ORIGINAL RESEARCH

### Diabetes mellitus predicts inferior survival in diffuse large B-cell lymphoma: a propensity score-matched analysis

This article was published in the following Dove Press journal: Cancer Management and Research

Rui Gao<sup>1</sup> Jin-Hua Liang<sup>2</sup> Tian-Shuo Man<sup>2</sup> Li Wang<sup>2</sup> Hua-Yuan Zhu<sup>2</sup> Wei Wu<sup>2</sup> Lei Fan<sup>2</sup> Jian-Yong Li<sup>2</sup> Tao Yang<sup>1,\*</sup> Wei Xu<sup>2,\*</sup>

<sup>1</sup>Department of Endocrinology, The First Affiliated Hospital of Nanjing Medical University, Jiangsu Province Hospital, Nanjing 210029, People's Republic of China; <sup>2</sup>Department of Hematology, The First Affiliated Hospital of Nanjing Medical University, Jiangsu Province Hospital, Collaborative Innovation Center for Cancer Personalized Medicine, Nanjing 210029, People's Republic of China

\*These authors contributed equally to this work

Correspondence: Tao Yang Department of Endocrinology, The First Affiliated Hospital of Nanjing Medical University, Jiangsu Province Hospital, 300 Guangzhou Road, Nanjing 210029, People's Republic of China, Tel +86 256 830 6466 Fax +86 256 830 6466 Email yangt@njmu.edu.cn

Wei Xu Department of Hematology, The First Affiliated Hospital of Nanjing Medical University, Jiangsu Province Hospital, 300 Guangzhou Road, Nanjing 210029, People's Republic of China, Tel +86 258 378 1120 Fax +86 258 378 1120 Email xuwei10000@hotmail.com



**Purpose:** Diabetes mellitus (DM) is associated with elevated cancer risk and poor survival. The objective of this study was to assess the prognostic value of DM in diffuse large B-cell lymphoma (DLBCL).

**Methods:** Five hundred and fifty-three newly diagnosed DLBCL patients whose treatments included rituximab were recruited. Propensity score-matched method was performed to balance baseline characteristics and eliminate possible bias. Multivariate Cox regression analyses screened the prognostic risk factors in relation to progression-free survival (PFS) and cancer-specific survival (CSS). Receiver-operator characteristic curves and the corresponding areas under the curve (AUC) assessed the predictive accuracy of international prognostic index (IPI) together with DM.

**Results:** One hundred and nine patients (19.71%) had pre-existing DM. In the propensitymatched cohort, DM was associated with unfavorable PFS and CSS in rituximab era, and it was an independent risk factor for both inferior PFS and CSS, especially in patients with age  $\leq 60$  years, IPI 0–2, B symptoms and lactate dehydrogenase  $\leq$ upper limit of normal. Prediabetics also demonstrated inferior prognostic outcomes compared to patients with no diabetic tendency. DM as one additional point to IPI had larger AUC compared with IPI alone in CSS prediction and could improve the prognostic capacity of IPI.

**Conclusion:** The results indicate that preexisting DM is an important risk factor for survival. It could help predict life expectancy and build refined prognostication models for DLBCL.

**Keywords:** diffuse large B-cell lymphoma, diabetes mellitus, progression-free survival, cancer-specific survival, propensity score-matched analysis

### Introduction

Diabetes and malignancy are multifactorial heterogeneous diseases, and both have witnessed a rapid increase in prevalence owing to environmental and lifestyle changes. Accumulating evidence suggested that diabetes mellitus (DM) was correlated with elevated cancer risks, including liver,<sup>1</sup> pancreatic,<sup>2</sup> colorectal<sup>3</sup> and breast cancer.<sup>4</sup> Concerning non-Hodgkin lymphoma (NHL), three meta-analyses of observational studies reported that the risk ratio (RR) of developing NHL in patients with diabetes was 1.19 (95% CI: 1.04–1.35),<sup>5</sup> 1.22 (95% CI: 1.07–1.39)<sup>6</sup> and 1.79 (95% CI: 1.30–2.47),<sup>7</sup> respectively. Possible underlying biological mechanisms of diabetes-promoting carcinogenesis were discovered, such as hyperglycemia, hyperinsulinemia and chronic inflammation.

Cancer Management and Research 2019:11 2849-2870

© 2019 Gao et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/terms.php and incorporate the Creative Commons Attribution – Non Commercial (unported, v3.0) License.(http://creativecommons.org/licenses/by-mc/3.0/). By accessing the work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 42 and 5 d our Terms (https://www.dovepress.com/terms.php). Diffuse large B-cell lymphoma (DLBCL), the most common lymphoid malignancy in adults, is characterized by wide clinical phenotypes and molecular heterogeneities.<sup>8,9</sup> Although the introduction of rituximab to CHOP (cyclophosphamide, doxorubicin, vincristine and prednisone) reformed the treatment of DLBCL, the long-term survival of one-third patients is still poor due to relapse and refractory after initial chemoimmunotherapies.<sup>10</sup> The international prognostic index (IPI), which consists of five clinical parameters, is the current standard prognostication system for DLBCL. However, it was proven insufficient in predicting the prognosis of highrisk patients in rituximab era<sup>11</sup> and revealing the underlying biological differences. Therefore, novel prognostic factors are urgently required in identifying high-risk DLBCL patients and facilitating treatments in personalized medicine.

The evidence regarding the prognostic impact of DM on lymphoma remains limited and inconclusive. No literature to date has focused on the correlation between DLBCL and DM, and the prognostic value of DM in DLBCL. To address this issue, our study indicated preexisting DM as a predictor of inferior prognosis in DLBCL by propensity score-matched (PSM) analyses. Considering the heterogeneity of different NHL subtypes, focusing on DLBCL would present more accurate and specific results.

### **Methods**

### Ethics approval and consent to participate

This study was approved by the Ethics Committee of the First Affiliated Hospital of Nanjing Medical University. All aspects of the study, including measurements for glucose levels and other periodical clinical and laboratorial checkups, were performed according to the principles of the Declaration of Helsinki (64th, 2013). Written informed consent and permissions for the possibility of utilizing their clinical data anonymously in the future were obtained from all subjects involved in this study at the time of first DLBCL admission.

#### Patients

Six hundred and twenty-eight consecutive subjects with newly diagnosed DLBCL between January 2008 and November 2016 from the First Affiliated Hospital of Nanjing Medical University were retrospectively enrolled. Seventy-five patients were excluded for not in accordance with the following criteria: 1) with complete clinical information and follow-up data; 2) without previous malignancies; 3) negative for HIV; 4) received chemoimmunotherapies as first-line treatment; 5) patients having dose reduction in chemoimmunotherapies (cyclophosphamide and anthracycline) owing to blood urea nitrogen (BUN) or creatinine (SCr) levels at diagnosis more than 1.5-fold upper limit of normal, or leftventricular ejection fraction below 55% were excluded to avoid confounding effect; 6) patients who died of accident or other medical conditions with no relation to DLBCL were excluded in order to analyze the prognostic value of DM on DLBCL-specific deaths. The flowchart of the study population is presented in Figure S1.

#### Data collection

All DLBCL patients were managed and treated in the inpatient department, while the follow-up examinations were conducted in the outpatient department. Baseline clinical characteristics, which consist of gender, age, IPI, Ann Arbor stage (I-IV), Eastern Cooperative Oncology Group (ECOG) performance status (PS), the number of extranodal sites, Hans classification, symptoms, previous cancer diagnosis, lymphoma treatments and response to therapy were available from medical records and hospital registries. Laboratorial and imageological data including blood biochemical examination (lactate dehydrogenase [LDH], albumin, etc.), renal function test (BUN, SCr, etc.), etiological test (hepatitis B virus, hepatitis C virus, treponema pallidum and HIV) and positron emission tomography-computed tomography (PET-CT) and echocardiography were routinely performed on every patient upon first DLBCL admission and were accessible from the hospital-based laboratorial and imageological service.

Among these 553 cases, 404 (73.06%) were treated with R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine and prednisone), 118 (21.34%) with dose-adjusted (DA)-EPOCH (etoposide, prednisone, vincristine, cyclophosphamide and doxorubicin) plus rituximab, 22 (3.98%) with R-CHOP like regimens including R-CDOP (rituximab, cyclophosphamide, pegylated liposomal doxorubicin, vincristine and prednisone) and R-CEOP (rituximab, cyclophosphamide, epirubicine, vincristine and prednisone) and 9 (1.63%) with R-mini CHOP (rituximab combined with low-dose CHOP). With regard to the relative dose intensity of chemotherapies, both diabetics and nondiabetics received standard treatments without delay or dose reduction (including steroid) due to hyperglycemia or other diabetic complications. The dose of rituximab is 375 mg/m<sup>2</sup> in all rituximab-based chemoimmunotherapies. In both groups for CHOP, the dose intensity is cyclophosphamide 750  $mg/m^2$ , doxorubicin 50 mg/m<sup>2</sup> and vincristine 1.4 mg/m<sup>2</sup>

(maximum 2 mg/m<sup>2</sup>) on day 1 and prednisone 60 mg/m<sup>2</sup> on days 1-5. EPOCH is composed of etoposide 50 mg/m<sup>2</sup>, vincristine 0.4 mg/m<sup>2</sup> and doxorubicin 10 mg/m<sup>2</sup> continuous infusion on days 1-4, cyclophosphamide 750 mg/m<sup>2</sup> on day 5 and prednisone 60  $mg/m^2$  on days 1–5. For CDOP, the specific dose is cyclophosphamide 600 mg/m<sup>2</sup>, pegylated liposomal doxorubicin 30 mg/m<sup>2</sup> and vincristine 1.4  $mg/m^2$  (maximum 2  $mg/m^2$ ) on day 1 and prednisone 60  $mg/m^2$  on days 1–5, while for CEOP the specific dose is cyclophosphamide 750 mg/m<sup>2</sup>, epirubicine 70 mg/m<sup>2</sup> and vincristine 1.4 mg/m<sup>2</sup> (maximum 2 mg/m<sup>2</sup>) on day 1 and prednisone 60 mg/m<sup>2</sup> on days 1-5. Moreover, the combination of mini-CHOP includes cyclophosphamide 400 mg/m<sup>2</sup>, doxorubicin 25 mg/m<sup>2</sup> and vincristine 1 mg on day 1,and prednisone 40 mg/m<sup>2</sup> on day 1-5. The numbers of patients according to diabetic status distributed by treatment types are presented in Table 1.

The final evaluation of response was performed after 6–8 cycles of chemotherapies, and lymphoma response was classified according to the International Workshop Criteria.<sup>12</sup> Diabetic and nondiabetic patients were given the comparable number of treatment cycles. Their antidiabetic medications before DLBCL admission were still used as basic glycemic control during the entire hospitalization. Insulin was also applied for poorly controlled hyperglycemic patients.

### Definition of preexisting DM and measurement for plasma glucose level

Preexisting DM referred to patients having any one of the following characteristics at diagnosis of DLBCL: diagnosis of DM from medical records of previous outpatient visit or hospitalization (based on the International Statistical Classification of Diseases and Related Health Problems, 10th revision (ICD-10) code E10, E11 and E14<sup>13</sup> or antidiabetic prescriptions); fasting plasma glucose (FPG) level  $\geq$ 7.0 mmol/l (126 mg/dl) at first hospital admission for DLBCL before administration of prednisone. Diabetes newly identified during the DLBCL follow-up period was ignored. Prediabetes was defined as having medical histories of impaired fasting glucose (IFG) or impaired glucose tolerance (IGT) (ICD-10 code R73.0) or FPG  $\geq$ 6.1 mmol/l (110 mg/dl) and <7.0 mmol/l (126 mg/dl) at DLBCL diagnosis. Diabetes, IFG and IGT were diagnosed according to the criteria established by the WHO. Diabetic duration calculated from the earliest applicable diagnosis of DM to the first DLBCL admission and antidiabetic treatments at enrollment were also collected from medical records and hospital registries.

FPG results were accessible from the hospital-based laboratorial service for all patients. Venous blood samples were collected between 7:00 am and 9:00 am after overnight fasting on the second day after the first admission. Within 2 h after blood sample collection, plasma glucose level was measured on an automatic enzymatic analyzer (Beckman Coulter, Brea, CA, USA) by means of glucose oxidase or hexokinase method under a stringent quality control mechanism.

### Follow-up and outcome measures

The follow-up events included progression-free survival (PFS) and cancer-specific survival (CSS). PFS was defined as the date of initial diagnosis to the date of disease progression or death caused by lymphoma or treatment toxicity. Disease progression was defined and evaluated according to nodal masses (appearance of a new lesion(s) >1.5 cm in any axis,  $\geq$ 50% increase in sum of the product of the diameters (SPD) of more than one node, or  $\geq$ 50% increase in longest diameter of a previously identified node >1 cm in short axis, lesions positron emission tomography (PET) positive if fluorodeoxyglucose (FDG)-avid lymphoma or PET positive prior to therapy), spleen and liver (>50% increase from nadir in the SPD of any previous lesions) and bone marrow (new or recurrent involvement).<sup>12</sup> For the accuracy of the standardized uptake value (SUV) measurement, the patient's blood glucose level was required to be controlled <150 mg/dL before receiving intravenous <sup>18</sup>F-FDG injections, otherwise rescheduling the PET-CT. Most cases of disease progression which were accessible by biopsy considering the location and operational risk were further pathologically confirmed. CSS was calculated as the interval between diagnosis and DLBCL-specific death or end of the follow-up (June 2017). Cause of death coded as C83.3 based on ICD-10 was classified as DLBCL-specific death.<sup>13</sup> Eleven deaths resulting from other causes, including four pneumonia (ICD-10 code J12.9 and J15.9), three lung cancers (ICD-10 code C34.9), three ischemic heart diseases (ICD-10 code I25.9) and one stroke (ICD-10 code I64) were excluded from the analyses. The follow-up included clinical and laboratorial checkups every 3 months for the first year and every 6 months from the 2nd to the 5th years at outpatient department. The patients were also required to be evaluated as soon as possible when having diseaserelated symptoms or signs of organ involvement. Specially, the examinations consisted of physical

matching
propensity
after J
and
before
status
diabetic
ording to
acc
haracteristics:
cal o
Clinic
e
Tabl

Table I Clinical chara	cteristics according to c	diabetic sta	itus before and after propen	ısity matching <sup>a</sup>					
Clinical parameter	S	Unmat	ched (complete) dataset			Propen	sity score-matched (1:1)	dataset <sup>b</sup>	
		Total	Diabetic (N=109) (%)	Nondiabetic (N=444)(%)	p-value	Total	Diabetic (N=109) (%)	Non-diabetic (N=109)(%)	p-value
Gender	Male Female	293 260	63 (21.50) 46 (17.69)	230 (78.50) 214 (82.31)	0.285	125 93	63 (50.40) 46 (49.46)	62 (49.60) 47 (50.54)	000.1
Age	≤60 years >60 years	324 229	42 (12.96) 67 (29.26)	282 (87.04) 162 (70.74)	100.0>	83 135	42 (50.60) 67 (49.63)	41 (49.40) 68 (50.37)	000.1
Idi	0-2 3-5	38I 172	67 (17.59) 42 (24.42)	314 (82.41) 130 (75.58)	0.065	138 80	67 (48.55) 42 (52.50)	71 (51.45) 38 (47.50)	0.673
Stage	1-11 111-1V	209 344	38 (18.18) 71 (20.64)	171 (81.82) 273 (79.36)	0.510	77 141	38 (49.35) 71 (50.35)	39 (50.65) 70 (49.65)	000.1
ECOG PS	0-1 2-4	471 82	87 (18.47) 22 (26.83)	384 (81.53) 60 (73.17)	0.097	180 38	87 (48.33) 22 (57.89)	93 (51.67) 16 (42.11)	0.372
Extranodal site(s)	⊽  ≺	403 150	78 (19.35) 31 (20.67)	325 (80.65) 119 (79.33)	0.720	159 59	78 (49.06) 31 (52.54)	81 (50.94) 28 (47.46)	0.761
Hans classification	GCB Non-GCB	211 342	37 (17.54) 72 (21.05)	174 (82.46) 270 (78.95)	0.324	71 147	37 (52.11) 72 (48.98)	34 (47.89) 75 (51.02)	0.773
Symptoms	No B symptoms B symptoms	342 211	64 (18.71) 45 (21.33)	278 (81.29) 166 (78.67)	0.509	132 86	64 (48.48) 45 (52.33)	68 (51.52) 41 (47.67)	0.678
ГДН	>∩LN (271U/L) ≤ULN (271U/L)	351 202	63 (17.95) 46 (22.77)	288 (82.05) 156 <i>(77</i> .23)	0.183	131 87	63 (48.09) 46 (52.87)	68 (51.91) 41 (47.13)	0.580
Albumin	<lln (40u="" l)<br="">≥LLN (40U/L)</lln>	332 221	69 (20.78) 40 (18.10)	263 (79.22) 181 (81.90)	0.448	128 90	69 (53.91) 40 (44.44)	59 (46.09) 50 (55.56)	0.216
Treatments	R-CHOP DA-EPOCH+R R-CHOP like regimens R-mini CHOP	404 118 22 9	80 (19.80) 21 (17.80) 5 (22.73) 3 (33.33)	324 (80.20) 97 (82.20) 17 (77.27) 6 (66.67)	0.692	164 40 9 5	80 (48.78) 21 (52.50) 5 (55.56) 3 (60.00)	84 (51.22) 19 (47.50) 4 (44.44) 2 (40.00)	0.917
Response to therapy <sup>c</sup>	CR or PR Non-CR or PR	392 138	70 (17.86) 30 (21.74)	322 (82.14) 108 (78.26)	0.314	147 58	70 (47.62) 30 (51.72)	77 (52.38) 28 ( <del>4</del> 8.28)	0.643
Notes: <sup>a</sup> The tests used wer- ECOG PS, extranodal site(s), unmatched dataset and 13 in <b>Abbreviations</b> : IPI, internat normal: R-CHOP, rituximab, combined with low-dose CH	e the Chi-square test or the Fi. Hans classification, symptoms ; propensity score-matched dat cional prognostic index; ECOG cyclophosphamide, doxorubici tOP; CR, complete remission;	sher's exact t and LDH leve taset. , Eastern Coo n, vincristine PR, partial re	est. <sup>b</sup> Propensity score-matched (I:1, <sup>c</sup> The evaluation of treatment respo operative Oncology Group; PS, perfe and prednisone; DA-EPOCH+R, dos mission.	, 109 pairs) analyses we anse was performed aftu ormance status; GCB, , se-adjusted (etoposide,	re performed er 6–8 cycles of germinal centei prednisone, vi	with a small chemothera c B-cell; LDH ncristine, cyc	aliper of 0.1 for matching potential pies. Patients who died during the th (, lactate dehydrogenase; ULN, uppe lophosphamide and doxorubicin) pli	confounders including : nerapies were excluded, er limit of normal; LLN, us rituximab; R-mini CF	iex, age, stage, including 23 in lower limit of IOP, rituximab

examination, a panel of blood tests, enhancement CT or PET-CT and bone marrow assessment if necessary. The medical staff in our center would contact the patients or their families for the reminding of scheduled follow-up checkups. Diabetic patients and nondiabetics were treated unbiasedly with uniformity of reassessment. For those who survived >5 years, the follow-up and mortality data were carefully retrieved from hospital records, death certificates in local disease control center or by interviewing (directly or by telephone) the patients, their families and personal physicians annually. The follow-up data were available for 553 patients finally enrolled, while two patients lost to follow-up were excluded from the study population.

#### Statistical analyses

All statistical analyses were performed using SPSS 23.0 (SPSS, Chicago, IL, USA), Graphpad Prism 6.01 (La Jolla, CA, USA) and R software 3.2.5 (http://www.r-project.org/) with MatchIt packages. Categorical variables were shown in percentage (%) and compared by the Chi-square test or the Fisher's exact test. Continuous variables were displayed as mean  $\pm$  standard deviation and determined as to whether the two populations were normally distributed using the Kolmogorov-Smirnov test or the Shapiro-Wilk test, and whether they had homogeneity of variances using the Levene's test. For data following normal distribution and having homogeneity of variances, results were compared by the Student's t-test. While for data following non-normal distribution or having heterogeneity of variances, the Mann-Whitney U test or the Kruskal-Wallis test was applied. Kaplan-Meier method was used to establish survival curves. The log-rank test was performed in subgroup analyses for each stratum investigating the interaction between clinical parameters and DM, and also in pairwise comparisons by pairwise over strata evaluating the difference between every two specific groups. Multivariate Cox regression analyses screened the independent prognostic factors for PFS and CSS, and variance inflation factor (VIF) was calculated to rule out multicollinearity between the included parameters. PSM analyses, using the 1:1 nearest neighbor technique with a small caliper of 0.1 were performed. Factors including sex, age, stage, ECOG PS, extranodal site(s), Hans classification, symptoms and LDH level were balanced to re-evaluate univariate and multivariate analyses in the matched couples. Receiver-operator characteristic curves (ROCs) and the corresponding areas under the curve (AUC) assessed the predictive accuracy of IPI and DM. The difference of AUCs was tested by a

nonparametric approach developed by DeLong et al<sup>14</sup>. Difference was considered significant with a two-sided *p*-value < 0.05.

### Results

## Patients' clinical characteristics in relation to diabetes

Five hundred and fifty-three newly diagnosed DLBCL subjects whose treatments included rituximab were retrospectively recruited. In total, 109 (19.71%) patients had preexisting DM, with 106 patients having a medical history of DM. Elevated FPG ( $\geq$ 7.0 mmol/l) was found in the other 3 patients at DLBCL diagnosis. Additionally, 43 (7.78%) patients had prediabetes, including 10 patients having a medical history of IFG or IGT, and the other 33 whose FPG level  $\geq$ 6.1 mmol/l and <7.0 mmol/l at enrollment. The mean FPG levels for diabetes and prediabetes were 7.235±2.115 mmol/l and 6.501±0.247 mmol/l, respectively. The median diabetic duration was 48 months (range, 0–360 months).

The baseline characteristics according to diabetic status are summarized in Table 1. In the complete dataset, DLBCL patients whose age >60 years were more likely to have comorbid DM (p<0.001). However, only a tendency toward significance was identified concerning IPI and ECOG PS that diabetic patients were more inclined to have high IPI and poor ECOG PS (p=0.065 and p=0.097, respectively). Moreover, no significant difference was detected in the distribution of DLBCL treatments (p=0.692) and response to therapy (p=0.314), and thus diabetic and nondiabetic patients received relatively comparable treatments.

To balance the characteristics between diabetics and nondiabetics, PSM analyses with 1:1 ratio were applied to minimize the differences in potential confounders including sex, age, stage, ECOG PS, extranodal site(s), Hans classification, symptoms and LDH level. After matching, the mean propensity scores for diabetic and nondiabetic groups were  $0.504\pm0.048$  and  $0.496\pm0.042$  (p=0.313). These clinicopathological parameters were adequately balanced and evenly distributed in the propensity-matched dataset as shown in Table 1 (all p>0.2).

## Prognostic value of preexisting DM in DLBCL

The last follow-up results of all 553 patients enrolled in our study were obtained in June 2017 with a median follow-up of 43 months (range, 7–114 months), during which 127 (22.97%) patients died, and 192 (34.72%) patients experienced disease progression. Event rates of PFS and CSS in subgroups stratified by different clinical parameters are displayed in Table S1.

The median survival time of the whole cohort was not reached for both PFS (range, 0–113 months) and CSS (range, 1–114 months) (Table S2). As for diabetes, a significant difference in PFS was identified, with median PFS of 56 months (range, 1–91 months) for diabetic group, while not reached (range, 0–113 months) for nondiabetics (p=0.001). Worse CSS was also observed in patients with DM (p<0.001). Although the median CSS was not reached for both groups, the 5-year CSS rate of diabetics (50.8%) (range, 1–91 months) was evidently lower compared with nondiabetics (77.8%) (range, 2–114 months) (Figure 1A and B).

Table 2 indicates the Cox regression analyses of prognostic factors for PFS and CSS (VIF<10 for both PFS and CSS). All the variables, regardless of significance in univariate analyses, further entered multivariate analyses to avoid confounding effect. For PFS of the complete cohort, five variables including age (>60 years) (HR=1.426; 95% CI: 1.058–1.920; p=0.020), advanced stage (III-IV) (HR=2.158; 95% CI: 1.437-3.241; p<0.001), Hans classification of non-GCB (HR=1.666; 95% CI: 1.208-2.298; p=0.002), LDH level (>ULN) (HR=2.041; 95% CI: 1.497 -2.782; p<0.001) and diabetic (HR=1.452; 95% CI: 1.033 -2.039; p=0.032) remained significant. For CSS, six variables were selected: age (>60 years) (HR=1.484; 95% CI: 1.027–2.143; *p*=0.036), advanced stage (III–IV) (HR=2.342; 95% CI: 1.390-3.946; p=0.001), ECOG PS (2-4) (HR=1.885; 95% CI: 1.241-2.861; p=0.003), Hans classification of non-GCB (HR=1.646; 95% CI: 1.106 -2.451; p=0.014), LDH level (>ULN) (HR=2.157; 95%) CI: 1.463-3.181; p<0.001) and diabetic (HR=2.267; 95% CI: 1.542-3.334; p<0.001). Consequently, DM was an independent risk predictor for both PFS and CSS.

Results observed in the complete cohort were corroborated using the PSM dataset. As exhibited in Kaplan– Meier survival curves for the matched groups (Figure 1C





Figure I Kaplan-Meier curves of PFS and CSS stratified by diabetic status before and after propensity matching. (A) PFS in unmatched (complete) dataset. (B) CSS in unmatched (complete) dataset. (C) PFS in propensity score-matched (1:1) dataset. (D) CSS in propensity score-matched (1:1) dataset. (Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival; DM, diabetes mellitus.

Clinic	al parameter	Unmatched (comp	lete) dataset			Propensity score-m	atched (I:I)	dataset <sup>a</sup>	
		Univariate analyse	s	Multivariate analys	es	Univariate analyses		Multivariate analys	es
		HR (95% CI)	p-value	HR (95% CI)	p-value	HR (95% CI)	p-value	HR (95% CI)	p-value
PFS	Male	1.169 (0.879–1.556)	0.284	1.010 (0.756–1.349)	0.946	0.776 (0.508–1.187)	0.243	0.703 (0.454–1.090)	0.115
	Age >60 years	1.678 (1.263–2.228)	<0.001	1.426 (1.058–1.920)	0.020	1.270 (0.811–1.987)	0.296	1.244 (0.787–1.967)	0.351
	Stage III–IV	3.311 (2.285–4.797)	<0.001	2.158 (1.437–3.241)	<0.001	2.717 (1.577–4.681)	<0.001	1.785 (0.963–3.308)	0.066
	ECOG PS 2-4	2.088 (1.486–2.935)	<0.001	1.419 (0.992–2.031)	0.055	1.763 (1.067–2.914)	0.027	1.334 (0.780–2.281)	0.293
	Extranodal sites >	1.897 (1.415–2.545)	<0.001	1.181 (0.860–1.620)	0.304	2.280 (1.476–3.522)	<0.001	1.723 (1.042–2.851)	0.034
	Non-GCB	1.904 (1.384–2.619)	<0.001	1.666 (1.208–2.298)	0.002	1.601 (0.985–2.603)	0.058	1.850 (1.125–3.044)	0.015
	B symptoms	1.912 (1.440–2.538)	<0.001	1.239 (0.912–1.682)	0.171	1.673 (1.094–2.557)	0.017	1.080 (0.655–1.783)	0.763
	LDH >ULN (271U/L)	3.041 (2.282-4.052)	<0.001	2.041 (1.497–2.782)	<0.001	2.459 (1.604–3.770)	<0.001	1.820 (1.090–3.039)	0.022
	Diabetic	I.744 (I.259–2.417)	0.001	1.452 (1.033–2.039)	0.032	1.607 (1.045–2.470)	0.03	1.665 (1.074–2.581)	0.023
CSS	Male	0.948 (0.669–1.344)	0.764	0.821 (0.577–1.169)	0.274	0.721 (0.448–1.161)	0.179	0.662 (0.408–1.073)	0.094
	Age ≥60 years	1.867 (1.316–2.647)	<0.001	1.484 (1.027–2.143)	0.036	1.388 (0.835–2.308)	0.206	1.308 (0.770–2.223)	0.320
	Stage III–IV	3.443 (2.135–5.551)	<0.001	2.342 (1.390–3.946)	0.00 I	2.676 (1.433–4.998)	0.002	1.727 (0.844–3.533)	0.135
	ECOG PS 2-4	2.730 (1.839-4.053)	<0.001	1.885 (1.241–2.861)	0.003	2.178 (1.266–3.746)	0.005	1.641 (0.915–2.942)	0.096
	Extranodal sites >	1.976 1.382–2.825)	<0.001	1.175 (0.799–1.727)	0.413	2.742 (1.698–4.429)	<0.001	1.908 (1.099–3.313)	0.022
	Non-GCB	1.896 (1.280–2.810)	0.001	1.646 (1.106–2.451)	0.014	1.816 (1.036–3.184)	0.037	1.995 (1.114–3.574)	0.020
	B symptoms	1.757 (1.241–2.489)	0.002	1.054 (0.722–1.537)	0.786	1.539 (0.956–2.477)	0.076	0.943 (0.540–1.648)	0.837
	LDH >ULN (271U/L)	3.341 (2.336–4.778)	<0.001	2.157 (1.463–3.181)	<0.001	2.621 (1.616-4.251)	<0.001	1.838 (1.028–3.286)	0.040
	Diabetic	2.713 (1.874–3.928)	<0.001	2.267 (1.542–3.334)	<0.001	2.145 (1.306–3.523)	0.003	2.057 (1.243–3.404)	0.005
Note: <sup>a</sup> P level. <i>p</i> -va Abbrevis	ropensity score-matched (1:1, 10 <sup>5</sup> lues in bold < 0.05. <b>utions:</b> PFS, progression-free sur <sup>-</sup>	9 pairs) analyses were perform vival; CSS, cancer-specific surv	ed with a small cali ival; ; ECOG, East	iper of 0.1 for matching poten ern Cooperative Oncology G	itial confounders ir iroup; PS, perform	ncluding sex, age, stage, ECOG 1ance status; GCB germinal ce	i PS, extranodal sit enter B-cell; LDH,	:e(s), Hans classification, symp lactate dehydrogenase; ULN,	toms and LDH upper limit of
normal.									

matching sitv and afte and a CSC hardene ï ĉ multivariate pue Table 2 Univariate and D), patients with DM demonstrated significantly poorer PFS (p=0.031) and CSS (p=0.003) compared with nondiabetics. Multivariate Cox regression analyses after PSM showed similar results that DM still remained as an independent prognostic factor for both PFS (HR=1.665; 95% CI: 1.074–2.581; p=0.023) and CSS (HR=2.057; 95% CI: 1.243–3.404; p=0.005).

Owing to the fact that excluding information from future events would generate selection bias, we analyzed the association between DM and all-cause mortality by censoring individuals at the time of death due to other causes rather than excluding them. Univariate and multivariate Cox regression analyses of PFS and overall survival (OS) were conducted in 564 patients including the 11 who died of other causes (Figure S2 and Table S3). DM was an independent factor correlated with worse PFS and OS both in the complete (HR=1.409; 95% CI: 1.012 -1.961; p=0.042 for PFS and HR=2.077; 95% CI: 1.432 -3.013; p<0.001 for CSS) and PSM datasets (HR=1.769; 95% CI: 1.144–2.735; p=0.010 for PFS and HR=2.134; 95% CI: 1.294–3.520; p=0.003 for CSS).

Cox regression analyses for PFS and CSS were also performed including chemoimmunotherapies received (Table S4). After adjusting for confounders including the treatment type (R-CHOP, DA-EPOCH+R, R-CHOP like regimens and R-mini CHOP), DM was still significantly associated with unfavorable PFS and CSS both in the complete (HR=1.457; 95% CI: 1.037–2.047; p=0.030 for PFS and HR=2.258; 95% CI: 1.535–3.324; p<0.001 for CSS) and PSM datasets (HR=1.672; 95% CI: 1.078 –2.592; p=0.022 for PFS and HR=2.052; 95% CI: 1.239 –3.397; p=0.005 for CSS).

## Subgroup analyses of preexisting DM in DLBCL

The results of subgroup analyses are shown in Table 3 (test for interaction: p < 0.05 for all subgroups). After PSM, patients with DM in low-risk grade (IPI 0-2) had evidently poorer PFS (p=0.011) and CSS (p < 0.001) compared with nondiabetics. Whereas in high-risk grade (IPI 3-5), no significance was identified between diabetic and nondiabetic DLBCL patients for both PFS (p=0.507) and CSS (p=0.370). Interestingly, similar results were noticed in subgroups stratified by age, stage, ECOG PS, extranodal sites involvement, Hans classification, symptoms and LDH level. Significant differences in PFS between diabetics and nondiabetics were shown in patients with younger age ( $\leq 60$  years), earlier stage (I–II), no B symptoms and lower LDH concentration ( $\leq$ ULN). While for OS, diabetic patients demonstrated significant inferior survival to nondiabetics in population with younger age ( $\leq 60$  years), better ECOG PS (0–1), fewer extranodal sites ( $\leq 1$ ), Hans classification of GCB, no B symptoms and lower LDH concentration ( $\leq$ ULN). These findings might indicate that DM as a valuable prognostic factor could further stratify the survival outcome of low-risk DLBCL patients.

### Prediabetes, diabetic duration, HbA1c and treatments in relation to DLBCL prognosis

Four hundred and forty-four nondiabetic patients were categorized into prediabetics (43 cases) and patients having no tendency toward DM (401 cases). As observed in Figure 2A and B, significant overall differences in PFS (p<0.001) and CSS (p<0.001) were identified among the three subgroups (log-rank test across all groups). Pairwise comparisons of every specific two groups exhibited that prediabetic patients were more inclined to have poorer PFS (p=0.001) and CSS (p=0.004) compared with those having no diabetic tendency. However, no significantly favorable survival outcome was detected in prediabetics compared with diabetics for both PFS (p=0.732) and CSS (p=0.277) (Table S5).

Diabetic duration was defined as the interval from the date of initial diagnosis as DM to the date of first DLBCL hospital admission. One hundred and nine diabetic patients were divided into four categories by quartiles (Q1=25%, Q2=50%, Q3=75%) of diabetic duration:  $\geq$ 120 months,  $\geq$ 48 and <120 months,  $\geq$ 12 and <48 months and <12 months. Three patients whose FPG >7.0 mmol/l at enrollment were also diabetics, and their diabetic durations were calculated as 0 month. In Figure 2C and D, longer diabetic duration was associated with worse PFS and CSS in DLBCL patients (p<0.001 for both PFS and CSS, log-rank test across all groups). Further comparisons by pairwise over strata showed diabetic patients with duration <12 months had evidently better prognosis than other subgroups (all p≤0.001) (Table S6).

Due to the retrospective nature of this study, test of HbA1c was only performed in 61 diabetic DLBCL cases. Diabetic patients with (61 cases) and without (48 cases) HbA1c result had no significant difference in the distribution of clinical characteristics (all p>0.05). Diabetic

Log-rank test (Mante	–Cox) of diabetes in	Unmatched (com	nplete) da	taset		Propensity score-	matched (	l:l) dataset <sup>a</sup>	
subgroups		PFS		CSS		PFS		CSS	
		HR (95% CI)	p-value	HR (95% CI)	p-value	HR (95% CI)	þ-value	HR (95% CI)	p-value
Age	≤60 years	2.165 (1.306–3.588)	<b>0.002</b>	3.662 (2.076–6.458)	<0.001	2.318 (1.074–5.000)	<b>0.027</b>	4.369 (1.606–11.884)	<b>0.002</b>
	>60 years	1.265 (0.821–1.948)	0.285	1.823 (1.115–2.981)	0.015	1.361 (0.804–2.304)	0.249	1.570 (0.873–2.822)	0.129
Id	0-2	2.024 (1.268–3.231)	<b>0.003</b>	3.406 (1.960–5.919)	<0.001	2.347 (1.189–4.632)	<b>0.011</b>	4.616 (1.839–11.589)	<b>&lt;0.001</b>
	3-5	1.321 (0.838–2.084)	0.229	1.903 (1.156–3.134)	0.010	1.213 (0.686–2.145)	0.507	1.321 (0.717–2.431)	0.370
Stage	I−I	3.336 (1.635–6.808)	<b>&lt;0.001</b>	6.063 (2.494–14.739)	<0.001	3.493 (1.100–11.090)	<b>0.024</b>	6.364 (1.384–29.255)	0.006
	III−IV	1.441 (0.996–2.086)	0.051	2.279 (1.507–3.447)	<0.001	1.355 (0.842–2.181)	0.208	1.813 (1.059–3.106)	0.028
ECOG PS	0-I	1.796 (1.237–2.609)	<b>0.002</b>	3.010 (1.956–4.631)	<b>&lt;0.001</b>	1.597 (0.980–2.603)	0.058	2.326 (1.301–4.158)	<b>0.003</b>
	2-4	1.324 (0.675–2.597)	0.412	1.641 (0.796–3.382)	0.175	1.560 (0.609–3.995)	0.35 I	1.463 (0.562–3.810)	0.433
Extranodal site(s)	VI .	1.687 (1.114–2.555) 1.879 (1.108–3.187)	0.012 0.017	2.686 (1.671–4.317) 2.799 (1.548–5.059)	<0.001 <0.001	1.696 (0.975–2.953) 1.455 (0.733–2.885)	0.059 0.281	2.835 (1.423–5.648) 1.424 (0.690–2.938)	<b>0.002</b> 0.337
Hans classification	GCB non-GCB	2.333 (1.277–4.261) 1.516 (1.028–2.238)	0.005 0.035	4.418 (2.218–8.799) 2.177 (1.399–3.389)	<0.001 <0.001	2.294 (0.933–5.640) 1.491 (0.904–2.460)	0.063 0.115	8.923 (2.010–39.602) 1.584 (0.912–2.751)	<b>0.1</b> 00
Symptoms	No B symptoms	2.125 (1.349–3.347)	<b>0.001</b>	3.331 (1.984–5.592)	<0.001	I.864 (I.014–3.426)	<b>0.042</b>	2.545 (1.257–5.150)	<b>0.007</b>
	B symptoms	1.346 (0.842–2.152)	0.212	2.077 (1.222–3.531)	0.006	I.303 (0.709–2.393)	0.392	1.728 (0.858–3.483)	0.121
НОТ	≤ULN (271U/L)	2.177 (1.329–3.566)	<b>0.002</b>	4.043 (2.252–7.258)	<0.001	1.965 (1.015–3.804)	<b>0.041</b>	3.328 (1.474–7.513)	<b>0.002</b>
	≤ULN (271U/L)	1.331 (0.862–2.057)	0.196	1.802 (1.114–2.918)	0.015	1.321 (0.746–2.339)	0.337	1.430 (0.759–2.692)	0.266
<b>Note</b> : <sup>a</sup> Propensity score-matche level. <i>p</i> -values in bold < 0.05. <b>Abbreviations</b> : PFS, progressio dehydrogenase; ULN, upper limit	d (1:1, 109 pairs) analyses were n-free survival; CSS, cancer-spe : of normal.	performed with a small cal scific survival; IPI, internati	iper of 0.1 for onal prognosti	matching potential confound c index; ECOG, Eastern C	ders including : poperative Or	iex, age, stage, ECOG PS, e cology Group; PS, perform	xtranodal site( iance status; G	s), Hans classification, sympt iCB, germinal center B-cell	coms and LDH ; LDH, lactate

Table 3 Prognostic value of diabetes in different subgroups before and after propensity matching

**Dove**Press



Figure 2 Prediabetes, diabetic duration and HbA1c level in relation to DLBCL survival. (A) PFS stratified by different diabetic status of T2DM, pre-diabetes and no T2DM tendency. (B) CSS stratified by different diabetic status of T2DM, pre-diabetes and no T2DM tendency. (C) PFS stratified by different diabetic durations. (D) CSS stratified by different diabetic durations. (E) PFS stratified by different HbA1c levels. (F) CSS stratified by different HbA1c levels.

Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival; DM, diabetes mellitus; HbA1c, hemoglobin A1c; DLBCL, diffuse large B- cell lymphoma.

patients were subdivided into three groups according to their HbA1c levels:  $\geq 8.0\%$ ,  $\geq 6.5\%$  and < 8.0% and < 6.5%. Figure 2E and F and pairwise comparisons presented that adequate glycemic control was associated with longer PFS and CSS in diabetic DLBCL patients (Table S7).

We also investigated the correlation between diabetic medications and prognosis of DLBCL. Pairwise comparisons exhibited that no significant difference was identified for both PFS and CSS between every two hypoglycemic treatments (oral antidiabetic drugs vs insulin; oral antidiabetic drugs vs nonmedication; insulin vs nonmedication). Further adjustments for other potential confounders including diabetic duration did not change the magnitude substantially (Table S8).

# DM together with IPI: a better prognostic index for DLBCL

For the reason that DM remained as an independent prognostic factor for both PFS and CSS in multivariate analyses, introducing a novel PI which included IPI together with DM might



Figure 3 DM together with IPI is a better prognostic index for DLBCL. (A) The AUC comparison between DM-PI, IPI and T2DM alone for PFS prediction. (B) Kaplan-Meier curves of PFS for three different DM-PI risk grades. (C) The AUC comparison between DM-PI, IPI and T2DM alone for CSS prediction. (D) Kaplan-Meier curves of CSS for three different DM-PI risk grades.

Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival; DM, diabetes mellitus; DLBCL, diffuse large B-cell lymphoma; IPI, international prognostic index; DM-PI, prognostic index including diabetes mellitus; AUC, area under the curve.

provide better predictive accuracy than IPI alone. PI was calculated as the sum of IPI and one additional point for patient's diabetic status. Interestingly as shown in Figure 3A and C, PI exhibited evidently larger AUC and greater accuracy than IPI alone in CSS prediction (0.752 vs 0.732, p=0.024). To further validate that PI was a better index for CSS prediction, we divided the entire cohort into three risk grades: low-risk group (DM-PI 0–1), intermediate-risk group (DM-PI 2–3) and high-risk group (DM-PI 4–6). A significant overall difference

in CSS can be observed from the three subgroups (p<0.001, log-rank test across all groups) that DLBCL patients in highrisk grade demonstrated the poorest CSS and those in low-risk grade had the most favorable survival outcomes (Figure 3B and D). The 3-year and 5-year PFS and CSS of the complete dataset stratified by the IPI and the DM-PI are demonstrated in Table 4. The DM-PI both widened the definition of low-risk and high-risk patients for 3-year and 5-year CSS and maintained equivalent accuracy to the IPI in PFS estimation.

Table 4 PFS and CSS according to the IPI and the DM-PI in the complete dataset

Risk group	IPI 3-year PFS (%)	IPI 5-year PFS (%)	IPI 3-year CSS (%)	IPI 5-year CSS (%)
Low (0-1) Intermediate (2-3) High (4-5)	85.2 58.6 39.3	83.9 54.8 39.3	92.8 72.4 51.8	91.5 70.9 50.0
Risk group	DM-PI 3-year PFS (%)	DM-PI 5-year PFS (%)	DM-PI 3 year-CSS (%)	DM-PI 5-year CSS (%)

Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival; IPI, international prognostic index; DM-PI, prognostic index including diabetes.

Additionally, pairwise comparisons as displayed in Table S9 indicated evident difference in CSS between every two specific risk grades, which suggested that PI could accurately differentiate DLBCL patients into different risk grades for CSS prediction.

### Discussion

To our knowledge, no previous study has focused on the correlation between DLBCL and DM, and this is the first article providing evidence that DM could serve as an independent risk factor for survival in patients with DLBCL.

Diabetes is associated with elevated risks in various malignancies, especially in gastric, pancreatic and hepatocellular cancers. Our study demonstrated that the prevalence of preexisting DM in DLBCL was 19.71%, which was evidently higher in comparison with the results from a Chinese national study of diabetes (9.70%), including representative samples of 46,239 adults.<sup>15</sup> Tseng et al concluded DM was associated with high risk of developing NHL (OR=1.51; 95% CI: 1.33-1.71) in Taiwan<sup>16</sup> and Yang et al demonstrated similar results with HR of 2.00 (95% CI: 1.32-3.03) in mainland China.<sup>17</sup> Additionally, Lin et al reported that preexisting DM was an independent risk factor for the occurrence of NHL (OR=1.88; 95% CI: 1.22–2.89), particularly for extranodal presentation, T-cell subtype and low-grade tumor.<sup>18</sup> However, a multinational investigation, namely European Prospective Investigation into Cancer and Nutrition study, revealed a lack of association.<sup>19</sup> This investigation did not separately analyze the incidence and mortality rate of NHL.

The pros and cons of PSM deserve mention and are listed as follows. Pros are: 1) equating groups at baseline especially in studies that do not use randomization; 2) accounted for "nuisance" factors that differ between the groups, so that any remaining difference is likely due to the variable of interest. Cons are: 1) difficult to include all the confounding covariates; 2) possibility of losing subjects depending on the original size of the pool of available people and the amount of overlap, further jeopardizing the results of the study; 3) many alternative methods (nearest neighbor, caliper matching, etc.) reflect that no single approach is ideal and each has limitations, thus sensitivity analyses are needed.

Our findings suggested that DM was associated with worse PFS and CSS both in the complete and propensitymatched cohorts, and it was an independent risk factor for both PFS and CSS. In subgroup analyses, DM was correlated with inferior survival only in patients with younger age ( $\leq 60$  years), lower IPI, no B symptoms and lower LDH concentration. The probable explanation for this phenomenon is that other strong prognostic factors such as older age, more advanced stage and higher IPI outweighed and covered up the effect of DM on prognosis. PFS and CSS of high-risk group may be greater influenced by lymphoma effect, and those of low-risk group may be more likely to be influenced by nonlymphoma comorbidities including DM. Accordingly, DM could be a valuable indicator improving the prognostic stratification of low-risk DLBCL patients.

We also demonstrated that prediabetics were more likely to have inferior prognosis compared to patients with no diabetic tendency. It may imply that DLBCL patients with prediabetes should be given equal concerns as those with comorbid DM because they might share inferior survival outcomes. Additionally, patients with shorter diabetic duration (<12 months) and adequate glycemic control (HbA1C <6.5%) presented better survival. Given that patients who died of non-DLBCL-related conditions including diabetic complications were excluded from the study, the above findings could directly indicate the impact of DM on DLBCL-specific survival.

We noticed that DM together with IPI (DM-PI) had larger AUC compared with IPI alone in CSS prediction. Although statistically significant, the inclusion of DM only resulted in minor improvement. To interpret our results more cautiously, DM-PI was not intended to replace IPI in clinical practice, but diabetes should be well-assessed and managed in DLBCL patient care. Due to the retrospective nature and lack of validation cohort, our results remain to be replicated and confirmed in epidemiologic studies with larger samples, longer follow-up periods and full adjustments for covariates to test the proposed prognostic score.

As previously reported in epidemiological evidence, diabetic NHL patients had evidently worse survival after approximately 1 year of follow-up, and it was an accelerated factor for the risk of death from causes related to non-Hodgkin's lymphoma.<sup>18</sup> Another cohort study among Taiwanese population suggested diabetics were at an increased risk of NHL mortality (HR=1.028; 95% CI: 1.005-1.051).<sup>20</sup> For other site-specific cancers, evidence was provided that preexisting diabetes was associated with poor disease-free survival (DFS) and overall survival (OS) in lung (HR=1.27; 95% CI: 1.07-1.50),<sup>21</sup> breast (HR=1.97; 95% CI: 1.45-2.68),<sup>22</sup> colorectal (HR=1.17; 95% CI: 1.09-1.25)<sup>23</sup> and prostate cancer (HR=1.56; 95% CI: 1.03-2.36).<sup>24</sup>

However, the potential etiology role of DM in DLBCL development has not been investigated, and it is difficult to tease out a possible causal sequence between DM and DLBCL. It was postulated that these two diseases might share common pathways and risk factors in the early stage of development, such as immunodeficiency and obesity.<sup>25</sup> DM is related to high oxidative stress, inflammatory status, and immune dysfunction which are implicated in the tumorigenesis of DLBCL. In preclinical experiments, a mouse DLBCL cell line with Chemokine (C-C motif) ligand 5 (CCL5) overexpression and knockdown subclones were subcutaneously injected into both BALB/c DM mice and normal mice. Zhang et al concluded that CCL5 could be one of the indispensable chemoattractant cytokines involved in DM with DLBCL, and high levels of CCL5 expression could accelerate the formation of DLBCL in DM mice.<sup>26</sup> Another study in vivo demonstrated G13997A mitochondrial DNA could regulate diabetes development and lymphoma formation via the overproduction of reactive oxygen species (ROS) or lactate.<sup>27</sup>

In the current clinical practice of DLBCL, little attention was paid to controlling the progression of comorbid DM and its complication. The implications of our study are: 1) concerns should be given to patients with diabetes or prediabetes at first diagnosis of DLBCL to help predict life expectancy; (2) blood glucose level should be routinely monitored for diabetic patients, because hyperglycemia could cause prednsolone dosing attenuated, it should be actively controlled using effective antidiabetic agents.

The limitations of our study are illustrated as follows: the restriction within one institution; limited number of patients enrolled; inability to account for unmeasured confounders despite using PSM analyses (eg, different types of rituximab combination chemotherapies including R-CHOP, DA-EPOCH+R, R-CHOP-ike regimes and R-mini CHOP; diabetes-related lifestyle) and the retrospective nature of this study with incomplete data on HbA1c. Owing to the fact that the informed consent was not restricted to this study and the research design was not established before the patients' enrollment, we still consider it as a retrospective one.

Our study concluded that DLBCL patients with preexisting DM had evident worse PFS and CSS than those without DM in rituximab era, and it was an independent risk factor for PFS and CSS both in the complete and PSM datasets. Prediabetics also demonstrated inferior prognostic outcomes compared to patients with no diabetic tendency. Adding the criterion of DM improved the prognostic capacity and risk stratification of IPI for CSS prediction. Further multi-central and multi-racial prospective studies with improved control of confounders are needed to confirm these findings.

### Acknowledgments

We are grateful to all the study participants, patients and their family members for their contributions and supports. This study was supported by National Natural Science Foundation of China (81370657, 81470328, 81600130, 81770166, 81720108002), Jiangsu Province's Medical Elite Programme (ZDRCA2016022), Project of National Key Clinical Specialty, National Science & Technology Pillar Program (2014BAI09B12), Jiangsu Provincial Special Program of Medical Science (BL2014086 and BE2017751) and National Science and Technology Major Project (2017ZX09304032).

### Disclosure

The authors report no conflicts of interest in this work.

### References

- Dyal HK, Aguilar M, Bartos G, et al. Diabetes mellitus increases risk of hepatocellular carcinoma in chronic hepatitis c virus patients: a systematic review. *Dig Dis Sci.* 2016;61(2):636–645. doi:10.1007/ s10620-015-3983-3
- Chari ST, Leibson CL, Rabe KG, et al. Pancreatic cancer-associated diabetes mellitus: prevalence and temporal association with diagnosis of cancer. *Gastroenterology*. 2008;134(1):95–101. doi:10.1053/j. gastro.2007.10.040
- Luo W, Cao Y, Liao C, et al. Diabetes mellitus and the incidence and mortality of colorectal cancer: a meta-analysis of 24 cohort studies. *Colorectal Dis.* 2012;14(11):1307–1312. doi:10.1111/j.1463-1318.2012.02875.x
- Zhou Y, Zhang X, Gu C, et al. Influence of diabetes mellitus on mortality in breast cancer patients. ANZ J Surg. 2015;85(12):972– 978. doi:10.1111/ans.12877
- Mitri J, Castillo J, Pittas AG. Diabetes and risk of non-hodgkin's lymphoma: a meta-analysis of observational studies. *Diabetes Care*. 2008;31(12):2391–2397. doi:10.2337/dc08-1034
- Castillo JJ, Mull N, Reagan JL, et al. Increased incidence of non-Hodgkin lymphoma, leukemia, and myeloma in patients with diabetes mellitus type 2: a meta-analysis of observational studies. *Blood*. 2012;119(21):4845–4850. doi:10.1182/blood-2011-06-362830
- Chao C, Page JH. Type 2 diabetes mellitus and risk of non-Hodgkin lymphoma: a systematic review and meta-analysis. *Am J Epidemiol.* 2008;168(5):471–480. doi:10.1093/aje/kwn160
- Martelli M, Ferreri AJ, Agostinelli C, et al. Diffuse large b-cell lymphoma. *Crit Rev Oncol Hematol.* 2013;87(2):146–171. doi:10.1016/j.critrevonc.2012.12.009
- Tilly H, Gomes Da Silva M, Vitolo U, et al. Diffuse large b-cell lymphoma (dlbcl): esmo clinical practice guidelines for diagnosis, treatment and follow-up. *Ann Oncol.* 2015;26(Suppl 5):v116–v125. doi:10.1093/annonc/mdv304
- Friedberg JW. Relapsed/refractory diffuse large b-cell lymphoma. *Hematology Am Soc Hematol Educ Program.* 2011;2011:498–505. doi:10.1182/asheducation-2011.1.498

- Goto N, Tsurumi H, Kasahara S, et al. Serum interleukin-18 level is associated with the outcome of patients with diffuse large bcell lymphoma treated with chop or r-chop regimens. *Eur J Haematol.* 2011;87(3):217–227. doi:10.1111/j.1600-0609.2011. 01649.x
- Cheson BD, Pfistner B, Juweid ME, et al. Revised response criteria for malignant lymphoma. J Clin Oncol. 2007;25(5):579–586. doi:10.1200/JCO.2006.09.2403
- National Center for Classification in Health. The International Statistical Classification of Diseases and Related Health Problems, Tenth Revision, Australian Modification (Icd-10, Am). 3rd ed. Sydney; 2012.
- DeLong ER, DeLong DM, Clarke-Pearson DL. Comparing the areas under two or more correlated receiver operating characteristic curves: a nonparametric approach. *Biometrics*. 1988;44(3):837–845.
- 15. Yang W, Lu J, Weng J, et al. Prevalence of diabetes among men and women in china. N Engl J Med. 2010;362(12):1090–1101. doi:10.1056/NEJMoa0908292
- 16. Tseng CH. Diabetes and non-hodgkin's lymphoma: analyses of prevalence and annual incidence in 2005 using the national health insurance database in taiwan. *Ann Oncol.* 2012;23(1):153–158. doi:10.1093/annonc/mdr334
- Yang WS, Li HL, Xu HL, et al. Type 2 diabetes and the risk of nonhodgkin's lymphoma: a report from two population-based cohort studies in china. *Eur J Cancer Prev.* 2016;25(2):149–154. doi:10.1097/CEJ.00000000000150
- Lin SY, Hsieh MS, Chen LS, et al. Diabetes mellitus associated with the occurrence and prognosis of non-hodgkin's lymphoma. *Eur J Cancer Prev.* 2007;16(5):471–478. doi:10.1097/01.cej.0000236253. 93984.8f

- Khan AE, Gallo V, Linseisen J, et al. Diabetes and the risk of nonhodgkin's lymphoma and multiple myeloma in the european prospective investigation into cancer and nutrition. *Haematologica*. 2008;93 (6):842–850. doi:10.3324/haematol.12297
- Tseng CH. Diabetes, insulin use, and non-Hodgkin lymphoma mortality in taiwan. *Metabolism*. 2012;61(7):1003–1009. doi:10.1016/j. metabol.2011.11.015
- Luo J, Hendryx M, Qi L, et al. Pre-existing diabetes and lung cancer prognosis. Br J Cancer. 2016;115(1):76–79. doi:10.1038/bjc.2016.141
- Mu L, Zhu N, Zhang J, et al. Type 2 diabetes, insulin treatment and prognosis of breast cancer. *Diabetes Metab Res Rev.* 2016;33:e2823.
- Mills KT, Bellows CF, Hoffman AE, Kelly TN, Gagliardi G. Diabetes mellitus and colorectal cancer prognosis: a meta-analysis. *Dis Colon Rectum.* 2013;56(11):1304–1319. doi:10.1097/ DCR.0b013e3182a479f9
- Polesel J, Gini A, Dal Maso L, et al. The impact of diabetes and other metabolic disorders on prostate cancer prognosis. J Diabetes Complications. 2016;30(4):591–596. doi:10.1016/j. jdiacomp.2016.02.008
- Ekstrom-Smedby K. Epidemiology and etiology of non-Hodgkin lymphoma–a review. Acta Oncol. 2006;45(3):258–271. doi:10.1080/ 02841860500531682
- 26. Zhang J, Luo J, Liu F, et al. Diabetes mellitus potentiates diffuse large bcell lymphoma via high levels of ccl5. *Mol Med Rep.* 2014;10 (3):1231–1236. doi:10.3892/mmr.2014.2341
- 27. Hashizume O, Shimizu A, Yokota M, et al. Specific mitochondrial DNA mutation in mice regulates diabetes and lymphoma development. *Proc Natl Acad Sci U S A.* 2012;109(26):10528–10533. doi:10.1073/pnas.1202367109

### Supplementary materials

Clinical parameters		Total	PFS		CSS	
			Event (N)	Rate (%)	Event (N)	Rate (%)
Gender	Male	293	109	37.20	67	22.87
	Female	260	83	31.92	60	23.08
Age	≤60 years	324	94	29.01	59	18.21
	>60 years	229	98	42.79	68	29.69
IPI	0-2	381	93	24.41	56	14.70
	3-5	172	99	57.56	71	41.28
Stage	I–II	209	34	16.27	20	9.57
	III–IV	344	158	45.93	107	31.10
ECOG PS	0-1	471	149	31.63	93	19.75
	2-4	82	43	52.44	34	41.46
Extranodal site(s)	≤	403	121	30.02	78	19.35
	>	150	71	47.33	49	32.67
Hans classification	GCB	211	52	24.64	34	16.11
	Non-GCB	342	140	40.94	93	27.19
Symptoms	No B symptoms	342	95	27.78	63	18.42
	B symptoms	211	97	45.97	64	30.33
LDH	≤ULN (271U/L)	351	82	23.36	49	13.96
	>ULN (271U/L)	202	110	54.46	78	38.61
Albumin	<lln (40u="" l)<="" td=""><td>332</td><td>132</td><td>39.76</td><td>92</td><td>27.71</td></lln>	332	132	39.76	92	27.71
	≥LLN (40U/L)	221	60	27.15	35	15.84
Treatments	R-CHOP	404	144	35.64	92	22.77
	DA-EPOCH+R	118	39	33.05	28	23.73
	R-CHOP like regimens	22	6	27.27	6	27.27
	R-mini CHOP	9	3	33.33	1	11.11
Response to therapy <sup>a</sup>	CR or PR	392	69	17.60	34	8.67
	Non-CR or PR	138	100	72.46	70	50.72
Diabetic status	Diabetic	109	49	44.95	43	39.45
	Nondiabetic	444	143	32.21	84	18.92

Table SI Event rates of PFS and CSS in subgroups stratified by different clinical parameters

Note: <sup>a</sup>The evaluation of treatment response was performed after 6–8 cycles of chemotherapies. Patients who died during the therapies were excluded. p-values in bold < 0.05. **Abbreviations:** PFS, progression-free survival; CSS, cancer-specific survival; IPI, international prognostic index; ECOG, Eastern Cooperative Oncology Group; PS, performance status; GCB, germinal center B-cell; LDH, lactate dehydrogenase; ULN, upper limit of normal; LLN, lower limit of normal; R-CHOP, rituximab, cyclopho-sphamide, doxorubicin, vincristine and prednisone; DA-EPOCH+R, dose-adjusted (etoposide, prednisone, vincristine, cyclophosphamide and doxorubicin) plus rituximab; R-mini CHOP, rituximab combined with low-dose CHOP; CR, complete remission; PR, partial remission.

Diabetic	Unmatc	hed (com	plete) dat	aset					Propensi	ty score-n	natched (	l:l) dataset <sup>®</sup>	-			
status	PFS			Median	css			Median	PFS			Median	css			Median
	l-year	3-year	5-year	PFS (months)	l-year	3-year	5-year	CSS (months)	l-year	3-year	5-year	PFS (months)	l-year	3-year	5-year	CSS (months)
Whole	76.9%	65.7%	59.3%		%0.16	76.2%	72.8%	ı	73.5%	61.1%	52.8%	70.03	87.8%	67.8%	61.1%	ı
cohort Diabetic	67.0%	54.6%	44.7%	56.30	84.9%	55.1%	50.8%		67.0%	54.6%	44.7%	56.30	84.9%	55.1%	50.8%	
group Nondiabetic	79.3%	68.4%	62.2%		92.5%	81.0%	77.8%		79.7%	67.4%	59.2%		90.7%	76.9%	71.6%	
group																
Note: <sup>a</sup> Propensity level. <i>p</i> -values in b Abbreviations: L	score-match old < 0.05. DH, lactate	ned (1:1, 109 dehydrogenas	pairs) analys se; ECOG, E	ses were perform astern Cooperat	ive Oncolog	aall caliper of y Group; PS,	0.1 for mat	ching potential c e status; PFS, pr	:onfounders i ogression-fre	ncluding sex, se survival; C	age, stage, F SS, cancer-sp	COG PS, extra becific survival.	nodal site(s),	Hans classif	ication, symp	toms and LDH

Table S2 Survival rate and median survival according to diabetic status before and after propensity matching

**Dove**Press

ching
:y mat
propensit
nd after
before a
nortality)
ll-cause n
iS (a
O P
S an
f PF
SS O
analyse
regression
Ň
Iriate (
multiva
and
Univariate
23 (
Table (

Clinic	cal parameter	Unmatched (comp	lete) dataset			Propensity score-m	atched (I:I)	dataset <sup>a</sup>	
		Univariate analyses		Multivariate analyse	es	Univariate analyses		Multivariate analys	es
		HR (95% CI)	p-value	HR (95% CI)	þ-value	HR (95% CI)	þ-value	HR (95% CI)	þ-value
PFS	Male	1.154 (0.874–1.524)	0.311	0.997 (0.753–1.321)	0.985	0.751 (0.494–1.141)	0.180	0.659 (0.428–1.016)	0.059
	Age >60 years	I.768 (I.341–2.331)	<0.001	1.534 (1.148–2.048)	0.004	1.322 (0.847–2.062)	0.219	1.285 (0.814–2.030)	0.282
	Stage III–IV	3.002 (2.116-4.258)	<0.001	1.957 (1.331–2.878)	0.001	2.523 (1.485–4.288)	0.001	1.600 (0.873–2.933)	0.128
	ECOG PS 2-4	2.075 (I.488–2.892)	<0.001	1.422 (1.002–2.017)	0.049	I.845 (I.128–3.020)	0.015	1.368 (0.808–2.316)	0.244
	Extranodal sites >	I.905 (I.432–2.534)	<0.001	1.227 (0.902–1.671)	0.193	2.444 (1.593–3.748)	0.000	1.892 (1.148–3.117)	0.012
	Non-GCB	I.804 (I.327–2.451)	<0.001	1.568 (1.151–2.137)	0.004	1.656 (1.020–2.687)	0.041	1.920 (1.167–3.160)	0.010
	B symptoms	1.830 (1.389–2.410)	<0.001	1.214 (0.901–1.635)	0.203	I.574 (I.035–2.393)	0.034	1.039 (0.635–1.698)	0.880
	LDH >ULN (271U/L)	2.912 (2.205–3.847)	<0.001	I.993 (I.475–2.693)	<0.001	2.502 (1.640–3.819)	0.000	1.870 (1.128–3.101)	0.015
	Diabetic	1.720 (1.251–2.366)	0.001	1.409 (1.012–1.961)	0.042	1.716 (1.119–2.629)	0.013	1.769 (1.144–2.735)	0.010
S	Male	0.950 (0.680–1.328)	0.765	0.832 (0.593–1.166)	0.285	0.695 (0.435–1.111)	0.129	0.637 (0.396–1.025)	0.063
	Age ≥60 years	2.010 (1.435–2.814)	<0.001	1.654 (1.162–2.355)	0.005	1.443 (0.871–2.391)	0.155	1.312 (0.775–2.221)	0.313
	Stage III–IV	2.932 (1.901–4.523)	<0.001	2.039 (1.266–3.284)	0.003	2.492 (1.363–4.556)	0.003	1.699 (0.854–3.383)	0.131
	ECOG PS 2-4	2.612 (1.783–3.829)	<0.001	1.820 (1.215–2.727)	0.004	2.218 (1.304–3.771)	0.003	1.590 (0.896–2.821)	0.113
	Extranodal sites >1	1.966 (1.394–2.771)	<0.001	1.222 (0.844–1.770)	0.289	2.917 (1.821–4.673)	0.000	1.998 (1.168–3.419)	0.012
	Non-GCB	1.756 (1.212–2.544)	0.003	1.502 (1.032–2.185)	0.033	1.882 (1.076–3.292)	0.027	2.039 (1.141–3.645)	0.016
	B symptoms	I.649 (I.180–2.303)	0.003	1.043 (0.726–1.499)	0.818	1.454 (0.909–2.326)	0.118	0.900 (0.518–1.563)	0.708
	LDH >ULN (271U/L)	3.094 (2.202–4.349)	<0.001	2.078 (1.435–3.009)	<0.001	2.674 (1.661–4.306)	0.000	1.934 (1.092–3.426)	0.024
	Diabetic	2.571 (1.797–3.679)	<0.00 I	2.077 (1.432–3.013)	<0.001	2.281 (1.394–3.734)	0.001	2.134 (1.294–3.520)	0.003
Note: <sup>a</sup> P level.	ropensity score-matched (1:1, 111	pairs) analyses were perform	ed with a small cal	iper of 0.1 for matching poten	tial confounders ir	ıcluding sex, age, stage, ECOG	PS, extranodal si	te(s), Hans classification, symp	toms and LDH

Abbreviations: PFS, progression-free survival; COS, overall survival; ECOG, Eastern Cooperative Oncology Group; PS, performance status; GCB germinal center B-cell; LDH, lactate dehydrogenase; ULN, upper limit of normal.

**Dove**Press

Table S	4 Univariate and multivar	iate Cox regression and	Ilyses of PFS a	and CSS including cheme	otherapy recei	ived before and after pr	opensity matc	hing	
Clinic	ıl parameter	Unmatched (comp	lete) dataset	t		Propensity score-m	atched (I:I)	dataset <sup>a</sup>	
		Univariate analyses		Multivariate analys	es	Univariate analyses		Multivariate analys	es
		HR (95% CI)	p-value	HR (95% CI)	p-value	HR (95% CI)	p-value	HR (95% CI)	p-value
PFS	Male	1.169 (0.879–1.556)	0.284	1.017 (0.761–1.359)	0.911	0.776 (0.508–1.187)	0.243	0.709 (0.457–1.102)	0.126
	Age >60 years	1.678 (1.263–2.228)	<0.001	1.409 (1.044–1.903)	0.025	1.270 (0.811–1.987)	0.296	1.230 (0.774–1.955)	0.380
	Stage III-IV	3.311 (2.285-4.797)	<0.001	2.163 (1.439–3.249)	<0.001	2.717 (1.577–4.681)	<0.001	1.794 (0.967–3.327)	0.064
	ECOG PS 2-4	2.088 (I.486–2.935)	<0.001	1.447 (1.006–2.081)	0.046	1.763 (1.067–2.914)	0.027	1.354 (0.786–2.332)	0.275
	Extranodal sites >	1.897 (1.415–2.545)	<0.001	1.176 (0.856–1.614)	0.317	2.280 (1.476–3.522)	<0.001	1.732 (1.046–2.868)	0.033
	Non-GCB	1.904 (1.384–2.619)	<0.001	1.660 (1.203–2.290)	0.002	1.601 (0.985–2.603)	0.058	1.857 (1.128–3.056)	0.015
	B symptoms	1.912 (1.440–2.538)	<0.001	1.235 (0.909–1.677)	0.177	1.673 (1.094–2.557)	0.017	1.091 (0.659–1.806)	0.735
	LDH >ULN (271U/L)	3.041 (2.282-4.052)	<0.001	2.048 (1.502–2.793)	<0.001	2.459 (1.604–3.770)	<0.001	1.808 (1.080–3.024)	0.024
	Treatments <sup>b</sup>	0.961 (0.755–1.222)	0.743	0.926 (0.721–1.191)	0.550	1.057 (0.753–1.484)	0.747	0.936 (0.634–1.383)	0.741
	Diabetic	1.744 (1.259–2.417)	0.001	1.457 (1.037–2.047)	0.030	1.607 (1.045–2.470)	0.03	1.672 (1.078–2.592)	0.022
CSS	Male	0.948 (0.669–1.344)	0.764	0.817 (0.573–1.164)	0.263	0.721 (0.448–1.161)	0.179	0.657 (0.404–1.069)	160.0
	Age ≥60 years	1.867 (1.316–2.647)	<0.001	1.500 (1.035–2.175)	0.032	I.388 (0.835–2.308)	0.206	1.320 (0.774–2.250)	0.308
	Stage III-IV	3.443 (2.135–5.551)	<0.001	2.340 (1.389–3.942)	0.001	2.676 (1.433–4.998)	0.002	1.722 (0.842–3.525)	0.137
	ECOG PS 2-4	2.730 (1.839-4.053)	<0.001	1.854 (1.213–2.835)	0.004	2.178 (1.266–3.746)	0.005	1.614 (0.890–2.929)	0.115
	Extranodal sites >	1.976 (1.382–2.825)	<0.001	1.179 (0.802–1.734)	0.402	2.742 (1.698–4.429)	<0.001	1.901 (1.094–3.303)	0.023

4 č 74 J DEC ( C4 1 PS **Notes:** <sup>a</sup>Propensity score-matched (1:1, 109 pairs) analyses were performed with a small caliper of 0.1 for matching potential confounders including sex, age, stage, ECOG PS, extranodal site(s), Hans classification, symptoms and LDH level. <sup>b</sup>Treatments referred to R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine and prednisone), DA-EPOCH+R (dose-adjusted (etoposide, prednisone, vincristine, cyclophosphamide and doxorubicin) plus rituximab), R-CHOP like regimens, and R-mini CHOP (rituximab combined with low-dose CHOP). *p*-values in bold < 0.05.

0.038

<0.001 0.076

> 2.621 (1.616-4.251) I.192 (0.824–1.724)

<0.001

1.539 (0.956–2.477)

1.816 (1.036–3.184)

0.014

1.650 (1.108-2.457) 1.054 (0.723-1.538) 2.156 (1.462-3.178) 1.068 (0.794–1.436) 2.258 (1.535-3.324)

0.00 0.002

1.896 (1.280-2.810) 1.757 (1.241–2.489) 3.341 (2.336-4.778)

Non-GCB

0.784

0.003

2.145 (1.306-3.523)

<0.001

<0.001

1.081 (0.810–1.442) 2.713 (1.874–3.928)

0.597

<0.001

(271U/L) LDH >ULN (271U/L) B symptoms

Treatments

Diabetic

0.665

0.351

0.812 0.021

1.990 (1.110-3.567) 0.934 (0.532-1.640) 1.851 (1.034-3.313) 1.059 (0.701-1.600) 2.052 (1.239-3.397)

0.037

0.005

0.784

Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival; ECOG, Eastern Cooperative Oncology Group; PS, performance status; GCB germinal center B-cell; LDH, lactate dehydrogenase; ULN, upper limit of normal.

Log-rank test (Mantel–Cox)	Diabetic status	No diabetic tender	ncy	Prediabetes		Diabetic	
		Chi-square	p-value	Chi-square	p-value	Chi-square	þ-value
PFS (overall p<0.001)	No diabetic tendency	-	-	11.777	100.0	14.787	<0.00 I
	Prediabetes	11.777	0.001	Ι	I	0.117	0.732
	Diabetic	14.787	<0.001	0.117	0.732	I	Ι
CSS (overall p<0.001)	No diabetic tendency	I	1	8.507	0.004	34.934	<0.001
	Prediabetes	8.507	0.004	Ι	Ι	1.179	0.277
	Diabetic	34.934	<0.001	1.179	0.277	I	I
<b>Note:</b> <i>p</i> -values in bold < 0.05.							

Table S5 Pairwise comparisons among different diabetic status for PFS and CSS

Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival.

Datients for PFS and CSS ine in diaheric shis diabetic duration Aiffo Table S6 Pairwise

	מווסוופ חוווכו כוור חומסכמר סמו ממסוו זמספו סמלאז ווו	י שומסכתים המתוכו							
Log-rank test (Mantel–	Diabetic duration subgroups	<12		≥I2 and <4		≥48 and <i2< th=""><th>0</th><th>≥<b> 20</b></th><th></th></i2<>	0	≥ <b> 20</b>	
Cox)	(months) <sup>4</sup>	Chi- square	þ- value	Chi- square	þ- value	Chi- square	þ- value	Chi- square	p- value
PFS (overall p<0.001)	<12	-	-	13.177	<0.001	14.850	<0.001	18.823	<0.001
	≥I2 and <48	13.177	<0.001	Ι	Ι	0.733	0.392	2.140	0.144
	≥48 and <120	14.850	<0.001	0.733	0.392	I	I	1.660	0.198
	≥I 20	18.823	<0.001	2.140	0.144	1.660	0.198	Ι	I
CSS (overall p<0.001)	<12	I	I	10.944	0.001	16.714	<0.001	20.524	<0.001
	≥I2 and <48	10.944	0.001	Ι	Ι	2.271	0.132	6.052	0.014
	≥48 and <120	16.714	<0.001	2.271	0.132	I	I	2.513	0.113
	≥I 20	20.524	<0.001	6.052	0.014	2.513	0.113	Ι	I
Note: <sup>a</sup> Diabetic duration subgroups were Abbreviations: PFS, progression-free su	e categorized by quartiles (Q1=25%, Q2=50%, Q3=75%) of rvival; CSS, cancer-specific survival.	disease duration in	diabetic patie	nts. p-values in bol	d < 0.05.				

Log-rank test (Mantel–C	ox) HbAIc levels	<6.5%		≥6.5% and <8.0%		≥8.0%	
		Chi-square	p-value	Chi-square <i>p</i>	-value	Chi-square	p-value
PFS (overall $p<0.001$ )	<6.5%	1	Ι	3.382 0.	.066	20.431	<0.001
	≥6.5% and <8.0%	3.382	0.066	1		14.328	<0.001
	≥8.0%	20.431	<0.001	14.328 <	0.001	I	
CSS (overall p<0.001)	<6.5%	Ι	-	5.645 0.	.018	22.044	<0.001
	≥6.5% and <8.0%	5.645	0.018			9.699	0.002
	≥8.0%	22.044	<0.001	0 6696	.002	Ι	I
Note: p-values in bold < 0.05. Abbreviations: PFS, progression-free Table S& Comparisons amon	e survival; CSS, cancer-specific survival; or antidiahotic tractments for	: HbAIc, hemoglobin AIc. PEC and CCC					
		) ) ) )					
Types of medication	No adjustment HR (95% Cl) p-value		Diabetic duratior HR (95% CI) p-value	n adjusted <sup>a</sup>	Multivar HR (95% <i>p</i> -value	iable adjusted <sup>t</sup> S CI)	
	PFS	css	PFS	CSS	PFS	CSS	
-		_					-

Types of medication	No adjustment HR (95% CI) p-value		Diabetic duratior HR (95% CI) p-value	ı adjusted <sup>a</sup>	Multivariable adju HR (95% CI) <i>p</i> -value	isted <sup>b</sup>
	PFS	CSS	PFS	CSS	PFS	CSS
Oral antidiabetic drugs vs Insulin	0.921 (0.450–1.886) <i>p</i> =0.822	0.677 (0.325–1.411) p=0.298	1.137 (0.542–2.386) b=0.735	0.863 (0.406–1.834) p=0.702	1.113 (0.500–2.477) b=0.793	0.844 (0.368–1.940) b=0.690
Oral antidiabetic drugs vs	0.868 (0.439–1.717) p=0.685	1.113 (0.532–2.326) p=0.776	0.602 (0.297–1.220) h=0 159	0.815 (0.387-1.717) p=0.590	0.623 (0.295–1.316) b=0.215	1.087 (0.491–2.409) b=0.837
Insulin vs nonmedication	0.953 (0.433–2.097) p=0.904	l.503 (0.641–3.527) p=0.349	p=0.093	0.785 (0.317-1.943) p=0.601	p=0.158 p=0.158	p=0.998 p=0.998
<b>Notes:</b> <sup>a</sup> Model adjusted diabetic dun	ation (median cutoff value: <48 and ≥	1 48 months) <sup>b</sup> Model further adjusted g	gender, age (≤60 and >60	rears), stage (I–II and III–IV), ECOG P	S (0–1 and 2–4), extranod	al sites (≤I and >I), Hans

classification (GCB and non-GCB), symptoms and LDH (≤271 and >271U/L). Abbreviations: PFS, progression-free survival; CSS, cancer-specific survival.

Cancer Management and Research 2019:11

Table 57 Fairwise comparisons of DIT-FITISK groups for FFS and CS.	Table S9	Pairwise	comparisons	of DM-PI	risk group	os for P	YFS and C	CSS
--	----------	----------	-------------	----------	------------	----------	-----------	-----

Log-rank test (Mantel–Cox)	DM-PI risk groups	Low risk (DM-PI 0-1)		Intermediate risk (DM-PI 2–3)		High risk (DM-PI 4–6)	
		Chi-square	p-value	Chi-square	p-value	Chi-square	p-value
PFS (overall p<0.001)	Low risk (DM-PI 0–1) Intermediate risk (DM-PI 2–3) High risk (DM-PI 4–6)	- 51.951 84.334	_ <0.001 <0.001	51.951 - 12.027	<0.001 - 0.001	84.334 12.027 -	<0.001 0.001 -
CSS (overall p<0.001)	Low risk (DM-PI 0–1) Intermediate risk (DM-PI 2–3) High risk (DM-PI 4–6)	- 39.720 107.797	- <0.001 <0.001	39.720 - 25.919	<0.001 - <0.001	107.797 25.919 	<0.001 <0.001 -

Abbreviations: DM-PI, prognostic index including diabetes mellitus; PFS, progression-free survival; CSS, cancer-specific survival. Note: p-values in bold < 0.05.



Figure SI Flowchart of the study population.

Abbreviations: DLBCL, diffuse large B-cell lymphoma; PFS, progression-free survival; CSS, cancer-specific survival.



Figure S2 Kaplan-Meier curves of PFS and OS (all-cause mortality) stratified by diabetic status before and after propensity matching. (A) PFS in unmatched (complete) dataset. (B) OS in unmatched (complete) dataset. (C) PFS in propensity score-matched (1:1) dataset. (D) OS in propensity score-matched (1:1) dataset. **Abbreviations:** PFS, progression-free survival; OS, overall survival.

**Cancer Management and Research** 

**Dove**press

Publish your work in this journal

Cancer Management and Research is an international, peer-reviewed open access journal focusing on cancer research and the optimal use of preventative and integrated treatment interventions to achieve improved outcomes, enhanced survival and quality of life for the cancer patient. The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http://www.dovepress.com/testimonials.php to read real quotes from published authors.

Submit your manuscript here: https://www.dovepress.com/cancer-management-and-research-journal