Introduction

Testicular cancer is the most common solid malignancy in men aged 15-35 years, 95% of which are germ cell tumors (GCT) (1). GCT is primarily located in the gonadal region and rarely occurs in an extragonadal area such as the mediastinum and retroperitoneum. Extranodal GCT (EGGCT) are formed by malignant transformation of gonadal cells that have not completed their migration during embryogenesis without a gonadal primary mass in ultrasonographic evaluation (2). EGGCT accounts for 2-5% of all GCT and has an approximate incidence of 1/1,000,000 (3). Although EGGCT has similar histological, serological and cytogenetic characteristics as gonadal GCT, their behavior is different clinically and biologically. EGGCT has a worse chemosensitivity and prognosis than gonadal tumors (4,5). The treatment of EGGCT patients is similar to the treatment of gonadal GCT, and after histological separation as seminoma and non-seminoma, chemotherapy (CT) is applied according to risk classification. Surgical resection is also performed in patients with residual tumors. This multimodal treatment increases efficacy and survival (6,7). There is no standard salvage CT treatment for relapsed/refractory EGGCT patients. Autologous...
stem cell transplantation (ASCT) therapy for EGGCT patients has started to increase gradually and is mostly applied as second or third salvage (8).

The aim of this study was to share our single center experience with patients who underwent salvage ASCT with the diagnosis of relapse/refractory EGGCT.

Methods

In this study, 30 patients with EGGCT who underwent high dose CT (HDCT) and ASCT between 4 February 1991 and 12 May 2015 at the University of Health Sciences, Gülhane Training and Research Hospital, Medical Oncology Clinic, Ankara, Turkey were evaluated retrospectively. The study was approved by the Ethics Committee of Gülhane Training and Research Hospital with the decision number 18/159. Patient interview information, patient files and electronic system data were used to obtain data. Patients’ demographic status, tumor localization, first recurrence dates, salvage CT protocols and treatment responses, HDCT and ASCT time, and HDCT and ASCT recurrence and final status were noted.

The primary endpoint is overall survey (OS) and progression free survey (PFS). The diagnosis date was accepted as the start date for the general OS value of the patients, the end point was the last control date for the surviving patients, and the exitus date for the ex-patients. In order to calculate OS values after HDCT and ASCT, start date HDCT and ASCT date is accepted. Endpoint for OS is the last control date for living patients, exitus date for ex patients.

The PFS values of the patients after the first step treatment were calculated as PFS1. In addition, the time to recurrence after HDCT and ASCT was calculated as PFS2; HDCT and ASCT dates were taken as the starting date, relapse date for the relapse as the endpoint, and the last control date for the non-relapse.

Patients with pathologic evidence of EGGCT who had relapsed after first-line treatment and who had HDCT and ASCT were included in the study. Patients who did not have HDCT and ASCT and whose files and follow-up information were missing were excluded from the study.

Statistical Analysis

Statistical Package for Social Sciences version 24.0 was used for conducting statistical analysis of data (IBM Corp, Armonk, NY, USA). Descriptive statistics for expressing continuous (quantitative) variables were mean, standard deviation, minimum and maximum values, while the categorical variables were expressed as number (n) and ratio (%). The suitability of the variables to the normal distribution was evaluated by visual and analysis methods and nonparametric tests were used because they did not fit the normal distribution. The chi-square and Fisher’s exact tests were used to determine the demographic characteristics of the patients. The Kaplan-Meier was used for univariate survey analysis and log rank test was used. In multivariate analyses, the Cox regression test was used. The Spearman’s rank correlation test was used for univariate correlation analysis. Statistically significant value was accepted as that less than 0.05.

Results

A total of 30 patients who underwent HDCT and ASCT were included in the study. The median age of the patients was 41 years (range: 21-60). First line 4 cycles of BEP (bleomycin, etoposide, cisplatin) were applied to all patients. During follow-up, all patients relapsed and the median PFS1 was 16 (range: 3-45) months. TIP (paclitaxel, ifosfamide and cisplatin) was given to 27 (90%) patients after the relapse and VIP (etoposide, ifosfamide and cisplatin) salvage CT was applied to 3 (10%) patients. After Salvage CT, complete response (CR) in 10 (33.3%), partial response (PR) in 12 (40%), stable response in three (10%) and progression in five (16.7%) patients were observed. The primary characteristics were summarized in Table 1.

<table>
<thead>
<tr>
<th>Table 1. Primary characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Primary site, n (%)</strong></td>
</tr>
<tr>
<td>Retroperitoneal</td>
</tr>
<tr>
<td>Mediastinal</td>
</tr>
<tr>
<td><strong>Pathology, n (%)</strong></td>
</tr>
<tr>
<td>Embryonal carcinoma</td>
</tr>
<tr>
<td>Teratoma</td>
</tr>
<tr>
<td>Mixed nonseminoma</td>
</tr>
<tr>
<td>Choriocarcinoma</td>
</tr>
<tr>
<td>Yolk sac</td>
</tr>
<tr>
<td><strong>Metastasis, n (%)</strong></td>
</tr>
<tr>
<td>Yes</td>
</tr>
<tr>
<td>No</td>
</tr>
<tr>
<td><strong>Metastasis-site, n (%)</strong></td>
</tr>
<tr>
<td>Brain</td>
</tr>
<tr>
<td>Bone</td>
</tr>
<tr>
<td>Lung</td>
</tr>
<tr>
<td>Liver</td>
</tr>
<tr>
<td>Multiple organs</td>
</tr>
<tr>
<td><strong>RPLND, n (%)</strong></td>
</tr>
<tr>
<td>Yes</td>
</tr>
<tr>
<td>No</td>
</tr>
<tr>
<td><strong>RPLND pathology, n (%)</strong></td>
</tr>
<tr>
<td>Necrosis</td>
</tr>
<tr>
<td>Viable tm</td>
</tr>
<tr>
<td><strong>Metastasectomy, n (%)</strong></td>
</tr>
<tr>
<td>Yes</td>
</tr>
<tr>
<td>No</td>
</tr>
</tbody>
</table>
All patients underwent HDCT and ASCT between 4 February 1991 and 12 May 2015. In our study, the follow-up period after diagnosis was 137 months (range: 30-353 months); the median follow-up period after HDCT and ASCT was 110 months (range: 29-327 months).

Patients who relapsed after the first step treatment were included in the study. The median PFS1 value of patients until the first relapse after the first cure was 16 months (range: 3-45 months). The patient relapsed for 9 months (30%) after HDCT and ASCT. The median value of PFS2 after HDCT and ASCT is 101 months (range: 26-235 months). All patients who recurred after HDCT and ASCT were ex.

According to our current data, 9 (30%) patients died and 21 (70%) patients were alive. The median OS diagnosis value after diagnosis was 136 months (range: 31 to 353 months), and the median OSASCT value after HDCT and ASCT was 103 months (range: 29-327 months).

The 2-year OS diagnosis value of the patients in our study group was 97.6%; the 5-year OS diagnosis value was 93.2% and the 10-year OS diagnosis value was 84.7%. The 2-year OSASCT value of our patients was 93.7%; the 5-year OSASCT value was 88.4% and the 10-year OSASCT value was 78.3%. The 2-year PFSASCT value of our patients was 92.6%; the 5-year PFSASCT value was 87% and the 10-year PFSASCT value was 77.5%.

Factors affecting OS diagnosis
The median OS diagnosis value of the patients after diagnosis was 136 (range: 31 to 353) months. The mediastinal or retroperitoneal (RP) localization of the disease did not affect OS diagnosis significantly (p=0.072). While the median OS diagnosis was found to be 140 months (range: 30-319) in patients with RP, the median was 133 months (66-353) in those with the mediastinum (Figure 1).

When the relationship between the pathologies and OS diagnosis was evaluated, the median was 186 months (range: 118-223) for embryonic carcinoma, the median was 136 months for teratoma (range: 30-218), the median was 130 months (range: 51-319) for mixed nonseminoma, the median was 107 months (range: 66-162) for patients with choriocarcinoma; the median was 222 months (206-240) in the yolk sac.

OS diagnosis was significantly affected by the relapse region of the patients (p=0.001). The median OS diagnosis was 135 months (range: 51-240) in patients with relapse RP, and 150 months (range: 86-353) in patients with recurrent mediastinal region, 118 months (range: 66-218) in patients with recurrence of the lung, 30 months only in the case of recurrence in the bone and 319 months in the only case of recurrence in the brain. When the subgroup analysis was performed, it was found that the relapse in the bone where the significance was caused by bone had a significantly lower OS diagnosis value than the others.

There was a significant relationship between recurrence...
after HDCT and ASCT and OS\textsubscript{diagnosis} (p=0.001). The median value of OS\textsubscript{diagnosis} after HDCT and ASCT was 136 months (range: 51-353) and the median OS\textsubscript{diagnosis} value was 130 months (range: 30-240).

**Factors affecting OS\textsubscript{ASCT}**

The median OS\textsubscript{ASCT} value after the ASCT date was 103 (range: 29-327) months. Primary mediastinal or RP did not significantly affect OS\textsubscript{ASCT} (p=0.075). While the median OS\textsubscript{ASCT} was 101 months (range: 29-304) in patients with RP and 103 months (49-327) in patients with primary mediastinum.

OS\textsubscript{ASCT} was found to have a significant effect on the relapse region (p=0.006). The median time was 114 months (range: 51-304) in patients with relapse RP and 90 months (range: 36-327) in patients with mediastinal area, 80 months (range: 45-153) in patients with lung recurrence, 29 months in a single case with bone and 50 months in a single case of recurrence in the brain. When the subgroup analysis was performed, it was found that the bone recurrent case had a significantly lower OS\textsubscript{ASCT} value than the others (Figure 2).

When the relationship between OS\textsubscript{ASCT} and the pathology of the patients was evaluated, the median was 172 months for embryonal carcinoma (range: 80-202), 109 months (range: 29-153) for teratoma, 88 months (range: 45-304) for mixed nonseminoma, 48 months (range: 36-117) for choriocarcinoma and 197 months (143-200) for yolk sac.

Unlike OS\textsubscript{diagnosis}, OS\textsubscript{ASCT} was significantly affected by the age of the patients. The median OS\textsubscript{ASCT} value of patients aged 40 years and under was 47 months (range: 28-118), whereas for those over 40 years, this value was 148 months (range: 50-327) (p=0.012) (Figure 3).

**Factors affecting PFS2 (PFS\textsubscript{ASCT})**

The median value of PFS2 (PFS\textsubscript{ASCT}) after HDCT and ASCT was 101 months (range: 26-235). All patients who recurred after HDCT and ASCT died. PFS2 did not significantly affect the mediastinal or RP primer (p=0.070). While the median PFS2 was 101 months (range: 26-304) in patients with RP primer, the median PFS2 was 101 months (45-327) in those with mediastinal primer.
The relapse region of the patients was significantly affected by PFS2 ($p=0.001$). The median PFS2 was 112 months (range: 27-27304) in patients with relapse RP, and 86 months (range: 35-327) in patients with mediastinal recurrence. It was 76 months (range: 40-153) in patients with lung recurrence, 26 months in bone recurrence, and 48 months in single brain recurrence. When subgroup analysis was performed, it was found that the recurrent bone case had a significantly lower PFS2 value than the others (Figure 4).

When the relationship between PFS2 and patients’ pathology was evaluated, it was found that the median PFS2 was 170 months (range: 76-201) in embryonal carcinoma, 109 months in teratoma (range: 26-153), 88 months in mixed nonseminoma (range: 43-304), 48 months in choricarcinoma (range: 36-117) and 197 months (143-200) in the yolk sheet.

PFS2 was significantly affected by the age of patients at the time of ASCT. The median PFS2 value was 45 months (range: 26-118) for patients aged 40 years and younger, and 147 months (range: 48-327) for those over 40 years of age ($p=0.010$).

**Discussion**

In relapsed/refractory GCT patients, HDCT and ASCT have been used as standard salvage treatment and are often used for second or third salvage purposes (9). However, due to the low number of relapsed/refractory EGGCT patients, the efficacy of HDCT and ASCT in these patients could not be clearly demonstrated due to the lack of randomized trials. In this study, we evaluated the demographic characteristics, progression-free and total survival data retrospectively in our center to demonstrate the efficacy of treatment in relapsed/refractory EGGCT patients who underwent HDCT and ASCT for second or third salvage purposes.

All of our EGGCT patients had histopathology in the non-seminomatous group and all of them were in the poor prognostic group according to IGCCG (9,10). We performed HDCT and ASCT as the second salvage in 10 of our patients and as the third salvage in 20 of our patients. There was no significant relationship between salvage application step and OS and PFS. There is a high chance of success in the studies performed in GCT patients in the third step and before, and it provides duration of response even in the next steps (11).

In many studies, VIP as the first salvage regimen is preferred primarily (12,13), but in our patients we preferred TIP treatment as the first salvage, as in the study of 69 patients included by Ko et al. (14). There is no study designed to compare the advantages of TIP and VIP regimens as salvage treatment.

Total survival in EGGCT patients varies according to primary tumor localization, relapse status, relapse duration and IGCCCG criteria. While 5-year OS is 65% in primary RP EGGCT, this rate decreases to 40-45% in mediastinal EGGCT patients (2,12,15,16). In a study conducted by Schmoll et al. (17), the disease-specific survival of RP EGGCT patients who underwent HDCT and ASC was to be be 76%. In our study, the 5-year OSASCT value was reported as 88.4% and clearly shows the efficacy of the treatment.

In our study, the mean age was 41 years and 15 years older than the median age in the literature (18,19). The average age is high compared to the literature, decreases bone marrow reserve, causes neutropenia and febrile neutropenia complications, and may lead to an increase in mortality rate. In addition, in the literature (20), the average rate of advanced disease was 30%, whereas in our study, this rate was 50% and the mortality risk was high. In our study, nine patients died and despite this high risk, this rate is significantly lower than in the literature. In addition, in our study, an unspecified result was obtained in the literature and OSASCT and PFS2 were significantly better in patients over 40 years of age, who underwent HDCT and ASCT.

Hege et al.’s (21) study revealed that treatment-related mortality was found to be 5.5% in HDCT and ASCT patients and Adra et al. (22) also presented treatment-related mortality as 2.4% (21). In our study, no patients died in association with the treatment.

Considering the limitations of our study, the study consisted of a report on rare case series that underwent AHSCT due to relapsed or refractory EGGCT; therefore, it included a small sample size of patients. Also, it had a heterogeneous patient population with regard to indications for AHSCT and it was a retrospective study.

**Conclusion**

As a result, for patients with relapsed/refractory EGGCT, high dose ifosfamide/carboplatin/etoposide regimen was safe and and an effective treatment choice. Although the chemosensitivity and prognosis of EGGCT patients are worse than GCT patients, their survival is significantly increased with multimodal treatments and ASCT. Future prospective randomized studies should reveal more reasonable and effective survival results.

**Acknowledgements**

We thank our patients and their families who participated in the research helpfully and devotedly without expecting material compensation. We are grateful to Prof. Dr. İlker Taşçı for critically editing of the manuscript.

**Ethics**

Ethics Committee Approval: The study was approved by the Ethics Committee of Gülhane Training and Research Hospital with the decision number 18/159.

Informed Consent: Retrospective study.
Published by Elsevier Inc. on behalf of the European Society for Blood and Marrow Transplantation on behalf of the European Society for Blood and Marrow Transplantation.

Peer-review: Externally peer-reviewed.

Authorship Contributions

Conflict of Interest: No conflict of interest was declared by the authors.

Financial Disclosure: The authors declared that this study received no financial support.

References