Towards personalized treatment of T2N0 rectal cancer: A systematic review of long-term oncological outcomes of neoadjuvant therapy followed by local excision

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Abstract

Background and Aim: Total mesorectal excision (TME) remains the treatment of choice in T2N0 tumors. However, evidence suggest that one-size-fits-all approach is not always beneficial for this group of patients. The aim of this study is to synthesize data on long-term outcomes after neoadjuvant therapy (NAT) followed by local excision (LE) in T2N0 rectal cancer patients in the perspective of a rectal-preserving strategy.

Methods: A systematic search of PubMed/MEDLINE, SCOPUS, and Web of Science databases was conducted until October 2021 to identify studies comparing LE after NAT and TME or reporting oncologic outcomes after conservative approach. A pooled analysis was conducted using a fixed-effect model in the case of non-significant heterogeneity (P > 0.1), and a random effect model (DerSimonian–Laird method) when significant heterogeneity was present (P < 0.1) CRD42022300344.

Results: Nine studies were included in the analysis. Three of them were comparative studies. The pooled 3-year DFS, 5-year DFS, 3-year OS, 5-year OS, local and distant recurrence rates were 92.8% (95% CI 81.6–99.5%), 91.3% (95% CI 88.3–94.3%), 96.1% (95% CI 90.5–100%), 72.6% (95% CI 57.5–87.7%), 4% (95% CI 18–63%), and 4.9% (95% CI 2–7.8%), respectively, in subjects treated with NAT followed by LE. No heterogeneity was found for all these analyses, except for the 5-year OS sub-analysis (P 95.5%, P < 0.001). Complete pathological response (ypT0) rate after NAT and LE ranges from 26.7% to 59%.

Conclusion: LE following neoadjuvant CRT may provide comparable survival benefit to radical surgery for patients with clinical stage T2N0 in selected patients although the evidence is still limited to provide solid recommendations. A personalized therapeutic approach taking into account tumor and patient-related factors should be considered.

Introduction

Early rectal cancer is defined as a cancer with good prognostic features that might be safely removed by transanal local excision (LE) preserving the rectum and that will have a very limited risk of relapse.1 According to the Association of Coloproctology of Great Britain & Ireland (ACPGBI), cT1-2N0M0 tumors are included in the early stage of rectal cancer.2 However, it is well established that LE has curative role only in T1 tumors with favorable pathologic characteristics (low-risk pT1) while radical surgery remains the treatment of choice in T2 rectal cancer because of a not negligible risk of local recurrence (from 26% to 47%) and occult nodal disease.3,4

Despite the benefits of the minimally invasive approaches for the surgical treatment of rectal cancer,5 an high risk of perioperative complications, permanent stoma, and functional impairments are still associated with radical surgery.6–8 In order to decrease morbidity related to major surgery, the use of organ-preserving strategies based on a multidisciplinary approach is gaining support among surgeons.9,10 Indeed, the possibility of avoiding a Total mesorectal excision (TME) with neoadjuvant chemoradiotherapy (CRT) followed by LE has been evaluated in the setting of clinical trials11,12 with similar oncological outcomes, or for patients unfit for surgery or refusing permanent ostomy.

T2 rectal cancer patients are probably the best candidate for this approach because smaller and more superficial cancers are more likely to exhibit better response to CRT and they are less likely to develop a tumor regrowth than ≥ cT3 tumors once they have achieved a complete clinical response.13,14 Downstaging and downsizing of the rectal lesion provide the opportunity to subsequently perform an LE to obtain a pathological evaluation of residual tumor with a low risk of nodes involvement.

Key words
local excision, neoadjuvant therapy, organ preservation, rectal cancer, T2 tumor.

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Previous systematic reviews evaluated outcomes of LE after CRT\textsuperscript{15–17}; however, they included studies with any preoperative tumor stage and merged results. Therefore, we aim to conduct a systematic review of the literature in order to assess long-term outcomes of LE following CRT in T2 rectal cancer patients exclusively. Evaluating the safety and effectiveness of the conservative management for cT2 rectal tumors could help shift the paradigm in favor of less invasive procedures without the postoperative inconveniences of major surgery.

**Methods**

**Literature search and selection of primary studies.** A systematic review of the existing literature was conducted in accordance with the preferred reporting items for systematic reviews and meta-analyses (PRISMA) guidelines\textsuperscript{18} to establish the evidence base regarding the use of neoadjuvant CRT followed by LE for the treatment of cT2 rectal tumors.

The systematic literature search was performed in PubMed/Medline, Scopus, and Web of Science databases to identify studies reporting oncologic outcomes from the beginning of indexing for each database till October 31, 2021. Bibliographic review of selected articles was assessed as secondary sources for full-length articles of studies. A literature search was performed using the following index terms: “T2 rectal cancer,” “early rectal cancer,” “neoadjuvant therapy,” “preoperative chemoradiotherapy,” “local excision,” “transanal endoscopic microsurgery,” “transanal minimally invasive surgery,” and “transanal excision.”

**Eligibility criteria.** Two reviewers (R. P. and M. M. D. N.) independently evaluated all the studies retrieved according to the eligibility criteria and any differences between the datasets were resolved by discussion. Studies were included if they met the following criteria: evaluation of oncologic outcomes in terms of recurrence rate or survival after neoadjuvant CRT followed by LE in non-comparative or in comparative studies with conventional treatment for cT2 rectal cancer. We excluded the articles if there was no sufficient documentation, if data were combined with those of other tumor stages, if sample size was ≤ 10 patients and if they were in languages other than English. Narrative reviews, duplicate publications, editorials, and abstracts were also excluded.

**Data extraction and management.** Data were extracted independently and entered into standardized Excel spreadsheets (Microsoft Inc., Redmond, Washington, USA). Data were presented as frequencies and percentages. Any disagreements were resolved through discussion. The following data were extracted from each study: first author, study period, study design, inclusion and exclusion criteria, number of participants, mean age, type of surgery ad platforms for LE, surgery indication, neoadjuvant regimen, ypT0 rate, survival, recurrence, and mortality rates. Study outcomes included disease-free survival or overall survival and local o distant recurrence rate.

**Assessment of the methodological quality of studies.** All studies were assessed for methodological quality. For randomized studies, the validated score described by Jadad et al.\textsuperscript{19} was used. The scale consists of three items pertaining to descriptions of randomization, masking, and dropouts and withdrawals in the report of an RCT. The scale ranges from 0 to 5, with higher scores indicating better reporting. High-quality trials scored more than 2 out of a maximum possible score of 5. Low-quality trials scored 2 or less out of a maximum possible score of 5. The Methodologic Index for Nonrandomized Studies (MINORS)\textsuperscript{20} tool was also used to assess quality of the studies included in the review. For noncomparative studies, a maximum score of 16 could be achieved using the MINORS tool, with a maximum score of 24 available for comparative studies.

**Statistical analysis.** Statistical analyses were performed by using Comprehensive Meta-analysis Software version 3.0 (Biostat, Englewood, New Jersey, USA). Heterogeneity was assessed by using chi-squared statistics and $I^2$ measure of inconsistency.

A pooled analysis was conducted using a fixed-effect model in the case of non-significant heterogeneity ($P > 0.1$), and a random effect model (DerSimonian–Laird method) when significant heterogeneity was present ($P < 0.1$). Corresponding forest plots were constructed for the pooled estimates of the abovementioned outcomes and weight of individual studies are represented by the size of individual squares.

A $P$ value < 0.05 was considered statistically significant for all outcomes.

**Results**

Figure 1 shows the PRISMA flow diagram of the literature selection process. The search strategy identified a total of 900 publications in the initial search. After the screening of title and abstract and removal of duplicates, 47 articles were selected for further review. After exclusion of 36 articles based on aforementioned criteria, 10 studies were initially included.\textsuperscript{21–30} However, two of them evaluated the results of the same national database during the same period.\textsuperscript{22,23} Therefore, we decided to exclude the study with fewer patients\textsuperscript{23} from quantitative analysis.

Only three retrospective studies\textsuperscript{22–24} and one RCT\textsuperscript{21} compared neoadjuvant therapy and LE with radical surgery. The remaining six studies were prospective\textsuperscript{25–29} and retrospective\textsuperscript{27–30} articles which investigated oncological outcomes of LE after neoadjuvant CRT. Two of them were multicenter studies.\textsuperscript{26,29} One study\textsuperscript{25} which considered T2 and T3s tumors was also included as T3s have the same conventional treatment as T2 cancers (TME).

Six\textsuperscript{24–28,30} of 10 studies have as indication for LE of T2 rectal tumor patients who refuse major surgery, or patients with poor performance status or complete response after CRT. Details and quality assessment of the studies are showed in Table 1.

**Oncologic outcomes and pooled analysis.** All studies which compared neoadjuvant therapy followed by transanal LE with transabdominal TME showed comparable survival outcomes for patients with cT2N0 rectal cancer (Table 2). In the only RCT,\textsuperscript{21} 5-year DFS and OS was 89% and 72% for organ-preserving treatment and 94% and 80% for radical surgery respectively, with no statistically difference. Likewise, local and distant recurrence rate did not differ between two groups (8% vs 6% and 4% vs 4%, respectively).
One study retrospectively reviewed American National Cancer Database (NCDB) to determine the effect of LE with preoperative CRT and major surgery. There were no difference in OS (77.7% vs 75.1\%) with similar rate in terms of 30- and 90-day postoperative mortality.

Finally, Lynn et al.\textsuperscript{24} compared 79 patients with cT2N0 rectal cancer treated with CRT and LE in ACOSOG Z6041 trial with a similar group of patients with pT2N0 tumors who underwent upfront TME in the Dutch TME trial. Even in this comparative study, no difference regarding 5-year DFS (88.2\% vs 88.3\%), 5-year OS (90.3\% vs 88.4\%) and LR rate (4\% vs 1.3\%) emerged between groups.

Among noncomparative studies, 3-year DFS and OS rates were reported in three articles\textsuperscript{25,26,28} ranging from 86.9\% to 97.1\% and from 73\% to 100\%, respectively. In the Korean Radiation Oncology Group (KROG) 12-06 study,\textsuperscript{27} 5-year DFS was 82\% for cT2 rectal cancer patients who refused radical surgery. Similarly, Guerrieri et al.\textsuperscript{30} had 93\% 5-year DFS and 50\% 5-year OS for stage T2 patients who underwent preoperative CRT and LE. Recurrence rate was reported in all noncomparative studies up to 11.7\% of cases.

Complete pathological response (ypT0) rate after CRT and LE was reported in eight studies,\textsuperscript{21,24–30} and it ranges from 26.7\% to 59\%.

The pooled 3-year DFS, 5-year DFS, 3-year OS, 5-year OS, local and distant recurrence rates were 92.8\% (95\% CI 81.6\%–99.5\%), 91.3\% (95\% CI 88.3\%–94.3\%), 96.1\% (95\% CI 90.5\%–100\%), 72.6\% (95\% CI 57.5\%–87.7\%), 4\% (95\%CI 18\%–63\%) and 4.9\% (95\%CI 2\%–7.8\%), respectively, in subjects treated with neoadjuvant therapy followed by transanal LE (Fig. 2). No heterogeneity was found for all these analyses, except for the 5-year OS sub-analysis ($I^2$ 94\%, $P < 0.001$).

**Discussion**

The results of this review indicate that LE after CRT confers equivalent survival advantages as radical surgery in cT2 rectal cancer patients. Pooled 5-year DFS rate was 91.3\%, 5-year OS was 73.3\% and LR rate was 4\% after treatment. Therefore, organ preservation seems a feasible alternative to TME in this setting.

Although local recurrences were less after TME in Dutch trial (1.3\%), 5-year DFS rate is similar (88.3\%) while 5-year OS is 88.4\%\textsuperscript{24,31} This slight difference in OS is probably due to the inclusion of patients with more severe comorbidities (unfit for major surgery) in the conservative treatment group of the selected studies.

In last few years, attention on organ preservation strategies with multimodal approach (CRT and LE) has increased. GRECCAR 2 is a prospective randomized multicenter trial which compared TME with LE in both after NAT. A three-step strategy was adopted to identify patients who can benefit from an organ-preserving treatment: selection occurs first at the moment of the initial clinical
Table 2  Summary of oncological outcomes

| Authors                  | Surgical procedure | No. of patients | Median age | ypT0 (%) | 5y-DFS (%) | 5y-OS (%) | 3y-DFS (%) | 3y-OS (%) | LR (%) | DR (%) | 30-day mortality | 90-day mortality |
|--------------------------|--------------------|-----------------|------------|----------|------------|-----------|------------|-----------|--------|--------|------------------|------------------|
| Lezoche et al. [21]      | NAT + LE           | 50              | 66 (58–70)| 14 (28)  | 89         | 72        | 4 (8)      | 2 (4)     | 0      |       |                  |                  |
|                          | TME                | 50              | 66 (60–69)| 13 (28)  | 94         | 80        | 3 (6)      | 2 (4)     | 0      |       |                  |                  |
| Jawitz et al. [22]       | NAT + LE           | 695             | 63 (16)   | 77.7     | 50.4 Gy + 5-FU       | 12/16     | 45 Gy + B6O or Capecitabine | 10/16     |
|                          | TME                | 6629            | 66 (20)   | 75.1     | 50.4 Gy + CAPOX; Short-course radiotherapy | 12/16     | 50.4 Gy + CAPOX | 12/16     |
| Lynn et al. [24]         | NAT + LE           | 79              | 62.7 (11.24)| 38 (50)  | 88.2       | 90.3      | 3 (4)      | 1 (1.3)   | 5 (0.7) | 10 (1.4) | 116 (1.3)       | 176 (2.7)        |
|                          | TME                | 79              | 64.4 (11.25)| 88.3     | 88.4       |           |           | 1 (1.3)   | 0       | 2 (4)   |                  |                  |
| Pericay et al. [25]      | NAT + LE           | 15              | 76 (57–87)| 4 (26.7) | 91         | 73        | 0          | 1 (6.7)   | 1 (1.7) | 2 (1.7) | 1 (2.9)          |                  |
| Garcia-Aguilar et al. [26]| NAT + LE           | 79              | 62 (30–83)| 38 (49)  | 96.9       | 95.7      | 3 (4)      | 2 (1.7)   | 2 (1.7) | 1 (2.9) | 1 (2.9)          |                  |
| Noh et al. [27]          | NAT + LE           | 17              | 63 (38–79)| 10 (59)  | 97.1       | 100       | 1 (2.9)    | 1 (2.9)   | 1 (5.5) |          |                  |                  |
| Shin et al. [28]         | NAT + LE           | 34              | 63.6 (36–83)| 19 (55.9)| 97.1       | 100       | 1 (2.9)    | 1 (2.9)   | 1 (5.5) |          |                  |                  |
| Yu et al. [29]           | NAT + LE           | 18              | 9 (45)    | 50 (4)   | 97.1       | 100       | 1 (2.9)    | 1 (2.9)   | 1 (5.5) |          |                  |                  |
| Guerrieri et al. [30]    | NAT + LE           | 185             | 68 (60–74)| 63 (34.1)| 93         | 50        | 24 (1.3)   | 50 (4)    | 1 (2.9) | 1 (2.9) | 1 (2.9)          |                  |

*Local and distant recurrences.

DFS, disease-free survival; DR, distant recurrence; LR, local recurrence; OS, overall survival.
Figure 2  Pooled analysis for Local Excision. (a) 3-year DFS. (b) 5-year DFS. (c) 3-year OS. (d) 5-year OS. (e) Local Recurrence. (f) Distant Recurrence.
staging, then at the restaging 8 weeks after CRT by pelvic MRI and finally at the evaluation of the pathological response. The authors found no significant difference in terms of survival and recurrence rates between patients who had a good clinical response after chemoradiotherapy for small T2T3 low rectal cancer at 5-year follow-up. Likewise, in CARTS study, including 55 patients with cT1-3N0 tumor, organ preservation was achieved in 35 patients (64%) with acceptable long-term oncological outcomes and health-related quality of life. In TREC trial, 55 patients with cT1T2N0 rectal cancer were randomly assigned to radical surgery group or short-course radiotherapy and LE group. Although primary endpoint was the feasibility of recruiting to a RCT comparing rectal-sparing strategy with TME, no difference in DFS and OS between groups emerged assuring high level of organ preservation with relatively low morbidity.

Despite the encouraging results of the aforementioned studies, patient selection for LE after CRT is still challenging due to balance between the risk of undertreatment and surgical morbidity. Pelvic MRI is the preferred modality when a T2 or larger tumor is suspected because it has higher accuracy than endorectal ultrasound (ERUS) for detection of mesorectal infiltration. High-resolution MRI staging allows to assess tumors infiltrating the muscularis propria (T2), to measure the depth of extramural spread (T3a-d) and to evaluate lymph node involvement. Recent technological advancements, e.g. diffusion-weighted imaging (DWI) and artificial intelligence-based reconstructions, may provide additional information to MRI and improve the accuracy of pre-treatment staging. ERUS should typically be considered complementary to MRI for purposes of clinical staging, and is most useful in differentiating between early T stages (i.e., T1 versus T2 tumors) or when MRI is contraindicated.

Upfront TME is standard for rectal tumors limited to the muscularis propria (T2) with negative nodes because the risk of recurrence after LE and of harboring occult nodal disease is not negligible. Indeed, compared with LE alone, radical surgery offers a significant decreased LR rate: 7% versus 13% in T2 rectal cancer. However, 30-day mortality rate after radical rectal excision is 2%, and it increases in elderly patients up to 6% after 75 years and 12% above 85 years. Furthermore, postoperative morbidity is present in 30–50% of cases with a significant adverse impact on quality of life, bowel, urinay, and sexual dysfunction, often including the need for a definitive or permanent stoma. Thus, it is not surprising that LE may seem a desirable alternative in selected patients such as the elderly and frail patients if oncologic outcomes are not compromised. The main characteristics that patients should fulfill in order to benefit from neoadjuvant CRT and LE include tumors < 10 cm from the anal verge, ≤ 4 cm in diameter, with low risk of positive lymph nodes on MRI imaging, and with a good clinical response after nCRT.

Although both immunological and individual factors can contribute to bowel diseases, preoperative CRT has certainly demonstrated its efficacy to decrease the local recurrence rate after rectal excision for locally advanced rectal cancer. Patients who have a good response after CRT have a better prognosis than those having a bad response. As found in the present study, complete pathological response (ypT0) rate in T2N0 is high: ≥ 50% in five studies and between 27% and 34% in three studies. Thus, in addition to LE, this rectal-preserving approach may avoid morbidity of major surgery without jeopardizing survival.

Furthermore, we assume that LE after neoadjuvant CRT in T2 cancers improves local control of the disease if compared with LE and adjuvant CRT. In fact, we found a pooled LR rate of 4% that is significantly lower than 12% and 14% reported by the Cancer and Leukemia Group B studies.

There are some limitations to our study. Few studies reported long-term oncological outcomes of LE after neoadjuvant CRT for T2 rectal cancer. Three of them were comparative retrospective studies and only one RCT. Thus, it is difficult to interpret the results with accuracy. However, all studies show similar outcomes compared with conventional surgical treatment. The selection criteria for conservative strategy were not standardized among studies as well as neoadjuvant therapy regimen and technical aspect to perform LE. As the comparison of different cancer treatments should be made for similar staging, we focused on cT2N0 rectal lesions avoiding merging both early and advanced tumors results as reported elsewhere. We are aware that further studies are warranted to confirm our findings. In this setting, preliminary results of a phase III multicenter RCT (NCT01308190) demonstrated no difference in terms of LR and DR between T2-T3s cancers treated with CRT and LE compared with TME at 2-year follow up.

Conclusion

LE following neoadjuvant CRT may provide comparable survival benefit to radical surgery for patients with clinical stage T2N0. This strategy may be considered as an alternative approach for patients unfit for major abdominal resection or who refuse surgery and it may also represent a viable treatment modality for all subjects. However, further studies are needed to draw firm conclusions.

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