OESOPHAGUS

Preoperative chemoradiotherapy for oesophageal cancer: a systematic review and meta-analysis

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Background: The benefit of neoadjuvant chemoradiotherapy in oesophageal cancer has been extensively studied but data on survival are still equivocal.

Objective: To assess the effectiveness of chemoradiotherapy followed by surgery in the reduction of mortality in patients with resectable oesophageal cancer.

Methods: Computerised bibliographic searches of MEDLINE and CANCERLIT (1970–2002) were supplemented with hand searches of reference lists.

Study selection: Studies were included if they were randomised controlled trials (RCTs) comparing preoperative chemoradiotherapy plus surgery with surgery alone, and if they included patients with resectable histologically proven oesophageal cancer without metastatic disease. Six eligible RCTs were identified and included in the meta-analysis.

Data extraction: Data on study populations, interventions, and outcomes were extracted from each RCT according to the intention to treat method by three independent observers and combined using the DerSimonian and Laird method.

Results: Chemoradiotherapy plus surgery compared with surgery alone significantly reduced the three year mortality rate (odds ratio (OR) 0.53 (95% confidence interval (CI) 0.31–0.93); p = 0.03) (number needed to treat = 10). Pathological examination showed that preoperative chemoradiotherapy downstaged the tumour (that is, less advanced stage at pathological examination at the time of surgery) compared with surgery alone (OR 0.43 (95% CI 0.26–0.72); p = 0.001). The risk for postoperative mortality was higher in the chemoradiotherapy plus surgery group (OR 2.10 (95% CI 1.18–3.73); p = 0.01).

Conclusions: In patients with resectable oesophageal cancer, chemoradiotherapy plus surgery significantly reduces three year mortality compared with surgery alone. However, postoperative mortality was significantly increased by neoadjuvant chemoradiotherapy. Further large scale multicentre RCTs may prove useful to substantiate the benefit on overall survival.

Methods

Selection of randomised trials

The primary source of the reviewed studies was MEDLINE and CANCERLIT, including non-English sources, with the following medical subject headings: oesophageal cancer, chemoradiotherapy, radiotherapy, chemotherapy, RCT, randomised, and clinical trial. The search included literature published through to December 2002. The computer search was supplemented with manual searches of reference lists for all available review articles, primary studies, abstracts from meetings, and bibliographies of books. We have contacted the investigators of an Australian trial, which was reported twice

Abbreviations: CRT, chemoradiotherapy; RCTs, randomised controlled trials; OR, odds ratio; NNT, number of patients needed to treat; NNH, number of patients needed to harm; BED, biologically effective dose; CT, computed tomography
as an abstract,\textsuperscript{19,} \textsuperscript{20} in order to obtain data on randomisation and survival.

Studies were included in the meta-analysis if they were RCTs comparing preoperative CRT plus surgery with surgery alone, if they included patients with resectable histologically proven squamous cell carcinoma or adenocarcinoma of the oesophagus without metastatic disease, and if mortality was assessed as an outcome measure of the effect of the treatment. Among the 11 studies reviewed, six RCTs met the inclusion criteria.\textsuperscript{21–23} Five studies\textsuperscript{16–18} were excluded because they were published as a preliminary report or abstract\textsuperscript{19,} \textsuperscript{20} and subsequently published as a final paper, or because they were published as an abstract, not reporting survival data.\textsuperscript{19,} \textsuperscript{20} As all the trials reported as abstracts\textsuperscript{21} \textsuperscript{22} were subsequently published as full papers, this meta-analysis included only peer reviewed reports.

This meta-analysis was performed according to the QUOROM statement.\textsuperscript{21}

\textbf{Review of the trials}

The trials were first reviewed using a list of predefined pertinent issues that concerned the characteristics of patients and treatments. To assess the methodological quality of RCTs, the two domains of blinding and handling withdrawals and dropouts, using the definitions given by Nicolucci and colleagues,\textsuperscript{26} were used, as suggested by Juni and colleagues.\textsuperscript{25} Each RCT was evaluated and classified by three independent investigators (FF, AV, DDB). Discrepancies among reviewers were infrequent (overall interobserver variations <10%), and were resolved by discussion.

\textbf{Statistical methods}

The crude rate of three year overall mortality was assessed as a measure of treatment effect. These data were available in three RCTs.\textsuperscript{15–17} In the remaining three trials\textsuperscript{12–14} we used the Kaplan-Meier estimates of the three year mortality in the treated and control groups reported in the text. Moreover, to assess the downstaging effect of CRT (that is, the probability of having a less advanced stage of the disease at pathological examination at the time of surgery), we separated patients with negative nodes and patients with positive nodes on pathological examination at the time of surgery. As a measure of treatment benefit, we compared the proportion of patients observed in the treated and control groups who were classified as stage 0, 1, or 2a according to the American Joint Committee on Cancer.\textsuperscript{25} Downstaging by chemoradiotherapy was also used as a measure of treatment effect on pathological response at the time of surgery. Furthermore, we analysed the 90 day hospital mortality (postoperative mortality). Evaluation of therapeutic effectiveness was performed by an intention to treat method. When not reported in the trial, response rate, according to the intention to treat method, was calculated. The number of patients who discontinued their original irradiation regimen because of side effects was also recorded. To combine results from individual trials, we used the proportion of events observed in the treated and control groups. With these observed proportions of events, the odds ratio (OR) was computed for each trial.

The overall OR among the events in both the chemoradiotherapy plus surgery group and the surgery alone group was calculated with models based on both fixed effects and random effects assumptions. In addition to within study variance, the random effects model considers heterogeneity among studies. Because of the different clinical settings and groups of subjects analysed and because the tests for heterogeneity lack statistical power due to the few studies included in this meta-analysis, we have presented the results of random effects models, according to DerSimonian and Laird.\textsuperscript{27} The 95% confidence interval (95% CI) of the odds ratio was also calculated. The overall OR was tested for significance using a Mantel-Haenszel \( \chi^2 \) test.\textsuperscript{28}

Moreover, we in turn excluded each study to ensure that no single study would be solely responsible for the significance of any result (so-called robust analysis). All our analyses were computed using Metaview 4.0. The number of patients needed to treat (NNT) to prevent one death, and the number of patients needed to harm (NNH) to prevent one death by postoperative mortality, which both derive from the inverse of the risk difference, were also used as a measure of treatment benefit and safety.\textsuperscript{29}

To improve the comparability of the different therapeutic regimens and to assess the relationship between radiation dose and survival benefit, the biologically effective dose (BED), corrected for time of the various radiation schedules, was estimated.\textsuperscript{30} In the Scandinavian trial,\textsuperscript{19} three different treatment arms were compared with the same surgery arm as the control. We included in the analysis only the chemoradiation treatment arm of this RCT, using effect size estimates calculated from observations on that measure. Therefore, the statistical analysis used only independent estimators of effect size.

Subgroup analyses were used to explore and explain the diversity among results of different studies. We used two stratifying variables: adenocarcinoma versus squamous cell carcinoma and BED >35 versus BED \(<35. A \chi^2\) test for interaction\textsuperscript{31} was used to examine whether the effect of treatment varied significantly between subgroups.

\textbf{Source of support}

This meta-analysis was entirely supported by the authors’ respective institutions.

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|l|l|l|l|l|}
\hline
\textbf{Study (reference)*} & \textbf{Accrual period (y)} & \textbf{Patients randomised (n)} & \textbf{Male (%)} & \textbf{Mean age (y)} & \textbf{Histology (%)} & \textbf{Preoperative stage (% of patients)} & \textbf{Source of support} \\
\hline
1 Nyaaga et al\textsuperscript{12} & 1983–88 & CRT+S & 53 & 70 & 60.1 & SCC & 100 & I–II (100) \\
2 Le Prise et al\textsuperscript{13} & 1988–91 & CRT+S & 41 & 93 & 56 & ADENO & 100 & I–II (100) \\
3 Apinop et al\textsuperscript{14} & 1986–92 & CRT+S & 34 & 76 & 59.6 & SCC & 100 & I (14.8), II (82.3); not specified (5.9) \\
4 Walsh et al\textsuperscript{15} & 1990–95 & CRT+S & 58 & 67 & 65 & ADENO & 100 & I (12.2), II (65.8); not specified (22) \\
5 Bosset et al\textsuperscript{16} & 1989–95 & CRT+S & 50 & 80 & 65 & SCC & 0 & I (112.5), II (55.6); not specified (8.9) \\
6 Urba et al\textsuperscript{17} & 1989–94 & CRT+S & 50 & 84 & 62 & ADENO & 0 & I (17.1), II (82.9); not specified (5.9) \\
\hline
\end{tabular}
\caption{Patients characteristics of trials included in the meta-analysis.}
\end{table}

*For expansion of the study names, see corresponding reference. CRT+S, chemoradiotherapy plus surgery; S, surgery; SCC, squamous cell carcinoma; ADENO, adenocarcinoma; NR, not reported.
RESULTS
Features of the RCTs
The main features of the trials included in the meta-analysis are shown in table 1. The six RCTs included 764 patients, 385 of whom received CRT before surgery. The percentage of males ranged from 78% to 93%. The criteria for inclusion were uniform in all but one RCT, which included only patients with locally advanced but otherwise operable tumours. Only two studies were multicentre trials. The sample size of each RCT varied greatly, ranging from 69 to 297 patients. No adequate blinding was used in two trials and the method of randomisation was not clearly reported in the remaining four studies. Three studies did not clearly define criteria for handling withdrawals.

The preoperative staging procedures were uniform in all RCTs. In four RCTs, all patients underwent computed tomography (CT) while in the study of Le Prise and colleagues and the Irish trial CT was performed in selected patients only. Data on preoperative staging of the tumour were similar in all RCTs, except the small study of Apinop and colleagues in which stage I oesophageal cancer was excluded (table 1). In all RCTs no significant differences were reported in the remaining four studies. Three studies did not clearly define criteria for handling withdrawals.

A transthoracic resection was performed in all RCTs but one in which a transhiatal oesophagectomy was employed. The rate of resection (curative or palliative) reported in all RCTs ranged from 82% to 97%. The proportion of patients who underwent resection classified as curative (defined by the surgeon as margins of the resected tissue free of tumour) was reported in only three RCTs ranging from 46% to 97%.

Pathological stage was assessed in all RCTs at the time of operation according to the American Joint Committee on Cancer. The rate of pathological stage III oesophageal cancer in patients treated by surgery alone was comparable in all RCTs, ranging from 50% to 78%. In four RCTs, only patients with squamous cell carcinoma were included, while in the study by Walsh and colleagues all subjects had adenocarcinoma, and in the RCT by Urba and colleagues patients with either squamous cell carcinoma or adenocarcinoma were included (table 1).

The therapeutic regimens of RCTs included in the meta-analysis are shown in table 2. Considerable heterogeneity was observed both in radiotherapy and chemotherapy protocols among the studies. A large variability in irradiation schedules between trials was found in: (a) the total dose, ranging between 20 and 45 Gy; (b) the daily dose, ranging between 1.75 and 3.70 Gy; and (c) the number of fractions, ranging between 10 and 30 given over 24 and 21 days, respectively.

Variability in the chemotherapy protocols between trials was found in: (a) the number and type of chemotherapeutic agents administered in combination with cisplatin; and

<table>
<thead>
<tr>
<th>Study (reference)</th>
<th>Total dose (Gy)</th>
<th>BED (Gy)</th>
<th>BED corrected by time (Gy)</th>
<th>Chemotherapy</th>
<th>Surgery</th>
<th>Interval between end of irradiation and surgery (weeks)</th>
</tr>
</thead>
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<tr>
<td>Nygaard et al</td>
<td>35</td>
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<td>1.75</td>
<td>CDDP 20</td>
<td>0-5</td>
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<td>10/12</td>
<td>2</td>
<td>CDDP 17.8</td>
<td>5</td>
<td>1-21</td>
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<td>Apinop et al</td>
<td>40</td>
<td>20/28</td>
<td>8</td>
<td>CDDP 33.6</td>
<td>100</td>
<td>1-21</td>
</tr>
<tr>
<td>Walsh et al</td>
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<td>15/21</td>
<td>2.67</td>
<td>CDDP 42.6†</td>
<td>75</td>
<td>1-5-25</td>
</tr>
<tr>
<td>Bosset et al</td>
<td>37</td>
<td>10/24</td>
<td>3.7</td>
<td>CDDP 38.4</td>
<td>80</td>
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<tr>
<td>Urba et al</td>
<td>45</td>
<td>30/21</td>
<td>1.5</td>
<td>CDDP 43.6†</td>
<td>20</td>
<td>1-5-17-21</td>
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</table>

*Tipot sqamo = 4.5 days, Tipotadeno = 6 days, α = 0.3
BED, biological equivalent dose; CDDP, cisplatin; BLM, bleomycin; 5-FU, 5-fluorouracil; VNB, vinblastin.
†For expansion of the study names, see corresponding reference.

Figure 1 Meta-analysis of six randomised controlled trials of preoperative chemoradiotherapy (CRT) for resectable oesophageal carcinoma using a random effects model. The odds ratio (OR) and 95% confidence interval (CI) for the effect of treatment on three year overall mortality are shown on a logarithmic scale. Studies are arranged by publication year.
(b) the dose and scheduling of chemotherapeutic drugs (table 2).

In one RCT, chemotherapy and radiotherapy were given sequentially with radiotherapy starting three weeks after completion of chemotherapy. In the remaining five RCTs, CRT was given simultaneously (concomitant strategy) with daily chemotherapy or with intermittent chemotherapy.

Figures 2 and 3.

**Overall mortality**

The effect of adjuvant CRT on overall mortality (six RCTs: 764 patients, 573 deaths) is shown in fig 1. Although the effect of treatment on total mortality favoured CRT in all the six trials, a statistically significant difference was observed in only one. However, the pooled estimate of the treatment effect was statistically significant (OR 0.53 (95% CI 0.31–0.92); z = −2.23; p = 0.03) (NNT = 10). Similar results were obtained when a fixed effect model was used (OR 0.61 (95% CI 0.43–0.85); z = −2.89; p = 0.004).

Robust analysis showed that the five trials remaining after exclusion of either the trial by Walsh and colleagues or by Urba and colleagues lost statistical significance for overall mortality (excluding the trial by Walsh: OR 0.73 (95% CI 0.51–1.05), z = −1.68 p = 0.093; excluding the trial by Urba: OR 0.55 (95% CI 0.29–1.06), z = −1.78; p = 0.074).

We performed subgroup analyses to evaluate whether there was evidence of a different effect of CRT in predefined subgroups of patients. For overall mortality, the pooled OR was statistically significant in patients with adenocarcinoma (OR 0.24 (95% CI 0.07–0.78); z = −2.36, p = 0.018) but not in those with squamous cell carcinoma (OR 0.81 (95% CI 0.55–1.19); z = −0.29, p = 0.77) with a significant interaction (χ² for interaction = 7.79; 1 df; p = 0.0055). When grouped by BED, analysis showed that the pooled OR was 0.40 (95% CI 0.13–1.22, z = −1.61, p = 0.11) in patients who received a BED greater than 35 Gy and 0.64 (95% CI 0.33–1.24, z = −1.32, p = 0.19) in those who received a BED of 35 Gy or less without a significant interaction (χ² for interaction = 0.03; 1 df; p = 0.86).

**Downstaging effect**

A total of four RCTs involving 573 patients were available for evaluating the downstaging effect of CRT (fig 2). Patients who received preoperative CRT were less likely to have an advanced stage of cancer at pathological examination than were controls. Preoperative CRT was superior to surgery alone in all studies, reaching statistical significance in two RCTs. The pooled OR was 0.43 (CI 0.26–0.72, z = −3.26, p = 0.001) (NNT = 5). Similar results were obtained when a fixed effect model was used (OR 0.44 (95% CI 0.31–0.62); z = −4.69, p = 0.00001). In all the robust analyses the pooled estimate of the treatment effect was significant.

**Compliance, postoperative complications, and mortality**

Compliance with treatment was generally satisfactory. Forty-five (11.7%) of 385 patients undergoing adjuvant treatment did not complete the planned protocol, and only 25 (6.5%) required a reduction in chemotherapeutic (21 patients) or irradiation dose (four patients).

The overall rate of postoperative adverse events was 39.4% (137/348) in the CRT group and 34.3% (123/358) in the surgery alone group (χ² = 1.90; df = 1; p = 0.16). The three most frequent adverse events were respiratory complications (19.9%), heart failure (6.9%), and anastomotic leak (6.9%).

The risk of postoperative mortality (within 90 days) was higher in the CRT group in five RCTs reaching statistical significance in one of these five trials in which a high dose per fraction radiation (3.7 Gy) was given (fig 3).
Combining the data from the six RCTs, which included 683 patients, showed a significant effect of CRT on postoperative mortality (OR 2.10 (95% CI 1.18–3.73); z = 2.53, p = 0.01) (NNH = 25). Similar results were obtained when a fixed effect model was used (OR, 2.15 (95% CI 1.23–3.74); z = 2.69; p = 0.007).

Robust analysis showed a loss of significance for postoperative mortality (OR 1.62 (95% CI 0.81–3.24); z = 1.36, p = 0.17) once the largest trial by Bosset and colleagues was excluded. Moreover, sensitivity analysis performed by excluding the two RCTs with a fraction per day dose >2 Gy showed a similar effect size as the robust analysis without statistical significance (OR 1.44 (95% CI 0.67–3.08) z = 0.94; p = 0.35).

**DISCUSSION**

This study has investigated the key clinical question of whether preoperative CRT is efficacious in treating oesophageal cancer. To our knowledge, no consensus on the type of patients amenable to chemoradiotherapy has been reached to date.

This meta-analysis of data from six RCTs showed that in resectable oesophageal cancer, preoperative CRT significantly improves three year survival versus surgery alone (NNT = 10). Moreover, an impressive reduction in the rate of advanced oesophageal cancer (stages IIB and III) was observed in almost all trials at the time of surgery (downstaging) (NNT = 5). Although there is evidence that chemoradiotherapy significantly increases postoperative mortality (NNH = 25), fewer patients need to be treated to benefit from the treatment long term than need to be treated to be harmed immediately post surgery.

The effect of preoperative CRT on overall survival is much more pronounced and statistically significant in patients with adenocarcinoma, that is now more prevalent than squamous cell carcinoma in the USA and Western Europe, with most tumours located in the distal oesophagus. However, the sample size of this subgroup analysis was small (data obtained from only two trials), and caution must be exercised when interpreting results from this exploratory analysis. Nevertheless, our data strongly indicate the need for designing future trials considering the clinical difference between adenocarcinoma and squamous cell carcinoma and its potential influence on patient response to therapy. Furthermore, the current staging system for oesophageal cancer is mostly applicable to patients with squamous cell carcinomas of the upper and middle thirds of the oesophagus. To better evaluate the differential therapeutic response to CRT of patients with adenocarcinoma, this staging system should be adapted to better fit patients with adenocarcinoma, who most often present with distal oesophageal and gastro-oesophageal junction tumours. In particular, patients with regional and/or coeliac axis lymphadenopathy should not necessarily be considered to have unresectable disease due to metastases.

Many efforts have been made to identify the optimal chemoradiotherapy regimen that would increase the cost effectiveness of therapy. There was considerable variation in the irradiation procedures, suggesting that worldwide accepted and standardised radiation techniques are needed to obtain comparable data on the efficacy and safety profile, particularly regarding the total and daily dose of radiation and the dose of cisplatin administered. We found that higher postoperative mortality was observed in the two RCTs in which a fraction per day dose >2 Gy was delivered. Furthermore, the risk of postoperative mortality was higher in the RCT by Bosset and colleagues in which a high dose of both radiotherapy and cisplatin was administered. On the other hand, the hyperfractionated irradiation schedule combined with a low dose of cisplatin administered in the trial by Urba and colleagues was more safe than the others, suggesting that a reduction of late effect achieves the best cost/effectiveness ratio. Finally, the trial by Urba and colleagues was the only one in which a transhiatal oesophagectomy was employed.

We believe the available information is inadequate to determine whether a concomitant regimen of chemoradiotherapy is better than induction chemotherapy followed by radiotherapy. Among the trials analysed, only that of Nygaard and colleagues delivered radiotherapy three weeks after completion of chemotherapy. The trials of Le Prise and colleagues and Bosset and colleagues delivered radiotherapy within one week after chemotheraphy (intermittent chemotherapy) while CRT was administered simultaneously in the remaining three RCTs. Two of these three RCTs (by Walsh and Urba) showed the highest therapeutic benefit. However, it has been postulated that induction chemotherapy may be more effective than the concomitant approach on the premise that areas of radiochemotherapy may harbour treatment resistant tumour cells and that combined toxic effects may limit the dose of drugs that can be given. Firm conclusions on the results of direct comparisons between chemoradiotherapy delivered sequentially or concomitantly are hampered by the fact that to date no trial has been performed to evaluate this variable.

Recently, a meta-analysis of individual patient data failed to show a statistically significant benefit of preoperative radiotherapy alone on survival. Moreover, the most recent large scale RCTs of preoperative chemotherapy as a single adjuvant treatment showed conflicting results. Our meta-analysis clearly shows that CRT as neoadjuvant chemotherapy as a single adjuvant treatment improves survival. We speculate that chemotherapy enhances the local effects of radiotherapy and thus decreases the likelihood of spread from the primary tumour prior to exposure of the patient to the tumour growth promoting stimulus of surgery.

The results of this retrospective analysis are subject to several limitations. Differences in the baseline severity of illness in the population of the RCTs, in the irradiation techniques, and in the chemoradiotherapy regimens may limit the accuracy of this meta-analysis. These summary results describe only between study, not between patient, variation because they reflect group averages rather than individual data. Lack of data on other potential confounders such as size and location of the tumour could also affect the accuracy of the results. The meta-analysis was performed using summary data, and more detailed treatment comparisons could be achieved with a meta-analysis of individual patient data.

Screening of the non-English literature, the extensive manual and computer searches for studies, in addition to the personal contacts made directly with principal investigators, make us confident that no important published trials were overlooked. Publication bias was probably not substantial and considered unlikely to change the direction of our pooled estimate of treatment effect. We should be particularly concerned about publication bias in settings in which small studies are being conducted.

Finally, the standard approach to inference for random effects model used in this meta-analysis including few RCTs can inflate the type I error rate.

The available evidence is sufficient to conclude the following: (1) preoperative CRT reduces the three year overall mortality compared with surgery alone. The magnitude of the overall effect is clinically relevant. Further large scale multicentre RCTs may prove useful to substantiate the benefit on overall survival; (2) the magnitude of the downstaging effect with CRT was large; and (3) postoperative mortality
was significantly increased by CRT. Studies addressing strategies that could potentially reduce the toxicity profile would have major implications for patients affected by oesophageal cancer.

Further RCTs in patients with oesophageal cancer investigating the efficacy and safety of induction chemotherapy in addition to preoperative chemoradiotherapy are underway by the RTOG Gastrointestinal Cancer Committee.

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