LONG-TERM OUTCOME OF PATIENTS WITH PERIPHERAL T-CELL LYMPHOMA TREATED WITH FIRST-LINE INTENSIVE CHEMOTHERAPY FOLLOWED BY AUTOLOGOUS STEM CELL TRANSPLANTATION

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Background. Peripheral T-cell lymphomas (PTCL) are infrequent subtypes of non-Hodgkin’s lymphomas. The clinical course is aggressive and the median survival is about 2–3 years. An optimal first-line chemotherapy protocol has not been established and the role of high-dose therapy with autologous stem cell transplantation (ASCT) is still unclear.

Aim. To analyze the long-term outcome of unselected PTCL patients treated with intensive first-line chemotherapy with high-dose therapy and ASCT.

Method. Here we report our experience with 29 patients with PTCL. The histological subtypes were as follows: peripheral T-cell lymphoma, not otherwise specified n=13; anaplastic large cell lymphoma (ALCL) ALK-negative n=5; ALCL ALK-positive n=3; ALCL with an unknown ALK status n=3; angioimmunoblastic lymphoma n=1; hepatosplenic lymphoma n=1; Sézary syndrome n=1; and enteropathy-associated T-cell lymphoma n=2. The median age at diagnosis was 48 years (29–64), most patients had advanced Ann Arbor stages (22 patients, 77%), IPI score ≥3 was found in 13 (45%) and PIT score ≥2 in 17 (59%) of the 29 patients. Eighteen patients received first-line high-dose therapy and autologous SCT consolidation; two patients were consolidated with allogeneic SCT in the 1st complete remission and one patient in the 1st relapse. Ten patients with FDG-avid lymphoma were examined with integrated Positron emission tomography/ Computed tomography (PET/CT) at the time of diagnosis and after first-line therapy, two other patients were assessed with Positron emission tomography/ Computed tomography (PET/CT) only at time of restaging.

Results. Nineteen (66%) patients achieved complete remission, 3 (10%) partial remission and 7 (24%) patients failed the treatment. The overall response rate was 76%. PET negativity (complete metabolic response) after therapy was achieved in 8/12 (75%) individuals. After a median follow-up of 55.1 months, 14 (48.3%) patients relapsed or progressed and nine patients died (lymphoma progression). Eleven patients (50% of chemosensitive patients) survived more than 50 months. Three of the long-term survivors were treated with allogeneic SCT. The 2-year event-free survival (EFS) was 52% (95% confidence interval [CI], 0.33–0.71); the 2-year overall survival (OS) rate reached 65% (95%CI, 0.47–0.84). PET negativity was associated with a lower probability of relapse (chi-square p=0.09).

Conclusion. Our data show that intensive first-line therapy with etoposide-doxorubicine-based regimens and ASCT consolidation may lead to long-term disease control in about a half of patients with chemosensitive PTCL. Achievement PET negativity is probably an essential prerequisite for long-term complete remission.

INTRODUCTION

Peripheral T-cell lymphomas (PTCLs) are infrequent types of non-Hodgkin’s lymphomas (NHLs). In Western countries they represent about 7% of NHLs1,2. The incidence of nodal T-cell lymphomas based on the data from the Czech Lymphoma Study Group (CLSG) registry is 6% (211 out of 3518 patients)3. The World Health Organization (WHO) classification recognizes three subgroups of PTCLs: nodal, extranodal and leukemic forms. Nodal PTCL’s are the most frequent and comprise peripheral T-cell lymphoma, not otherwise specified (PTCL, NOS), anaplastic large cell lymphoma (ALCL) and angioimmunoblastic lymphoma (AIL). With the exception of ALCL ALK (Anaplastic Lymphoma Kinase) positive cases is the overall prognosis of nodal PTCL’s poor4–6. Current conventional treatment modalities did not dramatically improved the outcome of patients and 5-year overall survival still remains between 30 % and 35 % using standard chemotherapy with second- and third-generation regimens7–9. The role and timing of high-dose therapy with autologous stem cell support (ASCT) remains unclear – some studies have confirmed the survival advantage of ASCT10,11 while others produced inconsistent results12,13. Data from the Czech national registry of (autologous) hematopoietic stem cell transplantations and the CLSG...
database show an 3-year-overall survival (OS) of 74% of PTCL patients while conventionally treated patients have a median overall survival of as few as 33 months14. Recently published phase II study data showed high efficacy of dose-dense etoposide-based regimen (CHOEP-14) with ASCT consolidation15. A high proportion of patients responded to induction therapy and received ASCT (70%), 3-year OS reached 57% and 3-year PFS 48%.

In the absence of large randomised trials for HDT-ASCT are prognostic factors points of interest to predict the potential benefit of patients receiving high-dose therapy. The prognostic scores like the International Prognostic Index (IPI) (ref.16,17) or PIT (Prognostic index for peripheral T-cell lymphoma-unspecified) (Table 2) (ref.18) showed a significant impact on the outcome after HDT-ASCT in most studies19–21. However, this only reflects the fact that high-risk patients remain at high-risk even after HDT-ASCT. In some studies, the IPI and PIT failed to demonstrate prognostic value22,23. Another major factor that seemed to be significantly associated with the outcome following HDT-ASCT was remission status at time of transplantation24,25. Most studies found remission status to be an independent prognostic factor. The crucial problem in remission assessment is to recognize patients at risk of early treatment failure. In 2007 were published revised response criteria, which included PET scan26. Achieving PET negativity (complete metabolic response, CMR) after therapy was found to have a strong prognostic impact in patients with diffuse large B-cell lymphoma and Hodgkin’s lymphoma27–29. Yet the use and predictive power of PET/CT imaging in T-cell lymphomas remains poorly studied30.

The objective of our retrospective single-center study was to analyze the prognostic scores (IPI, PIT) significance, role of PET assessment and overall outcome of unselected patients with PTCL treated with an intensive first-line chemotherapy protocol with high-dose therapy and autologous stem cell transplantation consolidation.

Table 1. Comparison of two most frequently used prognostic scores in T-cell lymphomas: IPI16,17 and PIT18. PIT is calculated from 4 variables: age above 60, low performance status, elevated lactate dehydrogenase level and bone marrow involvement.

<table>
<thead>
<tr>
<th>Risk group</th>
<th>IPI score</th>
<th>5-year OS</th>
<th>PIT score</th>
<th>5-year OS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>0–1</td>
<td>74%</td>
<td>0</td>
<td>62.3%</td>
</tr>
<tr>
<td>Low-intermediate</td>
<td>2</td>
<td>49%</td>
<td>1</td>
<td>52.9%</td>
</tr>
<tr>
<td>Intermediate-high</td>
<td>3</td>
<td>21%</td>
<td>2</td>
<td>32.9%</td>
</tr>
<tr>
<td>High</td>
<td>4–5</td>
<td>6%</td>
<td>3–4</td>
<td>18.3%</td>
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</tbody>
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PATIENTS AND METHODS

Here we report our experience with 29 patients with PTCL who were diagnosed in our center between the years 2000 and 2009. The histological subtypes were as follows: peripheral T-cell lymphoma, not otherwise specified (PTCL, NOS) n=13; anaplastic large cell lymphoma (ALCL) ALK-negative n=5; ALCL ALK-positive n=3; ALCL with an unknown ALK status n=3; angioimmunoblastic lymphoma n=1; hepatosplenic lymphoma n=1; Sézary syndrome n=1; and enteropathy-associated T-cell lymphoma (EATL) n=2. All biopsies were reviewed by a reference pathologist and final diagnoses were made in compliance with the published WHO classification of lymphoid tumours4. Patient baseline clinical characteristics are summarized in a Table 2.

Table 2. Patients baseline clinical characteristics (n = 29).

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<table>
<thead>
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<tbody>
<tr>
<td>Sex (male-to-female ratio)</td>
<td>20:9</td>
<td>2:2:1</td>
</tr>
<tr>
<td>Median age (range)</td>
<td>48 years</td>
<td>29–64</td>
</tr>
<tr>
<td>Performance status ≥2 (WHO)</td>
<td>7/29</td>
<td>24%</td>
</tr>
<tr>
<td>B symptoms</td>
<td>20/29</td>
<td>69%</td>
</tr>
<tr>
<td>Advanced disease stage (Ann Arbor stage III+IV)</td>
<td>22/29 patients</td>
<td>77%</td>
</tr>
<tr>
<td>Bone marrow involvement</td>
<td>9/29 patients</td>
<td>31%</td>
</tr>
<tr>
<td>Elevated lactate dehydrogenase level</td>
<td>26/29 patients</td>
<td>90 %</td>
</tr>
<tr>
<td>IPI score (≥3)</td>
<td>13/29</td>
<td>45%</td>
</tr>
<tr>
<td>PIT score (≥2)</td>
<td>17 patients</td>
<td>59%</td>
</tr>
<tr>
<td>Median follow-up (range)</td>
<td>55.1 months</td>
<td>4.4–120</td>
</tr>
</tbody>
</table>

Treatment strategy

Twenty-six patients underwent first-line therapy according to the sequential chemotherapy (SQ) protocol, two patients received the ProMACE-CytaBOM regimen and one patient was treated according to the modified SQ protocol. Nineteen patients received first-line high-dose therapy and autologous SCT consolidation (BEAM 200); two patients were consolidated with allogeneic SCT with reduced-intensity conditioning (Flu-Mel-ATG) in the 1st complete remission and one patient in the 1st relapse. Eight patients did not receive high dose therapy: five of them due to primary chemoresistant disease, one was poor stem cells mobilizer, one patient had severe co-morbidities (glomerulonephritis) and one patient was not indicated due to border line age and lymphoma subtype (ALCL ALK-1+).

Treatment protocols

ProMACE-CytaBOM and BEAM 200 regimens were administered as published before11,12. The sequential chemotherapy protocol consists of 3 cycles of CHOEP-21-like regimen (PACEBO), 1 cycle of an ifosfamide
Long-term outcome of patients with peripheral T-cell lymphoma treated with first-line intensive chemotherapy followed by autologous stem cell transplantation

The PACEBO regimen was administered as follows: doxorubicin 40 mg/m² intravenously day 1, cyclophosphamide 850 mg/m² intravenously day 1, etoposide 200 mg/m² intravenously day 1, bleomycin 10 mg/m² intravenously day 8, vincristine 1.4 mg/m² (maximum 2.0 mg) intravenously day 8 and prednisone 40 mg/m² orally days 1 to 14. The IVAM regimen consisted of ifosfamide 1500 mg/m² intravenously days 1 to 5, etoposide 150 mg/m² intravenously day 1 to 3, cytosine arabinoside 100 mg/m² intravenously day 1 to 3, methotrexate 3 g/m² intravenously day 5, mesna prophylaxis 1200 mg intravenously days 1 to 5, leucovorin rescue 25 mg/m² intravenously from day 6/7 until the plasma methotrexate level was below 0.05 μmol/l. The HAM regimen was administered as follows: cytosine arabinoside 2 g/m² twice daily intravenously days 1 and 2, mitoxantrone 10 mg/m² days 2 and 3. Stem cell mobilization was performed with 12 μg/kg of filgrastim given subcutaneously twice daily. The BEAM 200 conditioning regimen dosage was standard as previously published.

The treatment responses – complete response (CR), unconfirmed complete response (uCR), partial response (PR), stable disease and progressive disease – were defined according to the International Workshop NHL Response Criteria published by Cheson33. Twelve patients with FDG-avid lymphoma were examined with integrated PET/CT (Siemens Biograph Sensation 16) at the time of diagnosis and after first-line therapy. The PET scan was defined as positive if higher than mediastinal or background activity was observed. The PET/CT scans were assessed according to the Revised International Workshop Criteria26.

Statistical methods
Our data were analyzed using the Statistical Package for the Social Sciences (SPSS, version 16)34. Overall survival (OS) was defined as the time from first treatment to the date of last follow-up examination (censored) or the date of death (event) from any cause. Progression free survival (PFS) was defined as the date of first treatment to the date of documented disease progression or death (event) or the date of last follow-up examination (censored). The Kaplan-Meier method35 was used to calculate survival probabilities. The log-rank test was used to compare differences in survival times between patient subgroups. The significance level was set at a p= 0.05; 2-tailed tests were used in all calculations.

The EMBASE and PubMed databases were searched for literature reviews.

RESULTS
Analysis of response and survival
Nineteen (66%) patients achieved CR, 3 (10%) partial remission and 7 (24%) patients failed the treatment. The overall response rate was 76%. After a median follow-up of 55.1 months, 14 (48.3%) patients relapsed or progressed (8 PTCL NOS; 2 ALCL ALK-positive; 2 ALCL ALK-negative; 1 Sézary syndrome; 1 EATL; median time 16.1 months) and nine patients died (lymphoma progression).
PET negativity was associated with a lower probability of relapse or disease progression but the significance was borderline (chi-square p=0.09). The intensity of 18F-FDG accumulation may be semiquantitatively determined by calculating the so-called standardized uptake value (SUV). We analyzed SUV max. in ten patients at time of diagnosis. Median SUV max. in a whole group was 14.3 (mean 17.7, range 3.8–47.1). Four patients with ALCL have higher SUV max. (median 23; mean 25.95) than six non-ALCL patients (median 12.15; mean 13.8), (Fig. 5).

Toxicity of treatment
The toxicity of the SQ treatment protocol was tolerable. Most commonly, hematologic toxicity and infections of grades III-HV according to the NCI CTC were observed. No treatment toxicity-related death occurred. No persistent treatment-related disability or severe organ dysfunction were observed. High dose therapy and autologous stem cell transplant procedure was well tolerated. Except hematologic toxicity, only four patients experienced WHO-CTC grade III or IV (Common toxicity criteria) toxicity: two times was observed grade III gastrointestinal toxicity (mucositis, stomatitis), once grade IV gastrointestinal toxicity (stomatitis), three times grade III infection (G+ sepsis, pneumonia, fever of unknown origin). All patients received granulocyte colony-stimulated factors. Median time to engraftment of neutrophils (above 1.0x10^9/l) was 13 days (10–19), median time to engraftment of thrombocytes (above 20x10^9/l) was 11 days (8–19). So far, no secondary malignancies have occurred.

DISCUSSION AND CONCLUSION
PTCLs still represent a heterogeneous group of aggressive lymphomas with disappointing treatment results. In most cases, conventional chemotherapy is ineffective or leads only to response with limited duration. Gisselbrecht et al. reported a 5-year OS of 35% in PTCL patients9, Escalón et al. published even worse results both in patients treated intensively or with the CHOP regimen (3-year OS of 49% and 43%, respectively) (ref.33). The appropriate dose intensity of chemotherapy and the role of high-dose therapy with ASCT were unclear due to the heterogeneity of the studied sample and frequent inclusion of anaplastic PTCL or B-cell lymphoma subtypes25,36. Recently, numerous clinical studies have been published that show variable effects of intensive induction therapy followed by consolidation with high-dose chemotherapy and autologous stem cell transplantation. A large retrospective study carried out by the Lymphoma Working Party of the European Group for Blood and Marrow Transplantation have confirmed the benefit of this treatment modality in patients with AIL, especially if CR was achieved after induction treatment (OS 67% at 24 months and 58% at 48 months) (ref.73). In contrast, a prospective study by Mercadal et al. showed no clear benefit of
aggressive treatment approach in first-line therapy - the use of the high-dose CHOP regimen alternating with the platinum-based ESHAP regimen is associated with significant toxicity and no real increase in the rate of complete remission22. Rodriguez reported the effect of early ICE salvage therapy and autologous stem cell transplantation in high-risk PTCL patients, with gallium-positive scan after 3 cycles of treatment with the megaCHOP protocol and an optimistic OS at 3 years of 73% (ref. 39). Similarly favorable results are those in Rodriguez’s retrospective analysis of AIL patients with a 3-year OS of 60% (ref. 39).

According to our experience, a significant number of patients with nodal PTCL may benefit from the intensive first line therapy and early consolidation with autologous stem cell transplantation. In our group, the overall response rate reached 76% (CR rate 66%, PR rate 10%). The chemotherapy failed in a quarter of the patients. After initial treatment, relapse or lymphoma progression occurred in half of the patients, with a median of 17.7 months. Paradoxically, treatment failed in two patients with ALK-positive ALCL. Both patients suffered from mediastinal lymphoma with primary chemoresistant course. The 2-year progression-free survival (PFS) was 52%; the 2-year overall survival rate reached 65%. The toxicity of the sequential protocol was tolerable. Despite the administration of lower doses of cyclophosphamide and anthracyclines, and the absent platinum-based regimen, the protocol results in treatment outcomes comparable to those achieved by the MegaCHOP/ESHAP regimens. Beyond lower number of analyzed patients it appears that IPI score17 has more power to identify patients at risk of therapy failure, than PIT score. PET/CT assessment seems to be helpful in disease activity monitoring in T-cell lymphomas at time of restaging. Most of the T-cell lymphomas are FDG-avid and intensity of 18F-FDG accumulation at time of restaging. The chemotherapy failed in a quarter of the patients. After initial treatment, relapse or lymphoma progression occurred in half of the patients, with a median of 17.7 months. Paradoxically, treatment failed in two patients with ALK-positive ALCL. Both patients suffered from mediastinal lymphoma with primary chemoresistant course. The 2-year progression-free survival (PFS) was 52%; the 2-year overall survival rate reached 65%. The toxicity of the sequential protocol was tolerable. Despite the administration of lower doses of cyclophosphamide and anthracyclines, and the absent platinum-based regimen, the protocol results in treatment outcomes comparable to those achieved by the MegaCHOP/ESHAP regimens. Beyond lower number of analyzed patients it appears that IPI score17 has more power to identify patients at risk of therapy failure, than PIT score. PET/CT assessment seems to be helpful in disease activity monitoring in T-cell lymphomas at time of restaging. Most of the T-cell lymphomas are FDG-avid and intensity of 18F-FDG accumulation (standardized uptake value, SUV) is higher than average. Patients who do not achieve metabolic response progress very rapidly, on the other hand PET negative CR is prerequisite for long-term CR.

To conclude, the treatment approach to younger patients with PTCL is not unified. The current first-line treatment procedures range from the classical CHOP schemes, through the MegaCHOP/ESHAP/ICE intensive toxic regimens, to high-dose chemotherapy with autologous stem cell transplantation. The administration of our novel intensive first-line sequential chemotherapy protocol with consolidation with high-dose therapy and autologous stem cell transplantation is a safe and effective treatment modality in patients with PTCL. In about half of the patient population, intensive chemotherapy with autologous stem cell transplantation offers a chance for long-term survival and may lead to improved quality of treatment response or provide time for finding a donor for allogeneic transplantation40. The survival of primary chemoresistant patients is still poor. Future advances in treating these patients will not be possible without designing prospective studies, introducing new perspective agents, immunochemotherapy41 and implementing new prognostic schemes18,42 and PET/CT assessment.

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