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Superior outcomes associated with complete response in newly diagnosed multiple myeloma patients treated with nonintensive therapy: analysis of the phase 3 VISTA study of bortezomib plus melphalan-prednisone versus melphalan-prednisone

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Abstract

The phase 3 Velcade as Initial Standard Therapy in Multiple Myeloma: Assessment with Melphalan and Prednisone study in newly diagnosed multiple myeloma patients ineligible for high-dose therapy demonstrated that bortezomib-melphalan-prednisone (VMP) was superior to melphalan-prednisone across all efficacy end points. We assessed the prognostic impact of response on time-to-event parameters in the intent-to-treat population. Patients received nine 6-week cycles of treatment. Time to progression, time to next therapy, and treatment-free interval were associated with quality of response. When European Group for Blood and Marrow Transplantation criteria were used, complete response (CR) was associated with significantly longer time to progression (hazard ratio [HR] = 0.45, $P = .004$), time to next therapy (HR = 0.46, $P = .0004$), and treatment-free interval (HR = 0.38, $P < .0001$) versus partial response, but there was no significant difference in overall survival (HR = 0.87, $P = .54$); similar differences were seen with CR versus very good partial response by uniform criteria. Quality of response improved with prolonged VMP treatment, with 28% of CRs achieved during cycles 5-9. CR duration appeared similar among patients with “early” (cycles 1-4) and “late” CRs (cycles 5-9) and among patients receiving 9 versus < 9 cycles of bortezomib within VMP. These results highlight that CR is an important treatment goal and support prolonged VMP therapy to achieve maximal response. This study is registered at <http://www.clinicaltrials.gov> as NCT00111319.

Introduction

Multiple myeloma (MM) is an incurable plasma-cell neoplasm. The primary aim of current treatment strategy is to improve long-term outcomes, including progression-free survival (PFS) and overall survival (OS),¹ as well as health-related quality of life (HRQoL), which is affected by treatment effect, toxicity, and change in disease status.² The introduction of high-dose therapy and stem-cell transplant (HDT-SCT), which is now the standard of care for eligible patients ≤ 65 years of age, and the novel agents bortezomib, thalidomide, and lenalidomide has been associated with improved OS in the past 10 years.³⁻⁵

The authors of studies in previously untreated MM patients who received HDT-SCT have demonstrated that quality of response, notably achievement of complete response (CR) or at least very good partial response (VGPR), is prognostic for improved long-term outcomes.⁶⁻¹³ However, in patients ineligible for HDT-SCT, there is limited evidence for such a correlation between quality of response and long-term outcomes.¹⁴ This may be because until recently, standard frontline therapies for patients ineligible for HDT-SCT did not result in substantial rates of CR or CR/VGPR, thereby preventing meaningful analysis of outcomes according to

response. Now, novel agent-based treatment options for previously untreated nontransplantation MM patients are demonstrating high CR and CR/VGPR rates, superior to those achieved with previous standards of care.¹⁵ The use of bortezomib in combination with conventional therapies has resulted in increased CR rates associated with improved outcomes.¹⁵

The phase 3 Velcade as Initial Standard Therapy in Multiple Myeloma: Assessment with Melphalan and Prednisone (VISTA) study compared bortezomib plus melphalan-prednisone (VMP) versus melphalan-prednisone alone in previously untreated patients with MM who were ineligible for high-dose therapy.¹⁶ This study demonstrated that VMP was superior to MP across all response rate and time-to-event efficacy end points and resulted in a very high CR rate of 30%.^{16,17} Thus, we investigated whether, as seen in the transplant setting, achievement of CR was associated with improved outcome. Here, we assess the differential prognostic impact of best response on time-to-event parameters in VISTA in the substantial population of patients achieving CR and partial response (PR) and evaluate time-to-event distributions in the VMP arm by quality of response. In addition, we evaluate the impact of timing of CR on outcome in patients receiving VMP.

Methods

Study design

The VISTA study design has been described previously (<http://www.clinicaltrials.gov>, NCT00111319).¹⁶ In brief, previously untreated MM patients were randomized to receive 9 6-week cycles of VMP, comprising bortezomib 1.3 mg/m², days 1, 4, 8, 11, 22, 25, 29, and 32, cycles 1-4, and days 1, 8, 22, and 29, cycles 5-9; melphalan 9 mg/m², days 1-4, cycles 1-9; and prednisone 60 mg/m², days 1-4, cycles 1-9, or MP alone. The primary end point was time to progression (TTP). Response was determined by a prespecified computer algorithm and progression was determined by investigator assessment by use of the European Group for Blood and Marrow Transplantation (EBMT) criteria¹⁸; a central laboratory was used for M-protein assessment. In a post hoc analysis, response was determined by use of the International Myeloma Working Group (IMWG) uniform criteria.¹⁹ As previously reported,¹⁶ the VISTA study was stopped after the third interim analysis on the basis of an independent recommendation from the data monitoring committee (data cutoff, June 15, 2007). Centralized collection of M-protein data was discontinued; therefore, subsequent data on response rates and TTP were not updated; however, data collection continued for subsequent therapy use and OS.

The VISTA study was approved by the Independent ethics committee/institutional review board at all participating institutions and was conducted in accordance with International Conference on Harmonization Guidelines on Good Clinical Practice and the principles of the Declaration of Helsinki. All patients provided written informed consent.

Analysis of outcomes by response

Patients who had measurable disease at baseline and who received at least one dose of study drug were evaluable for response. For the purposes of the present analysis, patients were classified by either the major response achieved (CR or PR), or as those with minimal or no response (< PR). By EBMT criteria, 102 (30%), 136 (40%), and 95 (28%) patients in the VMP arm and 12 (4%), 103 (31%), and 212 (64%) patients in the MP arm achieved CR, PR, and < PR, respectively. By IMWG criteria, 111 (33%), 28 (8%), 112 (33%), and 82 (24%) patients in the VMP arm and 13 (4%), 13 (4%), 102 (31%), and 199 (60%) in the MP arm achieved

CR, VGPR, PR, and < PR, respectively.¹⁶ It was not possible to include the additional response category of near-CR²⁰ in the present analysis because of the small number of patients achieving this response.

TTP, time to next therapy (TNT), treatment-free interval (TFI), and OS were evaluated in the VMP arm according to response by EBMT and IMWG criteria. Long-term outcomes (TTP, TNT, TFI, OS, and time from response/CR to subsequent therapy) also were evaluated in VMP patients achieving early CR, during treatment cycles 1-4, or late CR, during treatment cycles 5-9. Response and TTP data were from the third interim analysis (median follow-up was 16.3 months).¹⁶ TNT, TFI, and OS data were from the most recent updated analysis, with a data cutoff of March 16, 2009, and median follow-up was 36.7 months. For the same reasons as for response and TTP data, response and CR duration could not be updated from the third interim analysis; therefore, time from response/CR to subsequent therapy was analyzed as a surrogate for response/CR duration by the use of subsequent therapy data from the updated analysis.

Statistical analysis

Response duration was defined as the time from first evidence of response to start of subsequent therapy, and CR duration was the time from first evidence of CR to start of subsequent therapy, as surrogates for time to relapse, as described previously. TTP was defined as time from randomization to progression, TNT was time from randomization to next therapy, and TFI was length of time after therapy without treatment. Associations between TTP, TNT, TFI, and OS and response were examined in the intent-to-treat population, which included both VMP and MP patients, by multivariate Cox regression analysis with time-dependent covariates, adjusted for stratification factors. Covariates included age, sex, race, MM type, baseline Karnofsky performance status, and number of bone lesions; *P* values for comparisons of outcomes by response were determined by stratified log-rank tests. TNT, TTP, OS, TFI, and time from response/CR to subsequent therapy distributions according to best response achieved were estimated for the response-evaluable population in the VMP arm by Kaplan-Meier methodology.

Results

Patients

A total of 682 patients were randomized to receive VMP (*n* = 344) or MP (*n* = 338). The response-evaluable population included 337 and 331 patients in the VMP and MP arms, respectively. Patient demographics and baseline disease characteristics for the overall population have been reported previously¹⁶ and were typical of a transplant-ineligible patient population. Patient demographics and baseline characteristics in the response-evaluable population in the VMP arm according to response by EBMT criteria are listed in Table 1. Characteristics appeared generally similar between response subgroups; however, the proportion of patients aged ≥ 75 years appeared greater in the < PR subgroup (42%) compared with the CR (27%) and PR (26%) subgroups. The proportion of patients who had a baseline β_2 -microglobulin level of > 5.5 mg/L appeared greater in those achieving CR versus PR or < PR (42% vs 26% or 32%). A greater proportion of patients achieving CR (42%) had ISS stage III disease, compared with patients achieving PR (29%) or < PR (33%). Among patients with ISS disease stage I who achieved CR, most were achieved early. In addition, the ratio of MM subtypes varied between response subgroups, with a lower proportion of immunoglobulin G MM and a greater proportion of immunoglobulin A MM patients within the CR subgroup compared with the PR or < PR subgroups.

Table 1

Table 1. Patient demographics and baseline characteristics in the VMP arm within subgroups defined by response and by timing of CR

Variable	Response subgroup				
	Early CR	Late CR	All CR	PR	< PR
No.	73	29	102	136	99
Male, %	48	34	44	50	60
White, %	81	86	82	91	90
Age, y					
Median	71	70	71	71	74
≥ 75, %	25	34	27	26	42*
KPS, %					
≤ 70	40	38	39	32	36
80	32	28	30	29	26
≥ 90	29	34	30	40	37
β₂-microglobulin, mg/L					
Median	4.40	5.10	4.65	4.00	4.10
< 2.5, %	10	7	9	13	13
2.5-5.5, %	51	45	49	61	55
> 5.5, %	40	48	42†	26	32
Albumin, g/L					
Median	34.0	32.0	33.0	33.5	33.0
< 3.5, %	53	76	60	57	58
ISS staging, %					
I	22	3	17	19	19
II	38	48	41	51	47
III	40	48	42‡	29	33
Region, %					
North America	15	7	13	9	7
Europe	70	76	72	82	82
Other	15	17	16	9	11
Type of MM, %§					
IgG	34	76	46	71	74
IgA	45	17	37	24	13
Light chain	16	3	13	3	9
Creatinine clearance, mL/min					
Median	61.7	56.7	60.2	57.9	57.6
≤ 30, %	7	7	7	5	5
31-60, %	41	48	43	50	48
> 60, %	52	45	50	45	46
High-risk cytogenetics [t(4;14), t(14;16), del(17p)], % of patients with cytogenetic information¶	15	17	15	16	13

Response is categorized by EBMT criteria (response-evaluable population, N = 337). Early CR occurred within 24 weeks of treatment; and late CR, after 24 weeks.

CR indicates complete response; EBMT, European Group for Blood and Marrow Transplantation; IgA, immunoglobulin A; IgG, immunoglobulin G; KPS, Karnofsky performance status; MM, multiple myeloma; PR, partial response; and VMP, bortezomib-melphalan-prednisone.

*P = .0067, Fisher exact test, for difference in rate of age ≥ 75 years between patients achieving CR/PR vs < PR.

†P = .0229, Fisher exact test, for difference in rate of β₂-microglobulin > 5.5 mg/L between patients achieving CR vs PR/< PR.

‡P = .0610, Fisher exact test, for difference in rate of ISS stage III disease between patients achieving CR vs PR/< PR.

§P = .0088, Cochran-Mantel-Haenszel test, for difference in MM type between patients achieving CR/PR versus < PR; P < .0001 for CR versus PR/< PR.

¶There were only 25 response-evaluable patients with high-risk cytogenetics, of whom 7 achieved CR (5 early, 2 late), 12 achieved PR, and 6 had < PR.

Analyses of outcomes by response

Results of multivariate Cox regression analysis of outcomes according to response by EBMT criteria are shown for the intent-to-treat population (all patients in both the VMP and MP arms) in Table 2. CR was associated with significantly longer TTP (hazard ratio [HR] = 0.45, $P = .004$), TNT (HR = 0.48, $P < .0001$), and TFI (HR = 0.42, $P < .0001$) versus PR. By contrast there was no significant difference between CR and PR in terms of OS (HR = 0.87, $P = .54$). Furthermore, CR and PR were associated with significantly longer TTP, TNT, TFI, and OS versus < PR. Similar associations between response and outcomes were seen in patient subgroups who had ISS stage I/II or ISS stage III MM (supplemental Tables 1 and 2, available on the *Blood* Web site; see the Supplemental Materials link at the top of the online article).

Table 2

Table 2. Multivariate analysis of outcomes according to best response in the intent-to-treat population, in VMP and MP patients, by EBMT criteria

	Hazard ratio (95% CI)	
	Versus PR	Versus < PR
TTP		
Patients achieving CR	0.45 (0.26-0.78), <i>P</i> = .004	0.31 (0.18-0.56), <i>P</i> = .0001
Patients achieving PR	—	0.69 (0.51, 0.95), <i>P</i> = .021
TNT		
Patients achieving CR	0.48 (0.34-0.68), <i>P</i> = .0001	0.33 (0.22-0.47), <i>P</i> < .0001
Patients achieving PR	—	0.68 (0.54-0.85), <i>P</i> = .0008
TFI		
Patients achieving CR	0.42 (0.30-0.60), <i>P</i> = .0001	0.27 (0.18-0.39), <i>P</i> < .0001
Patients achieving PR	—	0.63 (0.51-0.79), <i>P</i> < .0001
OS		
Patients achieving CR	0.87 (0.56-1.36), <i>P</i> = .54	0.52 (0.33-0.82), <i>P</i> = .005
Patients achieving PR	—	0.60 (0.44-0.81), <i>P</i> = .0009

CI indicates confidence interval; CR, complete response; EBMT, European Group for Blood and Marrow Transplantation; OS, overall survival; PR, partial response; TFI, treatment-free interval; TNT, time to next therapy; TTP, time to progression; VMP, bortezomib-melphalan-prednisone; MP, melphalan-prednisone; and —, not applicable.

Kaplan-Meier distributions of TTP, TNT, TFI, and OS in VMP patients achieving CR or PR by EBMT criteria reflect the findings of the multivariate analysis (Figure 1). Patients achieving CR had prolonged TTP, TFI, and TNT versus those achieving PR; OS appeared similar among patients achieving CR or PR to VMP although, as noted previously, by multivariate analysis, the HR was 0.87 in favor of CR.

Figure 1

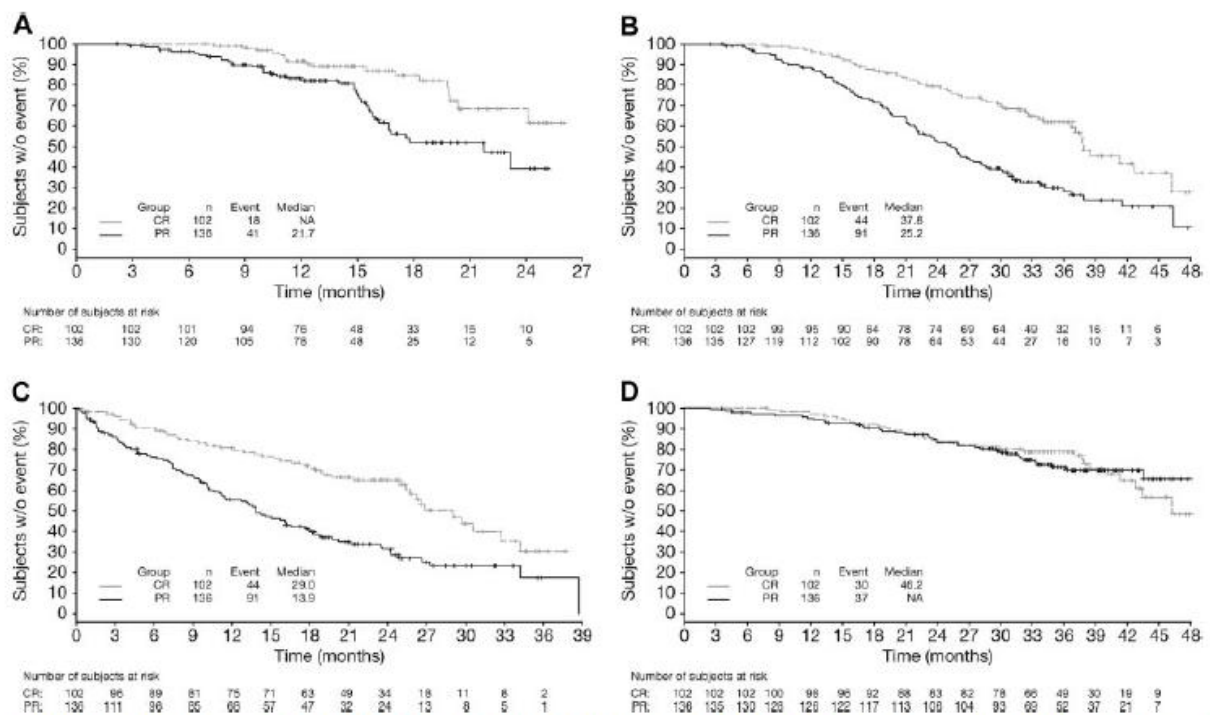


Figure 1. Time-to-event analyses in patients achieving CR versus PR by EBMT criteria with VMP. Kaplan-Meier analyses of (A) TTP, (B) TNT, (C) TFI, and (D) OS in patients achieving CR versus PR with VMP.

Multivariate Cox regression analysis of outcomes according to response by IMWG criteria (Table 3) showed that CR was associated with significantly longer TTP (HR = 0.45, $P = .019$), TNT (HR = 0.52, $P = .004$), and TFI (HR = 0.45, $P = .0008$) versus VGPR, whereas no significant differences were seen in outcomes between patients achieving VGPR and PR (Table 3). It should be noted that the number of patients achieving VGPR was small ($n = 41$). Analysis of outcomes in patients achieving CR versus VGPR + PR versus < PR by IMWG criteria demonstrated similar findings to the analysis of CR versus PR versus < PR by EBMT criteria (data not shown). In addition, CR + VGPR was associated with significantly longer TTP (HR = 0.60, $P = .025$), TNT (HR = 0.65, $P = .005$), and TFI (HR = 0.63, $P = .002$) compared with PR, but there was no significant difference in OS (HR = 0.88, $P = .54$). Kaplan-Meier distributions of TTP in patients achieving CR, CR + VGPR, VGPR, or PR by IMWG criteria with VMP demonstrated longer TTP (median not reached vs not reached vs 16.8 months vs 21.7 months) in patients who achieved CR or CR + VGPR (Figure 2).

Table 3

Table 3. Multivariate analyses of outcomes according to best response in the intent-to-treat population, by IMWG criteria

	Hazard ratios (95% CI)		
	Versus VGPR	Versus PR	Versus < PR
TTP			
Patients achieving CR	0.45 (0.24-0.88), $P = .019$	0.50 (0.29-0.87), $P = .013$	0.29 (0.17-0.52), $P < .0001$
Patients achieving VGPR	—	1.11 (0.63-1.96), $P = .73$	0.64 (0.36-1.15), $P = .14$
Patients achieving PR	—	—	0.58 (0.42-0.81), $P = .001$
Patients achieving CR + VGPR	—	0.60 (0.38-0.94), $P = .025$	0.37 (0.23-0.59), $P < .0001$
TNT			
Patients achieving CR	0.52 (0.34-0.81), $P = .004$	0.58 (0.41-0.81), $P = .001$	0.40 (0.28-0.57), $P = .0001$
Patients achieving VGPR	—	1.10 (0.74-1.65), $P = .64$	0.76 (0.50-1.16), $P = .20$
Patients achieving PR	—	—	0.69 (0.55-0.88), $P = .002$
Patients achieving CR + VGPR	—	0.65 (0.49-0.88), $P = .005$	0.47 (0.34-0.64), $P < .0001$
TFI			
Patients achieving CR	0.45 (0.28-0.72), $P = .0008$	0.50 (0.36-0.71), $P = .0001$	0.30 (0.21-0.43), $P < .0001$
Patients achieving VGPR	—	1.12 (0.73-1.72), $P = .61$	0.67 (0.44-1.04), $P = .072$
Patients achieving PR	—	—	0.60 (0.48-0.76), $P < .0001$
Patients achieving CR + VGPR	—	0.63 (0.46-0.85), $P = .002$	0.38 (0.28-0.52), $P < .0001$
OS			
Patients achieving CR	0.96 (0.53-1.75), $P = .90$	0.89 (0.58-1.36), $P = .58$	0.46 (0.31-0.67), $P = .0001$
Patients achieving VGPR	—	0.92 (0.52-1.62), $P = .78$	0.47 (0.27-0.82), $P = .008$
Patients achieving PR	—	—	0.52 (0.38-0.70), $P < .0001$
Patients achieving CR + VGPR	—	0.88 (0.59-1.32), $P = .54$	0.51 (0.34-0.75), $P = .0007$

CI indicates confidence interval; CR, complete response; IMWG, International Myeloma Working Group; OS, overall survival; PR, partial response; TFI, treatment-free interval; TNT, time to next therapy; TTP, time to progression; VGPR, very good partial response; and —, not applicable.

Figure 2

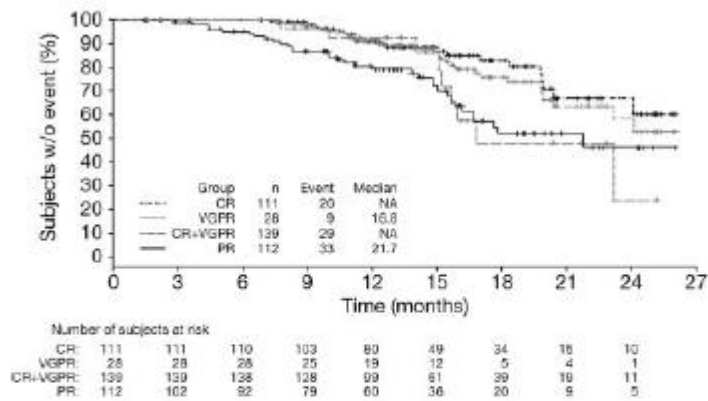


Figure 2. Time to progression in patients achieving CR versus VGPR versus PR by IMWG criteria with VMP. Kaplan-Meier analysis of time to progression according to response to VMP by IMWG criteria.

Impact of timing of CR on outcome

The median time to first response with VMP treatment was 1.4 months. Most first responses occurred early, but late de novo responses were seen, including 4% (9/238) that occurred beyond the first 24 weeks of therapy (ie, during cycles 5-9; Figure 3A). Quality of response improved with prolonged VMP treatment. Although most of patients' best responses occurred within the first 24 weeks of therapy (cycles 1-4), a substantial proportion (16%, 37/238) occurred later (after 24 weeks), including 28% (29/102) of CRs (Figure 3B).

Figure 3

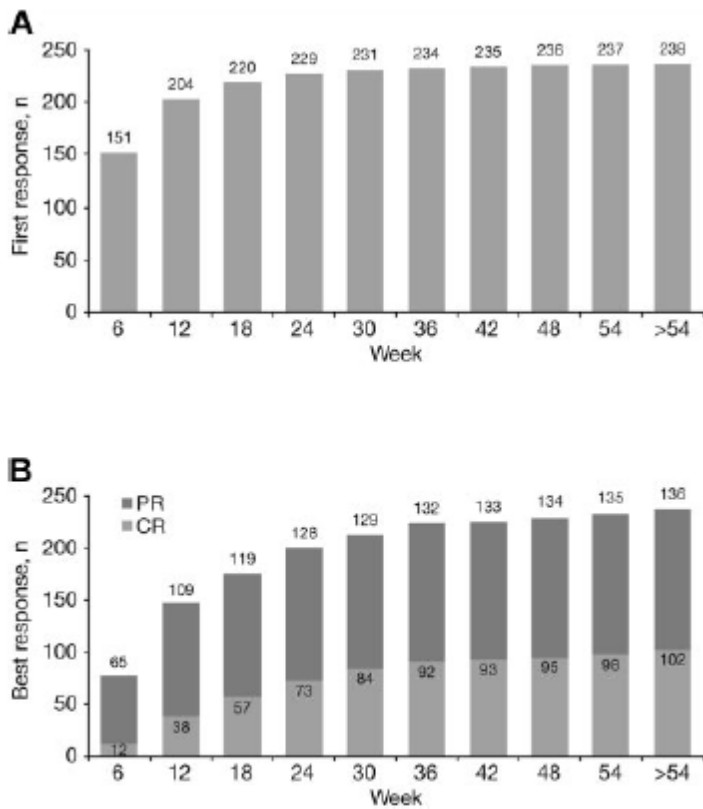


Figure 3. Achievement of first response and best response to VMP over time. Cumulative numbers of patients achieving their first (A) and best (B) responses to VMP by week of therapy.

Among patients achieving CR with VMP by EBMT criteria, CR duration (time from CR to subsequent therapy) appeared similar in patients achieving CR early (cycles 1-4, within 24 weeks) versus late (cycles 5-9, after 24 weeks, Figure 4A). TNT (Figure 4B; median 41.2 vs 37.8 months), response duration (Figure 4C; median 44.7 vs 36.7 months), and OS (Figure 4D) appeared somewhat longer in the patients who achieved CR late versus early. Because they were measured from the time of randomization, TNT and OS appeared somewhat prolonged with late versus early CRs, with similar CR duration. Mean duration of therapy was 8.4 versus 7.6 cycles of study treatment in the patients achieving CR late versus early.

Figure 4

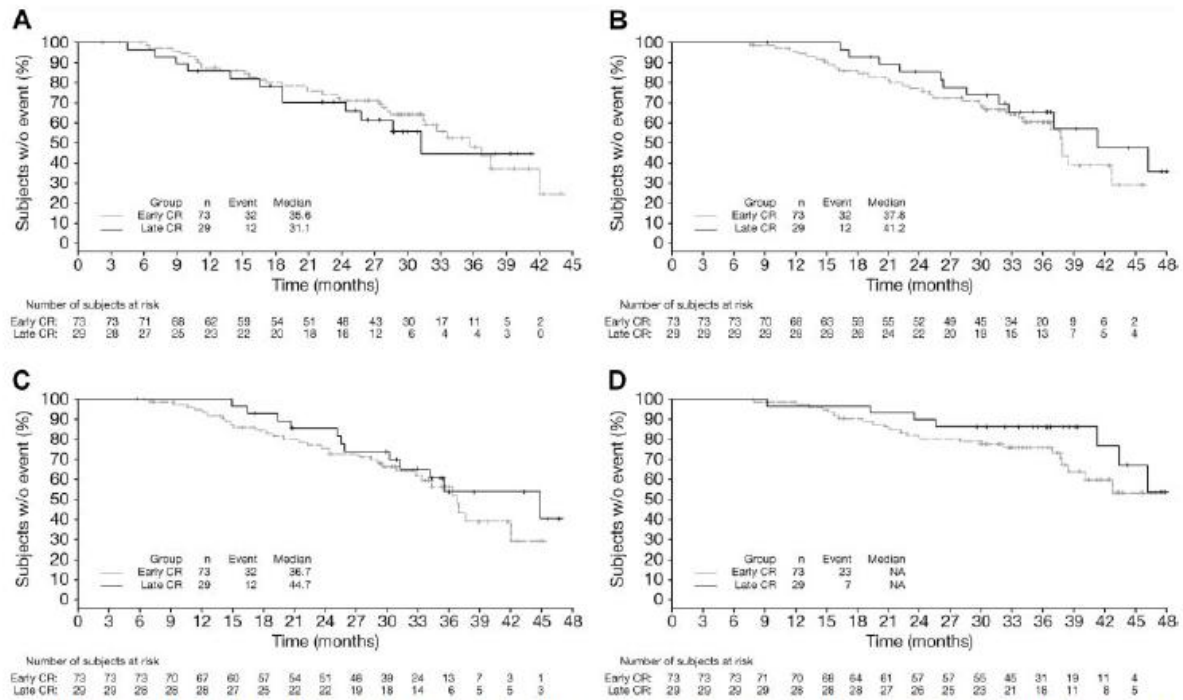


Figure 4. Time-to-event analyses in VMP patients achieving early or late CR. Kaplan-Meier analyses of time from CR to subsequent therapy (A), TNT (B), time from first response to subsequent therapy (C), and OS (D) in VMP patients achieving CR early (within 24 weeks) or late (after 24 weeks).

CR duration (time from CR to subsequent therapy) was analyzed among patients who received < 9 or all 9 protocol-specified cycles of bortezomib as part of VMP therapy, excluding those who stopped treatment early due to progression, and appeared similar in both groups (Figure 5A). This was also the case in an analysis restricted to those patients who achieved CR early (Figure 5B). Analyses of outcomes restricted to patients receiving all 9 cycles of bortezomib treatment demonstrated that CR duration (Figure 5C) appeared similar with late versus early achievement of CR, whereas TTP (data not shown), TNT (Figure 5D; median 46.2 vs 37.8 months), and response duration (data not shown) appeared somewhat longer in patients with late versus early CR, which was consistent with prior analyses (Figure 4). OS (data not shown) was similar between patients achieving late versus early CR in this subgroup receiving all 9 cycles of bortezomib.

Figure 5

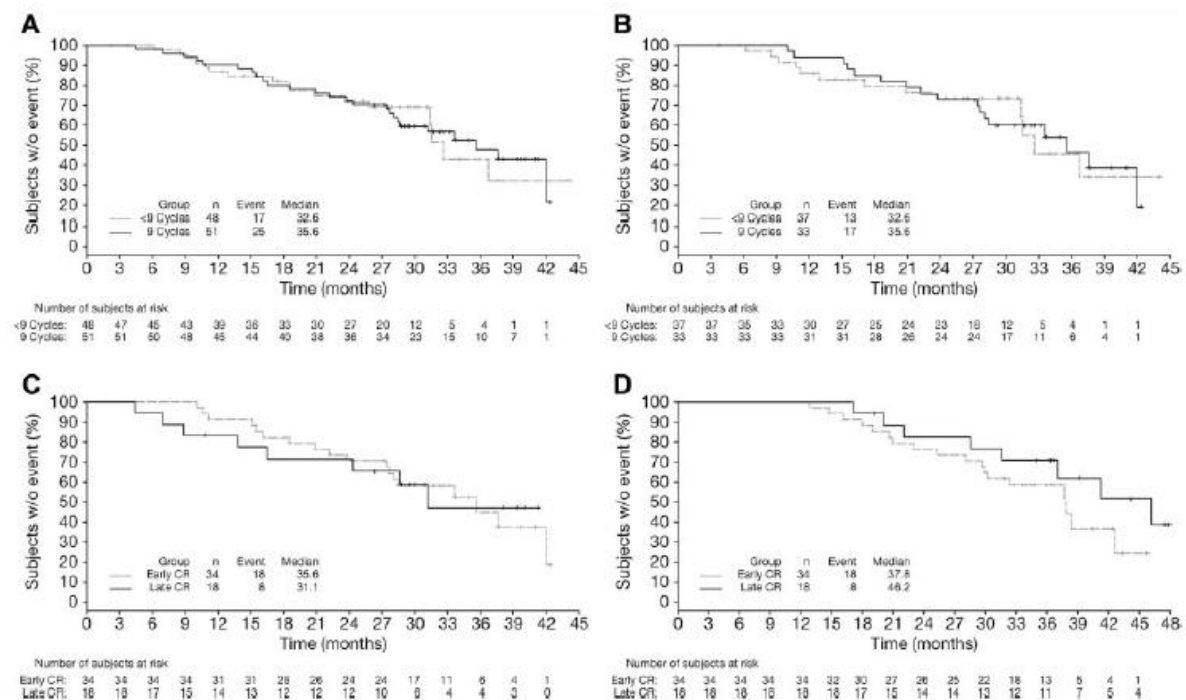


Figure 5. Time-to-event analyses according to duration of bortezomib therapy in patients achieving CR. Kaplan-Meier analyses of CR duration (time from CR to subsequent therapy) in (A) all patients achieving CR and (B) patients achieving early CR who received < 9 versus 9 cycles of bortezomib, excluding those who discontinued early because of PD, plus analyses of CR duration (C) and TNT (D) in patients achieving early versus late CR who completed all 9 cycles of bortezomib.

Discussion

This analysis of the VISTA phase 3 trial demonstrated the differential prognostic significance of quality of response on outcomes in the nontransplant setting. The large VISTA population and high CR rate with VMP has provided a large sample size to reliably explore this prognostic significance. In a previous analysis, San Miguel et al¹⁶ showed that OS is superior with VMP vs MP alone; here, we show achievement of CR by EBMT response criteria was associated with significantly longer TTP, TNT, and TFI versus PR. However, there was no statistically significant difference in OS with CR versus PR. With the increasingly widespread use of highly active novel agent-based regimens in the relapsed setting, analyses of OS according to parameters or response in the front-line setting are becoming problematic due to the confounding effect of achieving a high-quality response to salvage therapy.²¹ In the VISTA population overall, 411 of 682 patients (60%) had received subsequent therapy by data cutoff of March 2009, including 52% and 69% of patients in the VMP and MP arms, respectively.¹⁷ Among patients on the VMP arm who had received subsequent therapy, 76% had received bortezomib-, thalidomide-, or lenalidomide-based therapy.¹⁷ Thus, in patients with inherently sensitive MM (that is, those who responded to VMP), additional responses to subsequent lines of therapy, particularly novel agent-based therapy, may have resulted in the similar OS observed between patients achieving CR or PR to VMP, despite the other significant time-to-event and clinical benefits of CR versus PR. In addition, improved outcomes were associated with > PR versus < PR, with patients achieving significantly longer TTP, TNT, TFI, and OS. These analyses, by both EBMT and IMWG criteria, suggest that quality of response is associated with improved long-term outcomes with VMP treatment. In addition, in a separate analysis of HRQoL data from VISTA, CR achievement and duration were associated with clinically relevant improvements in patient HRQoL.²

These findings support the results of previous studies investigating the prognostic impact of CR versus PR on long-term outcomes in previously untreated MM patients, who were ineligible for, or did not undergo, HDT-SCT. In the ECOG E9486 trial, in which patients had active MM and received conventional chemotherapy, OS was 5.1 and 3.3 years in patients experiencing CR and PR, respectively ($P < .0001$).¹⁴ Similarly, in a retrospective analysis assessing clinical outcomes in 758 patients, Wang et al²² demonstrated longer OS after a 2-year landmark in patients who had achieved CR versus PR versus no response (12.4 years vs 3.7 years vs 2.1 years, respectively) at 2 years after receiving dexamethasone-based therapy.

Our analyses according to response by IMWG criteria showed improved outcomes with CR versus VGPR and with CR + VGPR versus PR. However, outcomes appeared similar with VGPR versus PR, indicating that there may not be a significant prognostic difference between VGPR and PR. This may be attributable to small numbers of patients achieving VGPRs ($n = 41$; 28 VMP, 13 MP) and these findings should be interpreted with caution. However, the superiority of CR over VGPR/PR is consistent with an analysis of outcomes and clinical benefit by quality of response in the bortezomib arm of the phase 3 APEX (Assessment of Proteasome Inhibition for Extending Remissions) trial, which demonstrated that CR was associated with significantly longer median TFI (24.1 vs 6.9/6.4 months) and time to alternative therapy (27.1 vs 13.6/14.0 months) versus VGPR/PR.²³

Two phase 2 studies of thalidomide-based regimens support our findings that CR + VGPR is superior to PR. Patients treated with pegylated liposomal doxorubicin, vincristine, dexamethasone, and thalidomide who achieved CR + VGPR demonstrated improved PFS ($P = .01$) and OS ($P = .03$) versus patients achieving PR/no response.²⁴ Similarly, patients treated with thalidomide, dexamethasone, and pegylated liposomal doxorubicin who achieved CR + VGPR had a significantly higher 3-year TTP (78% vs 40%, $P = .031$), event-free survival (78% vs 37%, $P = .021$), and OS (84% vs 61%, $P = .053$) rates versus patients achieving $< VGPR$.²⁵

Our analyses have demonstrated that the clinical benefit of CR with VMP appears similar, regardless of whether CR is achieved early or late. A substantial proportion of patients achieved CR late in their treatment course (after 24 weeks); these so-called “late CRs” may include patients improving from an initial response of PR. Thus, our data support continuation of VMP therapy beyond a lesser response in patients tolerating treatment, with the aim of achieving a CR. A lower proportion of early versus late CRs were achieved in elderly patients, and patients with β_2 -microglobulin > 5.5 mg/L and low albumin (< 3.5 g/dL), suggesting that these patients take longer to achieve CR. In contrast, patients with ISS stage I had a greater proportion of early versus late CRs compared with ISS stage II and III patients, which suggests a more rapid complete response in patients with low tumor burden. In patients receiving all 9 cycles of bortezomib with VMP therapy, long-term outcomes appeared either similar or slightly improved with late versus early CR. This may be because of the inherent bias of the additional time taken to achieve a CR from a lesser response. These results support findings from the APEX trial, in which response duration was not affected by rapid achievement of first response in patients receiving bortezomib.²⁶

The use of techniques such as serum free light chain assay, and immunohistochemistry or immunofluorescence have led to a more stringent definition of CR (sCR by IMWG criteria) than immunofixation.¹⁹ Furthermore, multiparametric flow cytometry²⁷ and polymerase chain reaction²⁸ have allowed an even more stringent definition of CR through evaluation of minimal residual disease. Indeed, the greater depth of response defined by these more sensitive techniques has been shown to be prognostic for improved outcomes.^{27–36} Thus, a CR defined by these more sensitive techniques may be the ultimate treatment goal, rather than achievement of an immunofixation-negative CR.

In conclusion, these results demonstrate that CR is prognostic for improved long-term outcomes compared with lesser responses and so achievement of CR, by the most stringent definition possible, should be considered an important treatment goal. Thus, our data also support continuation of VMP therapy in patients tolerating treatment to achieve maximal response and maximal clinical benefit.

References

1. Rajkumar SV, Kyle RA . *Multiple myeloma: diagnosis and treatment. Mayo Clin Proc 2005;80(10):1371-1382.*
2. Dhawan R, Meunier J, Regnault A, et al . *Impact of complete response on quality of life in newly diagnosed multiple myeloma patients [abstract]. Clin Lymphoma Myeloma 2009;9 suppl 1:S58. Abstract 352.*
3. Brenner H, Gondas A, Pulte D . *Recent major improvement in long-term survival of younger patients with multiple myeloma. Blood 2008;111(5):2521-2526.*
4. Kastiris E, Zervas K, Symeonidis A, et al . *Improved survival of patients with multiple myeloma after the introduction of novel agents and the applicability of the International Staging System (ISS): an analysis of the Greek Myeloma Study Group (GMSG). Leukemia 2009;23(6):1152-1157.*
5. Kumar SK, Rajkumar SV, Dispenzieri A, et al . *Improved survival in multiple myeloma and the impact of novel therapies. Blood 2008;111(5):2516-2520.*
6. Attal M, Harousseau JL, Stoppa AM, et al . *A prospective, randomized trial of autologous bone marrow transplantation and chemotherapy in multiple myeloma. Intergroupe Francais du Myelome. N Engl J Med 1996;335(2):91-97.*
7. Attal M, Harousseau JL, Facon T, et al . *Single versus double autologous stem-cell transplantation for multiple myeloma. N Engl J Med 2003;349(26):2495-2502.*
8. Cavo M, Tosi P, Zamagni E, et al . *Prospective, randomized study of single compared with double autologous stem-cell transplantation for multiple myeloma: Bologna 96 clinical study. J Clin Oncol 2007;25(17):2434-2441.*
9. Child JA, Morgan GJ, Davies FE, et al . *High-dose chemotherapy with hematopoietic stem-cell rescue for multiple myeloma. N Engl J Med 2003;348(19):1875-1883.*
10. Harousseau JL, Attal M, Avet-Loiseau H . *The role of complete response in multiple myeloma. Blood 2009;114(15):3139-3146.*
11. Harousseau JL, Avet-Loiseau H, Attal M, et al . *Achievement of at least very good partial response is a simple and robust prognostic factor in patients with multiple myeloma treated with high-dose therapy: long-term analysis of the IFM 99-02 and 99-04 trials. J Clin Oncol 2009;27(34):5720-5726.*

12. Lahuerta JJ, Mateos MV, Martinez-Lopez J, et al . *Influence of pre- and post-transplantation responses on outcome of patients with multiple myeloma: sequential improvement of response and achievement of complete response are associated with longer survival. J Clin Oncol 2008;26(35):5775-5782.*
13. van de Velde H, Liu X, Chen G, et al . *Complete response correlates with long-term survival and progression-free survival in high-dose therapy in multiple myeloma. Haematologica 2007;92(10):1399-1406.*
14. Kyle RA, Leong T, Li S, et al . *Complete response in multiple myeloma: clinical trial E9486, an Eastern Cooperative Oncology Group study not involving stem cell transplantation. Cancer 2006;106(9):1958-1966.*
15. Palumbo A, Rajkumar SV . *Treatment of newly diagnosed myeloma. Leukemia 2009;23(3):449-456.*
16. San Miguel JF, Schlag R, Khuageva NK, et al . *Bortezomib plus melphalan and prednisone for initial treatment of multiple myeloma. N Engl J Med 2008;359(9):906-917.*
17. Mateos MV, Richardson PG, Schlag R, et al . *Bortezomib plus melphalan and prednisone compared with melphalan and prednisone in previously untreated multiple myeloma: updated follow-up and impact of subsequent therapy in the phase III VISTA trial. J Clin Oncol 2010;28(13):2259-2266.*
18. Bladé J, Samson D, Reece D, et al . *Criteria for evaluating disease response and progression in patients with multiple myeloma treated by high-dose therapy and haemopoietic stem cell transplantation. Myeloma Subcommittee of the EBMT. European Group for Blood and Marrow Transplant. Br J Haematol 1998;102(5):1115-1123.*
19. Durie BG, Harousseau JL, Miguel JS, et al . *International uniform response criteria for multiple myeloma. Leukemia 2006;20(9):1467-1473.*
20. Richardson PG, Barlogie B, Berenson J, et al . *A phase 2 study of bortezomib in relapsed, refractory myeloma. N Engl J Med 2003;348(26):2609-2617.*
21. Anderson KC, Kyle RA, Rajkumar SV, et al . *Clinically relevant end points and new drug approvals for myeloma. Leukemia 2008;22(2):231-239.*
22. Wang M, Delasalle K, Feng L, et al . *CR represents an early index of potential long survival in multiple myeloma. Bone Marrow Transplant 2010;45(3):498-504.*
23. Niesvizky R, Richardson PG, Rajkumar SV, et al . *The relationship between quality of response and clinical benefit for patients treated on the bortezomib arm of the international, randomised, phase 3 APEX trial in relapsed multiple myeloma. Br J Haematol 2008;143(1):46-53.*
24. Hussein MA, Baz R, Srkalovic G, et al . *Phase 2 study of pegylated liposomal doxorubicin, vincristine, decreased-frequency dexamethasone, and thalidomide in newly diagnosed and relapsed-refractory multiple myeloma. Mayo Clin Proc 2006;81(7):889-895.*

25. Offidani M, Corvatta L, Piersantelli MN, et al . *Thalidomide, dexamethasone and pegylated liposomal doxorubicin (ThaDD) for newly diagnosed multiple myeloma patients over 65 years. Blood 2006;108(7):2159-2164.*
26. Richardson PG, Sonneveld P, Schuster M, et al . *Extended follow-up of a phase 3 trial in relapsed multiple myeloma: final time-to-event results of the APEX trial. Blood 2007;110(10):3557-3560.*
27. Paiva B, Vidriales MB, Cervero J, et al . *Multiparameter flow cytometric remission is the most relevant prognostic factor for multiple myeloma patients who undergo autologous stem cell transplantation. Blood 2008;112(10):4017-4023.*
28. Ladetto M, Donovan JW, Harig S, et al . *Real-time polymerase chain reaction of immunoglobulin rearrangements for quantitative evaluation of minimal residual disease in multiple myeloma. Biol Blood Marrow Transplant 2000;6(3):241-253.*
29. Corradini P, Voena C, Tarella C, et al . *Molecular and clinical remissions in multiple myeloma: role of autologous and allogeneic transplantation of hematopoietic cells. J Clin Oncol 1999;17(1):208-215.*
30. Corradini P, Cavo M, Lokhorst H, et al . *Molecular remission after myeloablative allogeneic stem cell transplantation predicts a better relapse-free survival in patients with multiple myeloma. Blood 2003;102(5):1927-1929.*
31. Fenk R, Korthals M, Sehnke N, et al . *Molecular level of minimal residual disease in bone marrow before high-dose therapy and autologous stem cell transplantation is a prognostic parameter in patients with multiple myeloma [abstract]. Blood 2007;110(11):448a. Abstract 1496.*
32. Martinelli G, Terragna C, Zamagni E, et al . *Polymerase chain reaction-based detection of minimal residual disease in multiple myeloma patients receiving allogeneic stem cell transplantation. Haematologica 2000;85(9):930-934.*
33. Martínez-Sánchez P, Montejano L, Sarasquete ME, et al . *Evaluation of minimal residual disease in multiple myeloma patients by fluorescent-polymerase chain reaction: the prognostic impact of achieving molecular response. Br J Haematol 2008;142(5):766-774.*
34. Rawstron AC, Davies FE, DasGupta R, et al . *Flow cytometric disease monitoring in multiple myeloma: the relationship between normal and neoplastic plasma cells predicts outcome after transplantation. Blood 2002;100(9):3095-3100.*
35. San Miguel JF, Almeida J, Mateo G, et al . *Immunophenotypic evaluation of the plasma cell compartment in multiple myeloma: a tool for comparing the efficacy of different treatment strategies and predicting outcome. Blood 2002;99(5):1853-1856.*
36. Sarasquete ME, Garcia-Sanz R, Gonzalez D, et al . *Minimal residual disease monitoring in multiple myeloma: a comparison between allelic-specific oligonucleotide real-time quantitative polymerase chain reaction and flow cytometry. Haematologica 2005;90(10):1365-1372.*

37. Palumbo A, Schlag R, Khuageva NK, et al . *Prolonged therapy with bortezomib plus melphalan-prednisone (VMP) results in improved quality and duration of response in the phase III VISTA study in previously untreated multiple myeloma (MM) [abstract]. Haematologica 2008;93(s1):83. Abstract 0207.*
38. Harousseau JL, Palumbo A, Richardson PG, et al . *Superior outcomes associated with complete response: analysis of the phase III VISTA study of bortezomib plus melphalan-prednisone versus melphalan-prednisone [abstract]. Blood 2008;112:959. Abstract 2778.*
39. Harousseau JL, Palumbo A, Niesvizky R, et al . *Prolonged therapy improves quality of response, and CR is associated with superior outcomes, with VMP in the phase 3 VISTA study [abstract]. Clin Lymphoma Myeloma 2009;9 suppl 1:S40. Abstract 235.*