Modern Treatment Outcomes for Early T-Stage Oropharyngeal Cancer Treated With Intensity-modulated Radiation Therapy at a Tertiary Care Institution

Eric J. Di Gravio  
University of Western Ontario: Western University

Pencilla Lang  
University of Western Ontario: Western University

Hugh Andrew Jinwook Kim  
University of Western Ontario: Western University

Tricia Chinnery  
University of Western Ontario: Western University

Neil Mundi  
University of Western Ontario: Western University

S. Danielle MacNeil  
University of Western Ontario: Western University

Adrian Mendez  
University of Western Ontario: Western University

John Yoo  
University of Western Ontario: Western University

Kevin Fung  
University of Western Ontario: Western University

Joe S. Mymryk  
University of Western Ontario: Western University

John W. Barrett  
University of Western Ontario: Western University

Nancy Read  
University of Western Ontario: Western University

Varagur Venkatesan  
University of Western Ontario: Western University

Sara Kuruvilla  
University of Western Ontario: Western University

Lucas C. Mendez  
University of Western Ontario: Western University
Research

Keywords: radiation, chemoradiation, oropharyngeal cancer, toxicity

DOI: https://doi.org/10.21203/rs.3.rs-83234/v1

License: This work is licensed under a Creative Commons Attribution 4.0 International License. 
Read Full License
Abstract

Background

Transoral surgery (TOS), particularly transoral robotic surgery (TORS) has become the preferred modality in the United States for the treatment of early stage oropharyngeal cancer, largely due to assumptions of fewer toxicities and improved quality of life compared to primary radiotherapy (RT). However, these assumptions are based on retrospective analysis, a subset of which utilize primary RT groups not limited to T1-2 stage tumors for which transoral robotic surgery is FDA approved. Thus, there is potential for underestimating survival and overestimating toxicity, including treatment related mortality, in primary RT.

Methods

Consecutive cases of early T stage (T1-T2) oropharyngeal cancer presenting to the London Health Sciences Centre between 2014-2018 treated with RT or chemoradiation (CRT) were reviewed. Patient demographics, treatment details, survival outcomes and toxicity were collected. Toxicities were retrospectively graded using the CTCAE criteria.

Results

A total of 198 patients were identified, of which 82% were male and 73% were HPV-positive. Sixty-eight percent of patients experienced a grade 2 toxicity, 48% a grade 3 and 4% a grade 4. The most frequent toxicities were dysphagia, neutropenia and ototoxicity. The rates of gastrostomy tube dependence at 1 and 2 years were 2.5% and 1% respectively. There were no grade 5 (fatal) toxicities. HPV-positive patients experienced improved 5-year overall survival (86% vs 64%, p=0.0026).

Conclusions

Primary RT or CRT provides outstanding survival for early T-stage HPV-positive disease, with low rates of severe toxicity and feeding tube dependence. This study provides a reference for comparison for patients treated with primary transoral surgery.

Background

Over the past few decades, there has been a dramatic rise in the incidence of oropharyngeal squamous cell carcinoma (OPSCC), largely due to increasing rates of infection with human papillomavirus (HPV)\[i\]. HPV-associated OPSCC patients tend to be younger and healthier than traditional smoking- and alcohol-related OPSCC patients, and have a significantly improved prognosis. The 5-year survival of HPV-associated OPSCC exceeds 80%, making quality of life after treatment increasingly important since these patients may now survive for many years with significant treatment toxicities\[ii\]. This has led to intense interest in treatment de-escalation, with the goal of reducing the toxicity of standard dose chemoradiation therapy\[iii\]. In the United States, transoral surgery (TOS), particularly transoral robotic surgery (TORS), has largely become the preferred treatment modality for early T-stage OPSCC, as retrospective data has
suggested a more favourable toxicity profile\[iv\]. However, there is a paucity of data directly comparing the two modalities making the choice of treatment for OPSCC highly controversial\[v\].

The drive to adopt TOS as the standard of care revolves around the assumption that it carries a lower toxicity profile, and therefore better quality of life, than standard chemoradiation\[v\]. The ORATOR study, a phase II trial of 68 patients, is the only randomized clinical trial to directly compare quality of life between primary radiotherapy and primary TOS approaches, and included patients regardless of HPV status\[vi\]. In this trial, toxicity profiles differed between the two modalities. In particular, swallowing-related quality of life as measured by the MD Anderson Dysphagia Inventory was statistically superior in the chemoradiation group as compared to the TORS group, although the difference did not represent a clinically meaningful change6.

This discrepancy with previously published retrospective data can partially be explained by two potential sources of bias in previous studies: the inclusion of advanced stage OPSCC in chemoradiation cohorts, as well improved toxicity profiles of modern radiation techniques\[vii\]. TORS is only FDA approved for treatment of early stage (T1-2) OPSCC. However, many retrospective studies comparing TORS to chemoradiation include chemoradiotherapy cohorts containing advanced T-stage disease even though these patients are generally not considered candidates for TORS\[vi\],[vii],[viii],[ix],[x]. Furthermore, advances in radiotherapy such as intensity-modulated radiation therapy (IMRT) allow for more conformal treatment plans, reducing prevalence and severity of side effects such as dysphagia\[xii\].

The purpose of this study is to examine the modern outcomes and toxicity in patients with early stage disease (T1-2, N1-2) treated with IMRT ± chemotherapy at a high-volume tertiary care cancer centre. We include only patients with T1-2 N1-2 disease as these patients would be considered candidates for treatment with primary transoral laser or robotic surgery.

**Methods**

*Study Participants and Clinical Features*

Research Ethics Board approval (17222E) was obtained. A retrospective chart review was performed of all patients with early stage OPSCC treated with curative-intent radiotherapy ± chemotherapy who presented to the London Health Sciences Centre (LHSC) between 2014 and 2018. Early stage was defined as American Joint Committee on Cancer 7th Edition stage T1-T2, N0-N2. Patient, tumour and treatment-related factors collected included: gender, age at diagnosis, smoking and alcohol history, site of primary tumour (base of tongue, palatine tonsil, soft palate, vallecula, lateral or posterior pharyngeal wall, or unknown), TNM stage, HPV status, and use of concurrent chemotherapy. HPV status was determined with p16 immunohistochemical analysis with strong and diffuse staining in >70% of tumours cells considered positive. Patients are reported with the chemotherapy protocol that they initiated treatment with regardless of whether they completed the full course or were switched to a different protocol during
treatment. Alcohol abuse was defined as a history of >20 alcoholic beverages per week. A significant smoking history was defined as a total of 10 or more pack-years.

Radiotherapy

All patients received definitive intent radiotherapy with IMRT with fixed-gantry or rotational techniques (tomotherapy or volumetric modulated arc therapy [(VMAT)]. Patients generally received 70 Gy in 35 fractions (5 daily fractions delivered per week) to the gross disease, and 56 Gy in 35 fractions to the elective nodal volume. Patients were generally treated with concurrent high dose cisplatin (100mg/m$^2$ given every three weeks), excluding patients aged ≥ 70, those with comorbidities or poor performance status or those who declined. Weekly cisplatin (40mg/m$^2$) could be used at the discretion of the medical oncologist. If patients were not suitable for concurrent cisplatin, alternative chemotherapy regimens could include the Calais regimen (carboplatin and 5-fluorouracil) or cetuximab. If patients did not receive concurrent systemic therapy an accelerated fractionation (6 fractions per week delivered over 5 days) was used at the discretion of the treating oncologist. Unilateral radiation was used for tonsil primaries with less than 1 cm extension into the tongue base or palate with ≤ 1 ipsilateral lymph node. All other patients received bilateral treatment.

Evaluation and follow up

After completion of radiotherapy all patients were seen at 6 weeks for a clinical assessment of response and treatment toxicity, and then every 3 months for the first 2 years, and every 4 to 6 months in the third to fifth years. A follow-up CT scan was obtained at 3 months, with additional imaging, including PET-CT, if there was an incomplete response or if clinically indicated. Salvage surgery was reserved for patients who had persistent disease on follow-up imaging or clinically suspicious findings. All recurrences were histologically confirmed if possible.

Toxicity Assessment

Treatment toxicities were graded using the Common Terminology Criteria for Adverse Events (CTCAE) version 5.0\[i]. Toxicities or death occurring during or within 30 days of the end of treatment were included. Toxicities assessed included dysphagia, neutropenia, febrile neutropenia, ototoxicity and acute kidney injury. Any use of nasogastric, gastrostomy, gastrostomy-jejunostomy tube or total parental nutrition (TPN) within this time frame was considered a grade 3 dysphagia. Any change in diet not requiring the aforementioned interventions were considered grade 2 dysphagia.

Statistical Analysis

All statistics were carried out in the R statistical environment (version 3.4.0). Overall survival (OS) and disease-free survival (DFS) outcomes based on HPV status were compared using the log-rank statistical approach using the R survival package (version 2.41-3). A Cox regression model was used for multivariate analysis to assess the association of baseline variables with OS and DFS. Backward step-
wise analysis was used to create the final multivariate model of overall and disease-free survival analyses were done using the `survival` package (Additional Files 1 and 2). A p-value of <0.05 was considered statistically significant.

Results

Patients and treatment characteristics

A total of 198 patients (163 male and 35 female) treated with chemoradiation or radiotherapy alone were included in this analysis. Figure 1 depicts a flow-chart of the screening process for inclusion. The mean follow up for surviving patients was 27.4 months from the completion of treatment (range 0-63). Baseline and treatment characteristics are summarized in Table 1. Median age at diagnosis was 61 years with 114/198 patients (58%) having greater than a 10 pack-year smoking history and 49/198 (25%) consuming more than 20 alcoholic beverages per week. One hundred and forty-four of 198 (73%) of patients had proven p16 positive disease and 140/198 (71%) and 125/198 (63%) had stage T2 and N2 disease respectively. One hundred and fifty-nine (80%) patients were treated with chemoradiation with the most common chemotherapy protocol being monotherapy with cisplatin. Of patients receiving cisplatin, 61.4% received high dose treatment, while 38.5% received weekly treatment (Table 1). Of patients receiving high dose cisplatin, 70/80 (88%) received 2 or more cycles with 54/80 (68%) completing all three. Of patients treated with weekly cisplatin, 45/55 (82%) completed 5 or more cycles with only 17/55 (31%) completing all 7 cycles.

Table 1. Baseline and treatment characteristics. Data are presented as number (%) unless otherwise specified.
|                                | All patients (n=198) |
|--------------------------------|----------------------|
| Age, median (IQR)              | 61 (54-66)           |
| Sex                            |                      |
| Male                           | 163 (82%)            |
| Female                         | 35 (18%)             |
| Total Pack Years               |                      |
| ≤10 pack years                 | 84 (42%)             |
| ≥10 pack years                 | 114 (58%)            |
| Alcohol Consumption            |                      |
| ≤20 drinks per week            | 149 (75%)            |
| ≥20 drinks per week            | 49 (25%)             |
| Primary Site                   |                      |
| Tonsil                         | 108 (55%)            |
| Base of Tongue                 | 76 (38%)             |
| Soft palate                    | 9 (5%)               |
| Vallecula                      | 3 (2%)               |
| Indeterminate                  | 2 (1%)               |
| Clinical T-Stage               |                      |
| T1                             | 58 (29%)             |
| T2                             | 140 (71%)            |
| Clinical N-Stage               |                      |
| N0                             | 21 (11%)             |
| N1                             | 52 (26%)             |
| N2                             | 125 (63%)            |
| p16 Status                     |                      |
| Positive                       | 144 (73%)            |
| Negative                       | 28 (14%)             |
| Unknown                        | 26 (13%)             |
| Treatment                      |                      |
| Treatment | Count | Percentage |
|-----------|-------|------------|
| RT        | 39    | 20%        |
| CRT       | 159   | 80%        |

Radiation Laterality

| Laterality | Count | Percentage |
|------------|-------|------------|
| Unilateral | 14    | 7.1%       |
| Bilateral  | 163   | 82.3%      |
| Unknown    | 21    | 10.6%      |

CT Simulation

| Simulation | Count | Percentage |
|------------|-------|------------|
| With contrast | 85 | 42.9% |
| Without contrast | 92 | 46.5% |
| Unknown | 21 | 10.6% |

Chemotherapy Regimen

| Regimen | Count | Percentage |
|---------|-------|------------|
| Cisplatin | 135 | 68%        |
| High dose | 83  | 61.4%      |
| Weekly  | 52   | 38.5%      |
| Carboplatinum + 5-Fluorouracil | 14 | 7% |
| Cetuximab | 5   | 2.5%       |
| Other   | 5    | 2.5%       |

RT=radiotherapy only. CRT=chemoradiation

Treatment Toxicity

Data on acute toxicity is summarized in Table 2. One hundred and thirty five of 198 (68%) patients experienced at least one grade 2 toxicity, 95/198 (48%) a grade 3 toxicity and 8/198 (4%) a grade 4 toxicity. Of note, no patients experienced a grade 5 (fatal) toxicity during treatment. The most common toxicity was dysphagia followed by neutropenia and ototoxicity. Patients treated with radiation alone experienced fewer toxicities compared to patients treated with chemoradiation (p<0.0001, Fisher's exact test). Most notably, in total, 56/198 (28%) of patients experienced grade 3 dysphagia and 37/198 (19%) required placement of G/GJ tube. However, the rate of G/GJ tube use during treatment was significantly greater in patients treated with chemoradiation compared to patients treated with radiation alone (p<0.02, Fisher's exact test). Among patients treated with radiation alone, rates of grade 3 dysphagia and G/GJ tube insertion during treatment were 4/39 (10%) and 2/39 (5.1%) respectively, compared to 52/159 (33%) and 33/159 (21%) for patients treated with chemoradiation. Two patients (1 treated with radiation alone...
and the other with chemoradiation) required G/GJ tube placement more than a month after completion of treatment. In total, the 1- and 2-year rates of gastrostomy/ gastrostomy-jejunostomy tube use were 2.5 and 1% respectively (5 and 2 patients respectively). Only 8 patients (4%) experienced a grade 4 toxicity (2 ototoxicity, 6 neutropenia).

| Toxicity          | Radiotherapy group (n=39) | Chemoradiation group (n=159) |
|-------------------|---------------------------|-------------------------------|
|                   | Grade 1-2 | Grade 3 | Grade 4 | Grade 5 | Grade 1-2 | Grade 3 | Grade 4 | Grade 5 |
| Dysphagia         | 26 (67%)  | 4 (10%) | 0       | 0       | 99 (62%)  | 52 (33%)| 0       | 0       |
| Neutropenia       | 0         | 0       | 0       | 0       | 64 (40%)  | 34 (21%)| 6 (4%)  | 0       |
| Febrile neutropenia | 0         | 0       | 0       | 0       | 0         | 14 (9%) | 0       | 0       |
| Ototoxicity       | 1 (3%)    | 0       | 46 (29%)| 34 (21%)| 2 (1%)    | 0       |         |         |
| Acute Kidney Injury | 0         | 0       | 0       | 0       | 0         | 10 (6%) | 0       | 0       |

**Table 2.** Summary of toxicities. Data are presented as numbers (%). Grading is consistent with CTCAE version 5.0.

**Predictors of overall and disease-free survival**

In total, 32/198 (16%) patients included in this study died. Seventeen of these 32 (53%) patients died with disease still present, either as a direct result of their disease or due to unrelated causes, while 15/32 (47%) died of unrelated causes while in remission. Furthermore, 32/198 (16%) of patients experienced disease recurrence, of which 27/32 (84%) were histopathologically proven. Nine of 32 (28%) recurrences were local, 13/32 (41%) were regional, 8/32 (25%) were distant, 1/32 (3%) was both local and regional and 1/32 (3%) was both regional and distant. Nineteen of 32 (59%) patients with recurrence required salvage surgery.

Thirty-two of 198 patients with incomplete clinical information were omitted from inclusion in the survival model. Univariable analysis revealed age, smoking status, alcohol use and p16- status as significant prognostic factors for overall survival (Additional File 1). With multivariable analysis, smoking history (HR 3.59 95% CI [1.02-12.7], p=0.0472) remained a significant prognostic factor for overall survival (Additional File 1). Age and alcohol use were predictors of disease-free survival in univariate and multivariate analysis (Additional File 2). p16-positive status was associated with improved overall survival (p<0.01) but not disease-free survival (p=0.17, Figure 2).

**Patterns of treatment failure**
Disease relapse stratified by p16 status is outlined in Figure 3. Consistent with the literature, p16-negative patients were more likely to experience locoregional relapse than p16-positive patients (7/28 vs 10/144, p< 0.01 Fishers Exact test), and significantly more likely to die from that recurrence (7/7 vs 3/10, p<0.01)\(^3\). There were nine distant metastatic failures in the p16-positive cohort including lesions in the lung, liver and skeleton, while none occurred in the p16-negative group, however this was not statistically significant (9/144 vs 0/28, p=0.36)

**Discussion**

In this study, we demonstrated that while IMRT-based (chemo)radiation can have short- and long-term toxicities, early-stage HPV-associated OPSCC patients experienced excellent survival with acceptable toxicities, low long-term gastrostomy dependency rates and negligible treatment-related mortality (0% in this study). In contrast, prior studies have reported higher toxicity rates, with treatment related deaths of up to 3%\(^2\).[i]. Similarly, a frequently cited meta-analysis of the Radiation Therapy Oncology Group treatment intensification trials by Machtay and colleagues reported that 43% of patients treated with primary chemoradiation for OPSCC suffered a severe major late toxicity, including a 10% rate of long-term feeding tube dependency\(^15\). This is relevant as these studies are often cited as a reference comparison for primary transoral surgery\(^4,8-11\). While TOS can be carried out for advanced stage disease, in most case series, >85% of tumours are limited T1-T2 stage\(^4,8-10\). In contrast, the chemoradiation studies included high rates of advanced T stage disease (>75% in one such study by Ang and colleagues)\(^2\), introducing bias into the comparison of historical series. Furthermore, many of these studies were carried out in the pre IMRT era, which would likely impact function and toxicity\(^2,14\). Patients eligible for TOS have low-volume disease that may allow for unilateral radiotherapy and improved normal tissue sparing compared to historical chemoradiation cohorts that include a wider range of patients. There is a deficiency in the literature of reported outcomes of solely early-stage patients. This study reports the outcomes and toxicities of early-stage oropharynx patients undergoing primary radiotherapy with modern techniques, and provides a historical comparison cohort for discussions around surgical outcomes. Rates of toxicity and treatment-related mortality are lower than in previously published cohorts.

Febrile neutropenia is a potentially life threatening and frequent complication of chemoradiation in many studies\[^ii\],[^iii\]. For example, a study by Bledsoe and colleagues reported a rate of febrile neutropenia of 26% in patients treated with chemoradiation with 2/32 (6.3%) of these patients dying as a direct result\(^15\). In contrast, 19.7% of our early-stage patients did not require chemotherapy at all and thus were not at risk of this complication. In the chemoradiation cohort specifically, the rate of febrile neutropenia was only 8.8% and there were no fatalities (Table 2).

One of the strongest predictors of poor patient quality of life following treatment for head and neck cancer is long term gastrostomy tube dependence\[^iv\]. Chemoradiation studies report rates of up to 10% long-term dependence while most TOS studies show lower rates\(^4, [v],[vi]\). In a previous systematic review, the majority of studies demonstrated gastrostomy rates of less than 4.5%, with many (6/13) showing
rates of 0% suggesting superior swallowing function with primary surgery\(^4\). However, this study has 2-year gastrostomy rates of 1%. This is similar to the results of the ORATOR trial, and suggest that swallowing outcomes are similar between the two treatment strategies\(^6\). Given the similar survival, treatment selection for early OPSCC should be an informed decision made between clinicians and patients.

In this study, we also attempt to describe potential prognostic indicators for overall and disease-free survival. It is well-established that HPV-associated OPSCC has a significantly better overall and disease-free survival than HPV-negative patients\(^3\). In our study, overall survival but not disease-free survival was found to be significantly better in HPV-associated OPSCC, possibly due to limited sample size (Figure 1). In early stage disease, chemoradiation appeared effective regardless of HPV status, and the differences in overall survival may be partially related to the presumed increased incidence of comorbidities in HPV-negative patients and subsequent non-cancer related death. Likely for similar reasons, on multivariate analysis, a history of smoking was associated with a worse overall survival but not disease-free survival (Additional File 1 and 2).

Concurrent chemoradiation with 3 cycles of high-dose cisplatin is currently the standard treatment for locoregionally advanced OPSCC\[^vii\]. However, due to short- and long-term toxicities, and due to the fact that HPV-associated OPSCC is more sensitive to chemotherapy and radiation than HPV-negative OPSCC, there has been much interest in treatment de-escalation\[^viii\].\[^ix\]. Some current strategies under investigation include weekly cisplatin instead of high-dose cisplatin, lower radiation dose, or decreased adjuvant radiation and/or chemotherapy after surgery\[^x\].\[^xi\].\[^xii\]. Other strategies, such as using the EGFR monoclonal antibody cetuximab, have conclusively been shown to provide inferior survival without meaningfully improving quality of life which only further highlights the importance of balancing toxicity with survival.\[^xiii\]\[^xiv\].

There are a number of limitations with this current study. First of all, it is inherently difficult to accurately grade toxicities retrospectively, precluding the inclusion of other common but less severe toxicities such as mucositis, xerostomia or peripheral neuropathy. The follow-up period was limited and thus may have led to an underestimation of long-term toxicities. This data is limited to a single centre and thus may not accurately portray the true variability in the patient population. Lastly, a number of important potential confounders such as socioeconomic factors and compliance with treatment were not addressed.

**Conclusion**

Due to the excellent prognosis for HPV-positive OPSCC, there is considerable interest in the de-escalation of therapy. Early-stage patients have excellent survival, and current assumptions about the toxicities associated with chemoradiation are likely over-stated for patients with early-stage disease. Early-stage patients have low rates of gastrostomy tube dependence and treatment-related mortality. The lower rate of toxicity in early-stage patients compared to historical series is important to keep in mind when
comparing toxicity profiles between chemoradiation and TOS. This study can provide a reference for comparison for patients treated with primary transoral laser or robotic surgery in future trials.

**Abbreviations**

CRT = Chemoradiation  
CTCAE = Common terminology criteria for adverse events  
DFS = Disease free survival  
HPV = Human papillomavirus  
IMRT = Intensity modulated radiotherapy  
LHSC = London Health Sciences Centre  
OPSCC = Oropharyngeal Squamous Cell Carcinoma  
OS = Overall survival  
RT = Radiotherapy  
TORS = Transoral robotic surgery  
TOS = Transoral surgery  
VMAT = Volumetric modulated arc therapy

**Declarations**

*Ethics approval and consent to participate:* Research Ethics Board approval (17222E) was obtained.

*Consent for Publication:* Not applicable

*Availability of data and materials:* All data generated and analyzed during this study are included in this published article (and its supplementary information files).

*Competing interests:* The authors declare that they have no competing interests

*Funding:* This work was supported by Canadian Institutes of Health Research grant MOP#142491 to JSM and ACN. ACN was supported by the Wolfe Surgical Research Professorship in the Biology of Head and Neck Cancers Fund.

*Author’s contributions:* EJD, ACN and DAP designed the study. EJD, PL, HAJK, TC, SM and ACN performed the analyses. All authors read and approved the final manuscript. EJD and ACN had full access to all the
data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

**Acknowledgements**: Not applicable.

**References**

1. Chaturvedi AK, Anderson WF, Lortet-Tieulent J, et al. Worldwide trends in incidence rates for oral cavity and oropharyngeal cancers. *J Clin Oncol* 2013; 31: 4550-59

2. Ringash J. Survivorship and quality of life in head and neck cancer. *J Clin Oncol* 2015; 33: 3322-27

3. Ang KK, Harris J, Wheeler R, et al. Human papillomavirus and survival of patients with oropharyngeal cancer. *N Engl J Med* 2010 362: 24-35

4. Yeh DH, Tam S, Fung K, et al. Transoral robotic surgery vs radiotherapy for management of oropharyngeal squamous cell carcinoma: a systematic review of the literature. *Eur J Surg Oncol* 2015; 41: 1603-14

5. Cracchiolo JR, Baxi SS, Morris LG, et al. Increase in primary surgical treatment of T1 and T2 oropharyngeal squamous cell carcinoma and rates of adverse pathologic features: National Cancer Data Base. *Cancer* 2016; 122: 1523–32.

6. Nichols AC, Theurer J, Prisman E, et al. Radiotherapy versus transoral robotic surgery and neck dissection for oropharyngeal squamous cell carcinoma (ORATOR): an open-label, phase 2, randomised trial. *Lancet* 2019; 20: 1349-59

7. de Almeida JR, Byrd JK, Wu R, et al. A systematic review of transoral robotic surgery and radiotherapy for early oropharynx cancer: a systematic review. *Laryngoscope* 2014; 124: 2096-102

8. Weinstein GS, O’Malley BW, Magnuson JS, et al. Transoral robotic surgery: a multicenter study to assess feasibility, safety and surgical margins. *Laryngoscope* 2012; 122: 1701-7

9. Weinstein GS, O’Malley BW, Cohen MA, et al. Transoral robotic surgery for advanced oropharyngeal carcinoma. *JAMA* 2010; 136: 1079-85

10. Moore EJ, Olsen SM, Laborde RR, et al. Long-term functional and oncological results of transoral robotic surgery for oropharyngeal squamous cell carcinoma. *Mayo Clin Proc* 2012; 87: 210-25

11. Holsinger FC & Ferris RL. Transoral endoscopic head and neck surgery and its role within the multidisciplinary treatment paradigm of oropharynx Cancer: robotics, lasers and clinical trials. *J Clin Oncol* 2015; 33: 3285-92
12. Gupta T, Kannan S, Ghosh-Laskar S, Agarwal JP. Systematic review and meta-analyses of intensity-modulated radiation therapy versus conventional two-dimensional and/or three-dimensional radiotherapy in curative-intent management of head and neck squamous cell carcinoma. *PLoS one* 2018; 13: e0200137.

13. S. Department of Health and Human Services, National Institutes of Health, National Cancer Institute. Common terminology criteria for adverse events (CTCadverse event) version 5.0. 2017.

14. Machtay M, Moughan J, Trotti A, et al. Factors associated with severe late toxicity after concurrent chemoradiation for locally advanced head and neck cancer: an RTOG analysis. *J Clin Oncol* 2008; 26:3582-89

15. Bledsoe TJ, Noble, AR, Hunter GK, et al. Oropharyngeal squamous cell carcinoma with known human papillomavirus status treated with definitive chemoradiotherapy: patterns of failure and toxicity outcomes. *Radiat Oncol* 2013; 174 doi:10.1186/1748-717X-8-174

16. Urban D, Corry J, Solomon B, et al. Weekly cisplatin and radiotherapy for low risk, locoregionally advanced human papillomavirus-positive oropharyngeal squamous cell carcinoma. *Head Neck* 2015; 38: E1117-21

17. Wells M, Swartzman S, Lanh H, et al. Predictors of quality of life in head and neck cancer survivors up to 5 years after end of treatment: a cross-sectional survey. *Support Care Cancer* 2016; 24: 2463-72

18. Chen AM, Daly ME, Luu Q, et al. Comparison of functional outcomes and quality of life between transoral surgery and definitive chemoradiotherapy for oropharyngeal cancer. *Head Neck* 2014; 37: 382-385

19. Sharma A, Patel S, Baik FM, et al. Survival and gastrostomy prevalence in patients with oropharyngeal cancer treated with transoral robotic surgery vs. chemoradiotherapy. *JAMA Otolaryngol Head Neck Surg* 2016; 142: 691-697

20. You EL, Henry M and Zeitouni AG. Human papillomavirus-associated oropharyngeal cancer: review of current evidence and management. *Curr Oncol* 2019; 26:119-123

21. Oosthuizen JC & Doody J. De-intensified treatment in human papillomavirus-positive oropharyngeal cancer. *Lancet* 2019; 393: 5-7

22. Price KAR, Nichols AC, Shen CJ, et al. Novel strategies to effectively de-escalate curative-intent therapy for patients with HPV-associated oropharyngeal cancer: current and future directions. *Am Soc Clin Oncol Educ Book* 2020; 40:1-13

23. Deschuymer S, Mahanna H & Nuyts S. Toxicity reduction in the treatment of HPV positive oropharyngeal cancer: emerging combined modality approaches. *Front Oncol* 2018; 8: 439
24. Hargreaves S, Beasley M, Hurt C, et al. Deintensification of adjuvant treatment after transoral surgery in patient with human papillomavirus-positive oropharyngeal cancer: the conception of the PATHOS study and its development. *Front Oncol* 2019; 9: 936

25. Nichols AC, Lang P, Prisman E, et al. Treatment de-escalation for HPV-associated oropharyngeal squamous cell carcinoma with radiotherapy vs. trans-oral surgery (ORATOR2): study protocol for a randomized phase II trial. *BMC Cancer* 2020; 20:125

26. Mehanna H, Robinson M, Hartley A, et al. Radiotherapy plus cisplatin or cetuximab in low-risk human papillomavirus-positive oropharyngeal cancer (De-ESCALaTE HPV): an open-label randomised controlled phase 3 trial. *Lancet* 2019; 393: 51-60

27. Gillison ML, Trotti AM, Harris J, et al. Radiotherapy plus cetuximab or cisplatin in human papillomavirus-positive oropharyngeal cancer (NRG oncology RTOG 1016): a randomised, multicentre, non-inferiority trial. *Lancet* 2019; 393: 40-50