Development and Validation a Nomogram for Predicting the Third Station Lymph Node Metastasis in Early Gastric Cancer

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Research article

Keywords: Stomach Cancer, Lymph Nodes, Nomogram

DOI: https://doi.org/10.21203/rs.3.rs-142894/v1

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Abstract

Background:

This study aimed to evaluate the value of radiomic nomogram in predicting lymph node metastasis in T1-2 gastric cancer according to the No. 3 station lymph nodes.

Methods:

A total of 159 T1-2 gastric cancer (GC) patients who had undergone surgery with lymphadenectomy between March 2012 and November 2017 were retrospectively collected and divided into a primary cohort (n = 80) and a validation cohort (n = 79). Radiomic features were extracted from both tumor region and No. 3 station lymph nodes (LN) based on computed tomography (CT) images per patient. Then, key features were selected using minimum redundancy maximum relevance algorithm and fed into two radiomic signatures, respectively. Meanwhile, the predictive performance of clinical risk factors was studied. Finally, a nomogram was built by merging radiomic signatures and clinical risk factors and evaluated by the area under the receiver operator characteristic curve (AUC) as well as decision curve.

Results:

Two radiomic signatures, reflecting phenotypes of the tumor and LN respectively, were significantly associated with LN metastasis. A nomogram incorporating two radiomic signatures and CT-reported LN metastasis status showed good discrimination of LN metastasis in both the primary cohort (AUC: 0.915; 95% confidence interval [CI]: 0.832-0.998) and validation cohort (AUC: 0.908; 95%CI: 0.814-1.000). The decision curve also indicated its potential clinical usefulness.

Conclusions:

The nomogram received favorable predictive accuracy in predicting No.3 station LN metastasis in T1-2 GC, and could assist the choice of therapy.

Background

Gastric Cancer (GC) is rampant around the world, especially in East Asia [1]. Surgery is the primary treatment for patients with early gastric cancer (EGC), however, a number of sequelae (indigestion, iron deficiency, etc.) can seriously reduce the patient's quality of life for open surgery [2]. In order to improve the prognosis, less invasive surgical alternatives, such as endoscopic submucosal dissection and endoscopic mucosal resection, are used for the treatment of EGC [3]. However, endoscopic resection is only considered for tumors with a low risk of lymph node (LN) metastasis [4]. Studies have found that LN metastases exist in 8.2%-19.7% EGC [5–8]. Sentinel lymph node (SLN) biopsy, an invasive method, was used for detection of metastatic LNs in GC [9]. Studies have shown that SLN biopsy was a promising tool to assess the LN metastases in T1-2 GC patients [10–12]. However, there are still debates regarding the effectiveness of LN detection techniques and oncological safety of biopsy. Kitagawa et al [13]. and Miyashiro et al [14]. used different SLN biopsy methods, but consequently obtained different false-negative rates (7% and 46.4%, respectively). Besides, noninvasive medical imaging like CT is routinely used to assess perigastric LNs. However, the accuracy of CT detection of LN in early GC is about 60%, which is not good enough [15]. At present, it is still unable to accurately predict LN metastasis of EGC preoperatively.

Radiomics, the technique of converting medical images into mineable data and high-dimensional features, has been proven to improve diagnostic and prognostic accuracy in oncology [16–19]. Especially, it had been successfully applied to the prediction of LN metastasis in colorectal cancer and occult peritoneal metastasis in advanced GC [20, 21]. Whether radiomics has an advantage in predicting LN metastasis in T1-2 GC remains to be further studied.
The LNs of the stomach are given station numbers as No.1-No.16 [22, 23]. Among them, the No.3 station LNs are frequently invaded by tumor cells relatively [24–26]. Therefore, we evaluate the value of radiomic nomogram in predicting lymph node metastasis in T1-2 GC according to the No. 3 station LNs.

**Methods**

**Patients**

The Institutional Review Board of our hospital approved this retrospective study and the requirement for informed consent was waived.

The inclusion criteria for the primary and validation cohorts were as follows: (a) patients who underwent surgery with curative intent for T1-2 GC and with pathological results; (b) LN dissection performed; (c) excisional LN with detailed grouping and pathological diagnosis; (d) standard contrast-enhanced CT performed less than 10 days before surgical resection. The exclusion criteria were: (a) hypotensive drug taboo (such as glaucoma, prostatic hypertrophy, etc.); (b) preoperative therapy (radiotherapy, chemotherapy, or chemoradiotherapy); (c) concurrent with other tumors or diseases; (d) patients with variation of the left gastric artery; (e) invisible lesions on CT images.

A total of 159 patients between March 2012 and November 2017 were enrolled in this study (113 males, 46 females; average age, 61.78 ± 10.47 years). All the patients were randomly divided into two independent cohorts: a primary cohort, containing 80 patients (53 males, 27 females; average age, 61.78 ± 11.11 years), and a validation cohort, containing 79 patients (60 males, 19 females; average age, 61.78 ± 9.77 years).

Clinical data, including gender, age, carcinoembryonic antigen (CEA: 0–5 ng/ml), carbohydrate antigen 19–9 (CA19-9: 0–40u/ml), cancer antigen 125 (CA125: 0–35u/ml), pathologic grade (see detailed description in Supplementary Table S1), CT-reported LN metastasis status from radiologist, and tumor infiltration depth, were obtained by reviewing the medical records.

**CT data acquisition**

All patients fasted for at least 4 hours, and 20 mg anisodamine (654-2) was administered intramuscularly to reduce gastrointestinal peristalsis 10 minutes prior to CT examination. 800–1000 mL warm water was drank to distend the stomach. CT was performed using a 256-Slice (Brilliance iCT, ROYAL PHILIPS, Eindhoven, Netherlands) or a 64-slice (SOMATON sensation64, SIEMENS Healthcare, Muenchhen, Germany) multi-slice spiral CT. Patients underwent both unenhanced and two-phase enhanced CT examinations (arterial phase: 35 s after injection; venous phase: 70 s after injection). The CT scans, covering the entire stomach region, were acquired during a breath-hold with the patient supine in all of the phases. During the enhanced CT scan, patients were infused with 1.5 mL/kg of the non-ionic contrast material (iohexol, Yangzi River Pharmaceutical Group, Jiangsu, China; iodine concentration: 300 mg/mL) with a pump injector (Ulrich CT Plus 150, Ulrich Medical, Ulm, Germany) at a rate of 3.0 mL/s into the antecubital vein. The imaging parameters were as follows: 120 kV; 220–250 mAs; rotation time: 0.5 s; detector collimation: 128 × 0.625 mm or 32 × 0.6 mm; field of view: 400 × 400 mm; matrix: 512 × 512; reconstruction slice thickness: 5 mm for axial plane, and 3 mm for coronal and sagittal plane.

**Pipeline**

The pipeline of this study includes five steps: lesion detection, region of interest (ROI) segmentation, radiomic feature extraction, radiomic signature building, and nomogram construction and evaluation (Fig. 1).

**Detection of Lesion on CT Images**
All CT images were reviewed by a radiologist with more than 10 years of experience in GC diagnosis. Localization of GC lesions: The 159 patients selected in this study all had the results of gastroscopy and CT examination. Combined with gastroscopy and CT images (axial, coronal and sagittal images), the lesions could be located. The diagnostic criteria of CT-reported LN metastasis-positive were shown as follows: short-axis diameter of LN ≥ 5 mm, the ratio of short diameter to long diameter of LN ≥ 0.7, and the plain CT value of LN ≥ 25 HU or venous phase CT value of LN ≥ 75 HU; or multiple LNs were fused together even if above conditions were not satisfied.

**ROI Segmentation on CT Images**

Two 2-dimensional ROIs were manually segmented by a radiologist with more than 10 years of experience in GC diagnosis. The first ROI (ROI-1) was delineated on the tumor in the slice with the maximum tumor lesion. The second ROI (ROI-2) was delineated on the region of No.3 station LNs around the lesser curvature of stomach. ROI segmentation was performed using ITK-SNAP software (version 2.2.0; [www.itksnap.org](http://www.itksnap.org)) on the venous phase CT images with axial view (see Supplementary A1 for detail).

**Extraction of Radiomic Features**

Two feature groups were extracted from two ROIs, with each group containing 273 features [27, 28]. These features were divided into 4 categories: (a) shape and size features, (b) gray intensity features, (c) texture features, and (d) wavelet features. The feature extraction was implemented using MATLAB (version 2014a; Mathworks, Natick, MA, USA). Radiomic features of all patients were standardized by the z-score method, based on the parameters calculated from the primary cohort. More information about the radiomic feature extraction is shown in Supplementary A2.

**Radiomic Signature Construction**

Radiomic feature selection and signature building were performed in the primary cohort for ROI-1 and ROI-2, respectively. More details are described as follows. In order to avoid model over-fitting and improve performance, feature selection was performed to match the sample size (Supplementary A3).

First, the minimum redundancy maximum relevance algorithm (mRMR) ranked each feature based on its relevance to LN metastasis status, and the ranking process was able to consider the redundancy of these features at the same time [29]. Since the number of predictors should be kept within 1/10 ~ 1/3 of the size of the group that contains the smallest cases in the primary cohort (LN metastasis-positive group, n = 22) [30], the number of potential features was limited to 7 or less in this study.

Second, five-fold cross-validation was performed multiple times on the primary cohort to find the optimal number of features with the best performance based on ranked features. Then a radiomic signature (RS1) reflecting phenotype of ROI-1 and a radiomic signature (RS2) reflecting phenotype of ROI-2, were built as independent predictors of LN metastasis using selected features, respectively. For each radiomic signature, the signature score was calculated to reflect the risk of LN metastasis. The predictive performance of the radiomic signatures were quantitatively tested using the area under the receiver operator characteristic (ROC) curve in both the primary and validation cohorts.

**Construction and Evaluation of Nomogram**

Univariate analysis and multivariate analysis were used to screen out significant clinical risk factors. For univariate analysis, continuous variables were assessed using independent t-test or Mann-Whitney U test for differences between different groups, and categorical variables were assessed by Chi-squared test. As for multivariate analysis, we performed multivariate logistic regression to screen out key factors. Furthermore, multivariate logistic regression was used to merge two radiomic signatures and clinical risk factors into a nomogram. Meanwhile, we performed variable selection according to the p-values of the logistic regression. After that, the calibration curves and Hosmer-Lemeshow test were used to assess the goodness-of-fit of the nomogram, and the AUC was used to quantify its predictive performance. For
assessing overfitting, DeLong test was adapted to compare AUCs between primary and validation cohorts. Moreover, we used net reclassification index (NRI) to compare the performance between nomogram and clinical risk factors, and quantify the improvement in predictive performance.

Furthermore, a stratified analysis was used to evaluate the influence of clinical factors to the nomogram. In addition, we performed a subgroup analysis to evaluate the additional value of the nomogram in the CT-reported LN metastasis-negative (CT-LNM0) subgroup. Since the number of metastasis in No.4 station LNs (left greater curvature) ranked only behind No.3 station LNs (Supplementary Table S2), we further validated our nomogram on No.4 station LNs.

Finally, to estimate the clinical utility of the nomogram, decision curve analysis (DCA) was performed by calculating the net benefits using a range of threshold probabilities.

**Statistical Analysis**

All statistical analysis was performed using R software (version 3.3.4; http://www.Rproject.org). A two-sided $P$ value $< 0.05$ was used to indicate statistical significance.

**Results**

**Clinical characteristics**

Table 1 summarizes the patients’ clinical risk factors in both the primary and validation cohorts. There is no significant difference in the probability of LN metastasis between the two cohorts ($P = 0.384$). Univariable analysis showed that CT-reported LN metastasis status from the radiologist were significantly correlated with pathological LN metastasis status ($P < 0.05$), while CA125 was significantly correlated with LN metastasis status only in the primary cohort and tumor infiltration depth in the validation cohort. After multivariable analysis we chose the CT-reported LN metastasis status to predict LN metastasis.
| Characteristic                  | Primary cohort | Validation cohort | $P$-value | Primary cohort | Validation cohort | $P$-value |
|-------------------------------|----------------|------------------|-----------|----------------|------------------|-----------|
|                               | LNM (+)        | LNM (-)          |           | LNM (+)        | LNM (-)          |           |
| Sex, No. (%)                  | 0.451          | 0.366            |           |                |                  |           |
| Male                          | 16 (72.73)     | 37 (63.79)       |           | 11 (64.71)     | 49 (79.03)       |           |
| Female                        | 6 (27.27)      | 21 (36.21)       | 0.276     | 6 (35.29)      | 13 (20.97)       | 0.173     |
| Age, mean ± SD, years         | 59.82 ± 11.93  | 62.52 ± 10.69    |           | 58.82 ± 8.83   | 62.60 ± 9.86     |           |
| CEA, No. (%)                  | 0.114          | 0.204            |           |                |                  |           |
| Median (IQR)                  | 2.19 (1.44–2.54)| 1.31 (1.02–2.44) |           | 2.46 (1.50–3.10)| 1.86 (1.14–2.76)|           |
| Normal                        | 22 (100.00)    | 55 (94.83)       |           | 17 (100.00)    | 59 (95.16)       |           |
| Abnormal                      | 0 (0.00)       | 3 (5.17)         |           | 0 (0.00)       | 3 (4.84)         |           |
| CA19-9, No. (%)               | 0.171          | 0.867            |           |                |                  |           |
| Median (IQR)                  | 11.95 (8.22–16.46)| 8.97 (6.24–15.59)|           | 12.30 (7.12–16.11)| 9.99 (6.56–15.81)|           |
| Normal                        | 21 (95.45)     | 57 (98.28)       |           | 17 (100.00)    | 60 (96.77)       |           |
| Abnormal                      | 1 (4.55)       | 1 (1.72)         |           | 0 (0.00)       | 2 (3.23)         |           |
| CA125, No. (%)                | 0.035*         | 0.308            |           |                |                  |           |
| Median (IQR)                  | 14.18 (8.70–22.66)| 8.81 (5.43–16.04)|           | 10.25 (8.69–13.10)| 8.85 (4.95–15.69)|           |
| Normal                        | 20 (90.91)     | 57 (98.28)       |           | 16 (94.12)     | 60 (96.77)       |           |
| Abnormal                      | 2 (9.09)       | 1 (1.72)         |           | 1 (5.88)       | 2 (3.23)         |           |
| Pathologic grade, No. (%)     | 0.349          | 0.410            |           |                |                  |           |
| Low grade                     | 12 (54.55)     | 22 (37.93)       |           | 6 (35.29)      | 20 (32.26)       |           |
| Median grade                  | 9 (40.91)      | 34 (58.62)       |           | 11 (64.71)     | 36 (58.06)       |           |
| High grade                    | 1 (4.54)       | 2 (3.45)         |           | 0 (0.00)       | 6 (9.68)         |           |
| CT-reported LNM status, No. (%)| < 0.001*       | < 0.001*         |           |                |                  |           |
| LNM0                          | 12 (54.55)     | 54 (93.10)       |           | 11 (64.71)     | 58 (93.55)       |           |
| LNM1                          | 10 (45.45)     | 4 (6.90)         |           | 6 (35.29)      | 4 (6.45)         |           |
| Tumor infiltration depth, No. (%)| 0.181          | < 0.001*         |           |                |                  |           |
| T1a                           | 4 (18.18)      | 20 (33.48)       |           | 0 (0.00)       | 20 (32.26)       |           |
| T1b                           | 9 (40.91)      | 25 (43.10)       |           | 4 (23.53)      | 31 (50.00)       |           |
### Establishment of Radiomic Signature

During the feature selection, mRMR selected top 10 radiomic features from ROI-1 and top 10 radiomic features from ROI-2 in the primary cohort, respectively. As shown in Supplementary Figure S1 and Supplementary Table S3, the cross-validation reserved 4 features from ROI-1 and 2 features from ROI-2. The heatmaps of these features and unsupervised cluster partitioning are shown in Supplementary Figure S2. Significant association was found between these features and LN metastasis status. Two radiomic signatures were built using linear combination of these radiomic features (4 features from ROI-1 for RS1 and 2 features from ROI-2 for RS2), and the signature score calculation are presented in Supplementary A4. As shown in Fig. 2 and Table 2, both of the two radiomic signatures showed significant predictive ability of LN metastasis in primary cohort (AUC of RS1: 0.831, 95% confidence interval [CI]: 0.725–0.937, and AUC of RS2: 0.761, 95% CI: 0.629–0.893) and validation cohort (AUC of RS1: 0.852, 95% CI: 0.742–0.962, and AUC of RS2: 0.763, 95% CI: 0.626-0.900).

| Characteristic | Primary cohort | Validation cohort | Validation cohort |
|----------------|----------------|------------------|------------------|
|                | LNM (+)        | LNM (-)          | LNM (+)          | LNM (-)          | P-value | P-value |
| T2             | 9(40.91)       | 13(22.41)        | 13(76.47)        | 11(17.74)        |         |         |

NOTICE. P-value was derived from the univariable association analyses between each characteristic and LNM status; For univariate analysis, independent t-test or Mann-Whitney U test were used for continuous variables and Chi-squared test for categorical variables. * means P-value < 0.05; LNM0 refers to CT-reported LNM-negative; LNM1 refers to CT-reported LNM-positive. Abbreviations: LNM, lymph node metastasis; SD, standard deviation; CEA, carcinoembryonic antigen; CA19-9, carbohydrate antigen 19 – 9; CA125, cancer antigen 125; CT, computed tomography.
Table 2
Performance evaluation of models.

| Index       | Primary Cohort | Validation Cohort |
|-------------|----------------|-------------------|
|             | Radiomic signature-1 | Radiomic signature-2 | CT-reported LN metastasis status | Nomogram | Radiomic signature-1 | Radiomic signature-2 | CT-reported LN metastasis status | Nomogram |
| TP          | 18             | 14                | 10                | 20        | 12               | 12                 | 6                 | 15        |
| TN          | 44             | 46                | 54                | 54        | 54               | 42                 | 58                | 56        |
| FN          | 4              | 8                 | 12                | 2         | 5                | 5                  | 11                | 2         |
| FP          | 14             | 12                | 4                 | 4         | 8                | 20                 | 4                 | 6         |
| Acc         | 77.50%         | 75.00%            | 80.00%            | 92.50%    | 83.54%           | 68.35%             | 81.01%            | 89.87%    |
| Sen         | 0.818          | 0.636             | 0.455             | 0.909     | 0.706            | 0.706              | 0.353             | 0.882     |
| Spe         | 0.759          | 0.793             | 0.931             | 0.931     | 0.871            | 0.677              | 0.935             | 0.903     |
| PPV         | 0.563          | 0.538             | 0.714             | 0.833     | 0.600            | 0.375              | 0.600             | 0.714     |
| NPV         | 0.917          | 0.852             | 0.818             | 0.964     | 0.915            | 0.894              | 0.841             | 0.966     |
| AUC         | 0.831          | 0.761             | 0.693             | 0.915     | 0.852            | 0.763              | 0.644             | 0.908     |

(95% CI) (0.725–0.937) (0.629–0.893) (0.581–0.804) (0.832–0.998) (0.742–0.962) (0.626–0.900) (0.523–0.765) (0.814–1.000)

NOTICE. Abbreviations: TP, true positive; TN, true negative; FN, false negative; FP, false positive; Acc, accuracy; Sen, sensitivity; Spe, specificity; PPV, positive predictive value; NPV, negative predictive value; AUC, area under curve; CI, confidence interval.

Construction of Nomogram

During the multivariate logistic regression analysis, the two radiomic signatures and one clinical risk factor (CT-reported LN metastasis status) were identified as independent predictors of LN metastasis in T1-2 GC patients (Supplementary Table S4). An individualized nomogram was built using the regression method to predict the LN metastasis probability (Fig. 3A).

Evaluation of Nomogram

As shown in Fig. 2 and Table 2, our nomogram reached an AUC of 0.915 (95% CI: 0.832–0.998) in the primary cohort and an AUC of 0.908 (95% CI: 0.814–1.000) in the validation cohort, which were better than CT-reported LN metastasis status, RS1, and RS2. The NRI also demonstrated that the nomogram had better predictive ability than the CT-reported LN
metastasis status in the primary cohort (NRI = 0.339, \(P < 0.001\)) and validation cohort (NRI = 0.301, \(P < 0.001\)). The DeLong test revealed that difference was not significant between AUCs of our nomogram in primary and validation cohorts (\(P = 0.908\)), further indicating the robustness of our nomogram. As shown in Fig. 3B and C, the calibration curves of the nomogram demonstrate a good fit of the nomogram in both the primary and validation cohorts. The Hosmer-Lemeshow test also showed good performance of our nomogram in the primary cohort (\(P = 0.147\)) and validation cohort (\(P = 0.903\)).

Notably, the subgroup analysis showed that our nomogram had a good discriminatory ability in the CT-LNM0 subgroup (\(n = 109, \text{AUC: 0.904; 95\% CI: 0.816–0.993; Fig. 4}\)).

We also implemented stratified analysis, more details were presented in Supplementary A5 and Supplementary Figure S3. The results showed that our nomogram worked well in gender, age, pathologic grade and tumor infiltration depth subsets (DeLong test, \(P > 0.05\)).

Moreover, we selected 9 patients with LN metastasis and 11 patients with non-LN metastasis at No.4 station as a validation set to further validate our nomogram. Interestingly, our nomogram also showed a good performance on this station (AUC, 0.824; 95\% CI, 0.517-1; Supplementary Figure S4).

The decision curve of the nomogram is presented in Fig. 5. With a threshold of 0 to 0.85, patients using nomogram will have more diagnostic benefits than all-metastasis or none-metastasis strategies.

**Discussion**

In this study, an easy-to-use radiomic nomogram was established to identify LN metastasis of T1-2 GC preoperatively. The nomogram, incorporating two radiomic signatures and CT-reported LN metastasis status, showed the best discrimination ability of LN metastasis in both the primary and validation cohorts. The nomogram could assist the formulation of clinical treatment scheme.

Although the lymphatic system around the stomach is very complex [31], the No.3 station LNs belong to the N1 LNS [2, 4]. Previous researches showed that the incidence rate of LN metastasis of EGC in No.3 station was the highest (No.3, 11.6\%; No.1, 2.5\%; No.2, 4.8\%; No.4, 6.5\%; No.5, 0.5\%; No.6, 7.6\%, respectively) (Supplementary Table S5) [24–26]. Therefore, the diagnosis of LN metastasis in No.3 LNs is clinically useful.

We analyzed the radiomic features in the two significant radiomic signatures. The radiomic features used in RS1 included: (1) 'X1_fos_skewness' describes the shape of a probability distribution of the voxel intensity histogram, and reflects the distribution symmetry. (2) 'X0_fos_variance' measures the spread of intensity distribution about the mean value, and reflects the uniformity of distribution. (3) 'X3_fos_root_mean_square' is the root mean square of the voxels intensity value. (4) High 'X1_GLCM_dissimilarity' means there is a great disparity in intensity value among neighboring voxels. These radiomic features might quantify intratumor heterogeneity, and thus could predict the invasiveness of the tumor and the probability of LN metastasis [32]. The final selected radiomic features of lymph nodes consisted of: (1) 'X1_GLRLM_energy' measures of the magnitude of voxel values in an image describes the overall density of the lymph volume, (2) High 'X1_GLCM_cluster_prominence' implies more asymmetry. These radiomic features might indicate the high image intensity and heterogeneity in the No.3 station LN region, and thus the sign of LN metastasis. We have showed two examples of patients with and without LN metastasis (Supplementary A6 and Supplementary Figure S5). The CT images also demonstrated that higher heterogeneity of the primary tumor and No.3 LN region led to higher probability of LN metastasis.

In this study, CT-reported LN metastasis status from the radiologist was significantly correlated with LN metastasis in univariable analysis. This subjective judgment was also included in our nomogram. We also found that CA125 was
significantly associated with LN metastasis in the primary cohort \( (P = 0.035) \), but had no significance in the validation cohort. This may be caused by the relatively small sample size and baseline deviation. Moreover, the positive rate of CA125 was very low in early GC [33].

We conducted some stratified analysis, the results showed that the performance of our nomogram was not affected by gender, age, pathologic grade and tumor infiltration depth factors. In addition, we tested the correlations between the radiomic features and clinical risk factors using Pearson correlation analysis (Supplementary Figure S6). There was no correlation between radiomic features and clinical risk factors, which pointed that the radiomic features might be a good supplement to clinical factors. The good performance of our nomogram in CT-LNM0 subgroup also demonstrated the additional value of the nomogram to the radiologists.

More interestingly, the nomogram trained from phenotype of No.3 station LNs also showed a positive role in predicting LN metastasis in No.4 station LNs. This finding indicated that the radiomic signature from the LN region did reflect the early change of phenotype of LNs. Thus, our nomogram may be used in other stations of LNs.

There are some limitations in this study. Firstly, the relatively small sample size of this study. Secondly, the lack of the external validation. Thirdly, the presence of lymphatic invasion and LN micrometastasis have also been considered as important risk factors for LN metastasis in EGC [34–36], however, these factors were not routinely collected in our center. Finally, cases with invisible lesions on CT images were excluded, so some patients could not use the nomogram. These problems need to be further studied.

**Conclusions**

In summary, the nomogram received favorable predictive accuracy in predicting No.3 station LNs metastasis in T1-2 GC, and thus may assist the formulation of clinical treatment scheme.

**Abbreviations**

GC  
Gastric cancer  
LN  
Lymph node  
CT  
Computed tomography  
AUC  
Area under the receiver operator characteristic curve  
CI  
Confidence interval  
EGC  
Early gastric cancer  
ROI  
Region of interest  
mRMR  
Minimum redundancy maximum relevance algorithm  
RS  
Radiomic signature  
ROC
Declarations

Ethics approval and consent to participate

This study was approved by the institutional review board at the corresponding author’s institution (The Affiliated People’s Hospital of JiangSu University). As this is a retrospective case-control study, the need for informed written patient consent was waived by the ethics committee. All patient data were analyzed anonymously.

Consent for publication

The manuscript is approved by all participants for publication.

Availability of data and materials

The datasets used and analyzed during the current study available from the corresponding author on reasonable request (13913433095@163.com).

Competing interests

The authors declare that they have no competing interests.

Funding

This work was supported by the National Key R&D Program of China (2017YFC1308700, 2017YFA0205200, 2017YFC1309100, 2017YFA0700401), Jiangsu Provincial Research Foundation for Basic Research of China (BK20151334), National Natural Science Foundation of China (81971776, 81771924, 81227901, 81501616), Zhenjiang Innovation Capacity Building Program (technological infrastructure) - R & D project of China (SS2015023), Jiangsu Provincial Key R & D Special Fund (BE2015666), the Beijing Natural Science Foundation (L182061), the Bureau of International Cooperation of Chinese Academy of Sciences (173211KYSB20160053), and the Youth Innovation Promotion Association CAS (2017175).

Author contributions

XXW, CL, DD, JT, and XHS: Conceptualization, Methodology;
XXW, DD, and CL: Data curation;
CL, FFH, MJF, LWZ and LZZ: Software, Validation, Visualization;
XXW, CL: Writing - original draft;
JT, XHS, DD: Funding acquisition, Project administration, Supervision.

Acknowledgements
Not applicable.

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