Impact of previous disease-modifying treatment on safety and efficacy in patients with MS treated with AHSCT

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ABSTRACT

Background Autologous haematopoietic stem cell transplantation (AHSCT) is a highly effective treatment for multiple sclerosis (MS). The impact of previous long-lasting disease-modifying treatments (DMTs) for safety and efficacy of AHSCT is unknown.

Objective To explore whether previous DMTs with long-lasting effects on the immune system (anti-CD20 therapy, alemtuzumab and cladribine) affect treatment-related complications, long-term outcome and risk of new MS disease activity in patients treated with AHSCT.

Methods Retrospective observational study of 104 relapsing remitting patients with MS treated by AHSCT in Sweden and Norway from 2011 to 2021, grouped according to the last DMT used ≤6 months prior to AHSCT. The primary outcomes were early AHSCT-related complications (mortality, neutropenic fever and hospitalisation length), long-term complications (secondary autoimmunity) and proportion of patients with No Evidence of Disease Activity (NEDA-3 status): no new relapses, no MRI activity and no disease progression during the follow-up.

Results The mean follow-up time was 39.5 months (range 1–95). Neutropenic fever was a common AHSCT-related complication affecting 69 (66%) patients. There was no treatment-related mortality. During the follow-up period, 20 patients (19%) were diagnosed with autoimmunity. Occurrence of neutropenic fever, hospitalisation length or secondary autoimmunity did not vary dependent on the last DMT used prior to AHSCT. A total of 84 patients (81%) achieved NEDA-3 status, including all patients (100%) using rituximab, alemtuzumab or cladribine before AHSCT.

Conclusion This study provides level 4 evidence that AHSCT in patients previously treated with alemtuzumab, cladribine or rituximab is safe and efficacious.

INTRODUCTION

Autologous haematopoietic stem cell transplantation (AHSCT) is a treatment option for patients with aggressive relapsing remitting multiple sclerosis (RRMS). The therapeutic rationale is based on giving immunoablative treatment causing depletion of autoreactive cells, followed by infusion of cryopreserved autologous haematopoietic stem cells to support immune reconstitution.1 AHSCT is primarily offered to patients with highly aggressive MS, mostly after failing other disease-modifying treatments (DMTs). Available DMTs vary according to potency, side effects and the route and frequency of administration. Rituximab, cladribine and alemtuzumab cause depletion of immune cells, and also have the potential to induce long-term drug-free remissions.1,2 The preferred first-line treatment choice and sequence of DMTs vary between treatment centres, but DMTs with high potencies are recommended for patients with aggressive disease.3 Most centres have strict criteria for AHSCT treatment, preserving it as a second-line or third-line therapy for patients with highly aggressive MS, but...
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sometimes it is offered as a first treatment choice. The results published after AH SCT in RRMS patients demonstrate excellent outcomes, with 60%–80% of patients attaining No Evidence of Disease Activity (NEDA-3)-status for >5 years.

Whether AH SCT should be offered to patients previously treated with DMTs with long-lasting effects on the immune system is debated, and currently no consensus has been reached. Some treatment centres and studies have excluded patients due to preceding use of DMTs, especially alemtuzumab. The reasons have been fear of prolonged aplasia and risk of infections after AH SCT in patients with pre-AH SCT lymphopenia due to DMTs, and development of secondary autoimmune diseases. However, a recently published case report of three patients indicated that AH SCT may be safe after previous alemtuzumab treatment in patients with MS. So far, no reports are available on the efficacy and safety after AH SCT following cladribine or anti-CD20 therapy. The use of DMTs with long-lasting effects on the immune system is increasing. Accordingly, it is of importance to evaluate the effect and safety of AH SCT for patients previously exposed to these treatment options. In this study, data regarding the previous use of DMTs were collected from patients treated at Haukeland University Hospital in Norway and at Uppsala University Hospital in Sweden between 2011 and -2021 to determine whether various DMTs had an impact on AH SCT-related complications and long-term prognosis in RRMS, with the main aim to clarify the safety of previous use of rituximab, alemtuzumab and cladribine.

MATERIAL AND METHODS

Study population, design and data sources

This is a retrospective observational cohort study of 104 RRMS (35 Norwegian and 69 Swedish) patients. The Swedish patients consisted of RRMS patients treated with AH SCT from 1 January 2011 to 31 December 2018 at Uppsala University Hospital. All patients with MS in Sweden are recorded in the Swedish Multiple Sclerosis Register, and the patients were identified through a register search and originally asked to participate in a comparative study of AH SCT and alemtuzumab. In Norway, all patients with MS treated with AH SCT at Haukeland University Hospital from January 2015 to February 2021 before or outside an ongoing randomised trial (RAM-MS=RANDOMised controlled trial comparing autologous haematopoietic stem cell transplantation versus altemtuzumab, cladribine or ocrelizumab in MS), were asked to participate, and 95% accepted. In both countries, AH SCT is offered to patients with highly active MS, most frequently after detection of new disease activity in spite of ongoing standard DMT. In Sweden, some patients who present with aggressive MS are offered AH SCT as their first treatment option. All patients had clinical follow-ups at least annually, including assessment of Expanded Disability Status Scale (EDSS), MRI and registration of adverse events. Medical records of all patients were assembled and evaluated.

Procedure

The treatment protocols were the same at both treatment centres and consisted of an intermediate intensity, lymphoablative/not-myeloablative conditioning regimen. Peripheral haematopoietic stem cells were mobilised by a single dose of cyclophosphamide (2 g/m²), followed by daily granulocyte colony-stimulating factor, 5–10 &/kg × 1 per day for 5–7 days. Patients were conditioned with a combination of cyclophosphamide and rabbit antithymocyte globulin (cyclophosphamide 200 mg/kg; rATG 6 mg/kg). The cryopreserved autologous stem cells (a minimum amount of 3×10⁶ CD34+ cells/kg) were reinfused without any graft manipulation. All patients received prophylactic antibiotic regimens with ciprofloxacin, valaciclovir and fluconazole.

Measures of disease activity

NEDA-3 was defined as a composite score comprising absence of clinical relapses, sustained disability progression and new MRI disease activity (new T1 gadolinium enhancing lesions or new/enlarging T2-lesions) on MRI examinations for the given period. A relapse was defined as the appearance of new symptoms or signs that lasted for more than 24 hours without concurrent fever or illness. Progression was defined as an increase in EDSS score of at least one point from baseline sustained between two follow-up visits separated in time by no less than 6 months (1.5 point if EDSS at baseline was 0, 0.5 points if the baseline EDSS ≥5.5).

Statistics

Baseline data and demographic results were assessed using descriptive statistics. For illustration of safety outcomes and long-term treatment results, we used Kaplan-Meier survival curves. The differences between groups were assessed by one-way analysis of variance. A p<0.05 was considered statistically significant. All analyses were assessed by SPSS Statistics V.26 (IBM). Data are described as means and percentages.

RESULTS

Patient characteristics

The cohort consisted of 104 RRMS patients (69 Swedish and 35 Norwegian) with a mean age of 30.8 years (10.2–58.8), and a female-male ratio of 2.7:1. All patients had active disease, and the mean ARR 1 year prior to AH SCT were 1.7. Baseline demographics are shown in table 1. Most patients had a history of suboptimal treatment responses to other DMTs, due to either side effects or clinical relapses, but 12 patients had no previous MS treatment. The mean number of previous DMTs was 2.1, while 17 patients (16.3

<table>
<thead>
<tr>
<th>Table 1 Demographic and clinical data at baseline</th>
</tr>
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<tbody>
<tr>
<td><strong>Patients, n Swedish/Norwegian</strong></td>
</tr>
<tr>
<td><strong>Gender, female/male</strong></td>
</tr>
<tr>
<td><strong>Age, years (mean/range)</strong></td>
</tr>
<tr>
<td><strong>Disease duration, years (mean)</strong></td>
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<tr>
<td><strong>EDSS at baseline (median/range)</strong></td>
</tr>
<tr>
<td><strong>No of previous treatments</strong></td>
</tr>
<tr>
<td><strong>Last treatment (≥6 months prior to HSCT)</strong></td>
</tr>
<tr>
<td><strong>No treatment</strong></td>
</tr>
<tr>
<td><strong>Standard DMT</strong></td>
</tr>
<tr>
<td>Interferons</td>
</tr>
<tr>
<td>Glatiramer acetate</td>
</tr>
<tr>
<td>Fingolimod</td>
</tr>
<tr>
<td>Natalizumab</td>
</tr>
<tr>
<td>Dimethyl fumarate</td>
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<tr>
<td>Teriflunomide</td>
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<tr>
<td><strong>DMT with long-lasting effect</strong></td>
</tr>
<tr>
<td>Alemtuzumab</td>
</tr>
<tr>
<td>Cladribine</td>
</tr>
<tr>
<td>Rituximab</td>
</tr>
<tr>
<td><strong>DMT, disease-modifying treatment; EDSS, Expanded Disability Status Scale; HSCT, haematopoietic stem cell transplantation.</strong></td>
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</tbody>
</table>
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% had used ≥4 previous DMTs. A total of 79 (76 %) of the patients had been exposed to a DMT the last 6 months prior to AHSCT. A total of 26 patients (25 %) had used DMTs with a long-term effect on the immune system; rituximab, alumtruzumab and cladribine. The rest of the patients had been treated with natalizumab, fingolimod, interferons, dimethyl fumarate, glatiramer acetate and teriflunomide. The mean follow-up time was 39.5 months (range 1–95).

Early adverse events

The patients had a mean 13 days of inpatient care (range 10–23). Time to engraftment did not vary according to the last DMT given prior to AHSCT. A total of 69 patients (66 %) had neutropenic fever, and were treated by intravenous antibiotics, of these 20 (29 %) patients had used DMTs with long-term effects on the immune system. There was no significant difference between the groups (p=0.3). One patient (1%) treated with natalizumab 3 months before AHSCT, had septic febrile neutropenia with hypotonia and Epstein-Barr virus reactivation, and was observed in an intensive care unit for 1 day. One patient without previous DMT developed fever and psychosis, was treated with intravenous antibiotics and steroids, and had a good recovery. One patient previously treated with fingolimod had a thoracic venous thrombosis during hospitalisation. There was no treatment-related mortality. All patients were discharged from the hospital within 23 days.

Late adverse events

During the follow-up period, 20 patients (19 %) had a secondary autoimmune disease. A total of 11 acquired hyperthyroidism (10 %), seven hypothyroidism (7 %), one patient hypothyroidism and psoriasis vulgaris (1 %), and one patient (1 %) autoimmune thrombocytopenic purpura. Among the patients treated with DMTs with a long-lasting effect (N=26), four patients (15 %) developed secondary autoimmunity. Two patients previously treated with alemtuzumab were diagnosed with hyperthyroidism. A total of seven patients (7 %) with secondary autoimmunity was not exposed to DMT the last 6 months prior to AHSCT. There was no statistically significant difference in the number of patients developing secondary autoimmunity between the different pre-AHSCT DMTs (table 2).

Table 2 Demographics and results according to previous treatment (<6 months prior to AHSCT)

<table>
<thead>
<tr>
<th>Whole cohort</th>
<th>None</th>
<th>INTF</th>
<th>GA</th>
<th>FTY</th>
<th>NTZ</th>
<th>DMF</th>
<th>TFM</th>
<th>ALEM</th>
<th>CLD</th>
<th>RTX</th>
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<tr>
<td>Patients, no</td>
<td>104</td>
<td>25</td>
<td>6</td>
<td>4</td>
<td>15</td>
<td>20</td>
<td>5</td>
<td>3</td>
<td>6</td>
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<tr>
<td>Age, mean</td>
<td>30.8</td>
<td>29.7</td>
<td>24.4</td>
<td>37.5</td>
<td>28</td>
<td>29</td>
<td>28.5</td>
<td>34.8</td>
<td>32.5</td>
<td>23.5</td>
</tr>
<tr>
<td>Gender, no (F/M)</td>
<td>76/28</td>
<td>19/6</td>
<td>4/2</td>
<td>2/2</td>
<td>12/3</td>
<td>13/7</td>
<td>4/1</td>
<td>2/1</td>
<td>6/0</td>
<td>1/1</td>
</tr>
<tr>
<td>Baseline EDSS, median</td>
<td>3.0</td>
<td>3.5</td>
<td>2.7</td>
<td>3.0</td>
<td>3.0</td>
<td>3.3</td>
<td>4.0</td>
<td>2.5</td>
<td>4.0</td>
<td>2.3</td>
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<tr>
<td>ARR, 1 year prior to treatment, mean</td>
<td>1.7</td>
<td>1.6</td>
<td>1.8</td>
<td>1.3</td>
<td>1.7</td>
<td>2.6</td>
<td>2.2</td>
<td>0.7</td>
<td>1.5</td>
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<tr>
<td>Washout duration last DMT, months, mean</td>
<td>–</td>
<td>–</td>
<td>1.9</td>
<td>2.5</td>
<td>3.4</td>
<td>3.6</td>
<td>5.3</td>
<td>3.5</td>
<td>–</td>
<td>–</td>
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<tr>
<td>Follow-up, months, mean</td>
<td>39.5</td>
<td>38.3</td>
<td>59.3</td>
<td>73.3</td>
<td>37.7</td>
<td>41.9</td>
<td>44.8</td>
<td>16.6</td>
<td>23</td>
<td>24.5</td>
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<tr>
<td>Neutropenic fever, n (%)</td>
<td>69 (66)</td>
<td>14 (56)</td>
<td>4 (66)</td>
<td>3 (75)</td>
<td>12 (80)</td>
<td>10 (50)</td>
<td>4 (80)</td>
<td>2 (66)</td>
<td>6 (100)</td>
<td>1 (50)</td>
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<tr>
<td>Hospitalisation, days, mean</td>
<td>13.1</td>
<td>12.8</td>
<td>13</td>
<td>12</td>
<td>12.9</td>
<td>13.2</td>
<td>12.6</td>
<td>14.3</td>
<td>14</td>
<td>12.5</td>
</tr>
<tr>
<td>Secondary autoimmunity, n (%)</td>
<td>20 (19)</td>
<td>7 (28)</td>
<td>1 (17)</td>
<td>1 (25)</td>
<td>2 (13)</td>
<td>2 (10)</td>
<td>2 (40)</td>
<td>1 (33)</td>
<td>2 (33)</td>
<td>0 (0)</td>
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<tr>
<td>New disease activity, n (%)</td>
<td>20 (19)</td>
<td>6 (24)</td>
<td>3 (50)</td>
<td>1 (25)</td>
<td>5 (33)</td>
<td>4 (20)</td>
<td>1 (20)</td>
<td>0 (0)</td>
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</table>

*1 missing.

AHSCT, autologous haematopoietic stem cell transplantation; ALEM, alemtuzumab (Lemtrada); ARR, annualised relapse rate; CLD, cladribine (Mavenclad); DMF, dimethyl fumarate (Tecfidera); DMT, disease-modifying treatments; FTY, fingolimod (Gilenya); GA, glatiramer acetate (Copaxone); INTF, interferons (Pledigry/Betaferon/Avonex); NTZ, natalizumab (Tysabri); RTX, rituximab (MabThera); TFM, teriflunomide (Aubagio).

Figure 1 Kaplan-Meier survival curve of relapse-free survival (A), EDSS progression-free survival (B) and disease-free survival (D) (patients with achieved NEDA-3 status) according to last DMT. AHSCT, autologous haematopoietic stem cell transplantation; DMT, disease-modifying treatment; EDSS, Expanded Disability Status Scale; NEDA, no evidence of disease activity.

Efficacy

A total of 20 patients (19 %) had evidence of disease activity during the follow-up period; 9 patients had new relapses and MRI activity, 8 had only new MRI activity and 3 had sustained disease progression. The patients with disease progression had a baseline EDSS of 4–5.5 that increased from 1 to 3 points. The time span before new disease activity varied from 1 to 64 months after AHSCT. The number of patients attaining sustained NEDA-3 status differed according to the last pre-AHSCT MS treatment (p<0.01) as illustrated in figure 1. A total of 14 (70 %) of the patients with new disease activity after AHSCT had used standard DMTs the last 6 months prior to AHSCT. All patients using rituximab (N=18), alemtuzumab (N=6) or cladribine (N=2) before AHSCT attained NEDA-3 status during the follow-up period (table 2).

DISCUSSION

In this retrospective study of 104 RRMS patients, former treatment with rituximab, alemtuzumab or cladribine were associated with the same frequency of early and late adverse events as pre-AHSCT treatment with other DMTs. Neutropenic fever was a common complication, but was not associated with specific
DMTs. None of the three patients with the most serious early adverse events had been treated with rituximab, alemtuzumab or cladribine before AHSCST.

Rituximab, cladribine and alemtuzumab cause immunosuppression by lymphocyte depletion. Rituximab treatment is particularly associated with increased risk of infections, but this has also been shown for alemtuzumab. Since alemtuzumab increases the risk of opportunistic infections, such as *Listeria meningitis*, cautious and restricted use are recommended. Interestingly, we did not find any increase in AHSCST-related infections in patients previously treated with rituximab, cladribine or alemtuzumab. Alemtuzumab has previously been shown to increase the risk of secondary autoimmune diseases, with thyroid autoimmune disease being the most frequent, affecting nearly half of the patients treated. A total of 20 patients were diagnosed with a secondary autoimmune disease during the follow-up period, two of which had used alemtuzumab. The impact of DMTs on the safety and efficacy of a subsequent AHSCST has not been previously explored in patients with MS, except for a case report of three AHSCST patients treated with alemtuzumab. The results were consistent with our findings, indicating that the risk of AHSCST is not relatively increased after previous treatment with alemtuzumab.

We found that all patients previously treated with rituximab, alemtuzumab and cladribine attained a sustained NEDA-3 status throughout the follow-up period. For patients with other DMTs or no treatment the last 6 months prior to AHSCST, the proportion with new MS disease activity varied. The impact of previous fingolimod use on disease activity after AHSCST has so far not been examined. However, a number of reports have explored the effects of treatment shifts for various sequences of DMTs. An observational study found that fingolimod treatment was less effective in patients discontinuing natalizumab. For fingolimod, there is a known risk of a rebound effect after discontinuation, and several reports have described a suboptimal disease control for treatment with alemtuzumab, rituximab and ocrelizumab in patients previously treated with fingolimod.

This may be explained by the mode of action of fingolimod, that is, sequestering the immune cells in the lymph nodes. Hence, autoreactive pathogenic B cells may be sequestered in secondary lymph nodes and return to the circulation after new treatment is started. In this study, a total of 15 patients used fingolimod as the last treatment before AHSCST, with a wash-out period ranging from 1.5 to 5.5 months. Interestingly, no impact of fingolimod was registered regarding the clinical effects or adverse events after AHSCST.

We found a significant difference in MS disease activity after AHSCST depending on the patients pre-AHSCST medication, in favour of alemtuzumab, rituximab and cladribine. One possible reason may be that previous DMTs with long-lasting effects could influence the amount of remaining autoreactive lymphocytes in the circulation and in the autologous stem cell product. This study examined a significant cohort of RRMS patients treated with AHSCST, constituting most patients treated with AHSCST in Sweden and Norway from 2011 to 2021, and with an average follow-up period of more than 3 years. All patients were treated with the same intermediate intensity conditioning regime consisting of cyclophosphamide and ATG. The patients studied had been treated by most of the available DMTs prior to AHSCST. Treatment with rituximab, alemtuzumab or cladribine was used for 26 patients before AHSCST, making this a cohort suitable to explore potential differences in AHSCST treatment effect and side effects between pre-AHSCST-treatments with long-lasting and short-lasting immunomodulatory effects. There are, however, some limitations. The groups of patients treated with each individual DMT was small, and data indicating a higher chance of obtaining NEDA-3 among patients given pre-AHSCST DMTs with long-term effects on the immune system should accordingly be interpreted with caution. Furthermore, we cannot rule out that a longer observation time could have yielded somewhat different results regarding the occurrence of secondary autoimmunity or the post-AHSCST MS disease activities.

DMTs with long-term effects on the immune system have pronounced effects on disease activity, and are increasingly preferred treatment choices for MS. In the upcoming years, a higher proportion of AHSCST patient candidates will have a medical history of prior usage of such DMTs. Our data indicate that previous treatment with alemtuzumab, cladribine or rituximab is safe, and associated with a high likelihood of sustained NEDA-3 after transplantation.

CONCLUSION

This study provides level 4 evidence that AHSCST in patients previously treated with alemtuzumab, cladribine or rituximab is safe and efficacious.

Correction notice This article has been corrected since it was first published. The open access licence has been updated to CC BY.

Contributors SASK: study concept and design, acquisition of data. JB: acquisition of data, critical revision of the manuscript for important intellectual content, study supervision. AKL: acquisition of data, critical revision of the manuscript for important intellectual content, study supervision. AT: acquisition of data, critical revision of the manuscript for important intellectual content. CZ: acquisition of data, critical revision of the manuscript for important intellectual content. GKM: critical revision of the manuscript for important intellectual content. LB: acquisition of data, critical revision of the manuscript for important intellectual content, study supervision. ØT: study concept and design, analysis and interpretation, acquisition of data, critical revision of the manuscript for important intellectual content, study supervision. SASK acts as guarantor and is responsible for the overall content.

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Patient consent for publication Not applicable.

Ethics approval The study was approved by the Regional Ethical Committee (REK 2018/377) of Norway.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request.

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