ORIGINAL ARTICLE

Romaine Viollier · Jakob Passweg · Michael Gregor · Geneviève Favre · Thomas Kühne · Catherine Nissen · Alois Gratwohl · André Tichelli

Quality-adjusted survival analysis shows differences in outcome after immunosuppression or bone marrow transplantation in aplastic anemia

Received: 26 April 2004 / Accepted: 22 July 2004 / Published online: 31 August 2004 © Springer-Verlag 2004

Abstract Bone marrow transplantation (BMT) and immunosuppression (IS) have improved the prognosis of aplastic anemia; both treatments have specific advantages and drawbacks but similar survival rates. Analysis of additional endpoints may help in treatment decisions. In a single-center study, patients with aplastic anemia treated with IS (n=155) or BMT (n=52) were compared for survival, event-free survival, and quality-adjusted time without symptoms and toxicity (Q-TWiST). Probability of overall and event-free survival at 15 years was similar among both groups (BMT 51±15% and 25±14%, IS 53 $\pm 10\%$ and $27\pm 8\%$), with more early deaths in the transplant group and more late deaths in the IS group. There were differences in terms of mean duration of seven analyzed health states: time with symptoms from treatment-related toxicity (IS 0.36 years, BMT 0.27), transfusion dependency (IS 0.66 years, BMT 0.1 years), partial remission (IS 3.27 years, BMT 1.42), and secondary clonal disorder (IS 0.68 years, BMT 0.04) was significantly longer for IS compared to BMT ($p \le 0.001$). Patients treated with BMT spent more time with extensive chronic graft-versus-host disease (GvHD) (IS 0 years, BMT 0.96, p<0.023) and in CR without drugs (IS 1.22 years, BMT 2.43, p=0.056). In conclusion, survival, event-free survival, and Q-TWiST are similar. BMT-treated patients had longer periods free from symptoms, while IS-treated patients needed closer medical care, transfusion support, and medications.

Keywords Aplastic anemia · Treatment decisions · Quality of life

Introduction

Severe aplastic anemia is a rare disease characterized by pancytopenia and bone marrow aplasia. The goal of the treatment is to improve peripheral blood counts in order to free the patients of transfusions and the risk of bleeding and opportunistic infections. Bone marrow transplantation (BMT) as well as immunosuppression are standard treatment modalities that have improved the previously poor prognosis [1-6]. After any of the two treatments, more than 80% of patients are free from transfusions and some of them even normalize their cell counts. BMT is the standard approach for younger patients with an HLAmatched sibling donor, while immunosuppression is the treatment of choice for patients not eligible for BMT. In patients over the age of 40, it is less clear whether they should receive a BMT or antithymocyte globulin (ATG) even if they do have a matched sibling donor. With BMT, the defective organ is replaced by healthy marrow. Immunosuppressive treatment on the other hand will not cure the disease. It aims at eliminating "autoaggressive" cells that are responsible for the aplastic marrow resulting in pancytopenia. Despite these differences, there is no clear-cut advantage in terms of overall survival for either treatment form. Each treatment modality has its advantages and drawbacks [7–12].

Overall survival does not sufficiently describe the outcomes of patients with aplastic anemia. Therefore, one might want to take additional endpoints such as event-free survival, the ability to tolerate treatment, and quality of life during different sequences of relevant health states into account in the decision making process, when different treatment options result in similar long-term

R. Viollier \cdot J. Passweg \cdot G. Favre \cdot C. Nissen \cdot A. Gratwohl \cdot A. Tichelli (\boxtimes)

Transplantation Unit, University Hospitals of Basel,

Petersgraben 4,

4031 Basel, Switzerland e-mail: tichelli@datacomm.ch

Tel.: +41-61-2654254 Fax: +41-61-2654455

M. Gregor

Hämato-Onkologie, Kantonsspital Luzern, Luzern, Switzerland

T. Kühne Universitätskinderspital beider Basel, Basel, Switzerland survival. In aplastic anemia, quality of life outcomes are affected by treatment toxicity, graft rejection, infections, and graft-versus-host disease (GvHD) after BMT [13–15], and by persisting cytopenia due to slow or incomplete responses, relapse of aplastic anemia, or the development of clonal disorders such as myelodysplastic syndrome (MDS) or paroxysmal nocturnal hemoglobinuria (PNH) after immunosuppression [16–18].

In order to integrate quality of life considerations into comparing the two treatment modalities, additional factors such as treatment toxicity, transfusion and drug requirement, chronic GvHD, or clonal evolution have to be considered. The quality-adjusted time without symptoms and toxicity (Q-TWiST) methodology splits up survival times into several distinct periods associated with relevant health states weighted by quality of life factors [19–23]. We compared in a retrospective, single-center study aplastic anemia patients treated with BMT to those treated with immunosuppression applying survival, event-free survival, and Q-TWiST methodology.

Patients and methods

Study population

Patients with aplastic anemia as defined by the criteria of Bacigalupo and Camitta [24, 25] were treated since 1976 in a prospective single-center study at the University Hospital of Basel. Patients younger than 40 years of age with an HLA-matched sibling donor received allogeneic

Table 1 Characteristics of the patients with aplastic anemia treated with bone marrow transplantation (BMT) or antithymocyte globulin (ATG)

BMT ATG p value Patients(n)52 155 Male gender 27 (52%) 85 (55%) 0.418 Age at first treatment Median age (range) (years) 19 (2-55) 23 (2-74) 0.039 Patients >40 years 3 (6%) 37 (24%) 0.047 Etiology Idiopathic 42 (81%) 125 (81) 0.828 Viral 3 (6%) 10 (6%) Toxic/drugs 7 (13%) 18 (12) 0 (0%) Others 2 (1%) Severity of aplastic anemia^a Non-severe 14 (27%) 50 (32%) 0.759 Severe 23 (44%) 65 (42%) Very severe 15 (29%) 40 (26%) Median blood values at diagnosis (range) Neutrophil counts ($\times 10^9 l^{-1}$) 0.45(0-3.0)0.46(0-2.1)0.536 Platelet counts (×10⁹ l⁻¹) 14 (1-97) 15 (1-218) 0.450 Reticulocyte counts ($\times 10^9 l^{-1}$) 14 (0-77) 18 (0-186) 0.384 MCV (fl) 96 (84-126) 97 (80-133) 0.948 Transfusion dependence 51 (98%) 141 (91%) 0.78 Splenectomy 2 (4%) 84 (54%) < 0.001 Median time interval (range) Diagnosis first-line treatment (months) 1.7(0.2-14)1.2 (0.03-378) 0.130

11.5 (2-22)

11.3 (0.2-22)

0.439

Follow-up (years)

BMT, older patients and patients without a marrow donor were treated with ATG-based immunosuppression. On 1 January 1999, 207 patients were included in the study. Fifty-two underwent marrow transplantation and 155 received ATG. Of the 155 patients receiving ATG as first-line treatment, 8 were subsequently transplanted. These eight patients are included in the ATG group.

The characteristics of the patients are shown in Table 1. Of the 207 patients, 55 fulfilled the criteria of very severe aplastic anemia, 88 had severe, and 64 patients non-severe aplastic anemia. Patients with non-severe aplastic anemia had either thrombocytes $<10\times10^9$ l⁻¹, or neutrophils $<0.5\times10^9$ l⁻¹, or depended on transfusions. Both treatment groups were comparable for gender, etiology, severity of the disease, and for blood values before first-line treatment. The median age at first-line treatment was 19 years (range: 2–55 years) for BMT compared to 23 years (range: 2-74 years) in ATG-treated patients (p=0.039). The difference in frequency of splenectomy is due to a protocol of splenectomy in nonresponders to ATG prior to salvage immunosuppressive retreatment [26]. All patients were either treated in laminar air flow rooms or in single rooms with reverse isolation. They all received standard supportive care, including transfusions of red blood cells and single-donor platelet concentrates, as well as application of broad-spectrum intravenous antibiotics in case of fever or infections. The median duration of observation since first-line treatment in surviving patients was similar in both treatment groups, i.e., 11.5 years (range: 2-22 years) for BMT patients and 11.3 (range: 0.2–22 years) for patients treated with ATG.

^aAccording to Bacigalupo et al. [24], Camitta et al. [25]

Treatment protocol

The treatment protocol and overall results of the study have been published previously [27–29]. In short, the ATG treatment protocol was as follows: from 1976 to 1991 equine antithymocyte globulin from the same manufacturer (Lymphoser, Swiss Serum and Vaccine Institute, Berne, Switzerland) was used in a dose of 40 mg/kg per day for 4 days. Since 1992 a different equine antithymocyte globulin preparation (Lymphoglobuline), provided by SangStat (Fremont, Calif., USA), was used in a dose of 15 mg/kg per day for 5 days. Methylprednisolone was given in a dose of 1 g/day for all patients treated between 1981 and 1992 and in a dose of 2 mg/kg per day thereafter. Cyclosporine in a dose of 5 mg/kg per day was added to the regimen in 1991 and continued for at least 1 year.

All patients treated with BMT were conditioned with cyclophosphamide at a dose of 50 mg/kg per day for 4 days. Twenty-one patients received unirradiated donor buffy coat after transplantation. Since 1994, ATG was added at a dose of 30 mg/kg daily for 3 days (Lymphoglobuline, SangStat, Fremont, Calif., USA). For prevention of GvHD patients received methotrexate alone until 1979. Since July 1979, patients were given cyclosporin A, and since 1990 a combined regimen of methotrexate and cyclosporine was applied except for children, who continued to receive cyclosporine alone. Bone marrow was used as a source of stem cells in 51 of the 52 transplanted patients and peripheral stem cells in one. The donor was an HLA-identical sibling in 51 and a syngeneic donor in one patient.

Data analysis and definitions

A detailed review of all patient charts was the basis of this report. Variables analyzed included disease-related information, treatment (first-line and subsequent), time to response, time to a first event, and time to death or last follow-up. Clinical state (transfusions and medication after first-line treatment) and biological investigations [blood counts, bone marrow examination, screening tests for PNH, including hemosiderin in the urine, Ham's (acid hemolysin), and sucrose test] were measured before, 3, 6, and 12 months after first-line treatment, and then annually until last follow-up.

Response to treatment was defined as follows: complete remission (CR): neutrophil counts ≥1.5×10⁹ l⁻¹, hemoglobin ≥120 g/l, and platelet counts ≥150×10⁹ l⁻¹; partial remission: patients not fulfilling the criteria of CR, but achieving transfusion independence; nonresponse: patients who never fulfilled criteria of complete or partial remission, died from aplasia, or needed a second-line treatment before fulfilling criteria of partial or complete remission. An event was defined as nonresponse, relapse of aplastic anemia, development of a clinically relevant clonal complication such as MDS, PNH, or solid tumor, extensive chronic GvHD, or death. Criteria for late clonal transformation have been published in detail [30]. Labo-

ratory signs of PNH without clinical manifestation, limited chronic GvHD, and silent nonmalignant tumors (liver adenomas) were not considered as an event.

Causes of death were subdivided into disease-related deaths, treatment-related deaths, event-related deaths, and deaths independent from disease or treatment. Bleeding and infections were considered disease related as long as they occurred during the aplastic phase. For ATG patients, death was treatment related when it occurred during or immediately after ATG infusion. For transplanted patients, treatment-related deaths included deaths due to toxicity, acute GvDH, infections occurring after hemopoietic regeneration, interstitial pneumonia, and early graft rejection. Death occurring after relapse, late graft rejection, MDS, PNH, or solid tumors were classified as event-related deaths. In all other cases, the cause of death was classified as not related to aplastic anemia or its treatment.

Statistical analysis

The chi-square test was used for comparisons of categorical data and the Mann–Whitney U test for continuous unrelated variables among groups. Survival probabilities were estimated using the Kaplan–Meier method [31]. The log-rank test was used to compare survival [32]. Outcomes analyzed between BMT- and ATG-treated patients were overall survival, event-free survival (considering death, extensive chronic GvHD, secondary malignancy, MDS or clinical PNH, and nonresponse to treatment or relapse of aplastic anemia as events), and quality-adjusted survival.

The O-TWiST is based on subdividing survival times in distinct successive time lengths of relevant health states. For comparison of two cancer treatment strategies usually three states are defined, time with treatment-related toxicity, time without symptoms and toxicity (TWiST), and time from treatment failure to death [33, 34]. In aplastic anemia the situation is more complex. We defined the following clinically relevant health states (Table 2): the time in treatment-related toxicity (TOX) was arbitrarily fixed for patients of both treatment groups and for each treatment course to 3 months. TWiST was the time an individual spent in complete remission, free from medication, and without an event, as described above. The other relevant health states were time with clonal disease (CLON), time with extensive chronic GvHD (GvHD), time with transfusion dependency (TRANS), time with partial remission (PR), or time with complete remission but still on medication (CR). A hierarchy of the different health states was established in case a patient was in two health states at the same time. Accordingly, time with CLON stood over all other health states. After that came in the following order: time with GvHD, time with TRANS, time with PR, time with CR, and finally time with TWiST. A utility factor was allocated to each health state varying on a scale from 0 (as bad as death) to 1 (TWiST) to reflect the health state's quality of life value relative to time in TWiST. The utility coefficient (μ) was defined for TWiST

as 1, for CR and PR as 0.75, for TOX as 0.25, and for all other states as 0.5. The weighting of the utility factor was based on the 30 years of clinical experience of the study center and was done like the weighting of the original groups of Q-TWiST. Mean Q-TWiST for each treatment modality was then calculated from the mean clinical health state duration as

$$\begin{aligned} \mathbf{Q} - \mathbf{TWiST} &= [\mu_{\text{TOX}} \times \mathbf{TOX}] + [\mathbf{TWiST}] + [\mu_{\text{CLON}} \times \mathbf{CLON}] \\ &+ [\mu_{\text{GvHD}} \times \mathbf{GvHD}] + [\mu_{\text{TRANS}} \times \mathbf{TRANS}] \\ &+ [\mu_{\text{PR}} \times \mathbf{PR}] + [\mu_{\text{CR}} \times \mathbf{CR}] \end{aligned}$$

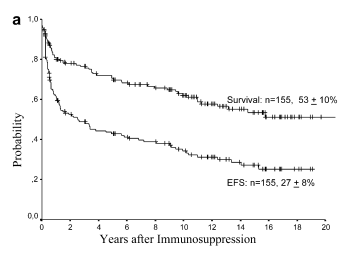
where μ_{TOX} is the utility coefficient for toxicity, μ_{CLON} for clonal complications, μ_{GvHD} for chronic GvHD, μ_{TRANS} for transfusion dependency, μ_{PR} for partial remission, and μ_{CR} for complete remission but still with drugs.

A threshold utility analysis comparing BMT with ATG treatment was performed for PR $(U_{\rm PR})$, which was the most relevant health state in terms of duration and differences between treatment groups. The threshold utility analysis was calculated for all possible values ranging from 0.0 to 1.0 (Fig. 2). Finally, Kaplan–Meier estimates for the time to events that signal transitions between clinical health states were used to partition the area under the overall survival curves separately for each treatment modality [35].

Results

Survival and event-free survival

At the time of last follow-up, 122 of 207 patients [59%; 95% confidence interval (CI) 52-65%] were alive and 85 (41%; 95% CI 34–48%) had died, 24 (46%; 95% CI 32–62%) after BMT and 61 (39%; 95% CI 32–47%) after ATG treatment (p=0.241). The probability of survival at 15 years was statistically similar between both treatment groups; it was $51\pm15\%$ after transplantation and $53\pm10\%$



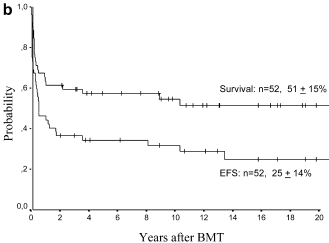


Fig. 1 a Actuarial survival and event-free survival (*EFS*) at 15 years according to Kaplan–Meier among 155 patients with aplastic anemia treated with ATG. **b** Actuarial survival and event-free survival (*EFS*) at 15 years according to Kaplan–Meier among 52 patients with aplastic anemia treated with BMT

after ATG (p=0.20), with more early deaths in the transplant group and more late deaths in the immunosuppression group (Fig. 1). There was an improvement in

Table 2 Definitions and criteria of health states for Q-TWiST analysis. MDS myelodysplastic syndrome, PNH paroxysmal nocturnal hemoglobinuria, GvHD graft-versus-host disease

| Name | Definition | Criteria | Utility coefficient |
|-------|--|---|---------------------|
| TWiST | Time without symptoms and toxicity (and without drugs) | Time in complete remission, without drugs, transfusions, clonal disease, without extended GvHD, and not in toxicity phase | 1 |
| TOX | Toxicity | Initial period of treatment, arbitrarily fixed for each course to 3 months | 0.25 |
| TRANS | Time with transfusion dependency | Time with transfusion dependency, but without clonal disease or extended GvHD, and not in toxicity phase | 0.5 |
| PR | Time in partial remission | Time not in complete remission, but without transfusion dependency, clonal disease, or extensive GvHD, and not in toxicity phase | 0.75 |
| CR | Time in complete remission | Time in complete remission, but still receiving drug medication, without clonal disease, without extended GvHD, and not in toxicity phase | 0.75 |
| GvHD | Time with extensive GvHD | Time with extensive chronic GvHD, but not in toxicity phase | 0.5 |
| CLON | Time with clonal disease | Time with a clonal disease, such as MDS, PNH, or solid tumor | 0.5 |

survival probabilities by decade for both treatment groups with probabilities at 5 years before 1980 of $40\pm22\%$ (BMT) and $55\pm18\%$ (ATG), between 1980 and 1990 of 60 $\pm22\%$ (BMT) and $74\pm9\%$ (ATG), and after 1990 of 83 $\pm21\%$ (BMT) and $75\pm18\%$ (ATG). This improvement by decade was statistically significant for the transplanted patients (p=0.05), but not for the ATG-treated group (p=0.1). The event-free survival at 15 years was similar between both treatment groups (p=0.12), with $25\pm14\%$ after BMT and $27\pm8\%$ after ATG treatment (Fig. 1).

Events and causes of death

Frequency and causes of events and deaths are listed in Table 3. Fifty-three events were observed in 52 patients treated with BMT and 196 events in 155 patients treated with ATG. Of the 52 patients treated with BMT, 21 patients had one and 16 patients two events. Of the 155 patients treated with ATG, 34 patients had one, 39 patients two, and 23 patients three or more events. During the study period, 15 of 52 (29%) BMT-treated and 59 of 155 (38%) ATG-treated patients never had an event. The overall frequency of events was not different between patients treated with BMT and ATG (p=0.302). However, there were differences in the type of an event between both groups. Primary refractory disease occurred significantly more often after immunosuppression (41 of 196 events, 21% vs 5 of 53 events, 9%, p=0.087), and MDS or leukemia (21 of 196, 11%, p=0.009) or PNH (18 of 196, 9%, p=0.016) were observed only after immunosuppression. Extensive chronic GvHD was observed only after BMT (12 of 53, 23%, p<0.001). There was no difference in the frequency of relapse/rejection or the occurrence of a solid tumor between both treatment groups. There were clear differences in cause of death between both groups (Table 3). Major cause of death in BMT patients was treatment related (46%), followed by disease-related deaths (33%). In contrast, after ATG treatment, the major cause of death was either disease related (41%) or event related (41%). Deaths due to toxicity were less frequent than in BMT patients (Table 3).

Q-TWiST analysis

The mean observation time per patient did not differ between the treatment groups. It was 6.41 years for transplanted patients and 7.80 years for ATG-treated patients (*p*=0.181). Important differences were found between both treatment modalities in terms of mean duration of various health states (Table 4). The mean time per patient spent with toxicity (ATG 0.36 years, BMT 0.27 years), with transfusion dependency (ATG 0.66 years, BMT 0.1 years), with partial remission (ATG 3.27 years, BMT 1.42 years), and with a clonal disorder (ATG 0.68 years, BMT 0.04 years) was significantly longer for ATG-treated patients compared to patients treated with BMT (*p*≤0.001). The difference in time spent with toxicity is

Table 3 Frequency and type of events, as well as causes of deaths in patients treated with bone marrow transplantation (BMT) or antithymocyte globulin (ATG)

| antitulyinocyte globuilli (ATG) | | | |
|-------------------------------------|----------|----------|---------|
| | BMT, | ATG, | p value |
| | n=52 | n=155 | |
| Patients with an event | 37 (71%) | 96 (62%) | 0.302 |
| With a single event | 21 (40%) | 34 (22%) | |
| With two events | 16 (31%) | 39 (25%) | |
| With three events | 0 (0%) | 23 (15%) | |
| Number/type of events | 53 | 196 | |
| Primary refractory | 5 (9%) | 41 (21%) | 0.087 |
| Relapse/late rejection | 10 (19%) | 50 (26%) | 0.411 |
| MDS/leukemia | 0 (0%) | 21 (11%) | 0.009 |
| Clinical PNH ^a | 0 (0%) | 18 (9%) | 0.016 |
| Extensive chronic GvHD ^b | 12 (23%) | 0 (0%) | < 0.001 |
| Malignant tumor ^c | 1 (2%) | 5 (2%) | 1.0 |
| Death | 24 (45%) | 61 (31%) | 0.077 |
| Others | 1 (2%) | 0 (0%) | 0.213 |
| Number/type of deaths | 24 (46%) | 61 (39%) | 0.241 |
| Treatment-related deaths | 11 (46%) | 8 (13%) | 0.003 |
| Early rejection | 3 | 1^{d} | |
| Acute GvHD | 7 | 0 | |
| Toxic | 1 | 7 | |
| Disease-related deaths | 8 (33%) | 25 (41%) | 0.686 |
| Bleeding | 1 | 14 | |
| Infection | 7 | 11 | |
| Event-related deaths | 3 (13%) | 25 (41%) | 0.024 |
| Chronic GvHD | 3 | 0 | |
| MDS or leukemia | 0 | 13 | |
| PNH | 0 | 2 | |
| Relapse of aplastic anemia | 0 | 9 | |
| Solid tumor | 0 | 1 | |
| Death without evident association | 2 (8%) | 3 (5%) | 0.618 |
| Car accident | 1 | 1 | |
| Heart disease | 1 | 1 | |
| Unknown | 0 | 1 | |

^aBiological paroxysmal nocturnal hemoglobinuria not included (*n*=12)

explained by the larger proportion of patients requiring retreatment in the ATG group. In contrast, patients treated with BMT spent more time with extensive chronic GvHD (0.96 years for BMT vs 0 for ATG, p<0.023) and in TWiST (BMT 2.43 years, ATG 1.22 years, p=0.056). None of the eight patients receiving ATG as a first-line treatment developed chronic GvHD after a subsequent transplant. The time spent in complete remission but still receiving medication was not statistically different between both groups, with 1.21 years for BMT and 1.60 years for ATG (p=0.431). Using the utility factors chosen previously, the Q-TWiST times were similar for the two treatment groups, i.e., 5.08 years after BMT and 5.72 years after ATG treatment (Δ -0.64, 95% CI -2.25 to +0.97, p=0.434). In terms of relative duration, there are

bLocalized cutaneous or mucosal chronic GvHD not included (n=5)

^cBenign tumors not included (*n*=3 liver adenomas) ^dAfter secondary bone marrow transplantation

Table 4 Quality-adjusted time without symptoms and toxicity (Q-TWiST) and drugs of patients with aplastic anemia treated with bone marrow transplantation (BMT) or antithymocyte globulin (ATG)

| Time interval | al BMT | | ATG | | Δ Between BMT and ATG | | |
|----------------------|----------------------------|----------------|----------------------------|----------------|------------------------------|-------------------------------|---------|
| | Mean years per patient (%) | Total years | Mean years per patient (%) | Total years | Mean years per patient | 95% CI | p value |
| Total years | 6.41 | 333.5 | 7.80 | 1,207.5 | -1.38 | -3.40/+065 | 0.181 |
| TWiST | 2.43 (38%) | 126.3 | 1.22 (16%) | 189.3 | +0.21 | +0.33/+2.45 | 0.056 |
| TOX | 0.27 (4%) | 14.3 | 0.36 (4%) | 55.8 | -0.09 | -0.12/-0.05 | < 0.001 |
| TRANS | 0.10 (1%) | 4.8 | 0.66 (8%) | 102.8 | -0.57 | -0.80/-0.35 | < 0.001 |
| PR | 1.42 (22%) | 73.8 | 3.27 (42%) | 507.0 | -1.85 | -2.93/-0.78 | 0.001 |
| CR with drugs | 1.21 (19%) | 63.0 | 1.60 (20%) | 248.3 | -0.40 | -1.37/+0.59 | 0.431 |
| GvHD | 0.96 (15%) | 50.0 | 0 | _ | +0.96 | +0.14/+1.78 | 0.023 |
| CLON | 0.04 (1%) | 2.0 | 0.68 (9%) | 104.5 | -0.64 | -0.88/-0.39 | < 0.001 |
| Q-TWiST ^c | 5.08 | 264.3 | 5.72 | 887.2 | -0.064 | / - 2.25/ +0.97 | 0.434 |

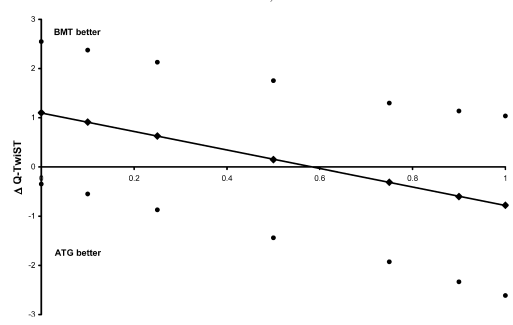
considerable differences between the health states: BMT patients spent for instance 38% of their time in TWiST and 19% in PR compared to 16% and 42%, respectively, for ATG-treated patients (Table 3).

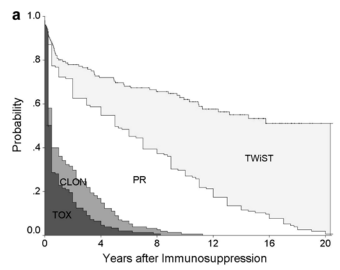
The threshold analysis of the PR utility factor is presented in Fig. 2. PR is the health state with the largest difference between the BMT and ATG groups. The plot illustrates the mean difference in Q-TWiST between BMT and ATG by varying the $U_{\rm PR}$ between 0 and 1. Irrespective of utility factor attributed to PR, there is no significant difference in Q-TWiST among both groups. Figure 3 shows the partitioning of survival times for each treatment group into the four clinically most relevant health states: TOX (including time with toxicity and time with transfusion dependency), PR (including time in partial remission and time in complete remission but requiring drugs), CLON for ATG group or GvHD for BMT group (including extensive GvHD and clonal complications), and TWiST.

Fig. 2 Threshold utility analysis was performed for the period "partial remission" (PR). The plot of mean difference in Q-TWiST between BMT- and ATG-treated patients using a utility factor (U_{PR}) between 0 and 1 is illustrated. The points on each side of the line representing the mean difference in Q-TWiST are the 95% confidence intervals

Discussion

BMT and immunosuppression are both effective treatment modalities for patients with aplastic anemia and 15-year survival probabilities are similar. More early deaths were observed in the transplant group and more late deaths in the ATG group. Event-free survival is an outcome measure that reflects the probability of a patient being alive without complications due to disease or treatment. This outcome measure has rarely been used in previous publications of aplastic anemia [36]. There were no significant differences in event-free survival at 15 years after BMT and ATG treatment. However, there were significant differences in outcomes affecting the quality of life of these patients. The O-TWiST analysis allows for a comparison of the time period spent in different health states when overall survival does not show a preference between the two treatment modalities. The quality-adjusted time without symptoms, toxicity and drugs is similar after BMT and ATG treatment, but both treatment modalities differ in





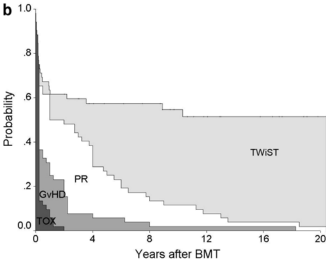


Fig. 3 a Partitioning of survival time for patients treated with ATG into the four clinically most relevant health states: TOX (including time with toxicity and time with transfusion dependency), PR (including time in partial remission and time in complete remission but necessitating drugs), CLON (including clonal complications), and TWiST. b Partitioning of survival times for patients treated with bone marrow transplantation into the four clinically most relevant health states: TOX (including time with toxicity and time with transfusion dependency), PR (including time in partial remission and time in complete remission but necessitating drugs), GvHD (including extensive GvHD and clonal complications), and TWiST

time periods spent in various health states. ATG-treated patients spent more time requiring transfusions and drugs, endure longer periods with abnormal blood values, and have a higher risk of clonal disorder. Patients in the ATG group required additional treatment more frequently than BMT patients due to more relapses and a higher incidence of refractory disease. In contrast, BMT patients spent nearly 60% of their time without symptoms and toxicity (in TWiST and CR with drugs).

The implications of these findings must be considered in the context of benefits of treatment over time. Although overall survival, event-free survival, or Q-TWiST do not favor one of the two treatments, BMT patients spent more time free from any medication and without symptoms of

the disease and toxicity. In contrast, ATG patients spent more time in health states other than TWiST and therefore had greater requirements for close medical care, transfusion support, and medication. Our results suggest that ATG patients spend more time with a reduced quality of life, and by spending more time in cost-intensive health states are likely to produce more costs due to medical care.

There is no patient-derived information available to indicate the appropriate utility factors for health states such as toxicity, transfusions, clonal disorders, GvHD, or partial remission. In our view, patients in partial remission have a fairly good quality of life, restricted by medications and frequent doctor visits; therefore, the coefficient of 0.75 was chosen for these periods, whereas for periods spent with transfusion requirements, extensive chronic GvHD, or clonal complications a coefficient of 0.5 was chosen reflecting considerable impairment of quality of life. A coefficient of 0.75 was chosen for the time spent in complete remission but still requiring regular medication, because taking drugs on a daily basis clearly interferes with quality of life. Yet different weighting of the utility coefficients might affect the results of the Q-TWiST analysis. Therefore, a threshold utility analysis was performed for the most relevant health states in terms of duration and differences between treatment groups. With this, we show that varying the utility factors for PR does not affect the results of the Q-TWiST analysis.

There are limitations to this study. Patients receiving immunosuppression were by study design significantly older. Therefore, we cannot exclude that age disparity could play a role in some of the differences observed between treatment groups. Furthermore, definitions of transition periods were arbitrary. There is heterogeneity in some of the transition periods; for instance, the time spent in partial remission includes patients who have close to normal blood counts and patients whose hemopoiesis is just sufficient to keep them free of transfusions. During the time spent in PR, more transplanted patients have nearly normal blood values, while ATG patients are more likely to present lower blood counts. In addition, the study period covers almost 3 decades. During the whole study period, changes in the protocols and progress in supportive care and treatment modalities have led to improvement of survival and event-free survival. However, there was no difference in Q-TWiST when comparing ATG-treated patients and patients treated with BMT (results not shown) by decade. Finally, Q-TwiST analysis attempts to provide information on quality of life without direct patientcentered questionnaires. Such an evaluation is easily performed in a study covering a limited time period [37]. A Q-TwiST evaluation is much more critical in studies with chronic disorders, such as aplastic anemia, where the longest observation time exceeds 20 years and the history of an individual patient is often complex [37, 38].

Even with these limitations, the Q-TWiST analysis is a methodology that allows evaluating retrospectively the time a patient spent at different health states, assuming that quality of life is dependant on each particular health state [39], and gives us a complementary view on the outcome

of the two standard treatment modalities. Despite similar survival and event-free survival probabilities, patients treated with BMT spend more time cured from their disease, while ATG-treated patients have greater requirements for close medical care, transfusion support, and medications and spend therefore more time in cost-intensive intervals. The results presented should be helpful for patient information and making decisions for patients with aplastic anemia.

References

- Bacigalupo A, Bnmo B, Saracco P, Di Bona E, Locasciulli A, Locatelli F, Gabbas A, Dufour C, Arcese W, Testi G, Broccia G, Carotenuto M, Coser P, Barbui T, Leoni P, Ferster A (2000) Antilymphocyte globulin, cyclosporine, prednisolone, and granulocyte colony-stimulating factor for severe aplastic anemia: an update of the GITMO/EBMT study on 100 patients. European Group for Blood and Marrow Transplantation (EBMT) Working Party on Severe Aplastic Anemia and the Gruppo Italiano Trapianti di Midollo Osseo (GITMO). Blood 95:1931–1934
- Doney K, Leisenring W, Storb R, Appelbaum FR (1997)
 Primary treatment of acquired aplastic anemia: outcomes with
 bone marrow transplantation and immunosuppressive therapy.
 Ann Intern Med 126:107–115
- Frickhofen N, Rosenfeld SJ (2000) Immunosuppressive treatment of aplastic anemia with antithymocyte globulin asnd clyclosporine. Semin Hematol 37:56–68
- 4. Marsh J, Schrezenmeier A, Marin P, Ilhan O, Ljungman P, McCann S, Socie G, Tichelli A, Passweg J, Hows J, Raghavachar A, Locasciulli A, Bacigalupo A (1999) Prospective randomized multicenter study comparing cyclosporin alone versus the combination of antithymocyte globulin and cyclosporin for the treatment pf patients with nonsevere aplastic anemia: a report from the European Blood and Marrow Transplant (EBMT). Blood 93:2191–2195
- Paquette RL, Tebyani N, Frane M, Ireland P, Ho WG, Champlin RE, Nimer SD (1995) Long-term outcome of aplastic anemia in adults treated with antithymocyte globulin: comparison with bone marrow transplantation. Blood 85:283– 290
- Young NS, Barrett AJ (1995) The treatment of severe acquired aplastic anemia. Blood 85:3367–3377
- Socie G, Stone JV, Wingard JR, Weisdorf D, Henslee-Downey PJ, Bredeson C, Cahn JY, Passweg JR, Rowlings PA, Schouten HC, Kolb HJ, Klein JP (1999) Long-term survival and late deaths after allogeneic bone marrow transplantation. Late Effects Working Committee of the International Bone Marrow Transplant Registry. N Engl J Med 341:14–21
- Transplant Registry. N Engl J Med 341:14–21 8. Deeg HJ, Leisenring W, Storb R, Nims J, Flowers ME, Witherspoon RP, Sanders J, Sullivan KM (1998) Long-term outcome after marrow transplantation for severe aplastic anemia. Blood 91:3637–3645
- Hows J, Marsh JC, Yin JL, Durrant S, Swirsky D, Worsley A, Fairhead SM, Chipping PM, Palmer S, Gordon Smith EC (1989) Bone marrow transplantation for severe aplastic anaemia using cyclosporin: long-term follow-up. Bone Marrow Transplant 4:11–16
- Najean Y, Haguenauer O (1990) Long-term (5 to 20 years) evolution of nongrafted aplastic anemias. The Cooperative Group for the Study of Aplastic and Refractory Anemias. Blood 76:2222–2228

- 11. Passweg JR, Socie G, Hinterberger W, Bacigalupo A, Biggs JC, Camitta BM, Champlin RE, Gale RP, Gluckman E, Gordon-Smith EC, Hows JM, Klein JP, Nugent ML, Pasquini R, Rowlings PA, Speck B, Tichelli A, Zhang MJ, Horowitz MM, Bortin MM (1997) Bone marrow transplantation for severe aplastic anemia: has outcome improved? Blood 90:858–864
- Tichelli A, Schrezenmeier H, Bacigalupo A (2000) Immunosuppressive treatment of aplastic anemia. In: Schrezenmeier H, Bacigalupo A (eds) Aplastic anemia: pathophysiology and treatment. Cambridge University Press, Cambridge, pp 154– 196
- 13. Locasciulli A, van't Veer L, Bacigalupo A, Hows J, Van Lint MT, Gluckman E, Nissen C, McCann S, Vossen J, Schrezenmeier A, et al. (1990) Treatment with marrow transplantation or immunosuppression of childhood acquired severe aplastic anemia: a report from the EBMT SAA working party. Bone Marrow Transplant 6:211–217
- Socie G, Henry-Amar M, Cosset JM, Devergie A, Girinsky T, Gluckman E.(1991) Increased incidence of solid malignant tumors after bone marrow transplantation for severe aplastic anemia. Blood 78:277–279
- 15. Storb R, Leisenring W, Anasetti C, Appelbaum FR, Buckner CD, Bensinger WI, Chauncey T, Clift RA, Deeg HJ, Doney KC, Flowers ME, Hansen JA, Martin PJ, Sanders JE, Sullivan KM, Witherspoon RP (1997) Long-term follow-up of allogeneic marrow transplants in patients with aplastic anemia conditioned by cyclophosphamide combined with antithymocyte globulin. Blood 89:3890–3891
- Schrezenmeier H, Marin P, Raghavachar A, McCann S, Hows J, Gluckman E, Nissen C, van't Veer-Korthof ET, Ljungman P, Hinterberger W et al (1993) Relapse of aplastic anaemia after immunosuppressive treatment: a report from the European Bone Marrow Transplantation Group SAA Working Party. Br J Haematol 85:371–377
- Socie G, Henry-Amar M, Bacigalupo A, Hows J, Tichelli A, Ljungman P, McCann SR, Frickhofen N, Van't Veer-Korthof E, Gluckman E (1993) Malignant tumors occurring after treatment of aplastic anemia. European Bone Marrow Transplantation-Severe Aplastic Anaemia Working Party. N Engl J Med 329:1152–1157
- Tichelli A, Gratwohl A, Nissen C, Wiirsch A, Signer E, Speck B (1988) Spätkomplikationen bei Patienten mit aplastischer Anämie. Schweiz Med Wochenschr 118:1528–1532
- Gelber RD, Goldhirsch A, Cole BF (1993) Evaluation of effectiveness: Q-TWiST. The International Breast Cancer Study Group. Cancer Treat Rev 19 [Suppl A]:73–84
- Gelber RD, Cole BF, Goldhirsch A, Gisselbrecht C, Sebban C, Morel P, Marit G, Bouabdallah R, Ravoet C, Salles G, Reyes F, Lepage E (1996) Adjuvant chemotherapy plus tamoxifen compared with tamoxifen alone for postmenopausal breast cancer: meta-analysis of quality-adjusted survival. Lancet 347:1066–1071
- 21. Mounter N, Haioun C, Cole BF, Gisselbrecht C, Sebban C, Morel P, Marit G, Bouabdallah R, Ravoet C, Salles G, Reyes F, Lepage E (2000) Quality of life-adjusted survival analysis of high-dose therapy with autologous bone marrow transplantation versus sequential chemotherapy for patients with aggressive lymphoma in first complete remission. Groupe d'Etude les Lymphomes de 1'Adulte (GELA). Blood 95:3687–3692
- 22. Parson SK, Gelber S, Cole BF, Ravindranath Y, Ogden A, Yeager AM, Chang M, Shuster J, Weinstein HJ, Gelber RD (1999) Quality-adjusted survival after treatment for acute myeloid leukemia in childhood: a Q-TWiST analysis of the Pediatire Oncology Group Study 8821. J Clin Oncol 17:2144–2152
- Zee B, Cole B, Li T, Browman G, James K, Johnston D, Sugano D, Pater J (1998) Quality-adjusted time without symptoms or toxicity analysis of interferon maintenance in multiple myeloma. J Clin Oncol 16:2834–2839

- 24. Bacigalupo A, Hows J, Gluckman E, Nissen C, Marsh J, Van Lint MT, Congiu M, De Planque MM, Ernst P, McCann S, et al. (1988) Bone marrow transplantation (BMT) versus immunosuppression for the treatment of severe aplastic anaemia: a report of the EBMT SAA Working Party. Br J Haematol 70:177–182
- 25. Camitta BM, Thomas ED, Nathan DG, Santos G, Gordon-Smith EC, Gale RP, Rappeport JM, Storb R (1976) Severe aplastic anemia: a prospective study of the effect of early marrow transplantation on acute mortality. Blood 48:63–70
- 26. Speck B, Tichelli A, Widmer E, Harder F, Kissling M, Wursch A, Stebler Gysi C, Signer E, Bargetzi M, Orth B, Gratwohl A, Nissen C (1996) Splenectomy as an adjuvant measure in the treatment of severe aplastic anaemia. Br J Haematol 92:818–824
- Speck B, Gratwohl A, Nissen C, Leibundgut U, Ruggero D, Osterwalder B, Burri HP, Cornu P, Jeannet M (1981) Treatment of severe aplastic anaemia with antilymphocyte globulin or bone marrow transplantation. Br Med J 282:860–863
- Speck B, Gratwohl A, Nissen C, Osterwalder B, Wursch A, Tichelli A, Lori A, Reusser P, Jeannet M, Signer E (1986) Treatment of severe aplastic anemia. Exp Hematol 14:126–132
- Tichelli A, Passweg JR, Nissen C, Bargetzi M, Hoffmann T, Wodnar-Filipowicz A, Signer E, Speck B, Gratwohl A (1998) Repeated treatment with horse antilymphocyte globulin for severe aplastic anaemia. Br J Haematol 100:393–400
- Tichelli A, Gratwohl A, Wiirsch A, Nissen C, Speck B (1988) Antilymphocyte globulin for myelodysplastic syndrome? Br J Haematol 68:139–140
- 31. Kaplan EL, Meier P (1958) Nonparametric estimation from incomplete observations. J Am Stat Assoc 53:457–481

- 32. Peto R, Pike MC, Armitage P, Breslow NE, Cox DR, Howard SV, Mantel N, McPherson K, Peto J, Smith PG (1977) Design and analysis of randomized clinical trials requiring prolonged observation of each patient. Br J Cancer 35:1–39
- 33. Cole BF, Gelber RD, Goldhirsch A (1995) A quality-adjusted survival meta-analysis of adjuvant chemotherapy for premenopausal breast cancer. International Breast Cancer Study Group. Stat Med 14:1771–1784
- Levy V, Porcher R, Delabarre F, Lepomer M, Cazin B, Chevret S (2001) Evaluating treatment strategies in chronic lymphocytic leukemia: use of quality-adjusted survival analysis. J Clin Epidemiol 54:747–754
- 35. Gelber RD, Gelber S (1995) Quality-of-life assessment in clinical trials. Cancer Treat Res 75:225-246
- 36. Frickhofen N, Kaltwasser JP, Schrezenmeier H, Raghavachar A, Vogt HG, Herrmann F, Freund M, Meusers P, Salama A, Heimpel H (1991) Treatment of aplastic anemia with antilymphocyte globulin and methylprednisolone with or without cyclosporine. N Engl J Med 324:1297–1304
- 37. Levy V, Porcher R, Leblond V, Fermand JP, Cazin B, Maloisel F, Harousseau JL, Remenieras L, Guibon O, Chevret S, French Cooperative Group on CLL and Macroglobulinemia (2001) Evaluating treatment strategies in advanced Waldenström macroglobulinemia: use of quality-adjusted survival analysis. Leukemia 15:1466–1470
- 38. Schwartz CE, Coulthard-Morris L, Cole B, Vollmer T (1997) The quality-of-life effects of interferon beta-lb in multiple sclerosis. An extended Q-TWiST analysis. Arch Neurol 54:1475–1480
- 39. Gelber RD, Cole BF, Goldhirsch A, Bonadonna G, Howell A, McArdle CS, Mouridsen HT, Rubens RD, Welvaart K (1995) Adjuvant chemotherapy for premenopausal breast cancer: a meta-analysis using quality-adjusted survival. Cancer J Sci Am 1:114