The role of posttransplantation maintenance chemotherapy in improving the outcome of autotransplantation in adult acute lymphoblastic leukemia

Ray Powles, Bhawna Sirohi, Jennifer Treleaven, Samar Kulkarni, Diana Tait, Seema Singhal, and Jayesh Mehta

Extending the principle of conventional acute lymphoblastic leukemia (ALL) therapy to transplantation, 77 adult patients receiving autografts in first remission after melphalan with or without total body irradiation were scheduled to receive 6-mercaptopurine (6MP), methotrexate (MTX), and vincristine-prednisone (VP) for 2 years after transplantation to reduce relapse. Seventy-one percent of patients receiving 6MP, 57% received MTX, and 38% received VP. Thirty patients had a relapse at 1.5 to 80 months (median, 12.5 months), 15 in the first year and 7 beyond 3 years. The cumulative incidence of relapse at 10 years was 42% (95% CI, 31%-55%). The 10-year probabilities of disease-free survival (DFS) and overall (OS) survival were 50% (95% CI, 38%-62%) and 53% (95% CI, 41%-65%), respectively. Age older than 30 years, more than 4 weeks to attain remission, and high-risk karyotypes, for example, t(9;22) or t(4;11), were adverse features contributing to the identification of 3 prognostic risk groups with 0, 1, and 2 adverse features, respectively: standard (47%), intermediate (36%), and high (17%). The 10-year cumulative incidences of relapse (20%, 48%, 85%; P < .0001) and probabilities of DFS (72%, 41%, 10%; P = .0003) were significantly different among these groups. In Cox analysis of the 71 patients alive and well 120 days after transplantation, those receiving 2 or 3 maintenance chemotherapy agents had significantly lower relapse rates and superior DFS compared with those receiving 0 or 1 agent. Our data suggest that maintenance chemotherapy improves the outcome of patients with ALL undergoing autografting. However, it is unlikely that autograft-based strategies are optimal for the high-risk group of patients who should be considered for alternative-donor allograft procedures.

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therapy according to the United Kingdom MRC’s adult UKALL XII chemotherapy protocol, which is more intense. Induction chemotherapy was administered either at the Royal Marsden Hospital or at other regional hospitals from where the patients were referred for autografting in first CR.

**CNS prophylaxis**

Patients treated with melphalan-TBI as part of the conditioning received an additional 800 cGy cranial (no CNS disease at presentation) or craniospinal (CNS disease at presentation) irradiation in 5 daily fractions the week before the TBI. Patients treated with melphalan alone received 2400 cGy cranial irradiation in 5 daily fractions the week before the TBI. Patients treated with melphalan-TBI as part of the conditioning received an additional 800 cGy cranial (no CNS disease at presentation) or craniospinal (CNS disease at presentation) irradiation in 15 fractions after hematologic recovery from intensification or phase II induction and before leukapheresis. No patient received testicular irradiation.

Patients without CNS disease received 6 injections of intrathecal MTX (usually 15 mg) over the treatment period prior to the transplantation, and those with CNS disease received triple intrathecal chemotherapy with MTX (usually 15 mg), cytarabine (usually 30 mg), and hydrocortisone (usually 100 mg) until 6 consecutive spinal fluid samples were free of disease. No intrathecal chemotherapy was administered after transplantation."
Table 4. Time to starting maintenance chemotherapy

<table>
<thead>
<tr>
<th>Drug</th>
<th>No. (%)</th>
<th>Marrow (%)</th>
<th>Blood (%)</th>
<th>P</th>
<th>Time to starting drug, d (range)</th>
<th>Probability of being on drug 1 y after transplantation</th>
</tr>
</thead>
<tbody>
<tr>
<td>6MP</td>
<td>55 (71)</td>
<td>25 (71)</td>
<td>30 (71)</td>
<td>.18</td>
<td>61 (15-678)</td>
<td>All, % 74, Marrow, % 84, Blood, % 67</td>
</tr>
<tr>
<td>MTX</td>
<td>44 (57)</td>
<td>18 (51)</td>
<td>28 (62)</td>
<td>.007</td>
<td>114 (34-891)</td>
<td>All, % 51, Marrow, % 60, Blood, % 46</td>
</tr>
<tr>
<td>VP</td>
<td>29 (38)</td>
<td>8 (23)</td>
<td>21 (50)</td>
<td>.015</td>
<td>83 (24-692)</td>
<td>All, % 38, Marrow, % 28, Blood, % 47</td>
</tr>
</tbody>
</table>

Supportive care

All patients were treated in protective isolation in rooms with positive-pressure ventilation. Blood products transfused were not screened for cytomegalovirus (CMV) antibody in CMV-seropositive patients. Antibiotic prophylaxis and therapy varied in accordance with prevalent practices and research programs. Broad-spectrum antibiotic therapy was started for fever in the neutropenic phase. Irradiated random platelets were transfused to maintain the platelet count at 20 × 10^9/L and packed cells to maintain the hemoglobin at 100 g/L.

Maintenance chemotherapy

Maintenance chemotherapy with daily 6MP was usually begun when the hemoglobin reached 100 g/L. Therapy was continued 2 years after starting therapy unless the dose of 6MP reached 75 mg. Weekly oral MTX at a dose of 5 mg (increasing in 2.5- to 5-mg steps) was added when the dose of 6MP reached 75 mg. All maintenance chemotherapy was discontinued 2 years after starting the first agent (6MP).

Drug doses were adjusted to maintain the absolute neutrophil count over 1 × 10^9/L. Initially, monthly vincristine (1.4 mg/m^2; maximum 2 mg) with prednisone (40 mg/m^2) was administered to patients who could not tolerate myelosuppressive chemotherapy. After 1995, this was added routinely to all patients. The target doses of 6MP and MTX were 80 mg/m^2 daily and 20 mg/m^2 weekly, respectively. Patients received folic acid and prophylaxis for Pneumocystis carinii pneumonia with oral trimethoprim-sulfamethoxazole (if on 50% of the target 6-MP dose) or aerosolized pentamidine (if on <50% of target 6-MP dose) while on maintenance chemotherapy. Acyclovir was administered to prevent reactivation of Varicella zoster virus.

The average dose of 6MP and MTX administered was determined by calculating the actual total amount of drug administered after the transplantation and dividing this amount by the number of days (6-MP) or weeks (MTX) from the date of the transplantation to the last date of the chemotherapy. The latter was the day on which chemotherapy was electively discontinued after completion of 2 years or the day of relapse or transplant-related mortality (TRM). The body surface area used for calculations was that at the time of the transplantation procedure.

Statistical analysis

The χ^2 test was used to compare categoric variables, and the Wilcoxon rank-sum test was used to compare continuous variables. The probabilities of disease-free survival (DFS) and overall survival (OS) were estimated by the Kaplan-Meier method, and compared using the log-rank test. The cumulative incidence of TRM and relapse was estimated using each type of event as a competing risk for the other. The significance of differences in TRM and relapse was calculated using the likelihood-ratio statistic for proportional-hazards regression models. Two patients dying in CR of causes obviously unrelated to the transplantation and the underlying disease (homicide and pre-existing ischemic heart disease at 8 and 36 months, respectively) were censored at the time of death for OS and DFS, and were considered competing events for calculating the cumulative incidence of TRM and relapse.

The disease risk stratification was based on age, cytogenetics, and the time taken to attain CR (Tables 2 and 3), and was a modification of the German classification proposed by Hoelzer et al. The following factors were analyzed in Cox proportional-hazards regression models for effect on relapse, DFS and OS: disease risk (standard versus intermediate versus high), sex, CR-transplantation interval (<4 versus ≥4 months), stem cell source (marrow versus blood), type of induction therapy (UKALL X or similar versus UKALL XII or similar), white blood cell (WBC) count at presentation (<30 versus ≥30 × 10^9/L), intensity of maintenance therapy (0/1 agent versus 2/3 agents), and immunophenotype (null versus others). The conditioning regimen was not included in the model because of 100% concordance between the stem cell source and conditioning.

Patient follow-up data are current through August 1, 2001, when the median, minimum, and maximum follow-up duration for surviving patients was 6.9, 2.5 (excluding the patient who was murdered 8 months after autografting), and 17 years, respectively.

Results

Neutrophil recovery to 0.5 × 10^9/L was complete and sustained in all patients, occurring earlier in PBSCT recipients than ABMT recipients. The stem cell source did not influence platelet recovery significantly.
TRM and morbidity

Six patients died due to transplant-related causes 2 to 8 months (median, 5.5 months) after the ABMT procedure, and none after the PBSCT procedure (P = .007; Fisher exact test). All deaths were due to interstitial pneumonitis and were most likely the result of TBI, which was used for ABMT but not for PBSCT (Table 1). The cumulative 10-year incidence of TRM was 8% (95% CI, 4%-17%).

Maintenance chemotherapy

Table 4 shows the time to starting each maintenance chemotherapy agent and the proportion of patients starting it for the entire group as well as for ABMT and PBSCT recipients separately. The only significant difference—the higher proportion of PBSCT recipients receiving VP—is the result of changed treatment strategy (routine use of VP rather than only in patients with poor marrow function). None of the 22 patients who did not receive 6MP received MTX, whereas 5 received VP.

Seventeen patients received no maintenance chemotherapy at all because of early TRM within 4 months (n = 3), pulmonary problems subsequently terminating in TRM beyond 4 months (n = 2), relapse within 4 months (n = 3), sluggish hematologic recovery terminating in relapse beyond 4 months (n = 3), unknown/patient preference (n = 5), and noncompliance/lifestyle (n = 1; this patient eventually became a homicide victim). Nine patients received only a single drug (5 VP and 4 6MP), 34 patients received 2 drugs (27 6MP-MTX and 7 6MP-VP), whereas 17 patients received all 3 drugs.

The average daily dose of 6MP administered was 3 to 109 mg/m² (median, 33 mg/m²). The average weekly dose of MTX administered was 0.2 to 18.1 mg/m² (median, 2.8 mg/m²). Maintenance chemotherapy was tolerated well, and other than transient myelosuppression requiring dose reduction or temporary cessation of therapy, no other side effects were seen.

Relapse and therapy after relapse

Thirty patients (39%) had a relapse at 1.5 to 80 months (median, 12.5 months); 15 in the first year, 7 in the second year, 1 in the third, 5 in the fourth, 1 in the fifth, and 1 at almost 7 years. All relapses were confirmed to be immunophenotypically and cytogenetically (where applicable) identical to the original disease. No case of second or secondary leukemia was seen. Figure 1 shows the cumulative incidence of relapse.

Eighteen patients died of relapsed disease or toxicity of salvage chemotherapy. Eleven patients underwent an allograft from an HLA-matched sibling (n = 6) or a matched unrelated donor (n = 5) in second CR, and 1 is awaiting allograft. Nine died of transplant-related toxicity (n = 7) or relapse (n = 2), and 2 are alive and well in second CR at 7 years (HLA-identical sibling) and 1.5 years (unrelated donor).

OS and DFS

As of August 1, 2001, 44 of 77 patients were alive and in remission 8 months to 17 years (median, 6.9 years) after the transplantation—41 in continuous first CR, 2 in second CR after allogeneic BMT, and 1 in second CR after chemotherapy (awaiting allograft). Figures 2 and 3 show the actuarial probabilities of DFS and OS. All patients are off maintenance chemotherapy.
Identification of risk groups

The 3 adverse risk factors, age older than 30 years, t(9;22), t(4;11), and more than 4 weeks to attain CR (Table 2), were identified to be high-risk factors for relapse, DFS, and OS (Table 3). As Table 2 shows, these factors could be combined to produce 3 risk groups. The outcome of these 3 groups in terms of relapse (Figure 4), DFS (Figure 5), and OS \((P = .003)\) was highly significantly disparate.

The effect of maintenance chemotherapy

Among patients alive and well at 120 days, those getting 2 or 3 maintenance chemotherapy agents had a significantly lower relapse rate (Figure 6), higher DFS (Figure 7), and OS \((P = .0008)\). This time point was chosen to eliminate the 6 patients dying early, 3 of relapse and 3 of TRM, before becoming eligible to receive maintenance chemotherapy.

In the same population of patients, there was no significant difference in the outcome of those getting no maintenance chemotherapy and those getting any maintenance chemotherapy (data not shown). There was no obvious relationship between the amount of chemotherapy given and outcome (data not shown).

Cox analysis

Table 5 shows the results of the multivariable analysis adjusted by survival on day 120 after transplantation. Standard-risk disease, longer CR-autograft intervals, and more intense maintenance chemotherapy were associated with lower relapse and higher DFS.

OS was beneficially affected by standard-risk disease and more intense maintenance chemotherapy.

Effect of maintenance chemotherapy by risk group

Table 6 shows that receiving more intensive maintenance chemotherapy made a significant difference in outcome in the standard-risk group and a less-marked difference to patients in the intermediate-risk group. On the other hand, it made no difference to patients in the high-risk group at all.

Discussion

Our data suggest that standard ALL-type maintenance chemotherapy can be administered safely after autotransplantation in patients with ALL. Although only a randomized study can answer the question of its benefit, if any, 2 observations in this series of patients suggest that maintenance chemotherapy may contribute to an improved outcome in terms of reduced relapse and increased survival. The first is that higher intensity of maintenance therapy was clearly associated with better outcome. The second is that the results reported here are better than most reported using the more conventional approach of autograft without any posttransplantation intervention.

An obvious shortcoming of the first observation here is that autografted patients with compromised myeloid reserve have a higher risk of disease recurrence. These are also the patients who are likely to be unable to start chemotherapy after transplantation. This obstacle was partially overcome by excluding patients having relapse early. However, even after excluding the 3 patients who had a relapse within 4 months, 4 additional patients continued to have poor counts and eventually had relapse without being able to start any maintenance therapy at all.

Another indirect observation that may support using posttransplantation therapy in ALL comes from the International Bone Marrow Transplant Registry, which showed that the use of methotrexate after transplantation as graft-versus-host disease (GVHD) prophylaxis decreased relapse in patients undergoing allografting for ALL independently of clinically obvious GVHD, perhaps secondary to a direct antileukemic effect.27

In a French cooperative group study that showed no impact of ABMT on survival compared to chemotherapy alone, late relapses were significantly less common among autografted patients,12 and relapse relatively early after the transplantation was an important cause of treatment failure. It is conceivable that posttransplantation
The data in Table 6 suffer from very small patient numbers in the various subgroups and must be interpreted with caution. They suggest that the outcome of patients with high-risk disease is dismal with autotransplantation whether posttransplantation chemotherapy is given. Maintenance chemotherapy in a setting like this may reduce the incidence of early posttransplantation relapse and improve DFS by eliminating clonogenic leukemic cells reinfused with the graft, or more likely, have remained viable in the patient despite the conditioning regimen.

Other posttransplantation treatments that have been used to decrease relapse rates after autotransplantation in ALL have included intensive chemotherapy, interleukin 2 infusions, and cell-mediated immunotherapy with haploidentical T lymphocytes. Reports describing these approaches have suffered from small patient numbers and inadequate follow-up; making any assessment difficult.

Patients with HLA-identical siblings in the ABMT era received allografts at the Royal Marsden Hospital. Those with HLA-identical siblings in the PBSCT era underwent autografting and the allograft was reserved for salvage therapy of relapse. The reasoning behind this was avoidance of toxic therapy (allograft) unless absolutely essential. Although treatment-related mortality was indeed reduced with the elimination of TBI and the use of blood-derived stem cells, this did not necessarily make the outcome of subsequent salvage therapy any easier or more successful; only 2 of 11 patients receiving allografts survive, with high TRM rates that were no different from rates we had observed in the past. These patients were conditioned using conventional-intensity regimens. It is possible that the outcome may have been better if reduced-intensity regimens had been used. In any case, we feel that an allograft from an HLA-identical sibling is probably the best treatment option in adult patients with ALL who require a transplantation in first CR, especially with the possibility of augmented graft-versus-leukemia reactions from the use of blood-derived stem cells and limited TRM with the use of an adequate number of CD34+ cells for the transplant.

What is the role of the pretransplantation conventional chemotherapy administered for induction and intensification in consolidation? It was disappointing to see that there was no difference in the outcome of patients receiving less intensive therapy initially (UKALL X and similar) compared with those getting more intensive therapy (UKALL XII). This is unlike in acute myeloid leukemia (AML) where in vivo purging before transplantation, achieved by administering at least 2 cycles of consolidation chemotherapy, makes a significant impact on outcome. Because graft source and conditioning regimen were changed simultaneously, and there was a strong association between these and the initial induction therapy, the relative contribution of each to outcome cannot be assessed. A beneficial effect of TBI of reduction of relapse cannot be ruled out because the higher-intensity induction chemotherapy that non-TBI patients received could have offset the disadvantage of not having received TBI. It is also possible that the difference between less and more intensive initial therapy is negated to some extent by posttransplantation maintenance therapy.

There has been a strong trend toward the use of alternative-donor transplants rather than autografts in ALL because of poor results of autotransplantation in ALL. However, high-risk allografts may not be the right approach for all patients. A recent registry comparison of unrelated donor transplants and autografts in first and second remission ALL found that although relapse rates were lower with allografts, the benefits of this were negated by significantly higher TRM. Because of this, survival was comparable with the 2 modalities of transplantation. In view of this, it is clearly important to identify patients who are likely to do poorly with an autograft and offer them alternative-donor transplantation. Those without features that increase the risk of relapse to prohibitive levels may be better off with an autograft and offer them alternative-donor transplantation.

The data in Table 6 suggest that the outcome of patients with high-risk disease is dismal with autotransplantation whether posttransplantation chemotherapy is given.
administered or not. This is the group of patients most likely to benefit from high-risk allograft approaches. What should be done for patients with standard-risk disease? It is tempting to suggest an autograft as the first approach even in patients who have an HLA-identical sibling donor because of the excellent outcome seen here. Relapsing patients can then be salvaged by an allograft in the second remission. However, we do not have adequate data to suggest an optimum approach in this situation, especially because of the poor outcome of allografted patients in this series. Patients with intermediate-risk disease ought to be allografted if an HLA-identical sibling is available. In the absence of a sibling donor, an autograft should be the preferred mode of therapy, to be followed by a salvage, high-risk, alternative-donor allograft in case of relapse.

Within the limitations of the data in Table 6 (small patient numbers), the suggestion that patients with the best disease also respond further to the maximum extent with posttransplantation therapy is interesting and is in keeping with observations in AML that patients with good-risk karyotypes gain the most from further intensification of therapy; either with chemotherapy or with autotransplantation.37

We conclude that autotransplantation followed by maintenance chemotherapy is an excellent way of intensifying therapy in adult patients with ALL unless the disease is high-risk. This is a strategy that should ideally be explored in a randomized trial. In the absence of such a study, it appears safe enough to use routinely in practice to improve outcome of autografted ALL patients.

References

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