

Treatment of relapsed refractory multiple myeloma: which new PI-based combination treatments do patients prefer?

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Background and objectives: This study describes preferences of German relapsed refractory multiple myeloma (RRMM) patients with novel proteasome inhibitor-based combination treatments.

Methods: Patients with a minimum age of 18 years and a diagnosis of RRMM were included. Their preferences were assessed using a discrete choice experiment design, which was developed based on a literature review and two patient focus group discussions. The final discrete choice experiment design consisted of four attributes, namely “therapy application regimen,” “time without progression of disease,” “possibility of grade ≥ 3 adverse events (AEs) affecting the blood,” and “possibility of grade ≥ 3 AE heart failure.”

Results: Analysis was based on 84 patients (36.9% females, mean age 62.7 years, mean multiple myeloma disease duration 5.5 years). Among the tested attributes, “therapy application regimen” was assigned the highest importance for treatment decisions (38.8%), the second important attribute was “time without progression of disease” (38.7%), followed by “possibility of AE heart failure” (13.9%) and “possibility of AEs affecting the blood” (8.6%). Patients preferred oral intake once a day and once a week over other application modes such as oral intake once a day and once a week plus twice-weekly infusions. Furthermore, they preferred longer disease progression-free time and lower risk of grade ≥ 3 AEs. The highest overall utility was derived for ixazomib + lenalidomide + dexamethasone (utility: 3.218), compared with lenalidomide + dexamethasone (2.769), and carfilzomib + lenalidomide + dexamethasone (1.928).

Conclusion: RRMM patients prefer treatments with an all-oral application, a longer disease-progression-free time, and a lower probability of AEs. If patients face tradeoffs, they accept a lower progression-free time and/or higher AE rates to get an all-oral therapy.

Keywords: MM patient’s treatment preferences, relapsed refractory multiple myeloma, RRMM, discrete choice experiment, DCE

Introduction

Immunomodulatory agents and proteasome inhibitors (PIs) such as bortezomib^{1,2} have led to an improvement of overall survival (OS) and health-related quality of life for relapsed refractory multiple myeloma (RRMM) patients and are a cornerstone of modern RRMM therapy. Treatments available or currently under development belonging to a new generation of PIs are, for example, carfilzomib, marizomib, oprozomib, or ixazomib,^{3–5} whereas the latter represents the first PI that is administered orally.⁵

Overall, when considering a therapy in a shared decision-making framework, treatments should be tailored as much as possible to the needs of the patients. Low acceptance of treatments by patients might be linked to a low treatment adherence

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and/or early treatment discontinuation due to patients' decisions and, finally, poorer OS.⁶ Therefore, it is important to collect reliable and valid data on patient preferences when providing therapies.

None of the previous patient preference studies in multiple myeloma (MM) addressed RRMM patients in particular and/or triple combination therapies.⁷⁻¹¹ However, RRMM patient preferences may differ considerably compared with newly diagnosed MM patient preferences, due to different treatment options available such as double/triple combination regimens, a higher number of treatment lines patients already experienced, longer disease duration, potential disappointments about previous treatment failure(s), or older age. Consequently, the objective of this study was to assess RRMM patient preferences with regard to selected new generation PI-based combination treatment options.

Methods

RRMM treatment options may be characterized by various characteristics (attributes) such as efficacy, safety, or treatment burden, which often require trade-off decisions of both physicians and patients. A discrete choice experiment (DCE) is a stated preference elicitation method that requires the respondents to evaluate such trade-offs when deciding about different treatment options, each consisting of numerous attributes, as it needs to be done in reality.¹² The basic assumption of a DCE is that rational individuals will always choose alternatives with higher levels of expected utility. This study, which is the first study using a DCE framework to assess RRMM patients' treatment preferences, followed a two-stage design, according to recommendations how to conduct DCE studies in health care.¹² In stage 1, a targeted review was conducted to generate a first list of potential attributes and respective attribute levels describing different PI-based RRMM combination treatment options. Here, summary of product characteristics, clinical trial data, as well as previously performed preference studies in MM were analyzed. With regard to adverse event (AE)-related attributes, only grade ≥ 3 AEs were taken into account as these are typically reducing the quality of life of patients most severely and are reported more consistently across different trials.

The whole list of attributes was presented to patients and discussed with them in two qualitative focus group discussions (FGDs). After a general discussion about potential additional attributes to be included, importance of each attribute was assessed by means of a dual questioning technique.¹³

Based on the final list of attributes, a factorial DCE design was generated (hypothetical choice sets). An additional choice set was added in which one previous choice set was doubled, in order to assess the consistency of a patient's response behavior. Finally, all choice sets were graphically visualized (choice cards) to facilitate the comprehensibility of the different attributes and attribute levels.

Patients were recruited in two different ways: 1) via German patient support groups (PSGs) and 2) via physician practices within Germany. The inclusion criteria for survey participants were defined as follows:

- Diagnosis of RRMM (please note that in case of patients included via PSGs, this information was obtained from the patient).
- Minimum age of 18 years.
- Ability and willingness to take part in an interview.

To collect DCE data, study patients were interviewed in stage 2 by trained interviewers, either face-to-face or via telephone. All phone interview patients received a postal mailing with the prepared choice cards and a brief description of the next steps.

Data analysis was only conducted for patients with consistent DCE data, based on the responses to the above-described test choice set.¹⁴ DCE data were analyzed using conditional logit regression models. The influence of different attribute levels on the probability of a patient's decision to choose a specific alternative as well as respective utility levels was estimated, considering interaction effects between attributes. Calculated utility levels were also applied to selected novel PI-based RRMM combination treatments. Analyses were done with Stata/MP 14 and Microsoft Excel 2016.

The study protocol was approved by the independent Ethics Committee of the Medical Faculty at the University of Rostock. The study was conducted in accordance with the Declaration of Helsinki. Participants provided written informed consent prior to their enrollment in the study.

Results

FGDs and final DCE design

In our targeted PubMed-based literature review, we identified 42 publications (Figure S1 for search strings and PRISMA chart); additionally, product characteristics of addressed agents were reviewed. Based on this review, 13 different attributes were discussed with eight RRMM patients in the FGDs (two FGDs with four patients each): progression-free survival; possible side effects: occurrence of thrombocytopenia, neutropenia, anemia, hypokalemia, cardiac failure, hypertension, ischemic heart disease, fatigue, diarrhea, renal

failure, rash; application of therapy. The dual questioning-ranking of these attributes resulted in the following seven top-ranked attributes (in that order): anemia as AE, time without progression of disease, neutropenia as AE, thrombocytopenia as AE, application mode of the treatment including number and duration of necessary physician visits, hypokalemia, and heart failure as AEs. The ranking showed that the four most relevant safety attributes were all connected to blood-related side effects. To reduce the number of attributes (and in this respect to optimize the DCE design), it was decided to merge these attributes to one overall attribute, defined as “possibility of AEs affecting the blood.”

Consequently, four attributes were used to generate the experimental design. To confront patients with realistic, although hypothetical choices, treatment-associated attribute levels were derived from either the SmPCs (application mode) or data reported in the respective clinical trials addressing new generation PIs (progression-free time and the MIN and MAX grade ≥ 3 AE frequencies).^{1,3,5} In the final DCE design, the following attributes/attribute levels were included: “therapy regimen/drug administration”: 1) oral intake once daily plus once per week with 2-hour physician visit once per month; 2) oral intake once daily plus twice per week with 2-hour physician visit once per month; 3) oral intake once daily plus once per week plus 3- to 4-hour physician visits twice per week including administration of one infusion per visit; “time without progression of disease”: 1) 17 months, 2) 20 months, 3) 26 months; “possibility of grade ≥ 3 AEs affecting the blood”: 1) 12 of 100 patients, 2) 19 of 100 patients; “possibility of grade ≥ 3 AEs in regard to heart failure”: 1) two of 100 patients, 2) four of 100 patients. The final DCE design consisted of ten choice sets. In each of the choice sets, the patient needed to decide between two different hypothetical treatment options (Figure 1: one example choice set).

Quantitative DCE survey

One hundred thirteen RRMM patients could be included in the study. Thirteen patients withdrew their informed consent after study inclusion or an interview appointment could not be made. Of the remaining 100 patients, 16 (16.0%) did not answer consistently and were thus excluded from all further analyses. Consequently, final analyses were carried out for 84 patients. Mean age of patients in the final analysis sample was 62.8 years, 36.9% were female, mean/median disease duration was 5.5/4.2 years, and the majority of patients had either an Eastern Cooperative Oncology Group (ECOG) performance status of 1 (45.2%) or 2 (35.7%) (Table 1); 83.3% of the included patients already received a stem cell

transplantation, and patients included via study sites, in comparison with those included via PSGs, were generally older and more female (Table 1).

Preference analysis was based on 756 different hypothetical treatment choices (nine decisions for each of the 84 patients). The attribute “therapy regimen/drug administration” influenced patients’ decisions with an extent of 38.8%. “Time without progression of disease” was almost equally important with an influence of 38.7%, whereas the attributes “possibility of side effect heart failure” with 13.9% influence and “possibility of side effects affecting the blood” with 8.6% influence were less important (Figure 2). Preferences observed in the total study population were widely consistent across subgroups according to age, gender, disease duration, and traveling distance to treating physician (Table S1).

Generally, utilities differed significantly between all tested attribute levels within the four attribute categories (Figure 2). So, for example, a treatment associated with 26-month progression-free time instead of 17 months was associated with an additional utility of 1.785, whereas oral intake once daily plus once weekly instead of oral intake once daily/once weekly plus an additional twice-weekly physician visit and infusion was associated with an additional utility of 1.794. Lower frequencies of grade ≥ 3 AEs were associated with positive utilities as well, although these were considerably lower, compared with the previously described attributes (Figure 2).

Table 2 shows the utilities as derived for the studied new generation PI-based combination regimens; utilities were derived from the sum of the attribute-level-specific utilities applicable for the respective treatment options: 1) carfilzomib plus lenalidomide plus dexamethasone (CRd), 2) ixazomib plus lenalidomide plus dexamethasone (IRd), and 3) lenalidomide plus dexamethasone (Rd). Total utilities associated with these treatments were calculated based on the median progression-free survival time as reported in the respective trials (17.6, 20.6, or 26.3 months, respectively), the respective application mode (1, 2, or 3), and the respective risk to experience severe AEs with respect to heart failure (2% or 3.8%) and blood-related side effects (ranging from 1.1% to 29.6%), as described in the respective clinical trials.^{3,5} Because the attribute “possibility of side effects affecting the blood” was a composite attribute (including thrombocytopenia, neutropenia, anemia, and hypokalemia), three scenarios were calculated: minimum AE risk, maximum risk, and mean AE risk across the four different grade 3+ AEs belonging to this category. When considering the mean values in this respect, highest utility for patients arose from

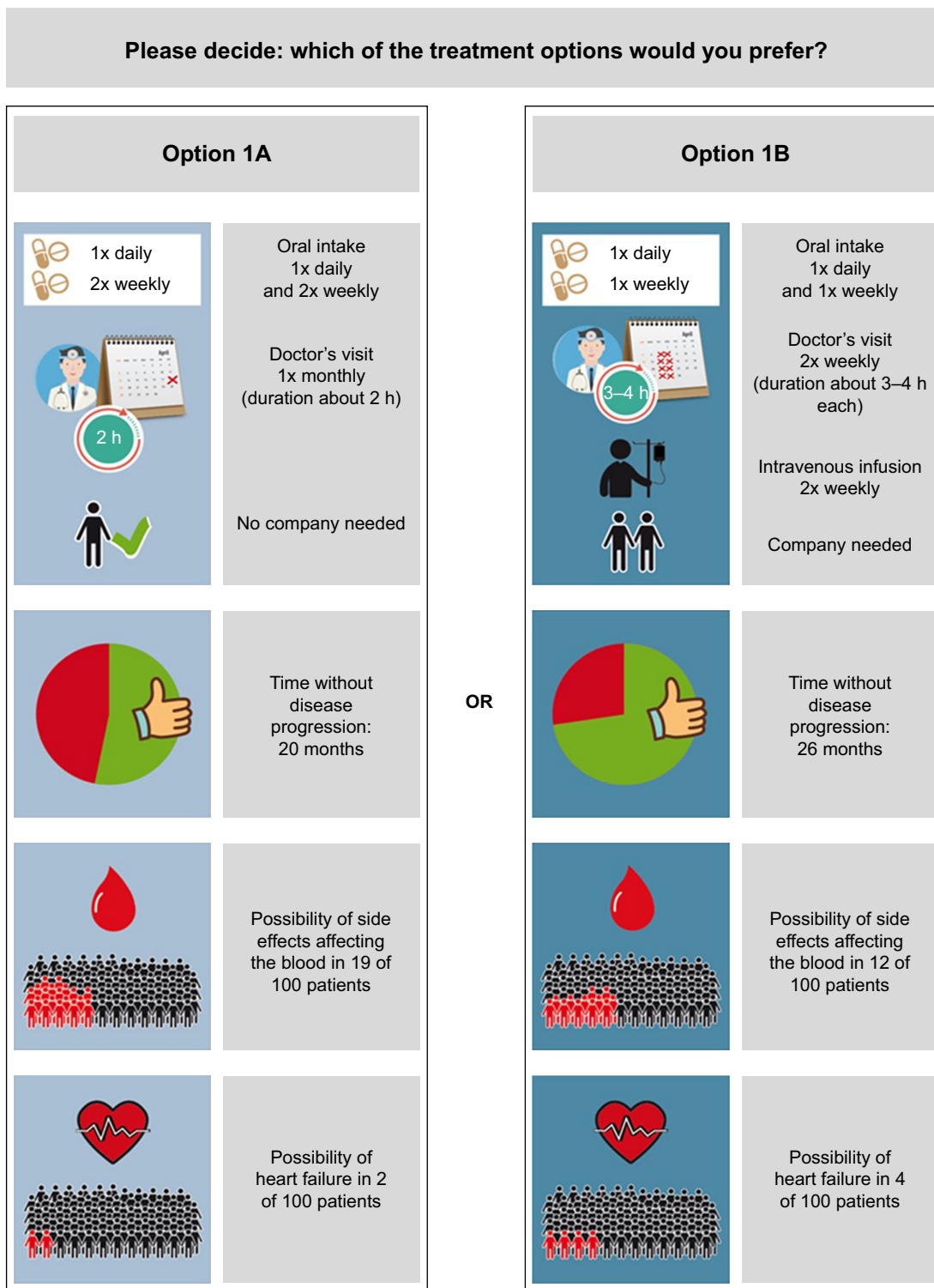


Figure 1 Example of a visualized choice card.

the treatment option IRd (3.218), followed by Rd (2.769) and CRd (1.928).

In a post hoc analysis, the length of progression-free time in months was calculated that patients would be willing to

“trade” to receive a certain attribute combination as treatment. Results show that patients would be willing to give up the highest number of progression-free months for treatment options incorporating all-oral application and/or lower probability

Table 1 Patient characteristics

Characteristics	Patients included in the DCE analyses	Patients recruited via PSGs (included for analyses)	Patients recruited via study sites (included for analyses)	Patients with inconsistent DCE answers (not included in analyses)
N	84	67	17	16
Female gender (%)	31 (36.9)	24 (35.8)	7 (41.2)	8 (50.0)
Age in years, mean (median/SD)	62.8 (63.5/9.7)	61.4 (62.0/9.5)	68.1 (68.0/8.6)	67.8 (65.0/10.3)
Disease duration in years, mean (median/SD)	5.5 (4.2/4.3)	5.6 (4.3/4.5)	5.2 (3.7/3.8)	6.2 (4.9/4.8)
Living situation, n (%)				
Patients living alone	11 (13.1)	8 (11.9)	3 (17.7)	5 (31.3)
Patients living together with a spouse/partner	70 (83.3)	56 (83.6)	14 (82.4)	10 (62.5)
Patients living together with children/others	3 (3.6)	3 (4.5)	0 (0.0)	1 (6.3)
Educational status, n (%)				
Untrained	1 (1.2)	0 (0.0)	1 (5.9)	1 (6.3)
Apprenticeship	29 (34.5)	21 (31.3)	8 (47.1)	11 (68.8)
University	26 (31.0)	26 (38.8)	0 (0.0)	1 (6.3)
Others	28 (33.3)	20 (23.8)	8 (47.1)	3 (18.8)
Employment status, n (%)				
Full time	7 (8.3)	7 (10.4)	0 (0)	0 (0.00)
Part time	8 (9.5)	7 (10.4)	1 (5.9)	1 (12.5)
Unable to work	10 (11.9)	9 (13.4)	1 (5.9)	2 (6.3)
Retired	50 (59.5)	36 (53.7)	14 (82.4)	12 (81.3)
Others	9 (10.7)	8 (11.9)	1 (5.9)	0 (0.00)
Ways in which patients get to their RRMM-treating physician, n (%)				
Public transportation	9 (10.7)	9 (13.4)	0 (0.00)	3 (25.0)
Car	45 (53.6)	40 (59.7)	5 (29.4)	6 (37.5)
Patient is taken by friends/family	10 (11.0)	8 (11.9)	2 (11.8)	4 (18.8)
Others	20 (23.8)	10 (14.9)	10 (58.8)	3 (18.8)
Distance to physician who treats RRMM in km, mean (median/SD)	37.2 (15.0/61.7)	43.2 (15.0/67.6)	13.6 (13.0/10.4)	27.9 (19.0/26.1)
Travel time to physician who treats RRMM in minutes, mean (median/SD)	39.4 (30.0/40.4)	44.1 (30.0/43.8)	20.9 (20.0/10.5)	33.8 (30.0/17.2)
Patients who received a treatment for RRMM at the time of interview, n (%)	55 (65.5)	42 (62.7)	13 (76.5)	8 (50.0)
Patients who already have experience with intravenous medication, n (%)	83 (98.8)	66 (98.1)	17 (100.0)	14 (87.0)
Patients experienced the intravenous application as ..., n (%)				
... very stressful	7 (8.3)	5 (7.5)	2 (11.8)	0 (0.0)
... stressful	21 (25.0)	19 (28.4)	2 (11.8)	5 (28.6)
... less stressful	28 (33.3)	22 (32.8)	6 (35.3)	7 (57.1)
... hardly stressful	27 (32.1)	20 (29.9)	7 (41.2)	2 (14.3)
Patient-reported average duration of an intravenous medication application in minutes, mean (median/SD)	101.1 (90.0/64.0)	105.8 (90.0/67.8)	85.3 (90.0/46.2)	96.8 (90.0/51.1)
Patients who received a stem cell transplantation, n (%)	70 (83.3)	58 (86.6)	12 (70.6)	10 (62.5)

(Continued)

Table 1 (Continued)

Characteristics	Patients included in the DCE analyses	Patients recruited via PSGs (included for analyses)	Patients recruited via study centers (included for analyses)	Patients with inconsistent DCE answers (not included in analyses)
ECOG-performance scale at the time of interview, n (%)				
Grade 0	11 (13.1)	9 (13.4)	2 (11.8)	3 (18.8)
Grade 1	38 (45.2)	26 (38.8)	12 (70.6)	6 (37.5)
Grade 2	30 (35.7)	27 (40.3)	3 (17.7)	5 (31.3)
Grade 3	5 (6.0)	5 (7.5)	0 (0.0)	2 (12.5)
Grade 4	0 (0.0)	0 (0.0)	0 (0.0)	0 (0.0)

Notes: Patient characteristics at the time of study inclusion are described. Characteristics of patients included in the final data analysis (n=84) with the respective subgroups of patients included via PSGs or study sites are shown. Moreover, characteristics of patients excluded from the final analysis set due to inconsistent DCE response behavior (n=16) are shown.

Abbreviations: DCE, discrete choice experiment; ECOG, Eastern Cooperative Oncology Group; PSGs, patient support groups; RRMM, relapsed refractory multiple myeloma.

of side effects. As an example, in order to receive an all-oral instead of an oral plus twice-weekly infusion therapy regimen combined with lower AE risk risks in regard to blood-related events and heart failure, RRMM patients would be willing to trade up to 14.4 months of progression-free time (Table S2).

Discussion

Our study shows that, when deciding on treatment options among hypothetical novel PI-based combination treatments that are characterized by the tested four attributes, patients

ranked the attribute “therapy regimen/drug administration” as the most important, closely followed by “time without progression of disease.” With regard to the therapy regimen, patients preferred application regimens based on an all-oral intake of drugs instead of lengthy infusions in physician’s practices, so that they take into account the effects of a therapy on their quality of life.

One of the main strengths of a DCE analysis is that different characteristics of specific treatments (here: attribute levels) can be transferred into utilities and, finally, overall

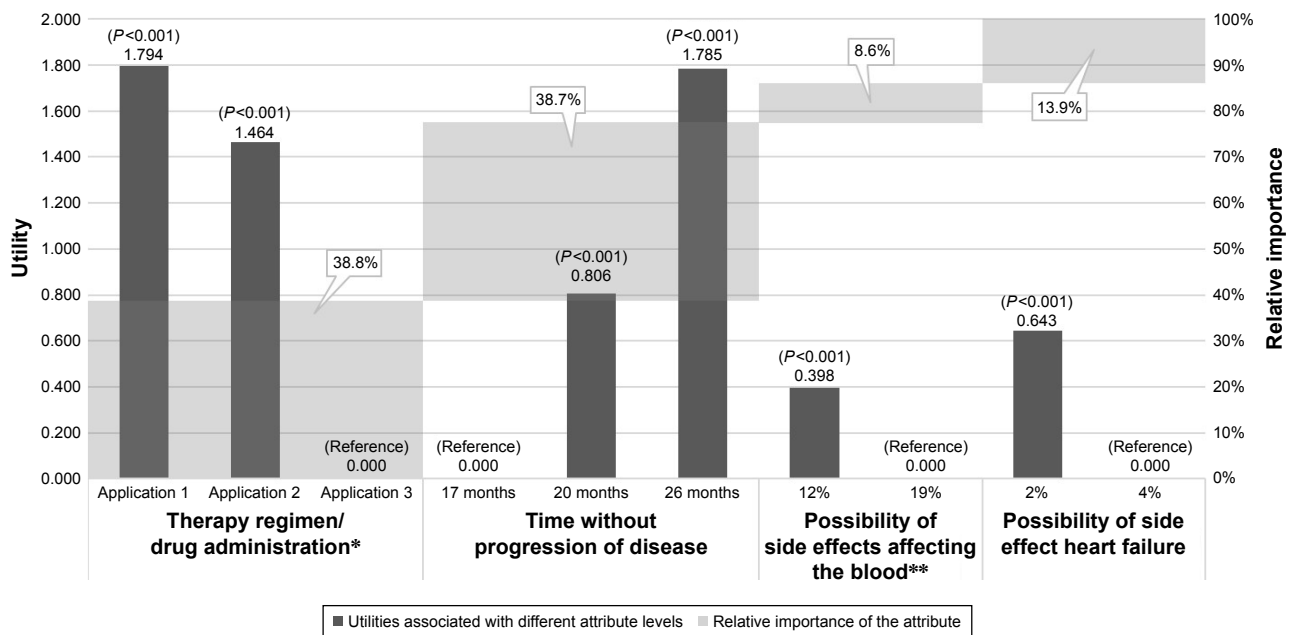


Figure 2 Relative importance of different treatment attributes and utilities of different treatment attribute levels, based on conditional logit regression analysis of the discrete choice experiment data.

Notes: *Application 1 describes the following therapy regimen/drug administration: oral intake once daily and once per week; physician visit once per month with a duration of approximately 2 hours; patient does not need to be accompanied; Application 2 describes the following therapy regimen/drug application: oral intake once daily and twice per week; physician visit once per month with a duration of approximately 2 hours; patient does not need to be accompanied; Application 3 describes the following therapy regimen/drug application: oral intake once daily and once per week; physician visit twice per week, including administration of one infusion per visit, with a duration per visit of approximately 3-4 hours; patient needs to be accompanied. **This attribute includes thrombocytopenia, neutropenia, anemia and hypokalemia.

Table 2 Utilities of different overall treatment options

	Progression-free survival in months (reference value: 17 months associated with utility=0.0)	Possibility of side effects affecting the blood (%) (reference value: 19% associated with utility=0.0)			Possibility of side effect "heart failure" (%) (reference value: 4% associated with utility=0.0)	Administration route (reference: application 3 is associated with utility=0.0)	Total utility based on assumed treatment profile
		Mean	Max	Min			
Carfilzomib ^a (+ lenalidomide + dexamethasone) Profile Related utility	26.3 1.834	18.38 ^a	29.6	9.4	3.8 0.062	Application 3 ^b 0.0	2.386 1.355 1.928
			-0.541				
		0.032					
Ixazomib ^c (+ lenalidomide + dexamethasone) Profile Related utility	20.6 0.904	13.80	23	4.2	2 0.615	Application 2 ^c 1.434	3.708 2.749 3.218
			-0.204				
		0.265					
Dual therapy ^b (lenalidomide + dexamethasone) Profile Related utility	17.6 0.161	13.50	26.5	1.1	2 0.615	Application 1 ^a 1.713	3.402 2.107 2.769
			-0.382				
		0.280					

Notes: ^aTreatment option described by the following attributes: progression-free survival of 26.3 months, mean possibility of side effects \geq grade 3 affecting the blood (including hypokalemia with 9.4%, neutropenia with 29.6%, thrombocytopenia with 16.6%, anemia with 17.9%) of 18.375%, possibility of heart failure (\geq grade 3) of 3.8, and application regimen 3 (oral intake once daily and once per week; physician visit twice per week, including administration of one infusion per visit, with a duration per visit of approximately 3–4 hours; the patient needs to be accompanied).³ ^bTreatment option described by the following attributes: progression-free survival of 17.6 months, mean possibility of side effects \geq grade 3 affecting the blood (including hypokalemia with 1.1% [5]/4.9% [3], neutropenia with 24.0% [5]/26.5% [3], thrombocytopenia with 9.0% [5]/12.3% [3], anemia with 13.0% [5]/17.2% [3]) of 13.50%, possibility of heart failure (\geq grade 3) of 2.0%, and application regimen 1 (oral intake once daily and once per week; physician visit once per month with a duration of approximately 2 hours; the patient does not need to be accompanied).^{3,5} ^cTreatment option described by the following attributes: progression-free survival of 20.3 months, mean possibility of side effects \geq grade 3 affecting the blood (including hypokalemia with 4.2%, neutropenia with 23.0%, thrombocytopenia with 19.0%, anemia with 9.0%) of 13.80%, possibility of heart failure (\geq grade 3) of 2.0%, and application regimen 2 (oral intake once daily and twice per week; physician visit once per month with a duration of approximately 2 hours; the patient does not need to be accompanied).⁵

utilities for complete treatment options can be derived. These take into account trade-offs patients make during their decisions. Among the tested treatments, an IRd treatment showed the highest utility, if compared with Rd or CRd. Utility values per attribute level can also be applied to other treatments, as long as these can be described by the four tested attributes in the study presented.

As far as the authors of this study are aware, five published studies so far dealt with patient preferences in MM.^{7–11} In terms of methodology (number and type of attributes tested, patient inclusion criteria, sample size, preference elicitation technique, multicenter vs single center), there are substantial differences between these studies which make it challenging to compare their results with those of our analysis. Three studies assessed patient preferences based on a DCE methodology.^{7,8,11} These studies confirmed, as our study did, that MM patients are able to express treatment preferences, even if different treatments are associated with specific advantages/disadvantages. Moreover, these studies,

as our study, confirmed that patients are more interested in a higher efficacy compared with safety. Our study differs from other studies in the fact that it addressed RRMM patients only and that it showed a higher importance of the application mode of the therapy than previous studies did.

Limitations

We acknowledge some limitations of our study. The design of the DCE used in this study consisted of complex attributes. Appropriate measures had been taken to increase user-friendliness, understandability, and convenience of the DCE choice sets. Nevertheless, it cannot completely be excluded that patients might have had difficulties in understanding the different treatment options presented on the choice cards.

Inclusion of all attributes describing the different RRMM treatment options was not possible because this would have resulted in too many choice cards. Only attributes ranked as most important in FGDs were selected for the DCE choice cards. However, both attributes in regard to blood-related AEs

and different aspects of drug application (frequency, necessity to visit physicians) were shown to patients as one respective attribute (“possibility of side effects affecting the blood,” “therapy regimen/drug administration”), so that a broader range of aspects of therapy options could be incorporated. This method may have led to an underrating of the importance of the single components of these attributes. Furthermore, it is highly probable that OS is also an attribute that would matter from a patient’s perspective. However, as it was the aim to compare new PI-based treatments and OS data were not available in sufficient maturity for these treatments at the time of the data collection, we needed to exclude this attribute. Nevertheless, in our view progression-free time is a very important outcome in MM as progression is often accompanied by bone pain and/or kidney failure, which reduces the quality of life of patients substantially. Additionally, as we aimed to present our patients easy to understand treatment choices, we used progression-free time as an attribute instead of the more difficult to understand outcome progression-free survival. We acknowledge that there exist differences in the definition of these outcomes. Moreover, due to the design of the study and associated data collection, we were not able to assess the current and previous treatment of interviewed patients. Both current and previous treatments might have influenced patients’ DCE responses. Finally, we included 67 out of 84 patients via PSGs, and their RRMM diagnosis was self-reported and thus, to a certain degree, uncertain. However, as 86.6% of them reported to have received a stem cell transplantation, we see a high probability that these patients indeed suffered from RRMM.

Conclusion

Because RRMM treatment options are constantly evolving and differ from each other in several attributes, it is important to investigate the preferences of patients in that respect. Our study shows that RRMM patients prefer effective and safe treatments, but are willing to accept a lower progression-free time if this is associated with an all-oral application mode of the therapy.

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Disclosure

TW has received honoraria from several pharmaceutical/consultancy companies, eg, Novo Nordisk, Abbvie; Merck;

GSK, BMS, LEO Pharma, Astra Zeneca, Bayer, Boehringer Ingelheim, Pharmerit. SM and SB participated in this study as a staff member of Ingress-health; the work of Ingress-health in this study was sponsored by Takeda Pharma Vertrieb GmbH & Co. KG. SP, LP, and BAR are employees of Takeda Pharma Vertrieb GmbH & Co. KG. HS received consultancy fees, project funding, and reimbursement of travel costs from different pharmaceutical companies, such as Celgene, Janssen-Cilag, Novartis, BMS, Amgen, Takeda. The authors report no other conflicts of interest in this work.

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Supplementary materials

Details regarding conducted targeted literature review

Date of literature search: 21 March 2016

Inclusion criteria:

- Article written in English.
- Article published during the last 10 years (related to the time of conducting the literature search).
- Answering the research question of the study.

Search strings as entered in PubMed:

“RRMM AND DCE,” “Multiple AND myeloma AND DCE,” “Multiple AND myeloma AND Discrete choice experiment,” “relapsed AND refractory AND multiple AND

myeloma AND discrete choice experiment,” “RRMM AND discrete choice experiment,” “relapsed refractory multiple myeloma AND treatment,” “RRMM AND treatment,” “attribute AND oncology AND preference,” “attribute AND cancer AND preference,” “relapsing refractory multiple myeloma AND ixazomib,” “RRMM AND ixazomib,” “discrete choice experiment AND oncology,” “discrete choice experiments AND Onkologie,” “discrete choice experiments AND Multiple Myelom,” “DCE AND Oncology AND cancer,” “oncology AND dce AND preference,” “oncology AND dce AND treatment,” “dce AND Multiple Myeloma.”

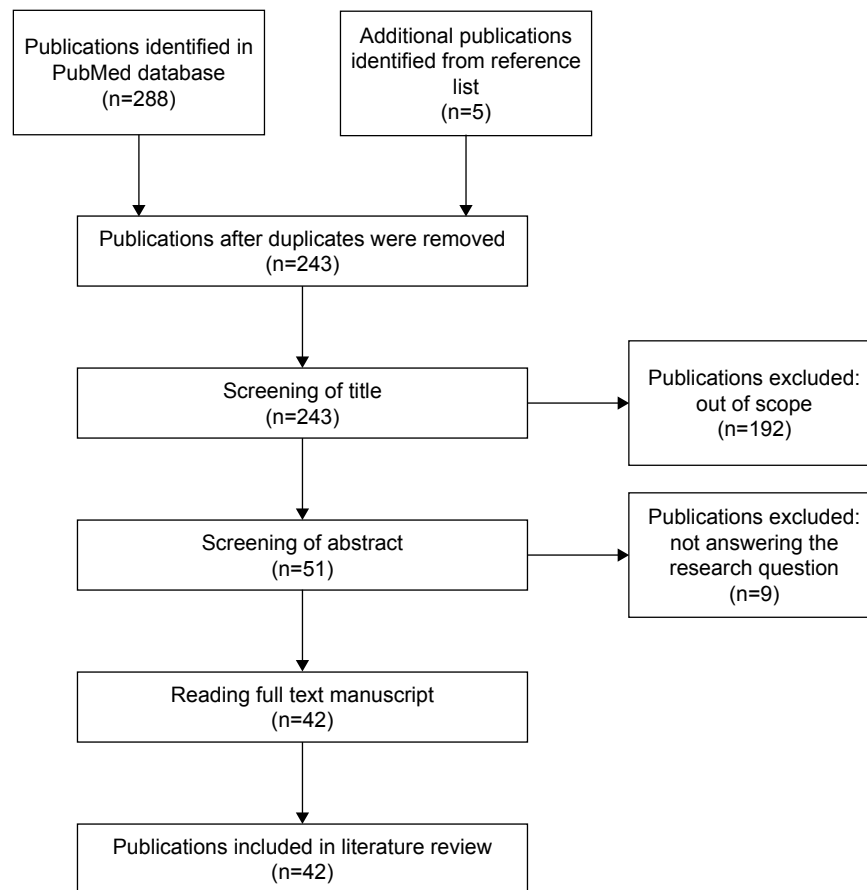


Figure S1 The PRISMA flowchart illustrating the publication selection process.

Table S1 Relative importance of different treatment attributes in defined RRMM patient

Attributes	Relative importance, expressed in % for "Gender"		Relative importance, expressed in % for "Age"		Relative importance, expressed in % for "Disease duration"		Relative importance, expressed in % for "Distance to treating physician"	
	Male (n=53)	Female (n=31)	≤63.5 years (n=42)	>63.5 years (n=42)	≤4.18 years (n=42)	>4.18 years (n=42)	≤15 km (n=45)	>15 km (n=39)
Therapy regimen/drug administration	39.20**	39.25**	37.66**	40.07**	38.43**	39.32**	35.44**	42.41**
Time without progression of disease	35.55**	43.20**	42.32**	35.21**	39.75**	37.38*	38.73**	38.95**
Possibility of side effects affecting the blood	8.38*	8.47*	4.64	12.00*	6.78#	10.60*	10.04*	7.04#
Possibility of side effect heart failure	16.86**	9.08**	15.37**	12.73**	15.04**	12.70**	15.79**	11.59**
Decision	100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00

Notes: Significance levels: **P<0.001; *P<0.05; #P<0.10 (significance of the respective attribute within the conditional logit model). This table gives an overview of the importance of treatment attributes for different patient. Separate conditional logit regression models were estimated for each of the presented subgroups. Threshold: median.

Abbreviation: RRMM, relapsed refractory multiple myeloma.

Table S2 Utilities for different hypothetical possible treatment options, expressed in months of progression-free survival that patients would be willing to trade

Therapy regimen/drug administration			Possibility of side effects affecting the blood ^a		Possibility of side effect heart failure		Utility (expressed in months of PFS patients would be willing to trade)	Rank
Application 1 ^b	Application 2 ^c	Application 3 ^d	12%	19%	2%	4%		
X			X		X		14.44	1
	X		X		X		12.94	2
X				X	X		12.52	3
X			X			X	11.13	4
	X			X	X		11.02	5
	X		X			X	9.63	6
X				X		X	9.21	7
	X			X	X	X	7.71	8
		X	X		X		5.22	9
		X	X	X	X		3.31	10
		X	X			X	1.92	11
		X	X	X		X	0.00	12

Notes: ^aThis attribute includes thrombocytopenia, neutropenia, anemia, and hypokalemia. ^bApplication 1 describes the following therapy regimen/drug application: "oral intake once daily and once per week; physician visit once per month with a duration of approximately 2 hours; patient does not need to be accompanied." ^cApplication 2 describes the following therapy regimen/drug application: "oral intake once daily and twice per week; physician visit once per month with a duration of approximately 2 hours; patient does not need to be accompanied." ^dApplication 3 describes the following therapy regimen/drug application: "oral intake once daily and once per week; physician visit twice per week, including administration of one infusion per visit, with a duration per visit of ~3–4 hours; patient needs to be accompanied." This table shows the results of the estimation of the patients' utilities for different combinations of the three treatment-describing attributes application, side effects affecting the blood, and side effect heart failure. In the table, the negative utility associated with a shorter progression-free survival (PFS) time was translated into a "willingness to accept" this shorter time to receive a treatment with more favorable attributes regarding administration regimen and side effect profile.

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