Phase II, single-arm trial of preoperative short-course radiotherapy followed by chemotherapy and camrelizumab in locally advanced rectal cancer

Zhenguo Hou, 6 Chii Ma, 6 Kunyu Yang, 1 Gang Wu, 1 Kaixiong Tao, 2 Tao Zhang 2

ABSTRACT

Background In locally advanced rectal cancer (LARC), preoperative short-course radiotherapy (SCRT) with delayed surgery has been shown to be as effective as long-course chemoradiotherapy, with only modest benefits. This study aimed to evaluate the efficacy and safety of preoperative SCRT combined with subsequent CAPOX (capecitabine and oxaliplatin) and the anti-PD-1 antibody camrelizumab in patients with LARC.

Methods This was a prospective, single-arm, phase II trial. Treatment-naive patients with histologically confirmed T3-4N0M0 or T1-4N+M0 rectal adenocarcinoma received 5×5 Gy SCRT with two subsequent 21-day cycles of CAPOX plus camrelizumab after 1 week, followed by radical surgery after 1 week. The primary endpoint was pathological complete response (pCR) rate. Biomarker analysis was performed to identify a potential predictor of pCR to treatment.

Results From November 7, 2019 to September 14, 2020, 30 patients were enrolled, and 27 patients received at least one dose of CAPOX plus camrelizumab. Surgery was performed in 27 (100%) patients. The pCR (ypT0N0) rate was 48.1% (13/27), including 46.2% (12/26) for proficient MMR tumors and 100% (1/1) for deficient MMR tumors. Immune-related adverse events were all grade 1–2, with the most common being reactive cutaneous capillary endothelial proliferation (81.5%). No grade 4/5 adverse events occurred. Biomarker analysis showed patients without FGFR1–3 deletions had a better tendency for pCR.

Conclusions SCRT combined with subsequent CAPOX plus camrelizumab followed by delayed surgery showed a favorable pCR rate with good tolerance in patients with LARC, especially in the proficient MMR setting. A randomized controlled trial is ongoing to confirm these results.

Trial registration number ClinicalTrials.gov identifier: NCT04231552.

INTRODUCTION

Rectal cancer, which accounts for more than one-third of colorectal cancer (CRC) cases, 1 has been challenging in terms of treatment and organ preservation, especially for locally advanced rectal cancer (LARC), because of the complex anatomical structures and high rates of postoperative complications and local recurrence. Currently, multidisciplinary therapy based on neoadjuvant long-course chemoradiotherapy (LC-CRT) with delayed surgery or short-course radiotherapy (SCRT) with immediate surgery is the predominant treatment strategy for LARC. 2 The Stockholm III trial of patients with LARC was the first to demonstrate that SCRT with delayed surgery was superior to SCRT with immediate surgery and long-course radiotherapy (LCRT) with delayed surgery in terms of pathological complete response (pCR) and tumor downstaging. 3 4 Recently, a novel treatment combination for LARC patients, namely, neoadjuvant short-course hypofractionated radiotherapy combined with subsequent chemotherapy followed by delayed surgery, led to a greater pCR and had a lower 3-year disease-related treatment failure rate than LC-CRT followed by delayed surgery. 3 6 However, unfortunately, the pCR rate is consistently below 30%, which is not good in terms of meeting treatment needs or enabling organ preservation for patients with LARC; therefore, there is a need to explore new neoadjuvant treatment options.

In 2015, immunotherapy was shown to provide significant clinical benefits to patients with metastatic CRC with a deficient mismatch repair (dMMR)/microsatellite instability-high (MSI-H) status, whereas no response was observed in patients with a proficient mismatch repair (pMMR)/ microsatellite stable (MSS) status, which accounts for approximately 96%–98% of metastatic CRCs and 85% of all CRCs. 7-9 Of
note, compared with patients with advanced-stage CRC, early-stage patients have higher CD8+ T-cell infiltration, an increase that could enhance the response to immune checkpoint inhibitors, as previously reported in melanoma. In addition, the preliminary data in the NICHE study demonstrated that 13% of patients with early-stage pMMR colon cancer could benefit from neoadjuvant dual immunotherapy, consistent with reports in other solid tumors that early-stage disease may be more responsive to immunotherapy, particularly as neoadjuvant therapy.

Numerous studies have demonstrated that the combination of checkpoint blockade immunotherapy and radiotherapy could generate synergistic antitumor effects against local and distant tumors. Recently, the clinical efficacy and safety of preoperative LC-CRT and subsequent nivolumab monotherapy in the treatment of rectal cancer were described in the VOLTAGE-A study, which enrolled and assessed 37 patients with MSS LARC. A pCR rate of 30% and a major pathologic response rate of 38% were observed, with tolerable toxicities. Given previous studies suggesting that short-course hypofractionated radiotherapy combined with subsequent chemotherapy is comparable with LC-CRT in terms of the effects of neoadjuvant therapy in rectal cancer and that the immune response was increased by hypofractionated radiotherapy plus programmed death 1 (PD-1) blockade, we hypothesized that neoadjuvant short-course hypofractionated radiotherapy combined with subsequent chemotherapy and immune checkpoint inhibitors followed by delayed surgery would afford clinical benefits for LARC.

We report the short-term results from a single-arm, single-center, phase II trial evaluating the efficacy and safety of preoperative short-course radiotherapy combined with subsequent chemotherapy (capcitabine and oxaliplatin) and the anti-PD-1 antibody camrelizumab in patients with LARC.

**METHODS**

**Study design and participants**

This was a prospective, single-arm, single-center, phase II trial performed at the Union Hospital affiliated to Tongji Medical College of Huazhong University of Science and Technology in China.

Eligible patients were aged 18 to 75 years, with histologically confirmed T3-4N0M0 or T1-4N+M0 rectal adenocarcinoma, inferior margin of ≤10 cm from the anal verge, treatment naïve, an Eastern Cooperative Oncology Group performance status of 0–1, and no severe hematologic, cardiac, pulmonary, hepatic, or renal functional abnormalities or immunodeficiency diseases. Laboratory tests were required to meet show the following: hemoglobin level ≥9 g/dL; white blood cell count ≥3×10^9/L; absolute neutrophil count ≥1.5×10^9/L; platelet count ≥100×10^9/L; bilirubin level <1.5× the upper limit of normal (ULN); aspartate aminotransferase and alanine aminotransferase (AST) levels ≤2.5× ULN; serum creatinine level ≤1.5× ULN or creatinine clearance ≥50 mL/min; thyroid-stimulating hormone level ≤1× ULN or T3 and T4 levels within normal limits. Patients were recruited regardless of their programmed death ligand 1 (PD-L1) expression level. Key exclusion criteria included previous exposure to any anti-PD-1 or anti-PD-L1 antibody, a history of pelvic radiation, treatment with corticosteroids or other immunosuppressive agents within 14 days prior to study drug administration, presence of autoimmune disease, and known interstitial lung disease.

**Procedures**

Eligible patients received SCRT (5×5 Gy over 5 days), followed 1 week later by two subsequent 21-day cycles of CAPOX (oxaliplatin 130 mg/m^2 intravenously, day 1; capcitabine 1000 mg/m^2 oral twice daily, days 1–14) plus camrelizumab (200 mg intravenous drip, day 1), followed by radical surgery after 1 week. Surgery was done according to total mesorectal excision principles. Low anterior resection was performed for middle and low rectal cancers with distal margins of more than 1 cm, and abdominoperineal resection was conducted by the surgeon for extremely low tumors.

Patients received a baseline assessment including collection of information on demographics, medical history, and disease characteristics before enrollment, and they underwent systematic physical examination and relevant laboratory and imaging (chest CT, liver MRI, abdominal and pelvic CT or MRI) tests before and after treatment. Patients discontinuing treatment for reasons other than progressive disease were followed up every 3 months until disease relapse or death.

Surgically resected specimens were processed and examined as previously reported. The extent of the residual tumor in the resected specimens was classified as per the 8th edition of the International Union Against Cancer TNM staging system. All resected specimens (including the whole tumor bed and surrounding tissues) were sampled and cross-sectioned consecutively. If no residual viable tumor was identified after examination, additional three-level sections were carried out. All sampled lymph nodes were examined based on standard procedures. All sections were evaluated independently by two senior pathologists. A pCR was defined as the absence of any remaining viable cancer cells in the resected primary tumor specimen and all sampled regional lymph nodes ( ypT0N0). The tumor regression grade (TRG) was assessed according to the criteria proposed by Ryan et al.

**Outcomes**

The primary endpoint was pCR rate, defined as the proportion of patients who had a pCR. Secondary endpoints included 3-year event-free survival rate (defined as the percentage of patients without disease recurrence or progression or death due to any cause at the 3-year follow-up), R0 resection rate (defined as the rate of negative margins microscopically), 3-year overall survival rate (defined as the percentage of patients alive at the 3-year follow-up), complication rate, safety, and
quality of life. Adverse events (AEs), recorded during the period when patients signed their informed consent forms to 90 days after surgery, were graded according to the National Cancer Institute Common Terminology Criteria for Adverse Events, version 5.0.

**Biomarker analysis**

Tissue samples or paraffin sections were required to be provided when patients were enrolled. Immunohistochemistry was performed to detect the expression of PD-L1, DNA mismatch repair (MMR) proteins (MSH6, MSH2, MLH1, and PMS2), CD4, and CD8. The PD-L1 combined positive score (CPS) was evaluated using the PD-L1 IHC 22C3 pharmDx assay (Agilent Technologies, Santa Clara, CA, USA), defined as the number of PD-L1 positive cells (tumor cells, lymphocytes, macrophages) as a proportion of the total number of tumor cells multiplied by 100. Positive PD-L1 expression was considered when the CPS was 1 or more. Comprehensive genomic profiling was conducted using targeted gene capture–based next-generation sequencing technology. Briefly, for formalin-fixed paraffin-embedded tissues, H&E staining was performed, and the stained sections were evaluated by a pathologist to ensure tumor cells ≥20%. DNA was extracted from the tumor tissues of patients using standard methods. A panel of 418 genes was captured and then sequenced through the Illumina NovaSeq 6000 platform (Illumina, San Diego, CA, USA) with 2×150 bp paired-end reads. The average sequencing depth of tumor tissues was ≥1000×. Genomic alterations, including tumor mutation burden (TMB), single nucleotide variants, short and long insertions and deletions, copy number variants, and gene fusions, were assessed.

**Table 1** Patient characteristics

| Characteristics                  | Patients (n=30) |
|----------------------------------|----------------|
| Age, years, median (range)       | 57 (31–73)     |
| Sex, n (%)                       |                |
| Male                             | 17 (56.7)      |
| Female                           | 13 (43.3)      |
| ECOG performance status, n (%)   |                |
| 0                                | 13 (43.3)      |
| 1                                | 17 (56.7)      |
| Clinical T category, n (%)       |                |
| cT3                              | 22 (73.3)      |
| cT4                              | 8 (26.7)       |
| Clinical N category, n (%)       |                |
| cN0                              | 4 (13.3)       |
| cN1                              | 16 (53.3)      |
| cN2                              | 10 (33.3)      |
| Clinical disease stage, n (%)    |                |
| Stage II (cT3-4N0)               | 4 (13.3)       |
| Stage III (cT1-4N1-2)            | 26 (86.7)      |
| CRM, n (%)                       |                |
| Positive                         | 21 (70.0)      |
| Negative                         | 9 (30.0)       |
| EMVI, n (%)                      |                |
| Positive                         | 12 (40.0)      |
| Negative                         | 18 (60.0)      |
| Distance from primary tumor to anal verge, n (%) | |
| Median (range), cm               | 4.7 (1.9–9.0)  |
| <5                               | 15 (50.0)      |
| 5–10                             | 15 (50.0)      |
| Length of tumor lesion, cm       |                |
| Mean (SD)                        | 5.5 (1.7)      |
| Median (range)                   | 5.4 (2.1–10.0) |
|Mismatch repair status, n (%)     |                |
| dMMR                             | 1 (3.3)        |
| pMMR                             | 28 (93.3)      |
| Unknown                          | 1 (3.3)        |
|PD-L1 expression, CPS, n (%)      |                |
| Negative                         | 20 (66.7)      |
| Positive (≥1)                    | 6 (20.0)       |
| Unknown                          | 4 (13.3)       |
|Tissue-based TMB (mut/Mb), n (%)  |                |
| <10                              | 7 (23.3)       |
| ≥10                              | 7 (23.3)       |
| Unknown                          | 16 (53.3)      |
|Baseline CEA level, n (%)         |                |
| Normal (≤5 ng/mL)                | 17 (56.7)      |
| Abnormal (≥5 ng/mL)              | 13 (43.3)      |

CEA, carcinoembryonic antigen; CPS, combined positive score; CRM, circumferential resection margin; dMMR, deficient mismatch repair; ECOG, Eastern Cooperative Oncology Group; EMVI, extramural vascular invasion; PD-L1, programmed death ligand 1; pMMR, proficient mismatch repair; TMB, tumor mutation burden.
Statistical analysis
According to previous studies, the pCR rate after preoperative CRT in patients with LARC is approximately 15%.\textsuperscript{22,23} We expected that the regimen of SCRT combined with subsequent chemotherapy and camrelizumab could increase the pCR rate from 15% to 40%. A sample size of 24 patients was required to provide at least 80% power to detect this estimated improvement in a one-sided $\chi^2$ test with a significance level of 2.5%, and a 20% dropout was estimated, resulting in a total sample size of 30 patients planned for this study.

Statistical analyses were conducted using SAS software (V.9.2; SAS Institute, Cary, NC, USA). Continuous variables were summarized using medians and ranges, and categorical variables were described using frequencies and percentages. Baseline and safety analyses were performed for all enrolled patients (intention-to-treat (ITT) population), and efficacy analyses were conducted for those who were administered at least one dose of camrelizumab (full analysis set). Comparisons between the groups were performed using Fisher’s exact test or the $\chi^2$ test. P values <0.05 were considered statistically significant.

RESULTS
Patient characteristics and compliance
From November 7, 2019 to September 14, 2020, the target number of eligible patients was enrolled (n=30). Three patients were not treated with CAPOX plus camrelizumab for the reasons given in figure 1 and were excluded from the efficacy analysis. Patient baseline and disease characteristics are summarized in table 1. Approximately half of the patients had at least one high-risk factor, including cT4 disease in 26.7% (8/30), cN2 disease in 33.3% (10/30), extramural vascular invasion in 40.0% (12/30), and tumors located within <5 cm from the anal verge in 50.0% (15/30).

During the treatment period, all patients received the full irradiation dose of 25 Gy (100%, 30/30), and 27 patients (90.0%, 27/30) received at least one cycle of CAPOX plus camrelizumab, of whom three patients did not complete the second cycle of CAPOX plus camrelizumab due to the impact of the COVID-19 pandemic. The median intervals from the completion of preoperative SCRT to receiving subsequent CAPOX plus camrelizumab and from the last dose of CAPOX plus camrelizumab to surgery were 12 (range, 4–18) days and 14 (range, 5–141) days, respectively.

Surgery and pathology
A total of 27 patients (100%, 27/27) underwent surgery in the full analysis set, with an R0 resection rate of 100% (27/27) and anal preservation rate of 88.9% (24/27). A total of four patients developed postoperative complications, including infection in three patients (11.1%, 3/27) and bleeding in one patient (3.7%, 1/27). No other complications were reported.
postoperative complications or treatment-related deaths occurred. The median interval from surgery to hospital discharge was 8 (range, 7–16) days.

The pCR (ypT0N0) rate was 48.1% (13/27), including 46.2% (12/26) for patients with pMMR and 100% (1/1) for those with dMMR. Negative nodes (ypN0) were reported in 19 (70.4%) of the 27 patients, 18 of whom were in the pMMR patient subset (69.2%, 18/26; table 2). As an example, imaging of patient 7 (a 63-year-old woman) with cT4N2bMx pMMR rectal cancer showed a visible tumor at baseline, the volume of which was significantly reduced after two cycles of treatment, and the histopathologic specimen revealed complete regression of the primary tumor, that is, pCR (figure 2A). Moreover, an obvious increase in CD4+ and CD8+ T-cell infiltration was observed in posttreatment tumor samples compared with pretreatment samples in patient 7 but not in patient 12 who did not achieve pCR (figure 2B). Notably, all

**Figure 2** Pathological response to preoperative short-course radiotherapy followed by CAPOX and camrelizumab. (A) Radiological and pathological response from patient 7. Left: MRI of the abdomen (upper row) and endoscopic imaging of the primary tumor (lower row) before and after treatment. Right: Postoperative specimen with mucosal retraction at the tumor site with a polypoid structure (upper row) and H&E staining showing CR of the primary tumor after treatment (lower row). (B) Representative CD8 and CD4 staining of pretreatment and posttreatment specimens of patient 7 and patient 12. Scale bars, 250 µm.
patients with PD-L1 CPS ≥1 were reported to be ypN0. The pCR of patients with PD-L1 CPS ≥1 was numerically higher than that of patients with CPS <1 (66.7%, 4/6 vs 45.0%, 9/20, p=0.2422).

**Safety**

Treatment-emergent AEs and immune-related AEs are summarized in table 3. At the data cut-off (January 15, 2021), the most common treatment-emergent AEs of any grade were leukopenia (80.0%, 24/30), reactive cutaneous capillary endothelial proliferation (73.3%, 22/30), and radiation proctitis (70.0%, 21/30). The most common grade 3 treatment-emergent AEs were neutropenia (13.3%, 4/30) and anemia (13.3%, 4/30). Immune-related AEs were all grade 1–2, and the most common was reactive cutaneous capillary endothelial proliferation, which occurred in 22 (81.5%) of 27 patients. Five patients experienced serious treatment-emergent AEs, among whom four were hospitalized or had prolonged hospitalization due to grade 2 radiation enteritis and one was hospitalized due to abdominal pain. No grade 4 or 5 AE occurred during the study.

**Results of biomarker analysis**

Next-generation sequencing was carried out in 53.8% (14/26) of patients with pMMR in this study; the remainder were not performed due to delivery of unqualified samples or patient refusal. As shown in figure 3A, the most frequently altered genes at baseline were TP53, APC, and KRAS. Our small-sample analysis did not identify any genes with significant differences in the frequency of genomic alterations between the pCR and non-pCR groups (figure 3B). Compared with patients with TMB <10, those with TMB ≥10 had a better tendency for pCR (42.9%, 3/7 vs 28.6%, 2/7, p=0.3671). Interestingly, according to baseline copy number variant, none of the five patients with FGFR1-3 deletions achieved pCR, while more than half of the nine patients without FGFR1-3 deletions achieved, although no statistically significant difference was observed (55.6%, 5/9 vs 0%, 0/5, p=0.086).

**DISCUSSION**

To the best of our knowledge, our study is the first to propose a new neoadjuvant therapy regimen of short-course hypofractionated radiotherapy combined with subsequent chemotherapy and anti-PD-1 antibody. In addition, this study provides preliminary evidence that the addition of camrelizumab to neoadjuvant SCRT followed by the CAPOX chemotherapy regimen results in a remarkable pCR for patients with LARC and is well tolerated, without new or unexpected safety issues.

As shown in previous studies, preoperative radiotherapy combined with chemotherapy resulted in tumor downstaging and reduced local recurrence, whereas pCR was observed only in 15%–30% of patients with rectal cancer. In this study, our pCR rate of 48.1% is encouraging, meaning that our innovative preoperative combination therapy strategy provides more opportunities for sphincter-preserving surgery and also raises the prospect that more patients with LARC, especially those with low rectal cancer, may achieve a clinical complete response and have a watch-and-wait strategy of nonsurgical treatment implemented to improve their quality of life.

Immunotherapy is generally ineffective in the pMMR/MSS tumors that constitute the majority of CRCs, which could be attributed to insufficient lymphocytic infiltration. Preclinical data have shown that radiotherapy can sensitize refractory tumors to PD-1/PD-L1 blockade by modulating the immunogenicity of tumor cells, enhancing antigen-specific CD8+ T-cell responses, and increasing PD-L1 expression on tumor cells and immune cells in the tumor microenvironment; in addition, chemotherapy can also upregulate PD-L1 on dendritic cells and increase immune-cell infiltration. Based on these rationales, immunotherapy strategies combined with chemoradiotherapy are being explored in patients

| Table 3 | Adverse events since the initiation of radiotherapy |
|---------|--------------------------------------------------|
| Treatment-emergent AEs | Any grade  | Grade 3 |
| Any AE, n (%) | 29 (96.7) | 8 (26.7) |
| Leukopenia  | 24 (80.0) | 1 (3.3) |
| Reactive cutaneous capillary endothelial proliferation | 22 (73.3) | 0 |
| Radiation proctitis | 21 (70.0) | 0 |
| Anemia | 20 (66.7) | 4 (13.3) |
| Peripheral neurotoxicity | 18 (60.0) | 0 |
| Neutropenia | 16 (53.3) | 4 (13.3) |
| Thrombocytopenia | 16 (53.3) | 0 |
| Fatigue | 13 (43.3) | 0 |
| Anorexia | 10 (33.3) | 0 |
| Alanine transaminase increased | 9 (30.0) | 0 |
| Aspartate transaminase increased | 9 (30.0) | 0 |
| Vomiting | 8 (26.7) | 0 |
| Nausea | 6 (20.0) | 0 |
| Hypothyroidism | 2 (6.7) | 0 |
| Abdominal pain | 2 (6.7) | 0 |
| Hand-foot syndrome | 1 (3.3) | 0 |
| Constipation | 1 (3.3) | 0 |
| Diarrhea | 1 (3.3) | 0 |
| Immune-related AEs, n (%) | 22 (81.5) | 0 |
| Reactive cutaneous capillary endothelial proliferation | 22 (81.5) | 0 |
| Hypothyroidism | 2 (7.4) | 0 |

*The frequency of immune-related AEs among 27 patients who received at least one dose of camrelizumab was calculated. AE, adverse event.*
Figure 3  Genetic analysis. (A) Overall frequency of gene alterations at baseline. (B) Frequency of genomic alterations between the pCR and non-pCR groups. pCR, pathological complete response.
with pMMR/MSS rectal cancer, especially in LARC setting. In the VOLTAGE study, the pCR rate was 30% in patients with MSS LARC receiving preoperative LC-CRT and sequential nivolumab. In our study, the pCR rate was 46.2% for patients with pMMR disease. In addition, the recently reported pCR rate was 37.5% among patients with locally advanced rectal adenocarcinoma receiving SCRT followed by mFOLFOX-6 plus avelumab (an anti-PD-L1 antibody) as neoadjuvant therapy in the Averectal study. When comparing the results of our study and the Averectal study, differential N staging was noted between the enrolled patients, with stage N1 patients being predominant in our study (53.3%) but stage N2 in the Averectal study (75.0%). In addition, a meta-analysis of randomized trials has indicated the superior efficacy of anti-PD-1 antibody over anti-PD-L1 antibody in solid tumors, regardless of monotherapy or combination strategies. Although the course of neoadjuvant immunotherapy plus chemotherapy is inconclusive, a short course of chemotherapy has been shown to improve the early efficacy of immunotherapy, as reported by Checkmate 9LA. Results from the NRG-G1002 study demonstrated neoadjuvant concurrent pembrolizumab and LC-CRT after FOLFOX induction did not significantly improve the short-term clinical outcomes of patients with LARC compared with FOLFOX and LC-CRT alone, with pCR rate of 31.9% in pembrolizumab group versus 29.4% in control group (p=0.75). Although cross-study comparisons should be made with caution, it is interesting that the pCR rate of 48.1% observed in our study compared favorably with the NRG-G1002 study. The differences may be explained by different neoadjuvant therapeutic strategies with the differential timing of anti-PD-1 antibody administration.

A time interval of at least 7 days after SCRT has been reported to be probably required to achieve a favorable immune response, suggesting that the choice of the timing of immunotherapy after SCRT is important. The PACIFIC trial reported that the initiation of durvalumab within 2 weeks after chemoradiotherapy (rather than ≥2 weeks) was linked to a higher clinical benefit. Therefore, we have reasons to believe that our choice of timing for the initiation of immune checkpoint inhibitor camrelizumab in this study (median 12 days) is appropriate.

The previous VOLTAGE-A study showed that positive PD-L1 expression and high TMB are good predictors of neoadjuvant efficacy from LC-CRT followed by nivolumab. Similarly, a trend toward a better pCR rate was observed in patients with PD-L1 CPS ≥1 or with TMB ≥10 in our phase II trial, although the differences were not statistically significant. Given that ours was an exploratory analysis with a small sample, we recommend that this finding be interpreted with caution. In addition, patients without FGFR1-3 deletions appear to be more likely to achieve clinical benefit from our neoadjuvant strategy than those with FGFR1-3 deletions (55.6% vs 0%). In urothelial carcinoma, patients with FGFR mutations have been reported to show a worse response to immunotherapy than those without FGFR mutations. A similar result was demonstrated in metastatic gastric adenocarcinoma, showing that the patient harboring MSI-H and FGFR2 alterations was resistant to anti-PD-1 monotherapy. If positively validated in the subsequent larger study that we plan to conduct, the presence of FGFR1-3 deletions may be a potent predictor of response to neoadjuvant immunotherapy in rectal cancer.

The limitations of this study consist of the small sample size of patients, lack of a control group, and insufficient postoperative follow-up time. Furthermore, whether improvement in pCR in our study can contribute to better survival needs further exploration. A large multicenter randomized phase III study is going to confirm the value of preoperative short-course hypofractionated radiotherapy combined with subsequent chemotherapy and camrelizumab regimen (NCT04928807).

In conclusion, SCRT combined with subsequent CAPOX plus camrelizumab followed by delayed surgery showed a favorable pCR rate with manageable toxicities, especially in the pMMR setting. This suggests that neoadjuvant SCRT combined with subsequent CAPOX plus camrelizumab is a promising feasible strategy and is expected to provide more opportunities for surgery to be delayed or avoided among patients with middle-low LARC and for organ function to be preserved in the future.

Author affiliations
1Cancer Center, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, China
2Department of Gastrointestinal Surgery, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, China
3Department of Digestive Surgical Oncology, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, China
4Department of Radiology, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, China
5Department of Pathology, Union Hospital, Tongji Medical College, Huazhong University of Science and Technology, Wuhan, China
6Department of Medical Affairs, Jiangsu Hengrui Pharmaceuticals Co., Ltd, Shanghai, China

Correction notice An error in Figure 2A has been corrected since this article was first published. The left image of H&E staining should have been labelled as post-treatment.

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Correction: Phase II, single-arm trial of preoperative short-course radiotherapy followed by chemotherapy and camrelizumab in locally advanced rectal cancer

Lin Z, Cai M, Zhang P, et al. Phase II, single-arm trial of preoperative short-course radiotherapy followed by chemotherapy and camrelizumab in locally advanced rectal cancer. *J Immunother Cancer* 2021;9:e003554. doi:10.1136/jitc-2021-003554

This article has been corrected since it was first published. There was an error in panel A of Figure 2; the left image of H&E staining is mistakenly described as pre-treatment, while in fact it is a post-treatment image. The figure has now been updated.

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