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RESEARCH ARTICLE

Double Relapsed and/or Refractory Multiple Myeloma: Clinical Outcomes and Real World Healthcare Costs

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Abstract

Double relapsed and/or refractory multiple myeloma (DRMM), MM that is relapsed and/or refractory to bortezomib and lenalidomide, carries a poor prognosis. The healthcare costs of DRMM have not previously been reported. We analyzed detailed medical resource utilization (MRU) costs, drug costs and outcomes for 39 UK patients receiving standard DRMM therapy. Median OS in this cohort was 5.6 months. The mean cost of DRMM treatment plus MRU until death was £23,472 [range: £1,411–£90,262], split between drug costs £11,191 and other resource use costs £12,281. The cost per assumed quality-adjusted life year (QALY) during DRMM was £66,983. These data provide a standard of care comparison when evaluating the cost-effectiveness of new drugs in DRMM.

Introduction

Patients with treatment-refractory malignancy have poor outcomes and high healthcare costs. In Multiple Myeloma (MM), the introduction of the proteasome inhibitor bortezomib and immunomodulatory drugs (IMiDs) thalidomide and lenalidomide has improved survival over the last decade [1], but increased the cost of treatment. While these drugs can result in remission, most patients will relapse with increasing symptom burden and worsening prognosis [2]. Double relapsed and/or refractory multiple myeloma (DRMM), MM that is relapsed and/or refractory to bortezomib and lenalidomide [3], carries a poor prognosis and therapeutic options remain limited. An IMWG retrospective analysis of patients who relapsed following bortezomib and at least one of the IMiDs showed a median overall survival (OS) and progression-free survival (PFS) of 9 months and 5 months respectively [4]. Only those potentially eligible for further clinical trials with Eastern Cooperative Oncology Group Performance Status

[research-training-fellowship-crtf/](#)). MS was supported by Oxford Cancer Research Centre Clinical Research Training Fellowship 2014–2017. Grant code OCRC-CRF14-MW (<http://www.cancercentre.ox.ac.uk/graduate-studies/clinical-research-training-fellowships/>). BresMed Health Solutions provided support in the form of salaries for authors AB, JE and DL, but did not have any additional role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript. The specific roles of these authors are articulated in the 'author contributions' section.

Competing Interests: AB, JE and DL are employees of BresMed, which receives consultancy fees from Celgene. SG, IL, MS, PR, JW, ED, KR are members of a hospital department that has received an unrestricted educational grant from Celgene. KR is in receipt of honoraria from Celgene. No author has any direct financial interests whose value could be affected by this publication. This does not alter the authors' adherence to PLOS ONE policies on sharing data and materials.

(ECOG PS) 0–2 were included, indicating survival may have been overestimated relative to the entirety of this heavily treated population.

Studies of healthcare costs for relapsed and/or refractory MM patients receiving bortezomib and/or lenalidomide based regimens have been undertaken in a variety of countries, using 'real-world' data and/or economic modelling, often from patients enrolled in clinical trials [5–9]. However, none of them address MRU for patients relapsed after bortezomib and lenalidomide. The third-generation IMiD pomalidomide and second-generation proteasome inhibitor carfilzomib have recently been licensed and have efficacy in DRMM [10]. The cost burden and clinical outcomes outside of trials of this phase of the disease has never been established, so new therapies with efficacy in DRMM have no published benchmark against which to judge cost-effectiveness. To ensure resources are allocated appropriately, the cost-effectiveness evaluation of these therapies in the DRMM setting must involve non-clinical trial, real world MRU data from relevant patients. Our data provides a standard of care comparison when evaluating cost-effectiveness of new drugs in DRMM.

Methods

Ethics Statement

All patients whose data were included in this study had provided written consent for the use of their anonymised data for the purposes of audit and service improvement by the Thames Valley Cancer Network, UK. This study was appropriately registered using the clinical audit project proposal system at Oxford University Hospitals NHS Trust, UK. The Clinical Audit Lead reviewed and approved the proposal, and in line with the UK NHS National Research Ethics Service (NRES) guidance, deemed it not to require IRB or ethics committee approval.

Anonymised data on clinical outcomes, anti-myeloma therapies prescribed and MRU were obtained for 39 DRMM patients pre-treated with or intolerant to bortezomib and lenalidomide in the Thames Valley Cancer Network, UK from 2011 to 2014 (Table A in [S1 File](#)). Based on the UK National Institute for Health and Care Excellence (NICE) guidelines, bortezomib-based therapy was used for first relapse unless contraindicated. Lenalidomide-dexamethasone combination was approved for second and subsequent relapse. Relapsed myeloma patients were identified using pharmacy-generated lists of all sequential lenalidomide recipients between January 2011 and July 2013 at Oxford University Hospitals and the Royal Berkshire Hospital, Reading, UK. The strategy of using hospital pharmacy dispensing data to identify subjects ensured all lenalidomide recipients during the study period were included in the analysis, as long as they had Multiple Myeloma and had progressed on or were refractory to lenalidomide, according to IMWG criteria [11]. 34 (87%) patients were pre-treated with lenalidomide and bortezomib, and had relapsed following, or failed to tolerate both therapies. 5 patients had not received bortezomib due to pre-existing peripheral neuropathy, sufficiently severe to contraindicate its use.

OS and PFS were calculated from the start date of the first therapy following relapse after lenalidomide ('1st DRMM therapy') until either IMWG criteria for progression/relapse were reached or death. The data censor date was 10 January 2014 for patients still alive. For each patient the following occurrences of MRU were retrieved from health care records from the start of each successive DRMM therapy until death or censoring: detailed drug regimens, outpatient clinic and chemotherapy unit attendances, inpatient/hospice admissions, supportive therapies, medical procedures, radiological investigations, blood product transfusions and blood tests. Costings were calculated using NHS reference costs 2012–13. These were combined to give one MRU cost from the start of DRMM therapy to death or censoring, using a micro-costing approach. Drug costs were separately calculated for each successive DRMM therapy.

Results

When offered DRMM therapy, 67% of patients preferred active treatment to palliative care (Table 1). The median age of those who chose palliative care was 73.2 years compared to 59.9 for active therapy ($p < 0.001$). The first DRMM treatment regimen was typically bendamustine, thalidomide and dexamethasone (43.6%) as published previously [12]. Retreatment with bortezomib (15.4%) or lenalidomide (33.3%) based regimens was used if poor bone marrow reserve precluded bendamustine use and suitable clinical trial alternatives were lacking.

Regimen choice was made by the treating clinician and based on ECOG PS and response/side effect profile of prior therapies. Third-generation IMiDs and second-generation proteasome inhibitors were not routinely available to this cohort, excepting pomalidomide in one patient. Two patients who had not previously received bortezomib due to neuropathy did receive it at DRMM with no recorded worsening of neuropathy; one had progressive disease despite its use and the other had a partial response of 6 months duration.

Median PFS was 5.2 months and median OS was 5.6 months from start of DRMM therapy (Fig 1). These statistics reflect a steep drop in survival early on with a few patients surviving significantly longer. 24/39 (61.5%) patients had died by the end of follow-up, of whom 79% died in hospital/ hospice. The cohort was deemed too small and heterogeneous for analysis of regimen effect on survival.

MRU was high in this cohort in comparison with that usually observed at earlier lines of therapy (Table 2). 60 inpatient hospital admissions occurred during DRMM therapy in 39 patients. Admissions lasted 9.3 days on average, for indications including pain management, renal failure and most commonly neutropenic fever, with some cases of culture-confirmed septicemia. Prolonged inpatient admissions, frequent AEs and high transfusion requirements reflect a poor quality of life (QoL) among these patients (Table 3). The most common recorded Grade 3–4 adverse events (AEs) were anaemia (43.6%), thrombocytopenia (28.2%) and bone pain (33.3%) (Table 3, Table B in S1 File).

The mean total MRU cost per patient from start of DRMM therapy until death or censor was £12,281 (Table 2). This comprises: clinic attendances £5,007 (41%); inpatient admissions £2,884 (20%); transfusions £2,479 (23%); supportive therapy £1,454 (12%); radiology/procedures £323 (3%); blood tests £151 (1%). The mean drug cost of DRMM therapy is estimated to be £11,191 per patient. The mean total cost of treatment plus MRU is therefore £23,472 [range: £1,411 - £90,262]; £760 per week of life with DRMM. Formal QoL data is lacking in this retrospective cohort but has been previously published for DRMM patients in the MM-003 trial [13]. Assuming that QoL was the same in this cohort (utility 0.59) and remained constant throughout patients' lifetimes, our analysis indicated a cost per QALY of £66,983.

Conclusion

Although the small sample size of this cohort limits the ability to draw definitive survival conclusions, PFS and OS were poor, with a wide range due to the inclusion of all patients whether treated with active or palliative intent, but were similar to published examples [4]. Despite the poor outcomes, up to two thirds of DRMM patients want to be treated with active intent to improve survival, reiterating the need to develop therapies that give patients an improved prognosis, whilst being cost-effective and well tolerated in a heavily pre-treated patient group. The heterogeneity of this cohort is acknowledged but intentional, representing a typical real world hospital cohort of patients, where a range of therapeutic options must be employed, constrained by varying patient-related factors such as ECOG PS, drug tolerance, social situation and patient choice.

Table 1. Patient characteristics.

Median age at diagnosis, years (range)*	64.3 (45–79)
Median age by treatment intent: Active/ Palliative	59.9 (n = 26)/ 73.2 (n = 13)
Isotope: IgA/ IgG/ Light chain	7 (51.3%)/ 20 (18.0%)/ 12 (30.8%)
ISS stage I at diagnosis	6 (15.4%)
ISS stage II at diagnosis	9 (23.1%)
ISS stage III at diagnosis	10 (25.6%)
ISS stage at diagnosis unknown	14 (35.9%)
Previous thalidomide-based treatment	34 (87.2%)
Previous bortezomib-based treatment	34 (87.2%)
Previous lenalidomide-based treatment	39 (100.0%)
Previous high dose melphalan with stem cell rescue	17 (43.5%)
Previous additional alternative treatment(s):	
Vincristine-based regime	7 (17.9%)
Melphalan-based regime	5 (12.8%)
Allograft	1 (2.6%)
Median years from diagnosis to DRMM point (range)*	4 years 9.5 months (6.5 mo– 10.5 y)
Median number regimes prior to DRMM (range)	4 (2–5)
1st DRMM therapy contained bendamustine	17 (43.6%)
1st DRMM therapy contained bortezomib	6 (15.4%)
1st DRMM therapy DT-PACE	1 (2.6%)
1st DRMM therapy contained lenalidomide	13 (33.3%)
No treatment given at DRMM	2 (5.1%)
2nd DRMM therapy (n = 7):	
Bendamustine, thalidomide, dexamethasone	2 (28.6%)
Melphalan, dexamethasone	2 (28.6%)
Thalidomide-based regime	2 (28.6%)
Pomalidomide, dexamethasone	1 (14.3%)
3rd DRMM therapy (n = 4):	
Bortezomib, cyclophosphamide, dexamethasone	2 (50.0%)
Bortezomib, melphalan, prednisone	1 (25.0%)
Bendamustine, thalidomide, dexamethasone	1 (25.0%)
Response to 1st DRMM therapy:	
CR	1 (2.6%)
VGPR	2 (5.1%)
PR	8 (20.5%)
SD	13 (33.3%)
PD or death within first month	14 (35.9%)
Unknown	1 (2.6%)
Duration of treatment (SD)	104.9 days (63.8)
Response to 2nd DRMM therapy:	
PR	3 (42.9%)
SD	2 (28.6%)
PD or death within first month	2 (28.6%)
Response to 3rd DRMM therapy:	
PR	2 (50.0%)
PD or death within first month	1 (25.0%)

(Continued)

Table 1. (Continued)

Unknown	1 (25.0%)
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*Date of diagnosis unavailable in 2 cases. ISS: International Staging System; DRMM: Double Relapsed and/or Refractory Multiple Myeloma; CR: Complete response; VGPR: Very good partial response; PR: Partial response; SD: Stable disease; PD: Progressive disease [11]

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This is the first report of non-clinical trial based ‘real-world’ MRU cost analysis in the setting of DRMM. These patients have high MRU costs. A comparable cost analysis report relates to a subset of 54 ‘4th line’ real world relapsed/refractory patients in a Netherlands study, recruited from a previous trial cohort [7]. However, it is likely that this cohort (data collected

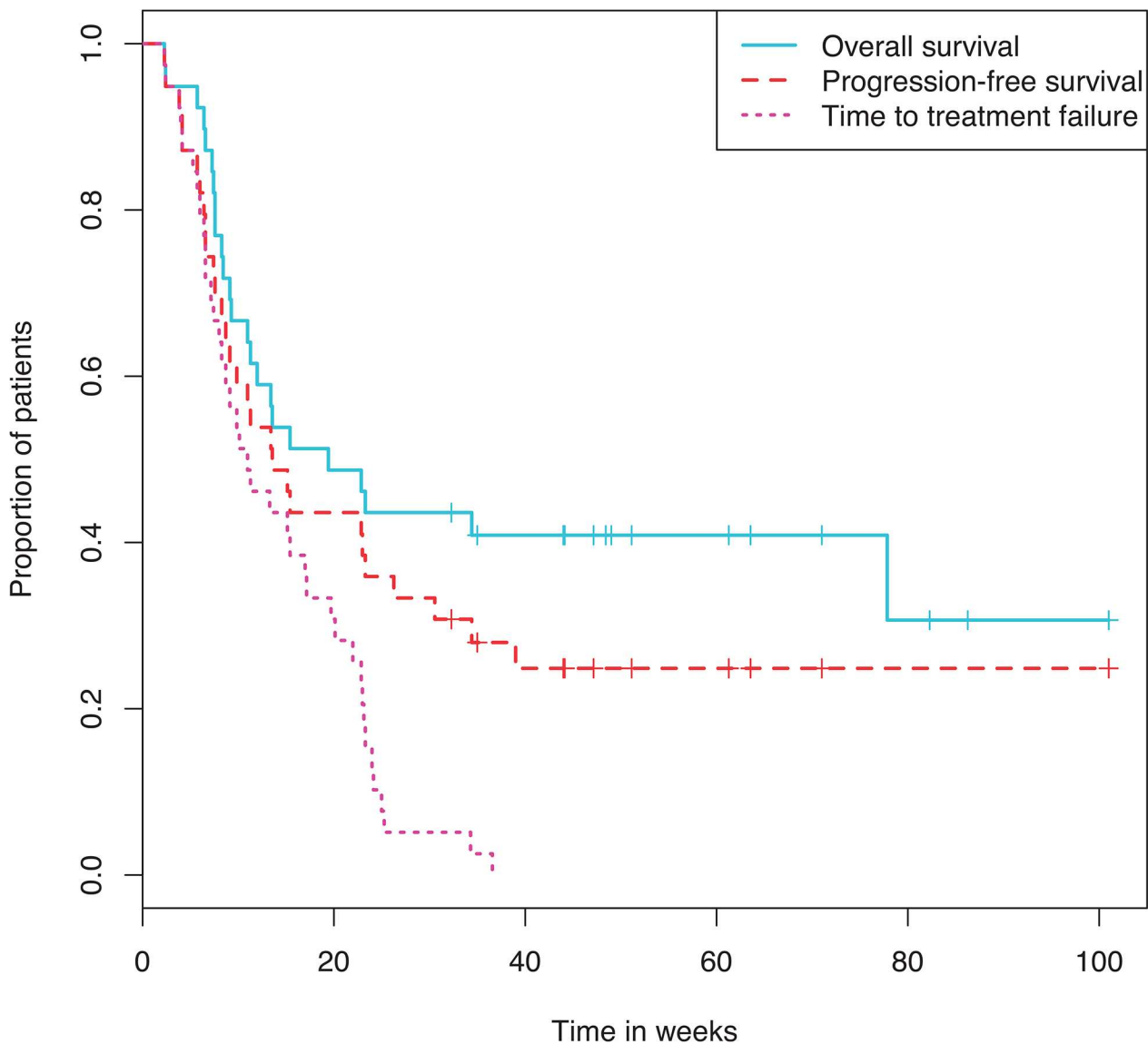


Fig 1. Kaplan Meier curve showing overall survival, progression free survival and time to treatment failure.

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Table 2. Medical Resource Utilisation Costs of Double Relapsed and/or Refractory Multiple Myeloma Therapy.

MRU Category	Treatment given	Occurrences	Cost per Patient		
Drug costs					
1st DRMM therapy (per 28 day cycle) ^a	Bortezomib-based	6/39	£4,022		
	Lenalidomide-based	13/39	£3,913		
	DT-PACE	1/39	£946		
	Bendamustine-based	17/39	£1,332		
	No active treatment	2/39	£0		
	Average		£2,532		
2nd DRMM therapy (per 28 day cycle) ^a	Ben/ Thal/ Dex	2/39	£853		
	Mel/ Dex	2/39	£133		
	Thal	2/39	£298		
	Pom/ Dex	1/39	£8,887		
	No 2 nd DRMM therapy	32/39	£0		
Average		£294			
3rd DRMM therapy (per 28 day cycle) ^a	Bor/ Cyc/ Dex	2/39	£4,118		
	Bor/ Mel/ Pred	1/39	£3,847		
	Ben/ Thal/ Dex	1/39	£1,983		
	No 3 rd DRMM therapy	35/39	£0		
Average		£361			
Total 1 st DRMM therapy drug costs (all cycles)			£9,527		
Total 2 nd DRMM therapy drug costs (all cycles)			£807		
Total 3 rd DRMM therapy drug costs (all cycles)			£857		
Total drug costs (all cycles)			£11,191		
		Occurrences during DRMM period ^d		Cost per Patient	Range
MRU Category	MRU Item	Mean	SD		
Other MRU costs					
Inpatient admissions	Night as inpatient	9.3	7.9	£2,463	£0, £16,169
	Outpatient	4.2	4.1	£630	£0, £3,012
Attendances	Day therapy unit	12.8	9.7	£4,331	£0, £11,848
	Triage, not admitted	0.4	0.9	£46	£0, £344
	CT scan	0.4	0.8	£42	£0, £435
Invasive and radiological procedures	MRI scan	0.3	0.6	£57	£0, £343
	X-ray	1.1	2.7	£31	£0, £311
	Maxillofacial	0.1	0.2	£21	£0, £406
	Other ^b	0.3	0.5	£172	£0, £3,225
Supportive therapy	Bisphosphonate ^c	2.6	2.5	£217	£0, £713
	Radiotherapy	1.1	3.6	£1,237	£0, £21,643
Transfusion	Red blood cells (units)	5.9	6.0	£1,684	£0, £5,993
	Platelets (units)	2.3	3.8	£1,200	£0, £6,784
	Full blood count	21.6	13.0	£65	£0, £196
Blood tests	Biochemistry	20.6	15.6	£26	£0, £110
	Immunology	4.4	2.9	£22	£0, £60
	Microbiology	5.6	7.0	£38	£0, £278
Total other MRU costs			£12,281		£995,

(Continued)

Table 2. (Continued)

		£40,274
Total (drug costs and other MRU costs)	£23,472	£1,411,
		£90,262

MRU: Medical Resource Utilization; DRMM: Double Relapsed and/or Refractory Multiple Myeloma; Ben: Bendamustine; Thal: Thalidomide; Dex: Dexamethasone; Mel: Melphalan; Pom: Pomalidomide; Bor: Bortezomib; Cyc: Cyclophosphamide; Pred: Prednisone.

^aDrug costs have been calculated as the average of all patients undertaking each regimen. Dosing changes have been incorporated where provided. Additional dosing regimen details have been taken from product SPCs. Costs are taken from BNF or eMIT; and are applied using the appropriate pack/vial size. The average surface area of a patient (used for IV therapies) is taken from the MM-003 clinical trial (approximately 1.86m²). Differences in costs of the same treatment between treatment lines are caused by differences in dosing for individual patients.

^bOther medical procedures consisted of: 1 vertebroplasty; 1 facet joint injection; 1 endoscopy; 1 bronchoscopy; 1 hip fracture repair under general anaesthetic; 2 PET scans and 4 ultrasound scans.

^cBisphosphonate costs calculated assuming all patients on bisphosphonates are on an average dose. The figure shows the approximate number of cycles for which patients are on bisphosphonate treatment.

^dAll MRU occurrences were recorded from initiation of 1st DRMM therapy until the end of follow up (or death)

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from 2001 to 2009) included patients who received bortezomib (n = 12) and/or lenalidomide (n = 20) for the first time at 4th line, as they were first made available during the study period. At €32,889 per patient (range: €1,055–€144,967), costs reported in that study are comparable to our findings. However an estimated cost per QALY of £66,983 is a significant increase on that reported by Brown et al in a UK study of the cost effectiveness of Lenalidomide-based therapies after one prior therapy [5], where cost was £30,153/QALY. The difference reflects the limited benefit to survival of any current therapy at DRMM, and the higher MRU costs at this later stage of disease.

Table 3. Surrogates of Quality of Life during Double Relapsed and/or Refractory Multiple Myeloma Therapy.

Surrogates of Quality of Life ^a	Number of patients
Grade 3–4 Adverse events during DRMM therapy:^b	
Anaemia	17 (43.6%)
Neutropaenia	7 (17.9%)
Thrombocytopenia	11 (28.2%)
Bleeding	1 (2.6%)
Febrile neutropaenia	6 (15.4%)
Bone pain	13 (33.3%)
Acute renal failure	4 (10.3%)
Dehydration / vomiting / diarrhoea	2 (5.1%)
Admissions during DRMM therapy (SD)	1.3 per patient (1.1)
Duration of admissions (SD)	9.3 days (7.9)
Outpatient clinic appointments during DRMM therapy (SD)	4.2 per patient (4.1)
Day therapy unit visits during DRMM, including CT & MRI (SD)	12.2 per patient (9.7)
RBC units during DRMM therapy (SD)	5.9 per patient (6.0)
Platelet units during DRMM therapy (SD)	2.3 per patient (3.8)

^aAll Surrogates of Quality of Life were recorded from initiation of 1st DRMM therapy until the end of follow up (or death)

^bAdverse events with no recorded grade are assumed to be grade 3 or 4.

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Compared with a decade ago, the price range of new anticancer agents has more than doubled [14], and the cost of care analysis of these agents is imperative. New MM therapies carfilzomib and pomalidomide have been priced significantly higher than currently available anti-myeloma drugs. Any subsequent cost benefit analysis comparisons performed in DRMM patients must be set in the context of the high background MRU as observed in our cohort. It is highly relevant that the cost per assumed QALY in this cohort is double that usually accepted by the UK National Institute for Health and Care Excellence [15]. In addition to improving survival, therapies that induce higher response rates or arrest disease progression could potentially increase therapy costs, but lower MRU costs and improve QoL if progression is halted. Biomarkers that focus use of new drugs to cohorts of patients where maximum benefit is obtained would improve cost-effectiveness further. New treatments should be compared with real-world non-trial outcomes such as that provided here, to give a realistic picture of the value of these new drugs.

Supporting Information

S1 File. Table A: Complete Data collected (excepting adverse events) on 39 patients with Double Refractory Multiple Myeloma. Table B: Adverse Events recorded for 39 patients with Double Refractory Multiple Myeloma.
(XLS)

Author Contributions

Conceived and designed the experiments: SG KR. Performed the experiments: SG IL MS ED KR. Analyzed the data: SG AB JE DL KR. Contributed reagents/materials/analysis tools: PR JW. Wrote the paper: SG DL KR.

References

1. Pozzi S, Marcheselli L, Bari A, Liardo EV, Marcheselli R, Luminari S, et al. Survival of multiple myeloma patients in the era of novel therapies confirms the improvement in patients younger than 75 years: a population-based analysis. *British journal of haematology*. 2013; 163: 40–46. doi: [10.1111/bjh.12465](https://doi.org/10.1111/bjh.12465) PMID: [23889344](https://pubmed.ncbi.nlm.nih.gov/23889344/)
2. Jordan K, Proskorovsky I, Lewis P, Ishak J, Payne K, Lordan N, et al. Effect of general symptom level, specific adverse events, treatment patterns, and patient characteristics on health-related quality of life in patients with multiple myeloma: results of a European, multicenter cohort study. *Supportive care in cancer: official journal of the Multinational Association of Supportive Care in Cancer*. 2013; 22: 417–426.
3. Meadows JP and Mark TM Management of Double-Refractory Multiple Myeloma. *Curr Hematol Malig Rep*. 2013; 8: 253–260. doi: [10.1007/s11899-013-0173-2](https://doi.org/10.1007/s11899-013-0173-2) PMID: [23975677](https://pubmed.ncbi.nlm.nih.gov/23975677/)
4. Kumar SK, Lee JH, Lahuerta JJ, Morgan G, Richardson PG, Crowley J, et al. Risk of progression and survival in multiple myeloma relapsing after therapy with IMiDs and bortezomib: a multicenter international myeloma working group study. *Leukemia*. 2012; 26: 149–157. doi: [10.1038/leu.2011.196](https://doi.org/10.1038/leu.2011.196) PMID: [21799510](https://pubmed.ncbi.nlm.nih.gov/21799510/)
5. Brown RE, Stern S, Dhanasiri S and Schey S Lenalidomide for multiple myeloma: cost-effectiveness in patients with one prior therapy in England and Wales. *Eur J Health Econ*. 2013; 14: 507–514. doi: [10.1007/s10198-012-0395-6](https://doi.org/10.1007/s10198-012-0395-6) PMID: [22572968](https://pubmed.ncbi.nlm.nih.gov/22572968/)
6. Durie B, Binder G, Pashos C, Khan Z, Hussein M and Borrello I Total cost comparison in relapsed/refractory multiple myeloma. *J Med Econ*. 2013; 16: 614–622. doi: [10.3111/13696998.2012.760159](https://doi.org/10.3111/13696998.2012.760159) PMID: [23281721](https://pubmed.ncbi.nlm.nih.gov/23281721/)
7. Gaultney JG, Franken MG, Tan SS, Redekop WK, Huijgens PC, Sonneveld P, et al. Real-world health care costs of relapsed/refractory multiple myeloma during the era of novel cancer agents. *J Clin Pharm Ther*. 2013; 38: 41–47. doi: [10.1111/jcpt.12020](https://doi.org/10.1111/jcpt.12020) PMID: [23126374](https://pubmed.ncbi.nlm.nih.gov/23126374/)
8. Gaultney JG and Uyl-de Groot CA Efficient allocation of novel agents in multiple myeloma: a work in progress. *The oncologist*. 2013; 18: 5–7. doi: [10.1634/theoncologist.2012-0484](https://doi.org/10.1634/theoncologist.2012-0484) PMID: [23299778](https://pubmed.ncbi.nlm.nih.gov/23299778/)

9. Teitelbaum A, Ba-Mancini A, Huang H and Henk HJ Health care costs and resource utilization, including patient burden, associated with novel-agent-based treatment versus other therapies for multiple myeloma: findings using real-world claims data. *The oncologist*. 2013; 18: 37–45. doi: [10.1634/theoncologist.2012-0113](https://doi.org/10.1634/theoncologist.2012-0113) PMID: [23299776](https://pubmed.ncbi.nlm.nih.gov/23299776/)
10. Lee HC, Shah JJ and Orlowski RZ Novel approaches to treatment of double-refractory multiple myeloma. *Am Soc Clin Oncol Educ Book*. 2013: 302–306.
11. Rajkumar SV, Harousseau JL, Durie B, Anderson KC, Dimopoulos M, Kyle R, et al. Consensus recommendations for the uniform reporting of clinical trials: report of the International Myeloma Workshop Consensus Panel 1. *Blood*. 2011; 117: 4691–4695. doi: [10.1182/blood-2010-10-299487](https://doi.org/10.1182/blood-2010-10-299487) PMID: [21292775](https://pubmed.ncbi.nlm.nih.gov/21292775/)
12. Lau IJ, Smith D, Aitchison R, Blesing N, Roberts P, Peniket A, et al. Bendamustine in combination with thalidomide and dexamethasone is a viable salvage option in myeloma relapsed and/or refractory to bortezomib and lenalidomide. *Annals of hematology*. 2014.
13. Palumbo A, Davies F, Lee D, Dhanasiri S, Facon T, Zaki M, et al. Quality of life weights (utilities) in refractory or relapsed and refractory multiple myeloma (RRMM) patients using EORTC-8D and EQ-5D. *Lymphoma and Myeloma 2013: An International Congress on Hematologic Malignancies 2013: Poster P-03*.
14. Kantarjian HM, Fojo T, Mathisen M and Zwelling LA Cancer Drugs in the United States: Justum Pretium—The Just Price. *Journal of Clinical Oncology*. 2013; 31: 3600–3604. doi: [10.1200/JCO.2013.49.1845](https://doi.org/10.1200/JCO.2013.49.1845) PMID: [23650428](https://pubmed.ncbi.nlm.nih.gov/23650428/)
15. National Institute for Health and Care Excellence. Guide to the methods of technology appraisal 2013 6. The appraisal of the evidence and structured decision-making. 2013; Available: <http://publications.nice.org.uk/guide-to-the-methods-of-technology-appraisal-2013-pmg9/the-appraisal-of-the-evidence-and-structured-decision-making>.