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Bone-independent extramedullary disease is associated with inferior overall survival in multiple myeloma patients: a single center, real-world experience

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Multiple myeloma (MM) is characterised by the accumulation of malignant plasma cells in the bone marrow resulting in lytic bone lesions, hypercalcaemia and anaemia.^{1,2} In up to 20% of myeloma cases at diagnosis, plasma cell aggregates occur outside of the bone marrow, referred to as extramedullary disease (EMD).³⁻⁶ Most extramedullary myeloma manifestations are contiguous with bone (paraosseous, bone-associated) but in a minority of cases, the infiltration occurs in anatomical sites independent of bone. Bone-Independent Extramedullary Disease (BI-EMD) is thought to confer a poorer prognosis compared with paraosseous EMD⁷ although this conclusion is mostly based on relatively small cohorts of patients from observational studies conducted prior to the widespread availability of novel myeloma agents. Most published studies on EMD focus on the clinical characteristics and outcomes of paraosseous EMD and only very few studies have specifically analysed BI-EMD cases with a view to determining associations with known high-risk features. The aim of this study was to determine the incidence, clinical characteristics, association with bone disease, skeletal related events (SRE) including pathological fractures, spinal cord compression, radiotherapy and surgical interventions, as well as response to treatment and overall survival of BI-EMD in a single centre cohort of newly diagnosed myeloma patients.

To determine the clinical characteristics and outcomes of myeloma patients presenting with BI-EMD at the time of diagnosis, we retrospectively analysed the clinical data of symptomatic adult MM patients, as defined by the International Myeloma Working Group (IMWG) criteria, who were treated and actively followed up at University College Hospital, London, United Kingdom, between 1 January 2018 and 31 December 2022. The data cut-off date was 3 May 2023. Bone-independent EMD (BI-EMD) in our study was defined as MM cases with plasma cell infiltration of an anatomical site distant from the bone marrow, diagnosed by whole body imaging. Patients with plasma cell leukaemia and those who did not meet the criteria for MM diagnosis were excluded from the study. P-values were

calculated using logistic regression or Fisher's exact test. This study was conducted in accordance with institutional requirements and the Declaration of Helsinki.

Thirty-three (33) of the 906 patients (3.6%) diagnosed with MM between 2004 and 2022 had BI-EMD at diagnosis (primary BI-EMD) while 22 (2.4%) cases were identified at relapse (secondary BI-EMD). The median duration of follow up was 45 months (range: 1 - 270 months). The clinical characteristics and outcomes of the primary BI-EMD cases are summarised in table 1. The commonest sites for primary BI-EMD were lymph nodes (45%), spleen (15%) and peritoneal nodules (9%). Other affected sites included thyroid, breast, testis, pancreas, larynx, pericardial effusion and pleural effusion. Compared with MM patients without BI-EMD, at least one Skeletal-Related Event (SRE) was present at diagnosis in 61% of patients with primary BI-EMD vs 44% without BI-EMD (p=0.07). There was no statistically significant difference for age, gender, ethnic origin, isotype, ISS stage, high-risk cytogenetics (FISH), bone disease or first line treatment regimen between patients with or without primary BI-EMD. Of the 22 patients with secondary BI-EMD, 12 had BI-EMD at first relapse and 10 at subsequent relapses. The most common secondary BI-EMD sites were lymph nodes (40%), spleen (14%), thyroid (14%), pancreas (14%), peritoneal nodules (9%), and liver (9%).

The induction treatment modalities for primary BI-EMD patients in this study were chemotherapy alone for 29 (88%) while 4 (12%) patients had radiotherapy in combination with chemotherapy. The majority of patients received proteosome inhibitor and immunomodulatory agents or a combination of these (72.8%). An autologous stem cell transplant (ASCT) was performed for 22 patients (67%) including 3 (9%) patients who received tandem ASCTs. Treatment response rates were 25% complete response (CR) or very good partial response (VGPR) for the BI-EMD group compared with 30% CR+VGPR in the patients without BI-EMD. The Overall Response Rate (ORR) was 83.3% (95% CI 62.6-95.3) for the BI-EMD patients and 95.8% (95% CI 93.7-97.3) for the non-BI-EMD patients (p=0.02). Fifteen (45%) primary BI-EMD patients relapsed during the follow up period and at

first relapse, 13 (87%) had non-EMD progression while 2 (13%) had bone associated EMD progression. None of the relapses presented as BI-EMD progression.

Univariate analysis showed inferior overall survival (OS) for MM patients with BI-EMD at diagnosis compared with those without BI-EMD; hazard ratio (HR) 4.62 (95% CI 1.99-10.77, p<0.01). After adjustment for age, disease isotype and SRE at diagnosis this difference remained statistically significant (HR 3.67, 95% CI 1.29-10.43, p=0.01). Kaplan-Meier analysis estimated 3-year Overall Survival rates of 87.7% (95% CI 66.0-95.9) for the BI-EMD group and 97.9% (95% CI 96.6-98.8) for the non-BI-EMD group. (Figure 1)

In this single-centre retrospective study, the rate of BI-EMD in MM patients was similar to previous studies.^{5,8} Usmani et al⁸ similarly analysed newly diagnosed myeloma patients who had baseline PET CT scans and reported EMD in 3.4% of MM patients at diagnosis. He et al³ also observed BI-EMD in 2.8% of cases by PET/CT at diagnosis. Weinstock et al reported 1.2% BI-EMD at the time of diagnosis in a cohort of patients who subsequently received an autologous or allogeneic transplant.⁹ In concordance with previously published observations, BI-EMD is less common (3.2%) at the time of diagnosis compared with paraosseous EMD (13.2%) however, unlike other studies, we did not observe a higher rate of BI-EMD in relapsed MM patients with a secondary BI-EMD rate of 2.4%. The use of FDG PET-CT scan definition for BI-EMD has a number of advantages and drawbacks. It allowed the identification of extramedullary disease in organs or tissues which would have been inaccessible to biopsy for histology. However, given the possibility of non-specific FDG uptake within certain tissues, specificity remains a challenge. It is noteworthy that histological confirmation was done in cases where biopsy was feasible. Nevertheless, correlation was observed in the resolution of extramedullary disease with overall myeloma disease response in individual patients. Interestingly, Gagelman et al⁵ observed from a large registry dataset, a significant increase in incidence rates of EMD from 6.5% to 23.7% between 2005 and 2014. While the authors did not specifically explore the reasons for this increase, it is possible that the recent adoption of sensitive imaging techniques such as PET-

CT and PET-MRI in myeloma may have contributed to this finding. In this retrospective analysis, fit MM patients with EMD were considered for treatment intensification at induction with regimens such as VTD-PACE and PET/CT imaging upstaging was essential for the identification of this group of patients. Evidently, well-designed clinical trials are needed to evaluate treatment regimens for this patient group in the current era of novel therapies including immunotherapies.

As plasma cells are usually dependent on the bone marrow microenvironment for survival, extramedullary MM occurs when malignant plasma cells evolve mechanisms to survive independent of the bone marrow stroma. Therefore, patients with BI-EMD at diagnosis (as opposed to paraosseous EMD and those without EMD) likely represent a group with unique fundamental biology. Several possible mechanisms have been postulated for the development of EMD including downregulation of chemokine receptors such as CCR1 and CXCR4 as well as decreased expression of adhesion molecules.¹⁰ Higher rates of chromosomal aberrations have also been identified in extramedullary plasma cells compared with bone marrow plasma cells in MM patients.¹¹ Given the small sample size, this study was not adequately powered to determine differences in presenting characteristics but interesting trends were observed. For instance, the rate of 17p deletion was 22.7% in MM patients with BI-EMD compared with 7.4% in patients without BI-EMD and this difference was statistically significant (Fishers exact test p=0.02). However, there was no significant difference when all high-risk cytogenetics were compared between the two groups (Table 1). The rate of stable disease (SD) following induction chemotherapy was also relatively higher in BI-EMD patients (16.7% vs 4.2%).

The presence of EMD regardless of the type is associated with inferior overall survival and a poor prognosis.^{3,4,8} Furthermore, findings from another single centre study reported inferior survival for MM patients with BI-EMD compared with paraosseous EMD⁷. Our single centre retrospective analysis showed inferior ORR and OS for BI-EMD patients despite the majority of patients receiving novel anti-myeloma agents (PI, IMid, anti-CD38 antibodies).

Progression-free survival analysis was however limited due to the relatively short duration of follow-up in this study. Evidently, investigation of novel agents is required to improve outcomes in EMD. A recent early-phase trial of mezigdomide, a cereblon E3 ligase modulator, in a heavily pretreated group of MM patients showed an overall response rate of 30% in a subgroup with EMD.¹² The activity of regimens with novel therapeutic agents including peptide-drug conjugates such as melflufen¹³ similarly need to be evaluated in myeloma patients with BI-EMD, particularly earlier in the disease course.

In summary, despite treatment advancements in MM¹⁴, primary BI-EMD remains a poor prognostic feature conferring inferior overall treatment response and overall survival. Data from this single-centre study suggests PET CT scans allow the identification of BI-EMD thereby enabling the possibility of intensifying treatment. More studies including translational studies and clinical trials are needed to elucidate the unique biology of BI-EMD and in developing targeted therapies for this group of MM patients.

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Characteristics	BI-EMD at diagnosis N=33	Without BI-EMD at diagnosis N=873	OR (95% CI)	p- value		
	N (%)	N (%)				
Age (years)						
<70	30 (90.9%)	738 (84.5%)	Reference			
70+	3 (9.1%)	135 (15.5%)	0.55 (0.16 - 1.82)	0.3		
Sex						
Male	20 (60.6%)	522 (59.8%)	Reference			
Female	13 (39.4%)	351 (40.2%)	0.97 (0.47 - 1.97)	0.9		
Ethnicity						
White	11 (50.0%)	443 (68.3%)	Reference			
Asian	4 (18.2%)	57 (8.8%)	2.83 (0.87 - 9.17)			
Black	5 (22.7%)	98 (15.1%)	2.05 (0.70 - 6.05)			
Chinese	0	4 (0.6%)	N/A	0.0*		
Mixed	0	12 (1.8%)	N/A	0.3"		
Other	2 (9.1%)	35 (5.4%)	2.30 (0.49 -			
	_ (*** / *)		10.79)			
Missing	11	224				
LDH (Units/L)						
<214	5 (45.5%)	172 (68.8%)	Reference			
>214	6 (54.5%)	78 (31.2%)	2.65 (0.78 - 8.93)	0.1		
Missing	22	623				
Haemoglobin (g/dL)						
<10	4 (26.7%)	160 (38.3%)	Reference			
10+	11 (73.3%)	258 (61.7%)	1.71 (0.53 - 5.45)	0.4		
Missing	18	455				
Adjusted Calcium (mmol/L)						
<2.6	7 (63.6%)	295 (67.0%)	Reference			
>2.6	4 (36.4%)	145 (33.0%)	1.16 (0.33 - 4.04)	0.8		
Missing	22	433				
Disease isotype						
IgG	17 (60.7%)	493 (59.9%)	Reference			
IgA	5 (17.9%)	133 (16.2%)	1.09 (0.39 - 3.01)			
LC	5 (17.9%)	194 (23.6%)	0.75 (0.27 - 2.05)	0.2*		
Other	1 (3.6%)	3 (0.4%)	9.67 (0.96 -	0.2		
N 41	-	50	97.80)			
Missing	5	50				
155	F (07.00()	005 (40.00()	Deference			
1	5 (27.8%) 7 (20.0%)	235 (42.9%)				
	7 (38.9%)	181 (33.0%)	1.82 (0.57 - 5.82)	0.4*		
	6 (33.3%)	132 (24.1%)	2.14 (0.64 - 7.13)			
Missing	15	325				
	7 (31.8%)	111 (20.0%)	Reference			
del(17p)	5 (22.7%)	41 (7.4%)				

	t(4;14)	3 (13.6%)	56 (10.1%)			
	t(14;16)	0	18 (3.2%)			
	Standard risk	15 (68.2%)	444 (80.0%)	1.87 (0.74 - 4.69)	0.2	
	Missing	11	318			
First line therapy						
	PI	6 (18.2%)	227 (26.1%)	Reference		
	IMiD	5 (15.2%)	80 (9.2%)	2.36 (0.70 - 7.96)		
	PI & IMiD	13 (39.4%)	307 (35.3%)	1.60 (0.60 - 4.28)		
	CC	0	25 (2.9%)	N/A		
	Anti-CD38 antibody + Pl/IMiD	4 (12.1%)	88 (10.1%)	1.72 (0.47 - 6.24)	0.7*	
	Immunotherapy other	0	0	1.32 (0.40 - 4.41)		
	Other	5 (15.2%)	143 (16.4%)	0.03 (0.01 - 0.06)		
	Missing	0	3			
Response to first line						
th	erapy			5 /		
	SD	4 (16.7%)	24 (4.2%)	Reference		
	PR	14 (58.3%)	371 (65.7%)	0.23 (0.07 - 0.74)		
	VGPR	4 (16.7%)	135 (23.9%)	0.18 (0.04 - 0.76)	0.06*	
	CR	2 (8.3%)	35 (6.2%)	0.34 (0.06 - 2.02)		
	Missing	9	308			
	Overall Response	83.3% (95% CI 62 6-95 3)	95.8% (95% CI 93 7-97 3)		0.02	
SRE at diagnosis						
	No	13 (39.4%)	484 (55.9%)	Reference		
	Yes	20 (60.6%)	382 (44.1%)	1.95 (0.96 - 3.97)	0.07	
	Missing	0	7			

Table 1. Patient characteristics with BI-EMD and without BI-EMD at diagnosis (P-values calculated using logistic regression).

First line therapy regimens: Proteasome Inhibitor PI based (Bortezomib Adriamycin and Dexamethasone; PAD); Immunomodulatory agents, IMiD based (Cyclophosphamide, Thalidomide and Dexamethasone, CTD; Melphalan, Prednisolone and Thalidomidde, MPT; Melphalan Prednisolone and Lenalidomide, MPR; Carfilzomib Lenalidomide and Dexamethasone, CRD; Dexamethasone Thalidomide Cisplatin Adriamycin Cyclophosphamide Etoposide, DT-PACE); PI & IMiD (Ixazomib Lenalidomide and Dexamethasone, IRD; Bortezomib Thalidomide and Dexamethasone, VTD); conventional chemotherapy (CC) (Etoposide methylprednisolone high dose Cytarabine cisplatin, ESHAP; Vincristine Adriamycin and Dexamethasone, VAD; Cyclophosphamide Vincristine Doxorubicin and methylprednisolone, C-VAMP); Anti-CD38 antibody +PI/IMiD (Isatuximab Pomalidomide Dexamethasone, Daratumumab-VTD); Immunotherapy other (teclistamab, talquetamab, CART, venetoclax, Daratumumab single agent).

SD Stable disease, PR Partial Response, VGPR Very Good Partial Response, CR Complete Response. LC Light Chain, SRE Skeletal-Related Event.

P-values calculated using Fisher's exact test (*).

Figure 1. Survival Outcomes of myeloma patients with Bone-Independent Extramedullary Disease (BI-EMD) compared with those without BI-EMD. (A) Overall Survival (OS). (B) Progression-Free Survival (PFS). HR: Hazard ratio, 95% CI: 95% Confidence Interval.



