Busulfan and melphalan conditioning is superior to melphalan alone in autologous stem cell transplantation for high-risk MM

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Since its introduction 3 decades ago, upfront autologous hematopoietic stem cell transplantation (auto-HCT) with melphalan (Mel) 200 mg/m² remains the standard of care in the treatment of patients with transplant-eligible, newly diagnosed multiple myeloma (MM).1-4 Several drug combinations for an optimal conditioning regimen have been tried without a convincing benefit. The combination of busulfan (Bu) and Mel is synergistic against MM cell lines.5 Previous studies with the combination of oral busulfan and Mel showed longer progression-free survival (PFS), albeit with a higher rate of veno-occlusive disease.6,7 Since then, the introduction of intravenous busulfan with linear pharmacokinetics and more reproducible systemic exposure has largely ameliorated veno-occlusive disease.8 We studied this combination regimen in a phase 3 trial in newly diagnosed MM patients receiving upfront auto-HCT. Our results showed that IV Bu plus Mel (Bu-Mel) was associated with a significantly better PFS than single-agent Mel (hazard ratio, 0.53; 3-year PFS, 72% vs 50%).5

Even with the use of novel drugs in the treatment of MM, the outcome in patients harboring high-risk (HR) cytogenetic abnormalities remains suboptimal.1,3 We hypothesized that myeloma patients with HR cytogenetics may particularly benefit from the intensification of conditioning in the Bu-Mel regimen. In a subset analysis of our trial, where 62 HR patients were evaluated separately, a significantly longer PFS in the Bu-Mel arm (median PFS not reached; \( P = 0.0087 \)) vs Mel arm (median 25 months) was observed. Here, we report the updated outcomes of HR patients enrolled in that trial.

Between 2011 and 2017, 205 patients were enrolled in the trial, 105 to the Bu-Mel arm and 100 to the Mel arm. HR MM was defined based on the presence of HR cytogenetic profile per International Myeloma Working Group criteria9 and included 17p deletion, t(4;14), t(14;16), t(14;20), and 1q gain detected by fluorescence in situ hybridization and del13q detected by karyotyping. Sixty-two patients had HR MM, 32 in the Bu-Mel arm and 30 in the Mel arm. Patients in the Bu-Mel arm received a test dose of 32 mg/m² Bu on day 28 if the participant was an outpatient or on day 29 if the participant was an inpatient, followed by pharmacokinetically adjusted doses of Bu on days 27, 26, 25, and 24, with a daily target area under the curve of 5000 mmol/min, and Mel was administered on days −7, −6, −5, and −4, with a daily target area under the curve of 5000 mmol/min, and Mel was administered on days −2 and −1 with a total Mel dose 140 mg/m². Patients in the Mel arm received Mel 200 mg/m² on days 2. The primary objective was to compare PFS in HR NDMM patients who underwent upfront auto-HCT with Bu-Mel vs Mel conditioning.

Baseline patient characteristics of HR MM arms are provided in Table 1. There was no significant difference in the hematopoietic cell transplantation–specific comorbidity index score, induction therapy, response to induction, number of patients receiving maintenance therapy, or other characteristics between the 2 treatment arms. The median age at auto-HCT was 61 years (range, 31.7-70.6 years) in the Bu-Mel arm and 60 years (range, 38.8-69.5 years) in the Mel arm. Overall, 32 (100%) and 29 (97%) patients received induction therapy with immunomodulatory drugs and/or proteasome inhibitors in the Bu-Mel and Mel arms, respectively (Table 1).
The median time to neutrophil engraftment (absolute neutrophil count ≥0.5 x 10^9/L) was 11 and 12 days in the Bu-Mel and Mel arms, respectively (P = .002). The median time to platelet engraftment (platelet count ≥20 x 10^9/L without transfusion support) was 10 and 12.5 days in the Bu-Mel and Mel arms, respectively (P < .001). There was no 100-day treatment-related mortality in either arm. There was an increased incidence of grade II to IV toxicities in the Bu-Mel arm (32/32; 100%) compared with the Mel arm (25/30; 83%) (P < .001). Details of these toxicities are provided in supplemental Table 1. These toxicities were mostly reversible and self-limited and included mucositis, febrile neutropenia, and transaminase elevation. There was no grade IV toxicity in either arm. Grade II to III mucositis was seen in 24 (75%) and 5 (17%) patients in the Bu-Mel and Mel arms, respectively (P = .049). The 3-year OS rates were 90% (95% CI, 50% to 82%) and 41% (95% CI, 22% to 59%) in the Bu-Mel and Mel arms, respectively (P = .005).

The median duration of follow-up from randomization was 42.6 months (range, 18.9-85.7 months) and 36.6 months (range, 3.4-74.9 months) in the Bu-Mel and Mel arms, respectively. The overall response rate, defined as partial response or better, at day 90 after auto-HCT was 100% (32/32) and 93% (28/30) of patients in the Bu-Mel and the Mel arm, respectively (P = .25). The survival outcomes are provided in supplemental Table 2. The 3-year PFS rates were 69% (95% confidence interval [CI], 50% to 82%) and 41% (95% CI, 22% to 59%) in the Bu-Mel and Mel arms, respectively. Median PFS was 44.7 (95% CI, 31.9 to not reached) months and 25.7 (95% CI, 16.5 to not reached) months in the Bu-Mel and Mel arms, respectively (P = .044; Figure 1A) (hazard ratio [95% CI], 0.48 [0.24-1.00]; P = .049). The 3-year overall survival rates were 90% (95% CI, 73% to 97%) and 87% (95% CI, 65% to 96%) in the Bu-Mel and Mel arms, respectively. The median overall survival was not reached.

### Table 1. Summary of patient and clinical characteristics

| Variables                      | Bu-Mel (n = 32) | Mel (n = 30) | P   |
|--------------------------------|----------------|-------------|-----|
| Age at auto-HCT, y             | 61.4 (31.7-70.6)| 59.8 (38.8-69.5) | .28 |
| Sex, n (%)                     |                |             |     |
| Male                           | 16 (50)        | 18 (60)     | .46 |
| Female                         | 16 (50)        | 12 (40)     |     |
| R-ISS, n (%)                   | 3 (9)          | 2 (7)       |     |
| I                              | 6 (22)         | 8 (38)      | .39 |
| II                             | 17 (63)        | 9 (43)      |     |
| III                            | 4 (15)         | 4 (19)      |     |
| Missing                        | 5              | 9           |     |
| HR features (IMWG), n (%)      |                |             |     |
| Individual                     |                |             |     |
| del(17p)                       | 11 (34)        | 8 (27)      |     |
| Gain 1q                        | 18 (56)        | 17 (57)     |     |
| Copy number 3                  | 14             | 8           |     |
| Copy number ≥4                 | 1              | 3           |     |
| Not available                  | 3              | 6           |     |
| t(4;14)                        | 6 (19)         | 4 (13)      |     |
| del(13q) (karyotype)           | 6 (19)         | 8 (27)      |     |
| Combined                       |               |             | .61 |
| del(17p)/Gain 1q (copy number ≥ 4)/(t(4;14)) | 17 (55) | 14 (47) |     |
| Other                          | 14 (45)        | 16 (53)     |     |
| Missing                        | 1              | 0           |     |
| Serum LDH                      |                |             |     |
| Normal                         | 19 (63)        | 19 (63)     | 1.00|
| Abnormal                       | 4 (17)         | 3 (14)      |     |
| Missing                        | 9              | 8           |     |
| HCT-CI, n (%)                  |                |             |     |
| 0                              | 11 (34)        | 14 (47)     | .23 |
| 1-2                            | 8 (25)         | 10 (33)     |     |
| ≥3                             | 13 (41)        | 6 (20)      |     |
| Response to induction, n (%)   |                |             |     |
| sCR/CR                         | 6 (19)         | 4 (13)      | .55 |
| nCR                            | 3 (9)          | 1 (3)       |     |
| VGPR                           | 9 (28)         | 11 (37)     |     |
| PR                             | 14 (44)        | 12 (40)     |     |
| SD                             | 0              | 2 (7)       |     |
| Maintenance regimen, n (%)     |                |             |     |
| Lenalidomide                   | 14 (44)        | 8 (27)      | .69 |
| Lenalidomide and elotuzumab    | 4 (13)         | 5 (17)      |     |
| Lenalidomide and ixazomib      | 1 (3)          | 3 (10)      |     |
| Lenalidomide and dexamethasone | 2 (6)          | 1 (3)       |     |
| Bortezomib                     | 3 (9)          | 7 (23)      |     |
| IRD                            | 2 (6)          | 1 (3)       |     |
| KRD                            | 1 (3)          | 2 (7)       |     |
| VRD                            | 2 (6)          | 1 (3)       |     |
| None                           | 3 (9)          | 2 (7)       |     |

### Table 1. (continued)

| Variables | Bu-Mel (n = 32) | Mel (n = 30) | P   |
|-----------|----------------|-------------|-----|
| Induction regimen, n (%) |                |             |     |
| VRD       | 19 (59)        | 17 (59)     | .99 |
| VCD       | 6 (19)         | 5 (17)      |     |
| KRD       | 4 (13)         | 3 (10)      |     |
| Vd        | 3 (9)          | 3 (10)      |     |
| CBAD      | 0              | 1 (3)       |     |
| Missing   | 0              | 1           |     |

CBAD, cyclophosphamide, bortezomib, adriamycin, and dexamethasone; CR, complete response; HCT-CI, hematopoietic cell transplantation–specific comorbidity index; IMWG, International Myeloma Working Group; IRD, irazomib, lenalidomide, and dexamethasone; KRD, carfilzomib, lenalidomide, and dexamethasone; LDH, lactate dehydrogenase; nCR, near CR; PR, partial response; sCR, stringent complete response; SD, stable disease; VCD, bortezomib, cyclophosphamide, and dexamethasone; Vd, bortezomib and dexamethasone; VGPR, very good partial response; VRD, .
in either arm ($P = .50$; Figure 1B) (hazard ratio [95% CI], 0.67 [0.20-2.20]; $P = .51$). In a fitted Bayesian piecewise exponential regression model, treatment with Bu-Mel was associated with superior PFS where the posterior probability of a beneficial effect was 0.90 (Bu-Mel arm vs Mel arm: mean hazard ratio, 0.62 [95% CI, 0.18-1.21]). Additional details are provided in supplemental Table 3.

Two patients (7%) in the Mel arm received consolidation therapy after auto-HCT compared with 5 patients (16%) in the Bu-Mel arm ($P = .62$). Fifty-seven patients received maintenance therapy during the trial (29 in the Bu-Mel arm [median duration, 14.0 months (range, 1.8-53.4 months)] and 28 in the Mel arm [median duration, 9.8 months (range, 0.3-38.8 months)]; $P = .19$). Of these patients, 12 (41%) in the Bu-Mel arm discontinued maintenance therapy because of disease progression or toxicity after a median duration of 10.3 months (range, 2.2-36.0 months) of maintenance compared with 16 (57%) patients in the Mel arm after a median duration of 9.4 months (range, 2.2-23.8 months) of maintenance ($P = .96$). More patients had early disease progression in the Mel arm than in the Bu-Mel arm, resulting in early discontinuation of maintenance therapy, which partly explains the shorter median duration of maintenance in the Mel arm. Furthermore, the median follow-up in the Mel arm was shorter compared with the Bu-Mel arm, which might also have contributed to a shorter duration of maintenance in the Mel arm.

This randomized trial showed an improvement in PFS with the Bu-Mel conditioning regimen in patients with newly diagnosed myeloma with HR cytogenetics. In addition, we showed that the combination of Bu-Mel led to a 52% reduction in the risk of disease progression or death compared with Mel alone. These results compare favorably with recent prospective randomized trials conducted in the era of novel therapeutics, which include BMTCTN 0702 and EMN02/HO95.5 The 3-year PFS for HR patients with single auto-HCT was 40% and ~45% in the BMTCTN0702 and EMN02/HO95 trials, respectively. Although it is always challenging to compare the results of different clinical trials, the PFS reported in these 2 trials was similar to the PFS seen in the Mel arm of our trial but significantly lower than what we reported in the Bu-Mel arm.

Interestingly, in the EMN02/HO95 trial, the 3-year PFS of 63% with tandem transplant in HR MM was comparable to what we reported in the Bu-Mel arm of our study. These data highlight that a risk-adapted approach using a higher intensity conditioning regimen, such as Bu-Mel, has the potential to significantly improve the outcomes in HR MM patients.11 Although the combination of Bu-Mel had caused more adverse events compared with Mel alone, most of the toxicities were reversible and self-limiting. We acknowledge that our trial is a single-center study, and we expect these results to be validated in a multicenter trial. A similar phase 3 randomized trial comparing IV Bu-Mel to Mel alone conditioning regimen has been completed (NCT019162520) by PETHEMA foundation group, and results are currently being awaited. In conclusion, our updated data continue to show that the Bu-Mel conditioning regimen leads to significantly better PFS when compared with Mel alone in HR MM patients.

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**Figure 1. Survival outcomes.** (A) PFS by treatment group. (B) Overall survival.
associated patents and an equity interest, though this technology does not bear on the current study. The remaining authors declare no competing financial interests.

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