Retrospective Study

Development and validation of a prognostic nomogram model for Chinese patients with primary small cell carcinoma of the esophagus

Dong-Yun Zhang, Gai-Rong Huang, Jian-Wei Ku, Xue-Ke Zhao, Xin Song, Rui-Hua Xu, Wen-Li Han, Fu-You Zhou, Ran Wang, Meng-Xia Wei, Li-Dong Wang

ORCID number: Dong-Yun Zhang 0000-0001-5885-9238; Gai-Rong Huang 0000-0002-4223-0539; Jian-Wei Ku 0000-0002-0015-4662; Xue-Ke Zhao 0000-0002-9036-6342; Xin Song 0000-0002-7680-4908; Rui-Hua Xu 0000-0002-6914-2168; Wen-Li Han 0000-0002-3516-2195; Fu-You Zhou 0000-0002-6548-4090; Ran Wang 0000-0002-2588-5768; Meng-Xia Wei 0000-0001-6602-8678; Li-Dong Wang 0000-0001-5103-8226.

Author contributions: Wang LD, Zhang DY, and Huang GR designed and wrote the paper; Ku JW, Xu RH, Han WL, Wang R, Wei MX, and Zhou FY performed the data collection and interpretation and follow-up; Zhang DY, Zhao XK, and Song X contributed to the data analysis; all authors approved the final manuscript.

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Abstract

BACKGROUND
Primary small cell carcinoma of the esophagus (PSCE) is a highly invasive malignant tumor with a poor prognosis compared with esophageal squamous cell carcinoma. Due to the limited samples size and the short follow-up time, there are few reports on elucidating the prognosis of PSCE, especially on the establishment and validation of a survival prediction nomogram model covering general information, pathological factors and specific biological proteins of PSCE patients.

AIM
To establish an effective nomogram to predict the overall survival (OS) probability for PSCE patients in China.

METHODS
The nomogram was based on a retrospective study of 256 PSCE patients.
Primary small cell carcinoma of the esophagus (PSCE) is a rare, highly invasive tumor associated with early metastasis, and its incidence accounts for only 0.05%-4% of all esophageal cancers[1]. Similar to small cell lung cancer, the diagnosis of PSCE mainly depends on immunohistochemical staining for several neuroendocrine markers, including synaptofysin (Syn), neuronal cell adhesion molecule 56 (CD56), and chromogranin A (CgA)[2]. However, their impact on the overall survival (OS) of PSCE patients has not been thoroughly studied. Related studies have reported some prognostic factors for PSCE, but the results are controversial. The main prognostic indicator is still the stage of the disease[3].

The tumor node metastasis (TNM) staging system by the American Joint Committee on Cancer (AJCC) is currently widely used in the clinical treatment and prognosis of cancer patients[4]. PSCE is generally staged based on the AJCC TNM classification of esophageal squamous cell carcinoma. A large amount of evidence shows that the staging system cannot well evaluate the clinical outcome of PSCE. Compared with the traditional staging system, our nomogram model can accurately evaluate the overall survival probability at 1, 3, and 5 years for PSCE patients in China.

INTRODUCTION

Primary small cell carcinoma of the esophagus (PSCE) is a rare, highly invasive malignant tumor associated with early metastasis, and its incidence accounts for only 0.05%-4% of all esophageal cancers[1]. Similar to small cell lung cancer, the diagnosis of PSCE mainly depends on immunohistochemical staining for several neuroendocrine markers, including synaptofysin (Syn), neuronal cell adhesion molecule 56 (CD56), and chromogranin A (CgA)[2]. However, their impact on the overall survival (OS) of PSCE patients has not been thoroughly studied. Related studies have reported some prognostic factors for PSCE, but the results are controversial. The main prognostic indicator is still the stage of the disease[3].

The tumor node metastasis (TNM) staging system by the American Joint Committee on Cancer (AJCC) is currently widely used in the clinical treatment and prognosis of cancer patients[4]. PSCE is generally staged based on the AJCC TNM classification of esophageal squamous cell carcinoma (ESCC)[5]. The 7th AJCC TNM system includes tumor invasion depth (T), lymph node invasion (N), detectable metastasis (M), and tumor location and grade. A previous study which focused on the OS effect of CgA on
According to the random numbers generated by the computer, 70% of the eligible 125 PSCE patients, demonstrated that the TNM staging system for ESCC may not be a good predictor of prognosis for PSCE[6]. Other clinicopathological factors that are not included in the 7th TNM staging may also affect OS[7]. Furthermore, a growing body of evidence has suggested that PSCE has a poorer prognosis compared to ESCC. A more accurate prognostic prediction model is needed for PSCE patients[8,9].

Nomograms are currently widely used in graphical representations of complex mathematical formulas[10]. A nomogram model has obvious advantages in quantifying risk compared with TNM staging, which generally takes all known clinicopathological variables into account and allows personalized prognosis prediction[11,12]. The first nomogram was developed in 2021 by Qie et al[12], based on the patients in the United States in the public surveillance, epidemiology, and end results (SEER) database to predict patient OS probability. Of note, this study did not involve the Chinese population and relevant neuroendocrine markers were not included in the model[13]. The present study, thus, aimed to build a prognostic predictive nomogram model including clinicopathological factors and neuroendocrine biomarkers for Chinese PSCE patients. It was also determined whether the nomogram model can predict OS more accurately than the 7th TNM staging system.

MATERIALS AND METHODS

Patients and study design
A retrospective observational study was conducted on 343 eligible patients, who were enrolled from our esophageal and gastric cardiac carcinoma database containing 500000 cases[14]. A total of 256 eligible patients were finally enrolled using the following inclusion criteria: Pathologically diagnosed with primary PSCE, no preoperative radiotherapy and/or chemotherapy, survival time more than 1 mo, and detailed clinical baseline records. Patients with other malignant disease or a history of anticancer treatment were excluded.

The demographic and clinicopathological parameters of the patients included gender, age, smoking history, alcohol history, histology type, tumor location, T stage, N stage, M, and treatment history. Immunostained biomarkers including CgA, Syn, and CD56 were also identified from clinical medical records. All eligible patients were followed every year by telephone or home interview. The latest follow-up was conducted at the end of June 2019. The OS was measured from the time of surgery to the most recent follow-up time or death. Survival status was determined on the last day of follow-up. The median follow-up time was 24 mo (range, 2 to 154.8 mo). The study protocol was approved by the Medical Ethics Committee of the First Affiliated Hospital of Zhengzhou University.

Statistical analysis
Statistical analyses were conducted using R version 4.0.3 (http://www.r-project.org/) and Stata version 15.0 (http://www.stata.com). The Kaplan-Meier (K-M) method was used to draw survival curves, which were compared by the log-rank test. Clinical related parameters (based on clinical experience and literature reports) and baseline variables with statistical correlation in univariate analysis (P < 0.2) were included in the multivariate Cox proportional hazard regression analysis[15,16]. A predictive nomogram of 1-year, 3-year, and 5-year OS based on the Akaike information criterion (AIC) was developed using the Cox regression model and the 'RMS' package in R[17]. The concordance index (C-index)[18], which was calculated by a bootstrap approach with 1000 resamples[19], was used to determine discrimination. By mapping the nomogram predicted probabilities of OS with the actual OS observed at 1, 3, and 5 years, the calibration curve was graphically evaluated, and the 45° line indicated the best predicted value[20]. Decision curve analysis (DCA) was used to evaluate the clinical application of the prediction model by quantifying net benefits[21]. In addition, we divided the model into three groups according to the cutoff points of the total prognostic scores (TPS), which were determined by the minimum P value obtained using the X-tile program[22].

RESULTS

Patient characteristics
According to the random numbers generated by the computer, 70% of the eligible
patients were randomly assigned to a training cohort (n = 179) and the remaining 30% to a validation cohort (n = 77). The clinical characteristics of these patients are summarized in Table 1. The male to female ratio in the training and validation cohorts was 1.84:1 (116/63) and 1.96:1 (51/26), respectively. The median age in each cohort was 62.19 years (range, 37-88 years) and 61.83 years (range, 46-77 years), respectively. The median survival time in the primary cohort was 23.59 mo (range, 1-105 mo) and the 1-, 3-, and 5-year OS rates were 67.6%, 35.1%, and 24.3%, respectively. In the validation cohort, the median survival time was 24.63 mo (range, 1-108 mo), and the 1-, 3-, and 5-year OS rates were 70.4%, 46.8%, 31.6%, respectively.

**Nomogram development**

Of the initial 13 variables, tumor location, alcohol consumption, and smoking history were excluded from the Cox proportional hazards regression analysis due to their weak and non-significant OS correlation in the univariate analysis ($P > 0.2$) (Figure 1). Based on the results of univariate analysis and clinical parameters, histology type, gender, age, T stage, N stage, M, operation, Syn, CgA, and CD56 were included in the Cox proportional hazards regression model. Finally, the most suitable nomogram model was determined using the backward step selection process with the smallest AIC, which included histology type, age, T stage, N stage, M, CgA, and CD56. The model suggested that N stage had the greatest influence on patient prognosis, followed by histological type and age. T stage, M, CD56, and CgA had moderate effects on OS. Each subtype of the variables was assigned a score on the scoring table. By adding the total score and positioning it in relation to the total subscale, a straight line was drawn to determine the estimated probability of survival at each time point. Figure 2 shows the nomogram model for predicting 1-, 3-, and 5-year OS in the training cohort, indicating that the higher the score, the worse the prognosis.

**Calibration and validation of the nomogram**

The calibration plots showed that the nomogram prediction was in good agreement with the actual observation of 1-, 3-, and 5-year OS in both cohorts (Figure 3). The accuracy of prognosis prediction was also compared between the nomogram model and the 7th TNM staging using the C-index (Table 2). In the training cohort, the C-index for OS prediction was 0.659 (95% CI: 0.607-0.712), which was significantly higher than that of the 7th TNM staging (0.591, 95% CI: 0.517-0.666, $P = 0.033$). This superior tendency in OS prediction was also verified using external validation in the validation cohort, with the C-index of the former (0.700, 95% CI: 0.622-0.778) also being higher than that of the latter (0.605, 95% CI: 0.490-0.721, $P = 0.041$). Figure 4 shows the results of DCA at 18 mo for PSCE, which indicated that our nomogram model had a higher overall net benefit than the 7th TNM staging within a wide range of threshold probabilities.

**Performance of the nomogram in stratifying patient risk**

As predicted by the nomogram model, the X-tile program was used to categorize the primary cohort into three groups: Low risk prognosis (TPS ≤ 13.2, 58 patients), medium risk prognosis (13.2 < TPS ≤ 22.8, 98 patients), and high risk prognosis (TPS > 22.8, 23 patients). The survival rate was highest in the low risk group (83.41%, 57.53%, and 33.40% for 1-, 3-, and 5-year survival rates), followed by the medium risk group (66.41, 31.71, and 21.14%) and the high risk group (34.78%, 4.35%, and 0%; Table 3). We then used cutoff values to plot K-M curves for both cohorts (Figure 5), which were also significantly correlated with OS ($P < 0.01$). Figure 6 shows the distribution of 5-year survival predicted by the nomogram for each stage of the 7th TNM staging in the primary and validation cohorts. And the results revealed that the higher the stage, the lower the survival rate.

**DISCUSSION**

A nomogram can be used to establish a statistical prognostic model to estimate the prognosis of cancer patients[10]. No studies have determined a prognostic nomogram for PSCE patients due to the rarity of the disease. In this study, we successfully constructed an effective nomogram model to predict PSCE prognosis. Combined with clinical correlations, we identified that age, histology type, T stage, N stage, M, CD56, and CgA were prognostic predictors of PSCE. However, tumor location and grade in the 7th TNM staging were not independent prognostic factors in this study. A previous study also failed to find a significant correlation of tumor location and grade with OS...
| Variable               | Primary cohort | Validation cohort |
|------------------------|----------------|-------------------|
|                        | n   | %   | n   | %   |
| Gender                 |     |     |     |     |
| Female                 | 63  | 35.2| 26  | 33.8|
| Male                   | 116 | 64.8| 51  | 66.2|
| Age, yr                |     |     |     |     |
| ≤ 60                   | 74  | 41.3| 32  | 41.6|
| > 60                   | 105 | 58.7| 45  | 58.4|
| Smoking history        |     |     |     |     |
| No                     | 122 | 68.2| 59  | 76.6|
| Yes                    | 57  | 31.8| 18  | 23.4|
| Drinking history       |     |     |     |     |
| No                     | 134 | 74.9| 64  | 83.1|
| Yes                    | 45  | 25.1| 13  | 16.9|
| Tumor location         |     |     |     |     |
| Upper                  | 23  | 12.9| 8   | 10.4|
| Middle                 | 111 | 62.0| 46  | 59.7|
| Lower                  | 45  | 25.1| 23  | 29.9|
| Histology              |     |     |     |     |
| Pure PSCE              | 106 | 59.2| 57  | 74.1|
| Mixed PSCE             | 73  | 40.8| 20  | 25.9|
| T stage                |     |     |     |     |
| T1a/T1b                | 34  | 19.0| 19  | 24.7|
| T2                     | 63  | 35.2| 31  | 40.2|
| T3/T4                  | 82  | 45.8| 27  | 35.1|
| N stage                |     |     |     |     |
| N0                     | 87  | 48.6| 35  | 45.4|
| N1                     | 64  | 35.8| 30  | 39.0|
| N2/N3                  | 28  | 15.6| 12  | 15.6|
| Metastasis             |     |     |     |     |
| No                     | 136 | 75.9| 60  | 77.9|
| Yes                    | 43  | 24.1| 17  | 22.1|
| Treatment methods      |     |     |     |     |
| Surgery                | 155 | 86.6| 68  | 88.3|
| Others                 | 24  | 13.4| 9   | 11.7|
| CD56                   |     |     |     |     |
| Negative               | 45  | 25.1| 15  | 19.5|
| Positive               | 124 | 74.9| 62  | 80.5|
| Syn                    |     |     |     |     |
| Negative               | 40  | 22.4| 17  | 22.1|
| Positive               | 139 | 77.6| 60  | 77.9|
| CgA                    |     |     |     |     |
Zhang DY et al. Prognostic nomogram model for PSCE

| Variable       | Primary cohort C-index (95%CI) | P value | Validation cohort C-index (95%CI) | P value |
|----------------|-------------------------------|---------|----------------------------------|---------|
| Nomogram model | 0.659 (0.607-0.712)           | 0.033   | 0.700 (0.622-0.778)              | 0.041   |
| 7th TNM staging| 0.591 (0.517-0.666)           |         | 0.605 (0.490-0.721)             |         |

C-index: Concordance index; CI: Confidence interval; TNM: Tumor node metastasis.

Table 3 Cox regression analysis for groups based on the model in the primary cohort

| Groups               | OS mean (1-yr (%)) | 3-yr (%) | 5-yr (%) | Sig | HR (95%CI) |
|----------------------|--------------------|----------|----------|-----|------------|
| Low risk             | 71.7               | 83.4     | 57.5     | 33.4| -          |
| Medium risk          | 19.8               | 66.4     | 31.7     | 21.4| 0.004      | 1.93 (1.23-3.05) |
| High risk            | 10.03              | 34.8     | 4.4      | -   | 0.000      | 5.47 (3.08-9.73) |

OS: Overall survival; CI: Confidence interval; HR: Hazard ratio.

PSCE: Primary small cell carcinoma of the esophagus; T: Tumor invasion depth; N: Lymph node invasion; CD56: Neuronal cell adhesion molecule 56; Syn: Synaptophysin; CgA: Chromogranin A.

In Chinese PSCE patients[6]. In addition, the analysis of four non-TNM risk factors, including age, histology type, CgA, and CD56, showed significant increases in risk scores for patients over 60 years old (6.0 points), pure PSCE (6.0 points), positive CgA (4.0 points), and negative CD56 (5.0 points). The nomogram model is beneficial in the development of a personalized scoring system for patients[23].

Histopathologically, some PSCEs are mainly composed of neuroendocrine tumor cells and may have other cancerous components, such as adenocarcinoma or squamous cell carcinoma[24]. PSCE has been reported to be a highly metastatic disease with a poor prognosis. This estimate corresponds well with our study finding that 24.1% of patients presented with stage IV and 40.8% of patients had coexisting squamous cell carcinoma. Mixed PSCE may be derived from multipotential stem cells of the esophageal mucosa[25]. With regard to survival, mixed PSCE patients had a superior outcome with a median OS of 23.21 mo and a 5-year OS of 31.2%. Pure PSCE patients were associated with a median OS of 16.67 mo and a 5-year OS of 20.3%. CD56 is a type of neurocellular adhesion molecule associated with the prognosis of multiple myeloma[26]. CgA is an acidic glycoprotein, which belongs to a class of regulated secretory proteins[27] and is a prognosis factor for several types of small cell carcinomas[28,29].

In addition, several previous studies have shown that surgery can enhance the prognosis of some PSCE patients[30,31]. In our study, compared with patients who received other treatments, patients in the surgery group showed an increasing trend in clinical outcome, with a median OS of 21.11 vs 12.39 mo and a 5-year OS of 28.3% vs 15.8%, although these differences were not statistically significant. A meta-analysis by Raja et al[32] showed that surgery or radiotherapy combined with chemotherapy can significantly improve patient prognosis. However, since only 14 patients received surgery plus radiotherapy and 10 underwent surgery and chemotherapy in our training cohort, we were unable to conduct further survival analysis to determine any differences. Therefore, treatment is not considered to be a potential prognostic factor in our nomogram model.

As far as we know, this is the first nomogram model of PSCE based on seven important prognostic factors. The discrimination and calibration were evaluated by internal and external validation[33]. In our study, the C-index of the nomogram was
Zhang DY et al. Prognostic nomogram model for PSCE

Figure 1 Survival analysis in the primary cohort. A: Univariate analysis in the primary cohort; B: Multivariate analysis in the primary cohort. T: Tumor invasion depth; N: Number of positive lymph nodes; CD56: Neuronal cell adhesion molecule 56; CgA: Chromogranin A; Syn: Synaptophysin.

| Points | Pure PSCE | Mixed PSCE |
|--------|-----------|------------|
| Histology | N0 | N1N1 |
| N stage | N2/N3 |
| T stage | T1 | T2 | T3/T4 |
| Age, years | ≤ 60 | > 60 |
| CD56 | Negative | Positive |
| Metastasis | No | Yes |
| CgA | Negative | Positive |

Figure 2 Nomogram model for predicting the 1-, 3-, and 5-year overall survival in primary small cell carcinoma of the esophagus patients. N: Number of positive lymph nodes; T: Tumor invasion depth; CD56: Neuronal cell adhesion molecule 56; CgA: Chromogranin A.

higher than that of the 7th TNM staging, indicating higher discrimination of the model. We then used the optimal cut-off analyses to successfully classify the patients in the training cohort into three risk groups. Subgroup stratification of patients with different risk levels is beneficial for clinicians to carry out individualized treatment. The calibration curves indicated that there was good consistency between the predicted and actual values for 1-, 3-, and 5-year OS. DCA also suggested that the prognostic prediction model had greater potential for clinical application than the 7th TNM staging.
Figure 3 Calibration curves for predicting overall survival at (A) 1 year, (B) 3 years, and (C) 5 years in the primary cohort and at (D) 1 year, (E) 3 years, and (F) 5 years in the validation cohort. OS: Overall survival.

Figure 4 Decision curve analysis for the 18-mo survival predictions in primary small cell carcinoma of the esophagus patients. TNM: Tumor node metastasis.

However, this study also had several limitations. First, due to the retrospective nature of our research, potential biases were inevitable. Second, due to the low incidence of PSCE, other relevant clinical parameters (e.g., incidence area, family history, body mass index, and reproductive history) were unavailable. These confounding factors might affect treatment choice or survival, which should be considered in future related studies.
Figure 5 Kaplan–Meier curves for all three groups based on the nomogram prediction. A: Kaplan–Meier curves for all three groups in the primary cohort. B: Kaplan–Meier curves for all three groups in the validation cohort.

Figure 6 Distribution of the nomogram predicted 5-year survival rate according to 7th edition tumor node metastasis stages. A: Distribution in the primary cohort; B: Distribution in the validation cohort. TNM: Tumor node metastasis.

CONCLUSION
We have developed, for the first time, a nomogram model for predicting OS in Chinese patients with PSCE. The novel nomogram classifies patients into different risk subgroups and showed superiority in predicting survival compared with the 7th TNM staging. Therefore, our nomogram may help clinicians make individualized prognostic predictions and better treatment recommendations for PSCE patients in the future.

ARTICLE HIGHLIGHTS
Research background
Primary small cell carcinoma of the esophagus (PSCE) is a rare tumor, accounting for 0.05% to 3.1% of all esophageal malignancies and approximately 2% of extrapulmonary small cell carcinomas. PSCE patients seem to have earlier metastasis and a worse prognosis than those with esophageal squamous cell carcinoma, which requires a more accurate prognostic prediction model.

Research motivation
Several previous studies have reported the prognostic factors for PSCE with contro-
versial results, partly due to their small sample size. To date, only one nomogram has been used to predict the overall survival probability for PSCE patients in the United States. In addition, the model did not include relevant neuroendocrine markers.

**Research objectives**
The present study aimed to build a prognostic predictive nomogram model including clinicopathological factors and neuroendocrine biomarkers for Chinese PSCE patients. It was also determined whether the nomogram model can predict overall survival (OS) more accurately than the 7th tumor-node-metastasis (TNM) staging system.

**Research methods**
The nomogram was based on a retrospective study of 256 PSCE patients, derived from our esophageal and gastric cardia carcinoma database including 500000 cases (1973-2015), established by State Key Laboratory of Esophageal Cancer Prevention and Treatment and Henan Key Laboratory for Esophageal Cancer Research of the First Affiliated Hospital of Zhengzhou University in Henan, China. The predictive accuracy and discriminative ability of the nomogram were determined by the concordance index (C-index), calibration plot, and decision curve analysis (DCA), and the results were also compared with the 7th TNM staging.

**Research results**
The final nomogram model included histology type, age, tumor invasion depth, lymph node invasion, distant metastases, chromogranin A, and neuronal cell adhesion molecule 56. The C-index of the model had a prognostic superiority over the 7th edition TNM staging system in both the primary cohort [0.659 (95%CI: 0.607-0.712) vs 0.591 (95%CI: 0.517-0.666), P = 0.033] and the validation cohort [0.700 (95%CI: 0.622-0.778) vs 0.605 (95%CI: 0.490-0.721), P = 0.041]. Good calibration curves were observed for the prediction probabilities of 1-, 3- and 5-year OS in the primary and validation cohorts. DCA analysis showed that our nomogram model had a higher overall net benefit than the 7th TNM staging.

**Research conclusions**
We have developed and validated a nomogram for predicting 1-year, 3-year, and 5-year OS in Chinese PSCE patients. The new nomogram classifies patients into different risk subgroups and shows a superiority of survival prediction over the 7th TNM staging.

**Research perspectives**
The nomogram model can be used to predict the survival probability of PSCE patients, which might help clinicians to make individualized prognosis predictions and give better treatment recommendations for PSCE patients in China.

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