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REVIEW

Concurrent chemoradiation therapy tailored to the older adults with esophageal cancer: state of the art and the future

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Purpose: The aim of this study was to review the published literature addressing the question of whether geriatric assessment (GA) should be routinely applied in the treatment of older adults with esophageal cancer (EC) who have received definitive concurrent chemoradiotherapy (dCRT). Materials and methods: A literature search of PubMed, Embase, and Cochrane Library was performed. Studies that contained original data outlining the inclusion and exclusion criteria, treatment compliance rate, and severe toxicity reports were reviewed. Additionally, criteria from ongoing clinical trials in the World Health Organization and National Institutes of Health registries were reviewed to evaluate the utilization of GA-related domains in elderly EC patients who received dCRT.

Results: Twenty-nine studies were identified based on the selection criteria: five were singlearm prospective studies, and the other studies were retrospective studies. All studies used chronological age and performance status as basic descriptors for this subpopulation. The comorbidity index and the malnutrition level were mentioned in several studies. However, factors such as "Demographic data and social support," "Psychology," "Polypharmacy," and "Geriatric syndromes" were not described in any of the included studies. Unfortunately, the results were similar for the registered clinical trials. Finally, treatment compliance and toxicity profile were found to be acceptable in selected elderly EC patients.

Conclusion: The current experience for older adults with EC receiving dCRT is mainly based on the results of a series of retrospective studies. Ongoing clinical trials should routinely consider GA-related domains to select appropriate treatments for patients in the future.

Keywords: older adults, esophageal cancer, chemoradiotherapy, inclusion criteria, toxicity, geriatric assessment

Introduction

Esophageal cancer (EC) is a prevalent disease with a very poor prognosis. It is the fourth most common cause of cancer death in People's Republic of China.^{1,2} As life expectancy is expected to increase over time, the number of older adults with EC is likely to increase worldwide. However, due to the vast heterogeneity in the aging population, elderly patients are underrepresented in most clinical trials. How to best manage this specific population is still a great challenge. In a recently published study, 21,593 elderly (aged over 70 years) EC patients with clinical stage II or III cancer identified from the National Cancer Database were retrospectively analyzed. Demographic characteristics revealed that 37.1% (8,010/21,593) of patients had received definitive concurrent chemoradiotherapy (dCRT).³ Considering that some patients were diagnosed with clinical stage I or IVa cancer, we believe that more than 37.1%

Clinical Interventions in Aging 2018:13 2275-2287

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of elderly EC patients were recommended dCRT as the firstline of cancer therapy.

Currently, there is consensus within the geriatric oncology community that chronological age alone is a poor descriptor of the heterogeneity that exists within the aging process. Variation in physiological reserves, comorbidities, geriatric syndromes, and limited social support networks also contribute to the risk associated with aggressive treatment modalities. In light of these circumstances, the International Society of Geriatric Oncology (SIOG) and National Comprehensive Cancer Network established recommendations for geriatric assessment (GA) to help oncologists determine suitable treatment choices for older patients.^{4–6} However, the role of GA in the application of dCRT for elderly EC patients remains to be clarified.

Thus, the purpose of this study was to conduct a systemic review of all existing observational cohort studies on the use of GA in the treatment decisions for elderly EC patients receiving dCRT. However, due to the low number of previous studies that used GA in pretreatment evaluation, we summarized the inclusion and exclusion criteria and other important aspects related to GA, including treatment compliance and toxicity for older adults with EC, focusing on whether GA-related domains were routinely applied. We provide some potential suggestions for registered clinical trials in the future.

Materials and methods Search strategy and article selection

A systematic literature search for articles published between January 1, 1990 and December 31, 2017 was conducted in PubMed, Embase, and Cochrane Library with the assistance of a research librarian. The primary outcome was to identify all publications containing original data on the use of dCRT in older adults with EC, including study inclusion and exclusion criteria, treatment compliance, and reports of severe toxicity. The following terms were used as search terms: (esophageal or esophagus or oesophageal) and (cancer or carcinoma or neoplasm or tumor or tumour) and (aged or older or elderly) and concurrent and treatment. Exclusion criteria for search results were as follows: the study involved neoadjuvant chemoradiotherapy (nCRT) with surgery; the study was in Phase I; the study involved palliative chemoradiotherapy; the study was not published with the full text; the study was combined with nCRT or radiotherapy alone; and the study results were not reported exactly. The search did not restrict the type of publication but was limited to English language.

To identify ongoing clinical trials focusing on the efficiency and safety of dCRT for older adults with EC, the World Health Organization registry network (http://www.who.int/ ictrp/network/primary) and the National Institutes of Health (NIH) clinical trial registry (http://www.ClinicalTrials.gov) were searched on December 31, 2017, using the search terms "esophageal cancer" and "radiotherapy." The search was limited to only interventional trials currently recruiting elderly EC patients or intending to start recruitment.

Review methods

The abstract of each article was first reviewed by the corresponding author (SW). Irrelevant citations were removed according to the selection criteria, thereby creating a preliminary set of potentially relevant publications. Then, the fulltext articles were distributed to the review team along with an evaluation form customized for reviewing the GA-related domains previously mentioned: study selection criteria, reports of severe toxicity and treatment compliance (TS and MF). The evaluation results were compared and reevaluated until consensus was reached between the two reviewers. For ongoing clinical trials, the following data were also extracted from the registry website: registration number, country, chemotherapy regimen, inclusion age, and GA-related domains mentioned in the selection criteria.

To allow the combination of performance status data, a score of 100 on the Karnofsky Performance Scale Index (KPS) was considered equivalent to an Eastern Cooperative Oncology Group performance status (ECOG PS) scale score of 0; a score of 80-90 on the KPS was considered equivalent to an ECOG PS score of 1; and a score of 60-70 on the KPS scale was considered equivalent to an ECOG PS score of 2.7 GA-related domains were classified using the following categories: "demographic data and social support," "comorbidity," "functional status," "cognition," "psychology," "nutrition," "polypharmacy," and "geriatric syndromes," following the SIOG recommendations released in 2014.5 In the published reports, restrictions in selection criteria were labeled as "specific criterion mentioned in the report" or "not limited" or "not reported" for each category listed above. In the registered clinical trials, restrictions in selection criteria were labeled as "strict," "mentioned," or "not reported" for GA-related domains.

Results Description of studies and methodological issues

After examination of the relevant abstracts and full-text articles by the review group, a total of 29 published studies involving 1,487 EC patients were incorporated into the systematic review. A flow diagram of citation retrieval and selection is presented in Figure 1. The basic characteristics of patients and GA-related domains mentioned in the methods and results are summarized in Table 1.^{8–36} Of these 29 reports, 5 studies were single-arm prospective clinical trials and the other studies were retrospective studies. Of the five prospective studies, one study used radiation with concurrent targeted therapy of gefitinib. Patient sample size in the eligible studies ranged from 7 to 188. Nearly 84.4% (1,255/1,487) of patients were diagnosed with esophageal squamous cell carcinoma (ESCC). Eleven reports defined the chronological age of 70 as the cutoff age for older adults. Among studies, considerable variation was found across chemotherapy regimens, radiation therapy doses, and radiotherapy parameters. These methodological issues made comparing the results challenging both within study groups and between studies.

GA-related issues

Unfortunately, "demographic data and social support," "psychology," "polypharmacy," and "geriatric syndromes" categories were not described in any of the selected studies (Table S1). Only one prospective study conducted by Servagi-Vernat et al⁸ used Folstein's test as the GA for "Cognition." The other domains are presented in the following.

Functional status

ECOG PS and KPS were the two most commonly used tools of functional status assessment in daily oncological practice. Among the 29 selected studies, 2 studies did not clearly report the functional status of elderly patients. For the remaining 27 studies, 15 studies used the basic ECOG PS score of 0–2 for inclusion, and 6 studies restricted the ECOG PS score to 0–1 for inclusion. Among the 16 studies that clearly demonstrated the proportion of performance status scores across patients, 77.2% (586/759) of the patients were assessed at an ECOG PS score of 0–1, and 173 patients were assessed at an ECOG PS score of 2–3. It is worth mentioning that only one prospective study, also conducted by Servagi-Vernat et al,⁸ used Instrumental Activity Of Daily Living as an additional instrument for evaluating functional status.

Comorbidity

Comorbidity is a characterization factor for elderly cancer patients. In this review, 18 studies mentioned comorbidities in enrolled patients. Among them, the Charlson Comorbidity Index (CCI), or its age-adjusted version (ACCI), was the most common tool for assessing comorbidity. In the five prospective studies, two clinical trials, both conducted by Servagi-Vernat et al,⁸ used inclusion criteria of CCI

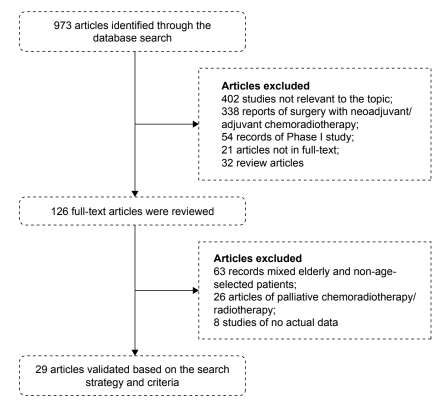


Figure I Schematic presentation of methodology used and selection criteria. Note: Search and selection criteria conducted in accordance with PRISMA statement criteria. Abbreviation: PRISMA, Preferred Reporting Items for Systematic Reviews and Meta-Analyses.

Study	Study design	Histology (SCC/AC/ other)	Age (range)	ст	Radiation dose (Gy)	Comorbidity	Functional status (ECOG PS)	Cognition	Nutrition	Compliance
Servagi-Vernat et al ⁸	Prospective	0/11/61	≥75 (79.4–92)	Single (DDP/OHP)	50	CCI ≤3	0-2 (23/7 0-1/2) and IADL	Folstein's test	Weight loss ≤15%	93.30%
Servagi-Vernat et al ⁹	Prospective	15/7/0	≥75 (75–89)	Single (DDP)	50	CCI ≤4	0-2 (16/6 0-1/2)	NR	Weight loss ≤15%	8001
Ji et al ¹⁰	Prospective	29/1/0	≥65 (65–80)	Single (S-I)	54	NR	0-2 (16/14 0-1/2)	NR	NL (main <10%)	89.00%
Xu et al''	Prospective	20/0/0	≥65 (65–83)	Single (gefitinib)	50.4	NR	5.986 pt	NR	NR	80%
Ohba et al ¹²	Prospective	15/1/0	≥70 (73–81)	Single (DTX)	60	NR	0-1	NR	NR	NR
Zhang et al ¹³	Retrospective	73/0/0	≥65 (65–89)	Double (TP/PF)	60	NL (median CCI 0)	02	R	NR	NR
Song et al ¹⁴	Retrospective	74/6/2	≥70 (70–89)	Double (TP)	60	NL (median	0-2 (63/19	NR	NL (main	67.10%
lvetal ¹⁵	Retrospective	63/5/0	>70 (70–88)	Single (S-1)	54-60	CCI 2) NI (median	0-1/2)	R	<10%) NI (main	70 60%
3					-	CCI 3)	0-1/2)		<10%)	
Xu et al ¹⁶	Retrospective	36/76/0	65-79	Double (5-FU + Tax)	50.4	NL	0-2	NR	NR	95%
Xu et al ¹⁶	Retrospective	16/40/0	≥80 (80–92)			NL	02	NR	NR	
Song et al ¹⁷	Retrospective	31/3/0	≥70	Double (TP)	60	NL (median	0-2 (22/12	NR	NL (main	67.60%
						CCI 2)	0-1/2)		<10%)	
Song et al ¹⁷	Retrospective	31/3/0	≥70	Single (erlotinib)	60	NL (median CCI 2)	0-2 (20/14 0-1/2)	NR	NL (main <10%)	91.20%
Zhao et al' ¹⁸	Retrospective	70/0/0	≥70	Single (S-I/Cape)	61.6	NL	0-3 (63/7	NR	NL (main	NR
							0-1/2-3)		<5%)	
Zhao et al ¹⁸	Retrospective	16/0/0	≥70	Double (DDP + S-I/Cape)	61.6	NL	0-1	NR	NL (main <5%)	NR
Münch et al ¹⁹	Retrospective	19/5/0	≥75 (78–85)	Single/double	7–65	NL (median ACCI 7)	0-2 (23/1 0-1/2)	R	NR	NR
Guo et al ²⁰	Retrospective	16/0/0	≥70 (73–84)	Single (nimotuzumab)	NR	NR	0-2 (12/4 0-1/2)	R	NR	%001
Lu et al ²¹	Retrospective	188/0/0	≥75 (75–107)	Double (PF)	50.7	NL	0-3	NR	NR	52.70%
Uno et al ²²	Retrospective	22/0/0	≥75 (75–85)	Single/double	≥50, m 60	NL	0-2 (20/2	R	R	86.40%
Nallapareddy et al ²³	Retrospective	2/28/0	≥70 (70–86)	Single/double	45-64.8	NR	NR	NR	NR	NR
Kosugi et al ²⁴	Retrospective	24/0/0	≥75 (75–85)	Single/double	56–71, m 67	NR	NR	NR	NR	NR
Xing et al ²⁵	Retrospective	40/0/0	≥ 65	Double (DDP + Cape)	54-60	NR	0-1 (30/10	NR	NL (main ≥5%)	NR
Go et al ²⁶	Retrospective	57/0/0	≥65 (65–84)	Double (mPF)	51	NL	1-2 (54/3 1/2)	NR	NR	75.40%

Takeuchi et al ²⁷	Retrospective	33/0/0	≥70 (71–79)	Double (PF)	60	NL	0-1	NR	R	66.70%
Wakui et al ²⁸	Retrospective	22/0/0	≥75 (75–85)	Double (platinum + 5-FU)	50.4	NR	0-2	NR	NR	NR
Li et al ²⁹	Retrospective	32/0/0	≥70 (70–90)	Single/double	50-60	NR	0-2	NR	NR	NR
Hagiwara et al ³⁰	Retrospective	38/0/0	≥75 (75–89)	Double (PF)	60	NL	01	NR	NR	NR
Tougeron	Retrospective	77/28/4	≥70 (70–88)	Double (PF/DDP + Irino)	49	NL (median	0-2 (87/22 0-1/2)	NR	NL (main <10%)	38.50%
Uno et al ³²	Retrospective	2/0/0	≥80 (80–85)	Single/double	50-60	NR	0-I	NR	NR	NR
Wang et al ³³	Retrospective	56/0/0	≥70 (70–87)	Double (DDP + S-1)	54	NR	0-I	NR	NL (main ≤10%)	67.90%
Anderson et al ³⁴	Retrospective	13/12/0	≥65 (66–88)	Double (5-FU + mitomvcine)	50.4	NL (median CCI 2)	12	NR	NL (main 5–30 lb)	88.00%
Chen et al ³⁵	Retrospective	49/0/0	⇒ 65	Double (DDP + Cape)	56-59.4	NL	0-2 (31/18	NR	NL (main <5%)	NR
Zhao et al ³⁶	Retrospective	52/0/0	≥75 (75–90)	Double (PF)	54-66	NL	0-1/2) 0-2 (46/6 0-1/2)	NR	NL (main =0%)	NR
Note: TP, paclita Abbreviations: EC, esophageal ca	Note: TP, paclitaxel and DDP; PF, 5-FU and DDP. Abbreviations: 5-FU, 5-fluorouracit; AC, adenoca EC, esophageal cancer; IADL, Instrumental Activiti	J and DDP. AC, adenocarcinon ental Activities of I	na; Cape, capecitabine; Daily Living; Irino, irinc	Note: TP, paclitaxel and DDP; PF, 5-FU and DDP. Abbreviations: 5-FU, 5-fluorouracit: AC, adenocarcinoma: Cape, capecitabine: CCI, Charlson Comorbidity Index: CT, chemotherapy; DTX, paclitaxel; ECOGPS, Eastern Cooperative Oncology Group performance status; DDP, cisplatin; EC, esophageal cancer; IADL, Instrumental Activities of Daily Living; Irino, irinotecan; mPF, modified 5-Fu and DDP; NL, not limited; NR, not reported; OHP, oxaliplatin; Ref., reference; SCC, squamous cell carcinoma; Tax, taxotere;	CT, chemotherapy; DP; NL, not limited;	DTX, paclitaxel; ECO NR, not reported; O	GPS, Eastern Coopera HP, oxaliplatin; Ref., r	tive Oncology Gr eference; SCC, so	oup performance s quamous cell carci	tatus; DDP, cisplatin; noma; Tax, taxotere;

dCRT	for	elderly	EC

scores <3 and <4. Among the 24 retrospective studies, 8 studies did not mention any information regarding the comorbidities of their enrolled patients. On the contrary, seven retrospective reports indicated the median CCI/ACCI clearly, with three publications reporting a median CCI of 2.

Nutrition

Malnutrition would certainly increase the risk of antitumor therapy and influence clinical outcomes, especially for elderly EC patients. In our systematic review, we found that the two prospective trials by Servagi-Vernat et al⁸ had criteria excluding patients with weight loss of more than 15% from baseline. In another Phase II trial conducted by Ji et al,¹⁰ weight loss was not used as an exclusion criterion, but over half of the patients (17/30, 56.7%) had <10% weight loss over nearly 6 months. Fourteen of the 24 retrospective studies did not reveal any results related to nutrition. The remaining 10 retrospective studies did not restrict the inclusion criteria in terms of malnutrition, with the main proportion of weight loss ranging from 0% to <10%.

Treatment compliance and severe toxicity

Different chemoradiation therapies could result in variable treatment compliance. In our systematic review, a total of 16 studies detailed the wide range of treatment compliance in elderly EC patients. The compliance rates ranged from 38.5% to 100%. Single-agent chemotherapy or targeted therapy had relatively higher treatment compliance rates (70.6%-100.0%) than did double-agent chemotherapies (38.5%-95.0%). The results of grade 3 or higher acute treatment-related toxicities are also presented in Table S2. Leukopenia was the most common acute toxicity reported in the literature, with incidences ranging from 0% to 72.7%. Ten studies also reported the incidence of severe neutropenia, ranging from 0% to 53.6%. The incidences of two other hematologic toxicities, anemia, and thrombocytopenia were 0%-45.5% and 0%-50.0%, respectively. Esophagitis was the most common radiation-related acute toxicity for elderly EC patients, ranging from 0% to 39.3%. Nausea/vomiting was the second most common during the acute phase. Other acute toxicities (grade >3) were rare with moderate tolerance.

Registered clinical trials

dCRT, definitive chemoradiotherapy

Nine registered clinical trials were found in the WHO and NIH registry network, based on our research methods (Table 2). Among these ongoing studies, six trials distributed across the mainland People's Republic of China. Trial NCT 02735057, conducted in France, was the only study that

₽	Country	Treatment protocol	Age	Demographic data and social support	Comorbidity	Functional status (ECOG PS)	Cognition	Psychological	Nutrition	Polypharmacy	G eriatric syndromes	Note
ChiCTR-OIC- I 701 1436	People's Republic of China	Lobaplatin + RT	≥ 65	NR	Strict	02	Σ	ж	Weight loss ≤10%	Σ	NR	
UMIN000020397	Japan	PTX + RT	70≤ age ≤80	R	Strict	0–2	NR	Σ	R	Σ	NR	
			80	NR	Strict	0-1	R	Σ	NR	Σ	NR	
UMIN000001846	Japan	Docetaxel + RT	70≤ age ≤80	R	Σ	0—1	NR	Σ	R	R	NR	
NCT02716688	People's Republic of China	S-I+RT	N_≤	NR	Strict	01	NR	NR	NR	NR	NR	
NCT02813967	People's Republic of China	S-I+RT	70≤ age ≤85	NR	Strict	10	NR	NR	R	NR	NR	
NCT02375581	People's Republic of China	EGFR-TKI + RT	70≤ age ≤85	NR	Strict	02	Σ	R	R	R	NR	
NCT02979691	People's Republic of China	S-I+RT	N_∕	NR	Strict and CCI ≤3	02	R	R	R	NR	NR	GA as end point
NCT02606916	People's Republic of China	S-I+ DDP + RT	70≤ age ≤80	NR	Strict and CCI ≤4	02	Σ	Σ	Weight Ioss ≤15%	R	NR	
NCT02735057	France	Cab + PTX + RT	75≤ age ≤80	Σ	Strict and CCI ≤3	0-2	Σ	Σ	Weight Ioss ≤15%	R	NR	Pre-GA
			≥80	Σ	Strict and CCI ≤2	0–2	Σ	Σ	Weight Ioss ≤15%	NR	NR	Pre-GA

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detailed their study protocol in another journal.³⁷ Among the nine studies, chronological age, comorbidity, and functional status were commonly established as notable markers for the assessment of the elderly. To our surprise, the inclusion criteria of malnutrition was only strictly restricted in three studies. Other meaningful markers such as "cognitive status," "polypharmacy," and "geriatric syndromes" were sporadically mentioned in these studies.

Discussion

Currently, high quality data from older adults with EC treated with dCRT are extremely limited, according to our systematic review. In the largest retrospective study to date, involving patients over 70 years of age diagnosed with esophageal cancer, Vlacich et al³ showed that over 37.1% of elderly patients received dCRT and had a satisfactory median overall survival (OS) time of 14.0 months (95% CI: 13.5-14.5).³ By contrast, for elderly patients treated with trimodality therapy, the median OS time was only increased to 19.3 months (95% CI: 17.1–21.5) with the added sacrifice of abnormal esophageal function that had an equally important impact on psychological and geriatric syndromes. Using multivariate logistic regression, they found that age over 80 years (odds ratio: 0.79) was a strong indicator of decreased OS likelihood and that patients with clinical stage III, tumors in the upper and middle third of the esophagus, a Charlson-Deyo comorbidity score <1, adenocarcinoma histology, and being treated at a non-academic cancer center were more likely to receive esophagectomy in the USA. Another large sample retrospective analysis from the National Cancer Database provided the opportunity to review treatment utilization and outcomes in patients \geq 80 years of age diagnosed with stage I esophageal cancer. Their analysis revealed that 22% of elderly patients received dCRT among the 923 patients. The survival curve demonstrated a remarkable benefit over observation alone (5-year OS: 20% vs 7%, P < 0.001) with the acceptable toxicity profile of dCRT.³⁸

Apart from known factors that are often associated with treatment disparities such as socioeconomic status and chronologic age, the functional and nutritional status at diagnosis influenced treatment utilization. In 2010, Tougeron et al³⁹ investigated the baseline parameters that influenced both therapeutic decisions and outcomes in 282 elderly EC patients. Their outcomes further suggested that elderly EC patients with good performance status, good nutritional status, and without serious comorbidities were able to benefit from curative treatment without severe adverse events.³⁹ In addition to functional and nutritional status for the elderly

patients, items of psychology and polypharmacy also play an important role in treatment decisions. In a randomized controlled study evaluating the psychological nursing interventions on personality characteristics and quality of life of EC patients, Cheng et al⁴⁰ proved that a psychological nursing intervention could affect the personality characteristics of EC patients and improve quality of life. Furthermore, analyses from elderly Danish cancer patients (including EC patients) showed that newly diagnosed elderly cancer patients (\geq 70 years) use more drugs than control patients (odds ratio: 1.76). Drug use increased markedly in the last 6 months prior to cancer diagnosis.⁴¹ This finding suggested an increased symptom burden in elderly patients and might serve as a warning signal for general practitioners.

In general, elderly EC patients were at increased risk of severe treatment-related toxicities and had a lower incidence of treatment compliance than younger patients. Based on the current review, compliance rates ranged from 38.5% to 100%. Single-agent chemotherapy or targeted therapy seemed to have higher treatment compliance (70.6%-100.0%) than did double-agent chemotherapies (38.5%-95.0%). Unlike combined therapies for non-age-selected patients, our recent Phase III trial demonstrated that dCRT (TP regimen and radiation therapy) with erlotinib could achieve significantly better OS and locoregional control than standard dCRT in patients with inoperable ESCC. Furthermore, this trial confirmed that, aside from rash and radiation esophagitis, the incidence of grade 3 or greater toxicities were manageable and did not differ across the four groups.42 Traditionally, it was thought that elderly patients have fewer body function reserves, shorter life expectancy, and more need for socioeconomic support than young patients. Therefore, this finding suggests that aggressive treatment decisions might not be suitable for this unique and challenging subpopulation. In a small-sample retrospective analysis, Wang et al³³ observed a significantly higher incidence of severe leukocytopenia in ESCC patients aged \geq 75 than in patients aged between 70 and 75 years (41.9% vs 72.0%, respectively, P=0.024). Hence, a detailed and full pretreatment evaluation, especially containing GA-related domains should be routinely applied to select appropriate elderly patients to receive dCRT.

Conclusion

Our systematic review fully demonstrates that the current experience of receiving dCRT in elderly patients is based on the results of a series of retrospective studies. High-level evidence of RCTs remains lacking. Ongoing prospective clinical trials should take GA-related domains into routine consideration to select appropriate care for patients in the future. Survival benefits and the toxicity profiles should also be balanced among these treatment strategies.

Acknowledgments

The authors thank Li Zhang, MD, and Zhao Jing, MD, for their help with translation and proofreading. This research received no specific grant from any funding agency in the public, commercial, or not-for-profit sectors.

Disclosure

The authors report no conflicts of interest in this work.

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Study	Study design	Histology (SCC/ AC/other)	Age (range)	сŢ	Radiation dose (Gv)	Demographic data and social support	Comorbidity	Functional status (ECOG PS)
Servagi-Vernat et al ¹	Prospective	0/11/61	≥75 (79.4–92)	Single (DDP/OHP)	50	NR.	CCI ≤3	0-2 (23/7 0-1/2) and IADL
Servagi-Vernat et al ²	Prospective	15/7/0	≥75 (75–89)	Single (DDP)	50	NR	CCI ≤4	0-2 (16/6 0-1/2)
Ji et al ³	Prospective	29/1/0	≥65 (65–80)	Single (S-1)	54	NR	NR	0-2 (16/14 0-1/2)
Xu et al ⁴	Prospective	20/0/0	≥65 (65–83)	Single (gefitinib)	50.4	NR	NR	0-2 (18/2 0-1/2)
Ohba et al ⁵	Prospective	15/1/0	≥70 (73–81)	Single (DTX)	60	NR	NR	0-1
Zhang et al ⁶	Retrospective	73/0/0	≥65 (65–89)	Double (TP/PF)	60	NR	NL (median CCI 0)	0-2
Song et al ⁷	Retrospective	74/6/2	≥70 (70–89)	Double (TP)	60	NR	NL (median CCI 2)	0-2 (63/19 0-1/2)
Lv et al ⁸	Retrospective	63/5/0	≥70 (70–88)	Single (S-1)	54-60	NR	NL (median CCI 3)	0-2 (42/26 0-1/2)
Xu et al ⁹	Retrospective	36/76/0	65–79	Double (5-FU + Tax)	50.4	NR	NL	0-2
Xu et al ⁹	Retrospective	16/40/0	≥80 (80–92)	Double (5-FU + Tax)		NR	NL	0-2
Song et al ¹⁰	Retrospective	31/3/0	≥70	Double (TP)	60	NR	NL (median CCI 2)	0-2 (22/12 0-1/2)
Song et al ¹⁰	Retrospective	31/3/0	≥70	Single (erlotinib)	60	NR	NL (median CCI 2)	0-2 (20/14 0-1/2)
Zhao et al ^{III}	Retrospective	70/0/0	≥70	Single (S-I/Cape)	61.6	NR	NL	0-3 (63/7 0-1/2-3)
Zhao et al''	Retrospective	16/0/0	≥70	Double (DDP + S-1/Cape)	61.6	NR	NL	0-1
Münch et al ¹²	Retrospective	19/5/0	≥75 (78–85)	Single/double	7–65	NR	NL (median ACCI 7)	0-2 (23/1 0-1/2)
Guo et al ¹³	Retrospective	16/0/0	≥70 (73–84)	Single (nimotuzumab)	NR	NR	NR	0-2 (12/4 0-1/2)
Lu et al ¹⁴	Retrospective	188/0/0	≥75 (75–107)	Double (PF)	50.7	NR	NL	0–3
Uno et al ¹⁵	Retrospective	22/0/0	≥75 (75–85)	Single/double	≥50, m 60	NR	NL	0-2 (20/2 0-1/2)
Nallapareddy et al ¹⁶	Retrospective	2/28/0	≥70 (70–86)	Single/double	4564.8	NR	NR	NR
Kosugi et al ¹⁷	Retrospective	24/0/0	≥75 (75–85)	Single/double	56–71, m 67	NR	NR	NR
Xing et al ¹⁸	Retrospective	40/0/0	≥65	Double (DDP + Cape)	54-60	NR	NR	0-1 (30/10 0-1/2)
Go et al ¹⁹	Retrospective	57/0/0	≥65 (65–84)	Double (mPF)	51	NR	NL	1-2 (54/3 1/2)
Takeuchi et al ²⁰	Retrospective	33/0/0	≥70 (71–79)	Double (PF)	60	NR	NL	0-1
Wakui et al ²¹	Retrospective	22/0/0	≥75 (75–85)	Double (platinum +5-FU)	50.4	NR	NR	02
Li et al ²²	Retrospective	32/0/0	≥70 (70–90)	Single/double	50-60	NR	NR	02
Hagiwara et al ²³	Retrospective	38/0/0	≥75 (75–89)	Double (PF)	60	NR	NL	0-1
Tougeron et al ²⁴	Retrospective	77/28/4	≥70 (70–88)	Double (PF/DDP + Irino)	49	NR	NL (median CCI I)	0-2 (87/22 0-1/2)
Uno et al ²⁵	Retrospective	7/0/0	≥80 (80–85)	Single/double	50-60	NR	NR	0-1
Wang et al ²⁶	Retrospective	56/0/0	≥70 (70–87)	Double (DDP + S-I)	54	NR	NR	0-1
Anderson et al ²⁷	Retrospective	13/12/0	≥65 (66–88)	Double (5-FU +	50.4	NR	NL (median CCI 2)	1–2
	_			mitomycine)				
Chen et al ²⁸	Retrospective	49/0/0	≥65	Double (DDP + Cape)	5659.4	NR	NL	0-2 (31/18 0-1/2)
Zhao et al ²⁹	Retrospective	52/0/0	≥75 (75–90)	Double (PF)	54-66	NR	NL	0-2 (46/6 0-1/2)

Table S1 Baseline characteristics of the published reports in elderly EC patients who received dCRT or concurrent radiation with

Supplementary materials

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Study	Cognition	Nutrition	Psychological	Polypharmacy	Geriatric syndromes	Compliance
Servagi-Vernat et al ^l	Folstein's test	Weight loss ≤15%	NR	NR	NR	93.30%
Servagi-Vernat et al ²	NR	Weight loss \leq 15%	NR	NR	NR	8001
Ji et al ³	NR	NL (main <10%)	NR	NR	NR	89.00%
Xu et al⁴	NR	NR	NR	NR	NR	80%
Ohba et al ⁵	NR	NR	NR	NR	NR	NR
Zhang et al ⁶	NR	NR	NR	NR	NR	NR
Song et al ⁷	NR	NL (main <10%)	NR	NR	NR	67.10%
Lv et al ⁸	NR	NL (main <10%)	NR	NR	NR	70.60%
Xu et al ⁹	NR	NR	NR	NR	NR	95%
Xu et al ⁹	NR	NR	NR	NR	NR	
Song et al ¹⁰	NR	NL (main <10%)	NR	NR	NR	67.60%
Song et al ¹⁰	NR	NL (main <10%)	NR	NR	NR	91.20%
Zhao et al''	NR	NL (main <5%)	NR	NR	NR	NR
Zhao et al''	NR	NL (main <5%)	NR	NR	NR	NR
Münch et al ¹²	NR	NR	NR	NR	NR	NR
Guo et al ¹³	NR	NR	NR	NR	NR	8001
Lu et al ¹⁴	NR	NR	NR	NR	NR	52.70%
Uno et al ¹⁵	NR	NR	NR	NR	NR	86.40%
Nallapareddy et al ¹⁶	NR	NR	NR	NR	NR	NR
Kosugi et al ¹⁷	NR	NR	NR	NR	NR	N R
Xing et al ¹⁸	NR	NL (main ≥5%)	NR	NR	NR	NR
Go et al ¹⁹	NR	NR	NR	NR	NR	75.40%
Takeuchi et al ²⁰	NR	NR	NR	NR	NR	66.70%
Wakui et al ²¹	NR	NR	NR	NR	NR	NR
Li et al ²²	NR	NR	NR	NR	NR	NR
Hagiwara et al ²³	NR	NR	NR	NR	NR	NR
Tougeron et al ²⁴	NR	NL (main <10%)	NR	NR	NR	38.50%
Uno et al ²⁵	NR	NR	NR	NR	NR	NR
Wang et al ²⁶	NR	NL (main ≤10%)	NR	NR	NR	67.90%
Anderson et al ²⁷	NR	NL (main 5–30 lb)	NR	NR	NR	88.00%
Chen et al ²⁸	NR	NL (main <5%)	NR	NR	NR	NR
Zhao et al ²⁹	NR	NL (main =0%)	NR	NR	NR	NR

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