Survival outcomes in esophageal cancer patients with a prior cancer

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Abstract
To achieve a deeper understanding of patients who developed esophageal cancer (EC) as a second primary malignancy, which may help guide in clinical practice for these patients in the future.

In the primary cohort, EC patients with a prior malignancy were identified from the surveillance, epidemiology, and end result database. The 8 most common types of prior cancers were picked out based on the frequency of occurrence. In addition, Kaplan-Meier and log-rank tests were performed to investigate the survival impacts of prior cancers on EC patients. Besides, a competing-risk model was constructed to explore the relationship between EC-treatment and EC-specific mortality. In the secondary cohort, patients with stage I–III (NOMO) EC from 2004 to 2014 were enrolled. After propensity score matching, univariate and multivariate Cox analyses were developed to determine the prognostic factors for EC patients.

A total of 1199 EC patients with a prior cancer were identified in the primary cohort. The 5 most common sites of prior cancers were prostate, female breast, lung and bronchus, and larynx. Kaplan-Meier analyses revealed that EC patients with prior prostate cancer and bladder cancer had the best overall survival (OS), while those with prior cancers of larynx and lung and bronchus had the worst OS. Fine and Gray competing risks analysis indicated that the administration of surgery was closely associated with better EC-specific survival (P < .001). In the secondary cohort, multivariate Cox analyses found that age at diagnosis, race, tumor grade, tumor extent, nodal status and metastasis stage, histology, and the administration of surgery were prognostic factors for OS and cancer-specific survival in EC patients. Besides, the existence of a prior cancer was an independent prognostic factor for cancer-specific survival.

EC remains to be the most important cause of death in EC patients with a prior cancer. EC related treatment should be actively adopted in patients with a prior cancer, as they were more likely to die from EC than the prior cancer. EC patients with a prior cancer had comparable OS than those without.

Abbreviations: AC = adenocarcinomas, COD = cause of death, CSS = cancer-specific survival, EC = esophageal cancer, ECSM = EC-specific mortality, IQR = interquartile range, IR = incidence rate, OS = overall survival, PEC = primary esophageal cancer, PSM = propensity score matching, SCC = squamous cell carcinomas, SEC = subsequent esophageal cancer, SEER = surveillance, epidemiology, and end result, SPM = second primary malignancy, US = United States.

Keywords: esophageal cancer, second primary malignancy, surveillance, epidemiology, and end result, survival

1. Introduction
Esophageal cancer (EC) is one the most common malignancies, the incidence rate (IR) ranked ninth of all malignant tumors worldwide in 2018.[1] In 2020, the estimated new cases and deaths were 18,440 and 16,170 in the United States (US).[2] Surgery and radiotherapy have been the standard treatment types of EC for many years. Nowadays, rapid development of immunotherapy and targeted therapy (such as trastuzumab) of EC has brought a tremendous promise in the treatment of EC.[3,4] Moreover, the 5-year survival rate of EC patients has increased from 10% to 25% due to the advancement of cancer detection and treatment.[5,6] Hence, more and more cancer survivors developed a second primary malignancy (SPM) because of the increasing IRs and improvement of survival outcomes.[7,8]

SPM is defined as a cancer which develops in a new tissue or organ after the initial diagnosis of the prior malignancy with a 6-month latency. Previous studies mainly focused on the risk of developing an SPM after a known malignancy. Liao et al.[9] discussed the main prognostic factors for oral cavity cancer patients with simultaneous SPM, and then developed a risk-stratification. Vassilev et al.[10] provided a historical risk estimation of developing an SPM in patients with metastatic castration-resistant prostate cancer. However, as far as we know, survival outcomes of patients with 1 known tumor as an SPM have not been well studied. Only a few published studies have discussed the risk of developing an SPM in primary cancer survivors.[11,12] Saad et al.[13] investigated the impact of the prior cancer on survival outcomes of stage IV EC patients, they found
that prior cancers did not adversely impact survival of EC patients with stage IV diseases. Besides, Chen et al. explored clinicopathological characteristics and prognosis of patients with EC as an SPM, they demonstrated that lower M stage, the administration of surgery, and chemotherapy were tightly related to better overall survival (OS) for patients with EC as an SPM.

In this study, patients diagnosed with EC as an SPM were extracted from the surveillance, epidemiology, and end result (SEER) database retrospectively. We aimed to achieve a deeper understanding of the outcomes of patients who developed EC as an SPM, which may help guide in clinical practice for these patients in the future.

2. Materials and methods

2.1. Database

Data were extracted from the SEER database retrospectively. It is a population-based registry sponsored by the US National Cancer Institute. The SEER database collects relevant information of cancer IR, baseline characteristics, treatment types and long-term follow-up, and covers approximately 34.6% of the US population till now (https://seer.cancer.gov/about/overview.html). We signed the Research Data Agreement before this study and got access to the database with the username of 11015-Nov2019. In addition, use of SEER registry was exempt by Institutional Review Board approval.

2.2. Primary cohort

In this section, we extracted EC patients with a prior malignancy from the SEER 18 program using the “multiple primary-standard incidence ratio" function by the SEER*Stat software (version 8.3.6; US National Cancer Institute, Bethesda, Maryland, USA). EC was diagnosed as the SPM with positive pathology. Furthermore, the exclusion criteria were as follows:

1. patients with more than 2 malignancies in total,
2. data were from autopsy or death certificate only,
3. year of diagnosis was not from 2004 to 2014,
4. patients with missing or unknown data,
5. interval between diagnosis of EC and the prior cancer was less than 6 months.

A 6-month latency was utilized to distinguish SPMs from simultaneous cancers. In our study, the International Classification of Diseases for Oncology third edition primary site codes for EC contained C15.0 (cervical esophagus), C15.1 (thoracic esophagus), C15.2 (abdominal esophagus), C15.3 (upper third of esophagus), C15.4 (middle third of esophagus), C15.5 (lower third of esophagus), C15.8 (overlapping lesion of esophagus), and C15.9 (esophagus, not otherwise specified). To be specific, codes C15.0 and C15.3 were used to identify upper esophageal tumors, while C15.4 was for middle esophageal tumors and C15.2 and C15.5 were for lower esophageal tumors. Moreover, histologic recode broad groupings were applied for the classification of histological subtypes (codes 8140-8389 were for adenomas and adenocarcinomas (AC), codes 8050-8089 were for squamous cell carcinomas (SCC) and all other remaining codes as other histology.

Then, demographic characteristics and clinical data for each patient were collected, including age at diagnosis (both prior cancer and EC), sex, race, histological type, primary sites of EC, American Joint Committee on Cancer 6th tumor extent, nodal status and metastasis (TNM) stage, diagnosis intervals, the administration of surgery, radiotherapy and chemotherapy, vital status, cause of death (COD) and follow-up. Age at diagnosis was categorized into <65 and ≥65 years old. Furthermore, CODs were classified into 3 groups: died from EC, died from the prior cancer, and died from other causes.

First of all, we picked out the 5 most common types of prior cancers based on the frequency of occurrence. Then, Kaplan-Meier and log-rank tests were performed to investigate the survival impacts of prior cancers on EC patients. Afterward, the percentage of EC-related and prior cancer-related deaths in patients with different prior malignancies were calculated, and the ratios of EC deaths to prior cancer deaths were obtained, further stratified by EC TNM stage and histological type. Finally, to explore the relationship between the administration of surgery and EC-specific mortality (ECSM), we constructed a competing model after taking died from other causes/prior cancers as a competing event.

2.3. Secondary cohort

In the secondary cohort, we identify patients with stage I-III (NOMO) EC from 2004 to 2014 in the SEER 18 database using the “case listing session” function. Based on the existence of a prior malignancy, all patients were then divided into “primary esophageal cancer (PEC)” and “subsequent esophageal cancer (SEC).” Propensity score matching (PSM) method was used to balance the basic characteristics of PEC and SEC patients with a ratio of 1:1. Survival discrepancies between PEC and SEC patients were compared before and after PSM. Lastly, univariate and multivariate Cox analyses were developed to discuss the prognostic factors which were significantly related to OS and cancer-specific survival (CSS) in patients with EC.

2.4. Statistical analysis

Student t test and Mann–Whitney U test were used for the comparisons of continuous variables. Chi-square analysis was utilized to make comparisons between categorical variables. The whole analysis was based on SPSS 23.0 (SPSS Inc, Chicago, IL) and R software (Version 3.4.1). A 2-sided P <.05 was considered significant.

3. Result

3.1. Baseline characteristics of the primary cohort

A total of 1199 EC patients with a prior cancer were eventually enrolled in the primary cohort. As shown in Table 1, the median (interquartile range [IQR]) ages at EC and the prior cancer diagnosis were 73.00 (66.00–80.00) and 64.00 (57.00–71.00) years old, respectively. Most patients were White (85.99%) and male (78.73%). The most common site of EC was lower esophagus (61.38%), 54.38% of the EC patients were with AC. The median (IQR) diagnosis interval between the prior cancer and EC was 91.00 (43.99–151.00) months. Moreover, the median (IQR) follow-up since EC diagnosis was 12.00 (4.00–30.00) months.

3.2. Survival outcomes in the primary cohort

The 5 most common sites of prior cancers were prostate (35.36%), female breast (8.42%), bladder (7.84%), lung and bronchus (5.75%), and larynx (4.50%) (Table 2). OS was
Table 1
Demographic and clinical factors of EC patients with a prior cancer (n = 1199).

| Variables                  | Value                      |
|----------------------------|----------------------------|
| Age, yr                    |                            |
| Mean (SD)                  | 63.23 (11.65)              |
| Median (IQR)               | 64.00 (57.00–71.00)        |
| Sex, n (%)                 |                            |
| Male                       | 944 (78.73)                |
| Female                     | 255 (21.27)                |
| Race, n (%)                |                            |
| White                      | 1031 (85.90)               |
| Black                      | 111 (9.26)                 |
| Other                      | 57 (4.75)                  |
| At EC diagnosis            |                            |
| Age, yr                    |                            |
| Mean (SD)                  | 73.46 (9.99)               |
| Median (IQR)               | 73.00 (66.00–80.00)        |
| Primary site, n (%)        |                            |
| Upper                      | 129 (10.76)                |
| Middle                     | 193 (16.10)                |
| Lower                      | 736 (61.38)                |
| Other                      | 141 (11.76)                |
| Histology, n (%)           |                            |
| AC                         | 652 (54.38)                |
| SCC                        | 453 (37.76)                |
| Other                      | 92 (7.87)                  |
| TNM stage, n (%)           |                            |
| I–II                       | 634 (52.88)                |
| III-IV                     | 565 (47.12)                |
| Interval between diagnoses, mo |                |
| Mean (SD)                  | 110.77 (87.27)             |
| Median (IQR)               | 91.00 (43.00–151.00)       |
| Time from EC diagnosis to death or end of study, mo | |
| Mean (SD)                  | 22.76 (27.67)              |
| Median (IQR)               | 12.00 (4.00–30.00)         |

AC = adenocarcinoma, EC = esophageal cancer, IQR = interquartile range, SCC = squamous cell carcinoma, SD = standard deviation, TNM = tumor extent, nodal status and metastasis.

significantly different in EC patients with different prior malignancies (P < .0001, Fig. 1). EC patients with prior prostate cancer and bladder cancer had the best survival outcomes (3-year OS rates were 27.7% and 29.2%, respectively), while those with prior cancer of larynx and lung and bronchus had the worst OS (3-year OS rates were 12.5% and 11.0%, respectively).

In the analysis of COD, 65.51% of EC patients died from EC and 16.75% of patients died from the prior cancer (Fig. 2). EC patients with prior cancers of lung and bronchus had the highest prior cancer-related death rate (26.15%) and the lowest EC-related death rate (58.46%). Furthermore, the ratios of prior cancer-related deaths to EC-related deaths were calculated. As shown in Figure 3, the ratios were less than 1 regardless of the histological type (Fig. 3A) or TNM stage (Fig. 3B) of EC. Hence, conclusion could be drawn that EC patients were more likely to die of EC regardless of the cancer types of prior cancers and EC.

Compared with patients who died from the prior cancer, those who died from EC had older ages at cancer diagnosis (both EC and the prior cancer) (all P < .05, Table 3). In addition, the proportions of AC and N1 diseases (all for EC) were significantly higher in patients who died from EC. The median interval between diagnosis of 2 cancers was significantly longer in patients who died from EC than that in patients who died from the prior cancer (92.00 vs 66.00 months, P < .001). Notably, the percentage of radiotherapy in patients who died from EC was significantly higher than those who died from the prior cancer (62.93% vs 53.85%, P = .031). To explore the prognostic role of cancer treatments, Fine and Gray competing risks analyses were developed. As shown in Figure 4, the administration of surgery was tightly related to better EC-specific survival (P < .001).

3.3. Survival of patients with EC as the prior cancer or subsequent primary cancer in the second cohort

From 2004 to 2014, a total of 7230 patients with stage I-III EC were enrolled in the secondary cohort, including 5281 (73.04%) patients had EC as the only malignancy (PEC) and 1949 (26.96%) patients with EC following a prior cancer (defined as SEC) (Table 4). SEC patients had significantly older age than PEC patients (265 years old: 77.68% vs 58.97%, P < .001). Furthermore, the proportions of male patients, lower esophageal tumors, histology of AC, higher stage (II–III) diseases, and the administration of surgery/radiotherapy/chemotherapy were significantly higher in SEC patients when compared with these in PEC patients (all P < .05). Therefore, a 1:1 PSM was applied to minimize the difference between SEC and PEC patients in baseline characteristics and treatment types. Eventually, a total of 1949 pairs of EC patients were included.

Supplemental Digital Content (Figure S1, http://links.lww.com/MD/F706) shows the comparisons of survival outcomes between SEC and PEC patients. After matching, there was no significant difference in OS between patients in 2 groups (Fig. 5A, P > .05). However, SEC patients had better CSS than PES patients (Fig. 5B, P < .05). Furthermore, subgroup analyses based on different histological types (AC and SCC) revealed the same results (Fig. 5C–F).

Multivariate Cox analysis indicated that age at diagnosis, race, tumor grade, TNM stage, histology, and the administration of surgery were prognostic factors for OS and CSS in EC patients (Tables 5 and 6). Besides, the existence of a prior cancer (PEC vs SEC) was an independent risk factor for CSS (P < .001).

4. Discussion

In recent years, the number of cancer survivors is rapidly increasing due to the improvement of cancer screening and treatment. Hence, the risk of developing an SPM in cancer survivors has also been increasing. It was reported that there was a 2% annual increase for the cancer survivor population in the US, and about 18% of cancer survivors developed an SPM during the rest of their lifetime according to the SEER registry. Furthermore, the history of a prior cancer played a critical role in making clinical decision, especially for those who participated in clinical trials. In many clinical trials, history of a prior cancer was a strict exclusion criterion for potential candidates, which may be
Figure 1. Overall survival of esophageal cancer patients with a prior cancer.

Figure 2. Distribution of causes of death in the top 5 most common sites of developing SPMs in EC patients. EC = esophageal cancer, SPM = second primary malignancy.
due to the survival impacts of the prior cancers.\textsuperscript{[16]} Although there was no powerful evidence supporting the hypothesis that exclusion of these patients could balance the outcomes and validity of clinical trials,\textsuperscript{[13]} many published trials excluded patients with a prior cancer routinely.\textsuperscript{[17]-[19]} A previous study revealed that there were approximately 20\% of lung cancer patients were excluded because of this restrictive exclusion rule.\textsuperscript{[18]} This study was to investigate the survival outcomes of EC patients with a prior cancer and to identify prognostic factors for EC patients.

In this study, the most common prior malignancy in EC patients was prostate cancer, followed by female breast cancer, bladder cancer, and lung cancer. Interestingly, these cancers are also the most common cancers as single malignancy in general. Hence, we guessed that there was no enrichment for a cancer type that may increase the risk of developing EC as an SPM. Similarly, Zhu et al.\textsuperscript{[20]} reported that the most common types of prior cancers in larynx cancer patients were from prostate, lung and bronchus, urinary bladder, and breast. Laccetti et al.\textsuperscript{[21]} found that prostate, gastrointestinal, breast, and other genitourinary were the most common types of prior cancer in locally advanced lung cancer.

Comparisons in survival outcomes of EC patients with different prior cancers showed significant statistical difference. EC patients with prior cancers of prostate cancer and bladder cancer had significant better OS than those with prior cancers of lung and bronchus. The survival discrepancy may be due to the level of threat to life of prior cancers. Moreover, EC patients were more likely to die of EC regardless of the cancer types of prior cancers and EC. Lastly, multivariate Cox analyses found that age, race, tumor grade, TNM stage, histology, and the administration of surgery were independent prognostic factors for OS and CSS in EC patients, and the existence of a prior cancer was an independent risk factor for CSS.

Most patients died from EC rather than the prior cancer (65.51\% vs 16.75\%) with a median follow-up of 12.00 months. Furthermore, subgroup analyses based on TNM stage and histology (AC and SCC) revealed the same results. Moreover, Kaplan–Meier analysis showed that PEC patients had similar OS.
### Table 3
Clinical and demographic factors associated with EC death versus prior cancer death.

| Characteristics                          | Died from prior cancer | Died from EC | P-value |
|------------------------------------------|------------------------|--------------|---------|
| Number of patients                       | 169                    | 661          |         |
| Age at EC diagnosis, median (IQR), yr    | 70.00 (62.00–76.00)    | 75.00 (67.00–81.00) | < .001  |
| Age at prior cancer diagnosis, median (IQR), yr | 62.00 (54.00–70.00)    | 65.00 (57.50–72.00) | .021    |
| EC, histology, n (%)                     |                        |              | < .001  |
| AC                                       | 59 (34.91)             | 345 (52.19)  |         |
| SCC                                      | 104 (61.54)            | 252 (38.12)  |         |
| Other                                    | 6 (3.55)               | 64 (9.68)    |         |
| EC, surgery treated, n (%)               |                        |              | .572    |
| No                                       | 133 (78.70)            | 533 (80.64)  |         |
| Yes                                      | 36 (21.30)             | 128 (19.36)  |         |
| EC, radiotherapy treated, n (%)          |                        |              | .031    |
| No/unknown                               | 78 (46.15)             | 245 (37.07)  |         |
| Yes                                      | 91 (53.85)             | 416 (62.93)  |         |
| EC, chemotherapy treated, n (%)          |                        |              | .074    |
| No/unknown                               | 74 (43.70)             | 240 (36.31)  |         |
| Yes                                      | 95 (56.21)             | 421 (63.69)  |         |
| EC, TNM stage, n (%)                     |                        |              | .063    |
| I-II                                     | 91 (53.85)             | 303 (45.84)  |         |
| III-IV                                   | 78 (46.15)             | 358 (54.16)  |         |
| EC, Tx/N1/Mx, n (%)                      |                        |              | .010    |
| No/unknown                               | 71 (42.01)             | 351 (53.10)  |         |
| Yes                                      | 42 (24.85)             | 194 (29.35)  | .247    |
| EC, grade I-II, n (%)                    |                        |              | .077    |
| 127 (75.15)                              |                         | 537 (81.24)  |         |
| Interval between diagnoses, median (IQR), mo | 66.00 (32.50–112.00)   | 92.00 (48.00–163.50) | < .001  |

AC = adenocarcinoma, EC = esophageal cancer, IQR = interquartile range, SCC = squamous cell carcinoma, SD = standard deviation, TNM = tumor extent, nodal status and metastasis.

Figure 4. Estimates of overall cumulative incidence of developing a second malignancy, taking surgery as a competing event.
Saad et al. [13] found that stage IV EC patients with a prior cancer had comparable OS with those had EC as their only malignancy. In that study, Saad et al. only focused on the survival impact of prior cancers on the advanced EC patients, rather than all EC patients. Similarly, Chen et al. [14] investigated the clinicopathological characteristics and survival outcomes of EC patients with a prior cancer, they found that the most common prior malignancy in EC patients was from genital system (about 43.5%). Moreover, EC patients with a prior cancer had comparable OS when compared with only primary EC patients. However, previous studies did not investigate the EC-specific survival. In our study, SEC patients had significant better CSS than PEC patients after matching. Better CSS could be attributed to the fact that cancer survivors receiving a stricter screening and care or being more cautious on healthy problems. Furthermore, Wang et al. [22] reported that nasopharyngeal carcinoma patients with a prior cancer had better CSS than those without a prior cancer. However, study conducted by Ji et al. [23] and Al-Husseini et al. [24] reached the opposite conclusions that breast cancer or glioblastoma patients with a prior malignancy had worse CSS than those had breast cancer or glioblastoma as their only malignancy.

In our study, the proportion of surgery was comparable in patients who died from EC with that in patients who died from the prior cancer. Interestingly, Fine and Gray competing analysis showed that the administration of surgery was closely related to a reduction of ECSM. Our findings strongly indicated that surgery was still an optional alternative for EC patients with a prior cancer. First, most EC patients with a prior cancer died from EC rather than the prior cancer, regardless of the clinical characteristics of the prior cancer and EC. Second, prolonged CSS was detected in SEC patients when compared with PEC patients. Dinh et al. [12] found that treatment for patients with high stage and high-grade prostate cancer was related to a decreased risk of prostate cancer-specific mortality.

Cox regression analyses revealed that age at diagnosis, race, tumor grade, TNM stage, histology, and the administration of surgery were prognostic factors for OS and CSS in EC patients.

### Table 4

Baseline characteristics of patients with PEC or SEC from the SEER database 2004-2014.

| Variables       | PEC before PSM | SEC before PSM | P-value | PEC after PSM | SEC after PSM | P-value |
|-----------------|----------------|----------------|---------|---------------|---------------|---------|
| N               | 5281           | 1949           |         | 1949          | 1949          | .393    |
| Age (yr)        |                |                | <.001   |                |                | .112    |
| <65             | 2167 (41.03)   | 435 (22.32)    |         | 413 (21.19)   | 435 (22.32)   |         |
| ≥65             | 3114 (58.97)   | 1514 (77.68)   |         | 1536 (78.81)  | 1514 (77.68)  |         |
| Race            |                |                |         | .299          | .299          |         |
| White           | 4442 (84.11)   | 1660 (85.17)   |         | 1678 (86.10)  | 1660 (85.17)  |         |
| Black           | 583 (11.04)    | 217 (11.13)    |         | 190 (9.75)    | 217 (11.13)   |         |
| Other           | 256 (4.85)     | 72 (3.69)      |         | 81 (4.16)     | 72 (3.69)     |         |
| Sex             |                |                | <.001   |                |                | 1.000   |
| Male            | 4066 (77.37)   | 1388 (71.22)   |         | 1388 (71.22)  | 1388 (71.22)  |         |
| Female          | 1195 (22.63)   | 561 (28.78)    |         | 561 (28.78)   | 561 (28.78)   |         |
| Location        |                |                | <.001   |                | .146          |         |
| Upper           | 461 (8.73)     | 322 (16.52)    |         | 278 (14.26)   | 322 (16.52)   |         |
| Middle          | 1055 (19.98)   | 444 (22.78)    |         | 461 (23.65)   | 444 (22.78)   |         |
| Lower           | 3765 (71.29)   | 1183 (60.70)   |         | 1210 (62.08)  | 1183 (60.70)  |         |
| Grade           |                |                | .464    |                | .702          |         |
| Grade I         | 494 (9.35)     | 170 (8.72)     |         | 155 (7.95)    | 170 (8.72)    |         |
| Grade II        | 2485 (47.06)   | 926 (47.51)    |         | 928 (47.61)   | 926 (47.51)   |         |
| Grade III       | 2217 (41.98)   | 830 (42.59)    |         | 837 (42.95)   | 830 (42.59)   |         |
| Grade IV        | 85 (1.61)      | 23 (1.18)      |         | 29 (1.49)     | 23 (1.18)     |         |
| Histology       |                |                | <.001   |                | .061          |         |
| AC              | 3156 (59.76)   | 967 (49.62)    |         | 1034 (53.05)  | 967 (49.62)   |         |
| SCC             | 1772 (33.55)   | 868 (44.54)    |         | 795 (40.79)   | 868 (44.54)   |         |
| Other           | 353 (6.68)     | 114 (5.83)     |         | 120 (6.16)    | 114 (5.83)    |         |
| TNM stage       |                |                | <.001   |                | .798          |         |
| I               | 2616 (49.54)   | 1102 (56.54)   |         | 1121 (57.52)  | 1102 (56.54)  |         |
| II              | 2176 (41.20)   | 682 (34.99)    |         | 671 (34.43)   | 682 (34.99)   |         |
| III             | 489 (9.26)     | 165 (8.47)     |         | 157 (8.06)    | 165 (8.47)    |         |
| Surgery         |                |                | <.001   |                | .383          |         |
| No              | 2858 (54.12)   | 1249 (64.08)   |         | 1275 (65.42)  | 1249 (64.08)  |         |
| Yes             | 2423 (45.88)   | 700 (35.92)    |         | 674 (34.58)   | 700 (35.92)   |         |
| Radiation       |                |                | .006    |                | .700          |         |
| No/unknown      | 2326 (44.04)   | 929 (47.67)    |         | 917 (47.05)   | 929 (47.67)   |         |
| Yes             | 2955 (55.96)   | 1020 (52.33)   |         | 1032 (52.95)  | 1020 (52.33)  |         |
| Chemotherapy    |                |                | <.001   |                | .949          |         |
| No/unknown      | 2487 (47.09)   | 1036 (53.16)   |         | 1034 (53.05)  | 1036 (53.16)  |         |
| Yes             | 2794 (52.91)   | 915 (46.84)    |         | 915 (46.95)   | 913 (46.84)   |         |

Data were n (%), unless otherwise specified. PEC = primary esophageal cancer, SD = standard deviation, SEC = subsequent esophageal cancer, SEER = surveillance, epidemiology, and end results, SPC = subsequent primary cancer, TNM = tumor extent, nodal status and metastasis. * Grade I = well differentiated, Grade II = moderately differentiated, Grade III = poorly differentiated, Grade IV = undifferentiated.
Figure 5. The comparisons of survival outcomes between SEC and PEC patients (after matching). OS in the whole population (A); CSS in the whole population (B); OS in patients with esophageal adenocarcinoma (C); CSS in patients with esophageal adenocarcinoma (D); OS in patients with esophageal squamous cell carcinomas (E); CSS in patients with esophageal squamous cell carcinomas (F). PEC = primary esophageal cancer, SEC = subsequent esophageal cancer.

Table 5
Uni- and multivariate Cox regression model analysis of OS.

| Variables       | Univariate | Multivariate |
|-----------------|------------|--------------|
|                 | HR         | 95% CI       | P-value     | HR         | 95% CI       | P-value     |
| Age, yr         |            |              |             |            |              |             |
| <65 Reference   | 1.462      | 1.337–1.599  | <.001       | Reference  | 1.409      | 1.286–1.544  | <.001       |
| ≥65 Reference   | 1.409      | 1.286–1.544  | <.001       | Reference  | 1.352      | 1.234–1.478  | <.001       |
| Race            |            |              |             |            |              |             |
| White Reference | 1.450      | 1.299–1.619  | <.001       | Reference  | 1.205      | 1.072–1.356  | <.001       |
| Black           | 1.409      | 1.286–1.544  | <.001       | Reference  | 1.072      | 0.946–1.210  | <.001       |
| Other           | 0.913      | 0.757–1.103  | <.001       | Reference  | 1.006      | 0.899–1.112  | <.001       |
| Sex             |            |              |             |            |              |             |
| Male Reference  | 1.047      | 0.968–1.131  | <.001       | Reference  | 1.171      | 1.088–1.260  | <.001       |
| Female          | 0.968      | 0.899–1.038  | <.001       | Reference  | 0.946      | 0.852–1.053  | <.001       |
| Grade           |            |              |             |            |              |             |
| Grade I-II      | Reference  | 1.203–1.386  | <.001       | Reference  | 1.088      | 1.002–1.177  | <.001       |
| Grade III-IV    | Reference  | 1.006–1.112  | <.001       | Reference  | 1.088      | 1.002–1.177  | <.001       |
| TNM stage       |            |              |             |            |              |             |
| I Reference     | 1.047      | 0.970–1.130  | <.001       | Reference  | 0.965      | 0.893–1.042  | <.001       |
| II              | 1.914      | 1.692–2.176  | <.001       | Reference  | 1.532      | 1.352–1.736  | <.001       |
| III             | 1.375      | 1.278–1.479  | <.001       | Reference  | 1.078      | 0.975–1.192  | <.001       |
| Histology       |            |              |             |            |              |             |
| SCC Reference   | 1.375      | 1.278–1.479  | <.001       | Reference  | 1.078      | 0.975–1.192  | <.001       |
| SCC             | 1.047      | 0.970–1.130  | <.001       | Reference  | 0.965      | 0.893–1.042  | <.001       |

(continued)
Many previous studies have explored the prognostic factors for OS and CSS in cancer survivors. Traditionally, age at diagnosis, tumor grade, TNM stage, and the administration of surgery were widely recognized risk factors for survival in many cancer types. In our study, the existence of a prior cancer (PEC vs SEC) was identified to be an independent prognostic factor for CSS, but not for OS. Some studies demonstrated that a prior cancer could seriously affect the survival of cancer survivors, and those with prior malignancies should be excluded from clinical trials. However, our data supported that careful selection of candidates for clinical trials should be performed in EC patients with a prior cancer, rather than excluding all patients.

### Table 5

(continued).

| Variables | Univariate | Multivariate |
|-----------|------------|--------------|
| Other     | 1.413      | 1.191        |
| Location  | .001       | .647         |
| Upper     | Reference  | Reference    |
| Middle    | 0.957      | 1.026        |
| Lower     | 0.768      | 1.059        |
| Diagnosis | .441       | .357         |
| PEC       | Reference  | Reference    |
| SEC       | 1.028      | 0.916–1.150  |
| Surgery   | <.001      | <.001        |
| No        | Reference  | 0.363        |
| Yes       | 0.336      | 0.333–0.395  |

CI = confidence interval, CSS = cancer-specific survival, HR = hazard ratio, PEC = primary esophageal cancer, SD = standard deviation, SEC = subsequent esophageal cancer, TNM = tumor extent, nodal status and metastasis.

*Grade I = well differentiated, Grade II = moderately differentiated, Grade III = poorly differentiated, Grade IV = undifferentiated.

### Table 6

Uni- and multivariate Cox regression model analysis of CSS.

| Variables | Univariate | Multivariate |
|-----------|------------|--------------|
| Age, yr   |            |              |
| <65       | Reference  | Reference    |
| ≥65       | 1.434      | 1.360        |
| Race      |            | .026         |
| White     | Reference  | Reference    |
| Black     | 1.464      | 1.187        |
| Other     | 1.023      | 0.907        |
| Sex       | .121       | .732–1.125   |
| Male      | Reference  | Reference    |
| Female    | 1.075      | 1.088        |
| Grade     |            | .952–1.245   |
| Grade I-II| Reference  | Reference    |
| Grade III-IV| 1.357  | 1.208        |
| TNM stage |            | .108–1.317   |
| I         | Reference  | Reference    |
| II        | 1.104      | 1.091–1.097  |
| III       | 2.132      | 1.678        |
| Histology |            | 1.455–1.935  |
| AC        | Reference  | Reference    |
| SCC       | 1.416      | 1.117        |
| Other     | 1.479      | 0.991–1.258  |
| Location  |            | .014–1.443   |
| Upper     | Reference  | Reference    |
| Middle    | 1.010      | 1.088        |
| Lower     | 0.783      | 0.978–1.306  |
| Diagnosis |            | .952–1.245   |
| PEC       | Reference  | Reference    |
| SEC       | 0.762      | 0.680–0.804  |
| Surgery   |            | .680–0.804   |
| No        | Reference  | Reference    |
| Yes       | 0.282      | 0.305        |

CI = confidence interval, CSS = cancer-specific survival, HR = hazard ratio, PEC = primary esophageal cancer, SD = standard deviation, SEC = subsequent esophageal cancer.

*Grade I = well differentiated, Grade II = moderately differentiated, Grade III = poorly differentiated, Grade IV = undifferentiated.
However, there were some limitations that should not be ignored. First, numerous data were lacking or missing in the SEER registry. Second, the nature of retrospective research led to the inevitable selection bias. Moreover, treatment strategies of prior cancers may have something to do with the occurrence and survival of SPM.\[25,26\] Therefore, further prospective and well-designed studies are needed to validate our findings.

5. Conclusions

In EC patients with a prior cancer, EC is the most important COD regardless of the clinical characteristics of the prior cancer and EC. Surgery for these patients decreased the risk of ECSM. These finding suggested that EC related treatment should be actively adopted in patients with prior cancers, as they were more likely to die from EC than the prior cancer. Lastly, age at diagnosis, race, tumor grade, TNM stage, histology, and the administration of surgery were found to be prognostic factors for OS and CSS in EC patients.

Author contributions

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References

[1] Bray F, Ferlay J, Soerjomataram I, et al. Global cancer statistics 2018: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin 2018;68:394–424.
[2] Siegel RL, Miller KD, Goding Sauer A, et al. Colorectal cancer statistics, 2020. CA Cancer J Clin 2020;70:145–64.
[3] Fatehi Hassanabad A, Chechade R, Breadner D, et al. Esophageal carcinoma: towards targeted therapies. Cell Oncol (Dordr) 2020;43:195–209.
[4] Schizas D, Charalampakis N, Kole C, et al. Immunotherapy for esophageal cancer: a 2019 update. Immunotherapy 2020;12:203–18.
[5] Fitzmaurice C, Allen C, Barber RM, et al. Global, regional, and national cancer incidence, mortality, years of life lost, years lived with disability, and disability-adjusted life-years for 32 cancer groups, 1990 to 2015: a systematic analysis for the global burden of disease study. JAMA Oncol 2017;3:524–48.
[6] Hyder J, Boggs DH, Hanna A, et al. Changes in neutrophil-to-lymphocyte and platelet-to-lymphocyte ratios during chemoradiation predict for survival and pathologic complete response in tridimensional esophageal cancer patients. J Gastrointest Oncol 2016;7:189–95.
[7] Wood ME, Vogel V, Ng A, et al. Second malignant neoplasms: assessment and strategies for risk reduction. J Clin Oncol 2012;30:1734–45.
[8] Youlden DR, Baade PD. The relative risk of second primary cancers in Queensland, Australia: a retrospective cohort study. BMC Cancer 2011;11:83.
[9] Liao CT, Fan KH, Kang CJ, et al. Clinical outcomes of patients with resected oral cavity cancer and simultaneous second primary malignancies. PloS One 2015;10:e0136918.
[10] Vassilev ZP, Gabarró MS, Kaye JA, et al. Incidence of second primary malignancies in metastastic castration-resistant prostate cancer: results from observational studies in three countries. Eur Urol 2020;16:3889–901.
[11] Schonfeld SJ, Morton LM, Berrington de González A, et al. Risk of second primary papillary thyroid cancer among adult cancer survivors in the United States, 2000-2015. Cancer Epidemiol 2020;64:101664.
[12] Diniz KT, Mahal BA, Zehir DR, et al. Risk of prostate cancer mortality in men with a history of prior cancer. BJU Int 2016;117:E20–8.
[13] Saad AM, Al-Husseini MJ, Elgebaly A, et al. Impact of prior malignancy on outcomes of stage IV esophageal carcinoma: SEER based study. Expert Rev Gastroenterol Hepatol 2018;12:417–23.
[14] Chen Z, Li M, Ma K, et al. Clinicopathological features and prognosis of patients with esophageal cancer as the second primary cancer: a large population-based analysis using the SEER program [2000-2015]. Transl Cancer Res 2020;9:1113–24.
[15] Smyth EC, Tarazona N, Peckitt C, et al. Exclusion of gastrointestinal cancer patients with prior cancer from clinical trials: is this justified? Clin Colorectal Cancer 2016;15:e53–9.
[16] Murphy CC, Gerber DE, Pruitt SL. Prevalence of prior cancer among persons newly diagnosed with cancer: an initial report from the surveillance, epidemiology, and end results program. JAMA Oncol 2018;4:832–6.
[17] Van Spall HG, Toren A, Kiss A, et al. Eligibility criteria of randomized controlled trials published in high-impact general medical journals: a systematic sampling review. JAMA 2007;297:1231–40.
[18] Gerber DE, Luccetti AL, Xuan L, et al. Impact of prior cancer on eligibility for lung cancer clinical trials. J Natl Cancer Inst 2014;106:41.
[19] Kotwall CA, Mahoney LJ, Myers RE, et al. Reasons for non-entry in randomized clinical trials for breast cancer: a single institutional study. J Surg Oncol 1992;50:125–9.
[20] Zhu K, Lin R, Zhang Z, et al. Impact of prior cancer history on the survival of patients with larynx cancer. BMC Cancer 2020;20:1137.
[21] Luccetti AL, Pruitt SL, Xuan L, et al. Prior cancer does not adversely affect survival in locally advanced lung cancer: a national SEER-medicare analysis. Lung Cancer 2016;98:106–13.
[22] Wang YQ, Lv JW, Tang LL, et al. Effect of prior cancer on trial eligibility and treatment outcomes in nasopharyngeal carcinoma: implications for clinical trial accrual. Oral Oncol 2019;90:23–9.
[23] JF, Yang CQ, Li XL, et al. Risk of breast cancer-related death in women with a prior cancer. Aging 2020;12:5894–906.
[24] Al-Husseini MJ, Saad AM, El-Sewy KM, et al. Prior malignancy impact on survival outcomes of glioblastoma multiforme; population-based study. Int J Neurosci 2019;129:447–54.
[25] Davis EJ, Beebe-Dimmer JL, Yee CL, et al. Risk of second primary tumors in men diagnosed with prostate cancer: a population-based cohort study. Cancer 2014;120:2735–41.
[26] Moschini M, Zaffuto F, Karakiewicz PI, et al. External beam radiotherapy increases the risk of bladder cancer when compared with radical prostatectomy in patients affected by prostate cancer: a population-based analysis. Eur Urol 2019;75:319–28.