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nomogram for predicting

# **BMJ Open** Competing-risks nomogram for predicting cancer-specific death in upper tract urothelial carcinoma: a populationbased analysis

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### ABSTRACT

Objective This study aimed to use a competing-risks model to establish a nomogram to accurately analyse the prognostic factors for upper tract urothelial carcinoma (UTUC) cancer-specific death (CSD).

Design Retrospective observational cohort study. Setting The programme has yielded a database of all patients with cancer in 18 defined geographical regions of the USA.

Participants We selected patients with UTUC from the latest edition of the Surveillance, Epidemiology, and End Results database from 1975 to 2016. After excluding patients with unknown histological grade, tumour size and lymph node status. 2576 patients were finally selected.

Primary and secondary outcome measures We used the Fine-Gray proportional subdistribution hazards model for multivariate analysis and compared the results with cause-specific hazards model. We finally constructed a nomogram for 3-year, 5-year and 8-year CSD rates and tested these rates in a validation cohort.

**Results** The proportional subdistribution hazards model showed that sex, tumour size, distant metastasis, surgery status, number of lymph nodes positive (LNP) and lymph nodes ratio (LNR) were independent prognostic factors for CSD. All significant factors associated with CSD were included in the nomogram. The 3-year, 5-year and 8-year concordance indexes were 0.719, 0.702 and 0.692 in the training cohort and 0.701, 0.675 and 0.668 in the validation cohort, respectively.

Conclusions The competing-risks model showed that sex, tumour size, distant metastasis, surgery status, LNP and LNR were associated with CSD. The nomogram predicts the probability of CSD in patients with UTUC at 3, 5 and 8 years, which may help clinicians in predicting survival probabilities in individual patients.

### INTRODUCTION

Urothelial carcinomas are the fourth most common type of tumour,<sup>1</sup> which is located in the upper or lower urinary tract. Upper tract urothelial carcinoma (UTUC), including the renal pelvis and ureter carcinoma, currently accounts for 5% of urothelial malignancies.<sup>2</sup> The annual incidence of UTUC is typically

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  Strengths and limitations of this study
  The study established the first competing-risks no-mogram for predicting the 3-year, 5-year and 8-year specific mortality probability for upper tract urotheli-al carcinoma (UTUC) based on a large retrospective sample, which can improve the ability of clinicians in predicting survival probabilities in individual patients.
  The established model is not comprehensive enough because the Surveillance, Epidemiology, and End Results database does not include all prognostic factors for UTUC.
  The data available on treatment status are not suffi-ciently detailed to distinguish the impact of various treatment plans.
  The model requires prospective studies to confirm its reliability.

Western countries.<sup>3</sup> However, the increasing  $\geq$ tra morbidity and mortality associated with UTUC<sup>4 5</sup> make research more important.

ing, A previous study showed that UTUC has unique prognostic factors which are different from bladder cancer and other urinary tract cancers.<sup>6</sup> Most studies analysing the prognostic factors for UTUC have adopted the Kaplan-Meier (K-M) method or Cox regression methods.<sup>7–9</sup> These methods only consider a single endpoint while determining survival parameters. However, in & clinical research, in addition to events of g interest, there are often competing events. Competing events for cancer deaths refer to death from other causes unrelated to primary cancer, such as other diseases, car collisions and suicide. In traditional survival analysis methods, these events were considered censored, making the cumulative incidence of cancer deaths overestimated. Applying standard survival analysis to competing-risks

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**Correspondence to** Dr Jun Lyu; lyujun2020@jnu.edu.cn data leads to false and biased results.<sup>10</sup> Although the use of all-cause death as the study endpoint does not cause a competing-risk bias, such an analysis cannot reflect the influence of factors on the specific endpoint of cancer death. Therefore, the cumulative incidence function (CIF) of UTUC cancer-specific death (CSD) needs to be calculated and the prognostic factors for UTUC analysed using the Fine-Gray proportional subdistribution hazards model.<sup>11</sup>

A nomogram is based on a prognostic model and graphically represents the predictive abilities of different prognostic factors as the length of line segments. This format makes it easy for clinicians to make rapid and comprehensive decisions and predict the probability of CSD, and has great clinical significance. Some studies have constructed competing-risks nomograms for cancers such as sarcoma and prostate cancer<sup>1213</sup>; however, there is a lack of studies related to UTUC.

The purpose of our research was to identify the prognostic factors for UTUC based on the competing-risks model and use them to construct a nomogram to predict the survival rates of patients at 3, 5 and 8 years.

### **METHODS**

### **Database and patients**

The Surveillance, Epidemiology, and End Results (SEER) program has yielded a database of all patients with cancer in 18 defined geographical regions of the USA collected by the National Cancer Institute. It is the largest cancer registry in the USA and includes information on approximately 28% of the US population. The SEER research data are publicly available; therefore, no informed consent or institutional review board approval is required when analysing the data. We additionally requested chemotherapy data for inclusion in our research and obtained a licence for using the SEER software.14 15

We selected patients with UTUC from the latest edition of the SEER database from 1975 to 2016. The primary sites were extracted using the SEER codes 'C65.9-Renal pelvis' and 'C66.9-Ureter'. Patients between 2004 and 2015 were included in the study. We included all of the histological subtypes of UTUC, according to the Third Revision of the International Classification of Diseases for Oncology. The following demographic indicators were selected: age at diagnosis, sex, race and marital status. Primary site, histological grade, tumour size, laterality, distant metastasis, surgery status, radiotherapy status, chemotherapy status, number of lymph nodes examined (LNE), lymph nodes positive (LNP) and lymph nodes ratio (LNR; calculated as the number of LNP divided by LNE) were also included as pathological characteristics. Tumour size was divided into three groups: <2, 2-4 and  $\geq$ 4 cm.<sup>1 16</sup> The study outcomes included CSD and death due to other causes (DOC). Survival time was reported in the available data in months.



Figure 1 Data selection flow chart. ICD-O-3, Third Revision of the International Classification of Diseases for Oncology: SEER, Surveillance, Epidemiology, and End Results; UTUC, upper tract urothelial carcinoma; LNP, lymph nodes positive; LNE, lymph nodes examined.

### **Exclusion criteria**

Protected by copyright, incl Our preliminary selection of the above methods initially identified 13581 patients. Then, to ensure the study's accuracy, the following were excluded: unknown histological grade, unknown tumour size and unknown lymph ð node status. The specific data selection process is shown r uses in figure 1. We finally chose 2576 patients for inclusion in the follow-up investigations.

### Statistical methods

to We randomly divided the 2576 eligible patients into two groups using R software (V.3.5.3, The R Foundation for te Statistical Computing, Vienna, Austria; http://www.rproject.org): 70% (n=1803) in the training cohort and 30% (n=773) in the validation cohort. We first described the basic composition of each factor in the two patient  $\mathbf{\vec{a}}$ cohorts using the R software. Age, LNE, LNP and LNR **E** were expressed as median and IQR values, while categorical variables were presented as percentages. We eval-≥ uated differences in patient characteristics between the two groups using the Student's t-test and  $\chi^2$  test.

We used the CIF to describe the probability of each event and also plotted the corresponding CIF curves. Moreover, Gray's test was used for univariate analysis to estimate the difference in CIF between the groups. Significant variables (p<0.05) were included in the multivariate regression model. The Fine-Gray proportional subdistribution hazards model was used for multivariate analysis and compared with the results of the cause-specific hazards model. Applying the standard Cox regression **B** method ignores the presence of competing risks and **g** hence overestimates the actual incidence of beneficial **3** events, leading to inappropriate risk stratification.<sup>17</sup> Several studies have confirmed that different approaches can be used in competing-risks settings for multivariate survival analysis. However, proportional subdistribution hazards model is the best method to predict the survival probability.<sup>18-20</sup>

Finally, the results of Fine-Gray proportional subdistribution hazards model were used to construct a nomogram of the 3-year, 5-year and 8-year CSD rates. We used

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the concordance index (C-index) and calibration plots to evaluate the differentiation ability and consistency of the established model in the training and validation cohorts.

All statistical tests were conducted using R software (V.3.5.3). Probability values of p<0.05 were considered statistically significant and all tests were two-sided. The SEER database can be accessed free of charge. This study was exempted from obtaining informed consent.

### Patient and public involvement

This study was conducted without patient involvement. Patients were not invited to comment on the study design and were not consulted to develop patient-relevant outcomes or interpret the results. Moreover, patients were not allowed to contribute to the writing or editing of this document for readability or accuracy.

### RESULTS

### **Patient characteristics**

The composition of each variable for the 2576 patients in the training and validation cohorts is presented in table 1. The median age was 71 years in the training and validation cohorts, respectively. The majority of patients were male (60.6% and 57.4%), white (86.2% and 82.5%) and married (86.8% and 87.6%). The main sites of UTUC were in the renal pelvis (63.9% and 62.7%, respectively, in the training and validation cohorts), with the rest in the ureter. The majority of patients were in the undifferentiated stage (58.1% and 55.6%), and most of the tumours in both cohorts were larger than 4cm. Most patients in both cohorts had received surgery, whereas a few patients had received radiotherapy or chemotherapy. Only about 9% of the patients had distant metastasis. The baseline characteristics of the training cohorts and validation cohorts were similar.

### **Univariate analysis**

We calculated the 3-year, 5-year and 8-year cumulative incidence rates of CSD and DOC. Year, laterality and marital status were not related to either outcome (p>0.05), while age, sex, histological grade, chemotherapy status and LNR were related to both outcomes (p<0.05). Race, primary site, tumour size, surgery status, radiotherapy status, distant metastasis, LNE and LNP were significantly related to CSD. The corresponding CIF curves are shown in figure 2. The cumulative incidences of CSD and DOC are compared in table 2.

### **Multivariate analysis**

Table 3 shows the comparison of the proportional subdistribution hazards model with the cancer-specific hazards model. The cancer-specific hazards model showed that sex, tumour size, distant metastasis, LNP and LNR were prognostic factors for UTUC (p<0.001). We constructed the Fine-Gray proportional subdistribution hazards model, which indicates that sex (HR=1.480 for female, 95% CI 1.241 to 1.764), tumour size (HR=1.556 for

2–4 cm, 95% CI 1.092 to 2.216; HR=2.205 for ≥4 cm, 95% CI 1.575 to 3.087), surgery status (HR=2.205 for no/ unknown surgery, 95% CI 1.292 to 3.761), distant metastasis (HR=2.414 for distant, 95% CI 1.842 to 3.163), LNP (HR=1.064, 95% CI 1.022 to 1.107) and LNR (HR=1.873, 95% CI 1.435 to 2.445) were prognostic factors affecting UTUC, as shown in table 3.

# Construction and verification of the nomogram

Figure 3 shows the nomogram constructed according to the results of the Fine-Gray proportional subdistribution hazards model for predicting CSD probabilities at 3, 5 and 8 years. LNP had the most significant impact on the probability of CSD, followed by distant metastasis, tumour J size, LNR, surgery and sex (figure 3).

We used both the training and validation cohorts to verify the nomogram after establishing it. The 3-year, 5-year and 8-year C-indexes were 0.719, 0.702 and 0.692 for the training cohort, respectively, and 0.701, 0.675 and 0.668 for the validation cohort. All of these values exceeded 0.6, indicating that the model had good discrimination ability. We then tested the prediction accuracy of g the model. As shown in figure 4, the 3-year, 5-year and uses related 8-year calibration plots for both cohorts were very close to the standard straight line, demonstrating that the model was well calibrated.

# DISCUSSION

The increasing incidence of UTUC<sup>21</sup> makes it necessary to further explore the prognostic factors for UTUC. The present study used a competing-risks model to accurately explore the prognostic factors for UTUC. The study used these factors to construct a nomogram that provides  $\Xi$ clinicians with direct guidance while making relevant decisions.

≥ The application of the study criteria resulted in the inclusion of 2576 patients from the SEER database, and 1542 of these patients died during follow-up. However, ğ only 750 of the deaths were related to UTUC. These results indicate that the number of DOC patients was almost the same as that for CSD. In this situation, if the traditional K-M or Cox survival analysis had been adopted, the DOC patients will be regarded as censored. This will lead to an overestimation of the cumulative incidence of CSD, which cannot truly reflect the prognosis.<sup>22 23</sup> We overcame Inol this shortcoming by using a competing-risks model which can adequately address the situation where the available  $\mathbf{g}$ data are related to multiple potential outcomes.<sup>24</sup> This **3** method was first proposed by Fine and Gray and applied in previous studies.<sup>17 25 26</sup> In the case of competing risks, there are usually two models. One is the cause-specific hazards model (CS), and the other is the proportional subdistribution hazards model (SD), also known as the Fine-Gray model. In the present study, two models were analysed and compared. CS is suitable for answering aetiological questions, and SD is suitable for establishing clinical prediction models and risk scores. Therefore, we

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Table 1      Basic characteristics of patients i	n this study		
Variables	Training cohort	Validation cohort	P value
Number of patients, n (%)	1803 (70)	773 (30)	
Age, median (IQR)	71.00 (64.00–78.00)	71.00 (63.00–78.00)	0.710
Sex, n (%)			0.150
Female	711 (39.4)	329 (42.6)	
Male	1092 (60.6)	444 (57.4)	
Race, n (%)			0.045
Black	80 (4.4)	50 (6.5)	
Other	169 (9.4)	83 (10.7)	
White	1554 (86.2)	640 (82.8)	
Marital status, n (%)			0.656
Married	1565 (86.8)	677 (87.6)	
Others	67 (3.7)	31 (4.0)	
Single	171 (9.5)	65 (8.4)	
Year, n (%)			0.813
2004–2006	346 (19.2)	159 (20.6)	
2007–2009	439 (24.3)	181 (23.4)	
2010–2012	479 (26.6)	198 (25.6)	
2013–2015	539 (29.9)	235 (30.4)	
Site, n (%)			0.609
Renal pelvis	1152 (63.9)	485 (62.7)	
Ureter	651 (36.1)	288 (37.3)	
Grade, n (%)			0.481
Grade I	47 (2.6)	16 (2.1)	
Grade II	149 (8.3)	69 (8.9)	
Grade III	559 (31.0)	258 (33.4)	
Grade IV	1048 (58.1)	430 (55.6)	
Size, n (%)			0.188
2–4	559 (31.0)	268 (34.7)	
<2	262 (14.5)	106 (13.7)	
≥4	982 (54.5)	399 (51.6)	
Laterality, n (%)			0.551
Left	995 (55.2)	416 (53.8)	
Right	808 (44.8)	357 (46.2)	
Surgery, n (%)			0.203
No/unknown	9 (0.5)	8 (1.0)	
Yes	1794 (99.5)	765 (99.0)	
Radiotherapy, n (%)			0.931
No/unknown	1676 (93.0)	720 (93.1)	
Yes	127 (7.0)	53 (6.9)	
Chemotherapy, n (%)			0.938
No/unknown	1243 (68.9)	531 (68.7)	
Yes	560 (31.1)	242 (31.3)	
Distant metastasis, n (%)			0.053
MO	1652 (91.6)	689 (89.1)	
M1	151 (8.4)	84 (10.9)	
LNE, median (IQR)	3.00 (1.00–7.00)	3.00 (1.00–7.00)	0.627
LNP, median (IQR)	0.00 (0.00–1.00)	0.00 (0.00–1.00)	0.542
LNR, median (IQR)	0.00 (0.00–0.50)	0.00 (0.00–0.33)	0.546

LNE, lymph nodes examined; LNP, lymph nodes positive; LNR, lymph nodes ratio.



Figure 2 CIF curves of CSD and DOC among patients with upper tract urothelial carcinoma. CIF, cumulative incidence function; CSD, cancer-specific death; DOC, death due to other causes.

used the CIF and the proportional SD model to explore the impact of various factors on the prognosis of CSD.

The results of the univariate analysis showed that age, sex, race, primary site, histological grade, tumour size, surgery status, radiotherapy status, chemotherapy status, distant metastasis, LNE, LNP and LNR were influencing factors for CSD, while age, sex, histological grade, chemotherapy status and LNR were influencing factors for DOC. The results of the CS model showed that age, sex, histological grade, tumour size, distant metastasis, LNP

tional SD model showed that sex, tumour size, surgery, distant metastasis, LNP and LNR are independent prognostic factors for CSD.

Age is generally considered to be a prognostic factor for most cancers and also for UTUC.<sup>27 28</sup> Our CS model showed that age was a predictor of CSD; however, it was not statistically significant in the SD model. This may be because the effect of age on DOC is higher than the CSD; namely, elderly patients are more likely to die of other

Table 2 Cum	ulative incidence of (	CSD and DOC amo	ng patients with upp	er tract ur	othelial carcinoma			
	CSD (%)				DOC (%)			
Variables	3 years (95% CI)	5 years (95% CI)	8 years (95% CI)	P value	3 years (95% CI)	5 years (95% CI)	8years (95% CI)	P value
Age				<0.001				<0.001
Sex				<0.001				<0.001
Male	22.903 (22.843 to 22.964)	27.131 (27.064 to 27.197)	29.457 (29.386 to 29.528)		25.697 (25.634 to 25.760)	33.645 (33.573 to 33.717)	40.755 (40.673 to 40.837)	
Female	33.320 (33.236 to 33.405)	38.157 (38.066 to 38.247)	40.339 (40.245 to 40.434)		18.710 (18.639 to 18.780)	24.144 (24.063 to 24.225)	29.031 (28.937 to 29.125)	
Race				0.008				0.057
White	25.881 (25.828 to 25.934)	29.921 (29.864 to 29.979)	31.856 (31.796 to 31.916)		23.502 (23.451 to 23.554)	30.839 (30.780 to 30.898)	37.596 (37.528 to 37.664)	
Black	35.688 (35.423 to 35.952)	44.479 (44.194 to 44.763)	48.470 (48.178 to 48.762)		22.782 (22.555 to 23.009)	27.889 (27.638 to 28.139)	30.253 (29.987 to 30.519)	
Other	33.577 (33.399 to 33.755)	39.945 (39.752 to 40.138)	43.895 (43.688 to 44.102)		17.728 (17.586 to 17.869)	22.120 (21.960 to 22.281)	25.822 (25.642 to 26.003)	
Marital status				0.578				0.888
Married	26.658 (26.605 to 26.711)	31.020 (30.962 to 31.077)	33.378 (33.317 to 33.440)		23.164 (23.113 to 23.215)	30.048 (29.990 to 30.106)	36.490 (36.423 to 36.557)	
Single	28.974 (28.808 to 29.140)	35.148 (34.964 to 35.331)	37.145 (36.956 to 37.334)		20.973 (20.824 to 21.122)	28.420 (28.246 to 28.593)	33.095 (32.902 to 33.287)	
Others	30.406 (30.132 to 30.681)	32.762 (32.475 to 33.048)	32.762 (32.475 to 33.048)		22.481 (22.231 to 22.730)	30.490 (30.186 to 30.794)	35.083 (34.731 to 35.436)	
Year				0.430				0.535
2004–2006	27.535 (27.424 to 27.646)	31.883 (31.767 to 31.999)	33.622 (33.505 to 33.739)		24.057 (23.951 to 24.164)	32.174 (32.058 to 32.290)	38.551 (38.430 to 38.672)	
2007–2009	29.665 (29.564 to 29.765)	34.035 (33.930 to 34.139)	36.463 (36.356 to 36.569)		22.606 (22.513 to 22.698)	28.351 (28.252 to 28.451)	34.151 (34.045 to 34.256)	
2010-2012	26.033 (25.940 to 26.126)	30.269 (30.170 to 30.368)	I		21.838 (21.751 to 21.926)	28.813 (28.715 to 28.910)	I	
2013-2015	25.678 (25.572 to 25.784)	1	1		23.766 (23.661 to 23.871)	1	1	
Site				<0.001				0.161
Renal pelvis	30.986 (30.921 to 31.051)	36.259 (36.189 to 36.329)	38.503 (38.430 to 38.577)		22.942 (22.883 to 23.001)	28.605 (28.539 to 28.672)	33.500 (33.425 to 33.576)	
Ureter	19.946 (19.870 to 20.021)	23.033 (22.951 to 23.114)	25.368 (25.279 to 25.456)		22.948 (22.868 to 23.028)	32.131 (32.037 to 32.225)	40.713 (40.604 to 40.822)	
Grade				<0.001				0.043
Well	13.707 (13.463 to 13.950)	13.707 (13.463 to 13.950)	13.707 (13.463 to 13.950)		18.635 (18.356 to 18.914)	21.710 (21.406 to 22.015)	40.163 (39.717 to 40.609)	
Moderate	10.664 (10.543 to 10.785)	13.393 (13.255 to 13.530)	15.852 (15.696 to 16.008)		10.147 (10.028 to 10.267)	18.691 (18.530 to 18.853)	27.546 (27.340 to 27.752)	
Poor	30.407 (30.315 to 30.498)	35.156 (35.059 to 35.253)	37.336 (37.236 to 37.437)		24.420 (24.335 to 24.506)	30.727 (30.633 to 30.822)	36.164 (36.060 to 36.269)	
								Continued

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Table 2 Cont	tinued							
	CSD (%)				DOC (%)			
Variables	3 years (95% CI)	5 years (95% CI)	8 years (95% CI)	P value	3 years (95% CI)	5 years (95% CI)	8years (95% CI)	P value
Undifferential	28.133 (28.066 to 28.200)	33.013 (32.939 to 33.086)	35.492 (35.413 to 35.572)		24.207 (24.142 to 24.271)	31.584 (31.510 to 31.658)	37.338 (37.252 to 37.424)	
Size				<0.001				0.733
<2	11.702 (11.603 to 11.801)	15.012 (14.896 to 15.129)	18.912 (18.770 to 19.054)		23.698 (23.568 to 23.828)	30.965 (30.815 to 31.114)	41.203 (41.019 to 41.386)	
2-4	20.339 (20.257 to 20.421)	25.952 (25.859 to 26.046)	27.404 (27.307 to 27.501)		21.685 (21.601 to 21.769)	29.655 (29.556 to 29.754)	38.855 (38.736 to 38.974)	
≥4	34.800 (34.728 to 34.872)	38.868 (38.792 to 38.944)	41.162 (41.083 to 41.242)		23.457 (23.393 to 23.521)	29.773 (29.701 to 29.845)	33.440 (33.361 to 33.519)	
Laterality				0.944				0.393
Left	26.970 (26.903 to 27.037)	31.835 (31.762 to 31.908)	33.636 (33.560 to 33.713)		22.586 (22.523 to 22.650)	29.551 (29.479 to 29.624)	35.006 (34.923 to 35.089)	
Right	27.092 (27.017 to 27.167)	31.010 (30.929 to 31.091)	33.867 (33.781 to 33.954)		23.335 (23.264 to 23.406)	30.305 (30.223 to 30.387)	37.564 (37.469 to 37.659)	
Surgery				0.001				0.980
Yes	26.715 (26.665 to 26.765)	31.206 (31.152 to 31.260)	33.490 (33.432 to 33.547)		22.940 (22.893 to 22.988)	29.950 (29.895 to 30.004)	36.234 (36.171 to 36.296)	
No/unknown	77.778 (76.910 to 78.646)	1	1		22.222 (21.522 to 22.923)	I	I	
Radiotherapy				<0.001				0.910
Yes	44.203 (43.994 to 44.413)	50.407 (50.191 to 50.623)	51.767 (51.548 to 51.986)		25.971 (25.788 to 26.155)	32.016 (31.816 to 32.217)	34.632 (34.422 to 34.842)	
No/unknown	25.670 (25.619 to 25.721)	29.951 (29.896 to 30.007)	32.292 (32.233 to 32.351)		22.699 (22.650 to 22.748)	29.733 (29.677 to 29.789)	36.285 (36.220 to 36.350)	
Chemotherapy				<0.001				0.003
Yes	36.276 (36.177 to 36.375)	41.751 (41.646 to 41.857)	43.283 (43.175 to 43.391)		20.156 (20.074 to 20.239)	26.807 (26.711 to 26.903)	30.702 (30.598 to 30.807)	
No/unknown	22.979 (22.922 to 23.035)	27.032 (26.970 to 27.094)	29.614 (29.547 to 29.681)		24.177 (24.119 to 24.235)	31.278 (31.212 to 31.344)	38.534 (38.457 to 38.611)	
Distant metastasis				<0.001				0.641
No	23.438 (23.388 to 23.488)	28.067 (28.012 to 28.122)	30.550 (30.491 to 30.609)		22.674 (22.624 to 22.723)	29.966 (29.909 to 30.024)	36.800 (36.734 to 36.866)	
Yes	65.586 (65.407 to 65.765)	67.967 (67.789 to 68.144)	67.967 (67.789 to 68.144)		25.548 (25.384 to 25.713)	28.750 (28.577 to 28.922)	28.750 (28.577 to 28.922)	
LNE	1	I	1	<0.001	1	1	1	0.941
LNP	I	I	I	<0.001	I	I	I	0.448
LNR	1	1	1	<0.001	1	1	1	<0.001
CSD, cancer-specific	c death; DOC, death due to o	ther causes; LNE, lymph noc	des examined; LNP, lymph noo	des positive; LN	NR, lymph nodes ratio.			

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Table 3 Multivariate analysis by proportional subdistribution hazards model and cause-specific hazards model for CSD among patients with upper tract urothelial carcinoma

	Proportional su	ubdistributior	n hazards model		Cause-speci	fic hazarc	ls model	
Variables	Coefficient	sdHR	95% CI	P value	Coefficient	csHR	95% CI	P valu
Age	-0.004	0.996	0.987 to 1.005	0.340	0.009	1.009	1.000 to 1.018	0.039
Sex								
Male	Reference				Reference			
Female	0.392	1.480	1.241 to 1.764	<0.001	0.301	1.351	1.134 to 1.611	<0.001
Race								
White	Reference				Reference			
Black	0.242	1.274	0.873 to 1.858	0.210	0.348	1.416	0.990 to 2.027	0.05
Other	0.201	1.223	0.930 to 1.607	0.150	0.164	1.178	0.899 to 1.544	0.23
Site								
Renal pelvis	Reference				Reference			
Ureter	-0.110	0.895	0.734 to 1.092	0.280	-0.106	0.899	0.732 to 1.105	0.31
Grade								
Well	Reference				Reference			
Moderate	-0.034	0.966	0.398 to 2.343	0.940	0.009	1.009	0.407 to 2.502	0.98
Poor	0.763	2.145	0.971 to 4.739	0.059	0.908	2.479	1.097 to 5.601	0.02
Undifferential	0.658	1.931	0.878 to 4.245	0.100	0.772	2.165	0.961 to 4.875	0.06
Size								
<2	Reference				Reference			
2–4	0.442	1.556	1.092 to 2.216	0.014	0.414	1.513	1.043 to 2.196	0.02
≥4	0.791	2.205	1.575 to 3.087	<0.001	0.881	2.414	1.691 to 3.447	<0.00
Surgery								
Yes	Reference				Reference			
No/unknown	0.791	2.205	1.292 to 3.761	0.004	0.752	2.120	0.990 to 4.539	0.05
Radiotherapy								
Yes	Reference				Reference			
No/unknown	-0.219	0.803	0.594 to 1.087	0.160	-0.240	0.787	0.595 to 1.040	0.09
Chemotherapy								
Yes	Reference				Reference			
No/unknown	0.025	1.025	0.829 to 1.269	0.820	0.171	1.187	0.972 to 1.450	0.09
Distant metastasis								
No	Reference				Reference			
Yes	0.881	2.414	1.842 to 3.163	<0.001	1.252	3.497	2.741 to 4.460	<0.00
LNE	-0.012	0.988	0.971 to 1.006	0.200	-0.013	0.987	0.972 to 1.002	0.09
LNP	0.062	1.064	1.022 to 1.107	0.002	0.069	1.072	1.032 to 1.113	<0.00
	0.627	1.873	1.435 to 2.445	< 0.001	0.934	2.544	1.965 to 3.294	<0.00

causes, which competitively leads to the fact that CSD incidence does not increase significantly with age. Sex and race have always been controversial prognostic factors. A previous study showed that race was a preoperative prognostic factor for patients with UTUC.<sup>29</sup> Moreover, another study found no statistically significant differences in survival between men and women.<sup>30</sup> However, the competing-risks model in our study showed that sex was a risk factor for UTUC, while race was not. This may be

because previous studies ignored the effect of competing risks. However, since most of the patients included in the SEER database are white, studies on different races need to be conducted.

Tumour size is also considered to be related to cancer prognosis. One study found 5-year recurrence-free survival rates of 46.9% and 25.8% in patients with UTUC with tumour sizes <3 cm and  $\geq 3 \text{ cm}$ , respectively.<sup>31</sup> The univariate and multivariate analyses performed in the



Figure 3 Nomogram based on the competing-risks analysis to predict CSD probabilities at 3, 5 and 8 years for patients with upper tract urothelial carcinoma. CSD, cancer-specific death; LNP, lymph nodes positive; LNR, lymph nodes ratio.

present study also indicated that tumour size was an influencing factor for CSD. The prognosis was worse for tumours larger than 2 cm. In addition, our research

also found that distant metastasis was an important risk factor for CSD. In terms of treatment methods, our study suggested that surgery status was a significant prognostic





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factor, which was consistent with the findings of Freifeld et al.<sup>32</sup> Surgery has long been considered the gold standard of UTUC treatment. However, radiotherapy status and chemotherapy status were not influencing factors for CSD in both competing-risks models. This result conflicted with some previous findings,<sup>33-35</sup> suggesting that traditional Cox regression analysis overestimated the effects of radiotherapy and chemotherapy. Obviously, the relative lack of information on the radiotherapy status and chemotherapy status in the SEER database may also lead to inaccurate results, and thus further exploration of these indicators is needed.

Some indicators related to lymph nodes (eg, distant lymph node metastasis, LNP and LNE) are important clinical information for cancer prognosis, but whether they are independent prognostic factors for UTUC has not been determined. One study found that lymph node metastases were significantly associated with reduced cancer-specific survival in univariate analysis.<sup>36</sup> It is worth noting that very few studies have investigated LNP, LNE and LNR. Our study is the first to use the SEER database to analyse the prognostic impact of these indicators on UTUC, and the results may be more accurate than those involving small samples. LNR is an emerging indicator that has been regarded as a prognostic factor in rectal cancer and breast cancer.<sup>37 38</sup> Our results also suggested that LNR was an important prognostic factor for UTUC. We found that LNE was an influencing factor for UTUC in the univariate analysis but not in the multivariate analysis. Moreover, both LNR and LNP entered the proportional SD model, suggesting that after adjusting for the effects of LNR and LNP, LNE was no longer an independent prognostic indicator. LNP was a prognostic factor in all of the analyses, indicating that it significantly influences the prognosis of UTUC.

We used the results from the above-mentioned proportional SD model to construct a nomogram that graphically represents the degree of influence of various prognostic factors. This nomogram can be used to predict the 3-year, 5-year and 8-year probabilities of CSD in patients with UTUC. The predictive function of the nomogram has been used for different types of cancer and has even been proposed as a new standard. The nomogram is easy to use. In order to calculate the CSD probability of a patient with UTUC, find the patient's sex (male or female) on the sex row, draw a vertical line on the dot row and obtain the sex score value. Repeat these steps for tumour size, M stage, surgery, LNP and LNR. Add the score values of each variable, find the total point on the total point axis, and draw a straight downward line to get the 3-year, 5-year and 8-year CSD probability of the patient with UTUC. For example, a woman (30 points) with a tumour size of 1.5 cm (0 points) at M1 stage (68 points) and had performed surgery (0 points), with an LNP equal to 5 (15 points) and LNR equal to 0.8 (45 points), and with a total score of 158 points, corresponds to 3-year, 5-year and 8-year CSD probability of 58%, 64% and 69%, respectively.

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The C-indexes for the nomogram all exceeded 0.6, demonstrating that the model provided a good fit to the available data. The prediction calibration curves in figure 4 were very close to the standard curve, indicating that the nomogram had good predictive ability. The results for the validation cohort also showed that the model was stable. Therefore, this model can help clinicians to quickly and easily determine the prognosis of individual patients and provide guidance in their clinical decision-making. However, the stability of the model

needs further verification. Our study used the large sample size and high-quality data from SEER database and competing-risks model, which guaranteed the accuracy of our study. However, 2 inevitably, our research had some limitations. First, the 8 established model is not comprehensive enough because the SEER database does not include all prognostic factors for UTUC. Second, the data available on treatment status are not sufficiently detailed to distinguish the impact of various treatment plans. Third, as a retrospective study, our results may be affected by confounding bias to some extent, so the conclusion needs to be further verified in future prospective studies. Fourth, Some of the causes of

future prospective studies. Fourth, Some of the causes of death in SEER may have been misclassified according to the death certificate report, which may also bring infor-mation bias to our study. **CONCLUSIONS** In summary, this study used a competing-risks model to determine the prognostic factors for UTUC. The proportional subdistribution hazards model showed that sex, tumour size, surgery, distant metastasis, LNP and LNR were associated with CSD, while LNE was not. The constructed nomogram can predict the 3-year, 5-year and mining, 8-year CSD probabilities of patients based on these rele-Al training, and similar technologies vant factors, which can support clinicians in making better decisions on the survival rates of individual patients.

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Patient consent for publication Not required.

Ethics approval All procedures performed in the present study were in accordance with the principles outlined in the 1964 Helsinki Declaration and its later amendments. Institutional review board approval and informed consent were not required in the current study because SEER research data are publicly available and all patient data are de-identified.

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