

Therapeutic Options in Unresectable Oral Squamous Cell Carcinoma: A Systematic Review

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Purpose: This review describes the current scientific evidence of therapeutic options in unresectable oral squamous cell carcinoma.

Methods: This study followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA). We searched MEDLINE (Via PubMed) to identify studies assessing treatments for unresectable oral squamous cell carcinoma. The methodological quality assessment of the included studies was performed using the Joanna Briggs Institute (JBI) checklist tool. The evidence was organized and presented using tables and narrative synthesis.

Results: Thirty-three studies met the eligibility criteria. Most studies had an observational design. The sample size varied from 16 to 916 participants. The methodology quality of the included studies ranged from 2.5 to 10 using the JBI tool. Overall, the optimal treatment of patients with unresectable oral cancer is challenging, so there is a sprinkling of studies assessing a variety of therapeutic options, such as radiotherapy, chemotherapy, concurrent chemoradiotherapy, immunotherapy, targeted therapy plus chemotherapy or radiotherapy, and gene therapy plus chemotherapy.

Conclusion: There is lacking evidence about the benefits of some therapeutic options for unresectable oral squamous cell carcinoma. Overall, these patients can be treated using a multimodal approach such as concurrent chemoradiotherapy or induction chemotherapy followed by chemoradiotherapy, which have shown good clinical outcomes. However, other options could be considered depending on the assessment of risk/benefits, tumor extension, and patient values and preferences.

Keywords: mouth neoplasms, unresectable oral cancer, oral squamous cell carcinoma, treatment, therapy

Introduction

Oral cancer is a health issue globally. It fully meets the criteria to be considered a public health problem such as high mortality rate, the impact of the condition on an individual level, and impact on wider society.¹ To illustrate, it has been reported that around 650,000 new cases are diagnosed annually; although it represents just 2% of the tumor incidence worldwide, the major reason for concern is its high mortality rate of around 50%.² Regarding the region-specific incidence age-standardized rates by sex for oral cancer in 2018, there is a higher incidence of this oral disease for men than women in all countries, being Melanesia, South Central Asia, Australia/New Zealand, Eastern Europe, Western Europe, and Northern America, the regions with the highest incidence.³ Likewise, this malignancy accounts for over 140,000 deaths per year and its age-standardized mortality rates can vary depending on geographical settings.³

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This disease stands amongst the six most common cancers worldwide, and about 40% of head and neck tumors are oral squamous cell carcinomas,⁴ which is the most common type of mouth cancers. Moreover, oral cancer has a substantial financial burden on the public healthcare system and produces both physical and psychological impacts on people suffering from this disease such as swallowing function, speech difficulties, and self-image concerns.⁵

Currently, there are important technological advances in cancer management and oncology research, but oral carcinoma still has a poor prognosis and its treatment involves usually severe physical and psychological after-effects.^{6,7} Amongst the main therapeutic options for oral cancer are surgery, radiotherapy (RT), and chemotherapy (CT),^{8,9} commonly locoregionally oral neoplasms are treated by surgical approach considered as the gold standard treatment,¹⁰ while those advanced or aggressive oral tumors with high probabilities of relapse after definitive treatment with surgery or RT are treated using a multimodal approach that combines surgery and pre/postoperative RT or CT.^{11,12} However, for those patients with unresectable disease, when the surgical approach is not feasible because of the extension of lesion, surgery is expected to result in poor functional outcomes, patients' poor status or patient values and preferences, the optimal therapeutic options are largely unclear.¹³

Likewise, it has been reported that the evidence on the benefits of therapeutic interventions for unresectable oral cancer is lacking.¹⁴ Thus, this review aimed to describe the current scientific evidence about therapeutic options in unresectable oral squamous cell carcinoma.

Methods

This review followed the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA).¹⁵ The aim and all methods used in this study were specified a priori in a protocol (available on request).

Search Strategy

We searched MEDLINE (via PubMed) to identify relevant studies assessing any therapeutic options in unresectable oral squamous cell carcinoma. We used MeSH descriptor, free text, and treasure terms such as “mouth neoplasms”, “oral cancer”, “oral carcinoma”, “buccal carcinoma”, “unresectable”, “advanced”, “inoperable”, “therapeutics”, “treatment”, “management” and “therapy” ([Supplementary Material 1](#)). There were no language restrictions, and the

search was filtered by the last 10 years in order to include the most updated evidence into the review. The last research was carried out on April 12, 2021. Also, a cited reference search was conducted.

Selection of Studies

Studies of different epidemiological designs (randomized controlled trial (RCT), clinical trial, cohort, and case-control studies) and published after 2010 were included. They had to evaluate any therapeutic option in individuals diagnosed with primary unresectable oral squamous cell carcinoma, defined as patients with advanced mouth neoplasm, no evidence of distant metastases, and unsuitable to surgical treatments for any reason. If a study was reported in more than one publication, only the most recent version was considered. Conversely, studies only on diagnosis and prognosis were excluded. Likewise, studies only focused on interventions before or after surgical treatments were excluded.

We managed all retrieved records using the Rayyan¹⁶ website. Initially, the duplicates automatically were removed, then at least two appraisers independently screened all titles/abstracts to exclude unrelated studies. Subsequently, full articles were obtained for a final decision. Detailed reasons for exclusion of any study considered relevant were clearly stated.

Methodological Critical Appraisal and Data Extraction

We critically appraised all included studies using the Joanna Briggs Institute (JBI)¹⁷ checklist for each study design included in this review. Overall, these checklists rate the quality of different factors such as selection, measurement, and comparability of groups. This tool gives a score for RCT (maximum of 13), clinical trial (maximum of 9), cohort (maximum of 11), and case-control (maximum of 10). There is no cut-off point, so a higher score indicates better methodology quality of the study.

At least two reviewers independently conducted all processes of study selection, methodological critical appraisal, and data extraction. If there were any disagreements, they were resolved by consensus, and when necessary, an additional reviewer participated in the discussion until an agreement was reached. We extracted data about general characteristics of the study (authors, publication year, design, country, aim, sample size, sample features, and risk factors reported) and characteristics of the therapeutic intervention (type, dose, comparison, outcomes, and conclusion

about its effectiveness). The evidence was organized and presented using tables and narrative synthesis.

Results

Selection of Studies

1887 records were identified after removing duplicates. After the title and abstract screening, 68 articles were obtained for final full-text review; 33 studies^{18–50} met the eligibility criteria (Figure 1). The list of excluded studies along with exclusion rationale is available in [Supplementary Material 2](#).

Characteristics of the Included Studies

All selected studies^{18–50} were published in English language between 2011 and 2020. There were 15 retrospective cohort studies,^{22,24–26,28,34,36–42,47,50} eight were prospective cohort studies,^{19,23,29,32,33,35,44,46} six were RCTs,^{18,21,27,30,31,43} four

were clinical trials,^{20,45,48,49} and none was case-control study. The sample size varied from 16²⁷ to 976⁴² participants. There were eight studies from India,^{19,21,34,36–38,43,46} six from Japan,^{26,27,32,33,40,45} four from Taiwan,^{20,42,47,48} three from the United States,^{25,35,41} two from China,^{30,31} two from Pakistan,^{28,49} while the other studies were one from each of the following: Canada,⁵⁰ Hungary,⁴⁴ Italy,²³ Iran,²⁹ Spain,³⁹ The Netherlands,²⁴ Thailand,²² and Ukraine.¹⁸ Only 14 studies^{20,22,24,26,28,34–38,41–43,46} reported oral cancer risk factors, such as smoking, alcohol consumption and betel chewing. All general characteristics of the selected studies are presented in [Supplementary Material 3](#).

The Methodological Quality of the Included Studies

Using the JBI's critical appraisal checklist tool, the mean score was 7.5 ± 2.7 (range from 2.5 to 10), 7.5

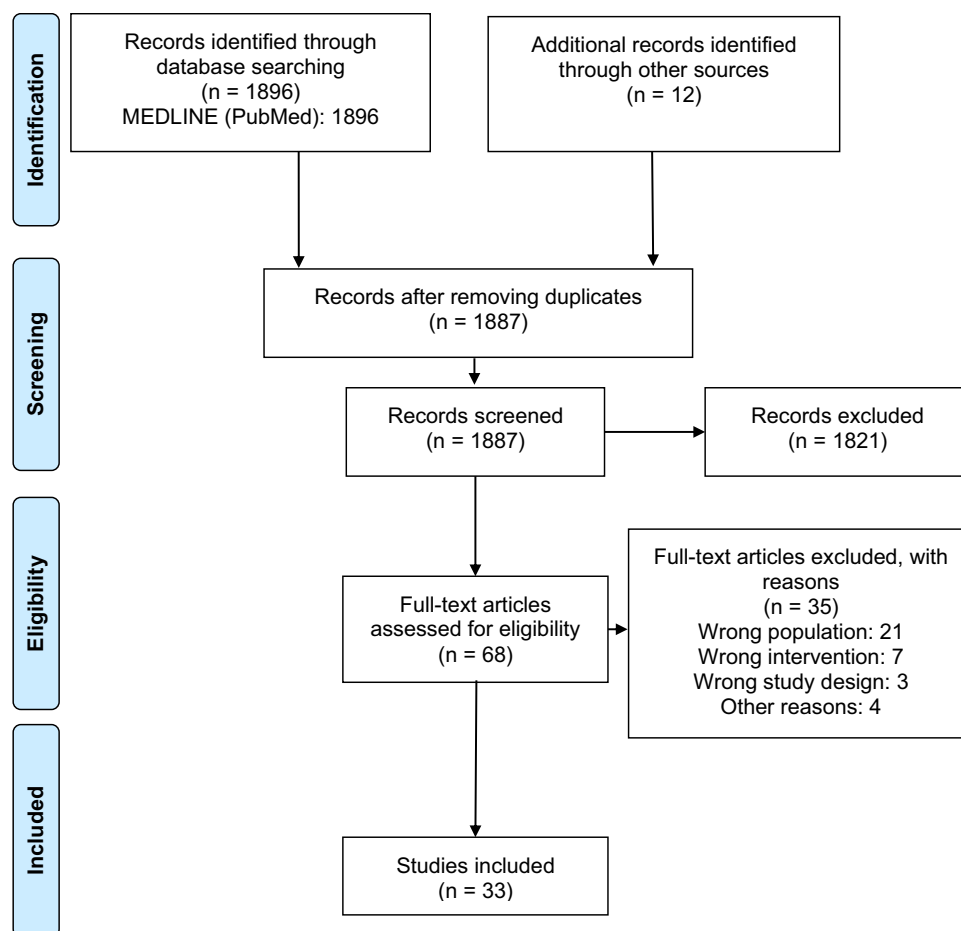


Figure 1 PRISMA flowchart describing the selection of studies.

Notes: Adapted from Page MJ, McKenzie JE, Bossuyt PM, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. *BMJ*. 2021;372:n71.¹⁵ Creative Commons Attribution (CC BY 4.0) license (<https://creativecommons.org/licenses/by/4.0/legalcode>).

Author and year	Study design	JBI'S TOOL													Overall
		Item 1	Item 2	Item 3	Item 4	Item 5	Item 6	Item 7	Item 8	Item 9	Item 10	Item 11	Item 12	Item 13	
Bazyka 2019	RCT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	8
Chhatui 2015	RCT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	8
Hino 2011	RCT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	2.5	
Li 2014	RCT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	10	
Meng 2014	RCT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	6	
Singh 2013	RCT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	9	
Chang 2017	CT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	-	-	7	
Takayama 2016	CT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	-	-	7.5	
Yen 2019	CT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	-	-	7	
Zaidi 2020	CT	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	-	-	7.5	
Biswas 2019	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7.5	
Chitapanarux 2017	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7.5	
Donato 2013	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7	
Elbers 2017	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7.5	
Foster 2018	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7.5	
Hayashi 2019	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	6	
Iqbal 2015	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	6.5	
Larizadeh 2012	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5	
Murakami 2017	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5	
Oyama 2020	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	6	
Patil 2014	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5.5	
Pederson 2011	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5.5	
Rewadkar 2017	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5	
Rudresha 2017a	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	3.5	
Rudresha 2017b	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	3.5	
Santos 2017	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5	
Sato 2019	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	5.5	
Scher 2015	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	6.5	
Shia 2020	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	10	
Takácsi-Nagy 2013	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7.5	
Vedasoundaram 2020	P. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7.5	
Wu 2014	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	7	
Zhang 2013	R. Cohort	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes	-	-	8	

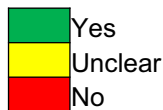


Figure 2 The methodological quality of the included studies by design.
Abbreviations: CT, clinical trial; P, prospective; R, retrospective; RCT, randomized controlled trial.

± 0.3 (range from 7 to 10), and 6.3 ± 1.5 (range from 3.5 to 10) for RCTs, clinical trials, and cohort studies, respectively. The results in detail for each study are shown in [Figure 2](#).

Therapeutic Options in Unresectable Oral Squamous Cell Carcinoma

Overall, included studies assessed a variety of therapeutic options in unresectable oral squamous cell carcinoma.

Twenty-seven studies^{18,19,21–29,32–42,44–46,49,50} assessed different therapies that in some way combined the use of CT and RT, three studies^{42,43,50} evaluated RT alone, two studies^{31,47} assessed CT alone, four studies^{20,31,43,48} examined the combination of targeted therapy plus CT or RT, and only one study for each of the interventions of immunotherapy²¹ and gene therapy.³⁰ The main outcomes reported were overall survival (OS), locoregional control (LRC), progression-free survival (PFS), disease-free survival (DFS), complete response (CR), and cancer-specific survival (CSS). All these studies used different approaches, doses, outcomes, etc. Thus, the scientific evidence about the effectiveness of the main treatments for unresectable oral cancer is summarized by groups as follows:

Concurrent Chemoradiotherapy

Concurrent chemoradiotherapy (CCRT) implies the use of CT delivered simultaneously with RT. This approach has multiple advantages, such as the improvement of the chances of LRC, OS rates, and organ-preserving intent. Moreover, CT given as part of CCRT could act systemically and possibly prevent distant metastases. Likewise, this therapeutic option improves function and cosmetic outcomes compared with surgical approaches. Fourteen studies^{19,21,24–28,32,33,35,40–42,50} evaluated the use of CCRT, two of them were clinical trials,^{25,27} one was a RCT,²¹ and the others were observational studies.^{19,24,26,28,32,33,35,40–42,50} The number of participants for each intervention group varied from 16²⁷ to 256⁴² people. Twelve studies^{19,21,24–26,28,32,33,35,40,42,50} only focused on patients with clinical stage III/IV oral cancer, whereas two studies^{27,41} also included other clinical stages. The main outcomes assessed were OS and LRC, which were evaluated in 10^{24–28,32,33,35,40,41,50} and seven^{24–26,33,35,40,41} studies, respectively (Table 1).

Ten^{19,21,24–28,33,35,40} out of 14^{19,21,24–28,32,33,35,40–42,50} studies suggested that CCRT is a therapeutic option in people with unresectable oral cancer, so this treatment should be considered when surgery is not feasible. Among those studies in favor of CCRT, the 5-year OS and LRC rates ranged from 22%²⁴ to 76%,³⁵ and 49%²⁴ to 90%,³⁵ respectively. Overall, cisplatin remains as the main chemotherapeutic drug used in CCRT for unresectable oral cancer treatment; three studies^{19,21,24} used cisplatin plus RT up to 70 Gy, one study⁵⁰ used cisplatin or carboplatin plus RT up to 70 Gy, while two studies^{27,32} used the S-1 (Taiho Pharmaceutical, Tokyo, Japan) plus

RT. Biswas¹⁹ evaluated the use of cisplatin plus RT 70 Gy between younger (<40 years) and older (>40 years) adults, concluding that there is a similar overall response (63.5% vs 65.9%) between these groups; thus, CCRT can be used both for young as the elderly population.

According to the use of S-1, Murakami³² reported a 3-year OS and PFS of 37% and 27%, respectively, employing S-1 plus RT up to 70 Gy. Similarly, Hino²⁷ assessed the use of S-1 plus RT 30 Gy and reported a median OS and PFS of 42.5 and 6.3 months, respectively, concluding that CCRT with S-1 is an effective treatment that can be safely conducted with minimal burden on patients. However, it is useful to highlight that that statement should be put into context because there are a small number of studies assessing this approach.

Regarding those studies that used 2–3 drug regimen CT, cisplatin was combined with docetaxel,^{26,40} carboplatin,⁴¹ gemcitabine,²⁸ 5-fluorouracil (5-FU),⁴¹ and paclitaxel.⁴¹ Hayashi²⁶ evaluated the use of intra-arterial CT (total, 150 mg/m² cisplatin and 60 mg/m² docetaxel) with daily conventional RT (total, 60 Gy/30 fr) for 6 weeks. At a median follow-up of 40 months, the 3-year OS and LRC were 64.3% and 84.3%, respectively. Similarly, Sato⁴⁰ evaluated the use of cisplatin combined with docetaxel plus RT up to 66 Gy, reporting a 3-year and 5-year OS and LRC of 52.9%, 33.0%, 50.9%, and 50.9%, respectively. Conversely, some studies^{41,50} have reported poor OS and LRC rates using CCRT in patients with unresectable oral cancer. To illustrate, Zhang⁵⁰ reported a 5-year OS of just 10% using cisplatin or carboplatin plus RT up to 70 Gy. Similarly, Scher⁴¹ reported a 5-year OS of just 15% using cisplatin combined with carboplatin and 5-FU or paclitaxel plus RT up to 70 Gy. Likewise, Shia⁴² stated that although CCRT has benefits compared to non-treatment, there are no survival differences between the use of RT alone and CCRT.

Overall, CCRT could be considered as the main therapeutic option in unresectable oral squamous cell cancer because there is scientific evidence supporting its use. However, many factors should be taken into account in order to improve the clinical outcomes of these patients, who often are considered beyond cure.

Induction Chemotherapy Followed by Chemoradiotherapy

Induction chemotherapy (ICT) refers to CT given before the definitive treatment, in this case, that treatment was chemoradiotherapy (CRT). Among the benefits of ICT are

Table I Summary of Evidence on Concurrent Chemoradiotherapy

Author and Year	Clinical Stages	Interventions	Follow-Up	Outcomes
Biswas 2019 ¹⁹	III IV	Arm 1 (n=20): Age <40 Y, CCRT (cisplatin + RT 70 Gy) Arm 2 (n=26): Age >40 Y, CCRT (cisplatin + RT 70 Gy)	11.1 M	OR: Arm 1, 63.5%; Arm 2, 65.9%
Chhatui 2015 ²¹	III IV	(n=25): CCRT (cisplatin + RT 70 Gy)	15 M	DFS: 69% CR: 52%
Elbers 2017 ²⁴	III IV	(n=100): CCRT (cisplatin + RT up to 70 Gy)	13 M	OS [5 Y]: 22% LRC [5 Y]: 49% DFS [5 Y]: 22% DSS [5 Y]: 39%
Foster 2018 ²⁵	III IV	(n=140): CCRT (CT + RT up to 75 Gy)	5.7 Y	OS [5 Y]: 63.2% LRC [5 Y]: 78.6% PFS [5 Y]: 58.7% DC [5 Y]: 87.2%
Hayashi 2019 ²⁶	III IV	(n=46): CCRT (docetaxel + cisplatin + RT up to 60 Gy)	40 M	OS [3Y]: 64.3% LRC [3Y]: 84.3%
Hino 2011 ²⁷	II III IV	(n=16): CCRT (S-I+ RT 30 Gy)	7.4 M	OR: 87.5% OS [Median]: 42.5 M PFS [Median]: 6.3 M
Iqbal 2015 ²⁸	III IV	(n=63): CCRT (gemcitabine + cisplatin + RT 55 Gy)	60 M	OS [5 Y]: 30% PFS [5 Y]: 49% DFS [5 Y]: 30%
Murakami 2017 ³²	III IV	(n=47): CCRT (S-I+ RT up to 70 Gy)	22 M	OS [3 Y]: 37% PFS [3 Y]: 27%
Oyama 2020 ³³	III IV	(n=37): CCRT (docetaxel + nedaplatin + RT up to 70 Gy)	40 M	OS [5 Y]: 64.5% DF [5 Y]: 59.9% LC [5 Y]: 85.5%
Pederson 2011 ³⁵	III IV	(n=20): CCRT (5-FU + hydroxyurea + RT up to 75 Gy)	60 M	OS [2 Y]: 76% OS [5 Y]: 76% DFS [2 Y]: 71% DFS [5 Y]: 71% LRC [2 Y]: 90% LRC [5 Y]: 90%
Sato 2019 ⁴⁰	III IV	(n=17): CCRT (docetaxel+ cisplatin + RT up to 66 Gy)	41 M	OS [3 Y]: 52.9% OS [5 Y]: 33.0% LRC [3 Y]: 50.9% LRC [5 Y]: 50.9%
Scher 2015 ⁴¹	I II III IV	(n=73): CCRT (cisplatin + carboplatin + 5-FU or paclitaxel + RT up to 70 Gy)	73.1 M	OS [5 Y]: 15% LRC [5 Y]: 37.4% DC [5 Y]: 70.2%
Shia 2020 ⁴²	III IV	Arm 1 (n=256): CCRT Arm 2 (n=227): Non-treatment	15 M	Death risk: Arm 1, 1 (Ref); Arm 2, 1.60 (1.30–1.97)
Zhang 2013 ⁵⁰	III IV	(n=10): CCRT (cisplatin or carboplatin + RT up to 70Gy)	3.52 Y	OS [2 Y]: 20% OS [5 Y]: 10% DSS [2 Y]: 26% DSS [5 Y]: 13% DFS [2 Y]: 13%

Abbreviations: CCRT, concurrent chemoradiotherapy; CR, complete response; CT, chemotherapy; DC, distant control; DF, disease free rate; DFS, disease-free survival; DSS, disease specific survival; FU, fluorouracil; LC, local control; LRC, locoregional control; M, months; OR, overall response, OS, overall survival; PFS, progression-free survival; RT, radiotherapy; Y, years.

to shrink the tumor, decrease the chances of distant metastases, increase the chances of organ preservation, and improving the outcomes such as OS and PFS. Seven studies^{29,34,36–38,45,49} assessed this approach, three studies^{29,45,49} were clinical trials and the rest of studies^{34,36–38} employed observational designs. The number of participants by group varied from 16²⁹ to 167³⁴ people. Three studies^{34,37,38} only involved patients with stage IV oral cancer, three studies^{29,45,49} recruited patients with clinical stages III/IV, and one study³⁶ did not report it. All studies^{29,34,36–38,45,49} used the 2 or 3 drug regimens CT, and the main outcome reported was OS (Table 2).

In a clinical trial by Larizadeh,²⁹ patients with locoregionally advanced oral cancer were enrolled. ICT comprise 3 cycles of cisplatin and 5-FU with or without docetaxel. The overall response rate after ICT was 68.4%, and OS rates after 2 and 3 years were 38% and 26%, respectively. This author concluded that the outcome of patients with unresectable oral cancer is poor, so the benefits of the use of this therapeutic intervention for these patients are unclear. Likewise, Patil³⁴ reported after 2 years an OS rate of 20% and an LRC of 15.0% for patients who underwent ICT with multiple drugs (paclitaxel or docetaxel plus cisplatin or carboplatin with or without 5-FU) followed by CCRT (cisplatin plus RT up to 70 Gy).

In terms of CR, two studies^{36,49} reported this outcome. Rewadkar³⁶ reported a CR of 84% using bleomycin, methotrexate, and cisplatin on day 1 and repeated at an interval of 21 days during 3 cycles; then, participants were treated with CCRT using cisplatin infusion. That study stated that this approach can be superior to other treatments such as ICT plus RT alone. Conversely, Zaidi⁴⁹ conducted an open-label, non-randomized trial and reported a complete response of just 10.5% using ICT with docetaxel plus cisplatin followed by cisplatin plus concurrent RT up to 60 Gy.

Takayama⁴⁵ evaluated a complex approach using ICT followed by CCRT in 33 patients with stage III–IVB tongue cancer. Briefly, after two systemic CT courses and whole-neck irradiation using 36 Gy in 20 fractions, CCRT was used comprising proton beam therapy with weekly retrograde intra-arterial CT by continuous infusion of cisplatin with sodium thiosulfate. At a median follow-up of 43 months, the 3-year OS, PFS, local control (LC) rates were 87.0%, 74.1% and 86.6%, respectively. Overall, although ICT followed by CCRT can be used as a therapeutic option in unresectable oral cancer, its potential benefits still are controversial.

Induction Chemotherapy Followed by Radiotherapy

Four studies^{18,36–38} assessed this approach, one of them was an RCT,¹⁸ and the others were retrospective cohort studies,^{36–38} the number of participants by intervention group varied from 8³⁷ to 99¹⁸ people. Two studies^{37,38} included patients with clinical stage IV oral cancer, while two studies^{18,36} did not report it. All studies^{18,36–38} used the 2 or 3 drug regimen involving chemotherapeutic drugs, such as cisplatin, carboplatin, 5-FU, bleomycin, polyplatinene, methotrexate, paclitaxel, and docetaxel. The main outcome was OS, which was evaluated in three studies^{18,37,38} (Table 3).

One RCT¹⁸ have examined the role the ICT followed by RT in patients with locally advanced oral cavity, suggesting in terms of OS, that the use of ICT with polyplatinene plus 5-FU followed by RT 70 Gy is better than using ICT with cisplatin plus 1,5-FU followed by RT 70 Gy (36 months OS= 58% vs 29%). Two observational studies^{37,38} reported only a median OS of 7.3 and 8.5 months for patients treated with ICT followed by RT, which suggests that other therapeutic options should be considered in order to improve the disease-related outcomes.

Radiotherapy with/without Chemotherapy

All studies assessing RT alone or RT with or without CT were included in this group. Overall, five studies^{22,23,42,43,50} assessed this approach, three studies^{42,43,50} assessed RT alone and two studies^{22,23} assessed RT with or without CT. There was an RCT,⁴³ a clinical trial²³ and three observational studies.^{22,42,50} The number of participants for each group ranged from 9²³ to 315²² people. All studies^{22,23,42,43,50} recruited patients with stage III/IV oral cancer (Table 4).

Among the studies assessing RT alone, one study⁵⁰ reported a 2-year and 5-year OS of just 18% and 10%, respectively. One study⁴³ reported a CR of 33%, whereas another study⁴² reported that the death risk for those receiving non-treatment was approximately 60% higher than those receiving RT alone. Among those evaluating the use of RT with or without CT, Donato²³ reported a 2-year OS and DFS of 55.6% and 75%, respectively. Conversely, Chitapanarux²² reported a 5-year OS of just 15.9%. Overall, there is no strong evidence to support this approach to treat patients suffering from unresectable oral cancer.

Table 2 Summary of Evidence on Induction Chemotherapy Followed by Chemoradiotherapy

Author and Year	Clinical Stages	Interventions	Follow-Up	Outcomes
Larizadeh 2012 ²⁹	III IV	Arm 1 (n=16): ICT with cisplatin + 5-FU followed by CCRT (cisplatin + RT) Arm 2 (n=41): ICT with cisplatin + 5-FU + docetaxel followed by CCRT (cisplatin + RT)	32 M	OR: 68.4% OS [2 Y]: 38% OS [3 Y]: 26% OS [Mean]: Arm 1, 17.1 M; Arm 2, 27.9
Patil 2014 ³⁴	IV	(n=167): ICT with paclitaxel or docetaxel + cisplatin or carboplatin ± 5-FU followed by CCRT (cisplatin + RT up to 70 Gy)	28 M	OS [2 Y]: 20% LRC [2 Y]: 15.0%
Rewadkar 2017 ³⁶	NR	(n=25): ICT with bleomycin + methotrexate + cisplatin followed by CCRT (cisplatin + RT)	NR	CR: 84%
Rudresha 2017a ³⁷	IV	(n=27): ICT with paclitaxel + carboplatin followed by CCRT	NR	OS [Median]: 11.8 M
Rudresha 2017b ³⁸	IV	(n=44): ICT with docetaxel + cisplatin + 5-FU or paclitaxel + carboplatin followed by CCRT	NR	OS [Median]: 9.4 M
Takayama 2016 ⁴⁵	III IV	(n=33): ICT with 5-FU + nedaplatin + RT 36 Gy followed by CCRT (5-FU + nedaplatin + RT 39.6 Gy + cisplatin)	43 M	OS [3 Y]: 87.0% PFS [3 Y]: 74.1% LC [3 Y]: 86.6%
Zaidi 2020 ⁴⁹	III IV	(n=35): ICT with docetaxel + cisplatin followed by CCRT (cisplatin + RT up to 60 Gy)	4 M	OR: 78.8% CR: 10.5% PR: 68.4%

Abbreviations: CCRT, concurrent chemoradiotherapy; CR, complete response; FU, fluorouracil; ICT, induction chemotherapy; LC, local control; LRC, locoregional control; M, months; NR, not reported; OR, overall response; OS, overall survival; PFS, progression-free survival; PR, partial response; RT, radiotherapy; Y, years.

Table 3 Summary of Evidence on Induction Chemotherapy Followed by Radiotherapy

Author and Year	Clinical Stages	Interventions	Follow-Up	Outcomes
Bazyka 2019 ¹⁸	NR	Arm 1 (n=99): ICT with cisplatin + 1.5-FU followed by RT up to 70 Gy Arm 2 (n=43): ICT with polyplatin + 5-FU followed by RT up to 70 Gy	NR	OS [36 M]: Arm 1, 29%*; Arm 2, 58%*
Rewadkar 2017 ³⁶	NR	(n=25): ICT with bleomycin + methotrexate + cisplatin followed by RT up to 70 Gy	NR	CR: 60%
Rudresha 2017a ³⁷	IV	(n=8): ICT with paclitaxel + carboplatin followed by RT	NR	OS [Median]: 8.5 M
Rudresha 2017b ³⁸	IV	(n=11): ICT with docetaxel + cisplatin + 5-FU or paclitaxel + carboplatin followed by RT	NR	OS [Median]: 7.3 M

Note: *Data extracted from a figure.

Abbreviations: CR, complete response; FU, fluorouracil; ICT, induction chemotherapy; M, months; NR, not reported; OS, overall survival; RT, radiotherapy.

Radiotherapy with or without Chemotherapy Followed by Brachytherapy

Three studies^{39,44,46} focused on the use of external beam RT (EBRT) with or without concurrent CT followed by brachytherapy (BT), two of them were clinical trials^{44,46}

and one was a retrospective cohort study.³⁹ The number of participants for each intervention group varied from 24³⁹ to 60⁴⁴ people. One study³⁹ included patients with clinical stage III/IV oral cancer, one study⁴⁶ only included patients with stage III oral cancer, and another study⁴⁴ also

Table 4 Summary of Evidence on Radiotherapy with/without Chemotherapy

Author and Year	Clinical Stages	Interventions	Follow-Up	Outcomes
Chitapanarux 2017 ²²	III IV	(n=315): RT 60–70 Gy ± CT	11 M	OS [5 Y]: 15.9% OS [10 Y]: 12.9%
Donato 2013 ²³	III IV	(n=9): RT up to 70 Gy ± CT	24 M	OR: 77.8% OS [2 Y]: 55.6% DFS [2 Y]: 75%
Shia 2020 ⁴²	III IV	Arm 1 (n=237): RT Arm 2 (n=227): Non-treatment	15 M	Death risk: Arm 1, 1.06 (0.87, 1.31); Arm 2, 1.60 (1.30, 1.97)
Singh 2013 ⁴³	III IV	(n=30): RT 70 Gy	20 M	CR: 33%
Zhang 2013 ⁵⁰	III IV	(n=28): RT up to 80Gy	3.52 Y	OS [2 Y]: 18% OS [5 Y]: 10% DSS [2 Y]: 21% DSS [5 Y]: 21% DFS [2 Y]: 21% DFS [5 Y]: 21%

Abbreviations: CR, complete response; CT, chemotherapy; DFS, disease-free survival; DSS, disease specific survival; M, months; OR, overall response, OS, overall survival; RT, radiotherapy; Y, years.

included other clinical stages. The main outcomes assessed were OS, LRC, and CSS, which were evaluated in two^{39,44} out of three^{39,44,46} studies (Table 5).

In a nonrandomized clinical trial by Takácsi-Nagy,⁴⁴ a high-dose-rate (HDR) BT boost with a mean dose of 17 Gy was delivered after 50–70 Gy locoregional EBRT. Moreover, around 30% of participants also received concurrent CT with cisplatin, reporting that the 5-year rate of LC, LRC, OS, and CSS was 57%, 50%, 47%, and 61%, respectively. Furthermore, OS was significantly better in patients receiving concurrent CT (69% vs 39%; $p=0.005$). Santos³⁹ assessed the use of EBRT up to 60 Gy plus concurrent CT followed by HDR-BT up to 24 Gy with a median follow-up of 44 months, reporting that the 4-year OS and LRC rate was 68% and 80%, respectively. Similarly, Vedesoundaram⁴⁶ assessed the use of EBRT 50 Gy plus CT with cisplatin followed by HDR-BT 21 Gy and reported a CR of 77.2%. Overall, this approach seems to be effective to treat patients with unresectable oral cancer, but more research about it is needed.

Chemotherapy

Two studies^{31,47} evaluated the use of CT, one of them was a controlled clinical trial,⁴⁷ and another was an

observational retrospective study.³¹ One study⁴⁷ included 21 participants in the intervention group, whereas the other study³¹ included just 8 people in the intervention group. All two studies^{31,47} focused on patients with clinical stage III/IV oral cancer.

Wu⁴⁷ aimed to assess the efficacy of intra-arterial infusion CT for patients with locally advanced oral cancer. Patients received continuously an infusion of methotrexate (50 mg/day) into the external carotid artery for 8 days, followed by a weekly intra-arterial bolus of 25 mg methotrexate for 10 weeks. Overall, CR and the partial response rate were 62% and 33%, respectively. At a median follow-up of 69 months, the 1-year, 3-year, and 5-year OS rates were 80%, 71%, and 64%, respectively. Similarly, Meng³¹ evaluated the use of the docetaxel-cisplatin-FU regimen, showing a response rate of 37.5% and a disease control rate of 62.5%. However, few studies are assessing the use of CT alone for unresectable oral cancer, so its effectiveness should be determined.

Targeted Therapy, Immunotherapy, and Gene Therapy

Six studies^{20,21,30,31,43,48} were included in this group, four^{20,31,43,48} of them assessed therapeutic options for unresectable oral cancer including at least a drug

Table 5 Summary of Evidence on Radiotherapy with or without Chemotherapy Followed by Brachytherapy

Author and Year	Clinical Stages	Interventions	Follow-Up	Outcomes
Santos 2017 ³⁹	III IV	(n=24): EBRT up to 60 Gy + CT followed by HDR-BT	44 M	OS [3 Y]: 68% OS [4 Y]: 68% CSS [3 Y]: 75% CSS [4 Y]: 68% LC [3 Y]: 80% LC [4 Y]: 80% LRC [3 Y]: 84% LRC [4 Y]: 76% DFS [3 Y]: 62% DFS [4 Y]: 48%
Takácsi-Nagy 2013 ⁴⁴	I II III IV	(n=60): EBRT up to 70 Gy ± CT with cisplatin followed by HDR-BT up to 30 Gy	121 M	OS [5Y]: 47% LRC [5Y]: 50% LC [5Y]: 57% CSS [5Y]: 61%
Vedasoundaram 2020 ⁴⁶	III	(n=57): EBRT 50 Gy + CT with cisplatin followed by HDR-BT up to 21 Gy	60 M	CR: 77.2%

Abbreviations: CT, chemoradiotherapy; CR, complete response; CSS, cancer specific survival; DFS, disease-free survival; EBRT, external beam radiotherapy; HDR-BT, high dose rate brachytherapy; LC, local control; LRC, locoregional control; M, months; OS, overall survival; RT, radiotherapy; Y, years.

considered as targeted therapy, one study²¹ assessed the use of immunotherapy plus ICT followed by CRT, and one study³⁰ evaluated the use of gene therapy plus CT. There were three RCTs^{21,30,43} and three clinical trials.^{20,31,48} The number of participants for each intervention group varied from 9³¹ to 43²⁰ people. Two studies^{20,48} only focused on patients with clinical stage IV oral cancer, whereas four studies^{21,30,31,43} focused on patients with stages III/IV. The main outcome assessed was OS, which was assessed in three^{20,30,48} out of six^{20,21,30,31,43,48} studies (Table 6).

Two studies^{20,48} included the use of cetuximab in their treatment regimen. An open-label Phase II trial by Chang²⁰ evaluated a regimen comprising cetuximab-docetaxel-cisplatin, and 5-FU followed by bio-CRT with cisplatin and cetuximab; the 1-year OS and PFS were 68% and 43%, respectively. This author stated that this approach is an effective and tolerable ICT regimen for inoperable oral cancer. Likewise, a phase II clinical trial by Yen⁴⁸ assessed the neoadjuvant cetuximab plus paclitaxel, and cisplatin followed by cetuximab-based RT 70 Gy; and reported an overall response rate of 70.2% and a median OS of 15.2 months. Another study³¹ tested the use of nimotuzumab combined with the docetaxel-cisplatin-FU regimen, reporting a response rate of 89.9% and disease control rates of 100%, suggesting that this regimen is effective and safe in the treatment of advanced oral

squamous cell carcinoma. Similarly, Singh⁴³ assessed the use of gefitinib plus RT 70 Gy, showing a CR of 60% and suggesting that this intervention has better outcomes compared to RT alone.

Chhatui²¹ conducted an RCT assessing the ICT with cisplatin plus 5-FU regimen for three cycles and interferon alpha 2b, which was subcutaneously given at the dose of 3MU, biweekly for three weeks. Then, the participants received CRT with cisplatin 30 mg/m²/week and RT 70 Gy. This author reported a CR and DFS of 64% and 87%, respectively; concluding that this approach may produce superior outcomes. However, it is useful to highlight that there is limited evidence about it. Thus, the effectiveness of treatments involving immunotherapy for unresectable oral cancer is uncertain.

A Phase III RCT³⁰ aimed to assess a combination of recombinant adenoviral p53 (rAd-p53) gene therapy and intra-arterial delivery of CT agents for the treatment of oral squamous cell carcinoma. In that study, 99 participants were recruited and randomly divided into three arms: arm I (n= 35; intra-arterial infusion of rAd-p53 plus CT), arm II (n = 33; intra-arterial infusion of rAd-p53 plus placebo CT), and arm III (n = 31; intra-arterial infusion of placebo rAd-p53 plus CT). The 5-year OS rate was 48.5%, 30% and 22.5% for arm 1, arm 2 and arm 3, respectively. These findings suggest that the use of

Table 6 Summary of Evidence on Targeted Therapy, Immunotherapy and Gene Therapy

Author and Year	Clinical Stages	Interventions	Follow-Up	Outcomes
Chang 2017 ²⁰	IV	(n=43): cetuximab + docetaxel + cisplatin + 5-FU followed by bio-CRT with cisplatin and cetuximab	15 M	OS [I Y]: 68% PFS [I Y]: 43%
Chhatui 2015 ²¹	III IV	(n=25): ICT with cisplatin + 5-FU + interferon α -2b followed by CRT (cisplatin + RT 70 Gy)	15 M	CR: 64% DFS: 87%
Li 2014 ³⁰	III IV	Arm 1 (n=35): rAd-p53 + CT Arm 2 (n = 33): rAd-p53 + placebo CT Arm 3 (n = 31): placebo rAd-p53 + CT	36 M	CR: Arm 1, 48.5%; Arm 2, 16.7%, Arm 3, 17.2%
Meng 2014 ³¹	III IV	(n=9): nimotuzumab + docetaxel + cisplatin + 5-FU	NR	RR: 89.9% DCR: 100%
Singh 2013 ⁴³	III IV	(n=30): gefitinib + RT 70 Gy	20 M	CR: 60%
Yen 2019 ⁴⁸	IV	(n=39): cetuximab + paclitaxel + cisplatin followed by BioRT (cetuximab + RT 70Gy)	6.5 Y	OR: 70.2% CR: 8.5% PR: 61.7% PFS [Median]= 10.3 M OS [Median]= 15.2 M

Abbreviations: CR, complete response; CRT, chemoradiotherapy; CT, chemotherapy; DCR, disease control rate; DFS, disease-free survival; FU, fluorouracil; ICT, induction chemotherapy; M, months; NR, not reported; OR, overall response; OS, overall survival; PFS, progression-free survival; PR, partial response; rAd-p53, recombinant adenoviral p53; RR, response rate; RT, radiotherapy; Y, years.

rAd-p53 gene therapy plus CT can improve the clinical outcomes for people suffering from unresectable oral cancer, but these results should be considered with caution since there is lacking evidence about the effectiveness of these treatment options.

Discussion

In order to describe the main therapeutic options in unresectable oral squamous cell carcinoma, we conducted an evidence-based comprehensive analysis. This review may be the first one focused on unresectable oral cancer since we did not find any previous report. Moreover, other reviews focused on oral cancer treatment usually based their conclusions on studies including a large proportion of patients with other types of cancers such as head and neck tumors, and just a small proportion of people suffering from oral cancer.

Our findings suggest that the optimal treatment of patients with unresectable oral cancer is challenging; thus, there is a sprinkling of studies proposing a range of therapeutic options, such as RT, CT, CCRT, immunotherapy, targeted therapy plus CT or RT, and gene therapy plus CT. However, it is useful to highlight that the scientific evidence supporting many of these approaches is limited. Overall, the use of CCRT, and ICT followed by CRT have

shown good clinical results such as improvement of overall response,⁴⁹ OS and LCR rate.³⁵ In this sense, most studies^{19,24–28,33,35,40} supported the use of the CCRT as a therapeutic option in people with unresectable oral cancer. Likewise, some studies^{36,45,49} indicated the benefits of the use of ICT followed by CRT for the treatment of these patients. Consequently, these therapeutic options can be useful when surgery is not feasible.

However, some factors should be considered to choose the optimal therapeutic options in unresectable oral squamous cell carcinoma. Firstly, the treatment side effects; to illustrate, Chhatui²¹ reported that those patients receiving ICT had more toxicities and treatment interruptions; among its side effects were skin reactions, mucositis, anemia, leukopenia, nausea, and vomiting. Similarly, Sato⁴⁰ reported that among the adverse effects of using CCRT are stomatitis, dermatitis, anemia, and liver dysfunction. Secondly, individual patient factors and their possible role on the treatment effect should be taken into account. Some reports suggest that differences in lifestyle, living environment, and race, may affect the therapy effectiveness in oncology.^{51–53} For example, diet management improves OS and other clinical outcomes in people with head and neck cancers.⁵¹ Likewise, after oral cancer treatment, black people usually have poorer

OS rates than whites.⁵² Moreover, changes in habits such as quitting smoking, alcohol drinking, and betel nut could have a considerable impact on therapeutic interventions, especially in patients with unresectable oral cancer.⁵³ However, we highlight that few studies^{20,22,24,26,28,34–38,41–43,46} in this review reported those factors, and only one study²² analyzed the influence of them on treatment response. Finally, the reasons given to determine whether the tumor was unresectable should be considered; since unresectable oral neoplasms due to technical reasons could have different treatment responses compared with those unresectable tumors due to patients' comorbidities or poor general health status.¹³ Overall, any treatment should be judged and discussed with a multidisciplinary team, evaluating its risks and benefits.

Our findings may be comparable with the results reported by Alzahrani,¹³ who narratively reviewed the evidence on the optimal care for people suffering from locally advanced oral cancer, concluding that when surgery is not recommended, these patients can be treated by curative CRT. In addition, this author suggested the use of ICT before surgery or CRT for unresectable oral cancers. However, these results should be taken with caution since there are some differences between these two reviews; firstly, most studies included in the Alzahrani¹³ review had focused on head and neck cancers, while our review included studies exclusively focused on oral cancer, or those studies showing results separately for this oral disease. Secondly, since our main goal was to describe the therapeutic options when surgery is not feasible, we did not consider interventions before or after surgical treatments, which intent to become an unresectable lesion to an operable one or to provide adjuvant therapy.^{54,55}

Similarly, our findings also should be put into context. So initially, as it has been previously reported, there is limited evidence of therapeutic options in unresectable oral cancer.¹⁴ Secondly, most studies^{19,22–26,28,29,32–42,44,46,47,50} included in this review had an observational design and many of them conducted a retrospective analysis,^{22,24–26,28,34,36–42,47,50} therefore their conclusions may be biased since observational studies are not the best design to assess the effectiveness of treatments; so high-quality RCTs must be conducted, which have major relevance for clinical practice.⁵⁶ Finally, the methodological quality of some studies^{26,27,29,31,32,34–38,40} was suboptimal using the JBI's tool. Thus, any therapeutic option in this review should be analyzed and interpreted considering the limitations of each selected study.

The main practical implications of this review are related to helping practitioners and patients in the decision-making

process. Given knowing the available evidence and its quality is so important to provide evidence-based health care, the findings of this review can be useful to improve the management of oral cavity cancer. In this sense, those interventions identified as beneficial could be considered into dental clinical practice to provide evidence-based dentistry. Similarly, those treatments that have been used for decades without evidence support, and have no potential benefits, should not be considered as options to treat unresectable mouth cancers. However, it is useful to highlight that this review does not pretend to replace any clinical practice guideline. Thus, any treatment should be adapted for each patient considering the clinical expertise, the available resources, their risk/benefit ratio, and other contextual aspects.⁵⁷

Another potential implication of this review is related to conduct high-quality research on those interventions with lacking evidence such as gene therapy, which had only one selected study.³⁰ In this sense, cancer gene therapy is considered a novel approach that may significantly improve clinical outcomes such as OS of patients suffering from cancers.^{58,59} Likewise, it is useful to mention that more research on targeted therapy is needed. All these new therapeutic approaches have been developed on a better understanding of molecular mechanisms involved in the cancer disease; thus, they are more selective against tumor cells, which leads to decrease in side effects. However, their clinical applicability to treat head and neck cancers still is unclear.⁶⁰

Some limitations in this review should be mentioned such as the language barrier, due to all evidence found was published in English, which eliminated the inclusion of available evidence published in any other language. However, it is useful to highlight that no restrictions about languages were performed; moreover, since most evidence is published in English, it is more likely that evidence meeting the eligibility criteria is published in this language.

Among the strengths of this review, we highlight that all methods were described in a protocol in advance. Moreover, a sensitive search strategy was carried out, so it is unlikely that any relevant evidence was missed. Similarly, at least two reviewers independently conducted the whole processes of selection, methodological quality assessment, and data extraction. All these processes provide reasonable confidence in our results.

Conclusion

There is lacking evidence about the benefits of some therapeutic options for unresectable oral squamous cell carcinoma. Overall, these patients can be treated using

a multimodal approach, such as CCRT or ICT followed by CRT, which have shown good clinical outcomes. However, other therapeutic options could be considered depending on the assessment of risk/benefits, tumor extension and patient values and preferences. In all cases, any treatment should be adapted for each patient considering the clinical expertise, the available resources, and other contextual aspects.

Data Sharing Statement

All data generated or analyzed during this study are included in this published article and its supplementary information files.

Author Contributions

Conceived the study: MM and LTA. Designed the study: MM, LTA and CLA. Analyzed the data: MM, LTA, CLA. Wrote the first draft of the manuscript: MM and LTA. Contributed to the writing of the manuscript: MM, LTA, CLA. All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; agreed to submit to the current journal; gave final approval of the version to be published; and agreed to be accountable for all aspects of the work.

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