

1 **TITLE** Bendamustine plus rituximab for the treatment of Waldenström Macroglobulinaemia:
2 patient outcomes and impact of bendamustine dosing

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7

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55 **ABSTRACT**

56

57 Bendamustine and rituximab (BR) therapy is commonly used in the treatment of Waldenström
58 Macroglobulinaemia (WM). The impact dose of Bendamustine dose on response and survival
59 outcomes is not well established, and the impact of its use in different treatment settings is not clear.
60 We aimed to report response rates and survival outcomes following BR, and clarify the impact of
61 depth of response and bendamustine dose on survival.

62 A total of 250 WM patients treated with BR in the frontline or relapsed settings were included in this
63 multicentre, retrospective cohort analysis.

64 Rates of partial response (PR) or better differed significantly between the frontline and relapsed
65 cohorts (91.4% vs 73.9%, respectively; $p < 0.001$). Depth of response impacted survival outcomes:
66 two-year predicted PFS rates after achieving CR/VGPR vs PR were 96% vs 82%, respectively ($p = 0.002$).

67 Total bendamustine dose was predictive of PFS: in the frontline setting, PFS was superior in the group
68 receiving $\geq 1000 \text{mg/m}^2$ compared with those receiving $800\text{-}999 \text{mg/m}^2$ ($p = 0.04$). In the relapsed
69 cohort, those who received doses of $< 600 \text{mg/m}^2$ had poorer PFS outcomes compared with those
70 who received $\geq 600 \text{mg/m}^2$ ($p = 0.02$).

71 Attaining CR/VGPR following BR results in superior survival, and total bendamustine dose
72 significantly impacts response and survival outcomes, in both frontline and relapsed settings.

73 **INTRODUCTION**

74 Waldenström macroglobulinaemia (WM) is an indolent lymphoma characterised by the infiltration
75 of tissues (bone marrow, lymph nodes and/or spleen) with clonal lymphoplasmacytic cells and
76 consequent monoclonal IgM paraprotein production¹. With a median age at diagnosis in the seventh
77 decade, patients' comorbidities and performance status become key considerations in treatment
78 choices.

79

80 Bendamustine is a cytotoxic agent with structural similarities to both alkylating agents and purine
81 analogues and displays non-cross-resistance with other alkylators². In combination with rituximab, it
82 is a common choice in the treatment of WM. International guidelines recommend its use in both
83 frontline and relapsed settings^{3, 4}, due to its efficacy and relatively favourable toxicity profile⁵.
84 Response and survival outcomes of bendamustine/rituximab (BR) appear superior compared with
85 rituximab monotherapy⁶, R-CHOP⁷ (rituximab, cyclophosphamide, doxorubicin, vincristine and
86 prednisone) and DRC (dexamethasone, rituximab and cyclophosphamide)⁸⁻¹⁰, although definitive
87 randomised data is limited for comparisons with regimens other than R-CHOP. BR is considered to
88 be especially useful (among the chemoimmunotherapeutic options) in patients in need of rapid
89 disease control, or with bulky nodal or extranodal disease^{11, 12}. As per international consensus
90 recommendations, the recommended dose of bendamustine (in combination with rituximab) in
91 indolent non-Hodgkin lymphoma is 90mg/m² on days 1 and 2 every 4 weeks for 6 cycles in the
92 frontline setting, and 70-90mg/m² on days 1 and 2 every 4 weeks for 4 to 6 cycles in the
93 relapsed/refractory setting¹³.

94

95 Despite its frequent use in WM, questions remain regarding the best use of the BR regimen, including
96 the benefit of achieving deeper responses for improved survival outcome, the optimal bendamustine

97 dose to maximise survival outcome while minimising toxicity, and its use in the elderly population.
98 Herein we report the response and survival outcomes of the largest published real-world experience
99 of WM patients following BR therapy, and identify the implications of dose for maximising favourable
100 outcomes.

101 **METHODS**

102 ***Study Design and Participants***

103 This analysis included unselected, consecutively treated patients with a confirmed diagnosis of WM
104 according to the Second International Workshop on Waldenström's Macroglobulinaemia (IWWM)
105 criteria¹⁴ who received bendamustine with rituximab between September 2010 and May 2020 in
106 frontline or relapsed settings. Data were collected from 17 sites across four countries (Table S1).

107

108 The following baseline clinical and biological parameters¹⁵ were retrospectively collected from the
109 time of treatment commencement: blood counts, cross-sectional imaging (for the presence of
110 adenopathy, splenomegaly and extranodal disease), bone marrow histology, serum protein
111 electrophoresis, total immunoglobulin levels, and Eastern Cooperative Oncology Group performance
112 score (ECOG score). Treatment data collected were: number and types of prior therapies for
113 previously treated patients, year of treatment commencement, center of treatment, number of
114 bendamustine cycles, total bendamustine dose received (in mg/m²), number of rituximab doses,
115 dose and cycle reductions due to toxicity, and use of granulocyte colony stimulating factor (GCSF).
116 The impact of the following prior therapies on depth of response were assessed: rituximab, purine
117 analogues, Bruton tyrosine kinase inhibitors (BTKi) and autologous stem cell transplant (ASCT) (Table
118 S2).

119

120 ***Outcome Measures***

121 The primary outcomes were best response, progression free survival (PFS) and overall survival (OS).
122 Depth of response was graded using the modified IWWM-6 response criteria¹⁶, with response in IgM
123 level measured at 4-6 months following final chemotherapy dose. Major response rate (MRR)
124 included patients who had achieved partial response (PR), very good partial response (VGPR) or

125 complete response (CR). Overall response rate (ORR) also included those who achieved minor
126 response (MR). PFS was defined as the time from commencement of cycle 1 of BR treatment to the
127 earliest event of disease progression (by IWWM criteria¹⁶), or commencement of next treatment, or
128 death due to disease or treatment. Patients who did not have documented disease progression at
129 the time of data collection were censored on the date of their last recorded hospital contact. OS was
130 defined as the time from commencement of cycle 1 of BR to death from any cause, with living
131 patients censored at the time of last recorded hospital contact.

132 The primary outcomes were also assessed based on total bendamustine dose and number of
133 rituximab doses received. As per recommendations for bendamustine dose of 70-90mg/m² on days
134 1 and 2 for 4-6 cycles¹³ (i.e. total bendamustine doses of 560mg/m², 720mg/m², 840mg/m² and
135 1080mg/m², respectively), total bendamustine dose was categorised, for more direct clinical
136 application, into the following dose categories: <600mg/m², 600-799mg/m², 800-999mg/m² and
137 ≥1000mg/m². Rates of toxicity related bendamustine (dose/cycle) reduction and GCSF use were
138 assessed. Bendamustine starting dose was at the discretion of the treating physician.

139

140 ***Statistical Analysis***

141 Survival analysis was undertaken using the Kaplan Meier method,¹⁷ with survival distributions
142 compared using log-rank testing. Associations between baseline independent and outcome variables
143 were assessed with the Chi-square and Fisher's exact tests for categorical variables, and Wilcoxon-
144 Mann-Whitney test and Kruskal-Wallis ANOVA for numerical variables, as appropriate. Univariable
145 and multivariable binary logistic regressions were performed for attainment of CR/VGPR and for
146 toxicity related bendamustine reduction. Predictors of progression were identified with univariable
147 and multivariable Cox proportional hazard models, stratified according to treatment setting

148 (frontline vs relapsed). Statistical analyses were performed using IBM SPSS Statistics software,
149 version 27.

150 RESULTS

151 *Patient and Disease Characteristics*

152 A total of 250 patients with WM were treated with BR; 139 patients (55.6%) were treated in the
153 frontline setting, and 111 patients (44.4%) had received one or more prior therapies for WM before
154 receiving BR for relapsed disease (none of this cohort were treated for refractory disease). Baseline
155 characteristics are shown in Table 1. Seven patients with non-IgM-secreting lymphoplasmacytic
156 lymphoma were included in the analysis: five patients had an IgG paraprotein (two treated in the
157 frontline setting and three treated in relapsed setting), and two patients had an IgA paraprotein (one
158 patient treated in each of frontline and relapsed settings). Frontline and relapsed cohorts were
159 similar in terms of sex, age at the commencement of BR, ECOG score, and the following baseline
160 parameters: haemoglobin, bone marrow infiltration with LPL, and presence of adenopathy,
161 splenomegaly and extranodal disease.

162

163 *Depth of Response*

164 Overall, 209 patients (83.6%) achieved a major response and 229 patients (91.6%) achieved an
165 objective response. Three patients (1.2%) died of progressive disease during treatment, before
166 response assessment could be undertaken.

167 Depth of response was significantly superior in frontline vs relapsed cohorts (Table 2): CR/VGPR was
168 achieved in 66 patients (47.4%) vs 27 patients (24.3%), respectively ($p < 0.001$); major responses were
169 seen in 127 patients (91.4%) vs 82 patients (73.9%), respectively ($p < 0.001$); overall responses were
170 obtained by 136 patients (97.8%) vs 93 patients (83.8%), respectively ($p < 0.001$).

171 Depth of response did not differ within the relapsed cohort based on number of prior lines of therapy
172 (Figure S1A): major responses were seen in: 73.6% of patients (39/53) who had received one prior
173 line, 71.4% (20/28) who had received two prior lines, and 76.7% (23/30) who had received ≥ 3 prior

174 lines ($p=0.9$), with CR/VGPR achieved in 26.4% (14/53), 28.6% (8/28) and 16.6% (5/30), respectively
175 ($p=0.5$). The type of prior therapy also did not impact rates of CR/VGPR, major response or objective
176 response (Figure S1B). Depth of response was unaffected by year of treatment commencement and
177 centre of treatment.

178

179 On univariable analysis, age, sex, total bendamustine dose, and number of Rituximab doses
180 significantly impacted upon depth of response. Of 203 patients with a baseline ECOG score of 0 or 1,
181 176 (86.7%) achieved a major response, compared with 33 (70.2%) of the 47 patients with a baseline
182 ECOG score of ≥ 2 ($p=0.006$). In the frontline cohort, patients aged <70 years achieved higher rates of
183 CR/VGPR (45/75, 60%) than subjects aged ≥ 70 years (21/64, 32.8%; $p=0.001$). Multivariable binary
184 logistic regression – adjusted for sex, ECOG score, bendamustine and rituximab doses, haemoglobin,
185 platelet count, bone marrow infiltration, and extranodal disease – demonstrated older age,
186 treatment in the relapsed setting, and higher baseline paraprotein to be the only significant
187 predictors of non-attainment of CR/VGPR (Table S3).

188

189 ***Survival***

190 At a median follow up of 37 months, disease progression had occurred in 25 patients (18.0%) treated
191 in the frontline setting and 48 patients (43.2%) in the relapsed cohort ($p=0.008$). Death due to all
192 causes had occurred in 16 frontline patients (11.5%) and 40 relapsed patients (36.0%; $p<0.001$).

193 In the frontline cohort, the median OS and PFS were not reached; two-year and five-year predicted
194 OS/PFS rates were 94%/89% and 77%/60%, respectively (Figures 1A and 1D). In the relapsed cohort,
195 median OS was 58 months and median PFS was 50 months, with two-year and five-year predicted
196 OS/PFS rates of 80%/67% and 43%/42%, respectively (frontline vs relapsed OS: HR 2.8, $p=0.001$;
197 frontline vs relapsed PFS: HR 2.43, $p<0.001$).

198

199 The type of prior therapy did not impact on PFS or OS, although there was a trend towards shorter
200 PFS in those who had prior rituximab therapy compared with rituximab naïve patients ($p=0.087$).

201

202 Depth of response was an important predictor of both OS and PFS. As there was no significant survival
203 difference between the CR and VGPR groups, these groups were analysed together. Likewise, there
204 was no significant survival difference between the groups who did not achieve an overall response
205 (stable disease [SD], progressive disease [PD] and subjects who died before response was
206 assessable); these groups were therefore analysed together. Two-year predicted PFS rates were 96%
207 in those achieving CR/VGPR, 82% in those achieving PR, and 49% in those achieving MR (CR/VGPR vs
208 PR, $p=0.002$); five-year predicted PFS rates were 71%, 48% and 31% in the CR/VGPR-, PR-, and MR-
209 attaining cohorts, respectively. Median PFS was 53 months in the PR cohort and was not reached in
210 the CR/VGPR cohort. Median OS was 83 months after achieving CR/VGPR, 65 months after PR/MR
211 and 28 months after SD/PD ($p<0.001$). These differences were maintained when frontline and
212 relapsed cohorts were analysed separately (Figures 1B, 1E, 1F). An ECOG score of ≥ 2 was associated
213 with worse OS (Figure 1C) and PFS (Figure 1G), with similar differences observed when frontline and
214 relapsed cohorts were analysed separately. There was no PFS or OS difference between ECOG scores
215 of 0 and 1.

216

217 Cox proportional hazards regression models for PFS are shown in Table 3. Univariable analysis
218 demonstrated no impact on PFS of age, sex, haemoglobin, platelet count, paraprotein level, bone
219 marrow infiltration level, or presence of extranodal disease. Factors that significantly impacted on
220 PFS on univariable analysis – ECOG score, depth of response, total bendamustine dose, and number
221 of rituximab doses – as well as age (due to its potential impact on bendamustine dose), were included

222 in a multivariable model. When these variables were adjusted for, ECOG score of ≥ 2 , achievement of
223 PR or less, and total Bendamustine dose of $< 1000\text{mg}/\text{m}^2$ (see below) were all independently
224 associated with poorer PFS in the frontline setting, but number of Rituximab doses was not. In the
225 relapsed setting, ECOG score of ≥ 2 , achievement of PR or less, and receiving ≤ 3 doses of rituximab
226 were all independently associated with poorer PFS.

227

228 ***Impact of Bendamustine Dose on Outcomes***

229 Starting bendamustine doses were similar between frontline and relapsed cohorts, with 78.4% and
230 75.7% of patients, respectively, commencing treatment at a dose of $\geq 90\text{mg}/\text{m}^2$ on days 1 and 2
231 ($p=0.6$). There was significant variation in bendamustine starting dose choice between centres noted
232 in the frontline cohort; starting dose was independently affected by age and ECOG score. Patients in
233 the frontline cohort received higher total bendamustine doses than those in the relapsed setting,
234 due to higher rates of cycle truncation and dose reduction in the relapsed setting (median total
235 bendamustine dose $1080\text{mg}/\text{m}^2$ vs $720\text{mg}/\text{m}^2$, $p<0.001$; Table 1).

236

237 Total bendamustine dose received, stratified into dose categories (see Methods), significantly
238 impacted on MRR as well as PFS. In the frontline setting, MRR was highest in the top dose category:
239 80/81 patients (98.8%) who received $\geq 1000\text{mg}/\text{m}^2$ achieved a major response, compared with 27/33
240 patients (81.8%) who received $800\text{-}999\text{mg}/\text{m}^2$, and 20/25 patients (80%) who received $< 800\text{mg}/\text{m}^2$
241 ($p=0.001$). CR/VGPR rates in the aforementioned three dose categories were 53.1% (43/81), 45.5%
242 (15/33) and 32% (8/25), respectively ($p=0.17$). PFS was significantly longer in patients who received
243 $\geq 1000\text{mg}/\text{m}^2$ compared with those receiving smaller doses (Figure 1H), including when adjusted for
244 age, ECOG score and depth of response (Table 3). In the relapsed cohort, there were no significant
245 differences in response (MRR, ORR or CR/VGPR rates) between the largest 3 dose categories.

246 Similarly, there was no appreciable PFS difference based on total bendamustine dose if $\geq 600\text{mg}/\text{m}^2$
247 was received (Figure 11). Those who received total doses of $< 600\text{mg}/\text{m}^2$ (i.e. $70\text{mg}/\text{m}^2$ on days 1&2
248 for 4 cycles, or less) had significantly poorer PFS compared with those who received $\geq 600\text{mg}/\text{m}^2$,
249 with two-year predicted PFS rates of 46% and 78%, respectively ($p=0.004$).

250

251 **Toxicity**

252 Twenty-four frontline patients (17.3%) had toxicity related bendamustine reduction (both dose
253 reductions and cycle truncation) compared with 39 relapsed patients (35.1%; $p<0.001$);
254 myelosuppression accounted for most of the toxicity related reductions in treatment. Of the 109
255 frontline patients who commenced treatment at a dose of $\geq 90\text{mg}/\text{m}^2$, 20 (18.3%) subsequently
256 underwent bendamustine reduction due to toxicity, compared with 26/84 relapsed patients (31%;
257 $p=0.04$). Multivariable binary logistic regression analyses for toxicity related bendamustine dose
258 reduction were performed in both frontline and relapsed settings, and included age, ECOG score, and
259 bendamustine starting dose (≥ 90 or $< 90\text{mg}/\text{m}^2$). The rate of toxicity related dose reduction was
260 affected only by ECOG score in the frontline setting (ECOG score 0-1 vs ECOG score ≥ 2 : OR 3.63, 95%
261 CI 1.37-9.65, $p=0.01$), and not by age or starting dose. None of the variables affected rates of
262 bendamustine reduction in the relapsed setting. Rates of GCSF use did not differ between frontline
263 vs relapsed cohorts (30.9% vs 36.9%; $p=0.34$); rates of GCSF use also did not differ based on starting
264 doses (32.3% for $\geq 90\text{mg}/\text{m}^2$ vs 38.6% for $< 90\text{mg}/\text{m}^2$, $p=0.43$).

265

266 Older subjects received lower total bendamustine doses. In the frontline cohort, median total
267 bendamustine dose was $1080\text{mg}/\text{m}^2$ among subjects < 70 years of age ($n=75$) and $990\text{mg}/\text{m}^2$ in the
268 ≥ 70 -year group ($n=64$) ($p=0.051$). Of the 17 frontline patients aged ≥ 80 years, 13 (76.5%) received
269 total bendamustine doses of $\geq 720\text{mg}/\text{m}^2$ and 10 (58.5%) received total doses of $1080\text{mg}/\text{m}^2$. In the

270 relapsed cohort, median bendamustine dose received by subjects aged <70 years (n=59) was
271 840mg/m² compared with 585mg/m² for those aged ≥70 years (n=52; p=0.024).

272

273 Rates of secondary malignancies were assessed. Two patients (0.8%) developed therapy-related
274 myeloid neoplasms (t-MN), diagnosed at three and six years, respectively, after receiving BR; both
275 had also received Fludarabine prior to being diagnosed with T-MN (and, in one case, prior to receiving
276 BR). Rates of new solid tumour diagnoses were comparable pre- and post-BR (4.8% vs 3.6%).

277 **DISCUSSION**

278 Herein we report real-world experience of BR in the treatment of WM in both frontline and relapsed
279 settings, in the largest such series published to date. This analysis reflects the experience of academic
280 institutions as well as secondary care hospitals (Table S1). Prior to this analysis, evidence for the use
281 of BR in WM had been largely obtained from small retrospective series (Table S4). We demonstrate
282 excellent outcomes in unselected patients with WM treated with BR and address outstanding
283 questions regarding the best use of this regimen. Our cohort included elderly patients as well as
284 heavily pre-treated patients (12% of the total cohort received BR after 3 or more prior lines of
285 therapy). A slightly larger number of patients were treated with BR in the frontline setting (55.6% of
286 total cohort), and this cohort demonstrated superior response rates, longer PFS, and improved
287 tolerability of BR compared with patients treated in the relapsed setting. PFS was found to be
288 dependent on both depth of response achieved and total bendamustine dose received.

289
290 The benefit of achieving deeper responses has not always been clear¹⁸, with a previous retrospective
291 series showing PFS benefit in achieving CR/VGPR following rituximab-based therapy¹⁹ and another
292 series showing no PFS benefit in achieving CR/VGPR following BR²⁰. This analysis demonstrated a
293 clear survival benefit with deeper responses, with the achievement of CR/VGPR being associated with
294 longer PFS and OS in both frontline and relapsed settings. While the CR- and VGPR-attaining groups
295 were analysed together in the survival analyses (as there was no significant survival difference
296 between these groups), it is possible that differences in survival outcome between the groups could
297 emerge with longer follow up. With attainment of deeper responses with BR, the resulting extension
298 of the treatment free interval could minimise the cumulative burden of treatment toxicity for an
299 individual. We therefore conclude that depth of response is an important treatment goal with BR
300 therapy.

301

302 A previous evaluation of frontline patients showed that prospective dose reduction of bendamustine
303 did not adversely affect the attainment of major response²¹. Within the limitations of its
304 retrospective non-randomised context, this study more clearly delineates the bendamustine doses
305 associated with superior response and PFS outcomes. In the frontline cohort, treatment with 6 cycles
306 of 90mg/m² on days 1 and 2 (i.e. total bendamustine dose of $\geq 1000\text{mg/m}^2$) appeared to produce
307 superior PFS than lower bendamustine doses, even when adjusted for patient age and fitness (i.e.
308 ECOG score); this finding therefore supports a starting dose of 90mg/m² on days 1 and 2 for all
309 frontline patients where possible, aiming to administer a total of 6 cycles. Conversely, in the relapsed
310 cohort, no additional benefit is gained either in response rates or in PFS when a total dose of
311 $>600\text{mg/m}^2$ was used, suggesting that 4 cycles of 90mg/m² of bendamustine on days 1 and 2 may
312 be sufficient in the relapsed cohort; in cases where this starting dose is not expected to be tolerated,
313 a starting dose of 70mg/m² on days 1 and 2 may be sufficient provided 5-6 cycles are administered.
314 It is important to note that Bendamustine start dose and dose reductions were at the discretion of
315 individual clinicians, with dose choices being made in accordance with available international
316 consensus guidelines.

317

318 Prior studies report between 34 and 53% of patients were not able to receive the intended 6 cycles
319 of treatment, with myelosuppression/haematologic toxicity being the most common reason for
320 treatment truncation^{9, 20, 22, 23}. In this study, only 25% of patients overall required reductions in
321 bendamustine due to toxicity, with treatment in the relapsed setting and a baseline ECOG score of
322 ≥ 2 in the frontline setting predicting for higher rates of bendamustine reduction. Although the
323 starting dose ($\geq 90\text{mg/m}^2$ vs $<90\text{mg/m}^2$ on days 1 and 2 of each cycle) was at the clinician's discretion,
324 it had no appreciable effect on the rates of toxicity related bendamustine dose reduction in both

325 treatment settings. Within the median follow up time of approximately 3 years in this study, rates of
326 secondary malignancy were low following BR therapy, although longer follow up may reveal higher
327 rates of t-MN.

328

329 This study did not evaluate time to best response as the BR regimen is already known to induce later
330 responses: progressive decline in IgM is seen for some months following treatment completion²⁰,
331 and the cumulative incidence of objective response increases for up to 18 months after treatment
332 initiation^{22, 24}. The impacts of MYD88 and CCXCR4 mutations were not assessed in this study
333 specifically; the presence of these mutations has previously been shown to not impact on disease
334 response or progression free survival outcomes following BR^{9, 22}. This study also did not assess the
335 impact of ISSWM due to lack of available biological data.

336

337 In current clinical practice, the choice of bendamustine therapy and dose needs to be considered in
338 conjunction with potential risks of the SARS-CoV-2 pandemic: studies have associated a total
339 bendamustine dose of ≥ 1080 mg/m² with delayed CD4 recovery and prolonged CD4 lymphopenia
340 identified as a risk factor for serious infection complications during follow-up after treatment²⁵.

341

342 This study presents robust retrospective evidence that the BR combination, with its excellent
343 response rates, long PFS intervals and favourable toxicity profile particularly in the frontline setting,
344 retains an important role in the treatment of WM. Additionally, BR continues to be useful in the
345 treatment of relapsed disease, with evidence from the current study that good responses are
346 achievable even in the extensively pre-treated cohort and irrespective of type of prior therapy.
347 Regarding the use of BR in the present era of increasing availability of Bruton tyrosine kinase
348 inhibitors (BTKi), the BR combination reserves an important role for patients for whom limited

349 treatment duration would be preferred over indefinite therapy, or for patients for whom BTKi are
350 contraindicated; the PFS in frontline patients in this study is indeed comparable to the PFS seen in
351 treatment-naïve patients on Ibrutinib monotherapy²⁶. In both frontline and relapsed settings,
352 attaining CR/VGPR results in superior PFS and OS. Total bendamustine dose significantly impacts
353 response and survival outcomes in both frontline and relapsed settings.

354

355 **AUTHOR CONTRIBUTIONS**

356 SA designed the research study, collected data, performed statistical analysis and drafted the paper;
357 DB, HG, AC, TM, RK, AON, JMIV, GP, DT, KM, KM, CA, MG, CK, MJK, KL and GF collected data;
358 DES, DT, HM and SDS reviewed the manuscript; EK performed statistical analysis; MJB contributed to
359 analysis design and critically revised the manuscript; ADW designed the research study and critically
360 revised the manuscript.

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Table 1: Baseline patient characteristics

Parameter	All patients (n=250)	Frontline (n=139)	Relapsed (n=111)	P
Median age at start of BR (IQR), years	69.0 (62.1-76.0)	69.3 (60.4-76.0)	69.0 (65.2-76.2)	0.25
Male, No. (%)	167 (66.8)	95 (68.3)	72 (64.9)	0.56
ECOG performance score:				0.13
0, No. (%)	80 (32.0)	48 (34.5)	32 (28.8)	
1, No. (%)	123 (49.2)	70 (50.4)	53 (47.7)	
≥2, No. (%)	47 (18.8)	21 (15.1)	26 (23.4)	
Median time, diagnosis to BR start, months	15.6	2.1	76.1	<0.001*
Year of BR commencement:				<0.001*
2010-2015, No. (%)	85 (34.0)	32 (23)	53 (47.8)	
2016-2017, No. (%)	89 (35.6)	53 (38.1)	36 (32.4)	
2018-2019, No. (%)	76 (30.4)	54 (38.9)	22 (19.8)	
Haemoglobin ≤110g/L, No. (%)	171 (68.4%)	98 (70.5%)	73 (65.8)	0.47
Platelet count ≥100 x10 ⁹ /L, No. (%)	41 (16.4%)	18 (12.9%)	23 (20.7%)	0.09
Median bone marrow infiltration with LPL, %	60	60	60	0.26
Median paraprotein, g/L	23.2	26.5	21	0.049*
Adenopathy present, No. (%)	103 (41.2)	56 (40.3)	47 (42.3)	0.70
Splenomegaly present, No. (%)	54 (21.6)	36 (25.9)	18 (16.4)	0.09*
Extranodal disease present, No. (%)	36 (14.4)	18 (12.9)	18 (16.4)	0.47
Total bendamustine dose, median (range), mg/m ²	900 (70-1200)	1080 (1040-1080)	720 (70-1200)	<0.001*
Total bendamustine dose:				<0.001*
≥1000mg/m ² , No. (%)	119 (47.6)	81 (58.3)	38 (34.2)	
800-999mg/m ² , No. (%)	47 (18.8)	33 (23.7)	14 (12.6)	
600-799mg/m ² , No. (%)	32 (12.8)	13 (9.4)	19 (17.1)	
<600mg/m ² , No. (%)	52 (20.8)	12 (8.6)	40 (36.0)	
Rituximab doses received:				0.002*
0-3, No. (%)	57 (22.8)	20 (14.4)	37 (33.3)	
4-5, No. (%)	61 (24.4)	39 (28.0)	22 (19.8)	
6 or more, No. (%)	132 (52.8)	80 (57.6)	52 (46.9)	

P values reflect differences between Frontline and Relapsed cohorts; IQR = interquartile range; BR = Bendamustine/Rituximab; * = statistical significance reached

Table 2: Response rates for patients with Waldenström Macroglobulinaemia treated with Bendamustine/Rituximab

Response	Total (n=250)	Frontline (n=139)	Relapsed (n=111)	P value
Major response rate, No. (%)	209 (83.6)	127 (91.4)	82 (73.9)	<0.001
Objective response rate, No. (%)	229 (91.6)	136 (97.8)	93 (83.8)	<0.001
Categorical response, No. (%)				
Complete	22 (8.8)	17 (12.2)	5 (4.5)	0.027
Very good partial	71 (28.4)	49 (35.3)	22 (19.8)	0.007
Partial	116 (46.4)	61 (43.9)	55 (49.5)	0.372
Minor	20 (8.0)	9 (6.5)	11 (9.9)	0.322
Stable disease, No. (%)	17 (6.8)	2 (1.4)	15 (13.5)	
Progressive disease, No. (%)	1 (0.4)	0	1 (0.9)	
Died before assessment, No. (%)	3 (1.2)	1 (0.7)	2 (1.8)	

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Table 3: Cox proportional hazard regression models exploring factors associated with higher risk of progression following bendamustine/rituximab					
<i>Frontline cohort</i>					
Variable	Category	Univariable		Multivariable	
		Hazard ratio (95% CI)	P value	Hazard ratio (95% CI)	P value
Sex	Female	Reference		-	-
	Male	2.03 (0.69-5.94)	0.19		
Age (years)*		1.03 (0.98-1.08)	0.16	1.01 (0.96-1.05)	0.77
ECOG score	0-1	Reference		Reference	
	2	3.55 (1.56-8.06)	0.03	2.73 (1.16-6.4)	0.02
Depth of response	CR/VGPR	Reference		Reference	
	Partial response or less	3.66 (1.37-9.78)	0.009	4.16 (1.45-11.95)	0.008
Bendamustine dose (mg/m ²)	Total dose*	0.997 (0.996-0.998)	<0.001	-	-
	≥1000	Reference		Reference	
	800-999	2.93 (0.98-8.75)	0.053	3.26 (1.05-10.17)	0.04
	600-799	9.36 (3.05-28.72)	<0.001	6.79 (1.78-25.94)	0.005
	<600	10.44 (3.13-34.81)	<0.001	8.54 (1.2-60.49)	0.03
Rituximab doses (No.)	≥6	Reference		Reference	
	4-5	2.24 (0.89-5.66)	0.09	1.04 (0.36-3.01)	0.94
	0-3	5.85 (2.13-16.03)	<0.001	1.34 (0.26-6.98)	0.73
Haemoglobin (g/dL)*		0.87 (0.72-1.06)	0.18	-	-
Platelet count (x10 ⁹ /L)*		0.996 (0.98-1.013)	0.65		
Paraprotein (g/dL)*		1.01 (0.84-1.22)	0.88	-	-
LPL % in bone marrow*		0.999 (0.98-1.01)	0.86	-	-
Extranodal disease	No	Reference		-	-
	Yes	1.347 (0.46-3.93)	0.58		
<i>Relapsed cohort</i>					
Variable	Category	Univariable		Multivariable	
		Hazard ratio (95% CI)	P value	Hazard ratio (95% CI)	P value
Sex	Female	Reference		-	-
	Male	1.25 (0.676-2.294)	0.48		
Age*		1.01 (0.977-1.037)	0.69	0.998 (0.97-1.03)	0.91
ECOG score	0-1	Reference		Reference	
	2	2.52 (1.391-4.556)	0.002	3.28 (1.67-6.44)	<0.001
Depth of response	CR/VGPR	Reference		Reference	
	Partial response or less	3.44 (1.458-8.141)	0.005	3.83 (1.58-9.26)	0.003
Bendamustine dose (mg/m ²)	Total dose*	0.999 (0.998-1.00)	0.004	-	-
	≥1000	Reference		Reference	
	800-999	0.47 (0.159-1.409)	0.18	0.31 (0.1-0.97)	0.04
	600-799	0.52 (0.193-1.404)	0.12	0.45 (0.15-1.34)	0.15
	<600	1.75 (0.926-3.304)	0.08	0.77 (0.29-2.05)	0.61
Rituximab doses (No.)	≥6	Reference		Reference	
	4-5	1.06 (0.46-2.43)	0.89	1.36 (0.56-3.27)	0.5
	0-3	2.05 (1.1-3.82)	0.02	2.82 (1.14-6.97)	0.02
Haemoglobin (g/dL)*		0.75 (0.64-0.88)	<0.001	-	-
Platelet count (x10 ⁹ /L)*		1.007 (0.984-1.032)	0.55		
Paraprotein (g/dL)*		0.96 (0.8-1.15)	0.65	-	-
LPL % in bone marrow*		1.004 (0.99-1.01)	0.43	-	-
Extranodal disease	No	Reference		-	-
	Yes	1.58 (0.78-3.17)	0.2		

*Hazards to survival are relative to a unit increase in continuous variable
CI = confidence interval

437 **FIGURE LEGEND**

438

Figure 1	<p>Kaplan-Meier estimates following BR therapy:</p> <ul style="list-style-type: none">(A) Overall survival (OS) according to number of prior therapies.(B) OS according to best response.(C) OS according to ECOG score pre-treatment.(D) Progression free survival (PFS) according to number of prior therapies.(E) PFS according to best response – frontline cohort.(F) PFS according to best response – relapsed cohort.(G) PFS according to ECOG score pre-treatment.(H) PFS according to total Bendamustine dose received – frontline cohort.(I) PFS according to total Bendamustine dose received – relapsed cohort. <p>CR = complete remission; VGPR = very good partial response; PR = partial response; MR = minor response; SD = stable disease; PD = progressive disease.</p>
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