

Risk Stratification Of Diffuse Large B-Cell Lymphoma With Interim PET/CT By Combining Deauville Scores And International Prognostic Index

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Purpose: Prognostic evaluation using interim positron emission tomography/computed tomography (interim PET/CT; I-PET) remains controversial. For any predictor, the prognosis of patients around its cutoff value is most uncertain. If the patients around the cutoff value could be subdivided by another factor, like the international prognostic index (IPI), it may improve the predictive power of I-PET. The combination of I-PET and IPI for risk stratification of patients was explored in this study.

Patients and methods: One hundred and eleven diffuse large B-cell lymphoma (DLBCL) patients treated with R-CHOP therapy were included retrospectively, 59 of whom underwent PET/CT after three or four cycles of treatment (I-PET). Fifty-two patients underwent PET/CT after five or six cycles of treatment (end of treatment; E-PET).

Results: When Deauville 5-point scale (5-DS) scores of 4–5 were classified as a positive scan (denoted by DS [score 4]), there was no significant difference in progression-free survival (PFS) between I-PET positive and negative patients ($P=0.151$). Further, patients with 5-DS score 3 and high IPI were stratified into I-PET positive-, whereas those with 5-DS score 3 and low IPI were classified into I-PET negative scan groups. Under this stratification, there was a significant difference in PFS between I-PET positive and negative patients ($P=0.001$). The sensitivity, positive predictive value, and negative predictive value for 2-year PFS for the combination score were higher than DS (score 4) alone (66.7% vs 33.3%, 50.0% vs 37.5%, 93.6% vs 88.2%) whereas specificity was almost the same (90.0% vs 88.0%).

Conclusion: Subdivision of patients with 5-DS score 3 by IPI improved prognostic prediction accuracy. The IPI adds strength to 5-DS in I-PET to detect patients with good or poor prognosis. Compared with other combinations of I-PET and IPI, dividing the patients around the cutoff value of 5-DS by IPI was easily accepted by clinicians and allowed them to decide on further treatment practically.

Keywords: Deauville 5-point scale, interim positron-emission tomography/computed tomography, IPI, cutoff value, subdivision

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Introduction

Diffuse large B-cell lymphoma (DLBCL) is the most common type of aggressive lymphoma. Some patients can be cured after standard treatment, but there remains up to 30–40% of the patients who suffer relapses that are refractory to first-line immunochemotherapy.¹ Early identification of such patients is very important in clinical treatment decisions. Treatment continuation or even de-escalation can be performed in patients with good prognosis to reduce toxicity without adversely affecting outcome.

Treatment intensification or modification can be proposed in those patients with poor prognosis to improve cure rates.²

The International Prognostic Index (IPI) based on age, performance status, lactate dehydrogenase (LDH) level, number of extranodal sites, and lymphoma stage is the most widely adopted prognostic tool for patients with DLBCL. However, within each IPI score, patient outcomes are still heterogeneous.³

Another method for assessment of the treatment response and prognosis of DLBCL patients is by positron emission tomography-computed tomography (PET/CT) after a few cycles of (immuno) chemotherapy, which is also known as interim FDG-PET (I-PET).^{4,5} Compared with the reported high predictive accuracy of end of treatment PET/CT (E-PET), an early response assessment using I-PET is more controversial due to variable results.^{2,6}

The Deauville 5-point scale (5-DS)⁷ is the most widely adopted criteria for both I-PET and E-PET.⁸ Meignan et al⁷ reported the consensus guideline for 5-DS classification but did not define the cutoff value of 5-DS for classification of positive or negative scans. In previous studies,^{9–12} 5-DS of scores of 1–3 were defined as negative scans, and scores of 4–5 were considered to be a positive result in both I-PET and E-PET. Kim et al⁸ evaluated the ability of I-PET to identify risk using different cutoff values of 5-DS and showed that a cutoff score of 5 at I-PET had an alternative clinical implication for identifying patients at significant risk.

IPI reflects pre-treatment factors relating to prognosis, whereas the I-PET result relates to chemosensitivity. 5-DS criteria were also combined with IPI to improve the predictive accuracy in some studies. Chow et al¹³ reported I-PET with IPI as a nomogram based on a bootstrapped Cox regression model to improve predictive accuracy. However, the cutoff value of this nomogram was not given, so it is hard for clinicians to discriminate between good- or poor-outcome patients for further treatment options. Nols et al¹⁴ reported that combination of the qualitative and quantitative indices of I-PET with IPI was highly predictive of outcome in DLBCL. This type of combination divided patients into three groups, double positive, where two indices are positive; double negative, where two indices are negative; and a group with only one positive index. For patients where only one index was positive, the prognosis (eg, 2-year progression-free survival (PFS) of I-PET- or IPI-positive patients were 60% and 69%, respectively) was usually between that of the double-positive (2-year 0% PFS) and double-negative (2-year 88% PFS) groups. But Nols et al¹⁴ did not classify this

group of patients as either a positive group with poor prognosis or a negative group with good prognosis; thus, it was still difficult for clinicians to decide on further treatment of these patients.

Therefore, it is clear that these two methods of combining I-PET and IPI were not practical for informing treatment decisions. For any predictor, the prognosis of patients around its cutoff value was most uncertain. Patients with 5-DS scores of 3 showed uptake intensity between the mediastinum and the liver, and were considered to have partially progressed. According to the results of a meta-analysis,⁶ if these patients were classified as positive, then the sensitivity of I-PET for prognosis increased but the specificity decreased. Conversely, if these patients were classified as negative, then sensitivity decreased but specificity increased. This illustrates that patients with 5-DS scores of 3 are heterogeneous, with differing prognoses. Subdividing patients with 5-DS scores of 3 based on other predictive factors may improve predictive accuracy.

In our study, subdivision of patients with 5-DS scores of 3 according to the International Prognostic index (denoted DS+IPI) was considered. DS+IPI combined the features of DS and IPI, resulting in an I-PET negative group consisting of patients with DS scores 1–2 or 3 but low IPI, and an I-PET positive group consisting of patients with DS scores 4–5 or 3 but high IPI. This combination of I-PET and IPI can classify patients with good or poor prognosis explicitly. The present retrospective study was undertaken to assess the clinical value of DS+IPI in predicting the prognosis of patients with DLBCL.

Materials And Methods

Patients

Our investigation was approved by the ethics committee of Shanxi Medical University (Taiyuan, China, ID number: 2018033). Informed consent was not obtained since this was a retrospective study. All personal data were strictly confidential. The study protocol conformed to the ethical guidelines of the Declaration of Helsinki. Our retrospective study included patients with DLBCL visiting the Tumor Hospital of Shanxi Province between April 2015 and April 2017. All patients underwent PET/CT imaging during the treatment or post-treatment period. For each patient, 2 weeks elapsed from the end of the preceding treatment to the scan. Inclusion criteria were as follows: 1) newly diagnosed with DLBCL by histopathological and

immunohistochemical examination; 2) treated with 6–8 cycles of R-CHOP with a treatment course of 21 days; 3) received at least one 18F-FDG PET/CT examination; and 4) other malignant tumors or chronic diseases were not identified during the follow-up period. In total, 111 patients were included, of whom 59 underwent PET/CT after 3–4 cycles of treatment (I-PET) and 52 underwent PET/CT after 5–6 cycles of treatment (end of treatment, E-PET). Clinical data obtained from all patients prior to treatment included sex, age, ECOG performance status, number of extranodal sites of involvement, Ann Arbor stage, and serum LDH level, allowing calculation of the IPI.

Clinical follow-up was carried out for all patients once every three months, including histopathological examination and imaging, such as ultrasound. PFS was calculated from the date of diagnosis to first relapse or progression, death of any cause, or last follow-up date. The range of follow-up time was 4–83 months. The median follow-up time was 24 months.

18F-FDG PET/CT Acquisition

All patients underwent ^{18}F -fluoro-2-deoxy-D-glucose (^{18}F -FDG) PET/CT imaging on a Discovery STE PET/CT system (GE Healthcare, USA), consisting of a 16-detector-row CT scanner. The ^{18}F -FDG PET/CT imaging agent was produced by the PET/CT center at the Tumor Hospital of Shanxi Province using ^{18}F produced with a MINitrace drug synthesis system (GE Healthcare, USA). Eventually, imaging agents were synthesized using a Trace Lab FXN multifunctional composite device. The chemical purity of the drug was greater than 90%.

Patients fasted for at least 4 hrs before intravenous administration of ^{18}F -FDG (5–6 MBq/kg) to ensure a serum glucose level below 11 mmol/L. PET/CT scanning was acquired at 1 hr after ^{18}F -FDG administration. Transmission data were acquired using a low-dose CT scan (120 kV, 180 mA, a 512×512 matrix, 3.75-mm slice thickness), and the scanning range was from vertex to the upper segment of the femur. PET images were acquired by the 3D model, and the scanning range was the same as for CT. Six to eight bed-boards were collected according to the different heights of patients, with one bed-scanning taking 3 mins. The images were reconstructed using a conventional iterative algorithm (OSEM). Multiplanar reformed PET and CT images were analyzed frame by frame on a Xeleris workstation (GE Healthcare, USA).

PET Scan Interpretation

PET/CT scans were read by two trained imaging physicians. If there was a dispute regarding any given image, it was read by the director and agreement was reached after discussion. Visual analysis of the image was adopted in this study. 5-DS criteria in the National Comprehensive Cancer Network clinical guidelines, 2016,¹⁵ were used to assess treatment response qualitatively as follows: score 1, no residual disease; score 2, uptake less than or equal to the mediastinal blood pool; score 3, uptake more than the mediastinal blood pool but less than or equal to the liver uptake; score 4, uptake was moderately greater than that of the liver; score 5, uptake markedly increased compared with the liver or new lesions were detected. Clinically, it is necessary to stratify patients who performed to I-PET into negative scan or positive scan, which indicates a good or bad outcome for informing treatment decisions. But cutoff value needs to be explored. In previous studies,^{9–12} 5-DS of scores of 1–3 were defined as negative scans, and scores of 4–5 were considered to be a positive result.

Statistical Analysis

PFS was described by median and range. Qualitative data were described by proportion and comparisons of qualitative data were performed by Chi-square test. Distribution of PFS was estimated by the Kaplan–Meier method, and comparisons of PFS were analyzed by log-rank test. Univariate and multivariate Cox's proportional-hazard model was used to estimate the hazard ratio and confidence interval of clinical prognostic factors. Data were analyzed using SPSS software (IBM Corp, Version 20.0, USA).

Results

Description Of Patient Characteristics

Patient demographics, clinical characteristics and 2-year progression status are displayed in [Tables 1](#) and [2](#).

As shown in [Table 1](#), 8 of 59 patients (12 undergoing I-PET after 3 and 47 after 4 cycles, respectively) showed disease progression or recurrence. Of these 59 patients, 31 patients (52.5%) were in advanced-stage disease (III/IV). Patients with high/medium-high IPI status accounted for 35.6%. The proportion of high-risk patients in this study with 5-DS of 4–5 was 13.6% after I-PET measurement. According to the log-rank test, there was a statistical difference in PFS for IPI. There was no statistical difference in PFS when age, gender, and stage were taken into

Table 1 Characteristics And 2-Year PFS Of 59 Patients Assessed By Interim PET/CT Scan After 3 Or 4 Cycles Of Therapy

Variable	Value	Progression/n	2-Year PFS (%)	X ²	P
Age (year)	>60	4/25	79.0	0.056	0.813
	≤60	6/34	79.8		
Gender	Female	4/34	83.9	1.193	0.275
	Male	6/25	73.1		
IPI	High (IPI=3,4,5)	7/21	58.6	5.161	0.023
	Low (IPI=0,1,2)	3/38	93.9		
Stage	High (Stage=3,4)	8/31	72.8	2.161	0.142
	Low (Stage=1,2)	2/28	91.5		
I-PET	Positive (DS=4,5)	3/8	57.1	2.063	0.151
	Negative (DS=1,2,3)	7/51	83.4		
Stime (months)	Median (min-max)	-	23 (4–76)	-	-
Progress or not	Progress	-	16.9 (10/59)	-	-

Abbreviations: 2-year PFS, two-year progression-free survival rate; IPI, International Prognostic Index; I-PET, interim positron emission tomography/computed tomography; Stime, survival time.

Table 2 Characteristics And 2-Year PFS Of 52 Patients Assessed By PET/CT Scan After 5 Or 6 Cycles Of Therapy

Variable	Value	Progression/n	2-Year PFS (%)	X ²	P
Age	>60	5/21	79.5	0.119	0.730
	≤60	6/31	76.5		
Gender	Male	5/23	78.2	0.059	0.808
	Female	6/29	69.0		
IPI	High (IPI=3,4,5)	8/22	49.1	6.598	0.010
	Low (IPI=0,1,2)	3/30	78.9		
Stage	High (Stage=3,4)	9/32	70.3	1.185	0.178
	Low (Stage=1,2)	2/20	94.1		
E-PET	Positive (DS=4,5)	6/11	34.6	11.46	0.001
	Negative (DS=1,2,3)	5/41	91.7		
Stime (Months)	Median (min-max)	-	22 (5–83)	-	-
Progress or not	Progress	11/52	21.2	-	-

Abbreviations: 2-year PFS, two-year progression-free survival rate; IPI, International Prognostic Index; I-PET, interim positron emission tomography/computed tomography; Stime, survival time; min, minimum; max, maximum.

account. Similar findings were identified for I-PET with 5-DS of 4–5 ($P=0.151$).

As shown in Table 2, 5 patients underwent E-PET after 5 cycles of treatment and 47 patients after 6 cycles. There were no same patients underwent I-PET and E-PET. Eleven of 52 patients experienced progression or recurrence. According to the log-rank test, there was no statistical difference in PFS when age, gender, and stage were accounted for. There was a statistical difference in PFS between high and low IPI ($P=0.01$). The 2-year PFS was statistically different ($P=0.001$) between positive and

negative E-PET scans. The 2-year PFS was 34.6% in the positive group and 91.7% in the negative group, in which 5-DS ≥ 3 was considered to be positive.

Predictive Values Of I-PET For The Prognosis Of DLBCL

The patients with 5-DS of 4–5 denoted by DS (score 4) were classified as having positive scans and were commonly used in the previous studies.^{9–12} There was no significant difference in PFS between negative and positive I-PET scans by DS (score 4) (Log-rank test, $P=0.151$).

Kaplan–Meier curves for all patients with 5-DS scores 1–2, score 3, and score 4–5 are displayed in Figure 1A. The PFS curves of all patients are displayed in two clustered prognostic groups. PFS was identical between those with 5-DS score 3 and those with scores of 4–5. However, patients with a score of 3 in I-PET showed inferior PFS compared with those with scores of 1–2.

Of 9 patients with 5-DS score 3, there were 5 with low IPI values and 4 with high IPI values. Of the 5 low IPI patients, no patient experienced progression, with a 2-year PFS of 100.0%, whereas 3 of the 4 patients with high IPI experienced progression, leading to a 2-year PFS of 25.0%, which is shown in Table 3. We further subdivided patients with 5-DS score 3 as positive I-PET scan when IPI was high and negative I-PET scan when IPI was low. This group division criterion combined 5-DS and IPI, resulting in the I-PET negative group consisting of patients with 5-DS score 1–2 or 3 but low IPI, while the I-PET-positive group consisted of patients with 5-DS score 4–5 or 3 but high IPI. Under this group division (denoted by DS+IPI), there was a significant difference in PFS between I-PET-positive and I-PET-negative patients ($P=0.001$), which is shown in Table 4. The 2-year PFS of the positive and negative scan groups was 95.3% and 45.5%, respectively (HR=6.36 (95% CI [1.79, 22.60]), derived by univariate Cox regression). Compared with DS (score 4), with the cutoff value at score 4, the combination

criteria of 5-DS and IPI were more sensitive in separating good from poor prognosis patients.

The predictive values of I-PET, according to different cutoff values, are summarized in Table 5. When 5-DS scores of 4–5 were classified as positive PET scans, denoted by DS (score 4), 8 positive cases (13.6%) were detected among 59 patients. Among these positive cases, 37.5% (3/8) experienced progression in 2 years. Negative predictive values of DS (score 4) were 88.2% (45/51), and area under curve (AUC) was 0.62 (0.40–0.84). For the combined index DS+IPI, the number of positive patients increased to 12, and of these 12 patients, 6 experienced progression in 2 years. The positive predictive value of DS+IPI was 50.0%. Among 47 patients with negative PET scans, only 3 experienced progression. In terms of predictive indices, sensitivity, the positive predictive values (PPV) and negative predictive values (NPV) of DS+IPI were all higher than DS (score 4) (66.7% vs 33.3%, 50.0% vs 37.5%, 93.6% vs 88.2%), whereas specificity was almost same as DS (score 4) (88.0% vs 90.0%). Compared with DS (score 4), the predictive accuracy of DS+IPI was greatly improved with AUC=0.77 (0.58–0.97).

Age, gender, IPI, stage, and DS+IPI were included in multivariate Cox regression models to evaluate significant risk factors for PFS for 59 patients, which are summarized in Table 6. As a result, only DS+IPI was found to be associated with PFS in multivariate analysis ($P=0.004$).

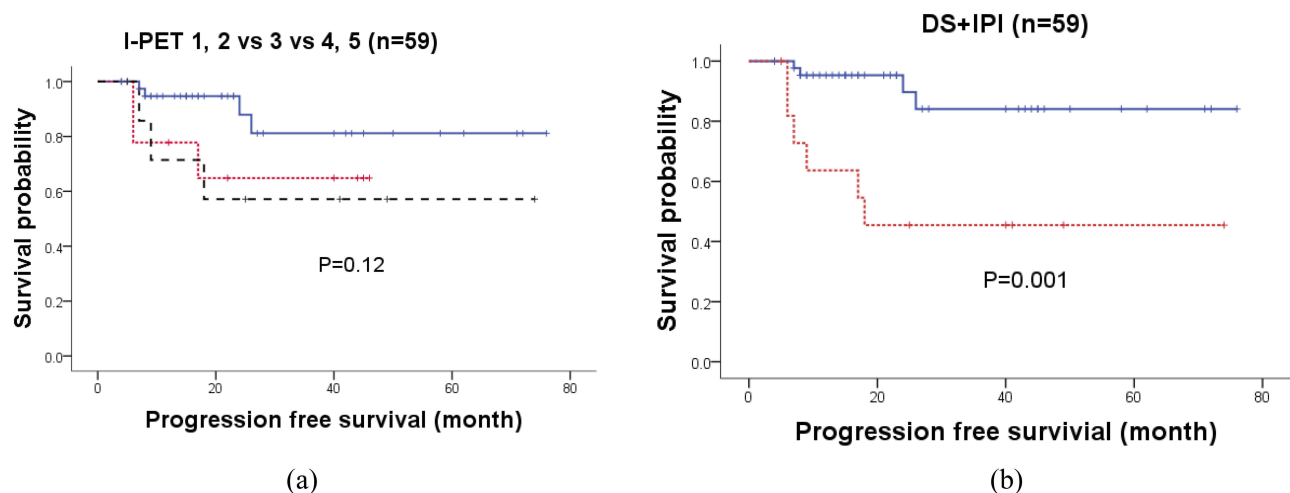


Figure 1 Kaplan–Meier curves for PFS of I-PET. **(A)** Kaplan–Meier curves for PFS of I-PET with cutoff at 1–2 vs 3 vs 4–5 for 59 patients treated by R-CHOP. The line represents 5-DS score 1 and 2, while the dashed line represents 5-DS score 3 and the dotted line represents 5-DS score 4 and 5. Log-rank test=4.25, $P=0.12$, HR=1.68 (95% CI [0.89, 3.19]). **(B)** Kaplan–Meier curves for PFS of patients stratified by the combination of I-PET and IPI. The line represents I-PET negative scan with 5-DS of 1–2 or 5-DS score 3 but low IPI, while the dotted line represents PET/CT positive scan with 5-DS score 4–5 or DS score 3 but high IPI. Log-rank test=10.86, $P=0.001$, HR=6.36 (95% CI [1.79, 22.56]).

Abbreviations: PFS, progression-free survival; I-PET, interim PET/CT; HR, hazard ratio; CI, confidence interval; IPI, International Prognostic Index; 5-DS, Deauville 5-point scale.

Table 3 Prognosis Of 9 Patients By Interim PET/CT Scan With 5-DS Of Score 3

Variable	Value	Progression/n	2-Year PFS (%)	P
IPI	High (IPI=3,4,5)	3/4	25.0	0.023
	Low (IPI=0,1,2)	0/5	100.0	

Abbreviations: 2-year PFS, two-year progression-free survival rate; IPI, International Prognostic Index.

Predictive Capacity Of E-PET For Prognosis Of DLBCL

Fifty-two patients underwent E-PET after five or six cycles of therapies in our study. When 5-DS of 4–5 were classified as having positive scans, there was a significant difference in PFS between patients with negative E-PET scans and those with positive E-PET scans (Log-rank test, $P=0.001$, HR=6.10 (95% CI [1.84–20.21]), derived by Cox regression). Kaplan–Meier curves for E-PET are shown in Figure 2. The predictive performance of E-PET for 2-year PFS is summarized in Table 7.

Age, gender, IPI, stage, and E-PET were included in multivariate Cox regression, which is summarized in Table 8. Only E-PET was found to be associated with PFS. A positive E-PET result had a higher risk of disease progression.

Discussion

Nowadays, 5-DS criteria for interpretation of response evaluation with PET/CT have been widely adopted in DLBCL.^{9–12,16} It represents simple, reproducible criteria

for I-PET interpretation.⁷ Barrington et al¹⁷ reported that validated 5-point London Criteria, which is the antecedent of the Deauville scoring system, were sufficiently robust to be used in a multicenter setting, both for the conservative (score 3 interpreted as “negative”) and sensitive readings (score 3 interpreted as “positive”). Another visual criteria for interpretation of ¹⁸F-FDG PET is the International Harmonization Project (IHP) which defines a positive FDG-PET scan as FDG uptake that is more intense than mediastinal blood pool activity in lesions larger than 2 cm and more than background in lesions smaller than 2 cm.¹⁸ That is, the cutoff value of considering as positive result by IHP is lower than DS (score 4). Some patients with score 3 of 5-DS criteria, which uptake more than the mediastinal blood pool but less than or equal to the liver uptake, would be divided into positive scan according to IHP criteria.

A recent meta-analysis⁶ of 20 studies comprising 2255 patients was conducted to assess the predictive value of visually assessed I-PET. Of 20 studies, 5-DS were used in 12 studies and IHP criteria were adopted in four. Other visual criteria which were not based on consensus guidelines were used in the four remaining studies. Burggraaff et al⁶ reported that higher sensitivity values seen in the ROC analyses for both IHP and custom criteria vs 5-DS might be explained by the lower threshold of test positivity with IHP vs 5-DS (using blood and liver pool activities as the reference tissues, respectively). To evaluate sensitivity and specificity comprehensively for 2-year PFS, we furthermore derived the Yoden Index (Yoden index=sensitivity+specificity-1) of 19 available studies in Burggraaff et al.⁶ The Yoden index of three studies (except one study without

Table 4 Univariate Analysis For PFS By Different Cutoff Value Of I-PET Scan

Variable	Value	Progression/n	2-Year PFS(%)	P	HR
DS (score 4)	Positive (DS=4,5) Negative (DS=1,2,3)	3/8 7/51	57.1 83.4	0.151	2.59 [0.67, 10.06]
DS+IPI	Positive (DS=4,5 or DS=3 and IPI=positive) Negative (DS=1,2 or DS=3 and IPI=negative)	6/12 4/47	45.5 95.3	0.001	6.36 [1.79, 22.60]

Abbreviations: 2-year PFS, two-year progression-free survival rate; IPI, International Prognostic Index; DS (score 4), Deauville 5-point scale (5-DS), scores of 4–5 were classified as a positive scan; DS+IPI, Combination of 5-DS and IPI in the way of dividing patients with 5-DS score 3 by IPI.

Table 5 Predictive Accuracy Of 5-DS (Score 4) And DS+IPI For 2-Year PFS

PET/CT	Sensitivity (%)	Specificity (%)	PPV (%)	NPV (%)	AUC (95% CI)
DS (score 4)	33.3 (3/9)	90.0 (45/52)	37.5 (3/8)	88.2 (45/51)	0.62 (0.40–0.84)
DS+IPI	66.7 (6/9)	88.0 (44/50)	50.0 (6/12)	93.6 (44/47)	0.77 (0.58–0.97)

Abbreviations: PPV, positive predictive value; NPV, negative predictive value; PFS, progression-free survival; DS (score 4) indicates DS≥4 as positive; DS+IPI, 5-DS of 1 to 2 as negative, 4 to 5 as positive, score 3 but high IPI as positive, score 3 but low IPI as negative; AUC, area under the curve.

Table 6 Multivariate Analysis For PFS In Patients With I-PET

Variables	B	Wald χ^2	P	HR	95% CI
DS+IPI	1.85	8.19	0.004	6.36	[1.79, 22.60]

Abbreviations: IPI, International Prognostic Index; DS+IPI, Combination of 5-DS and IPI in the way of dividing patients with 5-DS score 3 by IPI; HR, hazard ratio; CI, confidence interval.

reported sensitivity and specificity) with IHP criteria ranged from 28% to 42% and ranked 5th, 10th, and 13th of 19 studies from high to low according to the Yoden index. The 12 studies using 5-DS ranged from 13% to 58% in the Yoden index, indicating that the predictive accuracy of the IHP criteria was comparable with the Deauville system. The patients with 5-DS scores of 3 were those with uptake intensity between that of the mediastinum and the liver, which are considered to represent partial progression. According to the results of meta-analysis,⁶ if they were classified as positive, then sensitivity of I-PET for prognosis increased but specificity decreased. However, if they were classified as negative scans, then sensitivity decreased but specificity increased. This illustrates that patients with 5-DS scores of 3 are heterogeneous with differing prognoses. To subdivide patients with score 3 based on other predictive factors may improve the predictive accuracy.

In this study, for I-PET, there was no significant difference in PFS between I-PET positive and negative patients using the cutoff value of I-PET (5-DS score 4). The IPI is a well-established tool for the prediction of prognosis according to

pretreatment characteristics. In this study, patients with 5-DS score 3 were subdivided into I-PET-positive when their IPI was high, and I-PET-negative when their IPI was low. Under this subgroup division, there was a significant difference in PFS between I-PET-positive and I-PET-negative patients ($P=0.001$). Its sensitivity for 2-year PFS was higher than DS (score 4) (66.7% vs 33.3%), whereas its specificity was almost identical to DS (score 4) (88.0% vs 90.0%). Furthermore, the PPV and NPV of this combination were both higher than DS (score 4) (50.0% vs 37.5%, 93.6% vs 88.2%). Importantly, subdivision of patients with 5-DS score 3 by IPI improved the prediction accuracy.

In a previous study, Chow et al¹³ combined I-PET with IPI as a nomogram based on a bootstrapped Cox regression model. However, the cutoff value of this nomogram was not given, making it difficult for clinicians to discriminate between good- or poor-outcome patients. The clinical implications of the model were also hard for a clinician to understand. Nols et al¹⁴ divided patients into three groups: 1) double I-PET and IPI positive; 2) double I-PET and IPI negative; and 3) a group with only one positive index. Nols et al¹⁴ did not classify the group of patients for whom only one index was positive as either a positive group with poor prognosis or a negative group with good prognosis. Thus, it was still difficult for clinicians to decide on further treatment of these patients. Jiang et al¹⁹ divided patients into different risk groups using IPI and further subdivided them into risk groups using interim PET/CT. However, the authors encountered the same problem as Nols et al¹⁴ among patients with only one positive index (only IPI-positive or I-PET-positive); it was difficult for clinicians to discriminate good- or poor-outcome patients. Compared with other combinations of I-PET and IPI, dividing patients based on the cutoff value of 5-DS by IPI was easily accepted by clinicians and practical when considering further treatment options.

Instead of DS scores of 4–5 which are usually considered to be a positive result (denoted by DS (score 4)), Kim et al⁸ showed that a cutoff score of 5 by I-PET (denoted by DS (score 5)) had alternative clinical implications for identifying patients at significant risk. Changing the cutoff values of DS would result in increases in some predictive indices, but decreases in others. Kim et al⁸ found that compared with DS (score 4), DS (score 5) had higher specificity (96.4% vs 74.8%), but sensitivity decreased by more than 10% (26.6% vs 38.5%). That indicated that more than 10% of the poor-outcome patients could not be detected, which could confound further treatment decisions. Compared with DS (score

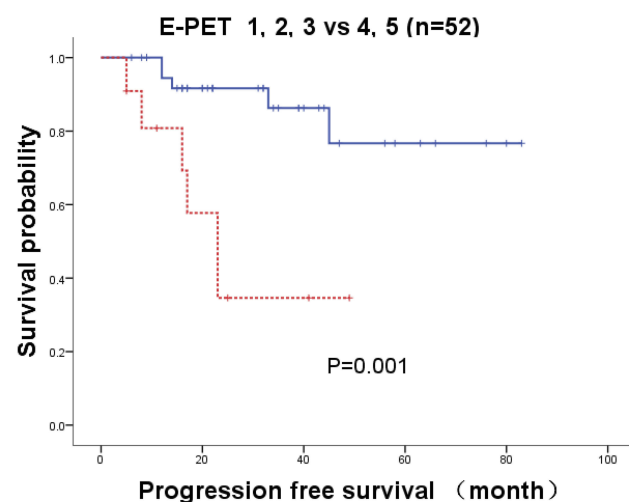


Figure 2 Kaplan-Meier curves for PFS of E-PET. Solid line represents negative E-PET scan with 5-DS score 1–3; dotted line represents positive E-PET scan with 5-DS score 4–5. Log-rank test =11.46, $P=0.001$, HR=6.10 (95% CI[1.84–20.21]). **Abbreviations:** PFS, progression-free survival; E-PET, patients underwent PET/CT after five or six cycles of treatment; HR, hazard ratio; CI, confidence interval.

Table 7 Predictive Accuracy Of E-PET For 2-Year PFS (n=52)

Variable	Sensitivity (%)	Specificity (%)	PPV (%)	NPV (%)	AUC (95% CI)
E-PET	66.7 (6/9)	88.4 (38/43)	54.5 (6/11)	92.7 (38/41)	0.78 (0.58–0.87)

Abbreviations: E-PET, patients underwent PET/CT after five or six cycles of treatment. AUC, area under the curve; PPV, positive predictive value; NPV, negative predictive value; PFS, progression-free survival; AUC, area under the curve.

Table 8 Multivariate Analysis Of Prognosis For Patients With E-PET (n=52)

Variables	B	Wald X ²	P	HR	95%CI
E-PET	1.81	8.75	0.003	6.10	1.84–20.21

Abbreviations: E-PET, patients underwent PET/CT after five or six cycles of treatment; HR, hazard ratio; CI, confidence interval.

4), we found that the predictive accuracy of DS (score 5) was not improved in our study, achieving slightly higher specificity (96.0% vs 90.0%) at the cost of much lower sensitivity (11.1% vs 33.3%). However, the sensitivity of DS+IPI was much higher than DS (score 4) (66.7% vs 33.3%), whereas the specificity was nearly the same (88.0% vs 90.0%). Thus, IPI adds value to DS in I-PET, and DS+IPI was superior to changing the cutoff values of DS for detection of patients with good or poor prognosis.

Meignan et al⁷ reported the consensus guideline of 5-DS classification but did not define the cutoff value of 5-DS for classification of positive or negative scans. They reported that for 5-DS of scores 2–4, correction methods using the Δ SUVmax should be investigated. Δ SUVmax²⁰ and SUVmax-liver²¹ at I-PET have been reported to be a more sensitive method of separating good- from poor-outcome patients. However, it may be less practical because of the demanding conditions required for accurate and reproducible results.²² Further validation is needed to determine whether such semiquantitative factors add strength to 5-DS to stratify patients accurately.

Some limitations of our study are worth noting. This is a single-center, retrospective study with relatively small sample size. However, our data suggest that combination of 5-DS score with IPI can improve the predictive value of PET/CT in terms of patient prognosis. The combined criteria of I-PET and IPI should be refined in prospective trials and may help inform clinical decisions in treatment of DLBCL.

Conclusion

In conclusion, in the way of dividing the patients around the cutoff value of 5-DS with I-PET by IPI, we were able to discriminate between DLBCL patients with good or poor

prognosis reliably. This approach was easily understood by clinicians and allowed them to decide on further treatment practically. Moreover, not only do this approach inform prognosis and further treatment, it provides a new way of combining I-PET criteria and other predictors to improve the predictive ability for the prognosis of DLBCL.

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Disclosure

The authors report no conflicts of interest in this work.

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