Combined PET and Biopsy Evidence of Marrow Involvement Improves Prognostic Prediction in Diffuse Large B-Cell Lymphoma

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Bone marrow is an important extranodal site in diffuse large B-cell lymphoma (DLBCL), and marrow histology has been incorporated into the new National Comprehensive Cancer Network international prognostic index. Marrow involvement demonstrated histologically confers poor prognosis but is identified by staging PET in more cases. How information from staging PET and biopsy should be combined to optimize outcome prediction remains unclear. Methods: The International Atomic Energy Agency sponsored a prospective international cohort study to better define the use of PET in DLBCL. As a planned subsidiary analysis, we examined the interplay of marrow involvement identified by PET and biopsy on clinical outcomes. Results: Eight countries contributed 327 cases with a median follow-up of 35 mo. The 2-y outcomes of cases with no evidence of marrow involvement (n = 231) were 81% (95% confidence interval [CI], 76%-86%) for event-free survival (EFS) and 88% (83%-91%) for overall survival (OS); cases identified only on PET (n = 61), 81% (69%-89%) for EFS and 88% (77%-94%) for OS; cases identified only on biopsy (n = 10), 80% (41%-95%) for EFS and 100% for OS; or cases identified by both PET and biopsy (n = 25), 45% (25%-64%) for EFS and 55% (32%-73%) for OS. The hazard ratios for PET-negative/biopsy-negative cases versus PET-positive/biopsy-positive cases were 2.67 (95% CI, 1.48-4.79) for EFS and 3.94 (1.93-8.08) for OS. Conclusion: This large study demonstrates that positive iliac crest biopsy histology only confers poor prognosis for patients who also have abnormal marrow 18F-FDG uptake identified on the staging PET scan. Abnormal 18F-FDG uptake in marrow, when iliac crest biopsy histology is normal, has no adverse effect on outcomes. Key Words: positron emission tomography; diffuse large B-cell lymphoma; bone marrow; International Prognostic Index

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At the same time, the authors highlighted the need for further studies to determine the relative role of PET and bone marrow biopsy (BMB) in determining prognosis (13).

We have recently completed a prospective multinational cohort study, sponsored by the International Atomic Energy Agency (IAEA), to investigate the application of PET for risk assessment of DLBCL. As a planned subsidiary analysis, we investigated marrow involvement identified by staging PET and routine iliac crest biopsy for predicting treatment outcomes and to inform international practice. Our hypothesis, based on recent literature (11) and the most recent international consensus guidance on the use of PET/CT for lymphoma staging (14), was that BMB is no longer necessary in the era of routine staging by PET.

MATERIALS AND METHODS

Major cancer centers in 8 countries (in São Paulo, Brazil; Santiago, Chile; Budapest and Debrecen, Hungary; Mumbai, India; Bologna, Italy; Seoul, South Korea; Manila, Philippines; Bangkok, Thailand) from 5 United Nations–defined regions participated in the study. Recruitment commenced in 2008 through September 2011.

Eligibility Criteria

Participating clinicians recruited sequential cases of DLBCL, with patients aged 16 y or older, provided that informed consent could be obtained before chemotherapy. Exclusions were central nervous system involvement, cancer within preceding 5 y, and steroid therapy before staging scan.

Study Protocol

The protocol was developed by the study lead clinicians from each country during 2 investigator meetings. Diagnosis of DLBCL was based on histology with standard immunohistochemistry to include, at minimum, CD20, CD3, and either Ki-67 or MiB1 (15). All patients were staged by PET/CT (6 centers) or PET and CT separately (Italy and Brazil) and by iliac crest biopsy. Recommended treatment was 6 cycles of R-CHOP (rituximab with cyclophosphamide, adriamycin, vincristine, prednisolone) at 21-d intervals. To accommodate local practice, R-CHOP up to 8 cycles was permitted. Omission of rituximab was allowed in recognition that some eligible patients might otherwise be excluded for financial reasons. All patients underwent PET scans after 2–3 cycles of chemotherapy and 4–8 wk after all chemotherapy as part of the study design to assess response at nodal and extranodal sites. Treatment modification in response to the midtreatment PET scan was not permitted, except in cases with confirmed disease progression.

Identification of Marrow Disease by PET. Marrow involvement on the staging PET or PET/CT (together referred to as PET throughout the manuscript) scan was classified as focal or diffuse as previously described (6). Focal was defined as 1 or more circumscribed areas of high 18F-FDG uptake within the skeleton and diffuse as uniform increased 18F-FDG uptake throughout the bone marrow space. Diffuse 18F-FDG uptake was not considered to represent lymphoma involvement unless supported by positive biopsy histology.

The criteria for focal 18F-FDG uptake in bone marrow to be classified as due to DLBCL were as previously reported by others (11,16). 18F-FDG uptake to be greater than the intensity of uptake in normal liver, with no anatomic changes to suggest alternative benign bone pathology. Bone or marrow involvement by spread from a contiguous nonskeletal site was excluded. Patients were considered to have bone marrow involvement by lymphoma if they had either histologic DLBCL in the marrow biopsy or focal PET-positive (PET+) marrow involvement irrespective of iliac crest biopsy histology.

Histologic Identification of Marrow Disease

All cases had untargeted unilateral iliac crest biopsy. BMB histology immunohistochemistry was required to include CD20, CD3, and CD10 for accurate detection of low-level marrow involvement (15).

Classification of Events

Study events were relapse after complete remission, death from any cause, treatment escalation for progressive disease while on treatment, and disease progression or failure to achieve complete remission at the end of chemotherapy based on the revised response criteria for lymphoma (17). The protocol required suspected residual disease to be confirmed by biopsy.

Data Quality Assurance

Each patient record was reviewed with the country chief investigator during the final collaborator meeting for consistency and accuracy.

Research Regulation and Data Protection

Each country gained research ethics approval for the study protocol and patient information from the appropriate national or local Ethics Review Board. Some countries imposed unusually strict recruitment criteria, which limited recruitment in some larger centers. Fully informed consent was an inclusion criteria for recruitment. Signed consent forms are kept by the local investigators. To ensure confidentiality while sharing data internationally, cases were assigned a numeric code, and only 2 identifiers for data validation, initials and date of birth, were recorded in the central database (18).

Statistical Methods

Follow-up was continued until 75% of patients had reached 2 y or died. Cases lost to follow-up were censored at date of last known disease status. Survival was estimated using Kaplan–Meier methods. Event-free survivals (EFS) and OSs are reported. Survival times are all reported at 2 y, using date of first chemotherapy as the origin. The

| TABLE 1

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>n</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male sex</td>
<td>173</td>
<td>53%</td>
</tr>
<tr>
<td>IPI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–1</td>
<td>135</td>
<td>41%</td>
</tr>
<tr>
<td>2</td>
<td>81</td>
<td>25%</td>
</tr>
<tr>
<td>3</td>
<td>68</td>
<td>21%</td>
</tr>
<tr>
<td>4+</td>
<td>43</td>
<td>13%</td>
</tr>
<tr>
<td>Stage</td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>22</td>
<td>6%</td>
</tr>
<tr>
<td>II</td>
<td>95</td>
<td>29%</td>
</tr>
<tr>
<td>III</td>
<td>64</td>
<td>20%</td>
</tr>
<tr>
<td>IV</td>
<td>146</td>
<td>45%</td>
</tr>
<tr>
<td>Performance status</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–1</td>
<td>276</td>
<td>84%</td>
</tr>
<tr>
<td>2</td>
<td>36</td>
<td>11%</td>
</tr>
<tr>
<td>3</td>
<td>9</td>
<td>3%</td>
</tr>
<tr>
<td>4</td>
<td>6</td>
<td>2%</td>
</tr>
<tr>
<td>LDH &gt; normal</td>
<td>166</td>
<td>51%</td>
</tr>
<tr>
<td>Extranodal sites ≥ 2</td>
<td>95</td>
<td>29%</td>
</tr>
</tbody>
</table>

Median age was 55 y (quartiles, 44, 63 y).
prognostic impact of bone marrow involvement, as identified by PET or biopsy, is estimated from a Cox proportional hazards model and hazard ratios (HRs) reported (19).

Multivariate analysis was undertaken to investigate whether marrow involvement identified by PET alone (focal or diffuse combined), by biopsy alone, or by both PET and biopsy was an independent predictor of outcome. The model included the component clinical indicators from the IPI (without National Comprehensive Cancer Network modifications), that is, age (>60), stage (III/IV), performance status ($\geq 2$), LDH greater than normal, and extranodal disease in 2 sites or more. The model also includes use or omission of rituximab from therapy. Analyses and graphs were produced using Stata 12 (StataCorp.).

RESULTS

Eight countries recruited 327 cases. The number of cases from each country was 61 from Brazil, 47 from Chile, 65 from Hungary, 32 from India, 49 from Italy, 9 from South Korea, 20 from Philippines, and 44 from Thailand. Patient characteristics are listed in Table 1. Median age was 55 y, and 35% of patients were older than 60 y; 280 (86%) patients received rituximab.

Identification of Marrow Disease

Staging was by PET/CT ($n = 217$) or separate PET and CT (Italy, Brazil; $n = 110$). Eighty-two of the 327 cases (25%) were considered to have lymphoma in marrow at diagnosis; 47 of 82 (57%) were identified by PET alone, 10 (12%) by BMB alone, and 25 (30%) by both PET and BMB (Table 2).

Of 241 judged to have normal marrow on PET, 10 (4%) had positive biopsies. Of these, 2 had deposits of indolent lymphoma, 6 had a low-level infiltrate (<10%) of DLBCL cells, and only 2 had a DLBCL infiltrate of more than 10% nucleated cells in the iliac crest sample. Of 68 focal PET+ cases, 21 (31%) had lymphoma detected by biopsy. Of 18 with diffuse $^{18}$F-FDG uptake throughout skeletal marrow including the pelvis, only 4 had histologic evidence of marrow disease on iliac crest biopsy (Table 2).

Marrow Disease and Stage

In the whole cohort, 146 (45%) of the cases were stage IV. PET identified marrow involvement not identified by marrow biopsy, which upstaged 23 patients to stage IV: 1 patient from I to IV, 11 from II to IV, and 11 from III to IV; all had focal marrow $^{18}$F-FDG uptake. No patients had their stage increased by iliac crest biopsy.

The distribution of extranodal disease in the 146 stage IV patients is shown in Table 3.

Marrow Disease and Survival

Median follow-up was 35 mo. Thirty-five cases had marrow involvement identified by iliac crest histology: 2-y EFS for BMB- positive (BMB+) cases, 56% (95% confidence interval [CI], 37%–71%), and for BMB-negative (BMB−) cases, 81% (76%–85%); and OS for BMB+, 68% (49%–82%), and BMB−, 88% (83%–91%). The HR for progression-free survival was 2.22 (1.31%–3.27) (Fig. 1).

Classified by both PET and biopsy status (Fig. 2), there were 231 patients who had neither increased $^{18}$F-FDG uptake in marrow on PET nor histologic evidence of lymphoma infiltration on biopsy (PET−/BMB−) and had a 2-y EFS of 81% (95% CI, 76%–86%) and an OS of 88% (83%–91%); 61 patients who had $^{18}$F-FDG uptake on PET but a negative biopsy (PET+/BMB−) and an EFS of 81% (69%–89%) and an OS of 88% (77%–94%); and 10 patients who had marrow involvement identified only by biopsy histology.

### Table 2

<table>
<thead>
<tr>
<th>PET</th>
<th>Total patients</th>
<th>BMB−</th>
<th>BMB+</th>
<th>Marrow involved by lymphoma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative</td>
<td>241</td>
<td>231</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Positive, focal</td>
<td>68</td>
<td>47</td>
<td>21</td>
<td>68</td>
</tr>
<tr>
<td>Positive, diffuse</td>
<td>18</td>
<td>14*</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Total</td>
<td>327</td>
<td>292</td>
<td>35</td>
<td>82</td>
</tr>
</tbody>
</table>

*These 14 cases were classified as not having marrow involvement by lymphoma.

### Table 3

<table>
<thead>
<tr>
<th>Extranodal sites</th>
<th>No. of patients</th>
<th>Total</th>
<th>Single extranodal organ involved</th>
<th>≥ 2 extranodal organs involved</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone marrow only*</td>
<td>23</td>
<td></td>
<td>23</td>
<td>—</td>
</tr>
<tr>
<td>Soft tissue (non–bone marrow) only</td>
<td>64</td>
<td>28</td>
<td>28</td>
<td>36</td>
</tr>
<tr>
<td>Soft tissue and bone marrow†</td>
<td>59</td>
<td></td>
<td>—</td>
<td>59</td>
</tr>
<tr>
<td>Total</td>
<td>146</td>
<td>51</td>
<td>95</td>
<td></td>
</tr>
</tbody>
</table>

*All cases for which bone marrow was identified as the only extranodal organ involved had marrow disease identified by PET but not by biopsy.
†All cases for which marrow biopsy histology was positive had at least one other, nonmarrow extranodal organ involved—that is, had at least 2 extranodal organs involved by lymphoma.
(PET–/BMB +), representing 29% of histology positive cases, with an EFS of 80% (41%–95%) and an OS of 100%. There was no significant difference in EFS or OS between cases with no evidence of marrow disease and those with marrow involvement identified by either PET or BMB alone.

In contrast, 25 patients with marrow involvement identified by both PET and iliac crest biopsy (PET+/BMB+) had significantly inferior 2-y outcomes, with an EFS of 45% (95% CI, 25%–64%) and an OS of 55% (32%–73%). HRs for double-positive (PET+/BMB+) versus uninvolved (PET-/BMB-) bone marrow were 2.67 (1.48–4.79) for EFS and 3.94 (1.93–8.06) for OS (Fig. 2).

The most frequent pattern of marrow involvement by DLBCL on staging PET is as multiple circumscribed, focal 18F-FDG–avid deposits. Because less than a third of focal PET-positive cases had disease detected histologically by iliac crest biopsy in this and other series, we explored the prognostic effect of focal disease and whether prognosis was influenced by the presence or absence of positive histology. Of the 68 cases in our series, 47 had negative biopsies (focal PET+/BMB–), EFS of 78% (95% CI, 63%–88%), and OS of 87% (73%–94%); and 21 had positive biopsies (focal PET+/BMB+), EFS of 46% (24%–65%), and OS of 57% (31%–76%). HRs were 2.47 (1.07–5.51) for EFS and 3.03 (1.10–8.40) for OS (Fig. 3).

Multivariate analysis was performed to explore the relative influence on survival of the IPI component clinical indicators, rituximab treatment, and marrow disease detected by only PET or biopsy or both PET and biopsy (Table 4). When adjusted for each of these covariates, the effect of marrow involvement detected by both PET and BMB together had a significant independent adverse impact on OS, with an HR of 2.26 (95% CI, 1.02–5.03) (P = 0.05). Marrow involvement identified by marrow biopsy histology alone had no impact on OS, with an HR of 0.49 (0.07–3.68) (P = 0.49). Whether rituximab was part of therapy had no effect (P = 0.90).

**DISCUSSION**

The purpose of this analysis of the IAEA Lymphoma Study data was to better define the relative importance of staging PET,
compared with routine iliac crest biopsy, for establishing the prognostic impact of marrow involvement in DLBCL. The analysis was based on cases of DLBCL from 8 countries in 5 geographic regions treated to a common protocol.

The IAEA cohort of 327 patients is as large as all 3 previous studies, which examined the prognosis of marrow involvement (10–12).

In the cohort, 25% were judged to have marrow involvement by lymphoma; 88% of these were identified by PET, 42% by histology (Table 2). These proportions are similar to a recent single-center U.K. study (11).

Cases with positive marrow histology had inferior outcomes, compared with those with normal biopsies, as reported by others (10–12). However, when information from staging PET and biopsy were considered together, it was apparent that histology-positive marrow disease did not uniformly predict for poor outcome.

When both PET and biopsy data were integrated, there were 3 separate diagnostic categories of marrow involvement (Fig. 2): marrow infiltration detected by histology with negative marrow on PET (PET+/BMB−), abnormal 18F-FDG uptake on PET with negative histology (PET+/BMB−), and abnormal 18F-FDG uptake in marrow with positive histology (PET+/BMB+). Cases with marrow involvement identified by a single diagnostic modality had a 2-y EFS of 80% and 81%, respectively, similar to patients with no evidence of marrow disease irrespective of stage, with an EFS of 81%. Only cases with marrow involvement identified by both PET and biopsy histology had significantly inferior survivals, with an EFS of 45%, OS of 55%, and HRs of 2.67 and 3.94, respectively, compared with patients with no marrow involvement.

Multivariate analysis demonstrated that the inferior survival of PET+/BMB+ cases was independent of the IPI and its components and independent of marrow involvement detected by either PET or biopsy histology alone. In this model, positive marrow histology did not by itself predict for poor EFS or OS.

The highly significant adverse effect on prognosis of marrow disease identified by both PET and histology, in contrast to the lack of effect when identified by only a single modality, needs further consideration.

Twenty-nine percent (10/35) of cases with positive marrow histology had normal marrow on staging PET and survivals that were not inferior to cases without extranodal spread to marrow. It has been previously reported that a marrow infiltrate of up to 10% DLBCL cells, or by indolent lymphoma cells, may not accumulate 18F-FDG above the metabolic activity of normal

| Variable                          | OS                     | EFS                    |
|-----------------------------------|------------------------|                       |
|                                   | HR (95% CI)            |          P            | HR (95% CI) | P          |
| Age ≥ 60 y                        | 1.51 (0.82–2.77)       | 0.18                  | 0.96 (0.60–1.54) | 0.86      |
| Ann Arbor stage III/IV            | 1.16 (0.54–2.50)       | 0.70                  | 1.21 (0.70–2.08) | 0.50      |
| >1 extranodal site                | 1.27 (0.66–2.46)       | 0.47                  | 1.20 (0.73–1.96) | 0.47      |
| LDH level normal                  | 2.42 (1.26–4.67)       | <0.01                 | 2.22 (1.39–3.56) | <0.01    |
| Eastern Cooperative Oncology Group performance status ≥ 2 | 1.55 (0.79–3.05) | 0.21      | 1.56 (0.93–2.64) | 0.10    |
| PET−/BMB+                         | 0.49 (0.07–3.68)       | 0.49                  | 1.13 (0.40–3.20) | 0.83     |
| PET+/BMB−                         | 0.74 (0.33–1.66)       | 0.46                  | 0.68 (0.37–1.27) | 0.23     |
| PET+/BMB+                         | 2.26 (1.02–5.03)       | 0.05                  | 1.61 (0.84–3.10) | 0.15     |

Data in parentheses are 95% CIs.
There is evidence from histopathologic studies that a DLBCL infiltrate of 10% cells in iliac crest marrow does not impact survival (2), and indolent lymphoma in the marrow of patients with DLBCL does not influence progression risk independent of the IPI (4). It is therefore to be expected that PET−/BMB+ cases in this cohort had survivals that were not inferior to cases without identified marrow disease.

It is well recognized that diffuse 18F-FDG uptake in marrow may be increased above background due to reactive myelopoiesis in both Hodgkin and non-Hodgkin lymphoma (6,22). Differentiation of reactive from pathologic diffuse PET positivity due to lymphoma may improve with experience (11). This study confirms that cases with diffuse 18F-FDG accumulation above background in marrow should always be biopsied to establish etiology.

Hence, data from our large cohort add support to the limited existing data that a positive marrow biopsy does not add useful information for treatment planning or outcome prediction when the marrow is negative on staging PET (13). However, the experience from this international cohort leads us to disagree with the conclusion of Adams et al. (13) that diffuse bone marrow 18F-FDG uptake throughout the skeletal marrow should be regarded as marrow involvement, without the need for corroboration by biopsy.

The prognostic impact of focal 18F-FDG uptake in marrow has, until now, remained unclear. Reported series to date have had insufficient numbers to analyze the outcome of this group separately and have reached opposing conclusions as to whether focal PET disease on its own is an independent adverse prognostic indicator (10–12). With this larger cohort we have, for the first time to our knowledge, been able to demonstrate that cases with focal 18F-FDG–avid deposits on PET, without positive histology, have EFSs and OSs that do not differ significantly from DLBCL cases without spread to marrow (EFS, 78%, vs. 81%, respectively). Focal PET-positive cases with positive histology had significantly inferior survival (EFS, 46%) (Fig. 3). The likely explanation is that the probability of an untargeted biopsy sampling a metastatic lymphoma deposit increases with the extent of marrow disease. Hence, it is this, rather than marrow involvement per se, that confers poor prognosis.

Our conclusion fits with that of Khan et al. who noted that PET/CT had high sensitivity for limited marrow diseases, whereas biopsy detects only the most extensive marrow involvement (11). Their conclusion was corroborated by finding that a positive marrow biopsy was strongly associated with spread to 4 or more extranodal organ sites. As the present study recorded only 1- versus 2-or-more-organ involvement, we were unable to provide confirmation that extensive extranodal disease on staging PET obviated the need for a marrow biopsy.

The unique international nature of this prospective, protocol-driven study brings with it both strengths and weaknesses. Geographic separation prohibited the sharing of histologic material for central review. PET scans were read centrally for treatment response but not consistently for bone marrow involvement. Less experienced centers appeared to overinterpret diffuse 18F-FDG uptake in marrow as lymphoma infiltration rather than reactive myelopoiesis. Nonetheless, our results have the unusual strength of being internationally applicable.

The new data from this large cohort provide a clearer answer to the question whether to use bone marrow staging by biopsy or PET for defining prognosis in DLBCL (23). The answer appears to be that both are important but not necessary for every patient. The patient with normal marrow on staging PET is unlikely to gain prognostic information from a routine marrow biopsy. However, when the marrow is abnormal on staging PET, the presence or absence of lymphoma in iliac crest histology has an important impact on outcome. When this algorithm is used, only a quarter of patients might need a biopsy.

CONCLUSION

We have demonstrated that neither bone marrow histology nor PET alone is a reliable indicator of poor risk marrow disease. The most consistent indicator is marrow involvement identified by both PET and histology. We suggest that the algorithm we propose above should be incorporated into practice guidelines and would strengthen the new IPI (3). The strategy we outline would reduce the burden on patients and improve the precise identification of poor prognosis DLBCL internationally.

DISCLOSURE

The costs of publication of this article were defrayed in part by the payment of page charges. Therefore, and solely to indicate this fact, this article is hereby marked “advertisement” in accordance with 18 USC section 1734. The IAEA funded three collaborator meetings and a writing group meeting. Some contribution was made to each participating center to partially defray the cost of additional study PET scans. The IAEA did not influence analysis or interpretation of the data or the decision to publish. No other potential conflict of interest relevant to this article was reported.

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REFERENCES


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