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REVIEW

Radiotherapy for esophageal carcinoma: dose, response and survival

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Abstract: Esophageal cancer (EC) is an extremely aggressive, lethal malignancy that is increasing in incidence worldwide. At present, definitive chemoradiotherapy is accepted as the standard treatment for locally advanced EC. The EC guidelines recommend a radiation dose of 50.4 Gy for definitive treatment, yet the outcomes for patients who have received standard-dose radiotherapy remain unsatisfactory. However, some studies indicate that a higher radiation dose could improve local tumor control, and may also confer survival benefits. Some studies, however, suggest that high-dose radiotherapy does not bring survival benefit. The available data show that most failures occurred in the gross target volume (especially in the primary tumor) after definitive chemoradiation. Based on those studies, we hypothesize that at least for some patients, more intense local therapy may lead to better local control and survival. The aim of this review is to evaluate the radiation dose, fractionation strategies, and predictive factors of response to therapy in functional imaging for definitive chemoradiotherapy in esophageal carcinoma, with an emphasis on seeking the predictive model of response to CRT and trying to individualize the radiation dose for EC patients.

Keywords: esophageal cancer, radiation dose, altered fractionation, predictive factors, individualization

Introduction

Esophageal cancer (EC) is a common cause of cancer death around the world.^{1,2} In East Asia, squamous cell carcinoma (SCC) is the most common type of EC, whereas adenocarcinoma is predominant in Western countries.^{3,4} These two types may represent different diseases, each with a distinct pathogenesis, epidemiology, prognosis, and tumor biology, including the pattern of lymph node metastasis.⁵

At present, definitive concurrent chemoradiotherapy (CCRT) is the basic strategy for locally advanced inoperable EC patients based on the results of the pivotal study of Intergroup Radiation Therapy Oncology Group (RTOG)-8501, which significantly improved the local control (LC) and overall survival (OS) with CCRT compared with radiotherapy (RT) alone.⁶ Furthermore, another RTOG study (#90-12) found that escalating the dose to 64.8 Gy did not confer a benefit compared with standard doses and may have contributed to a higher incidence of treatment-related death.⁷ In the landmark INT0123 trial, dose escalation from 50.4 to 64.8 Gy also did not increase OS; this may have been due to higher toxicities and no therapeutic gain in dose escalation for pathological complete responses (pCRs) after planned radiation.⁸ On the basis of these results, 50.4 Gy has become the accepted standard dose in Europe and North America

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for patients undergoing definitive CRT (DCRT). Although radiation dose escalation has failed to improve LC or OS, a dose of 60.0 Gy or more is more popular in Asian countries, where SCC is the predominant histological type.9,10 However, the prognosis for EC patients treated by standard-dose CRT was still disappointing, and >50% of such patients eventually developed recurrence or distant metastases and died as a result.¹¹ Many studies have been performed to describe patterns of recurrence after DCRT for EC, revealing that most failures in both early and advanced EC occurred in the gross target volume (GTV) (especially in the primary tumor) after CRT.^{12–18} Significantly, the recurrent site in the patient undergoing surgery occurred mainly in the locoregional node as well as distant metastasis, while in patients treated with DCRT it occurred mainly in the primary lesion.¹⁹⁻²¹ This indicates that the standard dose (50.4 Gy in 28 fractions), at least for some patients, may be inadequate to achieve a high probability of LC. Hence, for EC patients treated with DCRT, efforts should be made to intensify the treatment to improve the LC rate, which may lead to survival benefits.

In addition, on the basis of the theory of radiation biology, a radiation dose of 45-50 Gy is adequate to control microscopic tumors, 60 Gy or more is required to control gross tumors, and nearly 100 Gy is needed to cure solid tumors at conventional fractionation.²² In recent years, continuous advancements in RT technology over the past decades have allowed for EC now being able to be treated with 3-dimensional (3D) treatment planning, including intensity-modulated RT (IMRT), volumetric-modulated arc therapy (VMAT) and helical tomotherapy. These modern techniques allow the oncologist to deliver higher doses of radiation with more precision to the tumor and with less toxicity to the surrounding normal tissue, which has dramatically reduced morbidity.^{23,24} Thus, with the improvements in RT techniques and chemotherapy regimens, the question has been raised as to whether high-dose RT given concurrently with effective chemotherapy could achieve similar or better survival rates compared with the standard treatment, especially for esophageal SCC (ESCC).

The aim of this review is to evaluate the radiation dose, fractionation strategies, and predictive factors of response to therapy in functional imaging for definitive CRT in EC, with an emphasis on seeking the predictive model of response to CRT and trying to individualize the radiation dose for EC patients.

Higher dose RT for DCRT

Continuous advancements in RT technology allow the oncologist to deliver higher radiation doses to the tumor with less toxicity to the surrounding normal tissue. This has led to a dramatic decline in morbidity.^{23,24} Furthermore, the simultaneous integrated boost (SIB) technique has been applied to many tumors, including in EC. Use of an SIB technique enables delivery of a higher dose to the primary tumor (2.2 Gy/f), whereas lower doses to the subclinical disease (1.8 Gy/f).^{25,26}

With the improvements in treatment techniques, some researchers have tried to address the potential benefit of a higher radiation dose for EC. In the retrospective analysis by Suh et al the results also showed that patients who received a total dose \geq 50.4 Gy of RT had significantly better loco-regional control (LRC, 69% vs 32%, p<0.01) and progression-free survival (PFS, 47% vs 20%, p=0.01), than patients receiving <50 Gy when treated with concurrent chemotherapy. High-dose radiation ≥50.4 Gy showed no significant OS benefit for patients with EC (28 vs 18 months, p=0.26).²⁷ He et al used modern radiation delivery techniques to determine whether high radiation dose could confer benefits in terms of LC or OS.28 The results showed that high radiation dose provided a significant lower rate of local recurrence (17.9% vs 34.3%, p=0.024) compared with patients receiving low radiation dose. Furthermore, patients receiving high radiation dose did have a marginally better 5-year local-regional recurrence-free survival (68.7% vs 55.9%, p=0.052) than in the low-dose group. The 5-year OS rate showed no significant difference between the two groups (p=0.617). Although IMRT reduced the overall incidence of treatment-related toxicity, the rates of grade 3 or greater skin reaction and esophageal strictures were higher in the higher dose group. Considering the potential for better tumor LC but more severe toxicity with the higher radiation dose, He et al recommends that individualized treatment strategies be designed for patients treated with DCRT. Indeed, these studies found that radiation doses over 50.4 Gy could improve LC but not OS for EC patients.

However, some studies have shown that high-dose RT may provide a survival benefit. The study of Zhang et al found that the median survival time was 9 months for the lower dose group (\leq 51 Gy) and 14.5 months for the higher dose group (\geq 51 Gy) among 69 patients treated with CCRT (p=0.054).²⁹ In line with these findings, Kim et al evaluated the correlation between radiation dose and survival for EC patients treated with DCRT. The results showed that patients in the high-dose group (\geq 60 Gy) had significantly better 2-year LRC (69.1% vs 50.3%, p=0.002), median PFS (16.7 vs 11.7 months, p=0.029), and median OS (35.1 vs 22.3 months, p=0.043) than in the low-dose group (<60 Gy).³⁰ Similar to

these results, a retrospective study found that higher radiation dose may bring a survival benefit for EC patients.³¹ Wolf et al also concluded that the use of radiation doses exceeding 54 Gy was associated with better OS (p=0.002).³² Semrau et al in a retrospective analysis, showed that patients receiving high radiation dose had a greater 2-year OS (26.8%, vs 7.5%; p=0.0001), and PFS (17.4% vs 5.0%; p=0.0001) than the low-dose group.³³ A pooled analysis by Song et al investigated whether high-dose (>60 Gy) RT in DCRT could confer survival benefits compared with the low-dose RT for patients with EC.³⁴ Those results showed advantages in response rate, 5-year OS, locoregional recurrence, and distant failure compared with the standard RT arm. However, there is no evidence from prospective randomized trials to support that a higher radiation dose could bring survival benefit for EC patients.

In a recently published Phase I dose-escalation study, Yu et al suggest that it is feasible to deliver up to 70 Gy (2.8 Gy/F) to the GTV based on positron emission tomography/ computed tomography (PET/CT) in EC patients.³⁵ Thus, the use of a novel dose-escalation technique may be more likely than high-dose (>60 Gy) RT to lead to better LC and OS. A Phase II study by Chen et al evaluated the efficacy of radiation dose escalation using SIB in 60 EC patients.³⁶ RT consisted of 66 Gy at 2.2 Gy per fraction to the gross tumor and 54 Gy at 1.8 Gy per fraction to subclinical diseases simultaneously. The result showed that the 1-year LRC, distant metastasisfree survival (DMFS), disease-free survival (DFS), and OS rates were 87.6%, 78.6%, 86.0%, 80.5%, respectively. And the 2-year LRC, DMFS, DFS, and OS rates were 75.6%, 64.4%, 86.7%, 72.7%, respectively. This indicates that SIB combined with concurrent chemotherapy is feasible, with tolerable acute toxicities in EC patients, and a trend of significant improvements in LRC and OS was shown.

Radiation dose escalation for DCRT studies are summarized in Table 1. In summary, a few studies have demonstrated that higher doses than standard RT can improve local tumor control and may lead to better survival for EC patients. Although no evidence from Phase III prospective randomized trials support the additional survival benefit of dose-escalated RT in the whole population, these results may suggest that a subgroup of patients may benefit from limited dose escalation. Considering the potential survival benefits but more severe toxicity in the higher radiation dose group, the optimal radiation dose should be managed on an individual basis. Nevertheless, Phase III trials comprising a standard-dose arm using modern RT technique are warranted in the right subgroup population.

Dose fractionation strategies for DCRT

The prognosis for EC patients treated with conventional RT alone remains discouraging despite the advances in radiotherapeutic techniques. The low survival rate and high incidence of locoregional treatment failure in EC have initiated a modification of conventionally fractionated RT. Recent studies are consistent in revealing that the accelerated reproliferation of carcinoma stem cells after RT is an important reason for the failure of RT.^{37,38} Therefore, it is speculated that the rates of LC and survival can be improved by controlling the accelerated reproliferation of carcinoma stem cells after RT. In this respect, it has been revealed that 3–4 weeks after RT, carcinoma stem cells started to have accelerated reproliferation, which provides the theoretical principle for accelerated hyperfraction RT later in the treatment process.

Table I Studies regarding high-dose RT and/or conventional-dose RT in esophageal cancer

Authors	Radiation dosage	Radiation technique	No. of patients	LCR, %	p-value	OS	p-value
Minsky et al ⁸	50.4 Gy	2D-RT	109	52 (LFR)	>0.05	40% (2 year)	>0.05
	64.8 Gy		109	56		31%	
Zhang et al ²⁹	<51 Gy	2D-RT	43	19 (3 year)	0.011	3% (3 year)	0.054
	≥51 Gy		26	36		13%	
Suh et al ²⁷	<60 Gy	2D-RT	49	32 (2 year)	<0.01	18 months (MST)	0.26
	≥60 Gy		77	69		28 months	
He et al ²⁸	≤50.4 Gy	3D-RT	137	34.3 (LFR)	0.024	33.0% (5 year)	0.617
	>50.4 Gy		56	17.9		41.7%	
Kim et al ³⁰	<60 Gy	3D-RT or	120	37.3 (5 year)	0.02	22.3 months (MST)	0.043
	≥60 Gy	IMRT	116	59.7		35.1 months	
Chen et al ³¹	50–50.5 Gy	3D-RT	324	NR	NR	14% (5 year)	<0.05
	≥60 Gy		324	NR		22%	
Chen et al ³⁶	GTV 66 Gy /30f	SIB-IMRT	60	78.6% (2 year)	-	72.7% (2 year)	-
	CTV 54 Gy/30f						

Abbreviations: CTV, clinical target volume; GTV, gross target volume; IMRT, intensity modulated radiotherapy; LCR, local control rate; LFR, local failure rate; MST, median survival time; NR, not reported; OS, overall survival; RT, radiotherapy; SIB, simultaneous integrated boost.

Several studies have investigated whether it would be possible to achieve a better curative effect from RT by adopting a late-course accelerated hyperfraction (LCAF). A Japanese Phase II study of accelerated hyperfraction plus 5-fluorouracil/cisplatin chemotherapy showed a promising result.³⁹ Similarly, Shi et al⁴⁰ initiated a study of late-course accelerated hyperfractionated RT (LCAFRT) for ESCC treatment and it yielded encouraging results. They found that patients with the LCAFRT regime could achieve a better 5-year survival rate (34% vs 15% for patients with conventional fractionation) and LC rate (55% vs 21% for patients with conventional fractionation). A prospective study by Zhao et al also revealed that the LCAFRT regimen offers better LC and survival compared to standard chemotherapy plus RT, as in the RTOG 85-01 and 94-05 trials.⁴¹ A meta-analysis from China strengthened the evidence supporting the therapeutic benefits of LCAFRT compared with conventional fractionation for EC.42 In addition, a randomized controlled trial was recently undertaken to investigate whether LCAF 3D conformal RT could achieve better results than conventional fraction (CF).43 The resulting data showed that the 1-, 2- and 3-year survival rates were 79.2%, 56.3%, and 43.8%, respectively, in the LCAF group; and, in the CF group, the 1-, 2- and 3-year survival rates were 74%, 54%, and 36%, respectively (p=0.476). The 1-, 2- and 3-year LC rates were 81.3%, 62.5%, and 50%, respectively, in the LCAF group; in the CF group, the 1-, 2- and 3-year LC rates were 78%, 58%, and 42%, respectively (p=0.454). In the CF group, the incidence of radiation-induced esophagitis was lower than that in the LCAF group (72% vs 93.8%; p=0.008). It was concluded that EC patients in the LCAF group did have a slightly improved survival compared to those who received RT using conventional fractionation; the radiation toxicities, however, were greater in the LCAF group than those in the CF group. In a Phase III randomized study of LCAFRT plus concurrent chemotherapy for patients with ESCC, Zhao reported patients who received LCAFRT with concurrent chemotherapy had a tendency to better survival. But the incidence rates of grade 3 and 4 toxicities seemed higher in the LCAHRT+ CT arm (46%) than those in the LCAHRT arm (25%), and the grade 5 toxicities for the two group were 6% and 0%, respectively.44

These studies suggest that the accelerated hyperfractionated schedules were effective but with an increased incidence of acute III–IV grade toxicity, which limited the combination of concurrent chemotherapy with RT. The advent of modern RT techniques and low-toxicity chemotherapy drugs may improve the clinical efficacy. The medical effects and safety observations of this kind of combination should be verified in prospective trials.

Future thinking

Much evidence has shown that there is a positive correlation between OS rate and the scale of chemoradiation in the histopathological response of patients with EC.^{45–47} Patients with CRs had a 5-year OS rate of 61.6%, which is higher than patients with an incomplete response or no response.⁴⁵ We speculated that if more patients with incomplete or no responses after planned radiation could achieve CR through limited radiation dose escalation, their prognosis would be better. Besides, this scheme would avoid an increase in toxicity and a decrease in survival caused by dose escalation in patients with CRs.

But how to predict the response to therapy? A great number of studies have reported the response rates after CCRT in EC.^{6,8,16–18,48} The CR rate by stage was 89.7%–97% for T1, 50%-60% for T2 to T3, and 17%-39% for T4. For early-stage EC, most patients achieved a CR after CRT, while for locally advanced patients, CRT generally resulted in CR rates of 20%-50%. Apart from the tumor stage, patients with EC receiving RT also showed disparate treatment responses. Unfortunately, a large number of patients were resistant to CRT, which resulted in persistent disease or immediate local failure. For radiosensitive patients, the standard radiation dose of 50.4 Gy may be sufficient to obtain a pCR. But for the resistant lesions or advanced-stage EC, it is difficult to achieve a pCR at this dose. Even a dose escalation could not produce a better response; it may merely increase treatmentrelated toxicities in such refractory cases. We speculated that a subgroup of patients with partial remission after the standard radiation of 50.4 Gy could achieve better responses through limited radiation dose escalations without increased treatment toxicities. Hence, the identification of the predictive and prognostic factors will help to guide the oncologist in making informed decisions regarding the optimal radiation dose for treating ESCC, and indicate who have greater possibility benefit from limited radiation dose escalation.

Predictive value of fluorodeoxyglucose (FDG) PET/CT parameters

The ability to identify some factors to predict or assess treatment response at an early stage after the start of treatment would be of great value. Extensive research shows that metabolic-related parameters, such as standardized uptake value (SUV), metabolic tumor volume (MTV), and total lesion glycolysis (TLG) of the primary tumor have the potential to become valuable predictors and prognostic biomarkers in EC patients.⁴⁹⁻⁵²

Kato et al assessed the potential value of metabolicrelated parameters in predicting the response to treatment.53 The SUV_{mean} before CRT in the non-CR group and the CR group were 10.2 and 4.9, respectively. The SUV_{mean} after treatment in the two groups was 3.7 and 1.4, respectively. The changes in SUV of the CR patients were significantly lower than those of the non-CR patients (p < 0.05). The author concluded that the SUV_{mean} before CRT of the primary tumor has the potential to become a valuable predictor for response (p < 0.05). Similarly, Atsumi et al performed a study to assess the efficacy of metabolic-related parameters for the prediction of response in DCRT for EC.54 The results showed that the SUV_{max} values for the CR arm were higher than that in the non-CR arm, and all 18 patients in the low-SUV group had a CR. The data suggest that the SUV_{max} was a valuable predictor for response. However, Javeri et al evaluated the initial standardized unit value of FDGPET and its association with the degree of pathological response after CRT.55 Their work revealed that SUV higher than the median (10.1)was associated with a better pathological response (p=0.06). Similarly, Levine et al and Rizk et al also reported a high initial SUV_{max} was associated with better response.56,57 These conflicting results could be potentially attributed to differences in patient populations, tumor histology types, grouping criteria, as well as treatment, but they might also indicate that predictive value of SUV values are unreliable.

Recently, some studies have considered that SUV_{max}, which reflects only a single point in the tumor, may not always be representative of the whole tumor.⁵⁸ In contrast to SUV_{max}, volumetric parameters such as MTV and TLG represent the dual characteristics of tumor volume and the degree of tumor uptakes FDG.59 Therefore, volumetric parameters based on 18F-FDG PET/CT have been proposed as a more valuable biomarker for predicting survival or response to CRT than SUV_{max} in patients with EC. Roedl et al evaluated the value of volumetric parameters in predicting response to CRT in patients with EC. They found that a decrease in the metabolic tumor diameter between pre- and posttreatment was the single best predictor of tumor response and survival outcome.⁶⁰ In another study, Jayachandran et al evaluated the value of MTV on PET scanning in predicting response to CRT in patients with EC. They found that the MTV2.5 and TLG2.5 were valuable markers for predicting the tumor response.⁶¹ Another study also showed that the changes in MTV and TLG between pre- and posttreatment were more precise for predicting pathological response than ΔSUV_{max} .⁶² However, they showed that none of these parameters were very accurate in predicting a pCR and that the volumetric parameters had a marginally higher

accuracy than SUV_{max} in predicting treatment response to CRT. Overall, available data suggest that these metabolic parameters may be useful as predictors of treatment response, while the ability to predict accurately is still limited.

Predictive value of CT perfusion parameters

CT perfusion is a promising imaging tool in oncology; it can visualize changes in the tumor's vascular physiology and introduces elements of functional diagnostics in morphological imaging.^{63,64} Respecting this fact could be potentially useful in monitoring the response of the tumor to the CRT.

Stefanovic et al evaluated the value of the CT perfusion parameters in predicting response to CRT.⁶⁵ In their study, 40 patients with SCC were reevaluated after CRT. The CT perfusion parameter values after the CRT were significantly correlated with tumor regression grade. These results showed that CT perfusion imaging can predict the response to CRT. Hansen et al also reported that CT perfusion parameters could be an early predictor of treatment response to CRT in EC.⁶⁶

To further investigate the utility of each perfusion parameter for predicting histopathologic response in EC following chemoradiation, a great deal of research was performed. In the study performed by Hayano et al, they found that higher baseline blood flow (BF), higher baseline blood volume (BV) and low mean transit time (MTT) correlated significantly with a good response.⁶⁷ In another EC study, Makari et al examined changes in tumor perfusion before and after chemoradiation in ESCC.⁶⁸ The results showed that responders had a significantly higher baseline BF and a significantly shorter baseline MTT than non-responders, while BV did not in ESCC. Similar to the findings already mentioned, Djuric-Stefanovic et al questioned whether the CT perfusion parameters could be useful to predict the pCR of EC to CRT.⁶⁹ The results showed that BF_{post-CRT}? $BV_{post-CRT}$, and permeability surface_{post-CRT} were significantly lower and MTT_{post-CRT} was significantly higher in the pCR group. The investigators concluded that CT perfusion parameters enable accurate prediction of pCR of EC to CRT, which could be useful in improving patient selection for further treatment.

In conclusion, both FDG PET parameters and CT perfusion parameters could be a good predictor for treatment response. Such predictive factors could help to identify the subgroups that are more likely to benefit from radiation dose escalation.

Individualized radiation dose escalation based on decrease of tumor FDG uptake

The available data indicate that a decrease in tumor FDG uptake correlates with OS and pathological response for

patients with EC.⁷⁰ Cuenca et al performed a study showing that metabolic response during CRT for locally advanced EC has a great prognostic value.⁷¹ Using a 50% decrease in SUV_{max} as a cut-off, the 2-year OS in the good metabolic responders and poor responders was 62% and 27%, respectively (p=0.016). Similarly, Javeri et al declared that the higher the decrease of tumor FDG uptake after treatment, the better the survival of EC patients.72 Metabolic response using PET/ CT is a surrogate for histopathological response in predicting sensitivity to treatments of patients with EC.73 Evaluating the decrease in tumor FDG uptake could help to identify good responders to CRT. Thus, individualized radiation dose escalation based on decrease in tumor FDG uptake after standard radiation of 50.4 Gy could be feasible. According to the European Organization for Research and Treatment of Cancer criteria, metabolic response on FDGPET was divided into the following four types: complete metabolic response (CMR), partial metabolic response (PMR), stable metabolic disease, and progressive metabolic disease.74,75 CMR was defined as complete resolution of 18F-FDG uptake within the tumor volume so that it was indistinguishable from surrounding normal tissue. On the basis of our conjecture, patients with CMR may not need to receive dose escalation after the standard radiation of 50.4 Gy, which may reduce the treatment-related toxicities. Patients with PMR may benefit from dose escalation after a conventional radiation dose of 50.4 Gy. Of those with minor residual tumor after planned radiation, limited dose escalation would achieve CMR without increasing toxicities. These patients could benefit the most from dose escalation. For those with major residual tumor and no metabolic response after planned radiation, limited dose escalation may not change the persistence of local disease because of tumor resistance to CRT. Above all, it is necessary to determine the feasibility of individualized radiation dose escalation after planned chemoradiation based on the decrease in tumor FDG uptake.

Individualized radiation dose escalation based on flow-metabolic phenotypes

It has been reported that malignant tumors differ in terms of BF perfusion and glucose metabolism phenotype.^{76–78} For example, there are: 1) high-perfusion high-metabolism tumors; 2) low-perfusion low-metabolism tumors; 3) highperfusion low-metabolism tumors; and 4) low-perfusion high-metabolism tumors (Figure 1). A combined assessment of the flow-metabolic phenotype of EC using integrated 18F-FDG PET/perfusion CT may be of additional value in assessing the response to therapy as well as in identifying the patients who might be more likely to benefit from



Figure I Representative different flow-metabolic phenotypes. Notes: I: H-P and H-M; II: H-P and L-M; III: L-P and L-M; IV: L-P and H-M. Abbreviations: H-P, high-perfusion; H-M, high-metabolism; L-P, low-perfusion; L-M, low metabolism.

radiation dose escalation. Several studies have investigated the relationship between 18F-FDG PET and perfusion CT, demonstrating that the balance between tumor vascularity and glucose metabolism offers massive information regarding tumor treatment response.⁷⁶⁻⁷⁸ Tumors with matched high perfusion and glucose metabolism show a constitutive up-regulation of angiogenesis and metabolism; tumors with matched low perfusion and glucose metabolism likely reflect necrosis; whereas tumors that present with low perfusion and high glucose metabolism show an adaptation to hypoxia.⁷⁹ Thus, integrated 18F-FDG PET/perfusion CT makes it possible to distinguish different phenotypes; this may be useful in guiding individualized dose escalation based on functional imaging in patients with EC.

Conclusion

In conclusion, the available data reveal that LC after DCRT for EC remains a problem and that most local failures occur within the primary tumor. This indicates that the standard RT dose (50.4 Gy in 28 fractions) may be inadequate to achieve a high probability for LC for some subgroup patients. It is warranted to explore potential ways of improving LC, including IMRT and VMAT techniques, radiation dose escalation, and the use of more effective dose fractionation strategies. Some studies have found that the use of a higher dose and late course of accelerated hyperfraction radiation may lead to better LC and survival for EC patients undergoing CRT. Nevertheless, there is no evidence from Phase III randomized trials to support the additional benefit of dose-escalated RT. Although there is a potential for better tumor LC, there is also an associated higher incidence of toxicity. Therefore, the higher radiation dose should be used with caution on an individual basis in patients with EC.

It has been reported that EC probably has variable sensitivities to CRT. Therefore further studies will be required

to: 1) identify the factors involved in EC sensitivity to radiation; 2) determine the causes of recurrence and non-control from molecular biology perspectives; and 3) individually determine the target region of RT, fractionated dose, and total dose to increase both LC and survival rates and decrease the rates of metastasis in patients with EC.

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Disclosure

The authors report no conflicts of interest in this work.

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