Effectiveness of screening for oral cancer and oral potentially malignant disorders (OPMD): A systematic review

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ABSTRACT
Oral cancer (OC) is a debilitating disease with a high mortality rate when diagnosed in advanced stage. Conversely, early-stage OC has a high survival rate, supporting a need for early detection programmes. A previous systematic review of clinical trials evaluating efficacy of screening for OC was inconclusive. This systematic review aimed to determine the impact of screening for oral lesions on reducing mortality and incidence of OC by looking at a broader spectrum of evidence.

The search for randomized controlled trials and observational studies with a control group was conducted in PubMed, OVID, Cochrane, CINAHL and grey literature sources. Risk of bias for included studies was assessed with the tools developed by the Cochrane collaboration.

Six out of two identified randomized trials and five observational studies had moderate to high risk of bias. Nevertheless, the predictions on impact of OC screening on incidence and mortality were similar across the majority of the studies. The meta-analysis concluded on a 26% decrease in OC mortality, and an 19% decrease in advanced OC cases as a result of OC screening in high-risk population. Three out of four studies did not identify an impact of screening on OC incidence. No positive impact of OC screening on incidence or mortality among general population was identified in the only available randomized trial. Consistency in the outcomes and the limitations of the few available studies suggest a need for real-life setting research to evaluate the overall effectiveness of screening for OC in high-risk population.

1. Introduction
Definition of oral cancer (OC) varies in the literature, but for the purpose of early detection, OC is usually considered as a malignant neoplasia which arises on the lip or oral cavity. OC accounted for 2 % of all cancers, as well as 1.9 % of all cancer related deaths resulting in almost 355 000 new cases diagnosed and over 177 000 associated deaths (Miranda-Filho and Bray, 2020). While OC presents a five-year overall survival around 50 %, early OC diagnosis may increase it to 85 %. This supports the rationale of early detection contributing to better outcomes (National Cancer Institute. Browse the SEER Cancer Statistics Review (CSR) (2014)), with survival rates directly linked to the cancer stage at diagnosis (Strome et al., 2018).

OC is the 16th most common cancer with OC incidence varying widely around the world. Asia, Europe, and Oceania have the highest incidence rate in the world, while Asia has the highest OC mortality (Fig. 1). The incidence of OC in populations depends on the prevalence of the attributable risk factors such as tobacco chewing and smoking, betel quid usage, and alcohol consumption (Petti, 2009; Kumar et al., 2016). For instance, a greater incidence is noted in males, who culturally exhibit higher exposure to the risk factors (Rao et al., 2013; Zain, 2001), with a world age standardized incidence rate of 6 versus 2.3 per 100 000 people respectively, Fig. 1.

Screening programmes for other cancers sites demonstrated clinical

Abbreviations: CG, Control group; CI, Confidence interval; CINAHL, Cumulative Index to Nursing and Allied Health Literature; COE, Conventional oral examination; IG, Intervention group; ISRCTN, International Standard Randomised Controlled Trial Number; MSE, Mucosal self-examination; OC, Oral cancer; OR, Odds ratio; OSF, Oral submucous fibrosis; OPMD, Oral potentially malignant disorders; PYO, Person years of observation; RCT, Randomized clinical trial; ROB, Risk of bias; ROBINS-I, Risk of bias in non-randomized interventional studies; RR, Risk ratio/Relative risk; TB, Toluidine blue.

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benefits through detecting pre-malignant and early stage cancer lesions (Mandrik et al., 2019; Peirson et al., 2013). Motivated by the disease burden and conventional approach of community screening, a potential of different OC screening approaches to detect oral potentially malignant disorders (OPMDs) and invasive cancer has been investigated in randomised trials and the following systematic review (Brocklehurst et al., 2013). From the investigated approaches, the current standard of screening is through conventional oral examination (COE) under direct light. This is usually performed by a general dentist or doctor, but other healthcare workers such as nurses or community health workers have been known to assist in screening examinations with high efficiency noted after appropriate training (Birur et al., 2019). The healthcare worker redirects the suspected positive case to the diagnostic pathway including biopsy and histopathological confirmation (Higgins, 2021).

Considering potential benefits of OC screening, a few countries with high incidence of OC implemented national or pilot OC screening programmes targeting high-risk population, for instance Cuba, Taiwan (China), Kerala (India) and Sri Lanka. Meanwhile, commissioned by the Cochrane collaboration a systematic review on effectiveness of OC mortality, as well as downstaging (Appendix 1). We further hand-searched the citations of the retrieved eligible papers to identify additional publications that might have been missed during the initial search. We also searched clinicaltrials.gov for non-published studies.

2.2. Inclusion and exclusion criteria

This review considered quasi-experimental, randomized controlled (cluster and individually), case-controlled, cohort and cross-sectional studies with a control group.

b) Type of Participants.

Studies were considered eligible if included an adult population group (defined as anyone over 15 years of age) of any gender who attend OC screening programs. While the adult population is typically defined as over 18 years old, we extended the lower limit to 15-years old considering that the definition of adult population in the National Oral Cancer Screening programme in Cuba is 15 years and older (Birur et al., 2019). Symptomatic population, such as individuals with confirmed OC or a history of OC, were excluded. Studies on either general-risk or high-risk populations were included. High-risk population was broadly described as regular tobacco (any type or form – smoked or smokeless) and/or alcohol consumption.

c) Type of Intervention.

Studies investigating any screening method for OC or OPMD were eligible. OC screening comparing types of examination including, but limited to, the conventional oral visual examination, chemical staining, auto-fluorescence, biomarker analysis and chemiluminescence versus no screening or placebo, were also eligible. Self-examinations, conducted by patients under direction and supervision by healthcare workers, were excluded, considering the suggested low test accuracy (Ghani et al., 2019; Lee et al., 2019).

d) Comparator.

All studies considered eligible had to have a control group, including but not limiting to no screening, ‘usual care’ (e.g., opportunistic screening) or modified interventions. Studies which lacked a control group were excluded from this review.

e) Outcomes.

In order to be considered eligible, at least one of the following primary outcomes was required:

- OC mortality
- OC incidence
- Clinical stage at diagnosis

Studies identified in accordance with the above outcomes may also be subjected to extraction in terms of the following secondary outcomes:

- Sensitivity and specificity of screening programs or diagnostic examinations
- Overdiagnosis
- Other clinical benefits, such as incidental findings: detection of dental issues, other cancers, or systematic health problems.

As per conventional definition, OPMD has been defined as "a group of lesions and conditions characterized by a variably increased risk of developing cancers of the lip and the oral cavity" (Warnakulasuriya et al., 2021) such as leukoplakia, erythroplakia, lichen planus, and oral submucous fibrosis.
3. Risk of bias in included studies

Cochrane’s Risk of Bias (ROB-2) in RCTs tool and Cochrane’s Risk of Bias in Non-randomised Studies – of Interventions (ROBINS-I) was used to assess the risk of bias in studies independently by two reviewers resolving any disagreements through discussion.

3.1. Data analysis

Review Manager software was applied to synthesise the outcomes. Dichotomous outcomes, such as OC mortality, OC detection, advanced stage OC detection and OPMD detection were used to estimate the effect of screening as expressed by risk ratios with a confidence interval (95%).

We used the I² statistic to assess a level of heterogeneity and so to define a possibility for quantitative synthesis. A meta-analysis, with a weighted random effects model was performed to report on the effect of OC screening on mortality and advanced OC cases. Only studies reporting similar outcomes were included in the meta-analysis.

4. Results

Of the initial 12,276 records identified, seven studies published between 1995 and 2021 were selected for inclusion (Fig. 2). The characteristics of the included studies are reported in Table 1.

4.1. Quality of evidence

Five of the seven studies included in this review are judged to be at a high or serious risk of bias (16,20–23) (Table 2). One study was judged to be at a moderate risk of bias (Sankaranarayanan et al., 2013), and only one study was deemed to be at a low risk of bias (Su et al., 2010). Five of the included studies are observational studies, with a high risk of selection bias and confounding (Frenández Garrote et al., 1995; Chuang et al., 2017; Morikawa et al., 2021; Ho et al., 2019; Sankaranarayanan et al., 2002)(Appendix 3).

The overall quality of the evidence found is poor and limited. The lack of high-quality studies precluded us from only analysing studies deemed to be at a low or moderate risk of bias. However, the results presented above are consistent, and show a positive effect of OC
4.2. Effect on OC mortality

Three studies of different designs reported the effect of OC screening on OC mortality (Chuang et al., 2017; Ho et al., 2019; Sankaranarayanan et al., 2013). A meta-analysis of two studies reporting the number of deaths and the population size in the intervention and the comparator groups was performed based on the different subgroups as well as an overall statistic (Chuang et al., 2017; Sankaranarayanan et al., 2013). A mortality significant decrease of 26 % with minimal heterogeneity ($I^2 = 0$ %) was noted when analysing the high-risk group, but no difference was observed among the general population (Fig. 3). The limited number of studies, and methodological differences included in this meta-analysis should be noted. Pei-Shan Ho et al. (Ho et al., 2019), also reported the hazard ratio for mortality associated with OC screening of 0.92 in screened vs non-screened population (95 % CI, 0.84–1.00) (Ho et al., 2019).

Table 1

Characteristics of included studies.

| Author/year/Country | Inclusion criteria | Cluster randomization | Sample size | Number of screening rounds | Follow up for screen positive cases-definition | Compliance with intervention | Endpoints measured | Study design | Intervention | Control | Follow up compliance rate |
|---------------------|--------------------|----------------------|-------------|-----------------------------|-----------------------------------------------|-----------------------------|-------------------|-------------|-------------|---------|--------------------------|
| Garrote et al., 1995 | IG 1984–1990 | IG- 92 % | 12 990 677 | 0-2 | Not reported | Males- 11.9 % | OC incidence OC mortality | 40 years | National screening program, conducted by dentists | Routine care, data taken from national cancer registry | 77.60 % |
| Garrote et al., 1995 | CG 1984–1990 | CG for 1 round of screening- 46 % | 84 228 675 | 0-2 | Not reported | Females- 19.9 % | OC late-stage incidence | 40 years | Determination of screening history in advanced OC cases, part of the national screening program | Individuals without screening history who were reported to have OC | 77.60 % |
| Sankaranarayanan et al., 2013 | IG 1984–1990 | IG- 92 % | 200 | 1 -more than 3 | Not reported | IG- 91 % | OPMD incidence OPMD incidence | 40 years | Analysis of Taiwan Oral mucosal screening program | Data linked to National cancer registry used to identify cases in the control group, who did not attend screening | 77.60 % |
| Pei-Shan Ho et al., 2019 | CG 1984–1990 | CG- 91 % | 600 | 0 | Not reported | CG- 91 % | OPMD incidence OPMD incidence | 40 years | IG- late-stage OC CG- healthy individuals residing within 200 m of the matched OC case | OPMD incidence OC incidence | 77.60 % |

4.3. Effect on OC incidence

For OC detection in the general population, three studies (Frenández Garrote et al., 1995; Morikawa et al., 2021; Sankaranarayanan et al., 2013) were included, as well as two studies in the high-risk group (Chuang et al., 2017; Sankaranarayanan et al., 2013). None of the included studies reported a statistically significant decrease in OC incidence in the general population, whereas in the high-risk population one observational study (Chuang et al., 2017) found a statistically significant decrease in OC incidence of 30 % which decreased to 17 % after adjustment for self-selection bias (Fig. 4). No impact on incidence for OC screening was observed in randomised controlled trials (Sankaranarayanan et al., 2013). A meta-analysis could not be conducted for the effect of OC screening on OC incidence due to the high level of heterogeneity ($I^2 = >70$ %).

4.4. Effect on OC stage at diagnosis

Three studies reported increased detection of early-stage OC through the screening programmes. Pei-Shan Ho et al. (Ho et al., 2019) reported
an increase in early OC detection for confirmed OPMD’s (99 %), confirmed non OPMD’s (85 %) not referred (49 %) or did not comply with referral (25 %), which may lead to earlier treatment, resulting in lower advanced OC cases (Ho et al., 2019). Sankaranarayanan et al. (Sankaranarayanan et al., 2013) reported a higher percentage of early stage OC in the screened vs non-screened group (39.4 % vs 27 %) (Sankaranarayanan et al., 2013). Garrote et al. (1995) reported an increased proportion of early OC detection, 50 % in the assigned control year of 1983 compared to 64 % in 1989 (Frenández Garrote et al., 1995).

Four studies reported the reduction of advanced stage OC detected at diagnosis as outcome (defined as stage 3 and 4 by TNM classification-8th edition). For illustrative purposes, the reported relative risk of late stage diagnoses is presented on Fig. 5 (Chuang et al., 2017; Ho et al., 2019; Sankaranarayanan et al., 2013).

Fig. 5 shows four studies reporting a statistically significant reduction in incidence of advanced stage OC in high-risk populations. Chuang et al. (Chuang et al., 2017); recorded a 21 % reduction in advanced stage presentation amongst screening participants after adjustment for self-selection bias 0.79 (95 % CI, 0.76–0.82) (Chuang et al., 2017). Sankaranarayanan et al. (Sankaranarayanan et al., 2013) recorded a 21 %
decrease in advanced stage OC presentation amongst screening participants in India, RR = 0.79 (95 % CI, 0.68–0.91). Sankaranarayanan et al. (Sankaranarayanan et al., 2002) reported a 19 % decrease in advanced OC in screened individuals in Cuba. They also reported an OR of advanced OC from 0.67 (95 % CI, 0.46–0.95) after one screening to 0.41 (95 % CI, 0.24–0.68) in individuals who were screened two or more times, alluding to greater protection offered by subsequent screenings. The calculated RR based on reported distribution of advanced stage OC in the Pei-Shan Ho et al. (Ho et al., 2019) study, showed a statistically significant decrease in advanced stage OC presentation of 15 %, RR = 0.85 (95 % CI, 0.78–0.92) (Ho et al., 2019).

A meta-analysis showed a reduction in advanced stage OC of 19 % (95 % CI, 0.74–0.88) in the high risk screened groups with a minimal heterogeneity of 30 %.

4.5. Effect on OPMD incidence

Outcomes on OPMD were reported by 4 studies (Su et al., 2010; Chuang et al., 2017; Morikawa et al., 2021; Ho et al., 2019). Morikawa et al. (2020) reported a statistically significant 14 % decreased risk of developing OPMD in the countermeasure group (Morikawa et al., 2021). Chuang et al. (Chuang et al., 2017), reported a statistically significant 33 % increase in OPMD detection for a subsequent screening (Chuang et al., 2017). Pei-Shan Ho et al. (Ho et al., 2019); reported a statistically significant hazard ratio of 0.72 (0.64, 0.81) for individuals with confirmed OPMD in the screening group. This equates to a decreased OC mortality of 28 % for individuals with confirmed OPMD detected during OC screening (Ho et al., 2019). Su et al. (Su et al., 2010); reported a OPMD and malignant lesion detection risk ratio of 1.05 (95 % CI, 0.74–1.41), however this result was not statistically significant (Su et al., 2010).
4.6. OC screening-related harms

None of the included studies explicitly reported any data or analysis relating to overdiagnosis. Overdiagnosis is commonly measured in studies with the long follow-up and may be interpreted as a higher incidence of OPMD or early-stage OC in the intervention versus control group. A statistically non-significant but higher (31.2 vs 27.2) OC incidence per 100 000 was noted by Sankaranarayanan et al. (Sankaranarayan et al., 2013) in the screened vs non-screened group of general OC risk with 15 years of the follow-up and may be interpreted as a possibility for overdiagnosis related to OC screening. However, the authors mentioned mitigation of over-treatment, by conservative management of benign lesions and monitoring of OPMD (Sankaranarayanan et al., 2013).

5. Discussion

The objective of this systematic review was to determine the impact of OC screening on OC incidence, OC clinical stage at diagnosis, and OC mortality. Our meta-analysis demonstrated a risk reduction for OC mortality of 26% and for advanced OC cases of 19% among high-risk participants of the OC screening. Only one study (a RCT) assessed the impact of OC screening on advanced stage OC and mortality among general risk population and was not able to demonstrate a statistically significant impact of screening (Sankaranarayanan et al., 2013). To supplement these empirical findings, a recent analysis re-evaluated the data from Kerala trial using a Cox proportional hazards risk prediction model, assigning each person in the model the counterfactual hazard of OC mortality in the absence and presence of screening. The study concluded on 27% OC mortality reduction in the screening versus control arms (Cheung et al., 2021), providing proof of principle for risk-based OC screening.

Several studies reported on the effect of detection of OPMD through OC screening or a decrease in risk of developing OPMD when screened (Su et al., 2010; Chuang et al., 2017; Morikawa et al., 2021), though, considering that the natural history of OC is still not well explored, it is not clear how much OPMD detection will contribute to the final clinical endpoint (mortality decrement) or contribute to overdiagnosis. This issue may become especially important considering that three out of four included studies (two evaluating the same public screening programme in Taiwan) did not identify a statistically significant impact of OC screening on OC incidence. Overdiagnosis was not explicitly explored in the included studies and the risk and the magnitude of it needs to be addressed in further research.

The results from our systematic review strengthen the conclusions from the previous systematic review of (Brocklehurst et al., 2013) based only on one RCT (Kerala, India), that there is evidence of benefits of COE. In particular, the results from our review show that a screening program targeting individuals in a high-risk group decreases the OC mortality rate and incidence of late stage (stage 3 and 4) OC at diagnosis in the screened groups and improves detection of early-stage OC and OPMD, thus contributing to OC downstaging. While (Brocklehurst et al., 2013) concluded that there is insufficient evidence to recommend a population wide OC screening approach, our review, looking at the broader literature of different research designs, suggests the similarity in the risk reduction for clinical endpoints despite the methodological limitations and heterogeneity in designs and outcomes. Thus, we assume that implementation studies, i.e., well-designed pilot OC screening programmes in high-risk populations, will be necessary to support the existing evidence on effectiveness of OC screening. Our conclusion on usefulness of the available OC screening studies to inform policy questions albeit demonstrated weaknesses in their design, is aligned with the previous recent narrative reviews (D’Cruz and Vaish, 2021; Warnakulasuriya and Kerr, 2021). Despite the low quality of evidence and limited number of studies, we believe these results may inform public health policies, however, there is a significant need for further high-quality implementation research in this domain.

6. Limitations of the review

While the search strategy and databases used to conduct the search were comprehensive, it is possible that some studies may be missed considering that the abstracts were screened by one reviewer only. We did not have a language restriction, but no studies were found in languages other than English. While we were able to conduct a meta-analysis, the low number and lack of high-quality studies may reduce the impact of the findings in our review.

7. Research and information gaps

Pilot studies of prospective design with well-designed evaluation protocol should be conducted in countries with a high prevalence of tobacco and or alcohol consumption to provide local evidence for policy making in these nations. These studies should follow the structure of implementation research and supplement the assessments of screening benefits with evaluation of screening-related harms (overdiagnosis), and implementation outcomes, such as costs, acceptability and feasibility (Proctor et al., 2011). Due to the relatively low compliance rate and a rate of follow-up for the positive cases, reported in some of the current studies, incentives may be offered to improve compliance and uptake with these screening programs, as well as subsequent referrals. For instance, such motivational factors may include OC education, OC risk factor education, alcohol and tobacco use cessation therapy, and personalized or specific referral letters (Camilloni et al., 2013). More research is needed to conclude on efficacy of adjunctive technologies to COE.

8. Conclusion

OC screening via COE in high-risk populations results in a decrease in OC mortality (26%) and advanced OC cases at diagnosis (19%). There is no conclusive evidence on impact of OC screening on OC incidence as well as no evidence on impact of other OC screening methods on clinical outcomes. No studies were detected to assess a magnitude of over-diagnosis related to OC screening, ultimately motivating for more research to be done to address this issue.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

No data was used for the research described in the article.

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Appendix A. Supplementary data

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