

1 เอกสารกำกับยาภาษาอังกฤษ

2 **Summary of Product Characteristics**

3 **BORTOMA**

4 ^{Rx} **Bortezomib 3.5 mg Powder for Solution for Injection**

5 **1. Name of the medicinal product**

6 1.1 Product Name: BORTOMA

7 1.2 Strength: Bortezomib 3.5 mg

8 1.3 Pharmaceutical Dosage Form: Powder for Solution for Injection

9 **2. Qualitative and quantitative composition**

10 2.1 Qualitative declaration

11 Bortezomib is an antineoplastic agent

12 INN: Bortezomib.

13 2.2 Quantitative declaration

14 Each vial contains 3.5 mg bortezomib

15 For the full list of excipients, see section 6.1.

16 **3. Pharmaceutical form**

17 White to off-white lyophilized cake or powder

18 **4. Clinical particulars**

19 ([Ref.1: Velcade EU/1/04/274/001 page 1 - 20](#))

20 **4.1 Therapeutic indications**

21 ([Ref.1: Velcade EU/1/04/274/001 page 1](#))

22 BORTOMA as monotherapy or in combination with pegylated liposomal doxorubicin or dexamethasone is indicated
23 for the treatment of adult patients with progressive multiple myeloma who have received at least 1 prior therapy
24 and who have already undergone or are unsuitable for haematopoietic stem cell transplantation.

25 BORTOMA in combination with melphalan and prednisone is indicated for the treatment of adult patients with
26 previously untreated multiple myeloma who are not eligible for high -dose chemotherapy with haematopoietic stem
27 cell transplantation.

28 BORTOMA in combination with dexamethasone, or with dexamethasone and thalidomide, is indicated for the
29 induction treatment of adult patients with previously untreated multiple myeloma who are eligible for high -dose

30 chemotherapy with haematopoietic stem cell transplantation.
31 BORTOMA in combination with rituximab, cyclophosphamide, doxorubicin and prednisone is indicated for the
32 treatment of adult patients with previously untreated mantle cell lymphoma who are unsuitable for haematopoietic
33 stem cell transplantation.

34 **4.2 Posology and method of administration**

35 [\(Ref.1: Velcade EU/1/04/274/001 page 1 - 7\)](#)

36 BORTOMA treatment must be initiated under supervision of a physician experienced in the treatment of cancer
37 patients, however BORTOMA may be administered by a healthcare professional experienced in use of
38 chemotherapeutic agents .BORTOMA must be reconstituted by a healthcare professional (see section 6.6).

39 Posology for treatment of progressive multiple myeloma (patients who have received at least one prior therapy)

40 [\(Ref.1: Velcade EU/1/04/274/001 page 1\)](#)

41 *Monotherapy*

42 Bortezomib 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection at
43 the recommended dose of 1.3 mg/m² body surface area twice weekly for two weeks on days 1, 4, 8, and 11 in a
44 21-day treatment cycle .This 3-week period is considered a treatment cycle .It is recommended that patients
45 receive 2 cycles of Bortezomib following a confirmation of a complete response .It is also recommended that
46 responding patients who do not achieve a complete remission receive a total of 8 cycles of Bortezomib therapy .At
47 least 72 hours should elapse between consecutive doses of Bortezomib.

48 *Dose adjustments during treatment and re-initiation of treatment for monotherapy*

49 [\(Ref.1: Velcade EU/1/04/274/001 page 1 - 2\)](#)

50 Bortezomib treatment must be withheld at the onset of any Grade 3 non-haematological or any Grade 4
51 haematological toxicities, excluding neuropathy as discussed below. (see also section 4.4) Once the symptoms of
52 the toxicity have resolved, Bortezomib treatment may be re-initiated at a 25 %reduced dose, 1.3 mg/m² reduced
53 to 1.0 mg/m²; 1.0 mg/m² reduced to 0.7 mg/m² .(If the toxicity is not resolved or if it recurs at the lowest dose,
54 discontinuation of Bortezomib must be considered unless the benefit of treatment clearly outweighs the risk.

55 *Neuropathic pain and/or peripheral neuropathy*

56 [\(Ref.1: Velcade EU/1/04/274/001 page 1 - 2\)](#)

57 Patients who experience bortezomib-related neuropathic pain and/or peripheral neuropathy are to be managed as
58 presented in Table 1 (see section 4.4) Patients with pre-existing severe neuropathy may be treated with Bortezomib
59 only after careful risk/benefit assessment.

60 *Table 1: Recommended *posology modifications for bortezomib-related neuropathy*

Severity of neuropathy	Posology modification
Grade 1 (asymptomatic; loss of deep tendon reflexes or paresthesia) with no pain or loss of function	None
Grade 1 with pain or Grade 2 (moderate symptoms; limiting instrumental Activities of Daily Living (ADL)**)	Reduce Bortezomib to 1.0 mg/m ² or Change Bortezomib treatment schedule to 1.3 mg/m ² once per week
Grade 2 with pain or Grade 3 (severe symptoms; limiting self care ADL***)	Withhold Bortezomib treatment until symptoms of toxicity have resolved . When toxicity resolves re-initiate Bortezomib treatment and reduce dose to 0.7 mg/m ² once per week.
Grade 4 (life -threatening consequences; urgent intervention indicated) and /or severe autonomic neuropathy	Discontinue Bortezomib

61 *Based on posology modifications in Phase II and III multiple myeloma studies and post-marketing experience .
62 Grading based on NCI Common Toxicity Criteria CTCAE v 4.0.

63 ***Instrumental ADL* :refers to preparing meals, shopping for groceries or clothes, using telephone, managing money,
64 etc;

65 ****Self care ADL*: refers to bathing, dressing and undressing, feeding self, using the toilet, taking medicinal products,
66 and not bedridden.

67

68 *Combination therapy with pegylated liposomal doxorubicin*

69 ([Ref.1: Velcade EU/1/04/274/001 page 2](#))

70 Bortezomib 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection at
71 the recommended dose of 1.3 mg/m² body surface area twice weekly for two weeks on days 1, 4, 8, and 11 in a
72 21-day treatment cycle . This 3-week period is considered a treatment cycle . At least 72 hours should elapse
73 between consecutive doses of Bortezomib.

74 Pegylated liposomal doxorubicin is administered at 30 mg/m² on day 4 of the Bortezomib treatment cycle as a 1
75 hour intravenous infusion administered after the Bortezomib injection.

76 Up to 8 cycles of this combination therapy can be administered as long as patients have not progressed and
77 tolerate treatment . Patients achieving a complete response can continue treatment for at least 2 cycles after the
78 first evidence of complete response, even if this requires treatment for more than 8 cycles . Patients whose levels
79 of paraprotein continue to decrease after 8 cycles can also continue for as long as treatment is tolerated and they
80 continue to respond.

81 For additional information concerning pegylated liposomal doxorubicin, see the corresponding Summary of Product
82 Characteristics.

83 *Combination with dexamethasone*

84 [\(Ref.1: Velcade EU/1/04/274/001 page 2\)](#)

85 Bortezomib 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection at
86 the recommended dose of 1.3 mg/m² body surface area twice weekly for two weeks on days 1, 4, 8, and 11 in a
87 21 day treatment cycle .This 3-week period is considered a treatment cycle .At least 72 hours should elapse
88 between consecutive doses of Bortezomib.

89 Dexamethasone is administered orally at 20 mg on days 1, 2, 4, 5, 8, 9, 11, and 12 of the Bortezomib treatment
90 cycle.

91 Patients achieving a response or a stable disease after 4 cycles of this combination therapy can continue to receive
92 the same combination for a maximum of 4 additional cycles.

93 For additional information concerning dexamethasone, see the corresponding Summary of Product Characteristics.

94
95 *Dose adjustments for combination therapy for patients with progressive multiple myeloma*

96 For Bortezomib dosage adjustments for combination therapy follow dose modification guidelines described under
97 monotherapy above.

98
99 Posology for previously untreated multiple myeloma patients not eligible for haematopoietic stem cell transplantation

100 [\(Ref.1: Velcade EU/1/04/274/001 page 3 - 4\)](#)

101 *Combination therapy with melphalan and prednisone*

102 Bortezomib 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection in
103 combination with oral melphalan and oral prednisone as shown in Table 2 .A 6-week period is considered a
104 treatment cycle .In Cycles 1-4, Bortezomib is administered twice weekly on days 1, 4, 8, 11, 22, 25, 29 and 32 .In
105 Cycles 5-9, Bortezomib is administered once weekly on days 1, 8, 22 and 29. At least 72 hours should elapse
106 between consecutive doses of Bortezomib.

107 Melphalan and prednisone should both be given orally on days 1, 2, 3 and 4 of the first week of each Bortezomib
108 treatment cycle.

109 Nine treatment cycles of this combination therapy are administered.

110 *Table 2: Recommended posology for Bortezomib in combination with melphalan and prednisone*

Twice weekly Bortezomib (cycles 1 – 4)												
Week	1			2			3	4		5		6
Vc (1.3 mg/m ²)	Day 1	--	--	Day 4	Day 8	Day 11	rest period	Day 22	Day 25	Day 29	Day 32	rest period

M (9 mg/m ²) P (60 mg/m ²)	Day 1	Day 2	Day 3	Day 4	--	--	rest period	--	--	--	--	rest period
Once weekly Bortezomib (cycles 5 – 9)												
Week	1				2		3	4		5		6
Vc (1.3 mg/m ²)	Day 1	--	--	--	Day 8		rest period	Day 22		Day 29		rest period
M(9 mg/m ²) P (60 mg/m ²)	Day 1	Day 2	Day 3	Day 4	--	--	rest period	--	--	--	--	rest period

111 Vc =Bortezomib; M=melphalan, P=prednisone

112 Dose adjustments during treatment and re-initiation of treatment for combination therapy with melphalan and
113 prednisone.

114 Prior to initiating a new cycle of therapy:

115 • Platelet counts should be $\geq 70 \times 10^9/L$ and the absolute neutrophils count should be $\geq 1.0 \times 10^9/L$

116 • Non-haematological toxicities should have resolved to Grade 1 or baseline

117

118 *Table 3 :Posology modifications during subsequent cycles of Bortezomib therapy in combination with melphalan and*
119 *prednisone*

Toxicity	Posology modification or delay
<i>Haematological toxicity during a cycle</i>	Consider reduction of the melphalan dose by 25 % in the next cycle.
• If prolonged Grade 4 neutropenia or thrombocytopenia, or thrombocytopenia with bleeding is observed in the previous cycle	
• If platelet counts $\leq 30 \times 10^9/L$ or ANC $\leq 0.75 \times 10^9/L$ on a Bortezomib dosing day (other than day 1)	Bortezomib therapy should be withheld
• If several Bortezomib doses in a cycle are withheld (≥ 3 doses during twice weekly administration or ≥ 2) doses during weekly administration(Bortezomib dose should be reduced by 1 dose level (from 1.3 mg/m ² to 1 mg/m ² , or from 1 mg/m ² to 0.7 mg/m ²)
<i>Grade ≥ 3 non-haematological toxicities</i>	Bortezomib therapy should be withheld until symptoms of the toxicity have resolved to Grade 1 or baseline . Then, Bortezomib may be reinitiated with one dose level reduction (from 1.3 mg /m ² to 1 mg /m ² , or from 1 mg/m ²

	to 0.7 mg/m ²). For Bortezomib- related neuropathic pain and /or peripheral neuropathy, hold and /or modify Bortezomib as outlined in Table 1.
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120 For additional information concerning melphalan and prednisone, see the corresponding Summary of Product
121 Characteristics.

122

123 Posology for previously untreated multiple myeloma patients eligible for haematopoietic stem cell transplantation
124 (induction therapy)

125 [\(Ref.1: Velcade EU/1/04/274/001 page 4\)](#)

126 *Combination therapy with dexamethasone*

127 BORTEZOMIB 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection
128 at the recommended dose of 1.3 mg/m² body surface area twice weekly for two weeks on days 1, 4, 8, and 11 in
129 a 21-day treatment cycle . This 3-week period is considered a treatment cycle . At least 72 hours should elapse
130 between consecutive doses of Bortezomib.

131 Dexamethasone is administered orally at 40 mg on days 1, 2, 3, 4, 8, 9, 10 and 11 of the Bortezomib treatment
132 cycle

133 Four treatment cycles of this combination therapy are administered.

134

135 *Combination therapy with dexamethasone and thalidomide*

136 [\(Ref.1: Velcade EU/1/04/274/001 page 4 - 5\)](#)

137 BORTEZOMIB 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection
138 at the recommended dose of 1.3 mg/m² body surface area twice weekly for two weeks on days 1, 4, 8, and 11 in
139 a 28-day treatment cycle . This 4-week period is considered a treatment cycle . At least 72 hours should elapse
140 between consecutive doses of Bortezomib.

141 Dexamethasone is administered orally at 40 mg on days 1, 2, 3, 4, 8, 9, 10 and 11 of the Bortezomib treatment
142 cycle.

143 Thalidomide is administered orally at 50 mg daily on days 1-14 and if tolerated the dose is increased to 100 mg
144 on days 15-28, and thereafter may be further increased to 200 mg daily from cycle 2. (see Table 4)

145 Four treatment cycles of this combination are administered . It is recommended that patients with at least partial
146 response receive 2 additional cycles.

147

148 *Table 4 :Posology for Bortezomib combination therapy for patients with previously untreated multiple myeloma eligible*
 149 *for haematopoietic stem cell transplantation.*

Vc +Dx	Cycles 1 to 4				
	Week	1	2	3	
	Vc (1.3 mg/m ²)	Day 1, 4	Day 8, 11	Rest Period	
	Dx 40 mg	Day 1, 2, 3, 4	Day 8, 9, 10, 11	-	
Vc+Dx+T	Cycle 1				
	Week	1	2	3	4
	Vc (1.3 mg/m ²)	Day 1, 4	Day 8, 11	Rest Period	Rest Period
	T 50 mg	Daily	Daily	-	-
	T 100 mg ^a	-	-	Daily	Daily
	Dx 40 mg	Day 1, 2, 3, 4	Day 8, 9, 10, 11	-	-
	Cycles 2 to 4^b				
	Vc (1.3 mg/m ²)	Day 1, 4	Day 8, 11	Rest Period	Rest Period
	T 200 mg ^a	Daily	Daily	Daily	Daily
	Dx 40 mg	Day 1, 2, 3, 4	Day 8, 9, 10, 11	-	-

150 Vc =Bortezomib ; Dx=dexamethasone; T=thalidomide

151 ^a Thalidomide dose is increased to 100 mg from week 3 of Cycle 1 only if 50 mg is tolerated and to 200 mg from
 152 cycle 2 onwards if 100 mg is tolerated.

153 ^b Up to 6 cycles may be given to patients who achieve at least a partial response after 4 cycles

154 Dosage adjustments for transplant eligible patients

155 For Bortezomib dosage adjustments, dose modification guidelines described for monotherapy should be followed.

156 In addition, when Bortezomib is given in combination with other chemotherapeutic medicinal products, appropriate
 157 dose reductions for these products should be considered in the event of toxicities according to the recommendations
 158 in the Summary of Product Characteristics.

159 Posology for patients with previously untreated mantle cell lymphoma (MCL)

160 [\(Ref.1: Velcade EU/1/04/274/001 page 5 - 6\)](#)

161 *Combination therapy with rituximab, cyclophosphamide, doxorubicin and prednisone (VcR-CAP)*

162 Bortezomib 3.5 mg powder for solution for injection is administered via intravenous or subcutaneous injection at
 163 the recommended dose of 1.3 mg/m² body surface area twice weekly for two weeks on days 1, 4, 8, and 11,
 164 followed by a 10-day rest period on days 12-21 .This 3-week period is considered a treatment cycle .Six Bortezomib

165 cycles are recommended, although for patients with a response first documented at cycle 6, two additional
 166 Bortezomib cycles may be given .At least 72 hours should elapse between consecutive doses of Bortezomib.
 167 The following medicinal products are administered on day 1 of each Bortezomib 3 week treatment cycle as
 168 intravenous infusions :rituximab at 375 mg/m², cyclophosphamide at 750 mg/m² and doxorubicin at 50 mg/m².
 169 Prednisone is administered orally at 100 mg/m² on days 1, 2, 3, 4 and 5 of each Bortezomib treatment cycle.
 170 Dose adjustments during treatment for patients with previously untreated mantle cell lymphoma
 171 Prior to initiating a new cycle of therapy:

- 172 • Platelet counts should be $\geq 100,000$ cells/ μL and the absolute neutrophils count (ANC) should be $\geq 1,500$
- 173 cells/ μL
- 174 • Platelet counts should be $\geq 75,000$ cells/ μL in patients with bone marrow infiltration or splenic sequestration
- 175 • Haemoglobin ≥ 8 g/dL
- 176 • Non-haematological toxicities should have resolved to Grade 1 or baseline.

177 Bortezomib treatment must be withheld at the onset of any \geq Grade 3 Bortezomib- related non-haematological
 178 toxicities (excluding neuropathy) or \geq Grade 3 haematological toxicities (see also section 4.4) For dose adjustments,
 179 see Table 5 below.

180 Granulocyte colony stimulating factors may be administered for haematologic toxicity according to local standard
 181 practice .Prophylactic use of granulocyte colony stimulating factors should be considered in case of repeated delays
 182 in cycle administration .Platelet transfusion for the treatment of thrombocytopenia should be considered when
 183 clinically appropriate.
 184

185 *Table 5 :Dose adjustments during treatment for patients with previously untreated mantle cell lymphoma*

Toxicity	Posology modification or delay
<i>Haematological toxicity</i>	
<ul style="list-style-type: none"> • \geq Grade 3 neutropenia with fever, Grade 4 neutropenia lasting more than 7 days, a platelet count $< 10,000$ cells/μL 	<p>Bortezomib therapy should be withheld for up to 2 weeks until the patient has an ANC ≥ 750 cells/μL and a platelet count $\geq 25,000$ cells/μL.</p> <ul style="list-style-type: none"> • If, after Bortezomib has been held, the toxicity does not resolve, as defined above, then Bortezomib must be discontinued. • If toxicity resolves i.e .patient has an ANC ≥ 750 cells/μL and a platelet count $\geq 25,000$ cells/μL, Bortezomib may be reinitiated at a dose reduced by one dose level (from 1.3 mg/m² to 1 mg/m², or from 1 mg/m² to 0.7 mg/m²)

<ul style="list-style-type: none"> • If platelet counts < 25,000 cells/μL .or ANC < 750 cells/μL on a Bortezomib dosing day)other than Day 1 of each cycle(Bortezomib therapy should be withheld
Grade ≥ 3 non-haematological toxicities considered to be related to Bortezomib	Bortezomib therapy should be withheld until symptoms of the toxicity have resolved to Grade 2 or better . Then, Bortezomib may be reinitiated at a dose reduced by one dose level (from 1.3 mg/m ² to 1 mg/m ² , or from 1 mg/m ² to 0.7 mg/m ²) . For Bortezomib- related neuropathic pain and /or peripheral neuropathy, hold and /or modify Bortezomib as outlined in Table 1.

186 In addition, when Bortezomib is given in combination with other chemotherapeutic medicinal products, appropriate
187 dose reductions for these medicinal products should be considered in the event of toxicities, according to the
188 recommendations in the respective Summary of Product Characteristics.

189

190 Special populations

191 *Elderly*

192 [\(Ref.1: Velcade EU/1/04/274/001 page 6\)](#)

193 There is no evidence to suggest that dose adjustments are necessary in patients over 65 years of age with multiple
194 myeloma or with mantle cell lymphoma.

195 There are no studies on the use of Bortezomib in elderly patients with previously untreated multiple myeloma who
196 are eligible for high -dose chemotherapy with haematopoietic stem cell transplantation . Therefore no dose
197 recommendations can be made in this population.

198 *In a study in previously untreated mantle cell lymphoma patients, 42.9% and 10.4% of patients exposed to
199 Bortezomib were in the range 65-74 years and ≥ 75 years of age, respectively. In patients aged ≥ 75 years,
200 both regimens, VcR-CAP as well as R-CHOP, were less tolerated (see section 4.8)*

201

202 *Hepatic impairment*

203 [\(Ref.1: Velcade EU/1/04/274/001 page 6 - 7\)](#)

204 Patients with mild hepatic impairment do not require a dose adjustment and should be treated per the recommended
205 dose .Patients with moderate or severe hepatic impairment should be started on Bortezomib at a reduced dose
206 of 0.7 mg/m² per injection during the first treatment cycle, and a subsequent dose escalation to 1.0 mg/m² or further
207 dose reduction to 0.5 mg/m² may be considered based on patient tolerability (see Table 6 and sections 4.4 and
208 5.2)

209 *Table 6 :Recommended starting dose modification for Bortezomib in patients with hepatic impairment*

Grade of hepatic impairment*	Bilirubin level	SGOT (AST) levels	Modification of starting dose
Mild	$\leq 1.0 \times \text{ULN}$	$> \text{ULN}$	None
	$> 1.0 \times -1.5 \times \text{ULN}$	Any	None
Moderate	$> 1.5 \times -3 \times \text{ULN}$	Any	Reduce Bortezomib to 0.7 mg /m ² in the first treatment cycle . Consider dose escalation to 1.0 mg /m ² or further dose reduction to 0.5 mg /m ² in subsequent cycles based on patient tolerability.
Severe	$> 3 \times \text{ULN}$	Any	

210 Abbreviations : SGOT=serum glutamic oxaloacetic transaminase;

211 AST=aspartate aminotransferase; ULN=upper limit of the normal range.

212

213 *Renal impairment*

214 [\(Ref.1: Velcade EU/1/04/274/001 page 7\)](#)

215 The pharmacokinetics of bortezomib are not influenced in patients with mild to moderate renal impairment
 216 (Creatinine Clearance [CrCL] $> 20 \text{ ml/min/1.73 m}^2$); therefore, dose adjustments are not necessary for these
 217 patients .It is unknown if the pharmacokinetics of bortezomib are influenced in patients with severe renal impairment
 218 not undergoing dialysis (CrCL $< 20 \text{ ml/min/1.73 m}^2$). Since dialysis may reduce bortezomib concentrations,
 219 BORTEZOMIB should be administered after the dialysis procedure (see section 5.2).

220

221 *Paediatric population*

222 [\(Ref.1: Velcade EU/1/04/274/001 page 7\)](#)

223 The safety and efficacy of Bortezomib in children below 18 years of age have not been established (see sections
 224 5.1 and 5.2). Currently available data are described in section 5.1 but no recommendation on a posology can be
 225 made.

226

227 Method of administration

228 [\(Ref.1: Velcade EU/1/04/274/001 page 7\)](#)

229 Bortezomib 3.5 mg powder for solution for injection is available for intravenous or subcutaneous administration.

230 Bortezomib 1 mg powder for solution for injection is available for intravenous administration only.

231 Bortezomib should not be given by other routes .Intrathecal administration has resulted in death.

232

233 *Intravenous injection*

234 Bortezomib 3.5 mg reconstituted solution is administered as a 3 – 5 second bolus intravenous injection through a
235 peripheral or central intravenous catheter followed by a flush with sodium chloride 9 mg/ml (9%) solution for
236 injection. At least 72 hours should elapse between consecutive doses of Bortezomib.

237

238 *Subcutaneous injection*

239 Bortezomib 3.5 mg reconstituted solution is administered subcutaneously through the thighs (right or left) or
240 abdomen (right or left). The solution should be injected subcutaneously, at a 45 - 90° angle .Injection sites should
241 be rotated for successive injections.

242 If local injection site reactions occur following Bortezomib subcutaneous injection, either a less concentrated
243 Bortezomib solution (Bortezomib 3.5 mg to be reconstituted to 1 mg/ml instead of 2.5 mg/ml) may be administered
244 subcutaneously or a switch to intravenous injection is recommend.

245 *When Bortezomib is given in combination with other medicinal products, refer to the Summary of Product*

246 *Characteristics of these products for instructions for administration*

247

248 **4.3 Contraindications**

249 [\(Ref.1: Velcade EU/1/04/274/001 page 7\)](#)

250 Hypersensitivity to the active substance, to boron or to any of the excipients listed in section 6.1.

251 *Acute diffuse infiltrative pulmonary and pericardial disease.*

252 *When Bortezomib is given in combination with other medicinal products, refer to their Summaries of Product*

253 *Characteristics for additional contraindications.*

254

255 **4.4 Special warnings and precautions for use**

256 [\(Ref.1: Velcade EU/1/04/274/001 page 7 – 10\)](#)

257

258 Intrathecal administration

259 [\(Ref.1: Velcade EU/1/04/274/001 page 7\)](#)

260 There have been fatal cases of inadvertent intrathecal administration of Bortezomib. Bortezomib 1 mg powder for
261 solution for injection is for intravenous use only, while Bortezomib 3.5 mg powder for solution for injection is for
262 intravenous or subcutaneous use .Bortezomib should not be administered intrathecally.

263

264 Gastrointestinal toxicity

265 [\(Ref.1: Velcade EU/1/04/274/001 page 7\)](#)

266 Gastrointestinal toxicity, including nausea, diarrhea, vomiting and constipation are very common with Bortezomib
267 treatment . Cases of ileus have been uncommonly reported. (see section 4.8) Therefore, patients who experience
268 constipation should be closely monitored.

269
270 Haematological toxicity

271 [\(Ref.1: Velcade EU/1/04/274/001 page 8\)](#)

272 Bortezomib treatment is very commonly associated with thrombocytopenia and neutropenia. Platelets counts
273 decreased typically with dose-related and recovering before the initiation of the subsequent cycle in the clinical
274 studies in patients with previously untreated multiple myeloma, relapsed multiple myeloma, or mantle cell
275 lymphoma. There was no evidence of cumulative thrombocytopenia and neutropenia.

276
277 Herpes zoster virus reactivation

278 [\(Ref.1: Velcade EU/1/04/274/001 page 8\)](#)

279 Antiviral prophylaxis is recommended in patients being treated with Bortezomib.

280 In the Phase III study in patients with previously untreated multiple myeloma, the overall incidence of herpes
281 zoster reactivation was more common in patients treated with BORTEZOMIB +Melphalan+Prednisone compared
282 with Melphalan+Prednisone (14% versus 4% respectively).

283 In patients with MCL (study LYM-3002), the incidence of herpes zoster infection was 6.7% in the VcR-CAP arm
284 and 1.2% in the R-CHOP arm (see section 4.8).

285
286 Hepatitis B Virus (HBV) reactivation and infection

287 [\(Ref.1: Velcade EU/1/04/274/001 page 8\)](#)

288 When rituximab is used in combination with Bortezomib, HBV screening must always be performed in patients at
289 risk of infection with HBV before initiation of treatment. Carriers of hepatitis B and patients with a history of
290 hepatitis B must be closely monitored for clinical and laboratory signs of active HBV infection during and following
291 rituximab combination treatment with Bortezomib. Antiviral prophylaxis should be considered. Refer to the
292 Summary of Product Characteristics of rituximab for more information.

293
294 Progressive multifocal leukoencephalopathy (PML)

295 [\(Ref.1: Velcade EU/1/04/274/001 page 8\)](#)

296 Very rare cases with unknown causality of John Cunningham (JC) virus infection, resulting in PML and death,
297 have been reported in patients treated with Bortezomib. Patients diagnosed with PML had prior or concurrent
298 immunosuppressive therapy. Most cases of PML were diagnosed within 12 months of their first dose of
299 Bortezomib. Patients should be monitored at regular intervals for any new or worsening neurological symptoms or

300 signs that may be suggestive of PML as part of the differential diagnosis of CNS problems. If a diagnosis of PML
301 is suspected, patients should be referred to a specialist in PML and appropriate diagnostic measures for PML
302 should be initiated. Discontinue Bortezomib if PML is diagnosed.

303

304 Peripheral neuropathy

305 [\(Ref.1: Velcade EU/1/04/274/001 page 9\)](#)

306 Treatment with Bortezomib is very commonly associated with peripheral neuropathy, which is predominantly
307 sensory .However, cases of severe motor neuropathy with or without sensory peripheral neuropathy have been
308 reported .

309 It is recommended that patients be carefully monitored for symptoms of neuropathy such as a burning sensation,
310 hyperesthesia, hypoesthesia, paraesthesia, discomfort, neuropathic pain or weakness.

311 Patients experiencing new or worsening peripheral neuropathy should undergo neurological evaluation and may
312 require a change in the dose, schedule or route of administration to subcutaneous. (see section 4.2) Neuropathy
313 has been managed with supportive care and other therapies.

314

315 Seizures

316 [\(Ref.1: Velcade EU/1/04/274/001 page 9\)](#)

317 Seizures have been uncommonly reported in patients without previously history of seizures or epilepsy. Special
318 care is required when treating patients with any risk factors for seizures

319

320 Hypotension

321 [\(Ref.1: Velcade EU/1/04/274/001 page 9\)](#)

322 Bortezomib treatment is commonly associated with orthostatic/postural hypotension .Most adverse reactions are
323 mild to moderate in nature and are observed throughout treatment .Patients who developed orthostatic
324 hypotension on Bortezomib (injected intravenously) did not have evidence of orthostatic hypotension prior to
325 treatment with Bortezomib. Most patients required treatment for their orthostatic hypotension .A minority of
326 patients with orthostatic hypotension experienced syncopal events. Orthostatic/postural hypotension was not
327 acutely related to bolus infusion of Bortezomib. The mechanism of this event is unknown although a component
328 may be due to autonomic neuropathy. Autonomic neuropathy may be related to bortezomib or bortezomib may
329 aggravate an underlying condition such as diabetic or amyloidotic neuropathy. Caution is advised when treating
330 patients with a history of syncope receiving medicinal products known to be associated with hypotension; or who
331 are dehydrated due to recurrent diarrhoea or vomiting .Management of orthostatic/postural hypotension may
332 include adjustment of antihypertensive medicinal products, rehydration or administration of mineralocorticosteroids

333 and/or sympathomimetics. Patients should be instructed to seek medical advice if they experience symptoms of
334 dizziness, or fainting spells.

335

336 Posterior Reversible Encephalopathy Syndrome (PRES)

337 [\(Ref.1: Velcade EU/1/04/274/001 page 9\)](#)

338 There have been reports of PRES in patients receiving Bortezomib .PRES is a rare, often reversible, rapidly
339 evolving neurological condition, which can present with seizure, hypertension, headache, lethargy, confusion,
340 blindness, and other visual and neurological disturbances .Brain imaging, preferably Magnetic Resonance Imaging,
341 (MRI)is used to confirm the diagnosis .In patients developing PRES, Bortezomib should be discontinued.

342

343 Heart failure

344 [\(Ref.1: Velcade EU/1/04/274/001 page 9\)](#)

345 Acute development or exacerbation of congestive heart failure, and/or new onset of decreased left ventricular
346 ejection fraction has been reported during bortezomib treatment .Fluid retention may be a predisposing factor for
347 signs and symptoms of heart failure .Patients with risk factors for or existing heart disease should be closely
348 monitored.

349

350 Electrocardiogram investigations

351 [\(Ref.1: Velcade EU/1/04/274/001 page 9\)](#)

352 There have been isolated cases of QT-interval prolongation in clinical studies, causality has not been established.

353

354 Pulmonary disorders

355 [\(Ref.1: Velcade EU/1/04/274/001 page 9 - 10\)](#)

356 There have been rare reports of acute diffuse infiltrative pulmonary disease of unknown aetiology such as
357 pneumonitis, interstitial pneumonia, lung infiltration, and acute respiratory distress syndrome (ARDS) in patients
358 receiving Bortezomib (see section 4.8). **Some of these events have been fatal. A pre-treatment chest radiograph
359 is recommended to serve as a baseline for potential post-treatment pulmonary changes.**

360 In the event of new or worsening pulmonary symptoms, (e.g., cough, dyspnea) a prompt diagnostic evaluation
361 should be performed and patients treated appropriately .The benefit/risk ratio should be considered prior to
362 continuing Bortezomib therapy.

363 **In a clinical trial, two patients (out of 2) given high-dose cytarabine (2 g/m² per day) by continuous infusion over
364 24 hours with daunorubicin and Bortezomib for relapsed acute myelogenous leukaemia died of ARDS early in the
365 course of therapy, and the study was terminated. Therefore, this specific regimen with concomitant administration
366 with high-dose cytarabine (2 g/m² per day) by continuous infusion over 24 hours is not recommended.**

367 Renal impairment

368 ([Ref.1: Velcade EU/1/04/274/001 page 10](#))

369 Renal complications are frequent in patients with multiple myeloma .Patients with renal impairment should be
370 monitored closely. (see sections 4.2 and 5.2)

371

372 Hepatic impairment

373 ([Ref.1: Velcade EU/1/04/274/001 page 10](#))

374 Bortezomib is metabolised by liver enzymes .Bortezomib exposure is increased in patients with moderate or severe
375 hepatic impairment; these patients should be treated with Bortezomib at reduced doses and closely monitored for
376 toxicities. (see sections 4.2 and 5.2)

377

378 Hepatic reactions

379 ([Ref.1: Velcade EU/1/04/274/001 page 10](#))

380 Rare cases of hepatic failure have been reported in patients receiving Bortezomib and concomitant medicinal
381 products and with serious underlying medical conditions .Other reported hepatic reactions include increases in liver
382 enzymes, hyperbilirubinaemia, and hepatitis .Such changes may be reversible upon discontinuation of bortezomib)
383 (see section 4.8)

384

385 Tumour lysis syndrome

386 ([Ref.1: Velcade EU/1/04/274/001 page 10](#))

387 Because bortezomib is a cytotoxic agent and can rapidly kill malignant plasma cells and MCL cells, the
388 complications of tumour lysis syndrome may occur .The patients at risk of tumour lysis syndrome are those with
389 high tumour burden prior to treatment .These patients should be monitored closely and appropriate precautions
390 taken.

391

392 Concomitant medicinal products

393 ([Ref.1: Velcade EU/1/04/274/001 page 10](#))

394 Patients should be closely monitored when given bortezomib in combination with potent CYP3A4-inhibitors .Caution
395 should be exercised when bortezomib is combined with CYP3A4 -or CYP2C19 substrates (see sections 4.5).

396 Normal liver function should be confirmed and caution should be exercised in patients receiving oral hypoglycemics
397 (see sections 4.5).

398

399 Potentially immunocomplex-mediated reactions

400 ([Ref.1: Velcade EU/1/04/274/001 page 10](#))

401 Potentially immunocomplex-mediated reactions, such as serum-sickness-type reaction, polyarthritis with rash and
402 proliferative glomerulonephritis have been reported uncommonly. Bortezomib should be discontinued if serious
403 reactions occur.

404 **4.5 Interaction with other medicinal products and other forms of interactions**

405 [\(Ref.1: Velcade EU/1/04/274/001 page 10 - 11\)](#)

406 In vitro studies indicate that bortezomib is a weak inhibitor of the cytochrome P45 (CYP) isozymes 1A2, 2C9, 2C19,
407 2D6 and 3A4 .Based on the limited contribution (7%) of CYP2D6 to the metabolism of bortezomib, the CYP2D6
408 poor metaboliser phenotype is not expected to affect the overall disposition of bortezomib.

409 A drug -drug interaction study assessing the effect of ketoconazole, a potent CYP3A4 inhibitor, on the
410 pharmacokinetics of bortezomib (injected intravenously), showed a mean bortezomib AUC increase of 35% (CI
411 90% [1.032 to 1.772]) based on data from 12 patients .Therefore, patients should be closely monitored when given
412 bortezomib in combination with potent CYP3A4 inhibitors (e.g .ketoconazole, ritonavir).

413 In a drug -drug interaction study assessing the effect of omeprazole, a potent CYP2C19 inhibitor, on the
414 pharmacokinetics of bortezomib, (injected intravenously) there was no significant effect on the pharmacokinetics of
415 bortezomib based on data from 17 patients.

416 A drug-drug interaction study assessing the effect of rifampicin, a potent CYP3A4 inducer, on the pharmacokinetics
417 of bortezomib, (injected intravenously) showed a mean bortezomib AUC reduction of 45 % based on data from 6
418 patients . Therefore, the concomitant use of bortezomib with strong CYP3A4 inducers (e .g ., rifampicin,
419 carbamazepine, phenytoin, phenobarbital and St .John's Wort) is not recommended, as efficacy may be reduced.

420 In the same drug-drug interaction study assessing the effect of dexamethasone, a weaker CYP3A4 inducer, on the
421 pharmacokinetics of bortezomib (injected intravenously), there was no significant effect on the pharmacokinetics of
422 bortezomib based on data from 7 patients.

423 A drug-drug interaction study assessing the effect of melphalan-prednisone on the pharmacokinetics of bortezomib
424 (injected intravenously), showed a mean bortezomib AUC increase of 17 % based on data from 21 patients . This
425 is not considered clinically relevant.

426 During clinical trials, hypoglycemia and hyperglycemia were uncommonly and commonly reported in diabetic
427 patients receiving oral hypoglycemics .Patients on oral antidiabetic agents receiving Bortezomib treatment may
428 require close monitoring of their blood glucose levels and adjustment of the dose of their antidiabetics.

429

430 **4.6 Pregnancy and lactation**

431 [\(Ref.1: Velcade EU/1/04/274/001 page 11\)](#)

432 Contraception in males and females

433 Male and female patients of childbearing potential must use effective contraceptive measures during and for 3
434 months following treatment.

435

436 Pregnancy

437 No clinical data are available for bortezomib with regard to exposure during pregnancy.
438 In non-clinical studies, bortezomib had no effects on embryonal/foetal development in rats and rabbits at the highest
439 maternally tolerated doses. Animal studies to determine the effects of bortezomib on parturition and post-natal
440 development were not conducted (see section 5.3). Bortezomib should not be used during pregnancy unless the
441 clinical condition of the woman requires treatment with Bortezomib.
442 If Bortezomib is used during pregnancy, or if the patient becomes pregnant while receiving this medicinal product,
443 the patient should be informed of potential for hazard to the foetus.

444

445 Breast-feeding

446 It is not known whether bortezomib is excreted in human milk .Because of the potential for serious adverse reactions
447 in breast-fed infants, breast-feeding should be discontinued during treatment with Bortezomib.

448

449 Fertility

450 Fertility studies were not conducted with Bortezomib (see section 5.3).

451

452 **4.7 Effects on ability to drive and use machine**

453 (Ref.1: Velcade EU/1/04/274/001 page 11)

454 Bortezomib may have a moderate influence on the ability to drive and use machines .Bortezomib may be
455 associated with fatigue very commonly, dizziness commonly, syncope uncommonly and orthostatic /postural
456 hypotension or blurred vision commonly .Therefore, patients must be cautious when driving or using machines and
457 should be advised not to drive or operate machinery if they experience these symptoms (see section 4.8)

458

459 **4.8 Undesirable effects**

460 (Ref.1: Velcade EU/1/04/274/001 page 11 - 20)

461 Summary of the safety profile

462 Serious adverse reactions uncommonly reported during treatment with Bortezomib include cardiac failure, tumour
463 lysis syndrome, pulmonary hypertension, posterior reversible encephalopathy syndrome, acute diffuse infiltrative
464 pulmonary disorders and rarely autonomic neuropathy.

465 The most commonly reported adverse reactions during treatment with Bortezomib are nausea, diarrhoea,
466 constipation, vomiting, fatigue, pyrexia, thrombocytopenia, anaemia, neutropenia, peripheral neuropathy (including
467 sensory), headache, paraesthesia, decreased appetite, dyspnoea, rash, herpes zoster and myalgia.

468 Tabulated summary of adverse reactions

469 *Multiple Myeloma*

470 Undesirable effects in Table 7 were considered by the investigators to have at least a possible or probable
 471 causal relationship to Bortezomib. These adverse reactions are based on an integrated data set of 5,476 patients
 472 of whom 3,996 were treated with Bortezomib at 1.3 mg/m² and included in Table 7.

473 Overall, Bortezomib was administered for the treatment of multiple myeloma in 3,974 patients.

474 Adverse reactions are listed below by system organ class and frequency grouping. Frequencies are defined as:
 475 Very common ($\geq 1/10$); common ($\geq 1/100$ to $< 1/10$); uncommon ($\geq 1/1,000$ to $< 1/100$); rare ($\geq 1/10,000$ to $<$
 476 $1/1,000$); very rare ($< 1/10,000$), not known (cannot be estimated from the available data). Within each frequency
 477 grouping, undesirable effects are presented in order of decreasing seriousness. Table 7 has been generated
 478 using Version 14.1 of the MedDRA.

479 Post-marketing adverse reactions not seen in clinical trials are also included.

480 *Table 7: Adverse reactions in patients with Multiple Myeloma treated with BORTEZOMIB in clinical trials, and all*
 481 *post-marketing adverse reactions regardless of indication#*

System Organ Class	Incidence	Adverse reaction
Infections and infestations	Common	Herpes zoster inc disseminated & ophthalmic, Pneumonia*, Herpes simplex*, Fungal infection*
	Uncommon	Infection*, Bacterial infections*, Viral infections*, Sepsis inc septic shock*, Bronchopneumonia, Herpes virus infection*, Meningoencephalitis herpetic#, Bacteraemia inc staphylococcal, Hordeolum, Influenza, Cellulitis, Device related infection, Skin infection*, Ear infection*, Staphylococcal infection, Tooth infection*
	Rare	Meningitis inc bacterial, Epstein-Barr virus infection, Genital herpes, Tonsillitis, Mastoiditis, Post viral fatigue syndrome
Neoplasms benign, malignant and unspecified (incl cysts and polyps)	Rare	Neoplasm malignant, Leukaemia plasmacytic, Renal cell carcinoma, Mass, Mycosis fungoides, Neoplasm benign*
Blood and lymphatic system disorders	Very Common	Thrombocytopenia*, Neutropenia*, Anaemia*
	Common	Leukopenia*, Lymphopenia*
	Uncommon	Pancytopenia*, Febrile neutropenia, Coagulopathy*, Leukocytosis*, Lymphadenopathy, Haemolytic anaemia#
	Rare	Disseminated intravascular coagulation, Thrombocytosis*, Hyperviscosity syndrome, Platelet disorder NOS, Thrombotic microangiopathy inc thrombocytopenic purpura#, Blood disorder NOS, Haemorrhagic diathesis, Lymphocytic infiltration
Immune system disorders	Uncommon	Angioedema#, Hypersensitivity*
	Rare	Anaphylactic shock, Amyloidosis, Type III immune complex mediated reaction
Endocrine disorders	Uncommon	Cushing's syndrome*, Hyperthyroidism*, Inappropriate antidiuretic hormone secretion

	Rare	Hypothyroidism
Metabolism and nutrition disorders	Very Common	Decreased appetite
	Common	Dehydration, Hypokalaemia*, Hyponatraemia*, Blood glucose abnormal*, Hypocalcaemia*, Enzyme abnormality*
	Uncommon	Tumour lysis syndrome, Failure to thrive*, Hypomagnesaemia*, Hypophosphataemia*, Hyperkalaemia*, Hypercalcaemia*, Hyponatraemia*, Uric acid abnormal*, Diabetes mellitus*, Fluid retention
	Rare	Hypermagnesaemia*, Acidosis, Electrolyte imbalance*, Fluid overload, Hypochlorhaemia*, Hypovolaemia, Hyperchlorhaemia*, Hyperphosphataemia*, Metabolic disorder, Vitamin B complex deficiency, Vitamin B12 deficiency, Gout, Increased appetite, Alcohol intolerance
Psychiatric disorders	Common	Mood disorders and disturbances*, Anxiety disorder*, Sleep disorders and disturbances*
	Uncommon	Mental disorder*, Hallucination*, Psychotic disorder*, Confusion*, Restlessness
	Rare	Suicidal ideation*, Adjustment disorder, Delirium, Libido decreased
Nervous system disorders	Very common	Neuropathies*, Peripheral sensory neuropathy, Dysaesthesia*, Neuralgia*
	Common	Motor neuropathy*, Loss of consciousness inc syncope, Dizziness*, Dysgeusia*, Lethargy, Headache*
	Uncommon	Tremor, Peripheral sensorimotor neuropathy, Dyskinesia*, Cerebellar coordination and balance disturbances*, Memory loss exc dementia*, Encephalopathy*, Posterior Reversible Encephalopathy Syndrome#, Neurotoxicity, Seizure disorders*, Post herpetic neuralgia, Speech disorder*, Restless legs syndrome, Migraine, Sciatica, Disturbance in attention, Reflexes abnormal*, Parosmia
	Rare	Cerebral haemorrhage*, Haemorrhage intracranial inc subarachnoid*, Brain oedema, Transient ischaemic attack, Coma, Autonomic nervous system imbalance, Autonomic neuropathy, Cranial palsy*, Paralysis*, Paresis*, Presyncope, Brain stem syndrome, Cerebrovascular disorder, Nerve root lesion, Psychomotor hyperactivity, Spinal cord compression, Cognitive disorder NOS, Motor dysfunction, Nervous system disorder NOS, Radiculitis, Drooling, Hypotonia
Eye disorders	Common	Eye swelling*, Vision abnormal*, Conjunctivitis*

	Uncommon	Eye haemorrhage*, Eyelid infection*, Chalazion#, Blepharitis#, Eye inflammation*, Diplopia, Dry eye*, Eye irritation*, Eye pain, Lacrimation increased, Eye discharge
	Rare	Corneal lesion*, Exophthalmos, Retinitis, Scotoma, Eye disorder (inc. eyelid) NOS, Dacryoadenitis acquired, Photophobia, Photopsia, Optic neuropathy#, Different degrees of visual impairment up to blindness*
Ear and labyrinth disorders	Common	Vertigo*
	Uncommon	Dysacusis inc tinnitus*,Hearing impaired (up to and inc deafness), Ear discomfort*
	Rare	Ear haemorrhage, Vestibular neuronitis, Ear disorder NOS
Cardiac disorders	Uncommon	Cardiac tamponade#, Cardio-pulmonary arrest*, Cardiac fibrillation inc atrial, Cardiac failure inc left and right ventricular*, Arrhythmia*, Tachycardia*, Palpitations, Angina pectoris, Pericarditis inc pericardial effusion*, Cardiomyopathy*, Ventricular dysfunction*, Bradycardia
	Rare	Atrial flutter, Myocardial infarction*, Atrioventricular block*, Cardiovascular disorder inc cardiogenic shock, Torsade de pointes, Angina unstable, Cardiac valve disorders*, Coronary artery insufficiency, Sinus arrest
Vascular disorders	Common	Hypotension*, Orthostatic hypotension, Hypertension*
	Uncommon	Cerebrovascular accident#, Deep vein thrombosis*, Haemorrhage*, Thrombophlebitis inc superficial, Circulatory collapse inc hypovolaemic shock, Phlebitis, Flushing*, Haematoma inc perirenal*, Poor peripheral circulation*, Vasculitis, Hyperaemia inc ocular*
	Rare	Peripheral embolism, Lymphoedema, Pallor, Erythromelalgia, Vasodilatation, Vein discolouration, Venous insufficiency
Respiratory, thoracic and mediastinal disorders	Common	Dyspnoea*, Epistaxis, Upper/lower respiratory tract infection*, Cough*
	Uncommon	Pulmonary embolism, Pleural effusion, Pulmonary oedema inc acute, Pulmonary alveolar haemorrhage# , Bronchospasm, Chronic obstructive pulmonary disease*, Hypoxaemia*, Respiratory tract congestion*, Hypoxia, Pleurisy*, Hiccups, Rhinorrhoea, Dysphonia, Wheezing
	Rare	Respiratory failure, Acute respiratory distress syndrome, Apnoea, Pneumothorax, Atelectasis, Pulmonary hypertension, Haemoptysis, Hyperventilation, Orthopnoea, Pneumonitis, Respiratory alkalosis, Tachypnoea, Pulmonary fibrosis, Bronchial disorder*, Hypocapnia*, Interstitial lung disease, Lung infiltration, Throat tightness, Dry throat, Increased upper airway secretion, Throat irritation, Upper-airway cough syndrome

Gastrointestinal disorders	Very common	Nausea and vomiting symptoms*, Diarrhoea*, Constipation
	Common	Gastrointestinal haemorrhage inc mucosal*, Dyspepsia, Stomatitis*, Abdominal distension, Oropharyngeal pain*, Abdominal pain inc gastrointestinal and splenic pain*, Oral disorder*, Flatulence
	Uncommon	Pancreatitis inc chronic*, Haematemesis, Lip swelling*, Gastrointestinal obstruction inc small intestinal obstruction, ileus*, Abdominal discomfort, Oral ulceration*, Enteritis*, Gastritis*, Gingival bleeding, Gastroesophageal reflux disease*, Colitis inc clostridium difficile*, Colitis ischaemic#, Gastrointestinal inflammation*, Dysphagia, Irritable bowel syndrome, Gastrointestinal disorder NOS, Tongue coated, Gastrointestinal motility disorder*, Salivary gland disorder*
	Rare	Pancreatitis acute, Peritonitis*, Tongue oedema*, Ascites, Oesophagitis, Cheilitis, Faecal incontinence, Anal sphincter atony, Faecaloma*, Gastrointestinal ulceration and perforation*, Gingival hypertrophy, Megacolon, Rectal discharge, Oropharyngeal blistering*, Lip pain, Periodontitis, Anal fissure, Change of bowel habit, Proctalgia, Abnormal faeces
Hepatobiliary disorders	Common	Hepatic enzyme abnormality*
	Uncommon	Hepatotoxicity inc liver disorder, Hepatitis*, Cholestasis
	Rare	Hepatic failure, Hepatomegaly, Budd-Chiari syndrome, Cytomegalovirus hepatitis, Hepatic haemorrhage, Cholelithiasis
Skin and subcutaneous tissue disorders	Common	Rash*, Pruritus*, Erythema, Dry skin
	Uncommon	Erythema multiforme, Urticaria, Acute febrile neutrophilic dermatosis, Toxic skin eruption, Toxic epidermal necrolysis#, Stevens-Johnson syndrome#, Dermatitis*, Hair disorder*, Petechiae, Ecchymosis, Skin lesion, Purpura, Skin mass*, Psoriasis, Hyperhidrosis, Night sweats, Decubitus ulcer#, Acne*, Blister*, Pigmentation disorder*
	Rare	Skin reaction, Jessner's lymphocytic infiltration, Palmar-plantar erythrodysesthesia syndrome, Haemorrhage subcutaneous, Livedo reticularis, Skin induration, Papule, Photosensitivity reaction, Seborrhoea, Cold sweat, Skin disorder NOS, Erythrosis, Skin ulcer, Nail disorder
Musculoskeletal and connective tissue disorders	Very common	Musculoskeletal pain*
	Common	Muscle spasms*, Pain in extremity, Muscular weakness
	Uncommon	Muscle twitching, Joint swelling, Arthritis*, Joint stiffness, Myopathies*, Sensation of heaviness

	Rare	Rhabdomyolysis, Temporomandibular joint syndrome, Fistula, Joint effusion, Pain in jaw, Bone disorder, Musculoskeletal and connective tissue infections and inflammations*, Synovial cyst
Renal and urinary disorders	Common	Renal impairment*
	Uncommon	Renal failure acute, Renal failure chronic*, Urinary tract infection*, Urinary tract signs and symptoms*, Haematuria*, Urinary retention, Micturition disorder*, Proteinuria, Azotaemia, Oliguria*, Pollakiuria
	Rare	Bladder irritation
Reproductive system and breast disorders	Uncommon	Vaginal haemorrhage, Genital pain*, Erectile dysfunction
	Rare	Testicular disorder*, Prostatitis, Breast disorder female, Epididymal tenderness, Epididymitis, Pelvic pain, Vulval ulceration
Congenital, familial and genetic disorders	Rare	Aplasia, Gastrointestinal malformation, Ichthyosis
General disorders and administration site conditions	Very common	Pyrexia*, Fatigue, Asthenia
	Common	Oedema inc peripheral, Chills, Pain*, Malaise*
	Uncommon	General physical health deterioration*, Face oedema*, Injection site reaction*, Mucosal disorder*, Chest pain, Gait disturbance, Feeling cold, Extravasation*, Catheter related complication*, Change in thirst*, Chest discomfort, Feeling of body temperature change*, Injection site pain*
	Rare	Death inc sudden, Multi-organ failure, Injection site haemorrhage*, Hernia inc hiatus*, Impaired healing*, Inflammation, Injection site phlebitis*, Tenderness, Ulcer, Irritability, Non-cardiac chest pain, Catheter site pain, Sensation of foreign body
Investigations	Common	Weight decreased
	Uncommon	Hyperbilirubinaemia*, Protein analyses abnormal*, Weight increased, Blood test abnormal*, C-reactive protein increased
	Rare	Blood gases abnormal*, Electrocardiogram abnormalities inc QT prolongation*, International normalised ratio abnormal*, Gastric pH decreased, Platelet aggregation increased, Troponin I increased, Virus identification and serology*, Urine analysis abnormal*
Injury, poisoning and procedural complications	Uncommon	Fall, Contusion
	Rare	Transfusion reaction, Fractures*, Rigors*, Face injury, Joint injury*, Burns, Laceration, Procedural pain, Radiation injuries*
Surgical and medical procedures	Rare	Macrophage activation

482 Inc = including; exc = excluding

483 NOS=not otherwise specified

484 * Grouping of more than one MedDRA preferred term.

485 # Post-marketing adverse reaction regardless of indication

486

487 *Mantle Cell Lymphoma (MCL)*

488 (Ref.1: Velcade EU/1/04/274/001 page 16)

489 The safety profile of Bortezomib in 240 MCL patients treated with Bortezomib at 1.3 mg/m² in combination with
490 rituximab, cyclophosphamide, doxorubicin, and prednisone (VcR-CAP) versus 242 patients treated with rituximab,
491 cyclophosphamide, doxorubicin, vincristine, and prednisone [RCHOP] was relatively consistent to that observed in
492 patients with multiple myeloma with main differences described below. Additional adverse drug reactions identified
493 associated with the use of the combination therapy (VcR-CAP) were hepatitis B infection (< 1%) and myocardial
494 ischaemia (1.3%). The similar incidences of these events in both treatment arms, indicated that these adverse drug
495 reactions are not attributable to Bortezomib alone. Notable differences in the MCL patient population as compared
496 to patients in the multiple myeloma studies were a \geq 5% higher incidence of the haematological adverse reactions
497 (neutropenia, thrombocytopenia, leukopenia, anemia, lymphopenia), peripheral sensory neuropathy, hypertension,
498 pyrexia, pneumonia, stomatitis, and hair disorders.

499 Adverse drug reactions identified as those with a \geq 1% incidence, similar or higher incidence in the VcR-CAP arm
500 and with at least a possible or probable causal relationship to the components of the VcR-CAP arm, are listed in
501 Table 8 below. Also included are adverse drug reactions identified in the VcR-CAP arm that were considered by
502 investigators to have at least a possible or probable causal relationship to BORTEZOMIB based on historical data
503 in the multiple myeloma studies.

504 Adverse reactions are listed below by system organ class and frequency grouping. Frequencies are defined as:
505 Very common (\geq 1/10); common (\geq 1/100 to < 1/10); uncommon (\geq 1/1,000 to < 1/100); rare (\geq 1/10,000 to <
506 1/1,000); very rare (< 1/10,000), not known (cannot be estimated from the available data). Within each frequency
507 grouping, undesirable effects are presented in order of decreasing seriousness. Table 8 has been generated using
508 Version 16 of the MedDRA.

509 Table 8: Adverse reactions in patients with Mantle Cell Lymphoma treated with VcR-CAP in a clinical trial

System Organ Class	Incidence	Adverse reaction
Infections and infestations	Very Common	Pneumonia*
	Common	Sepsis inc septic shock*, Herpes zoster inc disseminated & ophthalmic, Herpes virus infection*, Bacterial infections*, Upper/lower respiratory tract infection*, Fungal infection*, Herpes simplex*

	Uncommon	Hepatitis B, Infection*, Bronchopneumonia
Blood and lymphatic system disorders	Very Common	Thrombocytopenia*, Febrile neutropenia, Neutropenia*, Leukopenia*, Anaemia*, Lymphopenia*
	Uncommon	Pancytopenia*
Immune system disorders	Common	Hypersensitivity*
	Uncommon	Anaphylactic reaction
Metabolism and nutrition disorders	Very Common	Decreased appetite
	Common	Hypokalaemia*, Blood glucose abnormal*, Hyponatraemia*, Diabetes mellitus*, Fluid retention
	Uncommon	Tumour lysis syndrome
Psychiatric disorders	Common	Sleep disorders and disturbances*
Nervous system disorders	Very Common	Peripheral sensory neuropathy, Dysaesthesia*, Neuralgia*
	Common	Neuropathies*, Motor neuropathy*, Loss of consciousness inc syncope, Encephalopathy*, Peripheral sensorimotor neuropathy, Dizziness*, Dysgeusia*, Autonomic neuropathy
	Uncommon	Autonomic nervous system imbalance
Eye disorders	Common	Vision abnormal*
Ear and labyrinth disorders	Common	Dysacusis inc tinnitus*
	Uncommon	Vertigo*, Hearing impaired up to and inc deafness
Cardiac disorders	Common	Cardiac fibrillation inc atrial, Arrhythmia*, Cardiac failure inc left and right ventricular*, Myocardial ischaemia, Ventricular dysfunction*
	Uncommon	Cardiovascular disorder inc cardiogenic shock
Vascular disorders	Common	Hypertension*, Hypotension*, Orthostatic hypotension
Respiratory, thoracic and mediastinal disorders	Common	Dyspnoea*, Cough*, Hiccups
	Uncommon	Acute respiratory distress syndrome, Pulmonary embolism, Pneumonitis, Pulmonary hypertension, Pulmonary oedema inc acute
Gastrointestinal disorders	Very Common	Nausea and vomiting symptoms*, Diarrhoea*, Stomatitis*, Constipation
	Common	Gastrointestinal haemorrhage inc mucosal*, Abdominal distension, Dyspepsia, Oropharyngeal pain*, Gastritis*, Oral ulceration*, Abdominal discomfort, Dysphagia, Gastrointestinal inflammation*, Abdominal pain inc gastrointestinal and splenic pain*, Oral disorder*
	Uncommon	Colitis inc clostridium difficile*
Hepatobiliary disorders	Common	Hepatotoxicity inc liver disorder
	Uncommon	Hepatic failure
Skin and subcutaneous tissue disorders	Very Common	Hair disorder*
	Common	Pruritus*, Dermatitis*, Rash*

Musculoskeletal and connective tissue disorders	Common	Muscle spasms*, Musculoskeletal pain*, Pain in extremity
Renal and urinary disorders	Common	Urinary tract infection*
General disorders and administration site conditions	Very Common	Pyrexia*, Fatigue, Asthenia
	Common	Oedema inc peripheral, Chills, Injection site reaction*, Malaise*
Investigations	Common	Hyperbilirubinaemia*, Protein analyses abnormal*, Weight decreased, Weight increased

510 Inc = including

511 * Grouping of more than one MedDRA preferred term.

512

513 Description of selected adverse reactions

514 (Ref.1: Velcade EU/1/04/274/001 page 18)

515

516 *Herpes zoster virus reactivation*

517 Multiple Myeloma

518 Antiviral prophylaxis was administered to 26% of the patients in the Vc+M+P arm. The incidence of herpes zoster
519 among patients in the Vc+M+P treatment group was 17% for patients not administered antiviral prophylaxis
520 compared to 3% for patients administered antiviral prophylaxis.

521

522 Mantle cell lymphoma

523 Antiviral prophylaxis was administered to 137 of 240 patients (57%) in the VcR-CAP arm. The incidence of
524 herpes zoster among patients in the VcR-CAP arm was 10.7% for patients not administered antiviral prophylaxis
525 compared to 3.6% for patients administered antiviral prophylaxis (see section 4.4).

526

527 *Hepatitis B Virus (HBV) reactivation and infection*

528 Mantle cell lymphoma

529 HBV infection with fatal outcomes occurred in 0.8% (n=2) of patients in the non- Bortezomib treatment group
530 (rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone; R-CHOP) and 0.4% (n=1) of patients
531 receiving Bortezomib in combination with rituximab, cyclophosphamide, doxorubicin, and prednisone (VcR-CAP).
532 The overall incidence of hepatitis B infections was similar in patients treated with VcR-CAP or with R-CHOP
533 (0.8% vs 1.2% respectively).

534

535 *Peripheral neuropathy in combination regimens*

536 Multiple Myeloma

537 In trials in which Bortezomib was administered as induction treatment in combination with dexamethasone (study
 538 IFM-2005-01), and dexamethasone-thalidomide (study MMY-3010), the incidence of peripheral neuropathy in the
 539 combination regimens is presented in the table below:

540
 541 *Table 9: Incidence of peripheral neuropathy during induction treatment by toxicity and treatment discontinuation due*
 542 *to peripheral neuropathy*

	IFM-2005-01		MMY-3010	
	VDDx	VcDx	TDx	VcTDx
	(N=239)	(N=239)	(N=126)	(N=130)
Incidence of PN (%)				
All GradePN	3	15	12	45
≥ Grade 2 PN	1	10	2	31
≥ Grade 3 PN	<1	5	0	5
Discontinuation due to PN (%)	<1	2	1	5

543 VDDx=vincristine, doxorubicin, dexamethasone; VcDx= Bortezomib, dexamethasone; TDx=thalidomide,
 544 dexamethasone; VcTDx= Bortezomib, thalidomide, dexamethasone; PN=peripheral neuropathy

545 Note: Peripheral neuropathy included the preferred terms: neuropathy peripheral, peripheral motor neuropathy,
 546 peripheral sensory neuropathy, and polyneuropathy.

547
 548 Mantle cell lymphoma

549 (Ref.1: Velcade EU/1/04/274/001 page 19)

550 In study LYM-3002 in which Bortezomib was administered with rituximab, cyclophosphamide, doxorubicin, and
 551 prednisone (R-CAP), the incidence of peripheral neuropathy in the combination regimens is presented in the table
 552 below:

553
 554 *Table 10: Incidence of peripheral neuropathy in study LYM-3002 by toxicity and treatment discontinuation due to*
 555 *peripheral neuropathy*

	VcR-CAP	R-CHOP
	(N=240)	(N=242)
Incidence of PN (%)		
All GradePN	30	29
≥ Grade 2 PN	18	9
≥ Grade 3 PN	8	4
Discontinuation due to PN (%)	2	<1

556 VcR-CAP= Bortezomib, rituximab, cyclophosphamide, doxorubicin, and prednisone; R-CHOP= rituximab,
557 cyclophosphamide, doxorubicin, vincristine, and prednisone; PN=peripheral neuropathy
558 Peripheral neuropathy included the preferred terms: peripheral sensory neuropathy, neuropathy peripheral,
559 peripheral motor neuropathy, and peripheral sensorimotor neuropathy
560

561 *Elderly MCL patients*

562 (Ref.1: Velcade EU/1/04/274/001 page 19)

563 42.9% and 10.4% of patients in the VcR-CAP arm were in the range 65-74 years and \geq 75 years of age,
564 respectively. Although in patients aged \geq 75 years, both VcR-CAP and R-CHOP were less tolerated, the serious
565 adverse event rate in the VcR-CAP groups was 68%, compared to 42% in the R-CHOP group.
566

567 *Notable differences in the safety profile of Bortezomib administered subcutaneously versus intravenously as single* 568 *agent*

569 (Ref.1: Velcade EU/1/04/274/001 page 19)

570 In the Phase III study patients who received Bortezomib subcutaneously compared to intravenous administration
571 had 13% lower overall incidence of treatment emergent adverse reactions that were Grade 3 or higher in toxicity,
572 and a 5% lower incidence of discontinuation of Bortezomib. The overall incidence of diarrhoea, gastrointestinal
573 and abdominal pain, asthenic conditions, upper respiratory tract infections and peripheral neuropathies were
574 12%-15% lower in the subcutaneous group than in the intravenous group. In addition, the incidence of Grade 3
575 or higher peripheral neuropathies was 10% lower, and the discontinuation rate due to peripheral neuropathies 8%
576 lower for the subcutaneous group as compared to the intravenous group.
577

578 Six percent of patients had an adverse local reaction to subcutaneous administration, mostly redness. Cases
579 resolved in a median of 6 days, dose modification was required in two patients. Two (1%) of the patients had
580 severe reactions; 1 case of pruritus and 1 case of redness.
581

582 The incidence of death on treatment was 5% in the subcutaneous treatment group and 7% in the intravenous
583 treatment group. Incidence of death from "Progressive disease" was 18% in the subcutaneous group and 9% in
584 the intravenous group.
585

586 *Retreatment of patients with relapsed multiple myeloma*

587 (Ref.1: Velcade EU/1/04/274/001 page 19)

588 In a study in which Bortezomib retreatment was administered in 130 patients with relapsed multiple myeloma,
589 who previously had at least partial response on a Bortezomib -containing regimen, the most common all-grade
590 adverse events occurring in at least 25% of patients were thrombocytopenia (55%), neuropathy (40%), anaemia

591 (37%), diarrhoea (35%), and constipation (28%). All grade peripheral neuropathy and grade ≥ 3 peripheral
592 neuropathy were observed in 40% and 8.5% of patients, respectively.

593

594 Reporting of suspected adverse reactions

595 Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows
596 continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to
597 report any suspected adverse reactions via ศูนย์เฝ้าระวังความปลอดภัยด้านผลิตภัณฑ์สุขภาพ กองแผนงานและ
598 วิชาการ สำนักงานคณะกรรมการอาหารและยา กระทรวงสาธารณสุข ถนนติวานนท์อำเภอเมือง จังหวัดนนทบุรี 11000
599 หรือ ผ่านระบบ ที่ <http://thaihpvc.fda.moph.go.th/thaihvc/Public/Webpage/main.jsf>

600

601 **4.9 Overdose**

602 (Ref.1: Velcade EU/1/04/274/001 page 20)

603 In patients, overdose more than twice the recommended dose has been associated with the acute onset of
604 symptomatic hypotension and thrombocytopenia with fatal outcomes.

605 There is no known specific antidote for bortezomib overdose. In the event of an overdose, the patient's vital signs
606 should be monitored and appropriate supportive care given to maintain blood pressure (such as fluids, pressors,
607 and/or inotropic agents) and body temperature. (see sections 4.2 and 4.4)

608

609 **5. Pharmacological Properties**

610 (Ref.1: Velcade EU/1/04/274/001 page 20 - 30)

611 **5.1 Pharmacodynamic properties**

612 (Ref.1: Velcade EU/1/04/274/001 page 20 - 29)

613 Pharmacotherapeutic group: Antineoplastic agents, other antineoplastic agents, ATC code: L01XG01.

614

615 Mechanism of action

616 (Ref.1: Velcade EU/1/04/274/001 page 20)

617 Bortezomib is a proteasome inhibitor . It is specifically designed to inhibit the chymotrypsin-like activity of the 26S
618 proteasome in mammalian cells . The 26S proteasome is a large protein complex that degrades ubiquitinated
619 proteins . The ubiquitin-proteasome pathway plays an essential role in regulating the turnover of specific proteins,
620 thereby maintaining homeostasis within cells . Inhibition of the 26S proteasome prevents this targeted proteolysis
621 and affects multiple signalling cascades within the cell, ultimately resulting in cancer cell death.

622 Bortezomib is highly selective for the proteasome .At 10 μ M concentrations, bortezomib does not inhibit any of a
 623 wide variety of receptors and proteases screened and is more than 1,500 -fold more selective for the proteasome
 624 than for its next preferable enzyme .The kinetics of proteasome inhibition were evaluated in vitro, and bortezomib
 625 was shown to dissociate from the proteasome with a $t_{1/2}$ of 20 minutes, thus demonstrating that proteasome
 626 inhibition by bortezomib is reversible.

627 Experiments have demonstrated that bortezomib is cytotoxic to a variety of cancer cell types and that cancer cells
 628 are more sensitive to the pro -apoptotic effects of proteasome inhibition than normal cells .Bortezomib causes
 629 reduction of tumour growth in vivo in many preclinical tumour models, including multiple myeloma.

630 Data from in vitro, ex -vivo, and animal models with bortezomib suggest that it increases osteoblast differentiation
 631 and activity and inhibits osteoclast function .These effects have been observed in patients with multiple myeloma
 632 affected by an advanced osteolytic disease and treated with bortezomib.

633

634 Clinical efficacy in previously untreated multiple myeloma

635 (Ref.1: Velcade EU/1/04/274/001 page 20 - 22)

636 A prospective Phase III, international, randomised (1:1), open-label clinical study (MMY-3002 VISTA) of 682 patients
 637 was conducted to determine whether Bortezomib (1.3 mg/m² injected intravenously) in combination with melphalan
 638 (9 mg/m²) and prednisone (60 mg/m²) resulted in improvement in time to progression (TTP) when compared to
 639 melphalan (9 mg/m²) and prednisone (60 mg/m²) in patients with previously untreated multiple myeloma. Treatment
 640 was administered for a maximum of 9 cycles (approximately 54 weeks) and was discontinued early for disease
 641 progression or unacceptable toxicity. The median age of the patients in the study was 71 years, 50% were male,
 642 88% were Caucasian and the median Karnofsky performance status score for the patients was 80. Patients had
 643 IgG/IgA/Light chain myeloma in 63%/25%/8% instances, a median hemoglobin of 105 g/l, and a median platelet
 644 count of 221.5 x 10⁹/l. Similar proportions of patients had creatinine clearance \leq 30 ml/min (3% in each arm).

645 At the time of a pre-specified interim analysis, the primary endpoint, time to progression, was met and patients in
 646 the M+P arm were offered Vc+M+P treatment. Median follow-up was 16.3 months. The final survival update was
 647 performed with a median duration of follow-up of 60.1 months. A statistically significant survival benefit in favour of
 648 the Vc+M+P treatment group was observed (HR=0.695; p=0.00043) despite subsequent therapies including
 649 Bortezomib -based regimens. Median survival for the Vc+M+P treatment group was 56.4 months compared to 43.1
 650 for the M+P treatment group. Efficacy results are presented in Table 11:

651 *Table 11: Efficacy results following the final survival update to VISTA study*

Efficacy endpoint	Vc+M+P n=344	M+P n=338
Time to progression		
Events n (%)	101 (29)	152 (45)
Median ^a (95% CI)	20.7 mo (17.6, 24,7)	15.0 mo (14.1, 17.9)

Hazard ratio ^b (95% CI)	0.54 (0.42, 0.70)	
p-value ^c	0.000002	
Progression-free survival		
Events n (%)	135 (39)	190 (56)
Median ^a (95% CI)	18.3 mo (16.6, 21.7)	14.0 mo (11.1, 15.0)
Hazard ratio ^b (95% CI)	0.61 (0.49, 0.76)	
p-value ^c	0.00001	
Overall survival* Events (deaths)		
n (%)	176 (51.2)	211 (62.4)
Median ^a (95% CI)	56.4 mo (52.8, 60.9)	43.1 mo (35.3, 48.3)
Hazard ratio ^b (95% CI)	0.695 (0.567, 0.852)	
p-value ^c	0.00043	
Response rate		
populatuion ^e n=668	n=337	n=331
CR ^f n(%)	102 (30)	12 (4)
PR ^f n(%)	136 (40)	103 (31)
nCR n(%)	5 (1)	0
CR+PR ^f n (%)	238 (71)	115 (35)
p-value ^d	<10 ⁻¹⁰	
Reduction in serum M-protein		
population ^g n=667	n=336	n=331
≥ 90% n (%)	151 (45)	34 (10)
Time to first response in CR + PR		
Median	1.4 mo	4.2 mo
Median^a response duration		
CR ^f	24.0 mo	12.8 mo
CR+PR ^f	19.9 mo	13.1 mo
Time to next therapy		
Events n (%)	224 (65.1)	260 (76.9)
Median ^a (95% CI)	27.0 mo (24.7, 31.1)	19.2 mo (17.0, 21.0)
Hazard ratio ^b (95% CI)	0.557 (0.462, 0.671)	
p-value ^c	< 0.000001	

652 ^a Kaplan-Meier estimate.

653 ^b Hazard ratio estimate is based on a Cox proportional-hazard model adjusted for stratification factors: β 2-
654 microglobulin, albumin, and region. A hazard ratio less than 1 indicates an advantage for VMP

655 ^c Nominal p-value based on the stratified log-rank test adjusted for stratification factors: β 2-microglobulin, albumin,
656 and region

657 ^d p-value for Response Rate (CR+PR) from the Cochran Mantel-Haenszel chi-square test adjusted for the
658 stratification factors

659 ^e Response population includes patients who had measurable disease at baseline

660 ^f CR=Complete Response; PR=Partial Response. EBMT criteria

661 ^g All randomised patients with secretory disease

662 ^h Survival update based on a median duration of follow-up at 60.1 month

663 mo: months

664 CI=Confidence Interval

665

666 *Patients eligible for stem cell transplantation*

667 (Ref.1: Velcade EU/1/04/274/001 page 22 - 24)

668 Two randomised, open-label, multicenter Phase III trials (IFM-2005-01, MMY-3010) were conducted to demonstrate
669 the safety and efficacy of Bortezomib in dual and triple combinations with other chemotherapeutic agents, as
670 induction therapy prior to stem cell transplantation in patients with previously untreated multiple myeloma.

671 In study IFM-2005-01 Bortezomib combined with dexamethasone [VcDx, n=240] was compared to vincristine-
672 doxorubicin-dexamethasone [VDDx, n=242]. Patients in the VcDx group received four 21 day cycles, each
673 consisting of Bortezomib (1.3 mg/m² administered intravenously twice weekly on days 1, 4, 8, and 11), and oral
674 dexamethasone (40 mg/day on days 1 to 4 and days 9 to 12, in Cycles 1 and 2, and on days 1 to 4 in Cycles 3
675 and 4).

676 Autologous stem cell transplants were received by 198 (82%) patients and 208 (87%) patients in the VDDx and
677 VcDx groups respectively; the majority of patients underwent one single transplant procedure. Patient demographic
678 and baseline disease characteristics were similar between the treatment groups. Median age of the patients in the
679 study was 57 years, 55% were male and 48% of patients had high-risk cytogenetics. The median duration of
680 treatment was 13 weeks for the VDDx group and 11 weeks for the VcDx group. The median number of cycles
681 received for both groups was 4 cycles.

682 The primary efficacy endpoint of the study was post-induction response rate (CR+nCR). A statistically significant
683 difference in CR+nCR was observed in favour of the Bortezomib combined with dexamethasone group. Secondary
684 efficacy endpoints included post-transplant response rates (CR+nCR, CR+nCR+VGPR+PR), Progression Free
685 Survival and Overall Survival. Main efficacy results are presented in Table 12.

686 *Table 12: Efficacy results from study IFM-2005-01*

Endpoints	VcDx	VDDx	OR; 95% CI; P value ^a
IFM-2005-01	N=240 (ITT population)	N=242 (ITT population)	
<i>RR (Post-induction)</i>			
*CR+nCR	14.6 (10.4, 19.7)	6.2 (3.5, 10.0)	2.58 (1.37, 4.85); 0.003

CR+nCR+VGPR+PR % (95% CI)	77.1 (71.2, 82.2)	60.7 (54.3, 66.9)	2.18 (1.46, 3.24); < 0.001
<i>RR (Post-transplant)^b</i>			
CR+nCR	37.5 (31.4, 44.0)	23.1 (18.0, 29.0)	1.98 (1.33, 2.95); 0.001
CR+nCR+VGPR+PR % (95% CI)	79.6 (73.9, 84.5)	74.4 (68.4, 79.8)	1.34 (0.87, 2.05); 0.179

687 CI=confidence interval; CR=complete response; nCR=near complete response; ITT=intent to treat; RR=response
688 rate; Vc= Bortezomib; VcDx= Bortezomib , dexamethasone; VDDx=vincristine, doxorubicin, dexamethasone;
689 VGPR=very good partial response; PR=partial response; OR=odds ratio.

690 * Primary endpoint

691 ^a OR for response rates based on Mantel-Haenszel estimate of the common odds ratio for stratified tables; p-values
692 by Cochran Mantel-Haenszel test.

693 ^b Refers to response rate after second transplant for subjects who received a second transplant (42/240 [18%] in
694 VcDx group and 52/242 [21%] in VDDx group).

695 Note: An OR > 1 indicates an advantage for Vc-containing induction therapy.

696
697 In study MMY-3010 induction treatment with Bortezomib combined with thalidomide and dexamethasone [VcTDx,
698 n=130] was compared to thalidomide-dexamethasone [TDx, n=127]. Patients in the VcTDx group received six 4-
699 week cycles, each consisting of BORTEZOMIB (1.3 mg/m² administered twice weekly days 1, 4, 8, and 11,
700 followed by a 17-day rest period from day 12 to day 28), dexamethasone (40 mg administered orally on days 1 to
701 4 and days 8 to 11), and thalidomide (administered orally at 50 mg daily on days 1-14, increased to 100 mg on
702 days 15-28 and thereafter to 200 mg daily).

703
704 One single autologous stem cell transplant was received by 105 (81%) patients and 78 (61%) patients in the
705 VcTDx and TDx groups, respectively. Patient demographic and baseline disease characteristics were similar
706 between the treatment groups. Patients in the VcTDx and TDx groups respectively had a median age of 57
707 versus 56 years, 99% versus 98% patients were Caucasians, and 58% versus 54% were males. In the VcTDx
708 group 12% of patients were cytogenetically classified as high risk versus 16% of patients in the TDx group. The
709 median duration of treatment was 24.0 weeks and the median number of treatment cycles received was 6.0, and
710 was consistent across treatment groups.

711
712 The primary efficacy endpoints of the study were post-induction and post-transplant response rates (CR+nCR). A
713 statistically significant difference in CR+nCR was observed in favour of the Bortezomib combined with
714 dexamethasone and thalidomide group. Secondary efficacy endpoints included Progression Free Survival and
715 Overall Survival. Main efficacy results are presented in Table 13.

716

717 *Table 13: Efficacy results from study MMY-3010*

Endpoints	VcTDx	TDx	OR; 95% CI; P value ^a
MMY-3010	N=130 (ITT population)	N=127 (ITT population)	
<i>*RR (Post-induction)</i>			
CR+nCR	49.2 (40.4, 58.1)	17.3 (11.2, 25.0)	4.63 (2.61, 8.22); < 0.001 ^a
CR+nCR+PR % (95% CI)	84.6 (77.2, 90.3)	61.4 (52.4, 69.9)	3.46 (1.90, 6.27); < 0.001 ^a
<i>*RR (Post-transplant)</i>			
CR+nCR	55.4 (46.4, 64.1)	34.6 (26.4, 43.6)	2.34 (1.42, 3.87); 0.001 ^a
CR+nCR+PR % (95% CI)	77.7 (69.6, 84.5)	56.7 (47.6, 65.5)	2.66 (1.55, 4.57); < 0.001 ^a

718 CI=confidence interval; CR=complete response; nCR=near complete response; ITT=intent to treat; RR=response
 719 rate; Vc= Bortezomib ; VcTDx= Bortezomib, thalidomide, dexamethasone; TDx=thalidomide, dexamethasone;
 720 PR=partial response; OR=odds ratio

721 * Primary endpoint

722 a OR for response rates based on Mantel-Haenszel estimate of the common odds ratio for stratified tables; p-
 723 values by Cochran Mantel-Haenszel test.

724 Note: An OR > 1 indicates an advantage for Vc-containing induction therapy

725

726 Clinical efficacy in relapsed or refractory multiple myeloma

727 (Ref.1: Velcade EU/1/04/274/001 page 24 - 25)

728 The safety and efficacy of Bortezomib (injected intravenously) were evaluated in 2 studies at the recommended
 729 dose of 1.3 mg/m²: a Phase III randomised, comparative study (APEX), versus dexamethasone (Dex), of 669
 730 patients with relapsed or refractory multiple myeloma who had received 1-3 prior lines of therapy, and a Phase II
 731 single-arm study of 202 patients with relapsed and refractory multiple myeloma, who had received at least 2 prior
 732 lines of treatment and who were progressing on their most recent treatment.

733 In the Phase III study, treatment with Bortezomib led to a significantly longer time to progression, a significantly
 734 prolonged survival and a significantly higher response rate, compared to treatment with dexamethasone (see Table
 735 14), in all patients as well as in patients who have received 1 prior line of therapy. As a result of a pre-planned
 736 interim analysis, the dexamethasone arm was halted at the recommendation of the data monitoring committee and
 737 all patients randomised to dexamethasone were then offered Bortezomib, regardless of disease status. Due to this
 738 early crossover, the median duration of follow-up for surviving patients is 8.3 months. Both in patients who were
 739 refractory to their last prior therapy and those who were not refractory, overall survival was significantly longer and
 740 response rate was significantly higher on the Bortezomib arm.

741 Of the 669 patients enrolled, 245 (37%) were 65 years of age or older. Response parameters as well as TTP
 742 remained significantly better for Bortezomib independently of age. Regardless of β 2- microglobulin levels at
 743 baseline, all efficacy parameters (time to progression and overall survival, as well as response rate) were
 744 significantly improved on the Bortezomib arm.

745 In the refractory population of the Phase II study, responses were determined by an independent review committee
 746 and the response criteria were those of the European Bone Marrow Transplant Group. The median survival of all
 747 patients enrolled was 17 months (range < 1 to 36+ months). This survival was greater than the six-to-nine month
 748 median survival anticipated by consultant clinical investigators for a similar patient population. By multivariate
 749 analysis, the response rate was independent of myeloma type, performance status, chromosome 13 deletion status,
 750 or the number or type of previous therapies. Patients who had received 2 to 3 prior therapeutic regimens had a
 751 response rate of 32% (10/32) and patients who received greater than 7 prior therapeutic regimens had a response
 752 rate of 31% (21/67).

753 *Table 14: Summary of disease outcomes from the Phase III (APEX) and Phase II studies*

	Phase III		Phase III		Phase III		Phase II
	All patients		1 prior line of therapy		> 1 prior line of therapy		\geq 2 prior lines
Time related events	Vc n=333 ^a	Dex n=336 ^a	Vc n=132 ^a	Dex n=119 ^a	Vc n=200 ^a	Dex n=217 ^a	Vc n=202 ^a
TTP, days [95% CI]	189 ^b [148, 211]	106 ^b [86, 128]	212 ^d [188, 267]	169 ^d [105, 191]	148 ^b [129, 192]	87 ^b [84, 107]	210 [154, 281]
1 year survival, % [95% CI]	80 ^d [74,85]	66 ^d [59,72]	89 ^d [82,95]	72 ^d [62,83]	72 ^d [62,83]	62 [53,71]	60
Best response (%)	Vc n=315 ^c	Dex n=312 ^c	Vc n=128	Dex n=110	Vc n=187	Dex n=202	Vc n=193
CR	20 (6) ^b	2 (< 1) ^b	8 (6)	2 (2)	12 (6)	0 (0)	(4)**
CR+nCR	41 (13) ^b	5 (2) ^b	16 (13)	4 (4)	25 (13)	1 (< 1)	(10)**
CR+nCR+PR	121 (38) ^b	56 (18) ^b	57 (45) ^d	29 (26) ^d	64 (34) ^b	27 (13) ^b	(27)**
CR+nCR+PR+MR	146 (46)	108 (35)	66 (52)	45 (41)	80 (43)	63 (31)	(35)**
Median duration Days (months)	242 (8.0)	169 (5.6)	246 (8.1)	189 (6.2)	238 (7.8)	126 (4.1)	385*
Time to response CR+PR (days)	43	43	44	46	41	27	38*

754 a Intent to Treat (ITT) population

755 b p-value from the stratified log-rank test; analysis by line of therapy excludes stratification for therapeutic history;
 756 p < 0.0001

757 c Response population includes patients who had measurable disease at baseline and received at least 1 dose of
758 study medicinal product.

759 d p-value from the Cochran Mantel-Haenszel chi-square test adjusted for the stratification factors; analysis by line
760 of therapy excludes stratification for therapeutic history

761 * CR+PR+MR **CR=CR, (IF-); nCR=CR (IF+)

762 NA=not applicable,

763 NE=not estimated

764 TTP-Time to Progression

765 CI=Confidence Interval

766 Vc= Bortezomib; Dex=dexamethasone

767 CR=Complete Response; nCR=near Complete response

768 PR=Partial Response; MR=Minimal response

769

770 In the Phase II study, patients who did not obtain an optimal response to therapy with Bortezomib alone were able
771 to receive high-dose dexamethasone in conjunction with Bortezomib. The protocol allowed patients to receive
772 dexamethasone if they had had a less than optimal response to Bortezomib alone. A total of 74 evaluable patients
773 were administered dexamethasone in combination with Bortezomib. Eighteen percent of patients achieved, or had
774 an improved response [MR (11%) or PR (7%)] with combination treatment.

775

776 *Clinical efficacy with subcutaneous administration of Bortezomib in patients with relapsed/refractory multiple myeloma*
777 [\(Ref.1: Velcade EU/1/04/274/001 page 25 - 26\)](#)

778 An open label, randomised, Phase III non-inferiority study compared the efficacy and safety of the subcutaneous
779 administration of Bortezomib versus the intravenous administration. This study included 222 patients with
780 relapsed/refractory multiple myeloma, who were randomised in a 2:1 ratio to receive 1.3 mg/m² of Bortezomib by
781 either the subcutaneous or intravenous route for 8 cycles. Patients who did not obtain an optimal response (less
782 than Complete Response [CR]) to therapy with Bortezomib alone after 4 cycles were allowed to receive
783 dexamethasone 20 mg daily on the day of and after Bortezomib administration. Patients with baseline Grade \geq 2
784 peripheral neuropathy or platelet counts < 50,000/ μ l were excluded. A total of 218 patients were evaluable for
785 response.

786 This study met its primary objective of non-inferiority for response rate (CR+PR) after 4 cycles of single agent
787 Bortezomib for both the subcutaneous and intravenous routes, 42% in both groups. In addition, secondary

788 response-related and time to event related efficacy endpoints showed consistent results for subcutaneous and
 789 intravenous administration (Table 15).

790 *Table 15: Summary of efficacy analyses comparing subcutaneous and intravenous administrations of Bortezomib*

	Bortezomib intravenous arm	Bortezomib subcutaneous arm
Response Evaluable Population	n=73	n=145
Response Rate at 4 cycles n (%) ORR (CR+PR)	31 (42)	61 (42)
p-value ^a	0.00201	
CR n (%)	6 (8)	9 (6)
PR n (%)	25 (34)	52 (36)
nCR n (%)	4 (5)	9 (6)
Response Rate at 8 cycles n (%) ORR (CR+PR)	38 (52)	76 (52)
p-value ^a	0.0001	
CR n (%)	9 (12)	15 (10)
PR n (%)	29 (40)	61 (42)
nCR n (%)	7 (10)	14 (10)
Intent to Treat Population ^b	n=74	N=148
TTP, months	9.4	10.4
(95% CI)	(7.6, 10.6)	(8.5, 11.7)
Hazard ratio (95% CI) ^c	0.839 (0.564, 1.249)	
p-value ^d	0.38657	
Progression Free Survival, months	8.0	10.2
(95% CI)	(6.7, 9.8)	(8.1, 10.8)
Hazard ratio (95% CI) ^c	0.824 (0.574, 1.183)	
p-value ^d	0.295	
1-year Overall Survival (%) ^e	76.7	72.6
(95% CI)	(64.1, 85.4)	(63.1, 80.0)

791 ^a p-value is for the non-inferiority hypothesis that the SC arm retains at least 60% of the response rate in the IV
 792 arm.

793 ^b 222 subjects were enrolled into the study; 221 subjects were treated with Bortezomib

794 ^c Hazards ratio estimate is based on a Cox model adjusted for stratification factors: ISS staging and number of
 795 prior lines.

796 ^d Log rank test adjusted for stratification factors: ISS staging and number of prior lines.

797 ^e Median duration of follow up is 11.8 months

798

799 *Bortezomib combination treatment with pegylated liposomal doxorubicin (study DOXIL-MMY-3001)*

800 (Ref.1: [Velcade EU/1/04/274/001 page 26 - 27](#))

801 A Phase III randomised, parallel-group, open-label, multicentre study was conducted in 646 patients comparing the
802 safety and efficacy of Bortezomib plus pegylated liposomal doxorubicin versus Bortezomib monotherapy in patients
803 with multiple myeloma who had received at least 1 prior therapy and who did not progress while receiving
804 anthracycline-based therapy. The primary efficacy endpoint was TTP while the secondary efficacy endpoints were
805 OS and ORR (CR+PR), using the European Group for Blood and Marrow Transplantation (EBMT) criteria.

806 A protocol-defined interim analysis (based on 249 TTP events) triggered early study termination for efficacy. This
807 interim analysis showed a TTP risk reduction of 45% (95% CI; 29-57%, $p < 0.0001$) for patients treated with
808 combination therapy of Bortezomib and pegylated liposomal doxorubicin. The median TTP was 6.5 months for the
809 Bortezomib monotherapy patients compared with 9.3 months for the Bortezomib plus pegylated liposomal
810 doxorubicin combination therapy patients. These results, though not mature, constituted the protocol defined final
811 analysis.

812 The final analysis for OS performed after a median follow-up of 8.6 years showed no significant difference in OS
813 between the two treatment arms. The median OS was 30.8 months (95% CI; 25.2- 36.5 months) for the Bortezomib
814 monotherapy patients and 33.0 months (95% CI; 28.9-37.1 months) for the Bortezomib plus pegylated liposomal
815 doxorubicin combination therapy patients.

816

817 *Bortezomib combination treatment with dexamethasone*

818 (Ref.1: [Velcade EU/1/04/274/001 page 27](#))

819 In the absence of any direct comparison between Bortezomib and Bortezomib in combination with dexamethasone
820 in patients with progressive multiple myeloma, a statistical matched-pair analysis was conducted to compare results
821 from the non 37inimizes37 arm of Bortezomib in combination with dexamethasone (Phase II open-label study MMY-
822 2045), with results obtained in the Bortezomib monotherapy arms from different Phase III 37inimizes37 studies
823 (M34101-039 [APEX] and DOXIL MMY-3001) in the same indication.

824 The matched-pair analysis is a statistical method in which patients in the treatment group (e.g. Bortezomib in
825 combination with dexamethasone) and patients in the comparison group (e.g. Bortezomib) are made comparable
826 with respect to confounding factors by individually pairing study subjects. This 37inimizes the effects of observed
827 confounders when estimating treatment effects using non-randomised data.

828 One hundred and twenty seven matched pairs of patients were identified. The analysis demonstrated improved
829 ORR (CR+PR) (odds ratio 3.769; 95% CI 2.045-6.947; $p < 0.001$), PFS (hazard ratio 0.511; 95% CI 0.309-0.845;
830 $p=0.008$), TTP (hazard ratio 0.385; 95% CI 0.212-0.698; $p=0.001$) for Bortezomib in combination with
831 dexamethasone over Bortezomib monotherapy.

832 Limited information on Bortezomib retreatment in relapsed multiple myeloma is available.

833 Phase II study MMY-2036 (RETRIEVE), single arm, open-label study was conducted to determine the efficacy and
834 safety of retreatment with Bortezomib . One hundred and thirty patients (≥ 18 years of age) with multiple myeloma
835 who previously had at least partial response on a Bortezomib – containing regimen were retreated upon
836 progression. At least 6 months after prior therapy, Bortezomib was started at the last tolerated dose of 1.3 mg/m²
837 (n=93) or ≤ 1.0 mg/m² (n=37) and given on days 1, 4, 8 and 11 every 3 weeks for maximum of 8 cycles either as
838 single agent or in combination with dexamethasone in accordance with the standard of care. Dexamethasone was
839 administered in combination with Bortezomib to 83 patients in Cycle 1 with an additional 11 patients receiving
840 dexamethasone during the course of Bortezomib retreatment cycles.

841 The primary endpoint was best confirmed response to retreatment as assessed by EBMT criteria. The overall best
842 response rate (CR + PR), to retreatment in 130 patients was 38.5% (95% CI: 30.1, 47.4).

843

844 Clinical efficacy in previously untreated mantle cell lymphoma (MCL)

845 (Ref.1: Velcade EU/1/04/274/001 page 27 - 28)

846 Study LYM-3002 was a Phase III, randomised, open-label study comparing the efficacy and safety of the
847 combination of Bortezomib , rituximab, cyclophosphamide, doxorubicin, and prednisone (VcRCAP; n=243) to that
848 of rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone (RCHOP; n=244) in adult patients with
849 previously untreated MCL (Stage II, III or IV). Patients in the VcR-CAP treatment arm received Bortezomib (1.3
850 mg/m²; on days 1, 4, 8, 11, rest period days 12- 21), rituximab 375 mg/m² IV on day 1; cyclophosphamide 750
851 mg/m² IV on day 1; doxorubicin 50 mg/m² IV on day 1; and prednisone 100 mg/m² orally on day 1 through day 5
852 of the 21 day Bortezomib treatment cycle. For patients with a response first documented at cycle 6, two additional
853 treatment cycles were given.

854 The primary efficacy endpoint was progression-free survival based on Independent Review Committee (IRC)
855 assessment. Secondary endpoints included, time to progression (TTP), time to next anti-lymphoma treatment (TNT),
856 duration of treatment free interval (TFI), overall response rate (ORR) and complete response (CR/CRu) rate, overall
857 survival (OS) and response duration.

858 The demographic and baseline disease characteristics were generally well balanced between the two treatment
859 arms: median patient age was 66 years, 74% were male, 66% were Caucasian and 32% Asian, 69% of patients
860 had a positive bone marrow aspirate and/or a positive bone marrow biopsy for MCL, 54% of patients had an
861 International Prognostic Index (IPI) score of ≥ 3 , and 76% had Stage IV disease. Treatment duration (median=17
862 weeks) and duration of follow-up (median=40 months) were comparable in both treatment arms. A median of 6
863 cycles was received by patients in both treatment arms with 14% of subjects in the VcR-CAP group and 17% of
864 patients in the R-CHOP group receiving 2 additional cycles. The majority of the patients in both groups completed
865 treatment, 80% in the VcR-CAP group and 82% in the R-CHOP group. Efficacy results are presented in Table 16:

866 **Table 16: Efficacy results from study LYM-3002**

Efficacy endpoint	VcR-CAP	R-CHOP	
n: ITT patients	243	244	
Progression free survival (IRC)^a			
Events n (%)	133 (54.7%)	165 (67.6%)	HR ^b (95% CI)=0.63
Median ^c (95% CI) (months)	24.7 (19.8; 31.8)	14.4 (12; 16.9)	(0.50; 0.79) p-value ^d < 0.001
Response rate			
n: response-evaluable patients	229	228	
Overall complete response (CR+CRu) ^f n(%)	122 (53.3%)	95 (41.7%)	OR ^e (95% CI)=1.688 (1.148; 2.481) p-value ^g =0.007
Overall response (CR+CRu+PR) ^h n(%)	211 (92.1%)	204 (89.5%)	OR ^e (95% CI)=1.428 (0.749; 2.722) p-value ^g =0.275

867 ^a Based on Independent Review Committee (IRC) assessment (radiological data only).

868 ^b Hazard ratio estimate is based on a Cox's model stratified by IPI risk and stage of disease. A hazard ratio < 1
869 indicates an advantage for VcR-CAP.

870 ^c Based on Kaplan-Meier product limit estimates.

871 ^d Based on Log rank test stratified with IPI risk and stage of disease.

872 ^e Mantel-Haenszel estimate of the common odds ratio for stratified tables is used, with IPI risk and stage of disease
873 as stratification factors. An odds ratio (OR) > 1 indicates an advantage for VcR-CAP.

874 ^f Include all CR+CRu, by IRC, bone marrow and LDH.

875 ^g P-value from the Cochran Mantel-Haenszel chi-square test, with IPI and stage of disease as stratification factors.

876 ^h Include all radiological CR+CRu+PR by IRC regardless the verification by bone marrow and LDH.

877 CR=Complete Response; CRu=Complete Response unconfirmed; PR=Partial Response; CI=Confidence Interval,
878 HR=Hazard Ratio; OR=Odds Ratio; ITT=Intent to Treat

879

880 Median PFS by investigator assessment was 30.7 months in the VcR-CAP group and 16.1 months in the R-CHOP
881 group (Hazard Ratio [HR]=0.51; p < 0.001). A statistically significant benefit (p < 0.001) in favour of the VcR-CAP
882 treatment group over the R-CHOP group was observed for TTP (median 30.5 versus 16.1 months), TNT (median
883 44.5 versus 24.8 months) and TFI (median 40.6 versus 20.5 months). The median duration of complete response
884 was 42.1 months in the VcR-CAP group compared with 18 months in the R-CHOP group. The duration of overall
885 response was 21.4 months longer in the VcR-CAP group (median 36.5 months versus 15.1 months in the R-CHOP

886 group). The final analysis for OS was performed after a median follow-up of 82 months. Median OS was 90.7
887 months for the VcR-CAP group compared with 55.7 months for the R-CHOP group (HR=0.66; p=0.001). The
888 observed final median difference in the OS between the 2 treatment groups was 35 months.

889

890 Patients with previously treated light-chain (AL) Amyloidosis

891 (Ref.1: Velcade EU/1/04/274/001 page 28)

892 An open label non randomised Phase I/II study was conducted to determine the safety and efficacy of Bortezomib
893 in patients with previously treated light-chain (AL) Amyloidosis. No new safety concerns were observed during the
894 study, and in particular Bortezomib did not exacerbate target organ damage (heart, kidney and liver). In an
895 exploratory efficacy analysis, a 67.3% response rate (including a 28.6% CR rate) as measured by hematologic
896 response (M-protein) was reported in 49 evaluable patients treated with the maximum allowed doses of 1.6 mg/m²
897 weekly and 1.3 mg/m² twice-weekly. For these dose cohorts, the combined 1-year survival rate was 88.1%.

898

899 Paediatric population

900 (Ref.1: Velcade EU/1/04/274/001 page 29)

901 The European Medicines Agency has waived the obligation to submit the results of studies with Bortezomib in all
902 subsets of the paediatric population in multiple myeloma and in mantle cell lymphoma (see section 4.2 for
903 information on paediatric use).

904 A Phase II, single-arm activity, safety, and pharmacokinetic trial conducted by the Children's Oncology Group
905 assessed the activity of the addition of bortezomib to multi-agent re-induction chemotherapy in paediatric and young
906 adult patients with lymphoid malignancies (pre-B cell acute lymphoblastic leukemia [ALL], T-cell ALL, and T-cell
907 lymphoblastic lymphoma [LL]). An effective re-induction multi-agent chemotherapy regimen was administered in 3
908 blocks. Bortezomib was administered only in Blocks 1 and 2 to avoid potential overlapping toxicities with
909 coadministered drugs in Block 3.

910 Complete response (CR) was evaluated at the end of Block 1. In B-ALL patients with relapse within 18 months of
911 diagnosis (n = 27) the CR rate was 67% (95% CI: 46, 84); the 4-month event free survival rate was 44% (95% CI:
912 26, 62). In B-ALL patients with relapse 18-36 months from diagnosis (n = 33) the CR rate was 79% (95% CI: 61,
913 91) and the 4-month event free survival rate was 73% (95% CI: 54, 85). The CR rate in first-relapsed T-cell ALL
914 patients (n = 22) was 68% (95% CI: 45, 86) and the 4-month event free survival rate was 67% (95% CI: 42, 83).
915 The reported efficacy data are considered inconclusive (see section 4.2).

916 There were 140 patients with ALL or LL enrolled and evaluated for safety; median age was 10 years (range 1 to
917 26). No new safety concerns were observed when Bortezomib was added to the standard pediatric pre B cell ALL
918 chemotherapy backbone. The following adverse reactions (Grade \geq 3) were observed at a higher incidence in the
919 BORTEZOMIB containing treatment regimen as compared with a historical control study in which the backbone

920 regimen was given alone: in Block 1 peripheral sensory neuropathy (3% versus 0%); ileus (2.1% versus 0%);
921 hypoxia (8% versus 2%). No information on possible sequelae or rates of peripheral neuropathy resolution were
922 available in this study. Higher incidences were also noted for infections with Grade ≥ 3 neutropenia (24% versus
923 19% in Block 1 and 22% versus 11% in Block 2), increased ALT (17% versus 8% in Block 2), hypokalaemia (18%
924 versus 6% in Block 1 and 21% versus 12% in Block 2) and hyponatraemia (12% versus 5% in Block 1 and 4%
925 versus 0 in Block 2).

926

927 **5.2 Pharmacokinetic properties**

928 (Ref.1: Velcade EU/1/04/274/001 page 29 - 30)

929

930 Absorption

931 Following intravenous bolus administration of a 1.0 mg/m² and 1.3 mg/m² dose to 11 patients with multiple myeloma
932 and creatinine clearance values greater than 50 ml/min, the mean first-dose maximum plasma concentrations of
933 bortezomib were 57 and 112 ng/ml, respectively . In subsequent doses, mean maximum observed plasma
934 concentrations ranged from 67 to 106 ng/ml for the 1.0 mg/m² dose and 89 to 120 ng/ml for the 1.3 mg/m² dose.
935 Following an intravenous bolus or subcutaneous injection of a 1.3 mg/m² dose to patients with multiple myeloma
936 (n= 14 in the intravenous group, n= 17 in the subcutaneous group), the total systemic exposure after repeat dose
937 administration (AUC_{last}) was equivalent for subcutaneous and intravenous administrations . The C_{max} after
938 subcutaneous administration. (20.4 ng/ml (was lower than intravenous 223 ng/ml) The AUC_{last} geometric mean
939 ratio was 0.99 and 90 %confidence intervals were 80.18% - 122.80%.

940

941 Distribution

942 The mean distribution volume (Vd) of bortezomib ranged from 1,659 L to 3,294 L following single -or repeated-
943 dose intravenous administration of 1.0 mg/m² or 1.3 mg m²to patients with multiple myeloma . This suggests that
944 bortezomib distributes widely to peripheral tissues . Over a bortezomib concentration range of 0.01 to 1.0 µg/ml,
945 the in vitro protein binding averaged 82.9% in human plasma .The fraction of bortezomib bound to plasma proteins
946 was not concentration-dependent.

947

948 Biotransformation

949 In vitro studies with human liver microsomes and human cDNA-expressed cytochrome P450 isozymes indicate that
950 bortezomib is primarily oxidatively metabolised via cytochrome P450 enzymes, 3A4, 2C19, and 1A2 .The major
951 metabolic pathway is deboronation to form two deboronated metabolites that subsequently undergo hydroxylation
952 to several metabolites .Deboronated-bortezomib metabolites are inactive as 26S proteasome inhibitors.

953

954 Elimination

955 The mean elimination half-life ($t_{1/2}$) of bortezomib upon multiple dosing ranged from 40 -193 hours .Bortezomib is
956 eliminated more rapidly following the first dose compared to subsequent doses . Mean total body clearances were
957 102 and 112 l/h following the first dose for doses of 1.0 mg/m² and 1.3 mg/m² respectively, and ranged from 15 to
958 32 l/h and 18 to 32 l/h following subsequent doses for doses of 1.0 mg/m² and 1.3 mg/m², respectively.

959

960 Special populations

961 [\(Ref.1: Velcade EU/1/04/274/001 page 30\)](#)

962 *Hepatic impairment*

963 The effect of hepatic impairment on the pharmacokinetics of bortezomib was assessed in a Phase I study during
964 the first treatment cycle, including 61 patients primarily with solid tumors and varying degrees of hepatic impairment
965 at bortezomib doses ranging from 0.5 to 1.3 mg/m².

966 When compared to patients with normal hepatic function, mild hepatic impairment did not alter dose -normalised
967 bortezomib AUC . However, the dose-normalised mean AUC values were increased by approximately 60 % in
968 patients with moderate or severe hepatic impairment . A lower starting dose is recommended in patients with
969 moderate or severe hepatic impairment, and those patients should be closely monitored. (see section 4 .2, Table
970 6)

971

972 *Renal impairment*

973 A pharmacokinetic study was conducted in patients with various degrees of renal impairment who were classified
974 according to their creatinine clearance values (CrCL) into the following groups : Normal (CrCL \geq 60 ml/min/1.73
975 m², n=12) Mild = (CrCL 40-59 ml/min/1.73 m², n=10), Moderate (CrCL =20-39 ml/min/1.73 m², n=9), and Severe
976 (CrCL < 20 ml/min/1.73 m², n=3). A group of dialysis patients who were dosed after dialysis was also included in
977 the study)n=8 .(Patients were administered intravenous doses of 0.7 to 1.3 mg/m² of Bortezomib twice weekly .
978 Exposure of Bortezomib) dose-normalised AUC and C_{max} (was comparable among all the groups (see section
979 4.2).

980

981 *Age*

982 The pharmacokinetics of bortezomib were characterized following twice weekly intravenous bolus administration of
983 1.3 mg/m² doses to 104 pediatric patients (2-16 years old) with acute lymphoblastic leukemia (ALL) or acute
984 myeloid leukemia (AML). Based on a population pharmacokinetic analysis, clearance of bortezomib increased with
985 increasing body surface area (BSA). Geometric mean (%CV) clearance was 7.79 (25%) L/hr/m², volume of
986 distribution at steady-state was 834 (39%) L/m², and the elimination half-life was 100 (44%) hours .After correcting
987 for the BSA effect, other demographics such as age, body weight and sex did not have clinically significant effects
988 on bortezomib clearance .BSA-normalized clearance of bortezomib in pediatric patients was similar to that observed
989 in adults.

990

991 **5.3 Preclinical safety data**

992 [\(Ref.1: Velcade EU/1/04/274/001 page 30\)](#)

993 Bortezomib was positive for clastogenic activity (structural chromosomal aberrations) in the in vitro chromosomal
994 aberration assay using Chinese hamster ovary (CHO) cells at concentrations as low as 3.125 µg/ml, which was
995 the lowest concentration evaluated .Bortezomib was not genotoxic when tested in the in vitro mutagenicity assay
996 (Ames assay) and in vivo micronucleus assay in mice.

997 Developmental toxicity studies in the rat and rabbit have shown embryo-fetal lethality at maternally toxic doses, but
998 no direct embryo-foetal toxicity below maternally toxic doses .Fertility studies were not performed but evaluation of
999 reproductive tissues has been performed in the general toxicity studies . In the 6-month rat study, degenerative
1000 effects in both the testes and the ovary have been observed . It is, therefore, likely that bortezomib could have a
1001 potential effect on either male or female fertility .

1002 In multi-cycle general toxicity studies conducted in the rat and monkey, the principal target organs included the
1003 gastrointestinal tract, resulting in vomiting and/or diarrhoea; haematopoietic and lymphatic tissues, resulting in
1004 peripheral blood cytopenias, lymphoid tissue atrophy and haematopoietic bone marrow hypocellularity; peripheral
1005 neuropathy (observed in monkeys, mice and dogs) involving sensory nerve axons; and mild changes in the kidneys .
1006 All these target organs have shown partial to full recovery following discontinuation of treatment.

1007 Cardiovascular safety pharmacology studies in monkeys and dogs show that intravenous doses approximately two
1008 to three times the recommended clinical dose on a mg/m² basis are associated with increases in heart rate,
1009 decreases in contractility, hypotension and death . In dogs, the decreased cardiac contractility and hypotension
1010 responded to acute intervention with positive inotropic or pressor agents .Moreover, in dog studies, a slight increase
1011 in the corrected QT interval was observed.

1012

1013 **6 .Pharmaceutical particulars**

1014 **6.1 List of excipients**

1015 Mannitol

1016 Nitrogen

1017

1018 **6.2 Incompatibilities**

1019 This medicinal product must not be mixed with other medicinal products except those mentioned in section 6.6.

1020

1021 **6.3 Shelf life**

1022 Unopened vial

1023 2 years

1024 Reconstituted solution

1025 ([Ref.1: Velcade EU/1/04/274/001 page 31](#))

1026 The reconstituted solution should be used immediately after preparation . If not used immediately, in-use storage
1027 times and conditions prior to use are the responsibility of the user . However, the chemical and physical in-use
1028 stability of the reconstituted solution has been demonstrated for 8 hours at 25°C stored in the original vial and/or
1029 a syringe . The total storage time for the reconstituted medicinal product should not exceed 8 hours prior to
1030 administration.

1031

1032 **6.4 Special precautions for storage**

1033 Do not store above 30°C.

1034 Keep the vial in the outer carton in order to protect from light.

1035

1036 **6.5 Nature and contents of container**

1037 Type I glass 10 ml-vial with a grey bromobutyl stopper and an aluminium seal, with a PP disc containing 3.5 mg
1038 bortezomib.

1039 Each pack contains 1 single-use vial.

1040

1041 **6.6 Special precautions for disposal and other handling**

1042 General precautions

1043 Bortezomib is a cytotoxic agent . Therefore, caution should be used during handling and preparation of Bortezomib .
1044 Use of gloves and other protective clothing to prevent skin contact is recommended.

1045 **Aseptic technique** must be strictly observed throughout the handling of Bortezomib, since it contains no
1046 preservative.

1047 There have been fatal cases of inadvertent intrathecal administration of Bortezomib . Bortezomib 1 mg powder for
1048 solution for injection is for intravenous use only, while Bortezomib 3.5 mg powder for solution for injection is for
1049 intravenous or subcutaneous use . Bortezomib should not be administered intrathecally.

1050

1051 Instructions for reconstitution

1052 BORTEZOMIB must be reconstituted by a healthcare professional.

1053 *Intravenous injection*

1054 Each 10 ml vial of Bortezomib must be carefully reconstituted with 3.5 ml of sodium chloride 9 mg/ml (0.9%)
1055 solution for injection, by using a syringe of the appropriate size, without removing the vial stopper . Dissolution of
1056 the lyophilised powder is completed in less than 2 minutes.

1057 After reconstitution, each ml solution contains 1 mg bortezomib .The reconstituted solution is clear and colourless,
1058 with a final pH of 4 to 7.

1059 The reconstituted solution must be inspected visually for particulate matter and discolouration prior to
1060 administration .If any discolouration or particulate matter is observed, the reconstituted solution must be discarded.

1061 *Subcutaneous injection*

1062 Each 10 ml vial of BORTEZOMIB must be carefully reconstituted with 1.4 ml of sodium chloride 9 mg/ml (0.9%)
1063 solution for injection, by using a syringe of the appropriate size, without removing the vial stopper .Dissolution of
1064 the lyophilised powder is completed in less than 2 minutes.

1065 After reconstitution, each ml solution contains 2.5 mg bortezomib .The reconstituted solution is clear and colourless,
1066 with a final pH of 4 to 7 .The reconstituted solution must be inspected visually for particulate matter and
1067 discolouration prior to administration .If any discolouration or particulate matter is observed, the reconstituted
1068 solution must be discarded.

1069

1070 Disposal

1071 Bortezomib is for single use only. Any unused medicinal product or waste material should be disposed of in
1072 accordance with local requirements.

1073

1074 **7. Marketing Authorization Holder**

1075 Imported by APL Pharma Thai Ltd, Bangkok

1076 Manufactured by:

1077 Eugia Pharma Specialities Limited,

1078 Survey No .550, 551 & 552, Kolthur Village,

1079 Shameerpet Mandal,

1080 Medchal-Malkajgiri District, Telangana,

1081 India.

1082

1083 **8. Marketing Authorization Number:** 1C...../.....(NG)

1084

1085 **9. Date of authorization:**

1086

1087 **10. Date of revision of the text:** [September 24, 2021](#)