Prognostic significance of tumor deposits in radically resected gastric cancer: a retrospective study of a cohort of 1915 Chinese individuals

Menglong Zhou1,2,3†, Wang Yang1,2,3†, Wei Zou1,2,3†, Jianing Yang1,2,3, Changming Zhou2,4, Zhiyuan Zhang1,2,3, Yaqi Wang1,2,3, Jing Zhang1,2,3, Yan Wang1,2,3, Guichao Li1,2,3, Zhen Zhang1,2,3* and Fan Xia1,2,3*

Abstract

Background: Tumor deposits (TDs) have been identified as an independent prognostic factor in gastric cancer (GC). However, the associated clinicopathological factors and how to simply and reasonably incorporate TD into the TNM staging system remain undetermined. The aim of the current study was therefore to assess the significance of TD among radically resected GC patients.

Methods: We retrospectively reviewed 1915 patients undergoing radical resection between 2007 and 2012. The patients were classified into two groups according to TD status (absent vs. present), and the clinicopathologic characteristics, DFS, and OS were compared. Associations of TD presence with other clinicopathologic factors were evaluated by logistic regression analysis. Univariate and multivariate Cox regression analyses were performed to determine the prognostic factors for DFS and OS in the primary cohort. Propensity score matching (PSM) was performed to reduce the possibility of selection bias according to the presence of TD. External validation of previously proposed modified staging systems incorporating TD was conducted.

Results: The detection rate of TD was 10.5% (201/1915). The presence of TD was significantly related to unfavorable clinicopathologic variables, including advanced T and N categories. According to the multivariate Cox regression analysis, the presence of TD was identified as an independent prognostic factor for DFS and OS in the primary cohort (both \( P \) < 0.001). In the after-PSM cohort, TD presence also significantly shortened DFS and OS. In the external validation, one system that incorporated TD into the pTNM stage had the best performance.

Conclusions: The presence of TD was significantly associated with poor survival in radically resected GC patients. The incorporation of TD into the TNM staging system can further improve the predictive capability. A multicenter cohort with a large sample size is needed to determine the appropriate method of incorporation.

Keywords: Gastric cancer, Tumor deposit, Prognosis, Propensity score

Introduction

According to the latest epidemiological data, gastric cancer (GC) ranks fifth and fourth in terms of the estimated number of new cases and deaths worldwide, respectively [1]. TNM stage is the most commonly used parameter for determining prognosis and for treatment
decision-making. Apart from TNM stage, there are other common clinicopathological factors significantly related to prognosis, such as lymphovascular invasion (LVI), perineural invasion (PNI), and tumor grade, which have also been identified in previous studies [2–5].

TD, as a typical histopathological feature of colorectal tumors, was first proposed by Gabriel W.B. as early as 1935 [6]. TDs are tumor-like nodular masses in addition to the primary tumor that are located in the fat tissue of the mesocolon and mesorectum. The role of TD in colorectal cancer has been widely studied [7]. Eventually, the association of poor prognosis with TD in colorectal cancer was confirmed [8–10], and TD status was included in the N1c category of colorectal cancer in the 7th edition of the AJCC staging system [11]. In addition, studies have shown that TD also exists in other cancers, such as breast cancer [12], thyroid cancer [13], lung cancer [14], gastric cancer (GC) [15–17], and pancreatic carcinoma [18].

In gastric cancer, with improvements in surgical and pathological detection technology, the number of detected TDs is gradually increasing, and its role in the staging and prognosis of GC has increasingly become a research focus. First, the presence of TD has been identified as an independent prognostic factor for GC by most previous studies, but the results regarding the clinicopathological factors associated with its presence have been inconclusive [16, 17, 19–28]. Second, whether there is a place for TD in staging and how to simply and reasonably incorporate TDs into the TNM staging system remain undetermined. TDs are likely to be considered metastatic lymph nodes [21, 26, 28], but several studies have indicated that TD should be regarded as serosal invasion [16, 23]. At present, the Japanese gastric cancer treatment guidelines recommend that each TD found in the lymphatic drainage area of the primary tumor should be included in the N category as a metastatic lymph node [29]. However, in the eighth edition of the American Joint Committee on Cancer (AJCC) GC staging system, TD first existed as one of the nineteen registry data collection variables, without mention of the role it may play in prognosis or its incorporation into staging [30]. Therefore, more studies are warranted to clarify the aforementioned aspects.

The current study retrospectively enrolled 1915 patients with resectable GC who underwent radical surgery in our center with the aim of comprehensively evaluating the effect of TD on resectable GC patients, including its association with clinicopathologic factors and its influence on prognosis. In addition, this study summarized and evaluated the existing methods of incorporating TD into the TNM staging system.

Materials and methods
Patient population
A cohort of 1915 GC patients who underwent radical resection between January 2007 and December 2012 at Fudan University Shanghai Cancer Center (FUSCC) was retrospectively identified. Patients were eligible if they met the following criteria: (1) histopathologically confirmed gastric or gastroesophageal junction adenocarcinoma; (2) no evidence of distant metastasis or peritoneal seeding on preoperative staging; (3) R0 resection with at least D1 lymphadenectomy; and (4) no preoperative chemotherapy or radiotherapy. The patients’ clinicopathologic and therapeutic factors were retrospectively collected. The current study was approved by the medical ethics committee of FUSCC, and the study was conducted in accordance with the Declaration of Helsinki.

Treatment delivery
Peritoneal washings were not routinely performed during the study period. Adjuvant treatment including chemotherapy (ChT) or ChT plus concurrent chemoradiotherapy (CRT) was recommended for all patients with stage II–III disease. The ChT regimens included single-agent fluoropyrimidines (tegafur gimeracil oteracil potassium capsules (S-1) or capecitabine), dual drug combinations (fluoropyrimidine plus platinum) or three drug combinations (fluoropyrimidine, platinum plus epirubicin, or taxanes).

Radiotherapy was given with 6 MV photons using either three-dimensional CRT or intensity-modulated radiation therapy (IMRT). Patients were treated with 25 to 28 fractions of 1.8 Gy for a total dose of 45 to 50.4 Gy (5 fractions/week). The clinical target volume (CTV) encompassed the preoperative tumor extension, tumor bed, anastomosis site, and regional draining LNs. The planning target volume (PTV) margin was 0.5 to 1.0 cm considering the individual uncertainties. The remnant stomach was not routinely included within the radiation field. Concurrent ChT regimens included (1) a continuous intravenous infusion of 225 mg/m² of 5-fluorouracil (5-FU) for 120 h each week and (2) 825 mg/m² of capecitabine twice daily from day 1 to 5 weekly or S-1 30 mg/m² twice daily from day 1 to 5 weekly.

Follow-up
All patients were followed-up every 3 months for the first 2 years, then every 6 months until the fifth year, and yearly thereafter. Follow-up examinations included a complete history and physical examination, measurement of serum tumor biomarkers, CT scans of the chest, abdomen and pelvis each time, and endoscopy each year. Disease-free survival (DFS) was defined as the time from surgery to death, locoregional recurrence, or
distant recurrence. Overall survival (OS) was defined as the interval from surgery to date of death from any cause or date of the most recent follow-up. Patients for whom none of these events were recorded were censored at the date of their last known contact. The median follow-up time for all the patients was 93.7 months (IQR 73.7–112.9 months).

Pathology and definition of TD
The histological sections of tumor specimens were reviewed independently by two pathologists, and disagreements were confirmed by a third pathologist. The tumor was classified according to the 8th edition AJCC staging system for GC. TDs are defined and evaluated as discrete foci of cancer cells found in the perigastric fat or adjacent ligament away from the primary lesion but within the area of locoregional lymph node stations. No identifiable lymph node tissue or vascular or neural structure should be found. The shape, contour and size of the deposits are not assessed in these designations.

Statistical analysis
Comparison of the clinicopathological characteristics between the TD-absent and TD-present groups was performed. Data are summarized as the mean (± standard deviation) or the median (with range) for continuous variables and numbers (percentages) for categorical variables. Continuous data were compared using the t-test/Wilcoxon rank sum test, whereas categorical data were analyzed using the chi-square test. The standardized mean difference (SMD) was reported to assess the balance of covariates between the two groups. Univariate and multivariate models using logistic regression were conducted to assess the relationship between TD status and other clinicopathological characteristics.

DFS and OS were calculated using the Kaplan-Meier method, and the log-rank test was employed to determine the significance. Associations between clinicopathologic features and survival were assessed with univariate analysis. The potentially relevant factors obtained from the univariate analysis were assessed in the multivariate model using Cox regression. Hazard ratios (HR) and 95% confidence intervals (CI) were calculated.

In the logistic and Cox regression analyses, age and tumor size were treated as categorical variables and dichotomized at the cohort median age of 58 years and tumor diameter of 3.2 cm. The number of retrieved lymph nodes was also treated as a categorical variable and dichotomized at the accepted cutoff of 15.

To reduce selection bias, a 1:3 propensity score-matching analysis was performed between the TD-absent and TD-present groups. Propensity scores were estimated using a logistic regression model and the following covariates: sex, age, tumor location, tumor size, histologic grade, vascular emboli, lymphatic and perineural invasion (VELIPI), T category, N category, number of retrieved lymph nodes, and adjuvant treatment. Using these propensity scores, patients with TDs (TD-present group) were individually matched to patients without TDs (TD-absent group).

The predictive abilities of the TNM staging system and other modified systems were evaluated by the χ² value, the area under the receiver operating characteristic curve (AUC), Harrell’s concordance index (C-index) and Akaike information criterion (AIC). Larger χ², AUC, and C-index values and a smaller AIC value indicate that the system has a better discriminative ability.

A two-sided significance level of 0.05 was applied. All statistical analyses were performed using the R statistical software package (version 4.2.1; R Project for Statistical Computing, Vienna, Austria).

Results
Clinicopathologic characteristics
In total, 1915 patients were enrolled in this cohort, and TD was present in 201 patients. Thus, the detection rate was 10.5%. The clinical and pathologic characteristics of the cohort were summarized according to TD status (Table 1). Of the 1915 patients, 1333 were men (69.6%), and 582 were women (30.4%), with a median age of 58 years (range, 19 to 84 years). The mean number of lymph nodes dissected per patient was 23.4 ± 8.4. Of these patients, 71.9% (1376/1915) received adjuvant treatment, including 1242 patients who received adjuvant ChT and 134 patients who received adjuvant CRT.

Among the 201 TD-present patients, a total of 329 TDs were detected, ranging from 1 to 12 TDs. The median and average numbers were 1 and 1.64, respectively. A total of 132 patients had 1 TD, 43 patients had 2 TDs, and 26 patients had ≥ 3 TDs. The common distribution areas were the lesser curvature omentum (n = 191), greater curvature omentum (n = 191), greater omentum (n = 80), and other areas (n = 16) (Supplementary Table 1).

The presence of TD was significantly associated with older age, distal tumor location, larger tumor size, advanced T category, advanced N category, and VELIPI. In multivariate logistic regression, older age, larger tumor size, advanced T category, and advanced N category were recognized as independent risk factors for the presence of TD (Table 2).

Survival analysis
According to the univariate analysis shown in Table 3, the following 10 clinicopathologic characteristics were demonstrated to be significantly associated with DFS and OS in the primary cohort: age (< 58 vs. ≥ 58,
Table 1  Comparison of clinicopathologic features between TD-absent and TD-present gastric cancer patients before and after PSM

| Variables                  | Before PSM |            | After PSM |            |
|---------------------------|------------|------------|-----------|------------|
|                           | TD-absent  | TD-present | P value   | SMD        |
|                           | n          | %          | n         | %          |
|                           |            |            |           |            |
| Overall                   | 1714       | 201        | 491       | 191        |
| Sex                       |            |            |           |            |
| Male                      | 1184       | 69.1       | 149       | 74.1       | 0.164 | 0.112 |
| Female                    | 530        | 30.9       | 52        | 25.9       | 1.36  | 0.277 |
| Age (years)               |            |            |           |            |
| < 58                      | 917        | 53.5       | 78        | 38.8       | < 0.001 | 0.298 |
| ≥ 58                      | 797        | 46.5       | 123       | 61.2       | 0.112  | 0.217 |
| Mean (SD)                 | 56.9 ± 11.1| 61.0 ± 10.6| 59.9 ± 10.6| 60.7 ± 10.6| 0.868  | 0.022 |
| Median (range)            | 58 (19-82) | 62 (33-84) | 61 (28-82) | 62 (33-84) | 0.798  | 0.029 |
| Tumor location            |            |            |           |            |
| GEJ + Upper 1/3           | 431        | 25.1       | 68        | 33.8       | 0.013  | 0.217 |
| Middle 1/3                | 417        | 24.3       | 51        | 25.4       | 0.141  | 0.122 |
| Lower 1/3                 | 866        | 50.5       | 82        | 40.8       | 0.868  | 0.022 |
| Tumor size                |            |            |           |            |
| <3.2 cm                   | 922        | 53.8       | 45        | 22.4       | < 0.001 | 0.683 |
| ≥ 3.2 cm                  | 792        | 46.2       | 156       | 77.6       | 0.142  | 0.023 |
| Mean (SD)                 | 3.6 ± 2.1  | 5.1 ± 2.3  | 4.7 ± 2.3 | 4.9 ± 2.1  | 0.964  | 0.023 |
| Median (range)            | 3.0 (0.3–16.0) | 4.5 (0.8–14.0) | 4.5 (0.5–14) | 6.0 (0.8–12) |
| Histologic grade          |            |            |           |            |
| Well-moderately           | 325        | 19.0       | 29        | 14.4       | 0.141  | 0.122 |
| Poorly                    | 1389       | 81.0       | 172       | 85.6       | 0.880  | 0.023 |
| T category                |            |            |           |            |
| T1                        | 521        | 30.4       | 2         | 1.0        | < 0.001 | 1.131 |
| T2                        | 255        | 14.9       | 11        | 5.5        | 0.868  | 0.022 |
| T3                        | 187        | 10.9       | 14        | 7.0        | 0.142  | 0.023 |
| T4a                       | 731        | 42.6       | 158       | 78.6       | 0.142  | 0.023 |
| T4b                       | 20         | 1.2        | 16        | 8.0        | 0.142  | 0.023 |
| N category                |            |            |           |            |
| N0                        | 790        | 46.1       | 16        | 8.0        | < 0.001 | 1.023 |
| N1                        | 308        | 18.0       | 36        | 17.9       | 0.868  | 0.022 |
| N2                        | 280        | 16.3       | 50        | 24.9       | 0.142  | 0.023 |
| N3a                       | 259        | 15.1       | 74        | 36.8       | 0.142  | 0.023 |
| N3b                       | 77         | 4.5        | 25        | 12.4       | 0.142  | 0.023 |
| TNM stage                 |            |            |           |            |
| I                         | 620        | 36.2       | 3         | 1.5        | < 0.001 | 1.131 |
| II                        | 398        | 23.2       | 22        | 10.9       | 0.868  | 0.022 |
| III                       | 696        | 40.6       | 176       | 87.6       | 0.868  | 0.022 |
| VELIPI                    |            |            |           |            |
| Negative                  | 996        | 58.1       | 54        | 26.9       | < 0.001 | 0.666 |
| Positive                  | 718        | 41.9       | 147       | 73.1       | 0.868  | 0.022 |
| No. of retrieved LNs      | < 15       | 331        | 19.3      | 36        | 17.9    | 0.702  | 0.036 |
| ≥ 15                      | 1383       | 80.7       | 165       | 82.1       | 0.702  | 0.036 |
| Mean (SD)                 | 21.3 ± 8.4 | 21.9 ± 8.4 | 21.9 ± 8.3 | 22.0 ± 8.5 | 0.868  | 0.022 |
| Median (range)            | 20 (2–74)  | 20 (8–62)  | 20 (5–55) | 20 (8–62)  | 0.868  | 0.022 |
| Adjuvant treatment        |            |            |           |            |
| No                        | 501        | 29.2       | 38        | 18.9       | < 0.001 | 0.388 |
| Single drug               | 304        | 17.7       | 40        | 19.9       | 0.868  | 0.022 |
| Double drugs              | 625        | 36.5       | 63        | 31.3       | 0.868  | 0.022 |
| Triple drugs              | 167        | 9.7        | 43        | 21.4       | 0.868  | 0.022 |
| CRT                       | 117        | 6.8        | 17        | 8.5        | 0.868  | 0.022 |

Abbreviations: CI confidence interval, GEJ gastroesophageal junction, LNs lymph nodes, PSM propensity score matching, SD standard difference, SMD standardized mean difference, TD tumor deposit, VELIPI vascular emboli, lymphatic, and perineural invasion
years), tumor location, tumor size (< 3.2 vs. ≥ 3.2, cm), histologic grade, T category, N category, VELIPI, adjuvant treatment, and TD status. After multivariate Cox proportional hazards model analysis, T category, N category, VELIPI, adjuvant treatment, and TD status remained independent prognostic factors for DFS, and age, T category, N category, adjuvant treatment, and TD status remained independent prognostic factors for OS (Table 4).

The 3-year DFS and OS rates of all enrolled patients were 76.2% (95% CI 74.3–78.2%) and 82.7% (95% CI 81.0–84.5%), respectively. There was a significant difference in the 3-year DFS rate between patients with TDs and those without (39.8% vs. 80.6%; HR: 1.75, 95% CI 1.42–2.14; P < 0.001; Fig. 1A). Additionally, the 3-year OS rate of the TD-present group was significantly lower than that of the TD-absent group (49.9% vs. 86.7%; HR: 1.93, 95% CI 1.57–2.38; P < 0.001; Fig. 1B).

In addition to TD status, the influence of TD number on prognosis was evaluated. This continuous variable was transformed into categorical variables by four different cutoff selections; however, no association with prognosis was found regardless of the cutoff selection (Supplementary Table 2).

### Table 2: Association of TD presence with clinicopathologic characteristics

| Variables | Univariate logistic regression | Multivariate logistic regression |
|-----------|-------------------------------|---------------------------------|
|           | Crude OR 95% CI | P value | Adjusted OR 95% CI | P value |
| Sex       | Male Ref. | Female 0.78 0.56–1.08 0.142 |       |       |
| Age (years) | < 58 Ref. | ≥ 58 1.81 1.35–2.46 < 0.001 | 1.48 1.07–2.05 0.018 |       |
| Tumor location | GEJ + Upper 1/3 Ref. | Middle 1/3 0.67 0.47–0.93 0.019 | 1.08 0.70–1.64 0.732 |       |
| Tumor size | < 3.2 cm Ref. | ≥ 3.2 cm 4.04 2.88–5.76 < 0.001 | 1.60 1.10–2.35 0.016 |       |
| Histologic grade | Well-moderately Ref. | Poorly 1.39 0.93–2.13 0.119 |       |       |
| T category | T1 Ref. | Ref. 11.24 2.99–72.96 0.002 | 6.34 1.65–41.65 0.018 |       |
|           | T2 | 19.50 5.38–124.96 < 0.001 | 6.96 1.82–45.84 0.013 |       |
|           | T4a | 56.31 17.89–341.41 < 0.001 | 17.78 4.86–105.80 < 0.001 |       |
|           | T4b | 208.40 54.60–1377.01 < 0.001 | 51.88 12.33–360.701 < 0.001 |       |
| N category | N0 Ref. | Ref. 5.77 3.21–10.83 < 0.001 | 3.10 1.68–5.94 < 0.001 |       |
|           | N1 | 8.82 5.05–16.21 < 0.001 | 3.56 1.98–6.73 < 0.001 |       |
|           | N2 | 14.11 8.29–25.49 < 0.001 | 4.99 2.83–9.31 < 0.001 |       |
|           | N3a | 16.03 8.28–31.89 < 0.001 | 5.03 2.50–10.41 < 0.001 |       |
|           | N3b | Ref. | Ref. |       |       |
| VELIPI | Negative Ref. | Positive 3.78 2.74–5.27 < 0.001 | 1.13 0.79–1.65 0.508 |       |

**Abbreviations:** CI confidence interval, GEJ gastroesophageal junction, VELIPI vascular emboli, lymphatic, and perineural invasion, TD tumor deposit, OR odds ratio, Ref. reference
Table 3  Univariate analyses for DFS and OS of GC patients

| Variables                        | Disease-free survival | Overall survival |
|---------------------------------|-----------------------|------------------|
|                                 | n                     | HR   | 95% CI     | P value | HR   | 95% CI     | P value |
| Overall                         | 1915                  |      |            |         |      |            |         |
| Sex                             |                       |      |            |         |      |            |         |
| Male                            | 1333                  | Ref. |            |         | Ref. |            |         |
| Female                          | 582                   | 0.93 | 0.77–1.11  | 0.400   | 0.87 | 0.72–1.06  | 0.161   |
| Age (years)                     |                       |      |            |         |      |            |         |
| < 58                            | 995                   | Ref. |            |         | Ref. |            |         |
| ≥ 58                            | 920                   | 1.40 | 1.19–1.65  | < 0.001 | 1.65 | 1.39–1.96  | < 0.001 |
| Tumor location                  |                       |      |            |         |      |            |         |
| GEJ + Upper 1/3                 | 499                   | Ref. |            |         | Ref. |            |         |
| Middle 1/3                      | 468                   | 0.82 | 0.66–1.01  | 0.063   | 0.81 | 0.65–1.00  | 0.053   |
| Lower 1/3                       | 948                   | 0.54 | 0.44–0.66  | < 0.001 | 0.55 | 0.45–0.67  | < 0.001 |
| Tumor size                      |                       |      |            |         |      |            |         |
| < 3.2 cm                        | 967                   | Ref. |            |         | Ref. |            |         |
| ≥ 3.2 cm                        | 948                   | 2.93 | 2.45–3.51  | < 0.001 | 3.05 | 2.53–3.68  | < 0.001 |
| Histologic grade                |                       |      |            |         |      |            |         |
| Well-moderately                 | 354                   | Ref. |            |         | Ref. |            |         |
| Poorly                          | 1516                  | 1.62 | 1.27–2.06  | < 0.001 | 1.51 | 1.19–1.92  | < 0.001 |
| T category                      |                       |      |            |         |      |            |         |
| T1                              | 523                   | Ref. |            |         | Ref. |            |         |
| T2                              | 266                   | 2.49 | 1.53–4.06  | < 0.001 | 2.62 | 1.56–4.41  | < 0.001 |
| T3                              | 201                   | 5.68 | 3.60–8.95  | < 0.001 | 6.11 | 3.77–9.92  | < 0.001 |
| T4a                             | 889                   | 11.93| 8.18–17.38 | < 0.001 | 13.05| 8.71–19.56 | < 0.001 |
| T4b                             | 36                    | 22.94| 13.34–39.45| < 0.001 | 28.20| 15.97–49.77| < 0.001 |
| N category                      |                       |      |            |         |      |            |         |
| N0                              | 806                   | Ref. |            |         | Ref. |            |         |
| N1                              | 344                   | 2.49 | 1.81–3.43  | < 0.001 | 2.68 | 1.94–3.71  | < 0.001 |
| N2                              | 330                   | 5.23 | 3.95–6.94  | < 0.001 | 5.08 | 3.79–6.82  | < 0.001 |
| N3a                             | 333                   | 10.23| 7.86–13.32 | < 0.001 | 10.34| 7.87–13.58 | < 0.001 |
| N3b                             | 102                   | 16.90| 12.30–23.22| < 0.001 | 18.88| 13.61–26.20| < 0.001 |
| VELIPI                          |                       |      |            |         |      |            |         |
| Negative                        | 1050                  | Ref. |            |         | Ref. |            |         |
| Positive                        | 865                   | 3.72 | 3.10–4.46  | < 0.001 | 3.43 | 2.86–4.12  | < 0.001 |
| No. of retrieved LNs            |                       |      |            |         |      |            |         |
| < 15                            | 367                   | Ref. |            |         | Ref. |            |         |
| ≥ 15                            | 1548                  | 1.22 | 0.98–1.51  | 0.078   | 1.17 | 0.93–1.45  | 0.175   |
| Adjuvant treatment              |                       |      |            |         |      |            |         |
| No                              | 539                   | Ref. |            |         | Ref. |            |         |
| Single drug                     | 344                   | 1.57 | 1.17–2.11  | 0.003   | 1.54 | 1.13–2.10  | 0.006   |
| Double drugs                    | 688                   | 2.13 | 1.66–2.71  | < 0.001 | 2.27 | 1.76–2.92  | < 0.001 |
| Triple drugs                    | 210                   | 3.24 | 2.44–4.31  | < 0.001 | 2.98 | 2.21–4.02  | < 0.001 |
| CRT                             | 134                   | 3.79 | 2.77–5.20  | < 0.001 | 3.58 | 2.58–4.97  | < 0.001 |
| TD                              |                       |      |            |         |      |            |         |
| Absent                          | 1714                  | Ref. |            |         | Ref. |            |         |
| Present                         | 201                   | 4.14 | 3.42–5.02  | < 0.001 | 4.45 | 3.66–5.42  | < 0.001 |

Abbreviations: CI confidence interval, DFS disease-free survival, GC gastric cancer, GEJ gastroesophageal junction, HR hazard ratio, LNs lymph nodes, OS overall survival, PSM propensity score matching, Ref. reference, TD tumor deposit, VELIPI vascular emboli, lymphatic, and perineural invasion
Propensity score-matching analysis
After 1:3 PSM, 191 TD-present patients and 491 TD-absent patients were obtained, and no significant differences were identified between the two groups in any of the baseline factors, which was demonstrated by both the P value and the SMD (Table 1). In the postmatched dataset, the DFS and OS of the TD-absent group were significantly longer than those of the TD-present group, which is consistent with the results of the survival analysis of the primary cohort. The median DFS was 87.9 months in the TD-absent group and 23.8 months in the TD-present group. The 3-year DFS rate was 61.5% in the TD-absent group and 40.8% in the TD-present group (HR 1.78, 95% CI 1.44–2.21; P < 0.001) (Fig. 1C).
The 3-year OS rate was 69.9% in the TD-absent group and 51.1% in the TD-present group (HR 1.92, 95% CI 1.54–2.39; P < 0.001) (Fig. 1D).

External validation of the previously proposed modified stage
The current primary cohort was utilized to compare the predictive capabilities of eight existing modified staging systems [16, 20, 21, 23, 26–28, 31] with that of the eighth AJCC TNM staging system (Table 5). Larger χ², AUC, and C-index values and a smaller AIC value indicate a better discriminative capability. Accordingly, the system proposed by Gu L. et al. [31] had the best performance. Among these systems, six had a better performance than the TNM staging system, except the system that suggests the presence of TD as T4a, which has a slightly worse performance than the 8th AJCC TNM staging system.

Fig. 1 Disease-free survival and overall survival between TD-absent and TD-present patients in the primary cohort (A, B) and after-PSM cohort (C, D). Abbreviations: PSM, propensity score matching; TD, tumor deposit
The incidence of TD in our cohort was 10.5%, which is similar to the previously reported range of the incidence rate (10.6–27.5%) [23, 26]. Moreover, we confirmed that the presence of TD was related to unfavorable clinicopathologic factors, reflecting a more progressive disease. In the primary cohort before PSM, the presence of TD was significantly associated with older age, distal GC, larger tumor size, advanced T category, N category, TNM stage, and VELIPI. Additionally, Lee H.S. et al. reported that TD-present status in the resection specimen was associated with the presence of synchronous distant metastasis [21]. Etoh T. et al. found that TD-present patients were more likely to present with peritoneal seeding at the time of surgery and develop peritoneal metastasis [17].

As TD is associated with unfavorable clinicopathologic factors, it is unsurprising that TD has been confirmed as an independent prognostic factor, together with T and N category, by many previous studies [17]. To estimate the prognostic influence of TD status on GC patients, univariate and multivariate Cox regression analyses were performed in the primary cohort, proving that TD was an independent prognostic factor for DFS and OS. Moreover, an after-PSM cohort was established by 1:3 PSM, and a negative impact of TD on DFS and OS still existed, which was consistent with the Cox regression results.

In addition to TD status (present vs. absent), which most studies evaluated, the evaluable parameters of TD also include the number of TDs and the categorization methods, the patterns of TD and the distribution area of TD. Wang W. et al. [20] transformed the number of TDs into categorical variables (0, 1, 2, and ≥ 3), and a correlation between this parameter and prognosis was found. Similar results were obtained by Sun Z. et al. [16] and Etoh T. et al. [17], but the transformation cutoff values were slightly different (1, 2–3, and ≥ 3 vs. 0, 1–4, and ≥ 5). However, both Anup S. et al. [23] and the current study did not find an association between the number of TDs and prognosis. Lee H.S. et al. [21] classified TD into five types (separate nodular, perivascular, perineural, lymphatic, and endovascular), but no association between TD types and prognosis in GC patients was found. The current study conducted a descriptive statistical analysis of the distribution areas of TD but did not correlate this variable with prognosis because there are various distribution areas of TD, and each patient may have more than one distribution area.

To date, the depth of tumor invasion (T category), nodal status (N category) and distant metastasis (M category),
which constitute the TNM stage used for prognostic prediction and guiding treatment, are the three most significant prognostic factors [30]. However, as previous and current studies have shown, the presence of TD is an indicator of poor prognosis in GC patients, and attempts have been made to incorporate TD into the TNM staging system. In this study, we evaluated seven previously presented proposals.

These proposals were suggested based on individual cohorts by comparing the prognosis of patients with TD with that of patients with different T or N categories to achieve the incorporation of TD. First, some proposals incorporated TD into T stage [16, 23]. Sun Z. et al. proposed that TD should be considered a form of serosal invasion (T4a), as they analyzed 2998 GC patients undergoing radical resection and found that no significant difference was observed between the prognosis of TD-present patients in the pT1-4a category and TD-absent patients in the pT4a category [16]. This finding was validated by another study in 2017 [23]. Second, suggestions that TD should be incorporated into the N category in different ways have been proposed by several studies [21, 26, 28, 32]. Kim et al. proposed that each TD should be treated as a positive lymph node, and a new N category was defined [21]. In contrast, Chen H. et al. proposed that the presence of TD should upstage the N category except for N3b [28]. Third, Wang W. et al. considered TD as a significant indicator and suggested incorporating TD into the N3 or M1 staging categories based on the number of retrieved TDs [20]. In addition, Etoh T. et al. suggested that TDs in GC may more closely resemble peritoneal metastasis than lymph node metastasis, but as there were long-term survivors with TD, they should be considered separately from peritoneal disease [17].

In addition to the different methods of integrating TD into the T, N or M categories, the status and number of TDs are another aspect to be considered. Some schemes considered only the status of TD (absent vs. present) [16, 23, 26–28]; some schemes counted one TD as a metastatic lymph node or a metastatic nodule [21]; and some schemes transformed the number of TDs into categorical variables [20]. In addition to the above considerations, most of the schemes did not change the division of the T and N categories, except Lee H.S. et al., who proposed a modified N category [21].

Based on the external validation results of the current dataset, the scheme proposed by Gu L. et al. [31] achieves the best performance. However, how to include TD in TNM staging remains unclear. First, the above seven proposals were presented based on their single-center databases, and external validation data were not provided. Unsurprisingly, the proposed systems performed well in

the internal validation. Second, the detection rate of TD in early T or N categories is very low, which is not conducive to evaluating the role of TD. Therefore, the extrapolation of the current existing schemes needs to be further validated. Additionally, a multicenter dataset with a large sample size is needed to verify these schemes or to create a more appropriate proposal.

Although the presence of TD was associated with unfavorable survival in GC patients, the survival rate and multivariate analysis results of our cohort suggest that patients with TD undergoing radical surgery combined with adjuvant therapy can obtain a satisfactory prognosis, indicating that en bloc clearance of adipose connective tissue by D2/R0 surgery is effective. Moreover, for patients with good performance and adherence, adjuvant treatment is highly recommended. Meanwhile, although many studies have confirmed that TD is an unfavorable clinicopathologic factor, adjuvant treatment is highly recommended. Meanwhile, although many studies have confirmed that TD is related to adverse clinicopathologic factors and that TD is an independent prognostic factor, further validations are still needed. Therefore, it is imperative to formulate corresponding standards so that TDs can be appropriately retrieved, examined and recorded.

The first limitation of this study is its retrospective nature. Only the number and anatomic location of the TD were recorded, and there were no data concerning TD patterns. Some variables, such as the gross type and Lauren’s classification of the tumor, were unavailable in some patients. Thus, these variables were not included in the analysis. Second, since this study focused on patients with locally advanced GC, it was not possible to compare the effect of the status and number of TDs on the prognosis of these patients with that of M1 patients. Therefore, this study evaluated only the impact of TD status on the prognosis of patients with different T and N categories. Third, all the patients were from a single center. Whether the results can be extrapolated to other patient groups needs to be further confirmed. Thus, prospective studies with larger sample sizes and more comprehensive information are needed to achieve more convincing conclusions.

Conclusions
In summary, the presence of TD was associated with unfavorable clinicopathologic factors, reflecting aggressive disease. Moreover, TD was identified as an independent unfavorable indicator of DFS and OS among radically resected GC patients. Incorporating TD into the TNM staging system can further improve the prognostic prediction accuracy, but the appropriate method of incorporation still needs to be explored and verified in prospective cohorts with larger sample sizes and more comprehensive information. In the future, corresponding standards must be formulated so that TDs can be appropriately retrieved, examined and recorded.
and the significance of TD in GC can be evaluated more comprehensively.

Abbreviations
AIC: Akaike information criterion; AJCC: American Joint Committee on Cancer; AUROC: area under the curve; C-index: concordance index; FUSCC: Fudan University Shanghai Cancer Center; GC: gastric cancer; IPTW: inverse probability of treatment weighting; OS: overall survival; PSM: propensity score matching; SMD: standardized mean difference; TD: tumor deposit; VELIP: Vascular emboli, lymphatic, and perineural invasion.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12957-022-02773-1.

Additional file 1: Supplementary Table 1. TD distribution areas and numbers. Abbreviations: TD, tumor deposit.

Additional file 2: Supplementary Table 2. Effect of number of TDs on OS in GC patients. Abbreviations: CI, confidence interval; DFS, disease-free survival; GC, gastric cancer; HR, hazard ratio; No., number; OS, overall survival; Pts, patients; Ref, reference; TD, tumor deposit.

Acknowledgements
The authors thank Dr. Cong Tan and Dr. Weimei Wang from the Department of Pathology for establishing the pathologic diagnosis. The authors thank American Journal Experts (English editor) for this editing of the submission.

Authors’ contributions
Fan Xia and Zhen Zhang contributed to the drafting, conception, and design of the study. Menglong Zhou, Wang Yang, Wei Zou, and Changming Zhou contributed to the manuscript writing and data analysis. Jia-ning Yang, Zhuyuan Zhang, Yaqi Wang, and Jing Zhang contributed to data collection. Yan Wang and Guichao Li contributed to patient follow-up. Fan Xia and Zhen Zhang contributed to supervision and paper revision. All authors helped to perform the research. The authors read and approved the final manuscript.

Funding
This study was supported by the National Natural Science Foundation of China (Grant No. 81773357, Grant No. 82003229) and Shanghai Anticancer Association EYAS Project (No. SACA-CY20B07; No. SACA-CY20C14). The funders had no role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Availability of data and materials
The datasets used during the current study are available from the corresponding author on reasonable request.

Declarations
Ethics approval and consent to participate
The retrospective study was approved by the Institutional Review Board of Fudan University Shanghai Cancer Center. The committee’s reference number is 1910208-11. The need for informed consent was waived because of the retrospective nature of the clinical study.

Consent for publication
Not applicable

Competing interests
The authors declare that they have no competing interests.

Author details
1 Department of Radiation Oncology, Fudan University Shanghai Cancer Center, 270 Dong’an Road, Shanghai 200032, People’s Republic of China. 2 Department of Oncology, Shanghai Medical College, Fudan University, Shanghai 200032, People’s Republic of China. 3 Shanghai Key Laboratory of Radiation Oncology, Shanghai 200032, People’s Republic of China. 4 Department of Cancer Prevention, Fudan University Shanghai Cancer Center, Shanghai 200032, People’s Republic of China.

Received: 24 April 2022 Accepted: 9 August 2022
Published online: 23 September 2022

References
1. Sung H, Ferlay J, Siegel RL, Laversanne M, Soerjomataram I, Jemal A, et al. Global cancer statistics 2020: GLOBOCAN estimates of incidence and mortality worldwide for 36 cancers in 185 countries. CA Cancer J Clin. 2021;71(3):209–49.
2. Hwang JE, Hong JY, Kim JE, Shim HJ, Bae WK, Hwang EC, et al. Prognostic significance of the coexistent existence of lymphovascular and perineural invasion in locally advanced gastric cancer patients who underwent curative gastrectomy and adjuvant chemotherapy. Jpn J Clin Oncol. 2015;45(6):541–6.
3. Blici A, Seker M, Ustaialooglu BB, Kefeli U, Yildirim E, Yavuzer D, et al. Prognostic significance of perineural invasion in patients with gastric cancer who underwent curative resection. Ann Surg Oncol. 2010;17(8):2037–44.
4. Peng F, Liu J, Wang F, Zheng G, Wang Q, Liu S, et al. Prognostic value of differentiation status in gastric cancer. BMC Cancer. 2018;18(1):865.
5. Veronesi N, Fassan M, Wood LD, Stubbs B, Solmi M, Capelli P, et al. Extracapsular extension of nodal metastases is a poor prognostic indicator in gastric cancer: a systematic review and meta-analysis. J Gastrointest Surg. 2016;20(10):1692–8.
6. Gabriel WB, Dukes C, Bussey HJR. Lymphatic spread in cancer of the rectum. Br J Surg. 1935;23(90):395–413.
7. Nagtegaal ID, Tot T, Jayne DG, McShane P, Nihlberg A, Marshall HC, et al. Lymph nodes, tumor deposits, and TNM: are we getting better? J Clin Oncol. 2011;29(18):2487–92.
8. Ono C, Yoshinaga K, Enomoto M, Sugihara K. Discontinuous rectal cancer spread in the mesorectum and the optimal distal clearance margin in situ. Dis Colon Rectum. 2002;45(6):744–9742.
9. Leno H, Mochizuki H, Shirozuki K, Kusumi T, Yamada K, Ikegami M, et al. Actual status of distribution and prognostic impact of extrarectal discontinuous cancer spread in colorectal cancer. J Clin Oncol. 2011;29(18):2550–6.
10. Shimada Y, Takii Y. Clinical impact of mesorectal extranodal cancer tissue in rectal cancer: detailed pathological assessment using whole-mount sections. Dis Colon Rectum. 2010;53(5):771–8.
11. Edge SB, Byrd DR, Carducci MA, Compton CC, Fritz AG, Greene FL, AJCC cancer staging manual. New York: Springer; 2010.
12. Mignano JE, Zaharakis ML, Chakravarty A, Pantadosi S, Dooley WC, Gage I. Significance of axillary lymph node extranodal soft tissue extension and indications for postmastectomy irradiation. Cancer. 1999;86(7):1258–62.
13. Sarogiou S, Akbulut N, Iplikci S, Aydin B, Dogan E, Unlu M, et al. Tumor deposits in head and neck carcinomas. Head Neck. 2016;38(Suppl 1):E256–60.
14. Lee YC, Wu CT, Kuo SW, Tseng YT, Chang YL. Significance of extranodal extension of regional lymph nodes in surgically resected non-small cell lung cancer. Chest. 2007;131(4):993–9.
15. Zhang HD, Tang R, Duan XF, Chen CG, Ma Z, Gao YY, et al. Extranodal metastasis is a powerful prognostic factor in patients with adenocarcinoma of the esophagogastric junction. J Surg Oncol. 2013;108(8):542–9.
16. Sun Z, Wang ZN, Xu YY, Zhu GL, Huang BJ, Xu Y, et al. Prognostic significance of tumor deposits in gastric cancer patients who underwent radical surgery. Surgery. 2012;151(6):871–81.
17. Etoh T, Sasaki M, Ishikawa K, Kata H, Sano T, Shimoda T. Extranodal metastasis is an indicator of poor prognosis in patients with gastric carcinoma. Br J Surg. 2006;93(3):369–73.
18. Puppia G, Ueno H, Kayahara M, Capelli P, Canzoneri V, Colombari R, et al. Tumor deposits are encountered in advanced colorectal cancer and other adenocarcinomas: an expanded classification with implications for colorectal cancer staging system including a unifying concept of intramyxoid metastases. Mod Pathol. 2009;22(3):410–5.
19. Tanaka T, Kumaqai K, Shimizu K, Masuo K, Yamagata K. Peritoneal metastasis in gastric cancer with particular reference to lymphatic advancement;
extranodal invasion is a significant risk factor for peritoneal metastasis. J Surg Oncol. 2000;73(3):165–71.

20. Wang W, Li Y, Zhang Y, Yuan X, Xu D, Guan Y, et al. Incorporation of extranodal metastasis of gastric carcinoma into the 7th edition UICC TNM staging system. PLoS One. 2011;6(6):e19657.

21. Lee HS, Lee HE, Yang HK, Kim WH. Perigastric tumor deposits in primary gastric cancer: implications for patient prognosis and staging. Ann Surg Oncol. 2013;20(5):1604–13.

22. Lee IS, Park YS, Ryu MH, Song MJ, Yook JH, Oh ST, et al. Impact of extranodal extension on prognosis in lymph node-positive gastric cancer. Br J Surg. 2014;101(12):1576–84.

23. Anup S, Lu J, Zheng CH, Li P, Xie JW, Wang JB, et al. Prognostic significance of perigastric tumor deposits in patients with primary gastric cancer. BMC Surg. 2017;17(1):84.

24. Paredes TO, Prado CS, Taxa RL, Luque-Vasquez C, Chavez I, Payet ME, et al. Clinicopathological factors associated with the presence of tumor deposits in resected gastric cancer patients. Helyon. 2021;7(6):e7185.

25. Wenquan L, Yuhua L, Jianxin C, Hongqing X, Kecheng Z, Jiyang L, et al. Tumor deposit serves as a prognostic marker in gastric cancer: a propensity score-matched analysis comparing survival outcomes. Cancer Med. 2020;9(10):3268–77.

26. Liang Y, Wu L, Liu L, Ding X, Wang X, Liu H, et al. Impact of extranodal tumor deposits on prognosis and N stage in gastric cancer. Surgery. 2019;166(3):305–13.

27. Tan J, Yang B, Xu Z, Zhou S, Chen Z, Huang J, et al. Tumor deposit indicates worse prognosis than metastatic lymph node in gastric cancer: a propensity score matching study. Ann Transl Med. 2019;7(22):671.

28. Chen H, Tang Z, Chen L, Li H, Wang X, Liu F, et al. Evaluation of the impact of tumor deposits on prognosis in gastric cancer and a proposal for their incorporation into the AJCC staging system. Eur J Surg Oncol. 2018;44(12):1990–6.

29. Association JGC. Japanese classification of gastric carcinoma: 3rd English edition. Gastric Cancer. 2011;14(2):101–12.

30. Amin MB, Edge SB, Greene FL, Byrd DR, Brookland RK, Washington MK, et al. AJCC Cancer Staging Manual. 8th ed. New York: Springer; 2017.

31. Gu L, Chen P, Su H, Li X, Zhu H, Wang X, et al. Clinical significance of tumor deposits in gastric cancer: a retrospective and propensity score-matched study at two institutions. J Gastrointest Surg. 2020;24(11):2482–90.

32. Japanese gastric cancer treatment guidelines 2010 (ver. 3). Jpn Gastric Cancer Assoc. 2011;14(2):113–23.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Ready to submit your research? Choose BMC and benefit from:

- fast, convenient online submission
- thorough peer review by experienced researchers in your field
- rapid publication on acceptance
- support for research data, including large and complex data types
- gold Open Access which fosters wider collaboration and increased citations
- maximum visibility for your research: over 100M website views per year

At BMC, research is always in progress.
Learn more biomedcentral.com/submissions