁰ A particular issue is whether radiotherapy should include the bilateral neck and pharyngeal mucosa or only the unilateral neck. Extensive radiotherapy may prevent recurrence in the contralateral neck and outgrowth of the occult primary tumor at the mucosal site but this is at the cost of significant increase of acute and late morbidity.^{7, 12-15} Some studies suggest that there is no difference in OS between patients treated with unilateral or bilateral radiotherapy and that patients can be spared the morbidity of bilateral treatment.^{12, 13} Recently, two reviews on treatment modalities in CUP also concluded that more evidence is needed regarding the extent of radiotherapy.^{9, 10}

The aim of this study is to determine the outcome of patients with cervical CUP in relation to the applied treatment in two Dutch head and neck clinics. Results of post-surgical unilateral versus bilateral post-operative irradiation and radiotherapy of the pharyngeal axis are compared in terms of disease-free survival (DFS), regional

recurrence rate (RCR) and OS. Also, the relation of HPV-detection in affected lymph nodes with outcome was investigated retrospectively.

6.3. MATERIAL AND METHODS

6.3.1. Patients

Data of patients presenting with cervical CUP at the departments of Otorhinolaryngology, Head and Neck Surgery of the Maastricht University Medical Centre (Maastricht UMC: Center 1) (n=60) and the Radboud University Medical Center Nijmegen (Radboud UMC: Center 2) (n=64), the Netherlands, from 1990 until 2009 were retrospectively assessed. In- and exclusion of patients with CUP are described in Figure 1. Approval by the ethics committee of both institutes was obtained.

All patients with CUP were discussed in the centers' multidisciplinary head and neck tumor boards: the most actual edition of the TNM-classification of the International Union Against Cancer was used to determine treatment plans. In the included era, HPV and EBV were not routinely tested.

Diagnostic work-up

The diagnostic work-up included a full history, a physical examination, ultrasonography (US) of the neck combined with fine needle aspiration cytology (FNAC), magnetic resonance imaging (MRI) of the head and neck region and/or computed tomography (CT) with (whole body) FDG-PET scan, introduced in the last decade. Also, a panendoscopy was performed together with systematic biopsies of suspect regions and blind biopsies of the nasopharynx and base of the tongue, as well as an ipsilateral or bilateral tonsillectomy. The contralateral tonsil was not removed routinely. If the patient had undergone tonsillectomy in the past, only biopsies of the tonsillar fossa were obtained.

6.3.2. Treatment

In both centers, the protocol for treatment of CUP syndrome is based on the Dutch national guideline: "Primary tumor unknown".¹⁶ Initially, treatment of CUP involved an ipsilateral neck dissection with adjuvant bilateral radiotherapy including radiotherapy of the pharyngeal axis. In the course of the last decade, the radiation target volume was reduced. The practice of adjuvant radiotherapy of the contralateral N0-neck, and radiotherapy of the pharyngeal axis were abandoned, and changed to post-operative ipsilateral radiation only.

Radiotherapy in Maastricht UMC: The elective radiation dose to the uninvolved neck regions and the pharyngeal axis was 46-50 Gy. The regions of the involved nodes were treated up to 66 Gy. Radiotherapy in Radboudumc: The median elective radiation dose to the uninvolved neck regions and pharyngeal axis was 50 Gy and the median dose delivered to the pathologically involved node level(s), was 64 Gy, range 56-70 Gy,

depending on histopathological criteria (higher dose if extranodal growth or close or positive resection margins).

Radiation therapy was administered using techniques that were available in those periods: in the beginning of the study, patients, undergoing bilateral neck irradiation including the mucosal axis, were treated with 2-D radiation with parallel opposing beams for the upper neck and a matching anterior lower neck field. Unilateral radiation for patients receiving ipsilateral neck radiation only was given by oblique wedge-pair beams. These techniques evolved into 3-D radiation and ultimately Intensity Modulated Radiation Therapy (IMRT).

Chemoradiation as part of the treatment protocol of CUP, was not used in both centers during the period of inclusion of this study. None of the included patients were therefore treated with adjuvant and/or concomitant chemotherapy.

6.3.3. Follow-up

Follow-up consisted of a periodic history and physical examination during five years in all patients and was scheduled every 2 months in the first year and extended to every 6 months during the fifth year after treatment of CUP. In case of suspect local, regional and/or distant failure, additional imaging tests and/or panendoscopy were performed, when considered necessary and when further treatment options were still present.

6.3.4. Statistics

Statistical analysis of the data was performed using SPSS software (v17.0). When comparing groups, the Pearson Chi-square test was used. Survival rates and data on disease-specific control in patients were calculated from the date of the first pathological confirmation of disease. There were no patients lost to follow-up. DFS, OS, RCR were computed with Kaplan Meier survival analysis.

DFS was defined as the survival until recurrence of disease locally, regionally and/or distantly. In order to report on the value of ipsilateral versus bilateral radiotherapy of the neck, the RCR is used. OS was defined as the survival until death. The log-rank test was used for univariate comparisons of the survival functions. Nominal two-sided p-values are reported, the significance level was set at $p \leq 0.05$.

6.3.5. HPV-status

The presence of HPV was retrospectively determined according to the algorithm described by Smeets et al.¹⁷ Thirty-two tumor samples in the center 1 and 40 samples in the center 2 were available in which immunohistochemical (IHC) detection of p16^{INK4A}, and/or HPV16 DNA polymerase chain reaction (PCR) were performed.

6.4. RESULTS

6.4.1. Study population

No differences were noted regarding the diagnostic work-up in the two centers. In total, 80 patients with CUP were analyzed in this study (Table 1). They all underwent a neck dissection followed by post-operative radiotherapy with curative intent (Table 2). Bilateral radiotherapy combined with irradiation of the pharyngeal axis was performed in 30 patients (38%) and ipsilateral radiotherapy without irradiation of the pharyngeal axis in 42 (52%).

Due to suspicious lesions found on imaging - but not histologically or cytologically confirmed, postoperative ipsilateral radiotherapy was combined with treatment of the pharyngeal axis in 2 patients, and another 5 patients received radiotherapy of the bilateral neck without irradiation of the pharyngeal axis. Finally, one patient with limited pN1-disease did not receive additional radiotherapy of the ipsilateral neck, but instead the pharyngeal axis was irradiated because of a suspected lesion found by imaging (Table 2).

6.4.2. Outcome

Five-year DFS, OS and RCR were 60%, 51.2% and 80%, respectively (Table 3). There were no significant differences in survival between the group of patients irradiated ipsilaterally and those treated bilaterally. Also, the 5-year regional control rates did not differ between both groups, resp. 77.3% and 82.9% ($p=.54$) (Figure 2), nor did the 5-year contralateral recurrence rate ($p=.23$).

In both groups, no primary tumors occurred in the pharyngeal axis during follow-up. Two primary tumors, both located in the floor of the mouth outside the pharyngeal axis, emerged during follow-up in the total population (in both patients 31 months after initial treatment). The first patient was initially treated with bilateral neck irradiation without the pharyngeal axis. The second patient was treated with radiotherapy of the bilateral neck and pharyngeal axis.

Twenty-three patients developed distant metastases during follow-up. This was not related to the extent of radiotherapy used.

6.4.3. HPV-status

In total, 4 out of 72 histopathological samples of cervical metastases tested positive for both p16^{INK4A}-expression and HPV DNA (5.7%), 3 patients of which were treated with radiotherapy of the pharyngeal axis, including the oropharyngeal mucosa. Five-year DFS, OS and RCR in the four HPV-positive patients were all 100%.

6.5. DISCUSSION

The objective of this study was to investigate the outcome of patients with CUP before and after de-escalation of post-surgical radiotherapy applied in two Dutch tertiary referral centers. In this study, no differences were found regarding survival and regional control rate in patients with CUP treated with neck dissection and post-operative bilateral radiotherapy including radiation of the pharyngeal axis (n=30) compared to patients with CUP treated with neck dissection and post-operative ipsilateral radiotherapy solely (n=42). Eight patients received additional therapy of the contralateral neck or the pharyngeal axis as a consequence of radiological suspicion of disease although this was not pathologically confirmed. In addition, no primary tumors occurred in the pharyngeal axis, even though radiotherapy to the pharyngeal axis was abandoned in 47 out of 80 patients.

6.5.1. Unilateral versus bilateral radiotherapy

In our study, no differences in survival rates and moreover no differences in regional control rates were found between the group of CUP-patients treated ipsilaterally and those treated with bilateral radiotherapy of the neck.

These results correspond with previous research by our group in which 29 patients of our cohort were compared with 22 patients with CUP in a German tertiary head and neck cancer referral center.¹⁸ In that study significant more contralateral recurrences were seen in the ipsilateral radiated patients compared to the bilateral radiated patients. In the current study, the above mentioned cohort of 29 patients was expanded to 80 patients which were homogeneously treated; the regional recurrence rate concerning contralateral relapses did not significantly differ.

In an early review of literature by Nieder et al. (2001), a median nodal relapse of 19% (range 8-45%) after comprehensive radiotherapy compared to 51.5% (range 31-63%) after ipsilateral radiotherapy was described.¹³ However, regarding 5-year overall survival rates no differences were noted between both groups (resp. 50%, range 34-63%, compared to 36.5%, range 22-41%). In a more recent meta-analysis by Liu et al (2016) of 16 studies that report outcome between bilateral versus ipsilateral radiotherapy,⁹ a significant reduced relative risk of 0.61 was described for nodal recurrence in patients treated with more comprehensive radiotherapy, however no differences between both groups were found for 5-year OS and DFS. This lack of difference in overall survival between both treatment groups was confirmed in a recent review of Müller von der Grün et al (2017).¹⁰ Our study also shows no significant differences regarding 5-year DFS and OS. Moreover, no differences were found regarding (ipsilateral and contralateral) regional control rates between both groups. In our study, the inclusion of the contralateral neck in 5 patients in which radiologically suspected lesions were found during radiation treatment planning without histologically or cytologically confirmation, may have contributed to the lack of differences in 5-year RCR between both groups. The strict evaluation criteria of CUP and uniform treatment (all patients underwent unilateral

neck dissection combined with post-operative radiotherapy) may have contributed to this favorable RCR when compared to other studies. Altogether, in our study metastatic disease (n=23) was a more common reason of disease failure than locoregional failure (n=18). The radiotherapy target volume was not related to the occurrence of distant metastases. This supports the current findings in literature that the possible benefit of extended volume radiotherapy on a slightly improved locoregional control, if present, cannot be translated into improved overall survival rates.^{10,19}

6.5.2. Radiotherapy of the pharyngeal axis

In our study, no primary tumors occurred in the pharyngeal axis regardless of inclusion of the pharyngeal mucosa in the radiation target volume. Two out of 80 patients developed a primary tumor during follow-up, both located in the oral cavity, which is generally not included in the target volume of pharyngeal axis irradiation. In a review of literature by Reddy et al (2001), a higher local (mucosal) failure is reported in patients who received treatment to the neck alone (44%) compared to those who received radiotherapy to the pharyngeal axis (8%).¹⁴ Also, lower primary tumor emergence rates were described for patients treated with bilateral radiotherapy when compared to patients treated with ipsilateral radiotherapy.^{7,14} A recent meta-analysis also reported a significantly lower 5-year primary tumor emergence rate (12%; RR=0.44) and a lower 5-year DFS rate when comprehensive radiation volumes were used.⁹ Again, the 5-year overall survival did not differ significantly for ipsilateral and comprehensive radiated patients, whereas acute severe toxicity and xerostomia were significantly increased in the latter group.

Contributing factors to the current reported low primary emergence rate might be the strict evaluation criteria of CUP in which only true-CUP-patients are selected, the comprehensive diagnostic work-up used in both centers in this study and the addition of radiotherapy of the pharyngeal axis in 3 patients in which lesions were suspected on imaging studies. The importance of a comprehensive diagnostic work-up is illustrated by the significant decrease of primary tumor emergence rates in literature after the introduction of PET-CT in the radiation treatment planning in CUP-patients.^{7,10,13} Nevertheless, the additional value of PET remains hard to quantify next to panendoscopy with blind biopsies of the base of tongue and tonsillectomy.²⁰

Furthermore in literature, the importance of selecting of irradiation volumes regarding the treatment of the primary site is emphasized with the introduction of IMRT, which allows preserving organs at risk (salivary tissue in particular), but on the other hand can miss the primary tumor that coincidentally would have been treated with older less sophisticated techniques.

6.5.3. HPV and CUP

In the 8th edition of the UICC's TNM-classification for CUP-patients, HPV- and EBV-associated lymph node involvement is staged separately corresponding to N-staging in HPV-associated oropharyngeal and EBV-associated nasopharyngeal carcinomas.

Inclusion of HPV- and EBV-testing in CUP may support prediction of the prognosis and the location of the primary tumor in the oropharynx and/or nasopharynx respectively.²¹ The prevalence of HPV-associated head and neck carcinomas still rises and the prognosis remains more favorable as in non-HPV-associated carcinomas. As patients treated with radiotherapy of the ipsilateral neck only were more often included in the second half of this study, it was interesting to investigate the possible presence of an HPV-endemic in this group and its influence on the presented outcome. Patients in our study cohort were treated before the introduction of HPV- and EBV-assessment to the routine diagnostic work-up and a retrospective analysis for HPV-presence could be performed in 72 out of 80 patients. Nevertheless only in 4 patients HPV/p16^{INK4a}-presence. Possibly, most HPV-positive primary tumor of the oropharynx were detected by tonsillectomy or blind biopsies of the oropharyngeal region resulting in low percentages of HPV-positive true CUPs.²²

6.5.4. Study design

In previous research we compared 29 patients of the currently presented cohort with 22 patients from a cohort in Germany.²² Due to a relative heterogenous treatment strategy in both centers, this study aimed to collect a larger patient group with a more homogenous therapeutic approach. Therefore, data was merged from two Dutch tertiary referral centers in which the diagnostic and therapeutic development evolved almost identically, in the period of inclusion of more than two decades. This retrospective study still encountered possible limitations to investigate the impact of radiotherapy target volume of the neck and pharyngeal axis on tumor control in CUP. The number of included patients is rather small (n=80). However the selected study group was derived from a collection of 2 cohorts which were homogeneously treated in 2 Dutch tertiary referral centers, and was the result of a meticulous exclusion of non-true-CUP. The inclusion period was extended over 2 decades: in the first decade of inclusion, radiotherapy of the bilateral neck and the pharyngeal axis was predominantly performed, whereas post-surgical ipsilateral radiotherapy was more often applied in the second decade of inclusion. This latter group, experienced a higher availability of IMRT compared to 2DRT in the first group, and was also more prone to have PET-CT included in the diagnostic work-up. The chance to coincidentally radiate a missed primary tumor with IMRT is nevertheless smaller than with older less sophisticated techniques. Regarding the role of PET, de Bree et al (2010) already mentioned the inability to quantify the additional value of PET next to a comprehensive diagnostic work-up including panendoscopy, tonsillectomy and blind biopsies of the base of tongue.¹⁹ The influence of a possible HPV-endemic in the latter half of the study, was also minimized as only 4 (out of 72) patients turned out to have HPV-associated disease. Again, the role of a comprehensive diagnostic work-up, particularly of the oropharyngeal region, may have led to this low prevalence of HPV in CUP.

In order to create the ideal world to compare the outcome of ipsilateral radiotherapy only with a comprehensive radiotherapeutic regime in CUP-patients, Nieder et al

recommended in 2001 a randomized controlled trial.¹³ It is reported that a similar trial which started in 2002, was never accomplished (EORTC-24001-22005) as a consequence of very limited patient enrollment.¹⁰ The low prevalence of patients with CUP and the heterogeneous treatment strategies as a consequence of the lack of well-designed studies, are important limiting factors to create a study design with an acceptable methodological work-up regarding this subject. A prospective multicenter approach in which homogenous therapeutic strategies are applied, are considered to be feasible.

6.6. CONCLUSION

In this study, omitting irradiation of the pharyngeal axis in patients with cervical lymph node metastases of unknown primary origin after performing a comprehensive diagnostic work-up, including PET-CT and panendoscopy with tonsillectomy and blind biopsies of the base of tongue, did not result in the emergence of a primary tumor in the pharyngeal axis during five years of follow-up. This can avoid acute and late toxicity of comprehensive radiotherapy of the pharyngeal mucosa with significant improvement of long-term quality of life of these patients. Also, the absence of post-surgical radiotherapy of the contralateral neck in CUP did not lead to a decrease of regional control rates nor of survival rates. The occurrence of distant metastases was the most important reason for failure of disease-free survival in this study. The true impact of radiotherapy target volume in CUP-patients still needs further investigation and at least requires a prospective multicenter approach.

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Figure 1. In- and exclusion criteria used for selecting patients with CUP in both participating centers.

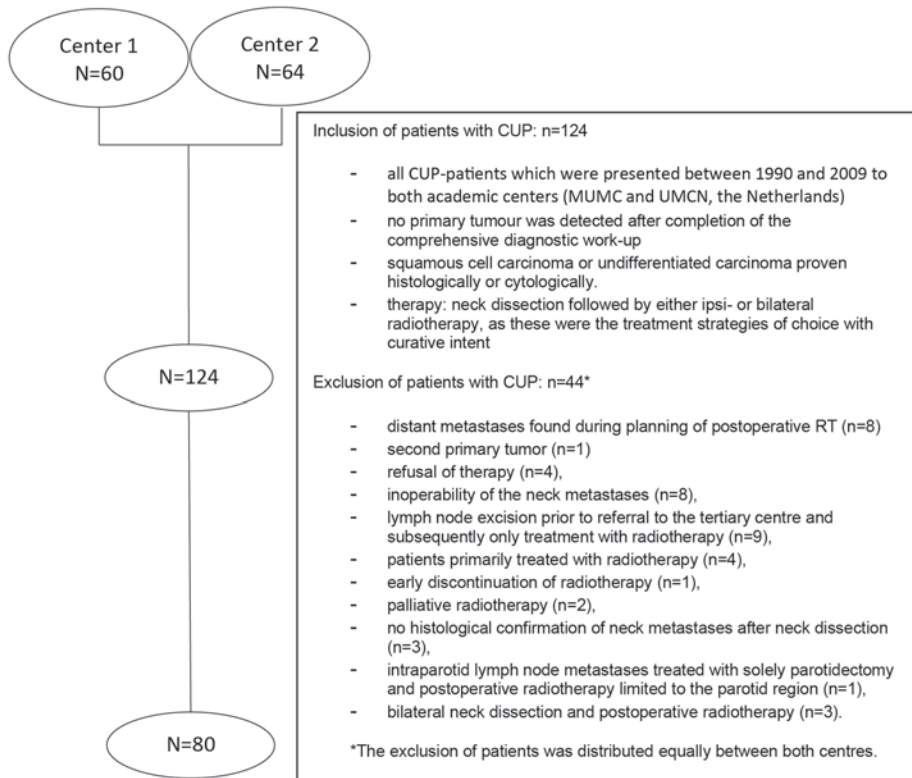
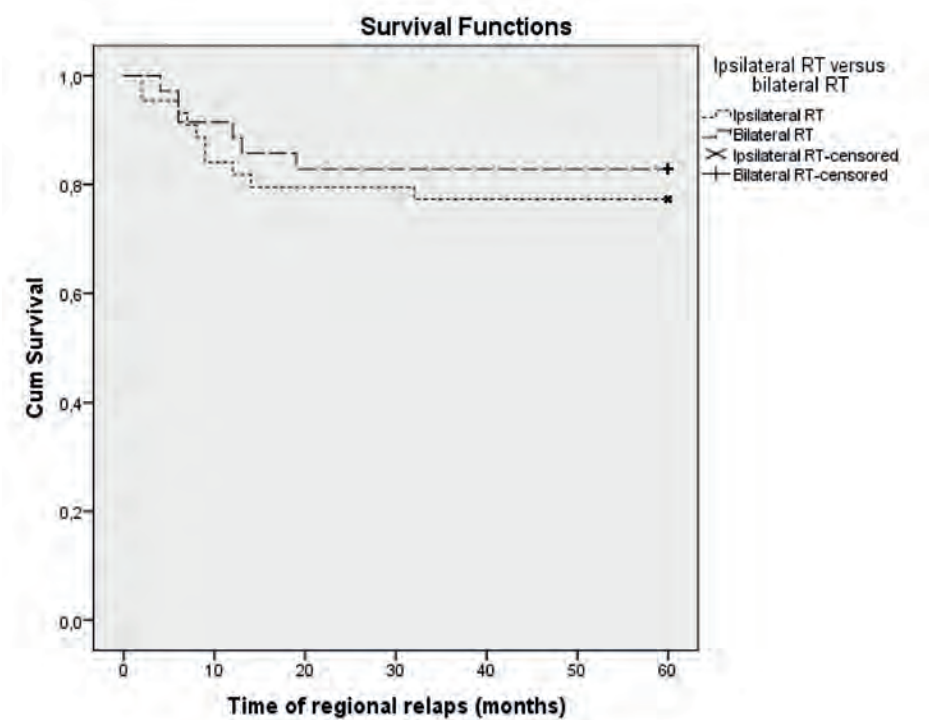


Figure 2. Regional control rate comparing patients treated with post-operative ipsilateral (n=44) versus bilateral radiotherapy of the neck (n=35)



Log-rank: p-value = NS

Table 1. Demographic data

		Total patient population	Center 1	Center 2	p-value
		n= 80	n= 43	n= 37	
Follow-up time (months)	mean	43.1	45.3	40.6	NS
	range	3-200	4-200	3-116	
Age (years)	mean	63.1	62,9	63.2	NS
	range	41-86	41-83	46-86	
Male/female		59/21	35/8	24/13	NS
Non-smoker	n (%)	1 (1.25)	1 (2,3)	0 (0)	NS
alcohol consumption ≤2 U/day	n (%)	36 (45)	18 (41,86)	16 (43.2)	NS
Histopathological data	squamous cell carcinoma	71	36	35	
	undifferentiated carcinoma	9	7	2	
pN-status (UICC version 7)	pN1	3	3	0	NS
	pN2a	17	8	9	
	pN2b	40	20	20	
	pN3	20	12	8	

Abbreviations: NS: not significant.

Table 2. Type of treatment based on pN-status

		number of patients	%	pN1	pN2a	pN2b	pN3	p-value
Type of neck dissection	Radical neck dissection	43	54	1	8	22	12	
	Modified radical neck dissection	22	28	1	6	12	3	
	Extended radical neck dissection	10	12	0	1	4	5	
	Selective neck dissection (regions I/II/III)	5	6	1	2	2	0	NS
Type of postoperative radiotherapy	Ipsilateral radiotherapy of the neck	44	55	2	11	21	10	NS
	Bilateral radiotherapy of the neck	35	44	0	6	19	10	NS
	Radiotherapy of the pharyngeal axis	33	41	2	6	16	9	NS
Type of combined therapeutical strategy	ND + RT-ipsi	42	52	1	11	20	10	
	ND + RT-bilat + RT-PA	30	38	0	6	15	9	
	ND + RT-ipsi + RT-PA	2	3	1	0	1	0	
	ND + RT-bilat	5	6	0	0	4	1	
	ND + RT-PA	1	1	1	0	0	0	NS
Total patient population		80	100	3	17	40	20	

Abbreviations: NS: not significant; ND: neck dissection; RT-bilat: radiotherapy of the bilateral neck; RT-PA: radiotherapy of the pharyngeal axis.

Table 3. Five-year overall survival, disease-free survival and regional control rate in the total patient population and in relation with pN-status and therapy

			Disease-free survival	Overall survival	Regional control rate
			N	%	%
Total patient population			80	60	51.2
pN-status	pN1	3	100	66.7	100
	pN2a	17	88.2	76.5	88.2
	pN2b	40	45	45	80
	pN3	20	60	40	70
	p-value		.014	NS	NS
Type of neck dissection	Radical neck dissection	43	53.5	41.9	76.7
	Modified radical neck dissection	22	72.7	63.6	86.4
	Extended radical neck dissection	10	40	50	70
	Selective neck dissection (regions I/II/III)	5	100	80	100
	p-value		.047	NS	NS
Post-operative ipsilateral versus bilateral radiotherapy	Post-operative ipsilateral RT	44	61.4	47.7	77.3
	Post-operative bilateral RT	35	57.1	54.3	82.9
	p-value		NS	NS	NS
Post-operative RT of the pharyngeal axis	Included	33	60.6	54.5	81.8
	Not included	47	59.6	48.9	78.7
	p-value		NS	NS	NS
Therapy strategy	ND + RT-ipsi*	42	61.9	50	78.6
	ND + RT-bilat-PA**	30	60	56.7	83.3
	p-value		NS	NS	NS

*ND + RT-ipsi: neck dissection and post-operative ipsilateral radiotherapy without radiation of the pharyngeal axis

**ND + RT-bilat-PA: neck dissection with post-operative radiotherapy including radiation of the pharyngeal axis.





Additional parameters to improve the prognostic value of the 8th edition of the UICC classification for HPV-related oropharyngeal tumors

Submitted. Straetmans J.M.J.A.A., Stuut M., Lacko M, Hoebers F, Speel E.M.J., Kremer B. Additional parameters to improve the prognostic value of the 8th edition of the UICC classification for HPV-related oropharyngeal tumors.

7.1. ABSTRACT

Background. This study explored the prognostic reliability of the TNM classification (8th edition) of the UICC for HPV-positive tonsillary cell carcinoma (TSCCs) compared to the previous (7th) edition. The aim was to see if using additional anatomical and non-anatomical parameters would improve its reliability.

Methods. 110 HPV-positive and 225 HPV-negative TSCC cases were retrospectively analyzed. HPV was determined by p16^{4INKA} immunohistochemistry, PCR and/or FISH. Survival was correlated with patient and tumor characteristics and with the tumor stages of the 7th and 8th UICC TNM classification system.

Results. The 8th edition of the UICC's HPV-associated TNM classification correlated better with prognosis in HPV-positive TSCCs than the 7th edition. In our HPV-positive patient group, however, smoking status was a stronger prognosticator of survival than UICC stage (8th edition). Non- or former smokers in our group had a 5-year overall survival of 95.1% regardless of tumor stage. Furthermore, age > 65 years, cN3, and M1 status were found to be significant prognostic factors.

Conclusion. In our patient population we found a significant improvement of the prognostic value of the 8th edition of UICC's staging system compared to the 7th edition. Nonetheless, further improvement would be possible by adding non-anatomical factors (smoking, age > 65y) and separating N0-N2 from N3.

Key words: HPV, tonsillar carcinoma, TNM classification

7.2. INTRODUCTION

In head and neck squamous cell carcinomas (HNSCCs), the UICC tumor staging system, based on T, N and M status, is crucial in predicting patient prognosis and guiding therapeutic decision-making.¹ Within this staging system, N status is considered to be the strongest prognosticator.²⁻⁴ However, for oropharyngeal tumors the prognostic value of N status has become questionable in light of the epidemic rise of HPV-associated carcinomas in recent decades.⁵⁻¹²

At diagnosis, HPV-positive tumors have smaller primary tumor sizes but equally affected lymph nodes (positive N status) compared to HPV-negative tumors. Moreover, their prognosis is more favorable than that of similarly staged HPV-negative tumors. The different clinical presentation and biological behavior of HPV-positive tumors, in combination with an increasing incidence of HPV, has shifted the prognostic value of “traditional” tumor classification models for HPV-dominant head and neck tumor sites. This became clear from our review of articles reporting the prognostic value of T and N status in tonsillar squamous cell carcinomas (TSCCs). We found 10 studies published prior to 1990 all reporting N status to be of prognostic importance (although only 2 studies provided results based on statistical analysis) while only 4 out of 12 studies published from 1990 onwards showed N status to be of prognostic relevance (see table 1).¹³⁻³⁴ In a previous study we tested the prognostic value of N status in a group of 81 HPV-positive and -negative TSCCs and found a decrease in prognostic value of N status in the whole group.³⁵ N status was of prognostic relevance only when tonsillar tumors were not HPV-associated. Similar findings have been reported by Klozar et al. (2013) in HPV-positive oropharyngeal SCCs and by Fritsch et al. (2013) in tumors located in HPV-dominant head and neck sites compared to non-HPV-dominant tumor sites.³⁶⁻³⁷

The issue of re-staging HPV-associated head and neck carcinomas using N status alone or in combination with other clinical parameters has been addressed from various perspectives in the literature. Ang et al. (2010) presented a predictive model on prognosis for stage III and IV oropharyngeal SCCs (OPSCCs) treated with concomitant chemoradiotherapy, whereby smoking status was combined with N stages “N0-2a” and “N2b-3”.³⁸ Spector et al. (2013) subdivided N status on the basis of diameter and number of nodes in 3 risk groups.³⁹ Huang et al. (2015) discriminated 4 prognostic groups in HPV-associated oropharyngeal carcinomas without hematogenous metastases, based on N status (N0-2c vs. N3), T status (T1-3 vs. T4), smoking behavior (fewer vs. more than 20 pack-years history) and age (younger vs. older than 70 years).⁴⁰ Dahlstrom et al. (2016) were not able to validate Huang’s results; consequently, they proposed an HPV-associated system in which N status was staged corresponding to nasopharyngeal carcinomas.⁴¹ Finally, the classification system of O’Sullivan et al. (ICON-S; 2016) was adopted for the 8th edition of the clinical TNM staging for HPV-related carcinomas (based on the sidedness and maximum diameter of the nodes rather than on the number of nodes).⁴² Regarding pathological staging, the system proposed by Haughey et al. (2016) was accepted for the 8th edition. In contrast to clinical staging, the number

of nodes (with a cut-off point of 4) determines N status (ranging from N0 to N2 without differentiation between N2a, N2b and N2c) for pathological staging.⁴³

Both the clinical and the pathological staging systems were recently validated by Cramer et al. in a population of more than 15,000 HPV-positive oropharyngeal carcinomas.⁴⁴ They have shown that the new 8th edition of the UICC TNM classification for HPV-related oropharyngeal squamous cell carcinomas overcomes some of the main shortcomings of the 7th edition.

However, given the findings cited from the literature, this study questions whether the 8th edition UICC classification system already reaches an optimum level to predict prognosis for HPV-positive OPSCCs. As mentioned above, the study of Ang et al. emphasized the role of both smoking and age.³⁸ Therefore, it is of interest to study the influence of tobacco use and senescence on tumor biology and thus on the validity of staging systems. Secondly, from 2009 onwards, as it became clear that the N-status is not a valid predictor in HPV-positive TSCCs, there were indications that the prognosis for N0 tumors was even worse than for N+ tumors.³⁵ It has been hypothesized that the presentation of a well treatable neck metastasis would lead to earlier detection of the primary tumor.⁴⁴ Thus, it is important to investigate the extent to which this is demonstrable in the currently adopted staging system (UICC 8th edition). A third factor that influences the prognostic reliability is the method for detection of HPV-association used in the 8th edition. The new system uses p16-IHC, which is widely available and frequently applied due to the low cost. However, it yields false positives in HPV-negative cases.

The primary aim of this study was to improve the classification system for HPV-associated tumors. We focused on a large series of 368 tonsillar SCCs which were subjected to HPV analysis using p16 overexpression, HPV-specific PCR and/or FISH. Tonsils were chosen for two reasons: they form the predominant oropharyngeal subsite where HPV-positive tumors develop; and different patterns of nodal dissemination have been reported between tonsillar SCCs and base of tongue carcinomas.⁴⁵ All cases were evaluated according to the 7th and the 8th edition UICC TNM classification. We examined the prognostic value of T, N and M status and investigate to what extent patient-associated clinical variables of age, smoking behavior, alcohol consumption, tumor differentiation grade, and treatment influence prognosis.

7.3. MATERIALS AND METHODS

7.3.1. Tumor material and patient data

The study population consisted of 368 TSCC patients diagnosed between 1987 and 2011 at the Maastricht University Medical Centre. That population was an expansion of our study group on which results were published in 2009 (n=81).³⁵ Their formaldehyde-fixed, paraffin-embedded archival biopsy and resection materials were classified by histopathology at the Department of Pathology, University Hospital Maastricht,

the Netherlands. The materials were analyzed for the presence of oncogenic HPV16 DNA by means of PCR and/or FISH as well as p16^{INKA} immunostaining in 335 available specimens.² Data on age, gender, TNM classification, tumor differentiation grade, tobacco and alcohol consumption, treatment modality, and follow-up (five years after treatment) were collected from the head and neck tumor database of our institute and from reviewing clinical, pathological, radiological and surgical reports. All tumors were re-classified according to the 7th and 8th edition of the UICC's TNM classification. Classification of smoking and alcohol consumption conformed to Hafkamp et al. (2002): patients were classified as daily smokers (≥ 1 cigarette, pipe and/or cigar per day), non-smokers (never smokers) or former smokers (those who had stopped smoking more than 10 years before the diagnosis of TSCC).⁶ During multidisciplinary counseling, treatment plans were based on tumor size, neck staging, presence of distant metastases, tumor histology and cytology of metastases, feasibility of surgery, clinical condition, co-morbidities and histopathology of resection specimens in case of surgery. Elective treatment of the neck was performed routinely - also in the N0 neck - because of the high incidence of occult metastases in TSCC.^{15, 19-21}

The investigation was conducted in accordance with the declaration of the 18th meeting of the world medical association in Helsinki 1964 and subsequent revisions. Approval for the study protocol was granted by the institutional ethical committee. Written informed consent was obtained from all included patients.

7.3.2. Statistical analysis

Variables considered were age at time of diagnosis, gender, TNM classification, tumor differentiation grade, smoking and alcohol consumption, T, N, and M status, and therapy.

The Youden Index was calculated to determine the optimal cut-off point in the ROC displaying the relation between sensitivity and specificity for the range of age value.⁴⁶ Survival analysis was performed; disease-free survival (DFS) and overall survival (OS) were calculated using the Kaplan-Meier method for each of the considered independent variables. Five-year survival (OS) was calculated from date of diagnosis until death or until discharge from follow-up. DFS was calculated from date of diagnosis until date of recurrence (local, regional or distant). Patients without recurrence were censored at date of last follow-up or death. Level of significance was determined at $p \leq 0.05$. Four patients who initially presented with distant metastases were excluded from the survival analysis. Multivariate analyses were performed using the Cox proportional hazards model. Variables remained in the model if p-values were below 0.10.² IBM® SPSS® Statistics version 20 was used for the statistical analysis.

7.4. RESULTS

7.4.1. Demographic data and outcome related to HPV association

In total, 368 patients with TSCC were included with an average age of 60.3 years (range 39-87). HPV association could be tested in 335 patient samples, of which 32.8% were HPV positive (n=110/335). An increasing prevalence of HPV-associated TSCCs during the last decade (since 2002) was observed. Prevalence rose from 21.4% in 2003 to 50% in 2011 (figure 1; the number of patients per year until 2002 was too small, so only the subsequent years were taken into account for this figure).

HPV-positive TSCCs were significantly related with a non- or former smoking status, with alcohol use of less than 2 units/day and a poorly differentiated tumor status. There was an equal distribution among age and gender groups regardless of HPV association.

No statistical differences were noted in the development of second primary tumors regardless of HPV association. Also, no differences were seen in prior history of head and neck cancer or cancer at other sites in both groups. For HPV-positive and -negative tumors, an equal number of patients presented with distant metastases. HPV status relative to patient characteristics is summarized in table 2.

7.4.2. Analysis of the 7th edition UICC tumor classification system for staging TSCCs

Compared to HPV-negative TSCCs, HPV-positive tumors were associated with significantly smaller primary tumors (Table 3). The extension of neck disease did not differ between both groups. As a consequence, HPV-positive and -negative TSCCs were equally distributed over UICC tumor stages I to IVc in the 7th edition. Within each tumor stage of the 7th edition, HPV-positive TSCCs were associated with a smaller T-status. For TSCCs staged Iva, for example, 38 out of the 61 HPV-positive tumors were classified T1-T2, compared to 31 out of 108 HPV-negative TSCCs (X^2 $p<0.001$).

In HPV-positive TSCCs, only N3 (affected lymph node >6cm) was associated with a worse survival (Cox regression, $p<0.001$; OR 8.233 (95%CI: 2.68-25.29)). No significant differences in survival were noted between N stages N0 to N2c, nor between the different stages when we classified tumors according to the 7th edition. Correspondingly, only TSCCs staged IVb or higher showed a worse survival (Cox regression, $p<0.001$), and no prognostic differences were noted between stages I and IVa, which altogether cover 90% of the HPV-associated TSCCs.

Among the largest subgroups, stage IVa, for example (n=181), an overall survival of 85% was found in HPV-positive TSCCs, a significantly higher rate than the 42.1% found in HPV-negative TSCCs (log rank, $p<0.001$). This difference may possibly reflect the association of HPV and smaller primary tumor size, given that in stage IVa the 5-year OS for T1-2 and T3-4 were resp. 62.3% versus 28.7%, independent of HPV status (log rank, $p<0.001$).

In contrast, in the HPV-negative population, a worse survival was associated with increasing T status, N status and UICC tumor stages (Cox regression, $p<0.001$).

Thus, in the 7th edition, HPV-positive tumors were mainly staged higher as a consequence of their nodal status. However, this higher stage seemed to have no prognostic implications.

7.4.3. Analysis of the 8th edition UICC tumor classification system for staging TSCCs

All HPV-positive tumors were re-staged in our study. In the 8th edition, a distinction is made between clinical and pathological staging. Because 38 out of our 110 HPV-positive TSCCs were treated surgically, pathological staging was possible in these 38 tumors (table 4).

Using the 8th edition of the UICC system for clinical staging, re-staging of the majority of HPV-positive tumors resulted in classifying only 3% of patients as having stage IV tumors, compared to 67% in the previous classification. Figure 2 depicts the shift between stages. Using the 8th edition, we found a decreasing survival with increasing clinical UICC tumor stage in HPV-positive tumors ($p < 0.001$) (table 4). In the group of HPV-positive TSCCs which were treated surgically and thus staged pathologically ($n = 38$), we found no decrease in survival with increasing tumor stage.

7.4.4. Univariate analysis of survival of T, N and M status (8th edition UICC TNM classification) IN HPV-positive TSCC patients

Only nodes larger than 6cm (cN3 status) were associated with a worse survival ($p < 0.001$), and no differences in survival were found for necks that were clinically staged cN0, cN1 and cN2 (table 5a). Remarkably, cN1 (unilateral nodes with a diameter of no more than 6cm) seemed to be associated with a better survival than when no lymph nodes were affected. The number of contralateral or bilateral involved necks in TSCCs was very low in this study.

Also, no correlation with survival was present when N status was defined pathologically, and no difference in survival was found between HPV-positive pathologically staged necks pN1 and pN2.

Altogether, only lymph nodes that were both larger than 6cm (cN3) or TSCCs that presented with distant metastases (M1) correlated with a poor prognosis in HPV-positive TSCCs.

Table 5b presents the odds ratios (and 95% confidence intervals) for each considered explanatory variable obtained from Cox regression modeling of OS.

7.4.5. Univariate analysis of survival of non-anatomical characteristics of HPV-positive TSCC patients Smoking, age >65yrs, cN3 status and M status were correlated with OS. Treatment category had no influence on OS, with the exception of palliative care for which a significantly lower OS was found.

The rationale for choosing age 65 years as the cut-off point is as follows. The Youden Index was calculated to determine the optimal cut-off point in the ROC curve that

displays the relation between sensitivity and specificity for the range of age value.⁴⁶ The optimal value was 2.22 for both the total data set and for the HPV+ cases. The Youden Index corresponded to the age of 63.5 for the total data set and 66.5 for the HPV+ patients. Age >65 years showed a significant correlation with a worse prognosis ($p<0.001$).

In the HPV-positive group, the 5-year OS in the non- or former smokers group ($n=41/110$) was 95%, compared to 66% in the smoking group ($n=62/110$) ($p<0.001$). The number of pack years had no further influence on prognosis. There was no difference in survival between never-smokers and former smokers (those having quit more than 10 years before diagnosis). Moreover, non- or former smokers were equally distributed over the different stages in the UICC 8th edition (figure 3). Interestingly, being a non- or former smoker had a favorable prognosis independent of tumor stage. In stage I the overall survival of non-smokers was 95% , in stage II 100%, and in stage III 90%.

An inverse relationship was found between age and smoking behavior in patients with HPV-positive TSCCs (X^2 , $p=0.036$): non-smokers were more often older than 65 years, whereas smokers developed a HPV-positive TSCC more often at or under age 65. Smokers older than 65 had a worse overall survival (log rank $p=0.049$). Smoking and age were not correlated to T status, N status and/or UICC tumor staging (8th edition).

7.4.6. Multiple regression analysis of survival including T, N and M status and non-anatomical characteristics of HPV-positive TSCC patients

The multiple regression analysis resulted in a model with M status, smoking category and age 65 years or older (Table 5c).

After analysis of the above-mentioned prognostic factors (smoking behavior, age and N3 status), we developed a predictive model of four groups of patients with HPV-positive TSCCs for patients without hematogenous metastases (figure 4). M status was not included as the 5-year OS was 0%. Group I ($n=41$) included non- or former smokers with a 5-year overall survival rate of 95.1%, regardless of age, number of pack years or T and/or N status, with the exception of cN3 status. Group II ($n=45$) included smokers of age 65 or younger with a 5-yr OS of 75.6.1% (HR 5.649, 95%CI 1,251-25,498). The third group ($n=13$) consisted of smokers older than 65 years of age with a 5-yr OS of 46.2% (HR 14.165, 95%CI 2,935-68,368). Survival in group II and III was determined regardless of TNM staging. The fourth group contains patients staged with a N3 staged neck This group showed a 5-yr OS of 33% ($n=6$). Interestingly, the two patients who survived were non-smokers who presented with a cN3-staged neck.

We then considered other prognostic classification models which utilize non-anatomical parameters, as introduced in literature after recursive partitioning analyses (RPA) and adjusted hazard ratios (AHR) by Ang et al., O'Sullivan et al. (RPA and AHR, i.e. ICON-S), and Haughey et al. When applying these to our HPV-positive tonsillar carcinoma patients, no relation between the proposed staging and outcome was found, except for the AHR based model by O'Sullivan that was used for the 8th edition UICC clinical staging system (figure 4).^{38, 42, 43} The model proposed by Ang et al. had been tested on

the stage III/IV tumors treated with concomitant chemotherapy, and also on the total population.³⁸

7.5. DISCUSSION

7.5.1. The implementation of the 8th edition of the UICC classification system is a step forward for staging HPV-associated OPSCCs

In the literature, the predictive value of the 7th edition of the TNM classification for oropharyngeal carcinomas, and for N status in particular, has shifted over time as a consequence of the epidemic of HPV-associated HNSCC's.¹³⁻³⁴ This shift prompted the publication of a new edition of the UICC staging system. The 8th edition introduced a separate classifying system for HPV-positive tumors. In this study we investigated an unselected group of 368 patients with carcinomas of the tonsil, which is the site associated with the highest prevalence of HPV. In total, 110 tumors tested HPV-positive with p16, PCR and/or FISH. The influence of HPV presence on the prognostic value of the 7th and 8th UICC tumor staging system as well as on individual T, N and M status was examined, taking into account the influence of patient-associated clinical variables including tumor differentiation grade, age, smoking behavior, alcohol consumption and treatment.

Altogether, when we investigated the prognostic value of T, N and M status in both classification systems (7th and 8th edition), only two criteria -- lymph nodes larger than 6cm (cN3), and the presence of distant metastases (M1) – were correlated with a poor prognosis in HPV-positive TSCCs.

Regarding the 7th classification system, our study did not reveal a difference in survival between tumor stages I to IVa, which are the staging groups that represent 90% of the HPV-positive tumors. This prognosis, which is favorable even within the more advanced stages, might be the result of a better response to treatment by HPV-positive tumors. Another explanation for the favorable prognosis might lie in the tumor biology which is associated with HPV. HPV-positive tumors are associated with smaller, less advanced T stages despite the presence of more advanced N stages.

For N status, no prognostic value in HPV-positive TSCCs could be demonstrated when using the 7th classification system. Only N3 status was correlated with a worse survival. Our review of the literature on outcome of treatment of TSCCs demonstrated the loss of prognostic value for N status over time (table 1). The causal relationship between this shift in prognostic value of N status and the rising incidence of HPV-associated carcinomas has been observed previously.^{35,36} Again, the HPV-associated tumor biology, with smaller primary tumors despite advanced-stage lymph nodes, may have a beneficial effect on the response to therapy.

Re-staging for HPV-associated carcinomas has been proposed in the literature since 2009. Spector et al. reported that the AJCC classification system for N status might

be unreliable in predicting outcome for HPV-positive oropharyngeal tumors.³⁹ A new classification system was proposed: “HPV+ N1” was defined as a single node <6cm, ipsilaterally or contralaterally; “HPV+ N2” was defined as a single node ≥6cm or ≥2 nodes ipsilaterally/contralaterally or ≥3 nodes bilaterally; “HPV+ N3” was defined as matted nodes. Applying that classification to our study population, no differences were found between the proposed “HPV+ N1”, “HPV+ N2” and “HPV+ N3” because the number of lymph nodes >6cm was relatively small and survival for HPV-positive N2b necks was good.

Ang et al. proposed a new classification system for oropharyngeal tumors.³⁸ In the HPV-positive groups, smoking status discriminated between mild and moderate risk. The same authors also reported a difference in outcome between N0-2a versus N2b-3. However, in the study of Ang et al. only stage III and IV OPSCCs were included, which were treated with radio- and chemotherapy. In our patient population, in contrast, we did not find a survival difference between N0-2a and N2b-3, nor did we find it when correcting for smoking behavior.

Huang et al. divided 537 oropharyngeal carcinomas without distant metastases into 4 groups: “T1-3/N0-N2c & ≤20 pack years of tobacco smoking”, “T1-3/N0-N2c & >20 pack years of tobacco smoking”, “T4 or N3 & age ≤70 years” and “T4 or N3 & age >70 years”.⁴⁰ The 5-year overall survival rates for the 4 groups were respectively 89%, 64%, 57% and 40%. Again, in our study no significantly different survival could be demonstrated between those groups.

Altogether, different cut-off points have been chosen in studies proposing an HPV-dependent tumor classification system. The results of the ICON-S study were finally adopted for the 8th edition of the UICC tumor staging system.⁴² Cramer et al. recently validated this classification system in a population of more than 15,000 patients (USA) and demonstrated a better stratification of the tumor staging for HPV-positive patients.⁴⁴ Also for T, cN and pN status, the prognostic value could be validated in the HPV-positive population for that study.

Unlike Cramer et al., we focused on tonsillar carcinomas, studied a smaller patient population, and used HPV16-DNA PCR and/or FISH in addition to p16-IHC, whereas P16-IHC was the only detection method used by Cramer et al. These methodological choices might lead to differences in the results. Even so, in our study population, the 8th edition of the UICC tumor staging system was associated with a better prognostic value for tumor stage. The 8th edition was also associated with a better stratification for evaluating the impact of anatomical tumor characteristics on survival.

However, our study showed that there were no prognostic differences between N stages cN0 to cN2 and pN0 to pN2. Only cN3 status and M1 status were significantly associated with prognosis, and no differences were found between clinical N stages cN0 to cN2. Remarkably, there even seemed to be a better prognosis for cN1 than for cN0. As this study population included TSCCs only, the amount of contralateral or bilateral involved lymph nodes (cN2-status) was very limited.

In literature, MacKenzie et al. also reported that only lymph nodes larger than 6cm (cN3) were associated with worse survival.⁴⁷ As mentioned, Cramer et al. had successfully validated the prognostic value of the clinical N-status (8th edition), however their tables did not show a significant difference in survival between cN stages cN0 versus cN1 and cN2 M, and even so reported a better prognosis for cN1 versus cN0.⁴⁴

All in all, the staging system for HPV-associated clinical N status as now included in the 8th edition UICC TNM classification still does not appear to be fully representative of HPV-associated tumor biology. Additional variables need to be addressed to allow even more adequate stratification of HPV-positive tumors. Possible opportunities in our study population to improve the prognostic value of the 8th edition of the HPV-associated UICC TNM classification are discussed below.

7.5.2. Role of non-anatomical characteristics

The 8th edition of the clinical classification system is the result of a study published by O'Sullivan et al. (ICON-S).⁴² The predictive staging model adopted in that study included only anatomical features of the HPV-associated tumors. The earlier recursive partitioning analysis RTOG (2010) by Ang et al. included non-anatomical parameters such as age and tobacco smoking and reported that tobacco smoking had important prognostic value.³⁸ Results of the study by Ang et al. were validated by others.⁴⁸⁻⁴⁹ Regarding smoking, Marur et al. noticed that treatment failures in a de-escalating regime of combining cetuximab with radiotherapy were seen in smokers (>10 pack years).⁵⁰ However, Haigentz et al. emphasized that including smoking in a predictive model has great limitations because of the lack of validated, prospective data and the subjectivity of data collection on tobacco use.⁵¹ In our predictive model, non- or former smokers had a very favorable prognosis of more than 95% 5-year OS, even in more advanced tumor stages. It need to be emphasized that this groups of non- or former smokers include also patients that quitted smoking for more than 10 years despite having smoked numerous packyears. Moreover, the number of pack years was probably therefore not a predictor in prognosis in our patient population. Concerning the role of age, both Ang et al. and Huang et al. found that age influenced prognosis in OPSCCs in their studies.^{38,40} In our data, we observed a prognostic role for age with a cut-off point of 65 years.

In the present study, outcome of HPV-positive TSCCs was not predominantly TNM-status dependent, even when using the 8th edition. The most significant prognostic factors in HPV-positive TSCCs were smoking, age, N3 status and the presence of distant metastases. Therefore, we developed a new prognostic model comprising four groups. The first group consisted of non- or former smokers (patients who had quit smoking more than 10 years prior to the diagnosis of TSCC), and was associated with a 5-yr OS of 95.1%. Group 2 included smokers aged 65 or younger with an associated overall survival rate of 75.6%. In group 3 patients who smoked and were older than 65 had a 5-yr OS of 46.2%. Patients with N3 and M1 status comprised group 4. Interestingly, the two patients who survived in this latter group were the only non-smokers with N3-staged

necks. Within the different groups, survival was not differentiated by T and/or N status. Therefore, this new model appears to provide a simple additional tool for predicting outcome in the clinical setting.

The role of non-anatomical parameters is not taken into account in the 8th version of the UICC classification system. The present study is the first in which both age and smoking behavior play a role in classifying HPV-associated tumors in a way that is almost without reference to TNM classification. Our findings suggest that outcome in HPV-positive TSCCs is influenced by patient characteristics including senescence and intoxication (factors such as tobacco use) regardless of tumor stage. Therefore, these factors are unavoidable when considering response to therapy and rate of survival.

7.5.3. How to address the N0 neck

In our patient population, a clinically negative neck status was not associated with a better prognosis than for N1. The cut-off point for prognostic value in our study of tonsillar carcinomas was N3, which meant that very few bilaterally involved (and thus N2) neck stages were diagnosed. The prognosis for the N0 neck was even worse than for the N1 neck. As already mentioned regarding the study by Cramer et al., in which the 8th edition was validated in a population of more than 15,000 HPV-positive oropharyngeal tumors, cN1-status was associated with a significantly better survival than N0. Moreover, the bilaterally involved neck (cN2 status) was not associated with a significantly worse survival than the clinically negative neck after adjustment for age, sex and race. Only cN3 status was significantly associated with a worse survival.⁴⁴ In previous research, it was noticed that patients with HPV-positive carcinomas more often had a lymph node as presenting symptom when compared to their HPV-negative counterparts. Presenting with that symptom may have led to an earlier discovery of the primary tumor.³⁵ It is therefore possible that these presenting “alarming” nodes are associated with a better prognosis, as we found in our study. Also Fritsch et al. and Ang et al. found that patients with an HPV-positive single neck node between 3 and 6cm in size (N2a, 7th edition) had a better outcome than patients without lymph node metastases.^{37,38} Fritsch et al. compared outcome based on N status between HPV-dominant (tonsillar fossa and base of tongue) and non-HPV-dominant oropharyngeal subsites in more than 15,000 oropharyngeal tumors.³⁷ In their HPV-dominant population, cN2a (7th ed.) had a better survival rate than the N0/1 patients. In the total population, no differences in outcome were noted as long as lymph node metastases were unilateral (<cN2c). Our study only included patients with TSCC; in this strongly HPV-dominant subgroup, similar results were found.

7.5.4. Role of oropharyngeal subsite

The influence of HPV on prognosis is often analyzed for all oropharyngeal subsites, without even discriminating between HPV-dominant and non-dominant subsites. For the present study, we selected a group of 110 patients with squamous cell carcinomas of the tonsil, the most HPV-dominant oropharyngeal subsite. According to Sood et al.,

a bilaterally involved neck status is predominantly seen in tumors at the base of the tongue, indicating that lymph node dissemination patterns differ even within the HPV-dominant oropharyngeal sites.⁴⁴ This may explain why only a few bilaterally involved necks are seen in our population of tonsillar carcinomas. It may also explain the lack of significance of the cN2 neck in our study where we classify bilateral neck involvement in accordance with the 8th edition.

7.5.5. HPV-detection

P16 immunohistochemistry is a widely available, low-cost test, unlike the more complex HPV-in situ hybridization. In our study, p16 was combined with HPV-DNA PCR and/or FISH. We found that 15 out of 124 p16-positive patients were not HPV-positive (12%).

Nauta et al. raised the issue of consensus on the exact definition of HPV-associated OPSCC.⁵² HPV infection alone is not sufficient to classify an instance of OPSCC as HPV-related since the presence of HPV-DNA could merely reflect a transient infection. Detection of p16 alone is not specific for HPV-activity. Smeets et al. described an algorithm in which p16-IHC was combined with HPV 16 DNA PCR.⁵³ Taberna et al. recently investigated the outcome of HPV-positive OPSCC in relation to the definition for HPV-positivity.⁵⁴ They concluded that definitions of HPV positivity have an impact on TNM classification and patients' survival. Bussu et al. recently confirmed that p16-IHC alone may not be specific enough to become the diagnostic standard from the perspective of treatment de-intensification. Standardization of clinical use and of detection methods for HPV as a marker for molecular characterization in head and neck oncology is warranted.⁵⁵

7.6. CONCLUSION

Altogether, our results are in line with those from other studies confirming that the introduction of the 8th HPV-associated tumor staging system is a step forward in staging HPV-associated OPSCC. However, our study indicates that the prognostic value is improved by including smoking history and age as additional prognostic factors. To further optimize stratification, issues such as the relevance of N status smaller than 6cm, influence of subsite, and HPV-detection method remain to be addressed.

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Figure 1. HPV-prevalence increased from 2002 until 2011

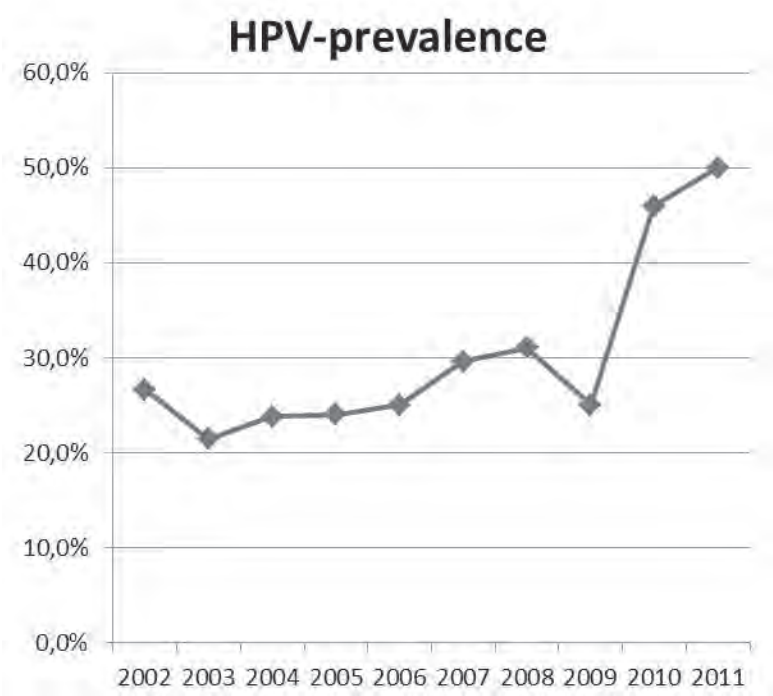
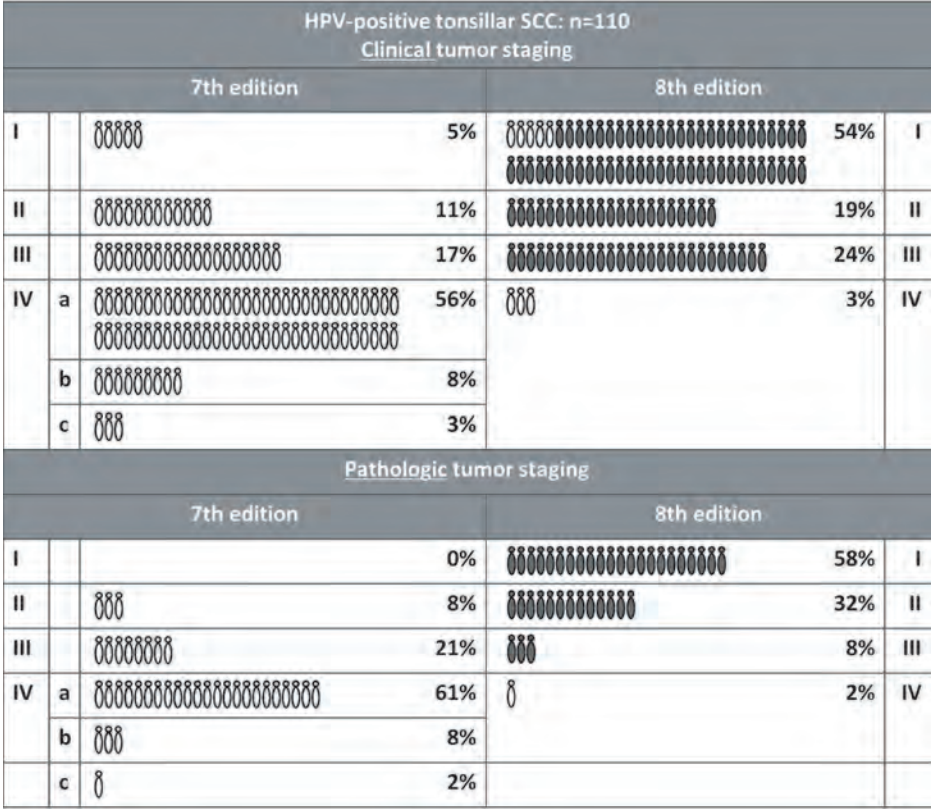


Figure 2. Changes in distribution of HPV-positive tonsillar SCCs between the 7th and 8th edition of the UICC tumor staging, resp. clinical and pathologic staging.



Legends: Changes in the distribution of patients with human papillomavirus-positive disease in the 7th and 8th edition staging systems. Each puppet represents a unique patient included in this study (n=110). The white color represents patients in the 7th edition guidelines who did not change stage groups. The gray color represents patients who changes stage groups in the 8th edition guidelines. In the bottom half of the figure, patients are included in which pathologic data are reported after neck dissection (n=38).

SCC: squamous cell carcinoma

Figure 3. Outcome of HPV-positive tonsillar SCCs staged with UICC clinical tumor staging 8th edition, related to smoking habits.

HPV-positive tonsillar SCC Clinical tumor staging: 8th edition			Non- or former smoker	Smoker
Stage	Number of patients: n=110	5-year OS	n=41	n=68
I	 n=60	83%	95%	77%
II	 n=21	76%	90%	64%
III	 n=25	72%	100%	53%
IV	 n=3	0%	-	0%
Total		77%	95%	66%

Legend: SCC: squamous cell carcinoma; OS: overall survival.

White colored puppets represents each single patient within the depicted tumor stages who were non- or former smokers (quitted more than 10 years before diagnosis). Gray colored (smoking) puppets represents respectively the smoking individuals included in this study.

5-year OS non- or former smokers versus smokers: p <0,001 (Log rank)

Figure 4a. Overall survival in presented predictive model

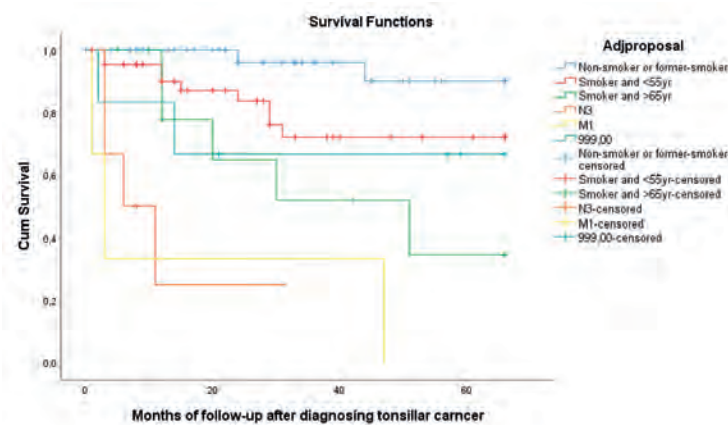


Figure 4b. Overall survival according to the predictive model based on the recursive partitioning analysis (RPA) published by Ang et al. (p-value: not significant)³⁸

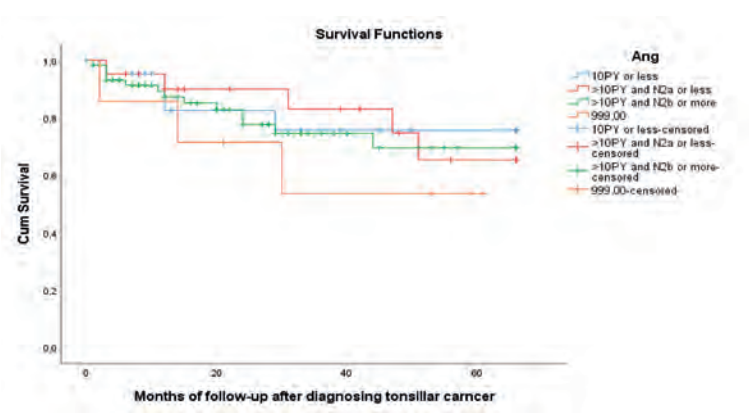


Figure 4c. Overall survival according to the predictive model based on the recursive partitioning analysis (RPA) published by O'Sullivan et al. (p-value: not significant)⁴⁰

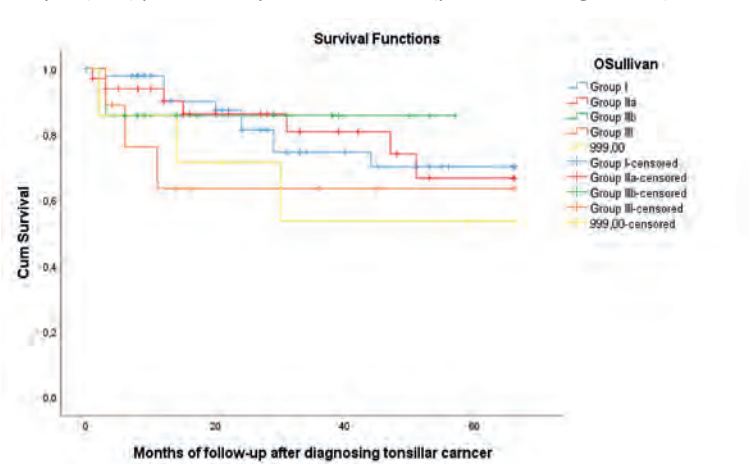
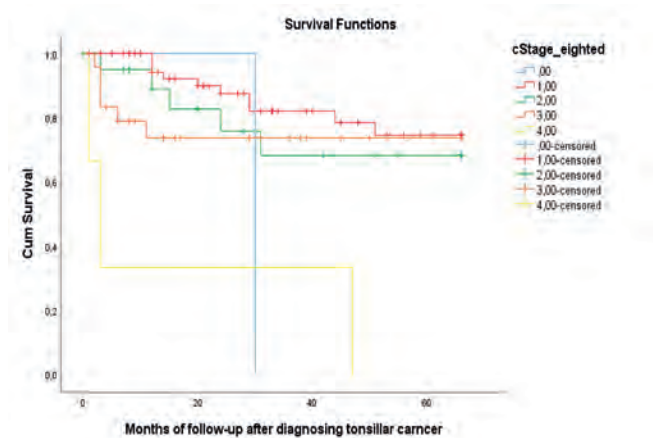
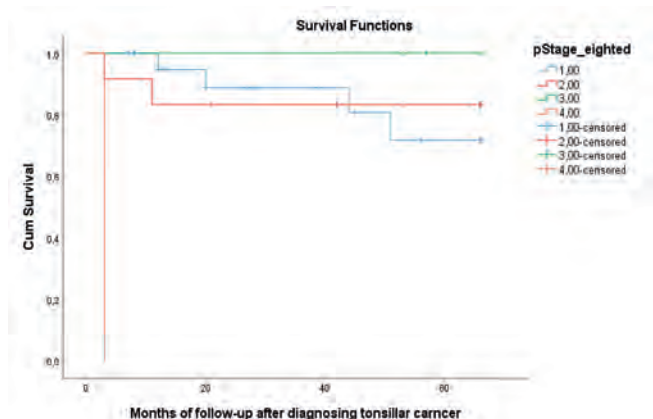


Figure 4d. Overall survival according to the predictive model based on the adjusted hazard ratio (AHR) published by O’Sullivan, defined as the ICON-S classification and used for the 8th edition UICC clinical tumor staging (p-value: $p < .001$).⁴⁰



- 1: Stage I (corresponding to T0-2 AND N0-1 (8th UICC))
- 2: Stage II (T0-2 AND N2; T3 AND N0-2)
- 3: Stage III (T1-4 AND N3; T4 AND N0-3)
- 4: Stage IV (any T AND any N AND M1)

Figure 4e. Overall survival according to the predictive model published by Haughey et al.: i.e. 8th edition UICC pathological tumor staging (p-value: not significant).⁴²



- 1: Stage I (corresponding to T1-2 AND N0-1 (8th UICC))
- 2: Stage II (T3-4 AND N0-1; T1-2 AND N2)
- 3: Stage III (T3-4 AND N2)
- 4: M1

Table 1. Prognostic value of N-status in TSCC in international literature between 1978-2010*

	Prognostic value of N-status		Site	Patients (number)	Inclusion UICC/AJCC		Therapie	Statistics	Survival tested based on N-status	P-value
	NO>N+	NO-1>N2-3			NO-1=N2-3	(years)				
Edström	1978	X*	tonsil	37	50-'70	AJCC '67	RT	none	5yr OS	unknown
Mantravadi	1978	X*	tonsil	117	55-'73	AJCC '67	combined	none	2-5yr DFS	unknown
Petrovich	1980	X*	tonsil	205	46-'76	AJCC '78	combined	none	5yr OS	unknown
Tong	1982	X*	tonsil	104	65-'75	AJCC '78	RT	none	3yr OS, DFS, LRC	unknown
Dubois	1983	X*	tonsil	215	70-'76	UICC/WHO '62-'79	RT	none	3-5yr OS	unknown
Orregia	1983	X	tonsil	79	60-'79	UICC '78	combined	life-table method, log rank	5yr OS	0,001
Amornmarn	1984	X*	tonsil	185	56-'77	AJCC '83	RT	none	5yr DFS, LC, RC	unknown
Mizono	1986	X*	tonsil	203	57-'79	AJCC '78	combined	none	5yr DFS	unknown
Vallis	1986	X	tonsil	87	62-'81	UICC '78	RT	life-table method, log rank	10yr OS, DFS	0,001
Lusinchi	1989	X	tonsil	193	70-'82	UICC '79	RT	Kaplan-Meier	3-5yr OS	unknown
Di Marco	1990	X (no N2)	tonsil	183	70-'84	UICC '78	RT	log rank	5yr OS	<0,005

Table 1. Continued.

	Prognostic value of N-status	Site			Patients (number)	Inclusion (years)	UICC/AJCC Therapie	Statistics	Survival tested based on N-status	P-value
	NO>N+	NO- 1>N2-3	NO=N+ 1=N2-3	NO- 1=N2-3						
Mak-Kregar	1990		X	X	92	66-'85	UICC '82 versus UICC '87/ AJCC '88	combined Kaplan- Meier, Cox regression	3yr DFS	NS
Al-Abdulwahed	1997		X		102	75-'95	UICC '87 (restaged)	combined Kaplan- Meier	5yr OS, 4yr DSS	0,02
Perez	1998	X			384	59-'91	Macomb & Fletcher '67	combined not described, (life-table method ?)	4-10yr DFS	multivariate DFS: 0,001
Friesland	1999		X		167	80-'95	AJCC '97	combined Kaplan- Meier, Cox regression	5-15yr OS, DSS	0,0057
Mellin	2000		X		60	84-'96	UICC '97	combined Kaplan- Meier, Cox regression	2-10yr DSS	NS
Charbonneau	2006	X			164	90-'99	AJCC '97	combined Kaplan- Meier, Cox regression	2-5yr OS	univariate OS: 0,0028
										multivariate OS: 0,043

Table 1. Continued.

		Prognostic value of N-status	Site	Patients (number)	Inclusion (years)	UICC/AJCC	Therapie	Statistics	Survival tested based on N-status	P-value
Pitkin	2007	X	tonsil N0-N2-3	84	90-'03	AJCC '97	combined	Kaplan- Meier, Cox regression	2-13yr OS, DSS, DFS	univariate DFS: 0,248 univariate DSS: 0,005 multivariate DSS: N1 0,521 N2 0,022 N3 0,832 univariate DSS: 0,557
Chien	2008		tonsil (HPV)	111	92-'05	AJCC '97	combined	Kaplan- Meier, Cox regression	5yr DSS	
Hafkamp	2008		tonsil	81	92-'01	AJCC '97	combined	Kaplan- Meier, Cox regression	5yr OS, DSS	NS, Unadjusted HR: 1,0 (95%CI: 0,5- 2,9)
Aziz	2010		tonsil	69	98-'00	AJCC '97	RT en brachy	Kaplan- Meier, Cox regression	3yr OS, DSS	univariate DSS: 0,304

Table 1. Continued.

	Prognostic value of N-status	Site	Patients (number)	Inclusion UICC/AJCC	Therapie	Statistics	Survival tested based on N-status	P-value
	N0>N+	N0- 1>N2-3	N0=N+ 1=N2-3					
Bachar	2010	X	tonsil 640	70-'90	niet beschreven	combined Kaplan- Meier, Cox regression	20yr OS, DSS	univariate OS: 0,0003 multivariate DSS N1 0,0654 N2 0,0483 N3 <0,0001

Results of the meta-analysis performed in Pubmed: inclusion criteria: studies 1960 - July 2013; search terms: "cancer", "neoplasm", "carcinoma*", "oropharyn*", "tonsil*", "lymph*", "node*", "nodal", "neck", "survival*", "prognos*", "mortality", "morbidity", "outcome". 634 abstracts were evaluated for inclusion, and in case of doubt the full text article was read. References of all read articles were checked. In total, 22 articles were selected for inclusion (publication years: 1978-2010; range in number of patients per study: 37-640; range in year of inclusion: 1950-2005).

Table 2. Characteristics of patients with tonsillar SCC: total study population and relation with HPV-status

Patient characteristics		Total (n=368)	%	HPV-positive* (n=110)	HPV-negative (n=225)	p-value
Sex	male	264	71.7	81	161	NS
	female	104	28.3	29	64	
Age	mean	60,32		60.1	60.55	NS
	range	39-87		39-84	41-87	
Age (years)	<55	111	30.2	36	66	NS
	55-65	159	43.2	38	103	
	65	98	26.6	36	56	
Smoking	non-smoker	34	9.2	21	9	<0.001
	former-smoker (#>10 years)	34	9.2	18	13	
	smoker	265	72	63	181	
	unknown	35	9.5	8	22	
Alcohol	none or <1 unit/day	74	20.1	34	36	0.003
	1-2 units/day	70	19	23	36	
	>2 units/day	187	50.8	45	129	
	unknown	37	10.1	8	24	
Tumor differentiation (n=191)	G1	24	6.5	7	16	0.041
	G2	108	29.3	24	79	
	G3	57	15.5	24	29	
	undifferentiated	2	0.5	1	1	
p16	positive	124	37	109	15	<0.001
	negative	211	62.9	1	210	
Previous cancer in medical history	head and neck carcinoma			3	16	NS
	other			2	4	
Distant metastases	present			3	3	NS
Second primary tumor				2	14	NS
Therapy	surgery	18	4.9	3	11	NS
	(surgery) + radiotherapy	(93) 237	(25.3) 64.4	(42) 78	(44) 139	
	(surgery) + concomittant chemoradiotherapy	(10) 79	(2.7) 21.5	(3) 21	(6) 49	
	none / palliative	23	6.1	7	16	
	unknown	11	3	1	10	

*HPV-association could be tested in the histologic specimens of 335 patients

Legends: SCC: squamous cell carcinoma

Table 3. Tonsillar SCC staging in the total study population and in relation to HPV using the 7th edition UICC tumor staging system

		Total population n=368		HPV- status n=335		p-value
		n	%	positive (n=110)	negative (n=225)	
7th edition tumor staging						
I	26	7.1	5	16	NS	
II	38	10.3	11	24		
III	78	21.2	20	49		
IV a	182	49.5	61	108		
IV b	32	8.7	9	21		
IV c	7	1.9	2	5		
?	5	1.4	2	2		
cT-status						
1	74	20.1	22	42	0.037	
2	112	30.4	44	57		
3	82	22.3	23	54		
4a	83	22.6	17	61		
4b	14	3.8	3	9		
?	3	0.8	1	2		
T1,2 vs T3,4			67 vs 42	99 vs 124	0.014	
cN-status						
0	113	30.7	27	73	NS	
1	60	16.3	16	38		
2a	14	3.8	5	9		
2b	112	30.4	43	62		
2c	39	10.6	11	24		
3	23	6.2	6	15		
?	7	1.8	2	4		
N0 vs N+			28 vs 81	73 vs 150	NS	
N0-1 vs N2-3			43 vs 66	112 vs 111	NS	
cM-status						
1	6	1.7	3	3	NS	

Legends: NS: not significant; SCC: squamous cell carcinoma

Table 4. Comparison of five-year overall survival rates in tonsillar squamous cell carcinomas (n=368) according to the 7th and 8th edition UICC tumor staging system

Five-year overall survival					
HPV-negative (n=225)			HPV-positive (n=110)		
	7th edition		7th edition	8th edition: clinical	8th edition: pathological
I	75.0%		80.0%	83.3%	81.8% I
II	79.2%		81.8%	76.2%	83.3 II
III	59.2%		75.0%	72,0%	100,0% III
IV	a	42.1%	85.0%	0,0%	0,0% IV
	b	33.3%	55.6%		
	c	20.0%	0.0%		
p-value	<0.001		<0.001	<0.001	<0.001

Table 5a. Univariate Cox regression analysis of overall survival (OS) in HPV-positive tonsillar SCC (n=110): T-status, clinical N-status (cN-status) and pathological N-status (pN-status) using the 8th edition UICC staging system.

T-status	p-value	HR		p-value	HR		p-value	HR
T1		1	cN0		1	pN0		1
T2	.295	1980	cN1	.534	.744	pN1	.966	95604.162
T3	.152	2686	cN2	.656	.699	pN2	.969	40630.224
T4	.459	1761	cN3	.004	6541			
T-status	.553*		cN-status	.003*		pN-status	.731*	

Legends: SCC: squamous cell carcinoma; HR: hazard ratio

*log rank, Kaplan-Meier

Table 5b. Univariate Cox regression of overall survival (OS) in HPV-positive tonsillar squamous cell carcinomas (n=110): non-anatomical variables.

		Hazard ratio	95% confidence interval
Age	> 65 years	2.30	1.05 - 5.06
Sex	male	1.27	0.51 - 3.19
Smoking status	Smokers versus non-smokers	4.64	1.10 -19.19
	Smokers versus non-and former smokers	5.32	1.58 -17.91
Alcohol intake	<1 unit per day	1	
	1-2 units per day	2.39	0.73 - 7.8
	> 2 units per day	1.87	0.65 - 5.40
Tumor differentiation	G1	1	
	G2	1.40	0.37 - 5.26
	G3	1.21	0.47 - 3.14
M-status	1	8.77	2.60 - 29.64
Treatment	Surgery as monotherapy	1	
	Surgery with radiotherapy	0.64	0.085 - 4.89
	Surgery with combined chemoradiotherapy	0.22	0.014 - 3.63
	Palliative treatment	11.5	1.15 - 98.25

Table 5c. Multiple regression analysis of overall survival (OS) in HPV-positive patients including the following variables: smoking status, age more than 65 years, M-status and lymph node size more than 6 cm.

	p-value	Hazard ratio
Smoking	.006	5723
Age > 65	.692	1225
M-status	.009	7118
N3 vs N012	.000	14862





Next-generation treatment strategies for human papillomavirus-related head and neck squamous cell carcinoma: where do we go?

Published: Olthof NC[†], Straetmans JM[†], Snoeck R, Ramaekers FC, Kremer B^{*}, Speel EJ^{*}. Next-generation treatment strategies for human papillomavirus-related head and neck squamous cell carcinoma: where do we go? Rev Med Virol 2012;22:88-105.

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8.1. SUMMARY

Oncogenic human papillomavirus (HPV) is currently recognised as a major risk factor for the development of head and neck squamous cell carcinomas (HNSCC). HPV is mostly detected in tumours arising from the oropharynx and more specifically from the tonsil. HPV-related tumours display clinical and molecular characteristics that are distinct from HPV-unrelated tumours, which are generally induced by alcohol and tobacco abuse. Detection of biologically active HPV in HNSCC has prognostic relevance, which warrants the separate classification of HPV-induced tumours and is a prerequisite for further optimisation of treatment protocols for this distinct group. Current guidelines for the treatment of oropharyngeal squamous cell carcinoma (OPSCC) have not incorporated specific treatment modalities for HPV-related tumours. The development of such treatment options is still in a preclinical phase or in early clinical trials. Recent data on treatment response of OPSCC have been obtained by retrospectively analysing HPV-status and indicate that patients with HPV-related tumours show a favourable prognosis, independent of the type of treatment. These patients may benefit from de-intensified treatment, which should be assessed in prospective clinical trials. The development and future use of new antiviral and immunomodulatory therapeutics may be instrumental in this approach to improve survival rates and decrease disease- and treatment-related morbidity. In this review we will focus on present therapeutic HPV-targeting strategies and discuss future directions for de-intensified treatment of HPV-positive HNSCC.

8.2. INTRODUCTION

Head and neck cancer is a serious health care problem in many parts of the world.¹ The vast majority of head and neck cancers are squamous cell carcinomas originating from the mucosal epithelium lining the oral cavity, nasal cavity, pharynx and larynx.² In 2008, head and neck squamous cell carcinomas (HNSCC) were estimated to cause 480 000 new cancer cases and 273 000 cancer deaths worldwide.¹ Despite the fact that advances have been made in diagnosis and treatment, mortality rates have only marginally decreased over the last decades and the 5-year survival rate currently ranges between 40%–60%.³ Approximately 80%–90% of HNSCC develop in patients with a history of alcohol and tobacco abuse, including tobacco and betel quid chewing and snuff dipping.⁴ These factors are also responsible for the process of ‘field cancerisation’ in the entire head and neck region,⁵ leading to multiple primary tumours in up to 40% of patients.⁶ Patients without exposure to these risk factors account for 10%–20% of HNSCC. These tumours are predominantly associated with viral carcinogenesis, including infection with EBV in nasopharyngeal carcinomas⁷ and, to a greater extent, infection with oncogenic human papillomavirus (HPV) in the oropharynx, in particular in the lingual and palatine tonsils. In the last decade, the incidence of HPV-related oropharyngeal squamous cell carcinoma (OPSCC) has increased relative to the total group of HNSCC.^{4, 8} Infection rates in OPSCC range from 20% to more than 90% in different studies, depending on geographical factors and the detection method used.^{9–12} In this review, we will present the clinical and molecular features of HPV-positive HNSCC. Subsequently we will focus on the current knowledge of potential anti-HPV strategies and discuss the most promising modalities for the treatment of HPV-positive HNSCC.

8.3. METHODS

Besides relevant articles selected from the general literature concerning HPV-related carcinogenesis and references therein, specific literature on treatment options for HPV-related HNSCC was obtained by a bibliographical search in PubMed, Medline and Embase, from inception to May 2011, using the search term (HPV OR papillomavirus OR papilloma) AND (HNSCC OR ‘head and neck cancer’ OR oropharyngeal OR oropharynx OR oral OR pharyngeal OR pharynx OR buccal OR base of tongue OR tongue OR tonsillar OR tonsil OR floor of mouth OR mouth OR vallecula) AND (treatment OR antiviral OR therapy) AND (cancer OR carcinoma OR tumour OR tumour OR neoplasm). This search yielded 1246 results in PubMed, 137 in Medline and 309 in Embase. Based on inspection of the title and/or abstract of these publications, 63 relevant papers on treatment options and some references therein were included in this review. Moreover, ongoing clinical trials concerning new therapeutic options for HPV-related HNSCC were identified from the Cochrane Controlled Trial Register and from the US National Institute of Health Clinical Trials (www.clinicaltrials.gov), yielding five relevant results.

8.4. HUMAN PAPILLOMAVIRUS AND TUMOURIGENESIS

8.4.1. Human papillomavirus

Human papillomaviruses are non-enveloped viruses, containing circular double-stranded DNA of approximately 8 kb, that are highly epitheliotropic and known to infect both mucosal and cutaneous epithelia.¹³ Papillomaviruses are species-specific and the human papillomavirus family can be classified into five genera and subdivided into 31 species and 120 types.¹⁴ A subgroup of 15 HPV types is linked to the development of malignant lesions of mucosal and cutaneous epithelia, and is considered to comprise high-risk (HR) HPVs.¹⁵ All HR-HPVs belong to the alpha-genus, including HPV-16 and HPV-18, which are found in ~50% and ~20% of cervical malignancies, respectively.¹⁶ Differences in the capacity to deregulate cellular protein function by viral oncogenes E6 and E7 account for the carcinogenic properties of HR-HPV in comparison with low risk (LR) HPVs.^{17, 18} LR-HPV types, such as HPV-6 and HPV-11, are often found in benign mucosal lesions and are only sporadically associated with carcinomas. Human lesions in which HPV types of the alpha-genus appear to be involved are summarised in Table 1.

8.4.2. Human papillomavirus replication and integration

The HPV life cycle is linked to the differentiation of the infected epithelial cell. HPV infection is initiated by binding of the virion L1 protein to heparan sulphate proteoglycans (HSPG) on segments of the basement membrane, which are exposed at sites of (micro)injury. This induces conformational changes and L2 cleavage finally resulting in binding of the L1 capsid protein to a so far undetermined cell surface receptor.¹⁹ The cell adhesion receptor α 6-integrin has been implicated to be this receptor,²⁰ but does not seem to be essential for HPV infection. However, α 6-integrin might be a matrix component closely associated with HSPG.²¹ The circular HPV DNA comprises 8 genes, coding for six early (E) and two late (L) proteins (Figure 1). The E-proteins regulate and facilitate virus-replication and are expressed early after infection.²² Oncoproteins E6 and E7 have a direct effect on several essential cellular processes, such as cell cycle and apoptosis regulation. E6 promotes degradation of p53 through interaction with E6-associated protein (E6AP), an E3 ubiquitin ligase, and subsequent ubiquitination and proteasomal degradation. Amongst others, this alters transcription of p53 target genes and activates human telomerase reverse transcriptase (hTERT), resulting in cell survival and ultimately in genetic instability.^{23, 24} The oncoprotein E7 binds to the unphosphorylated retinoblastoma tumour suppressor protein (pRb), which promotes the release of transcription factor E2F, leading to activation of the cell cycle and transition through the G1/S-phase, needed for DNA-replication.²⁵⁻²⁷ As a consequence, p16INK4A is upregulated but is unable to properly inhibit the cell cycle. Expression of oncoproteins E6 and E7 is tightly regulated by E2, the main regulator of viral gene transcription.²⁸ Molecular studies have shown that integration of HPV often leads to a

disruption in the E1/E2 open reading frame and concurrent loss of the E4 and E5 and parts of the E2 and L2 genes.²⁹ E2 function can moreover be abrogated by epigenetic alterations of the viral genome such as methylation of the E2 binding site in the long control region.³⁰ Absence of E2 function results in upregulation of the expression of oncoproteins E6 and E7, which in turn leads to uncontrolled cell cycle progression (see Figure 1).

The major structural protein L1 of the HPV capsid is sufficient for self-assembly into a capsid, but entry of the virus into the cell is co-dependent on L2, the minor structural protein.^{19, 31}

Under normal circumstances, HPV maintains an episomal state, and infection with HPV is transient. In a recent prospective cohort study, the reported average duration of active episomal infection in the uterine cervix appears to be approximately 8 months.³² Although uterine cervical HPV infection prevalence decreases with increasing age,³³ it is unclear whether age affects the duration of infection. Persistent infection, however, might lead to integration of the virus.^{34, 35} Numerous investigations have shown an etiological relationship between infection with HR-HPV infection and the development of uterine cervical squamous cell carcinomas (UCSCC) and other anogenital squamous cell carcinomas.^{36, 37} More than 90% of UCSCC contain and express HR-HPV sequences, which are predominantly present in an integrated form.^{38, 39} HPV-16 is the most common HPV type and is detected in more than 50% of UCSCC, followed by HPV-18, HPV-33 and HPV-45 (Table 1).^{37, 40}

The precise relationship between HR-HPV integration and head and neck carcinogenesis is less clear, partly because primary premalignant lesions of the oropharynx are seldom detected. Although controversial data have been reported,⁴¹⁻⁴⁴ integration of HR-HPV in OPSCC is a prevailing finding.

8.4.3. High-risk human papillomavirus in head and neck squamous cell carcinoma

Patients with a history of HPV-related anogenital carcinomas, patients seropositive for HPV-16, and husbands of patients diagnosed with uterine cervical dysplasia or carcinoma in situ all show increased risk rates for developing OPSCC.⁴⁵⁻⁴⁷

The involvement of HPV in head and neck tumourigenesis was first proposed by Syrjänen et al.,⁴⁸ who showed histopathological features of HPV infections in 40% of patients, and HPV-positive nuclei in 20% of patients using immunohistochemistry. Since then many studies have provided evidence that infection with HR-HPV is a significant independent risk factor for HNSCC and is associated with high-risk sexual behaviour.^{9, 10, 49-51} HR-HPV positive tumours are most frequently found in the oropharynx and are associated with HPV-16 in >90% of cases.^{9, 49, 52, 53} Because patients with OPSCC often present with metastatic disease at first diagnosis, information on the persistence of oropharyngeal HPV infections and premalignant lesions in this region is scarce.⁵⁴⁻⁵⁶ HPV prevalences of less than 1% have been found in tumour-negative tonsillar tissue samples, screened for HPV with PCR.^{51, 57, 58}

8.4.4. HR-HPV detection and tumour characteristics

The reported overall incidence of HPV in OPSCC ranges from less than 20% to more than 90% in different studies. This variation depends on several factors, including geographical features, sample preparation and detection methods used but also the amount and manner of tobacco consumption depending on geographical location.^{9, 12, 54, 59-61} It has been shown that not all tumours tested positive for HPV DNA can be regarded as etiologically HPV-related.^{50, 62} A clinically relevant infection, that is, a transcriptionally active infection should be present, which can be demonstrated by detectable expression of the viral oncogenes E6 and E7.⁶³ This correlates strongly with overexpression of the CDK inhibitor p16INK4A, which is considered a reliable surrogate marker for HR-HPV infection in most cases.^{41, 64} A reliable algorithm for HPV detection should thus start with p16INK4A detection, followed by in situ hybridisation (ISH) and/or RT-PCR analysis of E6/E7 transcripts after HPV typing,⁴¹ as suggested by two recent reports.^{59, 65} A representative example of these analyses is shown in Figure 2.

The HPV-associated OPSCC are now considered to comprise a separate entity with typical clinical and molecular features. Table 2 summarises the major differences between HPV-positive and HPV-negative OPSCC.

The HPV-positive OPSCC are characterised by overexpression of oncoproteins E6 and E7 leading to degradation of p53 and pRb, thereby inducing cell cycle and apoptosis deregulation. As a result, CDK inhibitors including p16INK4A, p14ARF, p18INK4C and p21Cip1/WAF1 are upregulated, which subsequently leads to downregulation of cyclin D1 and inhibition of complex formation with CDK4.^{22, 62, 66, 67} In HPV-negative tumours, cell cycle deregulation is established by p53 and pRb gene mutations, or alternatively by inactivation of p16INK4A and p14ARF gene expression through mutation, promoter hypermethylation or homozygous deletion,²² or activation of cyclin D1 expression via 11q13 amplification⁶⁸. High expression of EGFR by transcriptional upregulation is generally present in this OPSCC subgroup.^{22, 56, 62, 66-68} Upregulation of EGFR expression is usually not seen in HPV-positive OPSCC.^{56, 67, 69-71}

In addition, global genome and protein scanning approaches have been and are being used to unravel DNA, mRNA, microRNA, and protein signatures specific for HPV-positive and HPV-negative OPSCC. So far, these studies revealed that HPV-positive tumours exhibit a relatively stable genome with 11q and 16q loss,⁷²⁻⁷⁴ and upregulate transcriptional activity of cell cycle regulators (as mentioned previously), transcription factors (e.g. TFDP2, ZNF238, TAF7L and RPA2) and DNA repair proteins (e.g. RFC4 and RFC5). Also, HPV-positive tumours show decreased expression of genes involved in immune responses (e.g. IFIT1, IFITM1-3, IFI6-16, IFI44L, OAS2 and IFN- κ).^{68, 75-79} In addition, these tumours differentially express microRNAs, and for example upregulate miR-363 (belonging to the oncogenic miR-106a-363 cluster) and downregulate miR-218. A recent proteome analysis comparing HPV-positive and HPV-negative oral squamous cell carcinomas (OSCC) reported upregulation of thioredoxin and epidermal-fatty acid binding protein.⁸⁰ Thioredoxin is an important redox-mediator that stimulates cell growth and inhibits apoptosis under adverse conditions, apparently including HPV

infections, as also seen in cervical carcinomas. Epidermal fatty-acid binding protein, although, mainly involved in fatty acid uptake, transport and metabolism, also functions in cellular signalling, affecting differentiation, growth regulation and gene expression.⁸⁰ Although expression of 3q-specific genes has been reported as being specific for HPV-positive OPSCC, this finding remains to be confirmed, because extra copies of 3q-genes have been found in both HPV-positive and HPV-negative tumours.^{68, 77}

8.5. CURRENT TREATMENT OF OPSCC AND EFFECT OF HR-HPV STATUS ON TREATMENT RESPONSE

8.5.1. Current treatment modalities

Current international clinical guidelines for HNSCC treatment mention HPV as a risk factor for OPSCC. The American National Comprehensive Cancer Network has suggested to include HPV detection in the diagnostic work-up of these tumours.⁸¹ However, the treatment guidelines do not offer therapeutic modalities specific for HPV-related tumours. Current therapeutic options include surgery, radiotherapy, chemotherapy (CT), immunomodulatory therapies or combinations of the foregoing. Surgery as primary treatment avoids toxicity caused by radiotherapy and CT but causes loss of function, particularly in patients with larger tumours. The development of laser surgery and transoral robotic surgery (TORS) for OPSCC reduces functional morbidity as a consequence of three-dimensional visualisation and the ability to manipulate and perform reconstruction of the oropharynx without the need for an open surgical approach.^{82, 83} Regional infiltration of critical structures and unacceptable loss of function after surgery can classify a tumour as functionally or technically unresectable. In those cases, radiotherapy and/or CT is the treatment of choice⁸⁴ when aiming at restoring function, however with the disadvantage of therapy-related local and systemic side effects.⁸⁵ The final choice of treatment is based upon clinical variables such as tumour type, localisation and stage,⁸⁶ age of the patient, general medical and psychomedical condition⁸¹ and individual preferences of the patient.

8.5.2. Effect of HR-HPV status on outcome

In retrospective studies, HR-HPV- and/or p16INK4a positive tumours have been found to respond better to multimodal therapies as compared to HPV-negative tumours, thereby favouring patient survival.^{41, 42, 53, 64, 87, 88} More recent retrospective studies have shown that this favourable outcome is independent of treatment modalities.⁸⁹⁻⁹⁵ However, the heterogeneity of the HNSCC patient populations and consequent variability with regard to the HPV and/or p16INK4A status, as well as applied treatment protocols, has most probably negatively influenced the association between HR-HPV status and outcome in these studies. It can be anticipated that the actual difference in clinical outcome

between HPV-positive and HPV-negative cases will become even more pronounced when comparing a homogeneous population of OPSCC and application of reliable detection methods for clinically relevant HPV-infections. Prospective clinical trials are required to further validate HR-HPV presence as predictive factor for therapy outcome and to determine whether treatment de-intensification might improve quality of life while preserving the favourable clinical outcome in HPV-positive OPSCC patients.^{96, 97}

An explanation for the favourable response may lie in the fact that, although the pRb-pathway and p53-pathway are compromised in HPV-positive tumours, they retain some function, such that under the pressure of radiotherapy and/or CT, p53-mediated apoptotic pathways may still function. The presence of wild-type p53 in combination with low levels of Bcl-2/Bcl-xL and EGFR, which are features of HPV-positive tumours in non-smokers, may enhance this treatment advantage.^{9, 53, 63, 67, 69, 71} Moreover, limited tobacco and/or alcohol use reduces field cancerisation and the chance of developing a second primary tumour or distant metastasis in HPV-positive tumours,^{41, 53, 93} which underscores the need to investigate the effect of tobacco and alcohol exposure on the biological behaviour of HPV-positive OPSCC, as recently proposed.^{41, 94}

Besides the overall better survival of patients with HPV-positive tumours, their treatment may be further improved by the implementation of strategies that either 1) promote the immune response to eradicate the virus, 2) inhibit viral DNA replication, 3) specifically target viral oncoproteins or 4) have an effect on deregulated signal transduction pathways specific for HPV-positive tumour cells. In the following section, we review these strategies, their mode of action and possible benefits for patients with HPV-induced HNSCC.

8.6. PROPHYLACTIC AND THERAPEUTIC ALTERNATIVES FOR HPV-POSITIVE OPSCC

8.6.1. Immunomodulating therapies

Vaccination

Two prophylactic vaccines, containing recombinant virus-like particles (VLP) composed of the L1 proteins of the respective HPV types³¹ have been marketed recently, which are Cervarix® (GlaxoSmithKline, Brentford, Middlesex, TW8 9GS, United Kingdom) and Gardasil® (Merck, NJ, USA). Both vaccines have been FDA-approved for use in girls and young women^{98, 99} and Gardasil® has also been approved for use in men.¹⁰⁰

Cervarix® is a bivalent vaccine that protects against infection with HPV-16 and HPV-18, whereas the quadrivalent vaccine Gardasil® provides protection against HPV-6, HPV-11, HPV-16 and HPV-18. Reports indicate that Cervarix® also offers cross-protection against HPV-31, HPV-45 and HPV-52,^{101, 102} and Gardasil® possibly against HPV-31.¹⁰³ More robust cross-protection may be induced by adding L2 minor capsid proteins to the vaccine.¹⁰⁴

Both vaccines induce high antibody titers and seem to be well-tolerated and safe and provide >90% protection in HPV-16 and HPV-18 in naïve females when given in three doses within six months.^{102, 105} Currently, only young HPV-naïve females are vaccinated because vaccination of women actively expressing HPV-16 or HPV-18 at study entry did not result in decreased development of cervical intraepithelial neoplasia (CIN) lesions.^{102, 105} However, vaccination with Gardasil® also provided >90% protection in women with evidence of past infection (seropositive and HPV DNA negative) with one or more of the HPV-types against which the vaccine is directed.^{106, 107} Long-term benefits of vaccination are not yet known, but it is hypothesised that vaccination could also strongly reduce the number of HPV-related OPSCC. This would indicate that HPV-naïve boys and young men should be vaccinated as well, because HPV-related OPSCC is diagnosed in males more often than in females.^{61, 69} However, seeing that patients usually present with HPV-related head and neck tumours from the fifth decennium of life onwards, the efficiency of vaccination in these patients will only become evident within a few decades.

Patients with HPV-related disease may benefit from the development of therapeutic vaccines. These vaccines are designed to induce cell-mediated immunity against the overexpressed foreign viral oncoproteins, particularly E6 and E7. There are four classes of therapeutic vaccines: 1) live-vector based; 2) peptide/protein based; 3) nucleic acid based; and 4) whole cell vaccines (for a comprehensive review see ¹⁰⁸). In anogenital and uterine cervical lesions therapeutic vaccination has been shown to generate specific immunological and clinical responses, including complete regression of the lesion in 22% of patients with CIN III lesions as reported in a study using a fusion protein-based vaccine.¹⁰⁹⁻¹¹¹ A preclinical study using a DNA-based vaccine demonstrated that such an approach for therapeutic vaccination was efficacious in a mouse model of HPV-related HNSCC.¹¹² Clinical trials to evaluate the effectiveness of therapeutic vaccination in HPV-related HNSCC are ongoing.^{113, 114}

Interferon

Interferons are cytokines that are produced by many cell types in response to infection with bacteria, viruses and parasites.¹¹⁵ Two classes of IFNs can be distinguished: Class I consists of IFN- α and IFN- β , and Class II consists of IFN γ . Class I IFNs are secreted from infected cells and bind to the ubiquitously expressed heterodimeric interferon receptor. Binding of IFN α/β to the interferon receptor induces the transcription of several host cell proteins that inhibit viral replication in the infected epithelial cell, and leads to activation and the production of IFN γ in dendritic cells. IFN γ can also be produced by activated Th1 cells. Both classes of IFNs possess antiviral and antiproliferative properties. IFN γ can also activate macrophages and natural killer lymphocytes, and induce translocation of the major histocompatibility complex Classes I and II to the cell membrane.¹¹⁵ The interferon response, however, is suppressed upon HPV infection because several HPV proteins (E1, E6 and E7) interfere with the IFN signal transduction cascade by binding to, for example, Tyk2 kinase, IRF-1 and IRF-3, p48 and p56 leading to downregulation of

the levels of IFN-inducible genes, such as TNSFS10, IFIT1 and IFI54.^{78, 116, 117}

Despite this, a successful immune response to HPV is generally seen in healthy individuals, as for example reported in the studies of van der Burg and co-workers,^{118, 119} showing high frequencies of circulating CD4+ T-helper cells reacting with HPV16 E2 and E6, indicating a cell-mediated Th1 immune response. In persisting lesions, application of IFN therapy may restore antiviral defence mechanisms, thereby supporting effective treatment of HPV-infected lesions. IFN therapy proved to be beneficial in HPV infections such as condylomata acuminata,^{119, 120} whereas the use of IFN therapy in HPV-associated anogenital intraepithelial neoplasia has been assessed in several studies with contradicting results. Improved outcome for IFN-treated patients was shown in some studies,^{121, 122} whereas others reported no change in response rates between treated patients and controls.^{123, 124} This might be attributed to the fact that local application seems to achieve better responses than systemic application.¹²⁰ In addition, it seems that IFN therapy can eradicate episomal HPV infection but leads to growth advantage for cells containing integrated HPV.^{125, 126} IFN-induced upregulation of p56, which blocks HPV replication by binding to the E1 protein and inhibits its helicase activity, may explain this effect on episomal infection.^{127, 128} Loss of (parts of) E1 and E2 by viral integration, resulting in upregulation of E6 and E7 as stated previously could explain the lack of effect of IFN treatment in these cells and their selective growth advantage. On the contrary, IFN was also shown to increase viral early gene transcription in a cell model.¹²⁹ In recurrent respiratory papillomatosis (RRP) a long-term response to IFN- α therapy was seen in patients with HPV-6-related papillomas, but patients with HPV-11-related papillomas were much less responsive to IFN therapy.¹³⁰ In conclusion, the beneficial effects of IFN therapy seem to be limited to episomal infections, which limits the applicability of this therapy in HPV-positive carcinomas.

8.6.2. Antiviral therapy

Cidofovir

Cidofovir [(S)-1-(3-hydroxy-2-phosphonylmethoxypropyl)cytosine] (HPMPC) is a nucleoside analogue of deoxycytidine monophosphate with a remarkably broad spectrum of antiviral activities directed against DNA viruses, including HPV and polyoma.¹³¹ After intracellular double phosphorylation, the structure resembles dCTP and can act as a competitive substrate. After removal of the diphosphate group cidofovir can be incorporated into viral DNA during replication, resulting in selective antiviral activity for those viruses encoding their own DNA polymerase. Viral DNA polymerases, for instance cytomegalovirus, display greater affinity for cidofovir than human cellular DNA polymerases. Although HPVs do not produce viral DNA polymerases, cells infected with HPV show enhanced susceptibility to cidofovir-induced apoptosis as compared to non-infected cells for a yet unknown reason.^{132, 133}

In 1998, it was shown by Andrei et al. that acyclic nucleoside phosphonate (ANP) analogues, such as cidofovir, show a selective antiproliferative effect in HPV-bearing

tumour cell lines CK-1, SiHa, CaSki and HeLa.¹³⁴ This effect is partly induced by its non-selective toxicity to rapidly dividing cells.^{134, 135} Apoptosis might also be induced by accumulation of the tumour suppressor proteins p53 and p21Cip1/WAF1,^{132, 133} although an increase in p53 expression was not found in the HNSCC cell line UPCI:SCC090.¹³⁶ However, by combining cidofovir with radiotherapy, the radiosensitivity of UPCI:SCC090 and other HPV-containing cell lines could be enhanced in vitro,^{132, 136} as well as in vivo in nude mouse xenografts.¹³² CT combined with cidofovir also yielded a synergistic effect in an HNSCC cell line model.¹³⁷ One study expressed concern about using cidofovir for the treatment of RRP,¹³⁸ as it demonstrated high malignancy transformation rates in rats and cell lines. In humans, this effect has not been reported, and cidofovir is already applied as an effective adjuvant therapy for HPV-induced RRP in humans.¹³⁹

For the treatment of various HPV-related lesions the route of administration may be an important factor. Cidofovir can be applied systemically or topically or injected intralesionally. Although concern was raised about possible nephrotoxicity in systemic use, this side-effect can be greatly diminished by administration of probenecid and prehydration with saline solution.¹⁴⁰

In a clinical setting, it was shown that local therapy with cidofovir gel resulted in complete or partial regression of uterine CIN II and III lesions^{141, 142} as well as vulvar and other intraepithelial neoplasms.^{143, 144} On the other hand, intralesional treatment with cidofovir of one patient with an invasive carcinoma in the respiratory tract and a history of RRP only lead to minor clinical effects, limited to the superficial portion of the tumour.¹⁴⁵

Clinical trials using cidofovir as an adjuvant therapy in cervical cancer have started,¹⁴⁶ but trials for its application in HNSCC have to be initiated.

Interfering RNAs

The RNA interference (RNAi) can be used to inactivate gene expression and so far encouraging results have been reported for the treatment of HPV-related carcinomas in vitro as well as in vivo. Chen and co-workers, for example, reported a 50% reduction of E7 mRNA expression in HPV-6b/11 E7-expressing mouse tumour models.¹⁴⁷ RNAi against HPV-16 E6 and/or E7 has been shown to degrade these mRNAs leading to decreased expression of the gene products in both cervical as well as HNSCC cell line models. This resulted in restoration of pRb function and upregulation of p53 and p21Cip1/WAF1, leading to substantial apoptotic cell death.¹⁴⁸⁻¹⁵⁰ RNAi against HPV-18 E6 and E7 has also seemed to possess antitumour activity by retarding the growth of HeLa-cell induced tumours in NOD-SCID mice¹⁵¹ and to enhance the chemotherapeutic effect of cisplatin in HeLa cells in vitro.¹⁵²

8.6.3. Molecular therapy based on cellular targets

Because inactivated tumour suppressor gene products such as p53 and p16INK4A are difficult to restore by molecular therapy, many studies have focussed on the identification of oncogenes and deregulated cell signalling pathways in HNSCC. Key

pathways involved in HNSCC include EGFR, PI3K-PTEN-AKT, TGF β and NF- κ B signalling for which inhibitors are available, for example, the anti-EGFR antibody cetuximab¹⁵³, or being tested in several clinical trials (for reviews, see^{72, 73}).

In cervical cancer EGFR overexpression has been shown to negatively affect overall survival in patients treated with radiotherapy.¹⁵⁴ Anti-EGFR therapy using cetuximab lead to a therapeutic response in 12.5% of patients with uterine cervical SCC.¹⁵⁵ Also in HNSCC, including HPV-positive OPSCC, overexpression of EGFR correlates with poor prognosis,^{67, 70, 71} although only a small subgroup of HPV-positive OPSCC exhibit EGFR protein accumulation.^{67, 71} Large prospective trials with anti-EGFR therapy in HPV-positive HNSCC have been initiated,⁹⁷ although its efficacy is most probably limited to the small subgroup of EGFR-expressing tumours.

Alternatively, the PI3K-PTEN-AKT pathway might be an efficient target because HPV-positive OPSCC show extra copies of chromosome 3q in up to two-thirds of cases,⁶⁸ including the 3q26 locus, harbouring the PI3K gene.

Tumour angiogenesis and metastases are correlated to upregulation of the TGF- β and NF κ B pathways.^{156, 157} HPV positive OPSCC have been shown to metastasise in an earlier stage compared to HPV negative OPSCC,¹⁵⁸ indicating earlier endothelial-to-mesenchymal transition (EMT), which is characterised by the expression of vimentin, downregulation of E-cadherin and upregulation of β -catenin.¹⁵⁹ This suggests that EMT might be related to upregulation of these pathways and that particularly HPV-positive OPSCC might be a potentially interesting group for NF κ B-inhibitors, for which a clinical trial has recently started.¹⁶⁰

Increased degradation of cell cycle regulatory proteins p53 and pRb by the oncoproteins E6 and E7, can be inhibited by targeting the proteasomal pathway. Ritonavir, a protease inhibitor (PI) that is used in HIV-infected patients, inhibits the chemotryptic activity of the human cellular 20S proteasome while increasing the tryptic activity,¹⁶¹ resulting in reduced protein degradation. It was shown to enhance antitumour activity when combined with radiotherapy both in vitro and in vivo in a Hep-2 head and neck carcinoma model,¹⁶¹ later however shown to be contaminated with HeLa cells. The PI Lopinavir was shown to restore p53 expression and to induce apoptosis in SiHa cells.¹⁶² Although several clinical trials have evaluated the effectiveness of PIs in the treatment of HIV, clinical trials in the treatment of HPV-related disease have not been initiated.

Finally, replication of the HPV virus can be targeted. In episomal HPV infection, replication is initiated by binding of E2 to its origin of replication.⁵⁴ In human transcription factors, the most commonly found DNA binding motifs are zinc fingers. Recently, artificial zinc fingers (AZF) have been developed as a potent new inhibitor of HPV.¹⁶³ When linked to a cell-penetrating peptide (CPP) these AZF were shown to inhibit HPV-18 for 97%.¹⁶⁴ However, because the CPP-AZF is designed to prevent E2 from binding to its origin of replication, they are only effective in episomal HV infection.

8.7. DISCUSSION

The past decade has provided evidence for a biological association between oncogenic HPV and OPSCC. HPV-induced OPSCC show molecular and clinical features that are clearly different from tobacco-and-alcohol-induced tumours and these differences seem to underlie prognostic differences between both tumour subgroups. Independent of treatment modality, patients with HPV-positive tumours demonstrate up to 30% better survival rates. In the past decennia, intensification of treatment was the most important strategy to improve survival of patients with HNSCC. However intensification of treatment, combined with increased side effects, is finally reaching the maximum tolerance of the patient and this limits the intensity of treatment. Until now, no differentiation of therapeutic strategies has been made between the HPV-positive and HPV-negative subgroups in international guidelines on OPSCC treatment.⁸¹ Because of the clinical and molecular differences between both groups, the question arises whether HPV-positive tumours need equally intensive treatment protocols as their HPV-negative counterparts. Moreover, additional antiviral therapeutic strategies can possibly improve survival without increasing therapy-related morbidity in HPV-positive tumours. In current and future studies, we should, therefore, aim at improving the quality of life in patients by de-intensification regimens in selected cases. Next-generation treatment strategies for HPV-associated cancers should focus on decreasing adjuvant radiotherapy and chemotherapy, whether or not combined with therapeutic options specifically targeting HPV. Assessing which therapy is most effective will finally lead to a more personalised approach for individual patients.

Immunomodulating therapies, such as IFN therapy, may have beneficial effects, but this seems to be limited to episomal infections. This conveys the need to reliably establish the integration status of HPV infection. That this criterion has not yet been met becomes apparent when observing the reported integration frequencies, which range from 0% to 100%, depending on the population studied and methods used.^{41, 42}

Tumour-specific host responses could also be enhanced by depletion of CD4+/CD25+ regulatory T-cells (Tregs). Increased expression of Tregs was shown in patients with CIN and cervical cancer.^{165, 166} It is hypothesised that the enlarged population of Tregs suppresses HPV-specific immunity and inhibits tumour-specific T-cell responses. Upregulated Tregs have already been depleted using an anti-CD25 antibody, such as PC61.¹⁶⁷

Other immunomodulating therapies such as imiquimod, a topical immune response modifier that has successfully been used in the treatment of anogenital lesions with episomal HPV infections,¹⁶⁸ are thought to be unsuitable for application in HPV-related HNSCC and RRP. Application to cutaneous epithelia is known to induce local inflammatory responses and pain, which will be enhanced in mucosal epithelia. Moreover, the substance cannot be controlled to reach all tumour parts when topically applied, and, like IFN-therapy, will at best lead to eradication of only episomal HPV infections, whereas a large proportion of HPV-positive HNSCC show viral integration.¹⁶⁹

Antiviral therapies such as cidofovir and RNAi have already shown promising results and are expected to have progressive impact on the treatment of HPV-associated lesions. Cidofovir has been tested in cervical cancers and RRP, where it has been applied topically or intralesionally in most studies. It has been shown that combining IFN therapy with cidofovir could enhance the antiviral and antiproliferative effects of either substance alone, and it is postulated that adding IFN therapy could further improve the auspicious effects of cidofovir combined with CT and/or radiotherapy.¹⁷⁰ Furthermore, it is recommended to assess the effects of cidofovir as adjuvant therapy in the treatment of HPV-associated HNSCC in a larger, prospective clinical trial.

The RNAi treatment, although tested in mouse models, has not yet been evaluated for use in human HPV-associated HNSCC. Such studies can, however, be expected in the near future, judging from patents referring to the use of oligonucleotides in the treatment of HPV infections (see for example ¹⁷¹).

Therapeutic approaches based on the molecular profile of the tumours are emerging in an adjuvant setting. However, one of the major drawbacks of such an approach is that the applicability should be assessed for each individual patient. For example, cetuximab can only be applied in a small subgroup of patients with HNSCC because HPV-positive tumours tend to show a low EGFR expression. In the current practice, Cetuximab is already used for larger non-resectable head and neck tumours, irrespective of HPV status. With regard to PIs and AZFs, no clinical studies have yet tested the applicability of these therapeutic options in the treatment of HPV-related carcinomas.

Because therapeutic vaccination is expected to have minimal side effects it can be combined with other therapeutic approaches, such as radiotherapy and/or CT, to obtain synergistic effects. However, therapeutic vaccines are still in a developmental stage.

Although a significant reduction in the burden of HPV-related diseases can be anticipated if prophylactic vaccination will live up to its promises, only HPV-naïve females are currently vaccinated. We firmly believe that young HPV-naïve boys should also be vaccinated in order to achieve optimal protection, although it needs to be validated whether vaccination is cost-effective.

In conclusion, we can state that although it has become evident that HPV-positive HNSCC have a better prognosis than their HPV-negative counterparts, the choice of therapy for these two subgroups of HNSCC will strongly depend on the outcome of ongoing clinical trials, including de-intensification protocols and implementation of treatment options based on new insights into the molecular biology of HPV-infection.

Conflict of interest

The authors have no competing interest.

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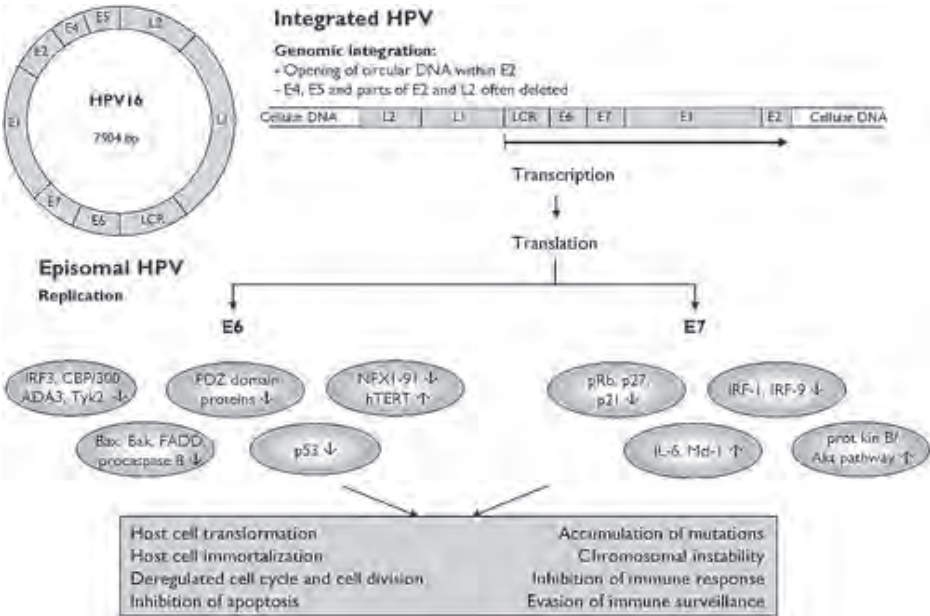
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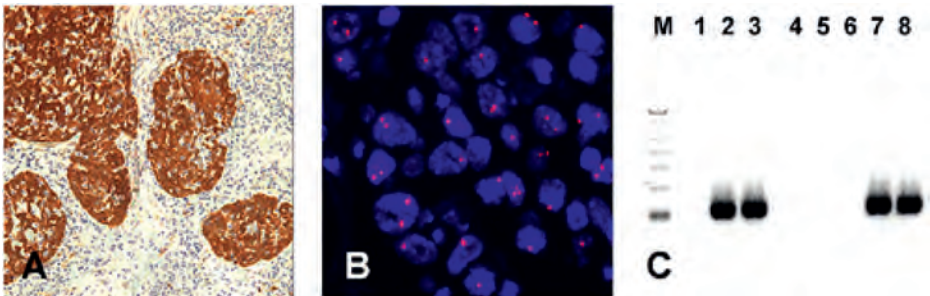
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Figure 1



Structure of circular episomal and linear integrated HPV DNA. The HPV genome is usually present in many episomal copies in the nucleus of infected cells. In the transition to cancer, viral DNA often integrates in 1 or more copies into the host genomic DNA. During this process, the ring structure of the HPV-DNA molecule is most often opened within the E2 reading frame, frequently leading to deletion of E4 and E5 and part of E2 and L2. The subsequent upregulation of E6 and E7 oncoproteins results in deregulation of cell signaling pathways, which, amongst others, leads to increased cellular proliferation and inhibition of apoptosis.¹⁷²⁻¹⁷⁵

Figure 2



Representative examples of strong nuclear and cytoplasmic p16INK4A immunostaining (A) and punctate nuclear HPV-16 FISH signals indicating viral integration (B) shown on paraffin embedded, formalin fixed tissue sections of oropharyngeal squamous cell carcinoma. An example of E6-specific HPV-16 RT-PCR products on a 1% agarose gel, on RNA extracted from cell lines and fresh-frozen oropharyngeal squamous cell carcinoma tumour tissues, are shown in (C)

Table 1. Involvement of human papillomavirus types of the alpha-genus in benign and malignant human lesions. The major human papillomavirus types for the different lesions are indicated in bold

Lesion	HPV types found	References
Head and neck benign		
Focal Epithelial Hyperplasia	13, 32	176, 177
Sinonasal papilloma	6, 11, 18	178, 179
Laryngeal papilloma and dysplasia	6 1), 11 , 16, 18	180, 181
Oral leukoplakia and lichen planus	6, 16, 18, 31, 33	182, 183
Head and neck malignant		
Oropharyngeal squamous cell carcinoma	6, 11, 16 , 18, 31, 33, 35	52, this review
Oral squamous cell carcinoma	16 , 18	52, 60
Laryngeal squamous cell carcinoma	6, 11, 16 , 30	52, 60
Sinonasal carcinoma	16 , 18	179
Anogenital		
Anogenital ²⁾ benign lesions ³⁾	6, 11, 16 , 18, 31, 33, 53, 56, 58, 66, 83	36, 184
Anogenital§ (squamous cell) carcinoma	6, 11, 16 , 18, 31, 33, 45	36
Cervical intraepithelial neoplasia and uterine cervical squamous cell carcinoma	6, 11, 16, 18 , 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, 66, 68, 70, 73, 82	16, 37
Adenocarcinoma in situ and uterine cervical adenocarcinoma	16 , 59 , 58 , 51 , 45 , 35 , 33 , 18	185, 186
Cutaneous		
Common skin warts	2, 3, 7, 10, 27, 28	15, 187
Periungual squamous cell carcinoma	16, 26, 33, 51, 56, 73	188, 189

1) The major HPV types for the different lesions are indicated in bold;

2) including anal, vulvar, vaginal and penile lesions;

3) including warts, lichen sclerosis, squamous cell carcinoma in situ , adenocarcinoma in situ and intraepithelial neoplasia.

Table 2. Clinical and molecular differences between OPSCC with or without HR-HPV *

	HPV-positive	HPV-negative
Clinical characteristics		
Preferred location	oropharynx	all sites
Degree of differentiation	poorly differentiated	moderately to well differentiated
Baseloid appearance	more often	less often
T-stage at diagnosis	T1-2	T3-4
Disease stage (TNM)	more advanced	less advanced
Average age	slightly younger than 60 years	slightly older than 60 years
Tobacco (ab)use	low	high
Alcohol (ab)use	low	high
5-year disease free survival	70-90%	30-60%
Second primary tumours within 5 years	0-10%	10-15%
Local recurrences within 5 years	10-20%	25-55%
Molecular characteristics		
E6/E7 expression	+	-
p53 downregulation	+	-
pRb downregulation	+	-
p16INK4A overexpression	+	- **
p14ARF overexpression	+	- **
p18INK4C overexpression	+	-
p21Cip1/WAF1 overexpression	+	-
Cyclin D1 overexpression	-	+ ***
EGFR overexpression	-	+ ***

* Summarised from 10, 22, 41, 56, 59, 62, 65-68, 92, 94, 158, 174, 190-192.

** Inactivated by gene mutation, hypermethylation or homozygous deletion.

*** Mainly induced by gene amplification or transcriptional upregulation.





General discussion and valorization

9.1. IMPLICATIONS OF HPV-RELATED TUMOR BIOLOGY ON TUMOR STAGING AND PROGNOSIS

In the past decade it has become clear that a biological association is present between oncogenic HPV and a subgroup of OPSCCs. HPV-induced OPSCCs show distinct molecular and clinical features that are different from tobacco-and- alcohol-induced tumors and these differences seem to underlie prognostic differences between both tumor subgroups.^{1,2} In chapter 2 we showed that HPV-positive tumors, which are associated with less smoking and alcohol consumption, have a different tumor biology. They have smaller primary tumor sizes, although regional lymph node involvement is comparable to HPV-negative tumors. In our study, this resulted in a different choice of treatment, but independent of treatment modality, HPV-positive tumors had a better prognosis.³ The prognostic value of nodal involvement is reduced by the presence of HPV. It was not until 2007 that the World Health Organization officially recognized HPV as causative agent in the development of head and neck cancer.⁴ Our study described in Chapter 2 was the first to address that lymph node involvement and extent of nodal disease do not have a reliable predictive value when the seventh edition UICC staging system for OPSCCs is used. We therefore advised an HPV-dependent staging system for the diagnostic work-up in oropharyngeal tumor staging. Similar results to our findings were found by the groups of Klozar et al. and Fritsch et al.⁵⁻⁶ When reviewing literature reporting on the prognostic value of N-status in tonsillar squamous cell carcinomas (TSCCs), 10 studies were published prior to 1990, which all reported that N-status was of prognostic importance (although only 2 studies provided results based on statistical analysis). In contrast, from 1990 onwards, only 4 out of 12 studies showed N-status to be of prognostic relevance.⁷⁻²⁸ This indicated that the prognostic shift of nodal status in TSCCs in literature runs parallel with the increase in HPV-prevalence in HNSCCs. In 2013, Dahlstrom et al. described in a US population of 3891 patients with OPSCCs collected between 1955 and 2004, a similar shift in prognostic reliability of TNM-status over time: compared to the previous study period, patients after 1995 were more often of younger age, had more tongue base or tonsillar tumors, were more often never smokers or former smokers and died only half as often.²⁹ The TNM-classification predicted the survival of patients treated prior to 1995 accurately, but lost its predictive value for patients treated between 1995 and 2004. They also noted an unusually favorable outcome for stage III and IV disease in the most recent decade. In 2015, data of the Surveillance, Epidemiology, and End Results (SEER) database for OPSCC – also without knowledge of HPV status - described similar changes in prognostic significance.³⁰ Similar to our results in chapter 2, the SEER-study noted a reduction of hazard ratio for survival for all N2 subcategories in the 7th edition UICC staging system compared with N0 disease.

In this period, it was recognized that in OPSCCs HPV-positivity devalued the prognostic value of N-status described in the 7th edition UICC staging system.²⁹ Therefore, a new staging system for HPV-positive OPSCCs was the next step. First, Ang et al. presented a risk-model to predict survival for stage III and IV OPSCCs treated with concomitant chemoradiotherapy, in which smoking status was combined with N stages “N0-2a” and “N2b-3”.³¹ In the HPV-positive group, smoking status (more or less than 10 pack years) discriminated between mild and moderate risk. Within the HPV-positive smoking group, difference in outcome was based on nodal status “N0-2a” and “N2b-3”, respectively, resulting in mild and moderate risk-associated groups. Spector et al. subsequently, subdivided N-status on the basis of diameter and number of nodes in 3 risk groups: a single node <6cm ipsilaterally or contralaterally as “HPV+ N1”, a single node ≥ 6cm or ≥ 2 nodes ipsilaterally/contralaterally or ≥3 nodes bilaterally as “HPV+ N2”. “HPV+ N3” was defined as neck node status with matted nodes.³² In 2015, Huang et al. studied a large group of 810 patients with OPSCCs and noted prognostic discrimination by using the 7th edition UICC staging system in HPV-positive tumors.³³ Compared to the HPV-negative tumors showing outcomes worsening from stage I to IV, their HPV-positive group showed no significantly different survival rates from stage I to IV. Two additional classifications were proposed using the existing T and N categories of the 7th edition UICC staging system for the HPV+ cohort. The first model, which was based on recursive partitioning (RPA), showed a better predictive value for prognosis (stage I: T1–3, N0–N2b; stage II: T1–3, N2c; stage III: T4 or N3, stage IV: M1 as stage IV). Interestingly, 56% of patients classified as stage III or IV according to 7th edition UICC staging system, changed to stage I when using the new model. The second model proposed by Huang et al. was based on adjusted hazard ratios (AHR): 4 prognostic groups in HPV-associated OPSCCs without hematogenous metastases were discriminated, based on N status (N0-2c vs. N3), T status (T1-3 vs. T4), smoking behavior (fewer vs. more than 20 pack-years history) and age (younger vs. older than 70 years).³³ Dahlstrom et al. (2016) were not able to validate Huang’s results; consequently, they proposed an own HPV-associated system, in which N status was staged corresponding to nasopharyngeal carcinomas.³⁴ Finally, the classification system suggested by O’Sullivan et al. (ICON-S; 2016) was used for the clinical TNM-classification in the 8th edition of the UICC staging system for HPV-positive carcinomas.³⁵ In this classification, N-status is based on the sidedness and maximum diameter of the nodes rather than on the number of nodes. It enhances the UICC staging system into more valid groups compared with the 7th edition to facilitate patient counseling, cancer surveillance, and translational research, and furthermore to optimize clinical trials design and outcome reporting.³⁶ For this multi-institutional analysis, also the data on HPV-positive OPSCCs of our study group were included. Besides the clinical staging system, a pathological staging system has been introduced to establish prognosis and guide adjuvant therapy decision in surgically-managed HPV-positive OPSCCs.³⁷ In contrast to clinical staging, the number of nodes (with a cut-off point of 4) determines N status (ranging from N0 to N2 without differentiation between N2a, N2b and N2c) for pathological staging.

This newly introduced 8th edition UICC staging system for HPV-positive OPSCCs was validated by others.³⁸⁻⁴² We also compared this 8th edition with the previous 7th edition UICC staging system for HPV-positive OPSCCs in this thesis. In our study only TSCCs were included, a very strict definition of HPV-positivity was used, and the added prognostic value of additional non-anatomical parameters was tested, as described in chapter 7 and further on discussed in this chapter in 7.6.

9.2. ADEQUATE DETECTION OF BIOLOGICALLY RELEVANT HPV-POSITIVITY IN OPSCCS

In chapter 2 we showed that the impact of HPV on tumor biology, prognosis, and choice of treatment is large. Therefore, detection of biologically relevant HPV-positivity is of increasing importance. Literature has addressed the issue of more consensus on the exact definition of HPV-associated OPSCCs.⁴⁴ HPV infection alone is not sufficient to classify an OPSCC as HPV-related, because the presence of HPV-DNA may only reflect a transient infection. In chapter 3, the reliability of p16INK4A immunohistochemistry (p16Ink4a-IHC) as surrogate marker for HPV was tested. We scored the p16Ink4a-IHC staining patterns according to the “block-type” immunopositivity approach, defined as p16Ink4a -IHC only being block positive if continuous (>70%) strong nuclear with or without cytoplasmic staining is present (in all head and neck lesions) and staining is observed in the basal cell layer with extension upwards (in benign and premalignant lesions).⁴⁵ Our results indicate that a strong nuclear and cytoplasmic p16Ink4a-IHC pattern can accurately predict the presence of HR-HPV16 in OPSCC and tonsillar dysplasias.⁴⁶ In low-risk-HPV6/11-positive benign and premalignant tonsillar and laryngeal lesions, however, the predictive value of p16Ink4a-IHC was lower and therefore caution is recommended when using this surrogate marker for HPV-infection. More recent studies have shown that in tumors outside the oropharynx, p16Ink4a-IHC has also less predictive value for HPV-presence.⁴⁶⁻⁴⁸

Despite the good correlation between P16Ink4a and the presence of HR-HPV16 in OPSCC, the decision to use p16Ink4a-IHC as sole diagnosticum to identify a HPV-positive OPSCC in the 8th edition UICC staging system may result in false-HPV-positive tumors. This was also recently emphasized by Bussu et al.⁴⁹ Moreover, differences in geographical patterns of HPV-prevalence implicate, that the positive predictive value of p16Ink4a-IHC as solitary diagnostic tool will drop if the a priori probability of having a HPV-positive OPSCC is lowered by 30 to 40%, as is the case in Europe compared to the US.⁵⁰ Throughout this thesis, therefore, p16Ink4a-IHC was combined with HPV-DNA PCR and/or Fluorescence In Situ Hybridization (FISH). We found that 14 out of 124 p16Ink4a-positive patients were not HPV-positive (11.2%). Taberna et al. recently investigated the treatment outcome of HPV-positive OPSCCs depending on the definition of HPV-positivity.⁵¹ They confirmed that definitions of HPV-positivity have

impact on TNM-classification and patients' prognosis and adequate testing with at least PCR for detecting HPV DNA next to p16Ink4a-IHC is emphasized.^{52, 53}

Nevertheless, p16Ink4a-IHC is a widely available, low-cost test in the general pathology laboratory in contrast to more expensive HPV DNA PCR and HPV-in situ hybridization analyses, which require a specialized laboratory and expertise. Therefore, it has been adopted as surrogate marker for HPV in the since 2018 used 8th edition UICC staging system for OPSCCs.⁵⁴

Prigge et al. suggested in their meta-analysis the most desirable technique to identify HPV-positive OPSCCs: the detection of HPV E6 and/or E7 oncogene transcripts of all HR-HPV types (a), in the form of all splice transcript variants (b), from fresh-frozen tumor tissue (c), and performed on isolated tumor cells (d), e.g. by means of tumor microdissection.⁵⁵ This "ideal" detection strategy is considerably laborious and, consequently, not realizable in the routine diagnostic setting.⁵⁵ Therefore, the combined p16Ink4a-IHC and HPV-DNA PCR assay significantly enhances specificity while maintaining high sensitivity. This diagnostic test combination thus represents an attractive testing strategy for the reliable diagnosis of HPV-positive OPSCCs in the clinical setting and may constitute an inclusion criterion for future therapeutic trials.⁴⁷

9.3. IS THERE ADDITIONAL VALUE FOR HPV-TESTING IN CUP-SYNDROME?

Because HPV-positive OPSCCs often spread to the lymph nodes already at low T-stages, a clear role for HPV-detection in the diagnostic work-up of cervical metastases of unknown primary tumors (CUP), to identify primary tumors in the oropharynx, has been advocated.⁵⁶ However, studies on the prevalence of HPV in lymph node metastases of which the primary tumor could not be detected after a comprehensive diagnostic workup, so-called "true" CUPs, are scarce and contradictory. In these studies, HPV prevalence rates range from 0% to 100% and were tested in very small sample numbers (range 1-25).⁵⁶⁻⁶² In chapter 4, we collected 29 true-CUP patients, of which the primary tumor was not present within 6 months of follow-up after treatment. Treatment consisted of a neck dissection and/or (chemo)radiotherapy. In this patient group 5/29 neck metastases were p16Ink4a-positive but in none of the specimen HPV DNA was detected by FISH and PCR (0%). No association between p16Ink4a-positivity and survival was found. All specimens were therefore regarded HPV-negative.⁶³ This indicates that the additional value of HPV-testing, next to a thoroughly performed diagnostic work-up, including panendoscopy, blind biopsies of tongue base and nasopharynx, bilateral tonsillectomy, and additional imaging, including PET-CT scanning, is limited.

9.4. SHOULD PATIENTS WITH CUP SYNDROME WITH OR WITHOUT HPV ALWAYS BE TREATED EXTENSIVELY, OR IS DE-ESCALATION AN OPTION?

More than a decade ago in the Netherlands, patients with CUP were treated extensively with postoperative radiotherapy of the bilateral neck and radiotherapy of the pharyngeal axis, in some cases combined with concurrent chemotherapy. The question arose whether de-escalation of therapeutic regimes in patients with CUP with or without HPV is possible. To enlarge our CUP patient group (see 7.3), we united the patient collectives of two European University medical centers (Cologne, Germany and Maastricht, the Netherlands; $n = 51$) and compared their data (chapter 5). Only true CUP patients (no primary tumor found within 6 month of follow-up) whose neck metastases were primarily treated surgically were included. The prevalence of HPV in this Maastricht-Cologne cohort of true CUP-patients was 7.8% (4/51 HPV-positive true CUPs).⁶⁴ No statistically significant difference of HPV-positivity between Maastricht and Cologne was found. Because of the low percentage of HPV-positivity, the influence of HPV positivity on outcome after therapy could not be assessed. Comparable to chapter 4, the added value of HPV testing in neck metastases of CUPs was found to be limited in chapter 5, especially when compared with OPSCCs in which the prevalence of HPV is much higher.⁵⁶

In the same study data about the therapeutic strategy for cervical metastasis of CUPs were retrospectively compared. All patients underwent a neck dissection, but the postsurgical management differed between patients and institutions. This enabled us to compare ipsilateral and bilateral radiotherapy, with or without radiation of the pharyngeal axis and with or without addition of chemotherapy. No significant differences in disease specific and overall survival between all subgroups was found, indicating that de-escalation might be a safe option in CUP-patients. Advanced nodal disease was, however, associated with a worse outcome, independent of treatment modality. The same was the case for the occurrence of regional recurrences after therapy: in almost half of the patients with regional recurrences during follow-up, distant metastases occurred ($n = 6/13$). Recently, also Sprave et al. found a high incidence of metachronous distant metastases in patients with regional recurrences.⁶⁵ In that study, the combined radiochemotherapy of the pharyngeal axis and bilateral cervical lymph nodes led to good results in case of specific risk factors (extra nodal spread and residual tumors),⁶⁵ which were not included in our study.

The presented data of the Maastricht-Cologne study contribute evidence to the ongoing discussion, in which the need for uniform international guidelines to treat patients with cervical CUP syndromes is claimed.⁶⁶ Our data indicate that the omission of radiotherapy of the pharyngeal axis and the contralateral neck after neck dissection,

as well as the omission of concomitant chemotherapy, might be safe under certain conditions. However, the variety of therapeutic strategies in both institutes in this complex population of patients with cervical CUP syndrome, prohibited any statistical evaluation or definitive conclusion in our study. Therefore, further research in a more homogeneous patient population was needed, addressed in chapter 6.

9.5. FURTHER EVIDENCE FOR SAFE DE- ESCALATION OF THERAPY FOR CUP PATIENT: NEED FOR TRIALS?

In order to homogenize treatment modalities of the neck and to exclude regional influences on the prevalence of HPV-positive CUPs, we merged our Maastricht database with a patient population collected at the Radboud University Medical center, Nijmegen, the Netherlands (n=80) (chapter 6). The protocols for diagnostics and treatment of CUP patients were similar in Nijmegen and Maastricht, and de-escalation of therapy in CUP patients was performed simultaneously in both institutes, starting in 2002. Again, in only 4 out of 72 neck dissection specimen presence of HPV DNA in tumor cells was found (6%) confirming previous results.⁶⁷ Therefore, also in this merged patient cohort HPV testing did not add value to the described diagnostic workup and did not influence therapeutic decision-making, for example, additional postsurgical radiotherapy of the oropharynx. This might be unexpected taking into consideration that small (T1-2 stage) HPV-positive OPSCC frequently are spreading to the lymph nodes and thus in principle could account for a substantial number of CUP. Probably, most HPV-positive primary tumors of the oropharynx are detected by tonsillectomy or “blind biopsies” of the oropharyngeal region, resulting in low percentages of HPV-positive true CUPs. Altogether, our studies in chapters 4-6 revealed that HPV-status was of insignificant importance to be used as in stratifying patients with CUP. Because of the low prevalence of HPV-positive CUPs, HPV-diagnostics improved neither the diagnostic and therapeutic work-up nor the outcome. As a result, the current treatment of patients with HPV-related CUP does not differ from the standard of care treatment.⁶⁶ Nevertheless, de-intensification of therapy remains an interesting option to be examined in the (small) group of HPV-positive CUP-patients.⁵⁶

Furthermore, in chapter 6, we confirmed our previous observation that omitting irradiation of the pharyngeal axis in patients with cervical true CUP syndrome does not result in the emergence of a primary tumor in the pharyngeal axis during five years of follow-up. This can avoid acute and late toxicity of comprehensive radiotherapy of the pharyngeal mucosa with significant improvement of long-term quality of life for these patients. Also, the absence of post-surgical radiotherapy of the contralateral neck in CUP did not lead to a decrease of regional control rates nor of survival rates.

In order to compare the outcome of ipsilateral radiotherapy solely with a comprehensive radiotherapeutic regime in CUP-patients, Nieder et al. recommended a randomized controlled trial already in 2001.⁶⁸ Only one randomized controlled trial was started since 2002, but was never completed (EORTC-24001-22005) as a consequence of limited patient enrollment.⁶⁹ Still, a prospective multicenter approach to analyze the true impact of radiotherapy target volume in CUP-patients is needed. However, the low prevalence of patients with CUP and the heterogeneous treatment strategies in different countries and regions are important limiting factors to design a large international study providing sufficient evidence for an international guideline.

For that reason, the American Society of Clinical Oncology convened an Expert Panel of medical oncology, surgery, radiation oncology, radiology, pathology, and advocacy experts to conduct a literature search including 100 relevant studies published from 2008 through 2019 including our presented results in this thesis.⁷⁰ Available evidence and informal consensus was used to develop evidence-based guideline recommendations about appropriate pre-operative evaluation, appropriate surgical diagnostic and therapeutic procedures, treatment considerations for surgical management, radiotherapy and chemotherapy in patients with CUP. Regarding radiotherapy of the occult primary tumor site, it was stated that solely radiotherapy of the ipsilateral oropharynx (i.e. ipsilateral tonsillar bed, ipsilateral soft palate and the mucosa of the entire tongue base) is recommended in patients with CUP treated with primary radiotherapy for one or more unilaterally located lymph nodes, not greater than 6 cm (AJCC 8th N1 HPV+ve and AJCC 8th N1-N2b HPV-ve), and in case of available PET-CT scan and performed contralateral tonsillectomy. Patients with CUP presenting with bilateral located lymph nodes not greater than 6cm (AJCC 8th N2 HPV+ve and AJCC 8th N2c HPV-ve), require bilateral treatment of the oropharyngeal mucosa. Patients with CUP presenting with nodes in the lower cervical stations (III and IV) should be considered for treatment of the larynx and hypopharynx, given the marginally higher risk of spread to stations III and IV from these organs.⁷⁰

The following argumentation was offered regarding radiotherapy of the unilateral versus bilateral neck: as bilateral neck irradiation for CUP has been considered standard of care historically, this approach is accompanied with considerable toxicity, including increasing dose to salivary glands, larynx, pharyngeal constrictors, mandible, hypopharynx, and esophagus. Following high-resolution imaging, ipsilateral only radiotherapy has been demonstrated to results in very acceptable rates of contralateral failure and reduced doses to the above-named structures. Ipsilateral neck irradiation is recommended in patients with CUP with a unilateral single node without extranodal extension and preferably in lymph node level II. In all other patients with CUP (multiple nodes, nodes, node(s) greater than 6 cm, level III or IV nodes, and/or clinical or radiologic ENE), bilateral neck treatment is recommended as in these patients higher rates of contralateral involvement are noted and prognosis is worse.⁷⁰

These above cited recommendations are in line with our presented results. However, the outcomes of interest in the above cited ASCO guidelines were survival, local and

regional disease control, and quality of life.⁷⁰ In our presented results in chapters 5 and 6, the occurrence of distant metastases was the most important limiting factor of survival. Again, this affected the more advanced staged necks, i.e. more than one lymph node or lymph node size more than 6cm (AJCC 7th N2b or higher). This underscores the importance of including distant disease control as primary endpoint in the evaluation of therapeutic strategies in CUP patients, as recently also was emphasized by Sprave et al.⁶⁵

9.6. THE PROGNOSTIC VALUE OF LYMPH NODE METASTASES IN HPV-POSITIVE TSCCS AND HOW IT INFLUENCED THE 7TH AND THE 8TH EDITION OF THE UICC CLASSIFICATION SYSTEM.

Chapter 7 describes that the predictive value of the 7th edition UICC tumor staging system for OPSCCs, and N-status in particular, has shifted over time as a consequence of the epidemic of HPV-associated HNSCCs. Since our study in 2009 (chapter 2), which was the first to report the direct correlation of HPV-status and the diminished prognostic value of N-status, different larger studies followed confirming our data.⁴³ This finally resulted in the new 8th edition UICC tumor staging system, in which a separate classification system for HPV-positive tumors has been introduced.⁵⁴ In chapter 7 we first summarized the changes of the 8th compared to the 7th edition of the UICC staging system: nodal stages underwent a transformation, clinical neck stages N1 until N2b were re-classified as cN1 including all ipsilateral neck metastases no greater than 6 centimeters, bilateral neck nodes were classified as cN2, and nodes greater than 6 cm as cN3. Along with the clinical staging, a pathological staging system was introduced based on the number of positive nodes identified by histopathological examination after neck surgery. These changes in tumor staging were then evaluated by testing the prognostic value of the different 7th- as well as 8th UICC tumor stages in HPV-positive OSCCs. The examination also included the separate T-, N-, and M-stages. An unselected population of 368 patients with SCCs of the tonsil, which is the site associated with the highest prevalence of HPV, was included in the study.⁷¹ Next to T-, N-, and M-stages the influence of patient-associated clinical variables including tumor differentiation grade, age, smoking behavior, alcohol consumption and treatment were taken into account. In total, 110 tumors were tested HPV-positive with p16Ink4a-IHC, HPV16-DNA PCR and/or FISH. Advanced stage HPV-positive tumors staged with the 7th edition UICC tumor staging system had a favorable prognosis. These tumors, however, had despite more advanced N-status (resulting in a higher overall stage) smaller primary tumors. When applying the 8th edition staging system we noted that 54% of all tumors were

classified as stage I compared to 5% of patients in the 7th edition. At the same time in the 7th edition 56% of the TSCCs were classified as stage IVa compared to only 3% of patients that were classified as stage IV in the 8th edition. We found that the 8th edition UICC tumor staging system, therefore, did better separate the different staged tumors in survival analysis. Our results were in line with other studies confirming that the introduction of the 8th HPV-associated tumor staging system is a step forward in staging HPV-associated OPSCCs.⁴³ However, when testing T-, c/pN- and M-status separately we found only cN3- and M1-status to be “anatomical” variables that significantly influence prognosis negatively. As a consequence of a favorable prognosis in HPV-positive T4-tumors, T-status in general was not associated with survival in our study. There was also no difference in survival dependent on pN-status. However, pN-status could be classified in only 38 HPV-positive TSCCs, because in those patients a neck dissection was performed.

Cramer et al. recently validated the 8th edition UICC staging system in a population of more than 15,000 patients (USA) and demonstrated an improved prediction of prognosis for HPV-positive OPSCC patients compared to the previous 7th edition UICC staging system.⁷² Also for T-, cN- and pN-status, the prognostic value could be validated in the HPV-positive population. In contrast to the validation study by Cramer et al., our study focused exclusively on SCCs of the tonsil, which is the subsite with the highest percentage of HPV-positive tumors. HPV-positivity was tested using HPV16-DNA PCR and/or FISH in addition to p16-IHC, which was the only test used in the study by Cramer et al. On the other hand, our study contained a much smaller patient population, although 110 unselected HPC-positive TSCCs were included. Despite these differences, the overall outcome of both studies were not similar. In our study population, the 8th edition of the UICC tumor staging system was also associated with a better prognostic value for tumor stage. However, in our study no significant prognostic value of N-stages cN0 to cN2 and pN0 to pN2 was found. Only cN3-status and M1-status were significantly associated with unfavorable prognosis. The cut-off point for a favorable prognostic value in our study of TSCCs was N3, however, only very few bilaterally involved (and thus cN2) neck stages were diagnosed. The study of MacKenzie et al. confirmed this observation and reported that only lymph nodes larger than 6cm (cN3) were associated with a worse survival.⁷³

A clinically negative (cN0) neck status was not associated with a better prognosis compared to N1 in our HPV-positive TSCC patient population, and showed even a worse prognosis than the N1 neck. In the validation study by Cramer et al., also no significant differences in survival were found between cN stages cN0 versus cN1 and cN2.⁷² Moreover, although the validation of the prognostic value of the clinical N-status (8th edition) was successfully performed, cN1-status was associated with a significantly better survival than N0 and also the bilateral involved neck (cN2-status) was not associated with a significantly worse survival than the clinically negative neck after adjustment for age, sex and race. Only cN3-status was significantly associated with a worse survival.⁷²

In previous research, it was noticed that patients with HPV-positive carcinomas more often had a lymph node as presenting symptom when compared to their HPV-negative counterparts. In HPV-positive carcinomas such a finding thus may guide the subsequent discovery of the primary tumor, and furthermore it led to the hypothesis that oncogenic HPV infection may play a substantial role in CUP-syndrome.³ As a consequence, patients presenting with these “alarming” nodes are expected to have a better prognosis upon treatment, which we did find in our study. Fritsch et al. and Ang et al. also described that patients with an HPV-positive single neck node between 3 and 6cm (N2a, 7th edition) had a better outcome than patients without lymph node metastases.^{6, 74} Fritsch et al. compared outcome based on N-status between HPV-dominant (tonsillar fossa and base of tongue) and non-HPV-dominant oropharyngeal subsites in more than 15,000 OPSCCs.⁶ In the HPV-dominant population, cN2a (7th ed.) was associated with a better survival than cN0/1 and in the total population no differences in outcome were noted as long as lymph node metastases were unilateral (<cN2c). In our study only patients with TSCCs were included. In this strongly HPV dominant subgroup similar results were found and a clinically negative neck (cN0) in HPV-positive TSCCs was not associated with a better survival than necks with lymph nodes smaller than 6 centimeter in diameter (i.e. cN1- and cN2-status). Possibly, unknown factors next to HPV-driven carcinogenesis play a role in the outcome of the clinical negative neck in HPV-positive OPSCCs, taken also into account the fact that the presentation of OPSCCs without involved neck nodes is atypical for HPV-associated tumoral behavior.

A point of discussion in the literature is that the influence of N-status on prognosis in HPV-positive tumors is often analyzed for all oropharyngeal subsites, without even discriminating between HPV-dominant or non-dominant subsites. In chapters 2 and 7, patients with TSCCs, the most HPV-dominant oropharyngeal subsite, were selected. Sood et al. described that a bilaterally involved neck status predominantly is seen in base of tongue tumors, indicating that lymph node dissemination patterns even differ within the HPV-dominant oropharyngeal sites.⁷⁶ This may explain the low number of bilaterally involved necks in TSCCs and the lack of significance of the cN2-stage classifying for bilateral neck involvement in the 8th edition which we found in our study. The location of the tumor in the different subsites of the oropharynx therefore likely plays a prominent role in the development of advanced (N-) tumor stages in HPV-positive OPSCCs.

All in all, the cN status classification assessed for HPV-positive OPSCCs according to the 8th edition UICC tumor staging system still turns out to be a suboptimal predictor of survival. In our study only cN3 and M1 were associated with a worse survival. Therefore, other, “non-anatomical” variables have been investigated to improve risk assessment of HPV-positive tumors.

9.7. ADDITIONAL NON-ANATOMICAL FACTORS IMPROVING PROGNOSIS IN HPV-RELATED OROPHARYNGEAL TUMOR STAGING.

The goal of implementing the 8th UICC staging system for HPV-positive OPSCCs was to more accurately represent the superior survival outcomes seen in these tumors and thereby to improve the prognostic value of the system and possibilities to guide treatment decisions.⁷⁷

As mentioned earlier, our results described in chapter 7 were in line with other studies confirming that the introduction of the 8th HPV-associated tumor staging system is a step forward in staging HPV-associated OPSCCs. However, in our study the survival of HPV-positive TSCCs was not predominantly dependent on TNM-status even when using the 8th edition. The most significant prognostic factors in HPV-positive TSCCs were smoking, age, N3-status, and the presence of distant metastases. Our study indicated that the prognostic value of the 8th edition UICC staging system can be improved by including smoking history and age with a cut-off point of 65 years as additional prognostic factors. Therefore, we have proposed a risk model for HPV-positive TSCCs based on smoking history, age, nodal size of ≥ 6 cm and presence of distant metastases resulting in 4 groups. The first group consisting of non- or former smokers (patients who quit smoking more than 10 years prior to the diagnosis of TSCC independent of the number of previously smoked pack years) was associated with a 5-yr OS of 95.1% even in advanced tumor stages. In groups 2 and 3, all patients smoked daily. Group 2 included smokers aged 65 or younger with an associated overall survival rate of 75.6%. In group 3 patients older than 65 who smoked had a 5-yr OS of 46.2%. Group 4 consisted of patients with N3- and M1-status (5-yr OS: 0%). Interestingly, two patients survived with a N3-staged neck, they were both non-smokers. Within the different groups, T- and/or N-status did not further differentiate between survival rate.

The issue of re-staging HPV-associated OPSCCs using N status alone or in combination with other clinical parameters has been addressed from various perspectives in the literature. Results of our research group on a prognostic model for OPSCCs was previously presented by Rios et al.⁷⁸ These results were validated by Rietbergen et al. in a larger cohort, showing the large impact of performance status on outcome in the whole patient group.⁷⁹ However, within the HPV-positive subgroup no further differentiation in risk profiles was provided.

In the recursive partitioning analysis of the radiation therapy oncology group by Ang et al. (2010), non- anatomic parameters such as age and tobacco smoking were included for the first time and an important prognostic value was found for tobacco smoking.⁷⁵ A predictive model for the prognosis of stage III and IV OPSCCs treated with concomitant chemoradiotherapy was presented, in which smoking status was combined with N stages “N0-2a” and “N2b-3”. In the HPV-positive groups, smoking status discriminated between mild and moderate risk. These results were validated by others.^{80, 81} Huang

et al. (2015) discriminated 4 prognostic groups in HPV-associated OPSCCs without hematogenous metastases based on N status (N0-2c vs. N3), T status (T1-3 vs. T4), smoking behavior (fewer vs. more than 20 pack-years history) and age (younger vs. older than 70 years), with associated 5-year overall survival rates of respectively 89%, 64%, 57% and 40%.⁸² Regarding smoking, Marur et al. noticed that treatment failures in a de-escalating regime of combining cetuximab with radiotherapy were seen in smokers (>10 packyears).⁸³ However, Haigentz et al. emphasized that including smoking in a predictive model goes along with great limitations as a consequence of the lack of validated, prospective data and the subjectivity of the data collection on tobacco use.⁸⁴ Further study by Rietbergen et al. showed no differences in outcome regarding smokers versus non-smokers.⁷⁹ In that study, smoking status was defined based on the number of pack years and no separate classification was performed for former-smoker status, which might have influenced the results for the smoking group.⁷⁹ Moreover, Broughman et al. postulated to leave the 10 packyear rule, proposed by Ang et al. as stratifier in HPV-positive OPSCCs, because of the favorable prognosis of former-smoking status independent of the number of pack years in their recent study.^{75, 85} These findings correspond with our results which show favorable outcomes in former-smokers with HPV-positive OPSCCs as presented in chapter 7. This stresses the importance of adequate history taking regarding smoking status in the work-up in a patient population with HPV-positive OPSCCs.⁸⁶

In conclusion, in our study the outcome of HPV-positive TSCCs was not predominantly dependent on TNM status even when using the 8th edition. The most significant prognostic factors in HPV-positive TSCCs were smoking, age, and also N3-status and the presence of distant metastases. In our predictive model, a prognostic role for age with a cut-off point of 65 years was observed. Non- or former smokers had a very favorable prognosis of more than 95% 5-year OS even in more advanced tumor stages and in former smokers who quit smoking longer than 10 years ago the number of pack years had no influence on prognosis. We think that this model could provide a simple additional tool for predicting the prognosis of HPV-positive TSCCs in the clinical setting.

9.8. HPV-POSITIVE HNSCC BIOLOGY AND DIRECTIONS FOR THERAPY.

The previous chapters of this thesis pointed out that detection of biologically active HPV in HNSCCs has prognostic relevance and, therefore, a separate classification of HPV-induced tumors has been introduced.^{3, 54} Further optimization of treatment protocols for this distinct group of HNSCCs is the next step. Data on treatment response of OPSCCs indicate that large low-risk subgroups of patients with HPV-positive tumors show up to 30% better survival rates than patients with HPV-negative tumors, independent of the type of treatment, as a consequence of their different tumor biology (Chapters 2 and

7).^{3, 71, 75} Due to these advances in insights in the clinical and molecular behavior of HPV-positive OPSCCs, the National Comprehensive Cancer Network (NCCN) Guidelines have made a distinction between treatment pathways for P16Ink4a-positive and -negative OPSCCs. However, the incorporated treatment strategies for both groups in the current guidelines remain almost identical and are mainly based on surgery, radiotherapy and chemotherapy.⁸⁷

To improve the efficacy of treatment while preventing increased side effects in patients with HPV-positive OPSCCs, the question arose whether treatment de-intensification and/or new HPV-targeted therapeutic options are possible. This question was addressed in chapter 8, in a large review on (future) therapeutic options for HPV-positive tumors.⁸⁸ Literature describes two main strategies for specific treatment of HPV-positive tumors. The first strategy focuses on the unique clinical behavior of HPV-associated HNSCCs and selects the patients based on risk-profiles to modulate and possibly de-intensify treatment. Do HPV-positive tumors need treatment protocols as intensive as their HPV-negative counterparts? Does the high chemo- and radiosensitivity of HPV-positive HNSCCs offer possibilities for de-escalation of therapy leading to reduced therapy-induced toxicities? In chapter 8, future directions for de-intensified treatment of HPV-positive HNSCCs were discussed.⁸⁸ Until now, mainly improved techniques associated with less treatment-related morbidity like IMRT/IMPT or TORS have been implemented in clinical practice.⁸⁹⁻⁹² Since the publication of our review, several de-escalation therapy trials for HPV-positive OSCCs have been performed, which can be summarized in three categories. In the first category, there are four large phase II studies, in which induction chemotherapy followed by reduced-dose RT led to less toxicities in p16Ink4a-positive and/or HPV-positive patients treated with the reduction dose RT: different low-risk profiles ('less than 20 pack years': Quarterback-trial, '< T4, < N2c, and ≤ 10 pack-year smoking history': ECOG E1308-study) and 'T1-T3, N0-N2b, and <10 pack-years': Optima II -trial) were associated with favorable outcomes in the reduction dose RT groups. These data support phase 3 clinical trials of radiation dose reduction after induction chemotherapy to be tested as a treatment strategy in HPV-positive OPSCCs and to quantify survival gain compared to standard of care.⁹³⁻⁹⁶

A second category is dose reduction of concurrent chemoradiotherapy. Chera et al. reported results of two performed phase II trials, in which patients with T0-T3, N0-N2c, M0, p16Ink4a-positive disease and a minimal smoking history were treated with 60 GY (16% less than standard dose) of intensity-modulated radiotherapy with concurrent weekly intravenous cisplatin (30 mg/m²; 40% less than standard dose). A good preservation of quality of life and an excellent 3-year tumor control and survival was found.⁹⁷ These first promising results have been reported in the second half of the last decade, and a phase III trial have to be awaited for.

In a third category, outcome was evaluated when concurrent cetuximab-based chemotherapy instead of cisplatin-based chemotherapy was administered. Phase III-trials by Gillison et al. and Mehanna et al. both noticed a higher locoregional control in the cetuximab-arm, however, overall survival was superior in the cisplatin-arm.⁹⁸⁻⁹⁹ Jones

et al. recently updated the results of this large phase III trial and concluded that the standard regimen of concurrent chemoradiation therapy being cisplatin-radiotherapy should not be replaced in HPV-positive OPSCCs.¹⁰⁰

On the basis of the above-mentioned conflicting results, there are no phase III trials to provide sufficient evidence that systemic therapy or radiotherapy may be de-intensified in HPV-positive HNSCC. Dose reduction of RT after a clinically good response to induction chemotherapy seems promising, particularly in well-defined low-risk subgroups of HPV-positive OPSCCs.

A second strategy of altering the therapeutic approach for HPV-positive HNSCC is to target HPV-specific molecular characteristics and search for new therapeutic options, in other words to provide HPV-positive tumors with another treatment protocol than their HPV-negative counterparts? In chapter 8, we, therefore, focused next on present therapeutic HPV-targeting strategies.

Different prophylactic and therapeutic alternatives for the current treatments of HPV-positive OPSCCs were discussed: 1) immunomodulating therapies including prophylactic and therapeutic vaccines. 2) antiviral therapies including interfering RNA and cidofovir; and 3) molecular therapy based on cellular targets including, protease inhibitors ritonavir and lopinavir, artificial zinc fingers, NFκB and anti-EGFR therapy.⁸⁸

Currently, these above mentioned targeted therapies for HPV-positive HNSCC seem promising, but still have not proven their efficacy in HPV-positive OPSCCs, i.e. data on vaccine efficacy in OPSCCs is lacking,¹⁰¹ or data is currently too preliminary.^{102, 103}

Interfering RNA has also not been tested in vivo in human HNSCCs up to now,¹⁰⁴ and cidofovir was recently tested in human HPV-positive HNSCC cell lines where it induced S- and G2/M phase arrest, resulting in mitotic catastrophe but not in apoptosis.¹⁰⁵ Protease inhibitors ritonavir and lopinavir have proven their efficacy in the treatment of HPV and although their therapeutic effect seems promising in cervical high-grade squamous intraepithelial lesions, clinical trials in HPV-positive OPSCCs are needed.¹⁰⁶ Regarding EGFR-targeted therapy, finally, the inferiority of cetuximab to cisplatin in concurrent radiotherapeutic regimes in overall survival in the general HPV-positive populations was described above.^{98, 99} A second EGFR-targeted therapeutic agent, panitumumab, did also not show better outcome in HPV-positive OPSCCs in the SPECTRUM and PARTNER trials.^{107, 108} The lack of overexpression of EGFR in a large group of HPV-positive OPSCCs may contribute to the failure of improving outcome based on EGFR-targeted therapies. Notwithstanding, narrowcasting EGFR-targeted therapy to the right selection of HNSCC patients still seems a serious option to explore.

A recent perspective regarding alternative therapies for HPV-positive HNSCC since the publication of our review is immunotherapy. Kim et al. recently classified OPSCCs immunologically into immune-rich (IR), mesenchymal (MS) and xenobiotic (XB) subtypes based on RNA-sequencing data.¹⁰⁹ All IR type tumors were HPV-positive, most XB types were HPV negative, and MS types showed both HPV-positive and -negative tumors. The IR type was associated with a favorable response signature during anti-PD-1/PD-L1

therapy, which seems a promising target in this HPV-positive OPSCC subgroup.¹¹⁰ The programmed death-1/programmed death ligand-1 (PD-1/PD-L1) pathway (referred to as the PD pathway) induces immunosuppression in the tumor microenvironment: tumor cells and other cells in tumor microenvironment can express high levels of PD-L1, which results in suppressed immunity upon interaction with PD-1. PD-L1-expressing cells use multiple mechanisms to suppress tumor immunity, e.g. PD-L1 on tumor cells can act as a receptor, and the signal delivered from PD-1 on T cells can protect tumor cells from cytotoxic lysis.¹¹⁰ Normal human tissues seldom express PD-L1 protein on their cell surface, with the exception of tonsillar tissue a.o.¹¹⁰ Based on durable objective response rates and a favorable safety profile (according to the results of the CheckMate 141 and KEYNOTE-048 trials), PD-L1-checkpoint inhibitors Nivolumab and Pembrolizumab (i.e. antibodies targeting the PD-1 receptor of lymphocytes) have been approved by the US Food and drug administration (FDA) for the treatment in HNSCCs.^{111,112} So far, however, no phase 3 trials were conducted regarding outcome after immunotherapy in HPV-positive OPSCCs. The implementation of immune checkpoint inhibitors revealed a new research field in cancer therapeutics which is evolving quickly. Besides anti-PD-1 and anti-PD-L1 therapy, other therapeutics which interfere with immune checkpoint are currently subject of ongoing studies in HNSCCs. Table 1 gives an overview of the ongoing studies related to anti-PD-1, anti-PD-L1, anti-CTLA4, anti-NKG2A and anti-P13K therapeutics in HNSCCs.

Overall, the development of new antiviral and immunomodulatory treatments may be instrumental in the future therapy management of HPV-positive HNSCCs to improve survival rates and decrease disease-and-treatment-related morbidity. There is rapidly increasing evidence for molecular and immunologic subgroups within the HPV-positive OPSCCs and also within HNSCCs in general, including the HPV-positive tumors. These subgroups most likely will show a different tumoral behavior and response to therapy. For example, Zhang et al. described different subgroups of HPV-positive HNSCCs that can be identified molecularly, i.e. HPV-KRT (upregulation of keratinization and oxidation-reduction process) and HPV-IMU (upregulation of mesenchymal and immune-response genes).¹¹³ They emphasize that further research is needed for a better understanding of these HPV-positive subgroups. The HPV-KRT group for example proved to be associated with the occurrence of *PIK3CA* mutations in the tumor. Recently, Beatty et al. performed two phase II trials in which the clinical significance of *PIK3CA* mutations in 77 HPV-associated OPSCC patients was studied. In these studies de-intensified CRT (60 Gy intensity-modulated radiotherapy with concurrent weekly cisplatin) was given.¹¹⁴ *PIK3CA* mutation was the only variable which was significantly associated with a worse disease-free survival (multivariate analysis: HR 5.71). Furthermore, recent research by Locati et al. reclassified these two molecularly defined subgroups (HPV-KRT and HPV-IMU) into three clusters.¹¹⁵ Cluster 1 is an immune-related subgroup characterized by high IFN γ signaling, associated with a good prognosis and probably the subgroup with a good response to immunotherapy. The HPV-KRT subgroup was further reclassified into two subgroups: Cluster 2 is an epithelial-mesenchymal transition-related subgroup (EMT),

characterized by fibroblast infiltration, increase in hypoxia and EMT-upregulation, and Cluster 3 is a proliferation-related subgroup, characterized by upregulation of E2F (G1 checkpoint transcription factor) and G2M checkpoint genes. HPV-positive tumors in Cluster 3 are associated with an intermediate risk profile and tumors in Cluster 2 with a high rate of integration of HPV DNA into the host genome and an unfavorable prognosis. HPV-positive tumors in this latter Cluster 2 subgroup are potential candidates for treatment intensification according to Locati et al.¹¹⁵

Further research must help to gain insight and develop tools for identification of risk-profiles based on clinical (smoking, age, cN3-status a.o.) and molecular characteristics (PIK3CA-mutation a.o.), which will play a key role in the stratification of patients for therapeutic decision making.

9.9. CONCLUSION: CHOOSING THE RIGHT THERAPY FOR THE RIGHT PATIENTS

HPV-positive OPSCCs have a different tumor biology and clinical behavior compared with their HPV-negative counterparts. The reported favorable prognosis despite frequent spread of HPV-associated tumors to the cervical lymph nodes have strongly influenced the discussion of adequate tumor staging in HPV-positive OPSCCs. This led to the implementation of a separate staging system for HPV-positive OPSCCs in the 8th UICC tumor staging edition. The associated favorable prognosis make HPV-related tumors more eligible for de-escalation of therapy to reduce treatment-related toxicities. However, HPV-positive tumors also prove to be a heterogeneous group of tumors and additional parameters are needed in stratifying risk groups, e.g. non- and former smoking status and age. Moreover, the rapidly increasing evidence of molecular and immunologic subgroups within the HPV-positive OPSCCs and within HNSCCs in general indicate that also other factors are influencing tumoral behavior and response to therapy. Therefore, when developing new HPV-targeted therapeutic strategies, these different molecular and biological characteristics must be taken into account.

The expected role of HPV infection in cervical lymph nodes of unknown primary origin was refuted in this thesis. Nevertheless, de-escalation of therapy in CUP still remains a serious option, independent of HPV, because it proved to be safe in a comprehensive, treated patient population described in this thesis.

Future trials on next generation treatment strategies for HPV-associated cancers should focus on reducing adjuvant radiotherapy and chemotherapy, whether or not in combination with therapeutic options specifically targeting HPV, HPV-related molecular biomarkers, and HPV-related subgroups defined by immunological and biological characteristics. This will enable the selection of superior treatment strategies for high-risk tumors and possible de-escalation therapies for low-risk groups to reduce toxic side-effects and minimize compromised functional outcome.

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Table 1. Ongoing clinical trials involving immunomodulation in HNSCC

Drugs	Drug types	Trial number	name	phase	status	Description
cemiplimab + ISA101B	anti PD-1 agent + therapeutic vaccine	NCT03669718		II	recruiting	Blinded, placebo-controlled, randomized, phase 2 study
cemiplimab + ISA101B	anti PD-1 agent + therapeutic vaccine	NCT04398524		II	not yet recruiting	Phase II study in Recurrent/Metastatic HPV16 Positive OPSCC with disease progression after prior anti-PD-1 therapy
avelimab + TG4001	anti PD-L1 agent + therapeutic vaccine	NCT03260023		Ib/II	recruiting	Phase Ib/II Trial in HPV-16 positive Recurrent or Metastatic Malignancies (incl. OPSCC)
durvalumab + MEDI0457	anti PD-L1 agent + therapeutic vaccine	NCT04001413		II	recruiting	Phase 2 trials in HPV-positive HNSCC.
pembrolizumab + PDS0101	anti PD-L1 agent + therapeutic vaccine	NCT04260126	Versatile-002	II	not yet recruiting	Phase 2, open-label, multicenter study in the first line treatment of HPV16 and PD-L1 positive recurrent or metastatic HNSCC: comparing results with KEYNOTE-048 study.
pembrolizumab + CUE-101	anti PD-L1 agent + a E7-pHLA-IL2- Fc Fusion Protein	NCT03978689	KEYNOTE KN-A78	I	recruiting	Phase1 study of CUE 101 in second line or CUE 101 with pembrolizumab in first line HPV+ R/M HNSCC. CUE-101 is a novel fusion proteinstimulating tumor specific T cells to eradicate HPV-driven malignancies.

Table 1. Continued.

Drugs	Drug types	Trial number	name	phase	status	Description
pembrolizumab + GSK3359609	anti PD-L1 + anti-ICOS agent	NCT04128696	INDUCE-3	II	recruiting	Randomized, double-blind, adaptive Phase II/III study comparing GSK3359609 inducible T cell co-stimulatory receptor (ICOS) agonist and pembrolizumab to pembrolizumab plus placebo in participants with programmed death receptor 1-ligand 1 (PD-L1) combined positive score (CPS) ≥ 1 R/M HNSCC.
nivolumab + ipilimumab + RT	anti PD-1 agent + anti-CTLA4 agent+ RT	NCT03799445		II	recruiting	Phase 2 Study in LAHPV-positive OPSCCs.
nivolumab + ipilimumab	anti PD-1 agent + anti-CTLA4 agent	NCT03003637	IMCISION	Ib/II	recruiting	Phase Ib/II trial in advanced stage HNSCC
durvalumab + tremelimumab + SBRT	anti PD-L1 agent + anti CTLA4 agent + SBRT	NCT03618134		Ib/II	recruiting	Phase Ib/II trial in HPV-positive OPSCCs.
monalizumab + cetuximab	anti NKG2a agent + anti EGFR-agent	NCT02643550	Interlink-1	Ib/II	recruiting	Phase 1b/2 trial in HPV+ and HPV- Recurrent or Metastatic HNSCC.

Table 1. Continued.

Drugs	Drug types	Trial number	name	phase	status	Description
monalizumab + durvalumab (among other cohorts)	anti NKG2a agent + anti PD-L1 agent	NCT03088059	UPSTREAM	II	recruiting	Biomarker-driven trial in recurrent or metastatic HNSCC progressing after first-line platinum-based chemotherapy. Based on potential biomarkers and molecular alterations identified in the biopsy from the central platform, patients will be allocated in different biomarker-positive and immunotherapy cohorts.
buparlisib (BKM120) + cisplatin + XRT	PI3K inhibitor + chemotherapy + radiotherapy	NCT02113878		Ib/II	active, not recruiting	Phase Ib study is combining standard chemoradiotherapy with weekly cisplatin and BKM120 in LAHNSCC.
alpelisib (BVL719)	PI3K inhibitor	NCT03601507		II	currently suspended (IRB consent forms)	Phase II trial studies of alpelisib in HPV-associated stage I-IVA HNSCC that can be removed by surgery.

(Status was checked on www.clinicaltrials.gov, August 2020)



10

Summary / Samenvatting

In this thesis the clinical features of HPV-related head and neck cancers and their implications for tumor staging and therapeutic strategies were studied. A couple of important, clinically relevant findings were made. Compared to HPV-negative tumors, HPV-positive tumors have different clinical and biological characteristics. The prognostic value of the severity of neck node involvement is significantly reduced in HPV-positive tumors, which we published as the first group worldwide. [Chapter 2]. A series of other and larger studies followed, which confirmed our observation and finally contributed to an adapted TNM classification for HPV-positive oropharyngeal squamous cell carcinomas (8th edition UICC tumor staging system OPSCCs), which is partly based on our data. Next, we showed that p16Ink4a-immunostaining is a strong surrogate marker for high risk-HPV16 in OPSCCs and tonsillar dysplasias [Chapter 3]. Definitions of HPV-positivity have impact on TNM-classification and patients' prognosis: adequate testing with at least PCR for detecting HPV DNA next to P16Ink4a-immunostaining is emphasized.

During our research the question arose whether the presence of HPV in neck metastases of cancers of unknown primary (CUP) could be used to identify the primary tumor location and to guide treatment. However, only a very low rate of HPV-positive metastases was found. [Chapter 4]. On the other hand we were able to show that de-intensified therapeutic regimens without postoperative radiation of the pharyngeal axis and the contralateral neck do not worsen prognosis, first in a smaller and more heterogeneous [Chapter 5] and subsequently in a larger and more homogeneous population [Chapter 6]. Results recently contributed to the publication of the first guideline on diagnosis and management of squamous cell carcinoma of unknown primary in the head and neck by the American Society of Clinical Oncology (ASCO).

In the recently introduced 8th edition UICC tumor staging system for HPV-positive tumors, still only anatomical parameters have been used to stage OPSCC, although in literature there were indications that non-anatomical parameters also are of significance to predict prognosis. We therefore tested the prognostic value of the 8th edition UICC tumor staging system for HPV-positive OPSCCs in a large series of exclusively tonsillar squamous cell carcinomas (TSCCs) and tested the prognostic value of other, non-anatomical factors [Chapter 7]. We have shown that the correlation between N-status and prognosis is still limited and that non-anatomical factors have a highly significant influence on prognosis (particularly non- and former smoking status and age). Based on this finding we proposed a further improved staging system for OPSCC.

The possibilities to improve existing and to develop new therapeutic options based on the different clinical and biological characteristics of HPV-positive compared to HPV-negative OPSCCs have been addressed in a review on the most promising approaches [Chapter 8]. Present therapeutic HPV-targeting strategies and future directions for de-intensified treatment of HPV-positive HNSCC were updated and are further debated in the general discussion [Chapter 9, point 9.8].

In dit proefschrift werden de klinische kenmerken van HPV-gerelateerde hoofdhalstumoren en hun implicaties voor tumorstadiëring en therapeutische

strategieën bestudeerd. Er werden een aantal belangrijke en klinisch relevante bevindingen gedaan. Vergeleken met HPV-negatieve tumoren hebben HPV-positieve tumoren andere klinische en biologische kenmerken. De prognostische waarde van de ernst van de betrokkenheid van lymfklieren is bij HPV-positieve tumoren significant verminderd, hetgeen wij als eerste groep wereldwijd publiceerden. [Hoofdstuk 2]. Een reeks andere en grotere studies volgden die onze observatie bevestigden, en uiteindelijk hebben geleid tot een aangepaste TNM classificatie voor HPV-positieve orofarynxcarcinomen (OPSCC's) (8e editie UICC tumor staging system), mede gebaseerd op onze data. Vervolgens toonden we dat p16Ink4a-immunostaining een sterke surrogaatmarker is voor hoog-risico-HPV16 in OPSCC's en tonsillaire dysplasieën [Hoofdstuk 3]. Definities van HPV-positiviteit hebben echter invloed op de TNM-classificatie en de prognose van patiënten: adequate tests met ten minste PCR voor het opsporen van HPV-DNA naast P16Ink4a-immunostaining wordt benadrukt.

Tijdens ons onderzoek kwam de vraag naar voren of de aanwezigheid van HPV in nekmetastasen van onbekende primaire tumoren gebruikt kan worden om de primaire tumorlocatie te identificeren en de behandeling te begeleiden. Er werd echter slechts een zeer laag percentage HPV-positieve metastasen gevonden. [Hoofdstuk 4]. Aan de andere kant konden we aantonen dat de prognose niet verslechterd wordt door minder intensieve therapeutische regimes zonder postoperatieve bestraling van de pharyngeale as en de contralaterale hals, eerst in een kleinere en meer heterogene populatie [Hoofdstuk 5] en later in een grotere en meer homogene populatie [Hoofdstuk 6]. De resultaten hebben recent bijgedragen aan de publicatie van de eerste richtlijn over de diagnose en behandeling van plaveiselcelcarcinoom van onbekende primaire tumoren in het hoofd en de nek door de American Society of Clinical Oncology (ASCO).

In de recent geïntroduceerde 8^e editie UICC tumor stadiëring voor HPV-positieve tumoren worden vooralsnog enkel anatomische parameters gebruikt om OPSCC's te classificeren, hoewel er in de literatuur aanwijzingen zijn dat ook niet-anatomische parameters van significante invloed zijn op de prognose. Daarom hebben we de prognostische waarde van de 8e editie TNM classificatie van OPSCC's getest in een grote reeks van uitsluitend tonsilcarcinomen en hebben we de prognostische waarde van andere, niet-anatomische factoren [Hoofdstuk 6] getest. We hebben aangetoond dat de correlatie tussen N-status en prognose nog steeds beperkt is en dat niet-anatomische factoren een zeer significante invloed hebben op de prognose (m.n. niet-roken en gestopt met roken, en leeftijd). Op basis hiervan hebben we een verbeterde stadiëring voor OPSCC's voorgesteld.

De mogelijkheden om bestaande en nieuwe therapeutische opties te verbeteren op basis van de verschillende klinische en biologische kenmerken van HPV-positieve in vergelijking met HPV-negatieve OPSCC's zijn onderzocht in een review van de hiervoor meest veelbelovende benaderingen [Hoofdstuk 7]. De huidige therapeutische strategieën gericht op HPV en toekomstige richtingen voor minder intensieve behandeling van HPV-positieve hoofd-halstumoren werden geüpdate en verder besproken in de algemene discussie [Hoofdstuk 9, punt 9.8].





Appendix

List of publication

Acknowledgments (Dankwoord)

Curriculum Vitae

LIST OF PUBLICATIONS

W1-publications

1. **Straetmans JMJA**, StuuT M, Wagemakers S, Hoebbers F, Kaanders JHAM, Speel EJM, Melchers WJG, Slootweg P, Kremer B, Lacko M, Takes RP. Tumor control of cervical lymph node metastases of unknown primary origin: the impact of the radiotherapy target volume. *Eur Arch Otorhinolaryngol*. 2020;277(6):1753-1761.
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1. **Straetmans JMJA**, Stuut M, Lacko M, Hoebers F, Speel EMJ, Kremer B. Additional parameters to improve the prognostic value of the 8th edition of the UICC classification for HPV-related oropharyngeal tumors.

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1. Decuypere FAG, Theunissen P, **Straetmans J**. Pilomatrixoma (calcifying epithelioma of Malherbe). *Eur. J. Pediatr. Dermatol*. 2014; 24:82-5.
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Jos Marie Jan Arthur Alfons Straetmans werd op 17 augustus 1981 geboren in Luik (Rocourt), België. Hij doorliep zijn middelbare onderwijs op het Onze-Lieve-Vrouwecollege in Tongeren waar hij in 1999 zijn humaniora afrondde in de afstudeerrichting Latijn-Wetenschappen. Aansluitend begon hij zijn studie geneeskunde aan de Universiteit van Maastricht die hij afrondde in 2005. In 2006 startte hij de opleiding tot KNO-arts aan het MUMC+ (opleiders: prof. dr. Bernd Kremer en prof. dr. Robert Stokroos) waarbij twee perifere stages werden doorlopen in resp. het Elkerliek Ziekenhuis Helmond (opleider: dr. Paul Schuil) en Catharina Ziekenhuis Eindhoven (opleider: dr. Frank Adriaansen). Aansluitend volgde hij een opleiding tot KNO-arts/Hoofd-halschirurg tijdens een fellowship Hoofd-halsoncologie dat hij doorliep in het MUMC+ (opleider: prof. dr. Bernd Kremer) en het NKI-AVL te Amsterdam (opleider: prof. dr. Bing Tan). Tijdens deze gehele opleidingsperiode werd de basis gelegd voor dit proefschrift. Hij bleef hierna iets meer dan een jaar verbonden aan het MUMC+ en startte parallel als chef-de-clinique in zowel het Laurentius Ziekenhuis in Roermond als het Atrium Medisch Centrum in Heerlen. Op 1 december 2014 werd hij voltijds maatschapslid van de vakgroep KNO in het Atrium MC te Heerlen, die opging in de vakgroep KNO Zuyderland Medisch Centrum, als gevolg van de fusie van het Atrium MC met het Orbis Medische Centrum Sittard. Vanaf 2017 was hij tevens bestuurslid bij het Medisch Specialistisch Bedrijf Zuyderland MC tot aan de oprichting van de huidige Bestuursraad medio 2019. Sinds 1 juni 2019 vervult hij de rol van voorzitter van de vakgroep KNO. Grensoverschrijdend Zuid-Limburg is zijn thuis waar hij woont met zijn vrouw Ilse en twee kinderen Camille en Julien. Samen genieten ze er van de mooie omgeving, lekker eten, en reizen naar het zuiden.



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