Prevalence and prognostic impact of retropharyngeal lymph nodes metastases in oropharyngeal squamous cell carcinoma: Meta-analysis of published literature

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Abstract

**Background:** This systematic review and meta-analysis aims to estimate the prevalence and prognostic impact of retropharyngeal lymph node metastases (RLNMs) in oropharyngeal squamous cell carcinoma (OPSCC).

**Methods:** This meta-analysis was conducted according to PRISMA guidelines. Inclusion criteria: studies with more than 20 patients reporting the prevalence or prognostic impact of RLNMs in OPSCC. Whenever available, data on HPV status and subsites were extracted.

**Results:** Twenty-two articles were included. The overall prevalence of RLNMs in OPSCC was 13%, with no significant differences depending on HPV status. The highest prevalence was observed for posterior pharyngeal wall SCC (24%), followed by soft palate (17%), palatine tonsil (15%), and base of tongue (8%). RLNMs were associated with a significantly higher risk of death (HR: 2.54; IC95% 1.89–3.41) and progression (HR: 2.44; IC95% 1.80–3.30).

**Conclusions:** The prevalence of RLNMs in OPSCC was 13%, being higher in tumors of the posterior pharyngeal wall. RLNMs were associated with unfavorable outcomes.

**KEYWORDS**
HPV, human papillomavirus, oropharyngeal cancer, retropharyngeal lymph nodes, retropharyngeal metastasis

1 INTRODUCTION

The management of retropharyngeal lymph nodes (RLNs) in head and neck squamous carcinoma (HNSCC) is still an open issue. Anatomically, RLNs are located within a thin fat pad in the retropharyngeal space which is delimited anteriorly by the pharynx and its buccopharyngeal fascia and posteriorly by the alar fascia which itself forms the anterior aspect of the danger space and the prevertebral plane. RLNs are classified in medial and lateral groups.
Robust data regarding the prevalence of RLN metastases (RLNMs) are lacking given that these lymph node chains are relatively inaccessible to clinical examination and fine needle aspiration biopsy, and they are not routinely removed in neck dissection. Therefore, there are insufficient clinicopathological correlations, such as size and shape of the nodes, to be able to define radiological criteria suggestive of their metastatic involvement. Consequently, the prevalence of RLNMs varies widely in the literature. Based on a few histological and radiological studies, rates of RLNMs in HNSCC ranges from 9% to 50%, with nasopharyngeal carcinoma (NPC) exhibiting the greatest propensity to metastasize to the RLNs (29–89%). Within non-nasopharyngeal HNSCCs, oropharyngeal squamous cell carcinomas (OPSCCs) have been reported to have the highest rate of RLNMs, although there is a wide range of prevalence across studies, ranging from 6% to 23%. Given the uncertainty regarding the real rate of RLNMs, its prognostic impact is controversial, especially considering the heterogeneous interpretations of results predominantly from small case series. Even more inconsistent results have been reported when considering the potential impact of human papillomavirus (HPV) status and the different oropharyngeal subsites on the rate of metastasis to the RLNs.

This systematic review and meta-analysis aims to summarize the current evidence on the prevalence of RLN metastasis in OPSCCs and on its prognostic relevance, with a focus on HPV status and different oropharyngeal subsites.

2 | METHODS

2.1 | Outcome measures

The primary outcome measure was the prevalence of metastases in RLNs in OPSCC, defined as the number of patients with metastatic involvement of the RLN/total number of patients. Pooled prevalence was estimated separately depending on the diagnostic tool used to assess the RLN metastatic involvement, that is, histopathological or imaging evaluation. The rate of RLNMs stratified by HPV status (as detected by p16 immunohistochemistry, HPV-DNA in situ hybridization or PCR) and by OPSCC subsites (tonsil, base of the tongue, lateral and posterior pharyngeal wall, and soft palate) were also explored. The secondary outcome of this meta-analysis was the prognostic impact of RLNMs for the following outcomes, (a) overall survival (OS), defined as the time from diagnosis or initiation of treatment to patient death, irrespective of cause; (b) disease free survival (DFS), defined as the time from diagnosis or initiation of treatment until tumor recurrence/progression or any-cause death; (c) loco-regional control (LRC), defined as the time from diagnosis or initiation of treatment to the first locoregional event; and (d) distant metastasis (DM), defined as the time from diagnosis or initiation of treatment to the first distant event.

2.2 | Search strategy

This systematic review and meta-analysis was conducted following the preferred reporting items for systematic reviews and meta-analysis (PRISMA) checklist. The databases Medline (via Ovid), Embase (via Ovid), Cochrane Library, Web of Science (core collection), and Scopus were searched from inception to May 2021. The search terms used are reported below. Prior to searching the databases, the search terms were peer reviewed by three authors (PBR, DB, and EZ) to ensure they conformed to PRESS guidelines. “Head and Neck Neoplasms” OR “Esophageal Neoplasm” OR “Facial Neoplasm” OR “Mouth Neoplasm” OR “Otorhinolaryngologic Neoplasm” OR “Tracheal Neoplasm” OR “Head and Neck neoplasm” OR “Carcinoma” OR “Squamous Cell” OR “Mouth Neoplasms” OR “Oral Cavity Neoplasm” OR “Oropharyngeal Neoplasms” OR “Oropharyngeal Neoplasm” OR “Hypopharyngeal Neoplasms” OR “Hypopharyngeal Neoplasm” OR “Head and Neck Cancer” OR “Esophageal Cancer” OR “Mouth Cancer” OR “Otorhinolaryngologic cancer” OR “Parathyroid Cancer” OR “Thyroid Cancer” OR “Trachea Cancer” OR “Squamous Cell Carcinoma” OR “Oropharyngeal Cancer” OR “Hypopharyngeal Cancer” AND “Hypopharyngeal Node.” The reference list of articles included in this review were also manually searched to minimize the risk of data loss. Two authors (DB and EZ) independently screened all titles and abstracts identified by the search and then evaluated the full text of the reports that respected the inclusion criteria. A third author (PBR) settled any disagreement between reviewers. A flow diagram illustrating all the steps that led to the selection of the reports eligible for meta-analysis is shown in Figure 1.

2.3 | Inclusion criteria

Studies were included in the analysis if they met the following criteria: (1) studies reporting the prevalence of RLNMs, based on imaging, that is, computed tomography (CT), magnetic resonance imaging (MRI), or positron emission tomography/CT (PET/CT), or pathological examination and/or the prognostic role of RLNMs in OPSCC; (2) studies in which RLN involvement was assessed radiologically or by histopathological
examination; (3) studies including at least 20 patients with OPSCC; and (4) studies reporting sufficient data for statistical analysis. Authors were contacted by email if a study met the inclusion criteria, but data were not immediately extractable. Non-English language studies were excluded. Studies containing aggregated data or duplicated data from previously published work were excluded, as were review articles, case reports, editorials, and letters. Two authors (JP, PBR) independently assessed the quality of the included studies with the Newcastle-Ottawa Scale. Low-quality articles (Newcastle–Ottawa Scale [NOS] score <6) were also excluded.

2.4 Data extraction and statistical analysis

The number of total cases and of positive RLNs were extracted from each study. When available, data were extracted according to HPV status and cancer subsite. To provide a summary estimate of the association between RLN-positivity and oncological outcomes, the hazard ratio (HR) was extracted, when reported. The standard error of the log hazard ratio (HR) was derived from the log confidence intervals (CIs). Summary estimates of proportion or hazard ratio (sHR), with corresponding 95% CI, were calculated according to random-effects models of DerSimonian and Laird, which incorporates both within- and between-study variability, as a weighted average giving each study a weight proportional to its precision. Analyses by strata according to HPV status and cancer subsite were conducted including studies with \( \geq 10 \) cases in the stratum under investigation. Statistical heterogeneity among studies was evaluated using the \( I^2 \) and \( \tau^2 \) statistics. Influence analysis was performed when the summary estimate was estimated from five or more studies: the summary estimate was calculated by omitting one study at a time. Publication bias was assessed through a funnel plot. The results of the meta-analysis were presented graphically using forest plots, plotting the estimates from individual papers, summary estimate, and 95% CI. Statistical significance was claimed for \( p < 0.05 \).

3 RESULTS

3.1 Search results and study characteristics

We identified a total of 949 potentially relevant articles with our initial search strategy. After screening the titles and abstracts of these articles, we excluded 922 studies because they were duplicated, or they did not meet the inclusion criteria (Figure 1). After reading 27 potentially

![FIGURE 1 PRISMA flow diagram](Color figure can be viewed at wileyonlinelibrary.com)
### TABLE 1  
Description of included studies

| Study                  | Study type | No. of patients | Cohort details               | RLN diagnosis     | Radiology criteria                                                                 | Treatment |
|------------------------|------------|----------------|------------------------------|-------------------|-------------------------------------------------------------------------------------|-----------|
| Rosen, 2021            | R          | 266            | OPSCC CT-PET/CT              | LN >8 mm, SUV > 2.5 | RT/RTCT                                                                             |           |
| Billfalk, 2019         | R          | 280            | cT1-T2 N1 OPSCC HPV+         | CT-MRI            | LN >8 mm                                                                             | RT/ CTRT |
| Iyizoba-Ebozue, 2020   | R          | 402            | OPSCC CT-MRI-PET/CT          | LN short axis ≥5 mm, necrosis and/or abnormal uptake on PET-CT | RT/RTCT |
| Lin, 2019              | R          | 796            | OPSCC HPV+                   | CT-MRI-PET/CT     | LN short axis ≥5 mm, long axis ≥ 10 mm, presence of any medial RLN; central necrosis; ≥2 clustered RPLNs; SUV >4.5 | RT/RTCT |
| Bhattassali, 2018      | R          | 238            | cT1-2 N1 OPSCC HPV +         | CT-MRI-PET/CT     | N/A                                                                                 | RT/ CTRT |
| Park, 2019             | R          | 71             | Tonsil cancer                | CT-MRI-PET/CT     | N/A                                                                                 | S         |
| Troob, 2017            | R          | 30             | OPSCC                        | CT-PET/CT         | N/A                                                                                 | S         |
| Spector, 2016          | R          | 205            | Stage III/IV OPSCC           | CT-PET/CT         | LN > 10 mm, abnormal SUV, cystic or necrotic, rENE                                   | RT/RTCT |
| Baxter, 2015           | R          | 165            | OPSCCs HPV+                  | CT-MRI-PET/CT     | LN abnormal SUV, LN short axis >6 mm, central necrosis or clustered.                | RT/ CTRT |
| Chung 2015             | R          | 54             | OPSCC                        | CT-MRI-PET/CT     | N/A                                                                                 | S         |
| Samuels, 2015          | R          | 231            | OPSCC HPV +                  | CT-MRI-PET/CT     | LN long axis >1 cm, necrotic/ cystic, abnormal SUV.                                  | RT/RTCT |
| Gunn, 2013             | R          | 981            | OPSCC                        | CT-MRI-PET/CT     | LN short axis ≥5 mm or long axis > 10 mm; necrosis, hypodensity; >1 lateral RLN, SUV >4.5; any medial RLN. | RT/RTCT |
| Moore, 2013            | R          | 72             | OPSCC                        | CT-MRI-PET/CT     | N/A                                                                                 | S         |
| Tang, 2013             | R          | 164            | OPSCC                        | CT-MRI-PET/CT     | SUV > 3 or LN short axis >6 mm.                                                     | RT/RTCT |
| Chung, 2011            | R          | 76             | Tonsil cancer                | CT-MRI-PET/CT     | N/A                                                                                 | S         |
| Chan, 2010             | P          | 102            | OPSCC                         | PET/CT            | Any medial RLN, abnormal SUV, LN short axis >5 mm                                    | RT/ CTRT |
| Tazzin, 2010           | R          | 53             | OPSCC                        | PET/CT            | SUV >3, LN ≥10 mm, any suspicious feature.                                            | RT/RTCT |
| Chu, 2009              | R          | 29             | OSCC, OPSCC, HPSCC           | CT-MRI-PET/CT     | LN >8 mm, abnormal density/ asymmetry/ enhancement. SUV >2,5                        | S         |
| Yoshimoto, 2007        | P          | 84             | OPSCC                        | Histopathology    | N/A                                                                                 | S OR RT/RTCT |
| Dirix, 2006            | R          | 208            | OPSCC                        | CT                | LN axis > 10 mm or central hypodensity                                              | S OR RT/RTCT |
| Shimizu, 2006          | R          | 77             | OPSCC                        | Histopathology    | N/A                                                                                 | S         |
| McLaughlin, 1995       | R          | 443            | H&N                          | CT-MRI            | LN > 10 mm, central hypodensity.                                                    | S OR RT/RTCT |

Abbreviations: LN, lymph node; P, Prospective; PET, Positron emission tomography; R, Retrospective; rENE, radiological extra nodal extension; RLN, Retropharyngeal lymph node; SUV, standardized uptakevalue; OPSCC, oropharyngeal squamous cell carcinoma; OSCC, oral squamous cell carcinoma; HPSCC, hypopharyngeal squamous cell carcinoma; H&N, head and neck.
eligible articles in detail, we determined that 16 studies met our inclusion criteria: the study by Bussels et al. was excluded because it reported the same cohort as of Dirix et al. Ten additional studies, of which six were eligible for meta-analysis, were identified through checking the reference lists of initial 27 eligible articles. Overall, 22 articles were included in the final analysis including 5027 patients with OPSCC. Among studies reporting the involved subsites of the oropharynx: 2242 subjects (52.1%) had palatine tonsil SCC, 1689 (39.3%) had SCC of the base of the tongue (BOT), 187 (4.3%) and 150 (3.4%) had posterior pharyngeal wall (PPW) and soft palate SCC, respectively. Thirteen authors were contacted in order to gather useful data to include in the metaanalysis but only one provided the requested information. The characteristics of the included studies are presented in Table 1.

### 3.2 | Quality assessment

The quality of included studies was high (Newcastle-Ottawa Scale score ≥7) in 18 (81%) of 22 studies, with a median of 7 (interquartile range: 7–8). A detailed report on the quality of included studies according to the Newcastle-Ottawa Scale is reported in Supplementary Table 1.

#### 3.3 | Prevalence of metastatic RLNs in OPSCC

All the eligible studies provided information about the prevalence of the metastatic involvement of RLN in OPSCC. When more subsites of the head and neck were analyzed, only data about OPSCCs were extracted. As shown in Figure 2, the pooled prevalence of RLN in OPSCC was 0.13 (CI 95% 0.10–0.16). The pooled prevalence of RLNMs was 0.13 (CI 95% 0.06–0.24) and 0.12 (CI 95% 0.10–0.14) when evaluated by histopathological analysis and imaging findings, respectively. Table 1 summarizes all criteria adopted by authors for this purpose. Among dimensional criteria, three authors used an 8 mm cut-off, seven 10 mm, and four a short axis >5 mm. As qualitative or metabolic features, central necrosis or hypodensity, clustered LN, and Standardized FIGURE 2 Forest plot showing the pooled prevalence of RLN metastasis in OSCC

| Study | RLN+ | OPSCC | Proportion (95% CI) |
|-------|------|-------|---------------------|
| (A) Radiology | | | |
| Rosen, 2021 | 37 | 266 | 0.14 (0.10-0.19) |
| Iyozra-Ebozue, 2020 | 40 | 402 | 0.10 (0.07-0.13) |
| Billfolk-Kelly, 2019 | 23 | 280 | 0.08 (0.05-0.12) |
| Lin, 2019 | 73 | 796 | 0.09 (0.07-0.11) |
| Bhattacharj, 2016 | 18 | 238 | 0.08 (0.05-0.12) |
| Spector, 2016 | 37 | 205 | 0.18 (0.13-0.24) |
| Baxter, 2015 | 16 | 165 | 0.10 (0.06-0.15) |
| Samuels, 2015 | 29 | 231 | 0.13 (0.09-0.18) |
| Gunn, 2013 | 94 | 981 | 0.10 (0.08-0.12) |
| Tang, 2013 | 19 | 164 | 0.12 (0.07-0.17) |
| Chan, 2010 | 25 | 102 | 0.25 (0.17-0.34) |
| Tauxin, 2010 | 11 | 53 | 0.21 (0.11-0.34) |
| Dix, 2006 | 34 | 208 | 0.16 (0.12-0.22) |
| McLaughlin, 1995 | 46 | 443 | 0.10 (0.08-0.14) |
| **Subtotal (Random effects)** | | | **4534** |
| Heterogeneity: $I^2 = 75\%$, $\tau^2 = 0.0974$, $p < 0.01$ | | | | |
| (B) Surgery/Histopathology | | | |
| Park, 2019 | 2 | 71 | 0.03 (0.00-0.10) |
| Troob, 2017 | 6 | 30 | 0.20 (0.08-0.39) |
| Chung, 2015 | 22 | 54 | 0.41 (0.28-0.55) |
| Moore, 2013 | 8 | 72 | 0.11 (0.05-0.21) |
| Chung, 2011 | 9 | 76 | 0.12 (0.06-0.21) |
| Chu, 2009 | 10 | 29 | 0.34 (0.18-0.54) |
| Yoshimoto, 2007 | 2 | 84 | 0.02 (0.00-0.08) |
| Shimizu, 2006 | 10 | 77 | 0.13 (0.06-0.23) |
| **Subtotal (Random effects)** | | | **493** |
| Heterogeneity: $I^2 = 84\%$, $\tau^2 = 1.0117$, $p < 0.01$ | | | | |
| **Total (Random effects)** | | | **5027** |
| Heterogeneity: $I^2 = 81\%$, $\tau^2 = 3.301$, $p < 0.01$ | | | | |
| Test for subgroup differences: $p = 0.85$ | | | | |
FIGURE 3 Forest plot showing the pooled prevalence of RLN metastasis in OSCC according to HPV-status

FIGURE 4 Forest plot showing pooled prevalence of RLN metastasis for different oropharyngeal subsites (A) Base of tongue (B) Tonsil (C) Soft palate (D) Posterior pharyngeal wall
Uptake Value (SUV) >4.5\textsuperscript{8,12} have been generally considered as pathologic. No medial RLNMs were reported in any of the seven studies providing data about their involvement.

### 3.4 | Prevalence of metastatic RLN according to HPV status

When stratified according to HPV status (Figure 3), the prevalence of RLN metastasis was 0.10 (95% CI 0.03–0.31) in HPV-negative cancers, while it was 0.12 (95% CI 0.10–0.15) in HPV-positive ones.

#### 3.5 | Prevalence of metastatic RLN according to oropharyngeal subsites

Oropharyngeal subsites were divided into tonsil, base of tongue and anterior pharyngeal wall, posterior and lateral pharyngeal wall, and soft palate or superior pharyngeal wall. Three studies did not provide a specific oropharyngeal localization.\textsuperscript{19,27,37} Figure 4 shows the prevalence of RLN metastasis according to oropharyngeal subsite. Posterior and lateral pharyngeal wall was the subsite with the highest prevalence of RLN metastasis (Figure 4D) with 0.24 (95% CI: 0.18–0.31), followed by soft palate (Figure 4C), 0.17 (0.08–0.30). The prevalence

![Figure 5](image) Forest plot showing hazard ratios for overall survival, disease free survival, locoregional recurrence and distant recurrence

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**TABLE 5**

| Study                        | RLN+ | RLN- | Hazard Ratio | HR (95% CI) |
|------------------------------|------|------|--------------|-------------|
| (A) Overall survival        |      |      |              |             |
| Rosen, 2021\textsuperscript{19} | 37   | 229  | 2.76         | (1.37-5.56) |
| Bhattachai, 2018\textsuperscript{21} | 18   | 220  | 3.32         | (1.19-9.26) |
| Samuels, 2015\textsuperscript{24} | 29   | 202  | 2.11         | (1.04-4.28) |
| Tang, 2013\textsuperscript{16}  | 19   | 145  | 2.40         | (0.90-6.40) |
| Total (Random effects)      | 103  | 796  | 2.54         | (1.89-3.41) |
| Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0.0048$, $p = 0.90$ | | | | |

| (B) Disease-free survival   |      |      |              |             |
| Bhattachai, 2018\textsuperscript{21} | 18   | 220  | 3.36         | (1.34-8.42) |
| Samuels, 2015\textsuperscript{24} | 29   | 202  | 2.25         | (1.08-4.69) |
| Tang, 2013\textsuperscript{16}  | 19   | 145  | 2.10         | (0.91-4.85) |
| Total (Random effects)      | 89   | 824  | 2.44         | (1.80-3.30) |
| Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0.0060$, $p = 0.89$ | | | | |

| (C) Local recurrence        |      |      |              |             |
| Dirix, 2006\textsuperscript{7} | 34   | 174  | 4.29         | (3.33-5.53) |
| Total (Random effects)      | 71   | 403  | 3.46         | (1.80-6.65) |
| Heterogeneity: $I^2 = 52\%$, $\tau^2 = 0.1396$, $p = 0.15$ | | | | |

| (D) Distant recurrence      |      |      |              |             |
| Bhattachai, 2018\textsuperscript{21} | 18   | 220  | 3.51         | (1.43-8.62) |
| Total (Random effects)      | 89   | 688  | 4.20         | (1.83-9.62) |
| Heterogeneity: $I^2 = 0\%$, $\tau^2 = 0.0300$, $p = 0.54$ | | | | |
of RLN metastasis was 0.15 (95% CI: 0.12–0.19) in tonsillar cancer (Figure 4B), while base of tongue (Figure 4A) showed the lowest frequency of involvement 0.08 (95% CI: 0.05–0.10).

3.6 | Outcome measures

Six articles were eligible for survival analysis. Among them only one did not perform multivariate analysis. All others specified which parameters were included in multivariate analysis. T category, N category, age, and smoking status were included in all studies which performed multivariate analysis. Four articles provided HRs and 95% CIs for OS. As shown in Figure 5, patients with metastatic involvement of the RLN had a significantly poorer OS than those without RLNMs (sHR = 2.54, 95% CI: 1.89–3.41). There was no significant heterogeneity among these four articles ($I^2 = 0\%$, $p = 0.90$). Four articles provided HRs and 95% CIs of DFS, $I^2 = 0\%$, $p = 0.89$). Only two articles provided HRs and 95% CIs of LRC: RLN positive patients were at higher risk for local(recional) recurrence (HR = 3.46, 95% CI: 1.80–6.65). Finally, concerning DM, three studies were included: RLN positive patients were also at higher risk for distant metastases (HR = 4.20, 95% CI: 1.83–9.62).

3.7 | Publication bias and sensitivity analysis

We inspected publication bias through a funnel plot. The result (Supplementary Figure 1A) indicated a lack of publication bias (test for asymmetry in funnel plot: $p = 0.295$). Furthermore, influence analysis was conducted on the estimation of prevalence of RLN metastases: no relevant impact emerged by any study (Supplementary Figure 1B), with prevalence estimates ranging from 12% to 13%.

4 | DISCUSSION

Combining data from 22 studies, the pooled prevalence of RLNMs in OPSCC was 13.0%. No significant differences emerged according to HPV status and the method used to assess RLN metastatic involvement. On the other hand, RLNMs were observed to be more prevalent in cancers of the posterior pharyngeal wall compared with other subsites. Finally, RLNMs were associated with an unfavorable prognostic outcome in patients with OPSCC, doubling the risk of death and progression and increasing by more than 3 times the risk of local and distant recurrence.

According to our pooled data, several individual studies were consistent in reporting a higher prevalence of RLNMs in patients with SCC of the posterior pharyngeal wall. Chung et al. suggested that this may be due to the contiguity of the retropharyngeal space with the pharyngeal wall. On the contrary, tonsil and BOT, which account for the vast majority of OPSCC cases, were less associated with RLNMs. The tonsil and BOT drain first to jugulodigastric nodes, and only then to the retropharyngeal space with only the involvement of the posterior pillar of the tonsil increasing the risk of RLNMs. The incidence of OPSCC is increasing worldwide, due to the emerging role of high risk strains of HPV in oropharyngeal carcinogenesis. The attributable fraction of HPV-driven OPSCC is globally 31% and it is higher for SCC arising from the palate tonsil and BOT as opposed to SCC arising from the soft palate and posterior pharyngeal wall, which are very rarely caused by HPV infection but are also less frequent (4.4% in the present systematic review). Thus, as the majority of HPV-negative OPSCC still arise from the tonsil and BOT, an expected at least indirect association between HPV-status and the risk of RLNMs was not observed in the present meta-analysis.

Among the seven eligible papers that reported information about medial RLN, none found medial RLN involvement. Bussels et al. found only one patient out of 208 (0.5%) with medial RLNMs. This patient also had a pathologic contralateral lateral RLN and the primary tumor was located in the posterior pharyngeal wall. Kim et al. found five medial RLNMs (11.6%) in a cohort of HNSCC where oropharyngeal and hypopharyngeal cancers counted for 54% and 39% of patients, respectively. Medial RLN are usually not evident in adults, therefore, their presence must be considered pathognomonic for metastatic involvement. Medial RLNMs are also less common than involvement of the lateral RLN in NPC, with one report identifying them only in six patients out of 3100 (0.2%). Unfortunately, no studies have been conducted specifically to investigate their clinical impact and prevalence in OPSCC; rather they have been reported only as collateral findings.

Only six articles reported sufficient survival data and could therefore be included in this meta-analysis. Results demonstrate a statistically significant trend towards unfavorable prognosis in patients with RLNMs in terms of
OS, DFS and recurrence. However, there is a lack of consensus in the literature on the topic. Indeed, several studies reported no statistical differences in terms of outcome. All of these had in common the fact that their cohorts were HPV positive. In the study by Tang et al., despite multivariate analysis not having been performed, OS and event-free survival became non-significant when only the HPV positive population was considered. On the other hand, in studies that did not consider HPV status, survival outcomes trended towards significantly worse outcomes even at multivariate analysis.

However, differences in reported rates of RLNMs may not be attributable to HPV status alone. As hypothesized by Gross et al., appropriate multimodal therapy may provide survival benefits overcoming any possible negative influence deriving from RLNMs. It is possible that since RLNMs are associated with higher N and T grade it may simply be a marker of advanced disease, although when multivariate analysis was performed, as in almost all the reports included in our metaanalysis, the effect of T and N category on prognosis faded, leaving RLNM as an independent prognostic factor towards worse outcomes. All of these factors may explain why there is still no sharp definition of the effect of RLNMs on prognosis.

This systematic review also highlights the difficulty in diagnosing RLNMs. CT alone is not sufficiently accurate, although the introduction of PET/CT has given rise to a powerful diagnostic tool based on multimodal imaging using CT, PET/CT and MRI. Indeed, PET/CT has been shown to increase the accuracy in diagnosing RLNMs to 86.7% from 60.6% in those imaged with CT and/or MRI, with similar improvements in sensitivity, specificity, positive predictive value, and negative predictive value. Meta-analysis of sensitivity, specificity, and accuracy of imaging studies has not been possible since different studies used different diagnostic tools, moreover different criteria have been used to address RLN as metastatic. Most investigators used a multimodality imaging strategy, which was proven to have a high accuracy in detecting RLN metastasis. This is consistent with the similar prevalence of RLNMs observed in imaging (12%) and histopathological studies (13%).
Some study limitations have to be acknowledged. Firstly, the included studies used a heterogeneous TNM classification, in fact, the time span of the eligible articles vary from AJCC 2nd ed. to AJCC 8th ed. Moreover, clinical records are not uniform across centers in terms of treatment and diagnostic criteria. RLN specimens for histologic examination are difficult to obtain, thus diagnosis relies primarily upon radiological assessment, with various criteria and modalities used by different centers. Reporting of N and T categories were only slightly heterogeneous, being included in the multivariate analysis in five out of six articles. In addition, the inclusion of previous studies was limited by the frequent lack of data required for the meta-analysis: indeed, a number of studies displayed Kaplan–Meyer curves without reporting hazard evaluations, thus considerably diminishing the amount of data available. Finally, few studies reported data regarding RLNMs according to primary tumor subsite, which limited this analysis.

In conclusion (Figure 6), this meta-analysis found that one in 10 OPSCC patients harbored RLNMs, with cancers arising from the posterior pharyngeal wall having the highest prevalence. RLNMs were associated with unfavorable prognostic outcomes. However, these results must be considered with caution since, given the paucity of data available in the literature, this meta-analysis was not based only on individual studies fully adjusted for possible confounding factors.

AUTHOR CONTRIBUTIONS
Giancarlo Tirelli, Nicoletta Gardenal, Enrico Zanelli, Daniele Borsetto, Veronica Philipps, Alberto Vito Marcuzzo, Jonathan Fussey, Jerry Polesel, and Paolo Boscolo-Rizzo contributed equally in conception, design, analysis, and interpretation of data.

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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REFERENCES
1. Doumas S, Fort-Schaale A, Skoulakis C, Moor J, Kanatas A. Current thinking on the management of abnormal retropharyngeal nodes in patients with oral, oropharyngeal, and nasopharyngeal squamous cell carcinoma: a structured review. Br J Oral Maxillofac Surg. 2019;57:515-528.
2. Rouvière H. Lymphatic system of the head and neck. In: Tobias MJ, ed. Anatomy of the Human Lymphatic System. Edwards Brothers; 1938:5-28.
3. He L-J, Xie C, Li Y, et al. Ultrasound-guided fine needle aspiration of retropharyngeal lymph nodes after radiotherapy for nasopharyngeal carcinoma: a novel technique for accurate diagnosis. Cancer Commun. 2018;38:20.
4. Morrissey DD, Talbot JM, Cohen JI, Wax MK, Andersen PE. Accuracy of computed tomography in determining the presence or absence of metastatic retropharyngeal adenopathy. Arch Otolaryngol Head Neck Surg. 2000;126:1478.
5. Li H, Liu X-W, Geng Z-J, Wang D-L, Xie C-M. Diffusion-weighted imaging to differentiate metastatic from non-metastatic retropharyngeal lymph nodes in nasopharyngeal carcinoma. Dentomaxillofac Radiol. 2015;44:20140126.
6. Coskun HH, Ferlito A, Medina JE, et al. Retropharyngeal lymph node metastases in head and neck malignancies. Head Neck. 2011;33:1520-1529.
7. Dirix P, Nuys S, Bussels B, Hermans R, van den Bogaert W. Prognostic influence of retropharyngeal lymph node metastasis in squamous cell carcinoma of the oropharynx. Int J Radiat Oncol Biol Phys. 2006;65:739-744.
8. Gunn GB, Debnam JM, Fuller CD, et al. The impact of radiographic retropharyngeal adenopathy in oropharyngeal cancer: retropharyngeal nodes in oropharynx cancer. Cancer. 2013;119:3162-3169.
9. Shimizu K-I, Inoue H, Saitoh M, et al. Distribution and impact of lymph node metastases in oropharyngeal cancer. Acta Otolaryngol. 2006;126:872-877.
10. Tang C, Komakula S, Chan C, et al. Radiologic assessment of retropharyngeal node involvement in oropharyngeal carcinomas stratified by HPV status. Radiother Oncol. 2013;109:293-296.
11. Billfalk-Kelly A, Yu E, Su J, et al. Radiologic extranodal extension portends worse outcome in cN+ TNM-8 stage I human papillomavirus–mediated oropharyngeal cancer. Int J Radiat Oncol Biol Phys. 2019;104:1017-1027.
12. Lin TA, Garden AS, Elhalawani H, et al. Radiographic retropharyngeal lymph node involvement in HPV-associated oropharyngeal carcinoma: patterns of involvement and impact on patient outcomes. Cancer. 2019;125:1536-1546.
13. Page MJ, McKenzie JE, Bossuyt PM, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. BMJ. 2021;372:n71. doi:10.1136/bmj.n71
14. McGowan J, Sampson M, Salzwedel DM, Cogo E, Foerster V, Lefebvre C. PRESS peer review of electronic search strategies: 2015 guideline statement. J Clin Epidemiol. 2016;75:40-46.
22. Chan S-C, Lin CY, Ng SH, et al. 18F-FDG PET for retropharyngeal nodes in squamous cell carcinoma of oropharynx: incidence, localization, and implications for target volume. *Int J Radiat Oncol Biol Phys*. 2006;65:733-738.

23. Bhattacharai O, Thompson LDR, Schumacher AJ, Iganej S. Radiographic nodal prognostic factors in stage I HPV-related oropharyngeal squamous cell carcinoma. *Head Neck*. 2019;41:398-402. doi:10.1002/hed.25504.

24. Panda S, Thakar A, Kakkar A, et al. Is the retropharyngeal lymph node the first echelon node for carcinoma tonsil? Prospective evaluation and literature review. *Acta Otolaryngol*. 2007;127:123-128.

25. Chan S-C, Lin CY, Ng SH, et al. 18F-FDG PET for retropharyngeal lymph node metastasis in oropharyngeal and hypopharyngeal cancers: impact on diagnosis and prediction analysis. *Nucl Med Commun*. 2010;31:260-265.

26. McLaughlin MP, Mendenhall WM, Mancuso AA, et al. Retropharyngeal adenopathy as a predictor of outcome in squamous cell carcinoma of the head and neck. *Head Neck*. 1995;17:190-198.

27. Kjems J, Gothelf AB, Håkansson K, Specht L, Kristensen CA, Friborg J. Elective nodal irradiation and patterns of failure in head and neck cancer after primary radiation therapy. *Int J Radiat Oncol Biol Phys*. 2016;94:775-782.

28. Nagatsuka M, Hughes RT, Shenker RF, Frizzell BA, Greven KM. Omitting elective irradiation of the contralateral retropharyngeal nodes in oropharyngeal squamous cell carcinoma treated with intensity-modulated radiotherapy. *Cureus*. 2019;11:e3825. doi:10.7759/cureus.3825.

29. Rosen BS, Wilkie JR, Sun Y, et al. CT and FDG-PET radiologic biomarkers in p16+ oropharyngeal squamous cell carcinoma patients treated with definitive chemoradiotherapy. *Radiother Oncol*. 2021;155:174-181.

30. Iyizoba-Ebozue Z, Murray LJ, Arunsingh M, Vaidyanathan S, Scarsbrook AF, Prestwich RJD. Incidence and patterns of retropharyngeal lymph node involvement in oropharyngeal carcinoma. *Radiother Oncol*. 2020;142:92-99.

31. Baxter M, Chan JYK, Mydlarz WK, et al. Retropharyngeal lymph node involvement in human papillomavirus-associated oropharyngeal squamous cell carcinoma: RPLN involvement in HPV-associated OPSCC. *Laryngoscope*. 2015;125:2503-2508.

32. Spector ME, Chinn SB, Bellile E, et al. Exploration for an algorithm for deintensification to exclude the retropharyngeal site from advanced oropharyngeal squamous cell carcinoma treatment. *JAMA Otolaryngol Head Neck Surg*. 2016;142:313-318.

33. Troob S, Givi B, Hodgson M, et al. Transoral robotic retropharyngeal node dissection in oropharyngeal squamous cell carcinoma: patterns of metastasis and functional outcomes. *Head Neck*. 2017;39:1969-1975.

34. Moore EJ, Ebrahimi A, Price DL, Olsen KD. Retropharyngeal lymph node dissection in oropharyngeal cancer treated with transoral robotic surgery: retropharyngeal nodes in oropharyngeal SCC. *Laryngoscope*. 2013;123:1676-1681.

35. Chung E-J, Kim GW, Cho BK, Cho SJ, Yoon DY, Rho YS. Retropharyngeal lymph node metastasis in 54 patients with oropharyngeal squamous cell carcinoma who underwent surgery-based treatment. *Ann Surg Oncol*. 2015;22:3049-3054.

36. Chu HR, Kim JH, Yoon DY, Hwang HS, Rho Y-S. Additional diagnostic value of 18F-FDG PET-CT in detecting retropharyngeal nodal metastases. *Otolaryngol Head Neck Surg*. 2009;141:633-638.

37. Spencer CR, Gay HA, Haughey BH, et al. Eliminating radiotherapy to the contralateral retropharyngeal and high level II lymph nodes in head and neck squamous cell carcinoma is safe and improves quality of life. *Cancer*. 2014;120:3994-4002.

38. Nguyen NP, Vock J, Vinh-Hung V, et al. Effectiveness of prophylactic retropharyngeal lymph node irradiation in patients with locally advanced head and neck cancer. *BMC Cancer*. 2012;12:253.

39. Kato H, Kanematsu M, Watanabe H, Mizuta K, Aoki M. Metastatic retropharyngeal lymph node metastases: comparison of CT and MR imaging for diagnostic accuracy. *Eur J Radiol*. 2014;83:1157-1162.

40. Giivi B, Troob SH, Stott W, Cordeiro T, Andersen PE, Gross ND. Transoral robotic retropharyngeal node dissection: Transoral robotic retropharyngeal node dissection. *Head Neck*. 2016;38:E981-E986.

41. Kim HJ, Choi KY, Lee SH, et al. The value of CT, MRI, and PET-CT in detecting retropharyngeal lymph node metastasis of head and neck squamous cell carcinoma. *BMC Med Imaging*. 2020;20:88.

42. Wong YK, Novotny GM. Retropharyngeal space: a review of anatomy, pathway, and clinical presentation. *J Otolaryngol*. 1978;7:528-536.

43. Boscolo-Rizzo P, Pawlita M, Holzinger D. From HPV-positive towards HPV-driven oropharyngeal squamous cell carcinomas. *Cancer Treat Rev*. 2016;42:24-29.
45. de Martel C, Plummer M, Vignat J, Franceschi S. Worldwide burden of cancer attributable to HPV by site, country and HPV type. *Int J Cancer*. 2017;141:664-670.

46. Silfverschiöld M, Sjövall J, Wennerberg J, Östensson E, Greiff L. Societal cost of oropharyngeal cancer by human papillomavirus status, cancer stage, and subsite. *PLoS One*. 2019;14:e0220534.

47. Zhang G-Y, Liu LZ, Wei WH, Deng YM, Li YZ, Liu XW. Radiologic criteria of retropharyngeal lymph node metastasis in nasopharyngeal carcinoma treated with radiation therapy. *Radiology*. 2010;255:605-612.

48. Mancuso AA, Harnsberger HR, Muraki AS, Stevens MH. Computed tomography of cervical and retropharyngeal lymph nodes: normal anatomy, variants of normal, and applications in staging head and neck cancer. Part I: normal anatomy. *Radiology*. 1983;148:709-714.

49. Wang XS, Yan C, Hu CS, et al. Study of the medial group retropharyngeal node metastasis from nasopharyngeal carcinoma based on 3100 newly diagnosed cases. *Oral Oncol*. 2014;50:1109-1113.

50. Gross ND, Ellingson TW, Wax MK, Cohen JI, Andersen PE. Impact of retropharyngeal lymph node metastasis in head and neck squamous cell carcinoma. *Arch Otolaryngol Head Neck Surg*. 2004;130:169-173.

**SUPPORTING INFORMATION**

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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