Comparison of different neoadjuvant treatments for resectable locoregional esophageal cancer: A systematic review and network meta-analysis

Yongxing Bao, Zeliang Ma, and Meng Yuan have contributed equally to this work.

Abstract

Purpose: The best pattern of neoadjuvant therapy for resectable locoregional esophageal cancer has not been determined. Our study evaluated the efficacy and postoperative events of different treatments using the Bayesian network meta-analysis.

Methods: We systematically tracked randomized clinical trials from the Medline, EMBASE, and Cochrane Library databases. The following treatments were included: neoadjuvant chemoradiation followed by surgery (NCRT + S), neoadjuvant chemotherapy followed by surgery (NCT + S), neoadjuvant radiotherapy followed by surgery (NRT + S), and surgery alone (S). The Revised Cochrane risk-of-bias tools were used to assess the quality of included trials. Overall survival (OS) and progression-free survival or disease-free survival (PFS/DFS) were assessed through hazard ratios (HR). Locoregional recurrence, distant metastasis, postoperative mortality, and postoperative morbidity were assessed through odds ratios (OR). These outcomes were compared between different treatments through Bayesian network meta-analysis.

Results: Twenty trials with 4384 patients were included. Compared with S, only NCRT + S could significantly improve OS for patients with esophageal cancer (HR = 0.78, 95% confidence interval [CI] 0.68–0.88). NCRT + S and NCT + S significantly improved PFS/DFS compared with S (NCRT + S vs. S, HR = 0.72, 95% CI 0.63–0.81; NCT + S vs. S, HR = 0.81, 95% CI 0.69–0.97). NCRT + S significantly reduced both locoregional recurrence (OR = 0.67, 95% CI 0.51–0.88) and distant metastasis (OR = 0.63, 95% CI 0.45–0.90) compared with S. There were no differences in postoperative morbidity between the four treatments. However, NCRT + S also increased postoperative mortality compared with S (OR = 1.77, 95% CI 1.09–2.82) and NCT + S (OR = 1.96, 95% CI 1.11–3.51).

Conclusion: NCRT + S is the most efficient neoadjuvant treatment for resectable locoregional esophageal cancer. However, NCRT + S increases the risk of postoperative mortality but not morbidity.

KEYWORDS
esophagus cancer, neoadjuvant chemotherapy, neoadjuvant radiotherapy, network meta-analysis
INTRODUCTION

Esophageal cancer leads to a poor prognosis and is the sixth most common cause of cancer-related deaths in the world.1 Surgery is an important treatment for resectable esophageal cancer, but the survival of patients receiving surgery alone is unfavorable, with a 5-year overall survival (OS) of 15–25%.2 Neoadjuvant therapy has been a vital pattern before surgery to eliminate micro-metastatic disease, reduce tumor burden, and improve surgery compliance, finally leading to better postoperative locoregional control, distant control, and OS.

However, the best pattern of neoadjuvant therapy has not been determined. Many randomized control trials have shown the significant advantage of neoadjuvant chemoradiotherapy followed by surgery (NCRT+S) compared with surgery alone,3–5 including a large phase III study from China.6 In addition, other research has shown the benefits of neoadjuvant chemotherapy followed by surgery (NCT+S) and neoadjuvant radiotherapy followed by surgery (NRT+S) compared to surgery alone.7–9 However, few studies have compared different neoadjuvant treatments to distinguish the best neoadjuvant therapies. Four previous randomized trials compared NCRT+S and NCT+S, and only one obsolete trial demonstrated the benefit of NCRT+S beyond NCT+S with just a positive p value10–13; no trials have directly compared NRT+S with other neoadjuvant treatments.

Previous network meta-analysis (NMA) suggested that NCRT+S might improve OS more than the other treatments, with some defects and insufficient evidence.14,15 Several randomized trials have reported their results in recent years,6,11,12 so the NMA needed to be updated.

The optimal choice among NCRT+S, NCT+S, and NRT+S remained unclear, and high-quality evidence is needed to direct clinicians. Our study evaluates the effect of neoadjuvant treatments for esophageal cancer and determines the best treatment pattern using the Bayesian NMA.

PATIENTS AND METHODS

Systematic review

Searching strategy

This study was performed following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses 2020 (PRISMA 2020) statement for network meta-analysis (Supplement 1 in Appendix S1). This NMA was registered in the Prospective Register of Systematic Reviews (Identification code: CRD42020170734).

Eligible data from the Medline, EMBASE, and Cochrane Library databases were systematically searched until April 30, 2021. The search terms are shown in Supplement 2 in Appendix S1.

Inclusion and exclusion criteria

The inclusion criteria were as follows:

1. Randomized clinical trials (RCTs).
2. Studies only including patients with esophagus cancer or esophagogastric cancer.
3. Full text published in English.
4. Studies comparing different neoadjuvant treatments or surgery alone.

The exclusion criteria were as follows:

1. Poor-qualities studies were defined by:
   1. Studies reporting Kaplan–Meier curves without censor, whether reporting censor or not in addition.
   2. Studies not reporting Kaplan–Meier curves or estimated hazard ratios (HR).
   3. Studies including sample size of less than 50.
   4. Studies with nonstandard and heterogeneous neoadjuvant treatments such as hyperthermia and radiation at low doses (below 30 Gy).
   5. Studies on perioperative and intraoperative treatments.

If several of the same trial publications were retrieved, the most recent information was extracted from these publications.

Data extraction and calculation

Two reviewers independently perused the full text of references meeting the aforesaid criteria. The discrepancies of the two reviewers were resolved through discussion, including a third reviewer. Outcomes assessed from the trials included OS, progression-free survival or disease-free survival (PFS/DFS), postoperative mortality, postoperative morbidity, locoregional recurrence (LRR), and distant metastases (DM). Postoperative mortality was defined as mortality within 1 or 3 months, or mortality during hospitalization after surgery. Postoperative morbidity was defined as surgery complications during hospitalization such as hemorrhage, heart failure, and fistula, and was calculated according to the number of incurred patients.

The OS and PFS/DFS were assessed by HRs and their standard error, which was extracted or calculated through the following approaches:

1. For studies directly reporting the HR and its 95% confidential interval (CI), the HR was collected and its standard error (SE) was calculated from the 95% CI.
2. For studies not reporting the HR but having a survival curve with an at-risk table, the HR and its SE were calculated using methods outlined by Parmar et al.16 The number of events and censors was extracted from the survival curve for every interval through WebPlotDigitizer version 4.2 software. Finally, the HR and its SE were calculated by combining all time intervals.
3. For studies not reporting the HR or at-risk tables but containing survival curves and follow-up information, the estimate of censor was approximated based on a linear pattern. Then the HR was calculated as the second approach.

The other outcomes were assessed by odd ratios (ORs) and their standard error, calculated from the number of events and sample size.

Quality assessment

Two reviewers independently assessed the quality of included trials through the revised Cochrane risk-of-bias tool for randomized trials (RoB 2). The discrepancies were resolved through discussion, including a third reviewer. The assessment scale was as follows: randomization process, deviations from intended interventions, missing outcome data, measurement of the outcome, selection of the reported result, and overall bias, with each domain judged as any of the three levels: low, high, or unclear.

Data synthesis and statistical analysis

Bayesian NMA was carried out to synthesize all therapeutic options within a mixed treatment comparison framework. The random-effects model was prioritized to address the trial-specific effects, which were components of the overarching distribution. Uninformative prior distribution was given to all parameters. The node-split method was used to assess the inconsistency. The estimates of relative effects and 95% CI were reported. The surface under the cumulative ranking (SUCRA) scores was also calculated. All statistical analysis was conducted through R 3.6.0.
RESULTS

Trial characteristics

In total 20 trials with 4384 patients were included in the final quantitative NMA. The search and selection process are displayed in a PRISMA 2020 flow diagram in Figure 1. The basic characteristics of the trials included are shown in Table 1. All trials were comparable in terms of clinical features, and the assumption of transitivity was acceptable. The network plot for treatment comparison of OS is shown in Figure 2, and the plots for other outcomes are shown in Supplement 4 in Appendix S1.

Quality assessment

The summary of the risk-of-bias assessment is shown in Supplement 3 in Appendix S1. No RCT had a high risk of bias. As expected, the blinding of participants was not explicitly indicated in most studies, especially in those involved in radiotherapy. We believe that it was unlikely that deviations would arise. However, the selection bias might exist, as the allocation concealment was not explicitly mentioned in most studies.

Inconsistency assessment

Four arms of different treatments were categorized to make comparisons. The node-split analyses for OS and other outcomes are shown in Supplement 5 in Appendix S1. There were no significant differences between direct and indirect evidence in all comparisons for all outcomes (p > 0.05), except for NRT + S versus NCT + S (p = 0.047) in the subgroup analysis of squamous cell carcinoma (SCC) for OS.

TABLE 1  Trial characteristics

| Trials     | Year | Designs          | N  | Histology | Location                  | Preoperative stage | RT schedule         | CT schedule | CT cycles |
|------------|------|------------------|----|-----------|---------------------------|--------------------|---------------------|-------------|-----------|
| Gignoux    | 1987 | NRT + S vs. S    | 208| SCC       | Thoracic esophagus        | T1-3N0-1M0         | 33 Gy/3.3 Gy/10f    | —           | —         |
| Nygaard    | 1992 | NCRT + S vs. NCT + S vs. NRT + S vs. S | 186| SCC       | Thoracic esophagus (below 21 cm) | T1-2N0-1M0         | 35 Gy/1.75 Gy/20f   | PB          | 2         |
| Walsh      | 1996 | NCRT + S vs. S   | 113| AC        | Esophagus (excluding cervical esophagus) | NA                | 40 Gy/2.67 Gy/15f   | PF          | 2         |
| Bosset     | 1997 | NCRT + S vs. S   | 282| SCC       | Thoracic esophagus        | I-II               | 37 Gy/3.7 Gy/10f    | P           | 2         |
| Law        | 1997 | NCRT + S vs. S   | 147| SCC       | Thoracic esophagus        | NA                 | —                   | PF          | 2         |
| Ancona     | 2001 | NCT + S vs. S    | 94 | SCC       | Esophagus                 | IIA-III            | —                   | PF          | 2         |
| Urba       | 2001 | NCRT + S vs. S   | 100| AC, SCC   | Esophagus                 | NA                 | 45 Gy/1.5 Gy/30f    | PFV         | 2         |
| Lee        | 2004 | NCRT + S vs. S   | 101| SCC       | Thoracic esophagus        | IIA-III            | 45.6 Gy/1.2 Gy/38f  | PF          | 2         |
| Burmeister | 2005 | NCRT + S vs. S   | 256| AC, SCC   | Thoracic esophagus        | T1-3N0-1M0         | 35 Gy/2.3 Gy/15f    | PF          | 1         |
| Kelsen     | 2007 | NCRT + S vs. S   | 243| AC, SCC   | Esophagus or GEJ          | I-III              | —                   | PF          | 3         |
| Tepper     | 2008 | NCRT + S vs. S   | 56 | AC, SCC   | Thoracic esophagus        | T1-3N0-1M0         | 50.4 Gy/1.8 Gy/28f  | PF          | 2         |
| Allum      | 2009 | NCT + S vs. S    | 802| AC, SCC, UC | Esophagus or GEJ         | NA                 | —                   | PF          | 2         |
| Lu         | 2010 | NCRT + S vs. S   | 238| SCC       | Thoracic esophagus        | IIA-III            | 40 Gy/2 Gy/20f      | TP          | 2         |
| Bonstra    | 2011 | NCT + S vs. S    | 169| SCC       | Thoracic esophagus        | I-III or M1a       | —                   | EP          | 2–4       |
| Burmeister | 2011 | NCRT + S vs. NCRT + S | 75 | AC        | Esophagus or GEJ          | IIA-III            | 35 Gy/2.3 Gy/15f    | PF          | 2         |
| Maritte    | 2014 | NCRT + S vs. S   | 195| AC, SCC   | Thoracic esophagus        | I-II               | 45 Gy/1.8 Gy/25f    | PF          | 2         |
| Shapiro    | 2015 | NCRT + S vs. S   | 368| AC, SCC, UC | Esophagus or GEJ          | T1N1M0 or T2–3N0–1M0 | 41.4 Gy/1.8 Gy/23f | TP          | 5*        |
| Stahl      | 2017 | NCRT + S vs. NCT + S | 119| AC        | GEJ                       | T3-4NxM0           | 30 Gy/2 Gy/15f      | PFFo/EP     | 5*        |
| Yang       | 2018 | NCRT + S vs. S   | 451| SCC       | Thoracic esophagus        | T1-4N1M0 or T4N0M0 | 40 Gy/2 Gy/20f      | NP          | 2         |
| VonDobeln  | 2019 | NCRT + S vs. NCT + S | 181| AC, SCC   | Esophagus or GEJ          | T1N1, T2-3N0-1 or M0-M1a | 40 Gy/2 Gy/20f | PF          | 3         |

Abbreviations: AC, adenocarcinoma; CT, chemotherapy; EP, cisplatin and etoposide; f, fractions; GEJ, gastroesophageal junction; NA, not available; NP, cisplatin and vinorelbine; P, single drug cisplatin; PB, cisplatin and bleomycin; PF, cisplatin and fluorouracil; PFo, cisplatin, fluorouracil, and folinic acid; PFV, cisplatin, fluorouracil, and vinblastine; RT, radiotherapy; S, surgery; SCC, squamous cell carcinoma; TP, cisplatin and paclitaxel; UC, undifferentiated carcinoma.

*aThe chemotherapy was given weekly.
Results of NMA

Survivals

The pooled treatment effects of OS for all the evaluated treatment options are shown in Table 2a. Compared with S, NCRT + S could significantly improve the OS for patients with esophageal cancer (HR = 0.78, 95% CI 0.68–0.88), while NCT + S and NRT + S failed to significantly improve the OS (NCT + S vs. S, HR = 0.90, 95% CI 0.78–1.02; NRT + S vs. S, HR = 0.86, 95% CI 0.65–1.10). Compared with NCT + S, NCRT + S failed to improve the OS with statistical significance (HR = 0.91, 95% CI 0.69–1.21). The ranking analysis based on SUCRA scores showed that NCRT + S was the most likely to be the best option in terms of OS benefit (Supplement 6a in Appendix S1).

In total 14 studies and three arms (NCRT + S, NCT + S, and S) were available for PFS/DFS analysis. The treatment effects of PFS/DFS are shown in Table 2b. Compared with S, NCRT + S and NCT + S significantly improved the PFS/DFS for patients with esophageal cancer (NCRT + S vs. S, HR = 0.72, 95% CI 0.63–0.81; NCT + S vs. S, HR = 0.81, 95% CI 0.69–0.97). NCRT + S improved the PFS/DFS from NCT + S without statistical significance (HR = 0.88, 95% CI 0.73–1.06). NCRT + S also reached the top of SUCRA scores in all treatments (Supplement 6b in Appendix S1).

Postoperative events

Eighteen studies were available for postoperative mortality and eight for postoperative morbidity. The treatment effects are shown in Table 2c,d. NCRT + S significantly increased postoperative mortality compared to S (OR = 1.77, 95% CI 1.09–2.82) and NCT + S (OR = 1.96, 95% CI 1.11–3.51). However, no significant difference in postoperative morbidity was detected between the four treatments. NCRT + S reached the lowest SUCRA scores of postoperative mortality and morbidity in all treatments (Supplement 6c,d in Appendix S1).

Failure patterns

Fourteen studies were available for LRR and DM. The treatment effects are shown in Table 2e,f. LRR was significantly lower in the NCRT + S group than in the S group (OR = 0.49, 95% CI 0.31–0.74). Compared with the S group, DM was significantly reduced in both the NCRT + S group and the NCT + S group (NCRT + S vs. S, OR = 0.67, 95% CI 0.51–0.88; NCT + S vs. S, OR = 0.63, 95% CI 0.45–0.90). No significant difference was detected between the other groups. NCRT + S reached the top of the SUCRA scores of both LRR and DM in all treatments (Supplement 6e,f in Appendix S1). Although NRT + S failed to reach significant differences in treatment effect analyses, it was the second group in ranking analysis for LRR and DM.

Subgroup analyses of pathological types

Fourteen studies were available for SCC and eight for adenocarcinoma (AC). The treatment effects of OS are shown in Table 3. For patients with SCC, both NCRT + S and NCT + S could significantly improve OS compared with S (NCRT + S vs. S, HR = 0.76, 95% CI 0.64–0.90; NCT + S vs. S, HR = 0.83, 95% CI 0.71–0.99). No significant difference was detected between the other groups. For patients with AC, no significant difference in OS was detected between the treatment groups, but both NCRT + S and NCT + S were apparently associated with better OS over S (NCRT + S vs. NCT + S, HR = 0.79, 95% CI 0.60–1.04; NCT + S vs. S, HR = 0.88, 95% CI 0.62–1.28). NCRT + S reached the top of the SUCRA scores in all treatments for both SCC and AC (Supplement 6g,h in Appendix S1).

DISCUSSION

This NMA of 4384 patients in 20 RCTs demonstrated that NCRT + S was the most effective neoadjuvant therapy pattern for resectable esophageal cancer. NCRT + S increased OS and PFS/DFS, and decreased LRR and DM compared with the other treatments without expanding the probability of postoperative morbidity. NCRT + S was the most
Effective treatment for both SCC and AC, although the OS effect over S was not statistically significant for AC. This NMA provides the best available evidence for the use of neoadjuvant CRT in esophageal cancer.

Several high-quality RCTs reported the efficiency and postoperative events of NCRT + S compared with S. The NEOCRTEC5010 study published in 2018 reported the OS and DFS benefits by NCRT + S compared with surgery alone in squamous esophageal cancer. The classical CROSS study also demonstrated the efficiency of NCRT + S over surgery alone. Our study confirms the role of NCRT in the neoadjuvant treatments of esophageal cancer.

While the survival benefit of NCRT over surgery alone has been established, the relative merits of radiotherapy and chemotherapy have not been determined. The long-term results of the POET study showed a borderline OS

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**TABLE 2 The treatment effect for all the evaluated treatment options**

|                | NCRT + S | NCT + S | NRT + S | S       |
|----------------|----------|---------|---------|---------|
| **a. Overall survival** | 1.15 (0.98, 1.35) | 1.1 (0.83, 1.45) | 1.29 (1.14, 1.47) | 1.29 (1.14, 1.47) |
| **b. Progression-free survival or disease-free survival** | 1.14 (0.95, 1.38) | 1.4 (1.24, 1.58) | 1.23 (1.03, 1.46) | 1.57 (1.35, 1.72) |
| **c. Postoperative mortality** | 0.9 (0.78, 1.02) | 0.86 (0.65, 1.1) | 0.9 (0.78, 1.02) | 0.9 (0.78, 1.02) |
| **d. Postoperative morbidity** | 0.78 (0.68, 0.88) | 0.72 (0.63, 0.81) | 0.51 (0.28, 0.9) | 0.57 (0.35, 0.92) |
| **e. Locoregional recurrence** | 1.47 (0.86, 2.43) | 1.23 (0.34, 4.49) | 0.61 (0.19, 2.09) | 0.61 (0.19, 2.09) |
| **f. Distant metastases** | 1.6 (1.11, 2.21) | 1.07 (0.77, 1.43) | 0.87 (0.4, 1.95) | 0.87 (0.4, 1.95) |

Note: Statistically confident results are in bold. Abbreviations: NCRT + S, neoadjuvant chemoradiotherapy followed by surgery; NCT + S, neoadjuvant chemotherapy followed by surgery; NRT + S, neoadjuvant radiotherapy followed by surgery; S, surgery.
Our study and previous meta-analyses did not find survival advantage for NCRT + S over NCT + S (p = 0.055). Another two phase II studies also failed to reach a statistically significant survival advantage for NCRT over NCT.\textsuperscript{10,12}

Our study and previous meta-analyses did not find survival differences between NCRT + S and NCT + S. The possible reason for this is that only a few trials directly compared NCRT + S and NCT + S (four studies with 472 patients in our study; Table 1). Further studies still need to be conducted to compare the efficiency of NCRT and NCT.

Interestingly, our study found that NCRT + S significantly increased the postoperative mortality of esophageal cancer compared with NCT + S or S alone. This result was consistent with a previously published NMA.\textsuperscript{14} The total postoperative deaths were 64 in the NCRT + S group and 35 in the S group. The main reasons for postoperative death in the NCRT + S group were respiratory complications (41% of all cases where specific complications were reported), severe infection and sepsis (23%), and anastomotic leak (17%). However, NCRT + S did not increase the postoperative morbidity compared with either NCT + S or S. Among the included 13 trials of NCRT + S, only one trial published in 1997 reported that the postoperative complications of NCRT + S were significantly higher than for S. The other 12 trials (including the CROSS and NEOCRT-TEC5010 studies) reported no significant difference between NCRT + S and other treatments. We further recalculated the postoperative morbidity as the number of events instead of incurred patients. We found no significant differences between the four treatments, consistent with postoperative morbidity calculated with the number of incurred patients. The possible reason for this could be that NCRT + S increased the severity of postoperative complications instead of the number. Postoperative complications should therefore be monitored and treated promptly when the patient has been treated with NCRT. Another interesting result of our study is that NCRT + S could significantly reduce DM compared with NCT + S. This could be because of the synergistic effect of radiotherapy and chemotherapy. Radiation could reduce tumor volume and eliminate tumor cells in local vessels, which could help concurrent chemotherapy to reduce DM.

Chan et al. reported another NMA of neoadjuvant therapy in 2018.\textsuperscript{14} The results also showed that NCRT + S was the most efficient treatment for esophageal cancer. However, this NMA had some drawbacks. The radiation dose of some included trials was 20Gy, which was insufficient. In addition, several exploratory and small-sized trials were included in the final analysis, reducing the homogeneity and credibility of results. Finally, this study did not analyze postoperative morbidity, and the results of postoperative mortality were uncertain (NCRT + S might increase the risk of postoperative mortality in esophageal cancer patients). Our NMA avoided these points, and added one published trial and two updated trials to expand the evidence base. Studies with nonstandard treatments or lower quality were excluded as the exclusion criteria mentioned above. Our results confirmed that NCRT + S increased postoperative mortality compared with NCT + S or surgery alone. We further analyzed postoperative morbidity.

There are a few limitations of this NMA. First, this analysis used published data instead of individual patient data, where some uncontrolled confounding factors could affect the results. Second, the sample sizes for NRT + S arm were limited with two studies (published in 1987 and 1992) of 150 patients and the radiotherapy techniques in the two studies were outdated traditional two-fields radiation. However, we could not find a trial with NRT + S using modern techniques, so we had to include the outdated studies. Thus, the results of NRT + S need to be treated with caution because of uncertainties and the wide 95% CIs. Third, the variability of included trials in patient populations, treatments, and procedures should not be ignored, even though the tests for inconsistency were negative. One of the important confounding factors was the location of esophageal cancer (gastroesophageal junction, EGJ or thoracic esophagus), which might induce a different response to neoadjuvant
Our study could not estimate the effect of localization on different neoadjuvant treatments because of insufficient evidence. Fourth, our analysis did not distinguish between chemoradiotherapy regimens or radiation doses. In most trials, platinum-based chemotherapy and a total radiation dose of 40–50 Gy were used. Finally, neoadjuvant immunotherapy was not included in this study. Although studies on neoadjuvant immunotherapy were in full flow, most were nonrandomized controlled studies and had not yet reported mature survival data.

In conclusion, NCRT + S could be the most efficient neoadjuvant treatment for resectable locoregional esophageal cancer. However, NCRT + S could increase the risk of postoperative mortality but not postoperative morbidity, and clinicians must be cautious in postoperative management.

**AUTHOR CONTRIBUTIONS**

Y.B., Z.M., M.Y., and Z.H. contributed to the conception and design of the study. Y.B., Z.M., and M.Y. performed the searches and collected raw data. Y.W. performed the statistical analysis. Y.B. wrote the main part of the manuscript. M.Y. and Z.M. wrote part of the manuscript. Y.M. and Z.H. reviewed and edited the manuscript. All authors contributed to the manuscript revision and approved the submitted version.

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**DATA AVAILABILITY STATEMENT**

All data were extracted from published articles.

**ORCID**

Yongxing Bao https://orcid.org/0000-0002-4178-4046
Zhouguang Hui https://orcid.org/0000-0002-7189-4692

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SUPPORTING INFORMATION
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