

เอกสารกำกับยาภาษาอังกฤษ

SUMMARY OF PRODUCT CHARACTERISTICS

DECINJET

^{Rx} Decitabine 50 mg Powder for Concentrate for Solution for Infusion (Single Dose Vial)

1. Name of the medicinal product

1.1 Product Name: DECINJET

1.2 Strength: Decitabine 50 mg

1.3 Pharmaceutical Dosage Form: Powder for Concentrate for Solution for Infusion

2. Qualitative and quantitative composition

Each vial of powder for concentrate for solution for infusion contains 50 mg Decitabine.

After reconstitution with 10 ml of water for injections, each ml of concentrate contains 5 mg of Decitabine.

Excipients with known effect

Each vial contains 0.29 mmol sodium

For the full list of excipients see section 6.1.

3. Pharmaceutical form

Powder for concentrate for solution for infusion (powder for infusion)

White to off-white lyophilized cake or powder

4. Clinical particulars

4.1 Therapeutic indications

Decitabine powder for concentrate for solution for infusion is indicated for the treatment of adult patients with newly diagnosed *de novo* or secondary acute myeloid leukaemia (AML), according to the World Health Organisation (WHO) classification, who are not candidates for standard induction chemotherapy.

4.2 Posology and method of administration

26 Decitabine powder for concentrate for solution for infusion administration must be initiated under
27 the supervision of physicians experienced in the use of chemotherapeutic medicinal products.

28 Posology

29 In a treatment cycle, Decitabine is administered at a dose of 20 mg/m² body surface area by
30 intravenous infusion over 1 hour repeated daily for 5 consecutive days (i.e., a total of 5 doses
31 per treatment cycle). The total daily dose must not exceed 20 mg/m² and the total dose per
32 treatment cycle must not exceed 100 mg/m². If a dose is missed, treatment should be
33 resumed as soon as possible. The cycle should be repeated every 4 weeks depending on the
34 patient's clinical response and observed toxicity. It is recommended that patients be treated for
35 a minimum of 4 cycles; however, a complete or partial remission may take longer than 4
36 cycles to be obtained. Treatment may be continued as long as the patient shows response,
37 continues to benefit or exhibits stable disease, i.e., in the absence of overt progression.

38 If after 4 cycles, the patient's haematological values (e.g., platelet counts or absolute
39 neutrophil count), have not returned to pre-treatment levels or if disease progression occurs
40 (peripheral blast counts are increasing or bone marrow blast counts are worsening), the
41 patient may be considered to be a non-responder and alternative therapeutic options to
42 Decitabine powder for concentrate for solution for infusion should be considered.

43 Pre-medication for the prevention of nausea and vomiting is not routinely recommended but
44 may be administered if required.

45 *Management of myelosuppression and associated complications*

46 Myelosuppression and adverse events related to myelosuppression (thrombocytopenia,
47 anaemia, neutropenia, and febrile neutropenia) are common in both treated and untreated
48 patients with AML. Complications of myelosuppression include infections and bleeding.
49 Treatment may be delayed at the discretion of the treating physician, if the patient
50 experiences myelosuppression-associated complications, such as those described below:

- 51 • Febrile neutropenia (temperature $\geq 38.5^{\circ}\text{C}$ and absolute neutrophil count $< 1,000/\mu\text{L}$)
- 52 • Active viral, bacterial or fungal infection (i.e., requiring intravenous anti-infectives or extensive
53 supportive care)
- 54 • Haemorrhage (gastrointestinal, genito-urinary, pulmonary with platelets $< 25,000/\mu\text{L}$
55 or any central nervous system haemorrhage)

56 Treatment with Decitabine powder for concentrate for solution for infusion may be resumed

57 once these conditions have improved or have been stabilised with adequate treatment (anti-
58 infective therapy, transfusions, or growth factors).

59 In clinical studies, approximately one-third of patients receiving Decitabine required a dose-
60 delay. Dose reduction is not recommended.

61 *Paediatric population*

62 Decitabine should not be used in children with AML aged < 18 years, because efficacy was
63 not established. Currently available data are described in sections 4.8, 5.1, and 5.2.

64 *Hepatic impairment*

65 Studies in patients with hepatic impairment have not been conducted. The need for dose
66 adjustment in patients with hepatic impairment has not been evaluated. If worsening hepatic
67 function occurs, patients should be carefully monitored (see sections 4.4 and 5.2).

68 *Renal impairment*

69 Studies in patients with renal impairment have not been conducted. The need for dose
70 adjustment in patients with renal impairment has not been evaluated (see section 4.4 and 5.2).

71 Method of administration

72 Decitabine powder for concentrate for solution for infusion is administered by intravenous
73 infusion. A central venous catheter is not required.

74 For instructions on reconstitution and dilution of the medicinal product before administration,
75 see section 6.6.

76 **4.3 Contraindications**

77 Hypersensitivity to Decitabine or to any of the excipients, listed in section 6.1.

78 Breast-feeding (see section 4.6)

79 **4.4 Special warnings and precautions for use**

80 Myelosuppression

81 Myelosuppression and complications of myelosuppression, including infections and bleeding
82 that occur in patients with AML may be exacerbated with Decitabine treatment. Therefore,
83 patients are at increased risk for severe infections (due to any pathogen such as bacterial,
84 fungal and viral), with potentially fatal outcome (see section 4.8). Patients should be monitored
85 for signs and symptoms of infection and treated promptly.

86 In clinical studies, the majority of patients had baseline Grade 3-4 myelosuppression. In
87 patients with baseline Grade 2 abnormalities, worsening of myelosuppression was seen in
88 most patients and more frequently than in patients with baseline Grade 1 or 0 abnormalities.
89 Myelosuppression caused by Decitabine is reversible. Complete blood and platelet counts
90 should be performed regularly, as clinically indicated and prior to each treatment cycle. In the
91 presence of myelosuppression or its complications, treatment with Decitabine may be
92 interrupted and/or supportive measures instituted (see sections 4.2 and 4.8).

93 Respiratory, thoracic and mediastinal disorders

94 Cases of interstitial lung disease (ILD) (including pulmonary infiltrates, organising pneumonia
95 and pulmonary fibrosis) without signs of infectious aetiology have been reported in patients
96 receiving Decitabine. Careful assessment of patients with an acute onset or unexplained
97 worsening of pulmonary symptoms should be performed to exclude ILD. If ILD is confirmed,
98 appropriate treatment should be initiated (see section 4.8).

99 Hepatic impairment

100 Use in patients with hepatic impairment has not been established. Caution should be exercised
101 in the administration of Decitabine to patients with hepatic impairment and in patients who
102 develop signs or symptoms of hepatic impairment. Liver function tests should be performed prior
103 to initiation of therapy and prior to each treatment cycle, and as clinically indicated (see
104 sections 4.2 and 5.2).

105 Renal impairment

106 Use in patients with severe renal impairment has not been studied. Caution should be
107 exercised in the administration of Decitabine to patients with severe renal impairment
108 (Creatinine Clearance [CrCl] < 30 ml/min). Renal function tests should be performed prior to
109 initiation of therapy and prior to each treatment cycle, and as clinically indicated (see section
110 4.2).

111 Cardiac disease

112 Patients with a history of severe congestive heart failure or clinically unstable cardiac disease
113 were excluded from clinical studies and therefore, the safety and efficacy of Decitabine in these
114 patients has not been established. Cases of cardiomyopathy with cardiac decompensation, in
115 some cases reversible after treatment discontinuation, dose reduction or corrective treatment,
116 have been reported in the postmarketing setting. Patients, especially those with cardiac disease
117 history, should be monitored for signs and symptoms of heart failure.

118

119 Differentiation syndrome

120 Cases of differentiation syndrome (also known as retinoic acid syndrome) have been reported
121 in patients receiving Decitabine. Differentiation syndrome may be fatal (see section 4.8).
122 Treatment with high-dose IV corticosteroids and haemodynamic monitoring should be
123 considered at first onset of symptoms or signs suggestive of differentiation syndrome.
124 Temporary discontinuation of Decitabine should be considered until resolution of symptoms and
125 if resumed, caution is advised.

126

127 Excipients

128 This medicine contains 0.5 mmol potassium per vial. After reconstitution and dilution of the
129 solution for intravenous infusion, this medicine contains less than 1 mmol (39 mg) of potassium
130 per dose, i.e. essentially 'potassium-free'.

131 This medicine contains 0.29 mmol (6.67 mg) sodium per vial. After reconstitution and dilution of
132 the solution for intravenous infusion, this medicine contains between 13.8 mg-138 mg (0.6-6
133 mmol) sodium per dose (depending on the infusion fluid for dilution), equivalent to 0.7-7% of
134 the WHO recommended maximum daily intake of 2 g sodium for an adult.

135 **4.5 Interaction with other medicinal products and other forms of interaction**

136 No formal clinical drug interaction studies with Decitabine have been conducted. There is the
137 potential for a drug-drug interaction with other agents which are also activated by sequential
138 phosphorylation (via intracellular phosphokinase activities) and/or metabolised by enzymes
139 implicated in the inactivation of Decitabine (e.g., cytidine deaminase). Therefore, caution should
140 be exercised if these active substances are combined with Decitabine.

141 Impact of co-administered medicinal products on Decitabine

142 Cytochrome (CYP) 450-mediated metabolic interactions are not anticipated as Decitabine
143 metabolism is not mediated by this system but by oxidative deamination.

144

145 Impact of Decitabine on co-administered medicinal products

146 Given its low in vitro plasma protein binding (< 1%), Decitabine is unlikely to displace co-
147 administered medicinal products from their plasma protein binding. Decitabine has been shown
148 to be a weak inhibitor of P-gp mediated transport in vitro and is therefore, also not expected to
149 affect P-gp mediated transport of co-administered medicinal products (see section 5.2)

150 **4.6 Fertility, pregnancy and lactation**

151 Women of childbearing potential/Contraception in men and women

152 Due to the genotoxic potential of Decitabine (see section 5.3), women of childbearing potential
153 must use effective contraceptive measures and avoid becoming pregnant while being treated
154 with Decitabine and for 6 months following completion of treatment. Men should use effective
155 contraceptive measures and be advised to not father a child while receiving Decitabine, and for
156 3 months following completion of treatment (see section 5.3).

157 The use of Decitabine with hormonal contraceptives has not been studied.

158 Pregnancy

159 There are no adequate data on the use of Decitabine in pregnant women. Studies have shown
160 that Decitabine is teratogenic in rats and mice (see section 5.3). The potential risk for humans
161 is unknown. Based on results from animal studies and its mechanism of action, Decitabine
162 should not be used during pregnancy and in women of childbearing potential not using effective
163 contraception. A pregnancy test should be performed on all women of childbearing potential
164 before treatment is started. If Decitabine is used during pregnancy, or if a patient becomes
165 pregnant while receiving this medicinal product, the patient should be apprised of the potential
166 hazard to the foetus.

167 Breast-feeding

168 It is not known whether Decitabine or its metabolites are excreted in breast milk. Decitabine is
169 contraindicated during breast-feeding; therefore, if treatment with this medicine is required,
170 breast-feeding must be discontinued (see section 4.3).

171 Fertility

172 No human data on the effect of Decitabine on fertility are available. In non-clinical animal
173 studies, Decitabine alters male fertility and is mutagenic. Because of the possibility of infertility
174 as a consequence of Decitabine therapy, men should seek advice on conservation of sperm
175 and female patients of childbearing potential should seek consultation regarding oocyte
176 cryopreservation prior to initiation of treatment.

177 **4.7 Effects on ability to drive and use machines**

178 Decitabine has moderate influence on the ability to drive and use machines. Patients should be
179 advised that they may experience undesirable effects such as anaemia during treatment.

180 Therefore, caution should be recommended when driving a car or operating machines.

181 **4.8 Undesirable effects**

182 Summary of the safety profile

183 The most common adverse drug reactions ($\geq 35\%$) reported are pyrexia, anaemia and
184 thrombocytopaenia.

185 The most common Grade 3-4 adverse drug reactions ($\geq 20\%$) included pneumonia,
186 thrombocytopaenia, neutropaenia, febrile neutropaenia and anaemia.

187 In clinical studies, 30% of patients treated with Decitabine and 25% of patients treated in the
188 comparator arm had adverse events with an outcome of death during treatment or within 30
189 days after the last dose of study drug.

190 In the Decitabine treatment group, there was a higher incidence of treatment discontinuation
191 due to adverse events in women compared to men (43% versus 32%).

192 Tabulated list of adverse drug reactions

193 Adverse drug reactions reported in 293 AML patients treated with Decitabine are summarised in Table
194 1. The following table reflects data from AML clinical studies and from post-marketing experience. The
195 adverse drug reactions are listed by frequency category. Frequency categories are defined as follows:
196 Very common ($\geq 1/10$), common ($\geq 1/100$ to $< 1/10$), uncommon ($\geq 1/1,000$ to $< 1/100$), rare (\geq
197 $1/10,000$ to $< 1/1,000$), very rare ($< 1/10,000$), not known (frequency cannot be estimated from the
198 available data).

199 Within each frequency grouping, adverse drug reactions are presented in order of decreasing
200 seriousness.

201

Table 1: Adverse drug reactions identified with Decitabine				
System Organ Class	Frequency (all Grades)	Adverse Drug Reaction	Frequency	
			All Grades^a (%)	Grades 3-4^a (%)
Infections and infestations	Very common	pneumonia*	24	20
		urinary tract infection*	15	7
		All other infections (viral,	63	39

		bacterial, fungal)*, b, c, d		
	Common	septic shock*	6	4
		sepsis*	9	8
		sinusitis	3	1
Neoplasms benign, malignant and unspecified (incl. cysts and polyps)	Not known	differentiation syndrome	Not known	Not known
Blood and lymphatic disorders	Very common	febrile neutropaenia*	34	32
		neutropaenia*	32	30
		thrombocytopaenia*, e	41	38
		anaemia	38	31
		leukopaenia	20	18
	Uncommon	pancytopaenia*	< 1	< 1
Immune system disorders	Common	hypersensitivity including anaphylactic reaction ^f	1	< 1
Metabolism and nutrition disorders	Very common	hyperglycaemia	13	3
Nervous system disorders	Very common	headache	16	1
Cardiac disorders	Uncommon	cardiomyopathy	< 1	< 1
Respiratory, thoracic and mediastinal disorders	Very common	epistaxis	14	2
	Not known	interstitial lung disease	Not known	Not known
Gastrointestinal disorders	Very common	diarrhoea	31	2
		vomiting	18	1
		nausea	33	< 1
	Common	stomatitis	7	1
	Not known	enterocolitis, including neutropaenic colitis, caecitis*	Not known	Not known
Hepatobiliary disorders	Very common	hepatic function abnormal	11	3
	Common	hyperbilirubinaemia ^g	5	< 1

Skin and subcutaneous tissue disorders	Uncommon	acute febrile neutrophilic dermatosis (Sweet's syndrome)	< 1	NA
General disorders and administration site conditions	Very common	pyrexia	48	9
<p>^a Worst National Cancer Institute Common Terminology Criteria for Adverse Events Grade.</p> <p>^b Excluding pneumonia, urinary tract infection, sepsis, septic shock and sinusitis.</p> <p>^c The most frequently reported "other infections" in study DACO-016 were: oral herpes, oral candidiasis, pharyngitis, upper respiratory tract infection, cellulitis, bronchitis, nasopharyngitis.</p> <p>^d Including enterocolitis infectious.</p> <p>^e Including haemorrhage associated with thrombocytopaenia, including fatal cases.</p> <p>^f Including preferred terms hypersensitivity, drug hypersensitivity, anaphylactic reaction, anaphylactic shock, anaphylactoid reaction, anaphylactoid shock.</p> <p>^g In clinical studies in AML and myelodysplastic syndrome (MDS), the reporting frequency for hyperbilirubinaemia was 11% for All Grades and 2% for Grade 3-4</p> <p>* Includes events with a fatal outcome. NA = Not applicable</p>				

202 Description of selected adverse drug reactions

203 *Haematologic adverse drug reactions*

204 The most commonly reported haematologic adverse drug reactions associated with
205 Decitabine treatment included febrile neutropaenia, thrombocytopaenia, neutropaenia,
206 anaemia and leukopaenia.

207 Serious bleeding-related adverse drug reactions, some of which lead to a fatal outcome,
208 such as central nervous system (CNS) haemorrhage (2%) and gastrointestinal (GI)
209 haemorrhage (2%), in the context of severe thrombocytopaenia, were reported in patients
210 receiving Decitabine.

211 Haematological adverse drug reactions should be managed by routine monitoring of
212 complete blood counts and early administration of supportive treatments as required.
213 Supportive treatments include, administration of prophylactic antibiotics and/or growth
214 factor support (e.g., G-CSF) for neutropaenia and transfusions for anaemia or
215 thrombocytopaenia according to institutional guidelines. For situations where Decitabine

216 administration should be delayed, see section 4.2.

217 *Infections and infestations adverse drug reactions*

218 Serious infection-related adverse drug reactions, with potentially fatal outcome, such as
219 septic shock, sepsis, pneumonia, and other infections (viral, bacterial and fungal) were
220 reported in patients receiving Decitabine.

221 *Gastrointestinal disorders*

222 Occurrences of enterocolitis, including neutropaenic colitis, caecitis have been reported
223 during treatment with Decitabine. Enterocolitis may lead to septic complications and may
224 be associated with fatal outcome.

225 *Respiratory, thoracic and mediastinal disorders*

226 Cases of interstitial lung disease (including pulmonary infiltrates, organising pneumonia and
227 pulmonary fibrosis) without signs of infectious aetiology have been reported in patients
228 receiving Decitabine.

229 *Differentiation syndrome*

230 Cases of differentiation syndrome (also known as retinoic acid syndrome) have been reported
231 in patients receiving Decitabine. Differentiation syndrome may be fatal and symptoms and
232 clinical findings include respiratory distress, pulmonary infiltrates, fever, rash, pulmonary
233 oedema, peripheral oedema, rapid weight gain, pleural effusions, pericardial effusions,
234 hypotension and renal dysfunction. Differentiation syndrome may occur with or without
235 concomitant leucocytosis. Capillary leak syndrome and coagulopathy can also occur (see
236 section 4.4).

237 *Paediatric population*

238 The safety assessment in paediatric patients is based on the limited safety data from a Phase
239 I/II study to evaluate pharmacokinetics, safety and efficacy of Decitabine in paediatric patients
240 (aged 1 to 14 years) with relapsed or refractory AML (n = 17) (see section 5.1). No new safety
241 signal was observed in this paediatric study.

242 Reporting of suspected adverse reactions

243 Reporting suspected adverse reactions after authorisation of the medicinal product is
244 important. It allows continued monitoring of the benefit/risk balance of the medicinal
245 product. Healthcare professionals are asked to report any suspected adverse reactions to
246 Thai FDA Health Product Vigilance Center (adr@fda.moph.go.th)

247 **4.9 Overdose**

248 There is no direct experience of human overdose and no specific antidote. However, early
249 clinical study data in published literature at doses greater than 20 times higher than the
250 current therapeutic dose, reported increased myelosuppression including prolonged
251 neutropaenia and thrombocytopaenia. Toxicity is likely to manifest as exacerbations of
252 adverse drug reactions, primarily myelosuppression. Treatment for overdose should be
253 supportive.

254 **5. Pharmacological properties**

255 **5.1 Pharmacodynamic properties**

256 Pharmacotherapeutic group: Antineoplastic agents, antimetabolites, pyrimidine analogues;
257 ATC Code: L01BC08

258 Mechanism of action

259 Decitabine (5-aza-2'-deoxycytidine) is a cytidine deoxynucleoside analogue that selectively
260 inhibits DNA methyltransferases at low doses, resulting in gene promoter hypomethylation
261 that can result in reactivation of tumour suppressor genes, induction of cellular
262 differentiation or cellular senescence followed by programmed cell death.

263

264 Clinical experience

265 The use of Decitabine was studied in an open-label, randomised, multicentre Phase III
266 study (DACO-016) in subjects with newly diagnosed *de novo* or secondary AML according
267 to the WHO classification. Decitabine (n = 242) was compared to treatment choice (TC, n =
268 243) which consisted of patient's choice with physician's advice of either supportive care
269 alone (n = 28, 11.5%) or 20 mg/m² cytarabine subcutaneously once daily for 10
270 consecutive days repeated every 4 weeks (n = 215, 88.5%). Decitabine was administered
271 as a 1-hour intravenous infusion of 20 mg/m² once daily for 5 consecutive days repeated
272 every 4 weeks.

273

274 Subjects who were considered candidates for standard induction chemotherapy were not
275 included in the study as shown by the following baseline characteristics. The median age
276 for the intent-to-treat (ITT) population was 73 years (range 64 to 91 years). Thirty-six
277 percent of subjects had poor-risk cytogenetics at baseline. The remainder of the subjects

278 had intermediate-risk cytogenetics. Patients with favourable cytogenetics were not included
279 in the study. Twenty-five percent of subjects had an ECOG performance status ≥ 2 .
280 Eighty-one percent of subjects had significant comorbidities (e.g., infection, cardiac
281 impairment, pulmonary impairment). The number of patients treated with Decitabine by
282 racial group was White 209 (86.4%) and Asian 33 (13.6%).

283

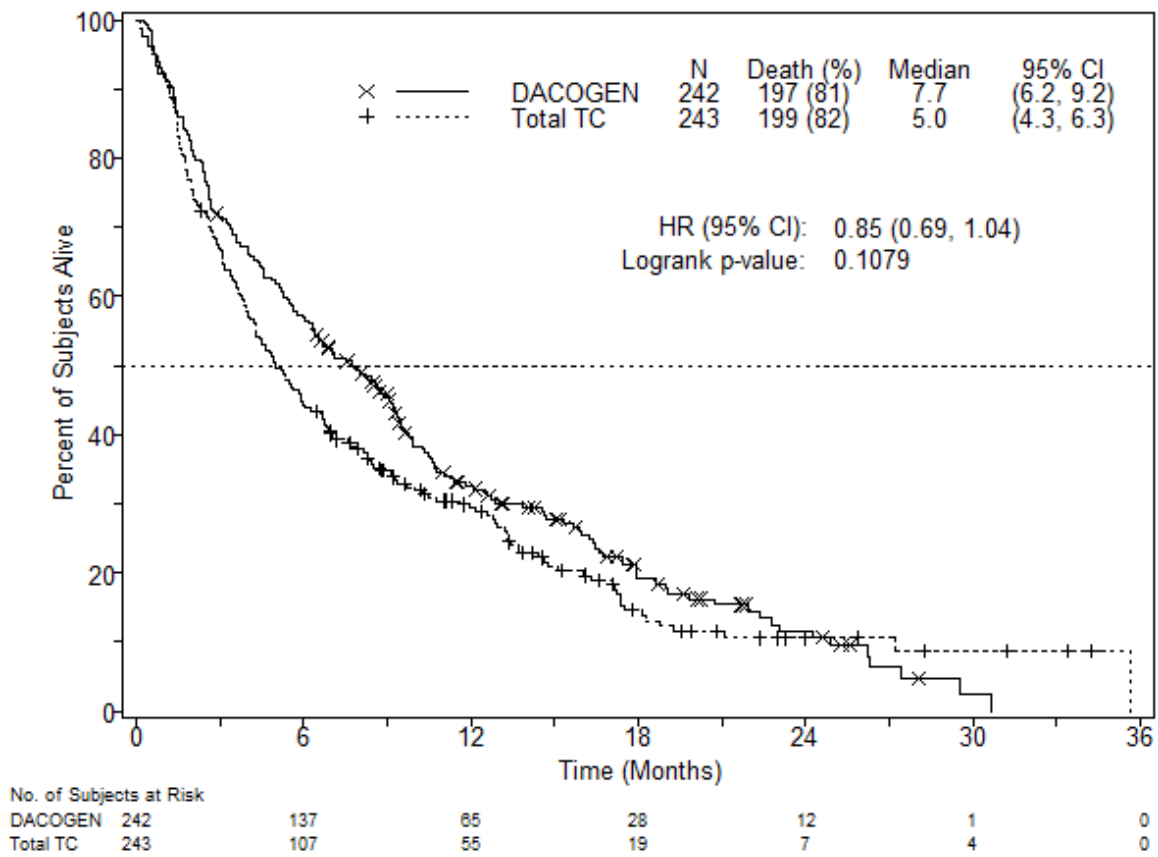
284 The primary endpoint of the study was overall survival. The secondary endpoint was
285 complete remission rate that was assessed by independent expert review. Progression-free
286 survival and Event-free survival were tertiary endpoints.

287

288 The median overall survival in the ITT population was 7.7 months in subjects treated with
289 Decitabine compared to 5.0 months for subjects in the TC arm (hazard ratio 0.85; 95% CI:
290 0.69, 1.04, $p = 0.1079$). The difference did not reach statistical significance, however, there
291 was a trend for improvement in survival with a 15% reduction in the risk of death for
292 subjects in the Decitabine arm (Figure 1). When censored for potentially disease modifying
293 subsequent therapy (i.e., induction chemotherapy or hypomethylating agent) the analysis
294 for overall survival showed a 20% reduction in the risk of death for subjects in the
295 Decitabine arm [HR = 0.80, (95% CI: 0.64, 0.99), p -value = 0.0437].

296

297 **Figure 1. Overall survival (ITT population).**

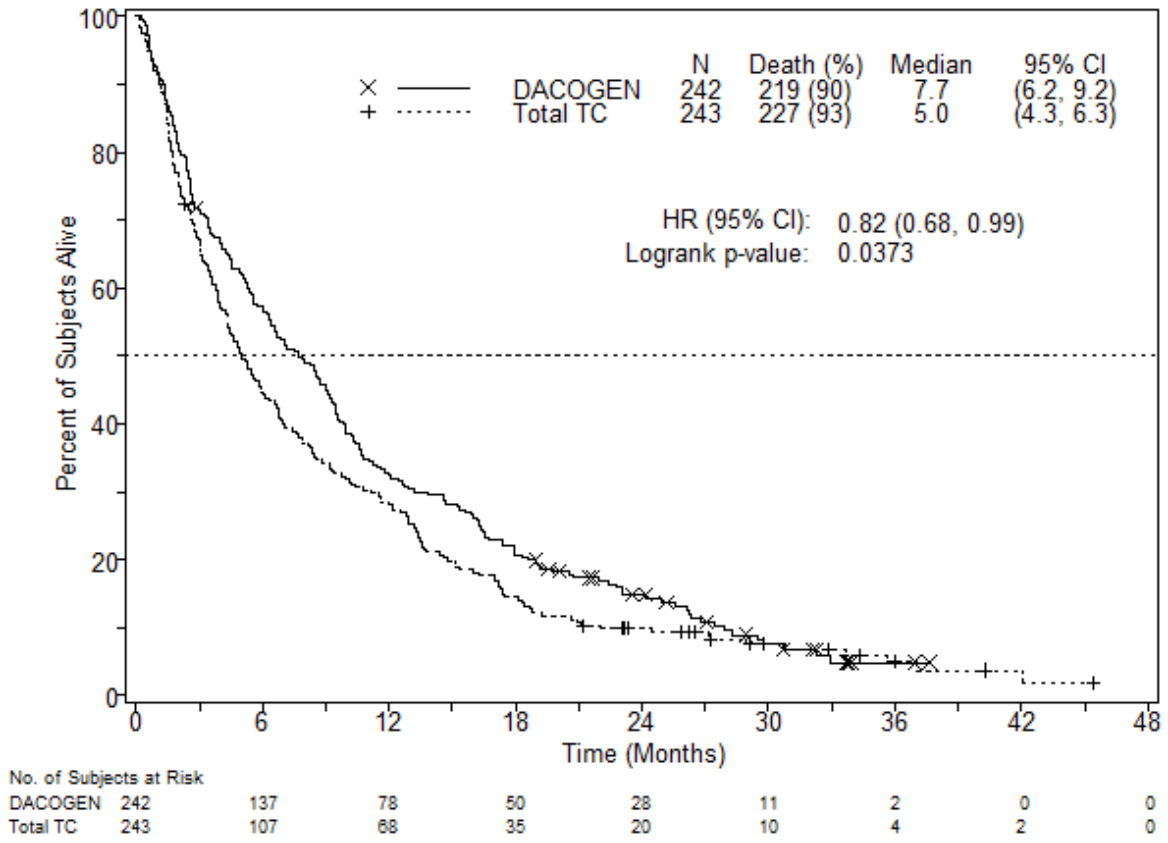


298

299 In an analysis with an additional 1 year of mature survival data, the effect of Decitabine on
 300 overall survival demonstrated a clinical improvement compared to the TC arm (7.7 months vs.
 301 5.0 months, respectively, hazard ratio = 0.82, 95% CI: 0.68, 0.99, nominal p-value = 0.0373,
 302 Figure 2).

303

304 **Figure 2. Analysis of mature overall survival data (ITT population).**



305

306 Based on the initial analysis in the ITT population, a statistically significant difference in
 307 complete remission rate (CR + CRp) was achieved in favour of subjects in the Decitabine arm,
 308 17.8% (43/242) compared to the TC arm, 7.8% (19/243); treatment difference 9.9% (95% CI:
 309 4.07; 15.83), p = 0.0011. The median time to best response and median duration of best
 310 response in patients who achieved a CR or CRp were 4.3 months and 8.3 months,
 311 respectively. Progression-free survival was significantly longer for subjects in the Decitabine
 312 arm, 3.7 months (95% CI: 2.7, 4.6) compared with subjects in the TC arm, 2.1 months (95%
 313 CI: 1.9, 3.1); hazard ratio 0.75 (95% CI: 0.62, 0.91), p = 0.0031. These results as well as
 314 other endpoints are shown in Table 2.

315

Outcomes	Decitabine n = 242	TC (combined group) n = 243	p-value
CR + CRp	43 (17.8%)	19 (7.8%)	0.0011
	OR = 2.5 (1.40, 4.78) ^b		

CR	38 (15.7%)	18 (7.4%)	-
EFS ^a	3.5 (2.5, 4.1) ^b	2.1 (1.9, 2.8) ^b	0.0025
	HR = 0.75 (0.62, 0.90) ^b		
PFS ^a	3.7 (2.7, 4.6) ^b	2.1 (1.9, 3.1) ^b	0.0031
	HR = 0.75 (0.62, 0.91) ^b		
CR = complete remission; CRp = complete remission with incomplete platelet recovery, EFS = event-free survival, PFS = progression-free survival, OR = odds ratio, HR = hazard ratio - = Not evaluable			
^a Reported as median months			
^b 95% confidence intervals			

316

317 Overall survival and complete remission rates in pre-specified disease-related sub-groups (i.e.,
318 cytogenetic risk, Eastern Cooperative Oncology Group [ECOG] score, age, type of AML, and
319 baseline bone marrow blast count) were consistent with results for the overall study population.

320

321 The use of Decitabine as initial therapy was also evaluated in an open-label, single-arm, Phase II
322 study (DACO-017) in 55 subjects > 60 years with AML according to the WHO classification. The
323 primary endpoint was complete remission (CR) rate that was assessed by independent expert review.
324 The secondary endpoint of the study was overall survival. Decitabine was administered as a 1-hour
325 intravenous infusion of 20 mg/m² once daily for 5 consecutive days repeated every 4 weeks. In the
326 ITT analysis, a CR rate of 23.6% (95% CI: 13.2, 37) was observed in 13/55 subjects treated with
327 Decitabine. The median time to CR was 4.1 months, and the median duration of CR was 18.2
328 months. The median overall survival in the ITT population was 7.6 months (95% CI: 5.7, 11.5).

329

330 The efficacy and safety of Decitabine has not been evaluated in patients with acute promyelocytic
331 leukaemia or CNS leukaemia.

332

333 Paediatric population

334 A Phase I/II open-label, multicentre study evaluated the safety and efficacy of Decitabine in
 335 sequential administration with cytarabine in children aged 1 month to < 18 years with
 336 relapsed or refractory AML. A total of 17 subjects were enrolled and received Decitabine
 337 20 mg/m² in this study, of which 9 subjects received cytarabine 1 g/m² and 8 subjects
 338 received cytarabine administered at the maximum tolerable dose of 2 g/m². All subjects
 339 discontinued the study treatment. The reasons for treatment discontinuation included
 340 disease progression (12 [70.6%] subjects), subjects proceeding to transplant (3 [17.6%]),
 341 investigator decision (1 [5.9%]), and “other” (1 [5.9%]). Reported adverse events were
 342 consistent with the known safety profile of Decitabine in adults (see section 4.8). Based on
 343 these negative results, Decitabine should not be used in children with AML aged < 18
 344 years, because efficacy was not established (see section 4.2).

345 5.2 Pharmacokinetic properties

346 The population pharmacokinetic (PK) parameters of Decitabine were pooled from 3 clinical
 347 studies in 45 patients with AML or myelodysplastic syndrome (MDS) utilizing the 5-Day
 348 regimen. In each study, Decitabine PK was evaluated on the fifth day of the first treatment
 349 cycle.

350 Distribution

351 The pharmacokinetics of Decitabine following intravenous administration as a 1-hour infusion
 352 were described by a linear two-compartment model, characterised by rapid elimination from the
 353 central compartment and by relatively slow distribution from the peripheral compartment. For a
 354 typical patient (weight 70 kg/body surface area 1.73 m²) the Decitabine pharmacokinetic
 355 parameters are listed in the Table 3 below.

Table 3: Summary of population PK analysis for a typical patient receiving daily 1-hour infusions of Decitabine 20 mg/m² over 5 days every 4 weeks		
Parameter ^a	Predicted Value	95% CI
C _{max} (ng/ml)	107	88.5 – 129
AUC _{cum} (ng.h/ml)	580	480 – 695
t _{1/2} (min)	68.2	54.2 - 79.6
Vd _{SS} (L)	116	84.1 – 153
CL (L/h)	298	249 – 359
^a The total dose per cycle was 100 mg/m ²		

356 Decitabine exhibits linear PK and following the intravenous infusion, steady-state concentrations
357 are reached within 0.5 hour. Based on model simulation, PK parameters were independent of
358 time (i.e., did not change from cycle to cycle) and no accumulation was observed with this
359 dosing regimen. Plasma protein binding of Decitabine is negligible (< 1%). Decitabine V_{dSS} in
360 cancer patients is large indicating distribution into peripheral tissues. There was no evidence of
361 dependencies on age, creatinine clearance, total bilirubin, or disease.

362 Biotransformation

363 Intracellularly, Decitabine is activated through sequential phosphorylation via phosphokinase
364 activities to the corresponding triphosphate, which is then incorporated by the DNA
365 polymerase. *In vitro* metabolism data and the human mass balance study results indicated
366 that the cytochrome P450 system is not involved in the metabolism of Decitabine. The primary
367 route of metabolism is likely through deamination by cytidine deaminase in the liver, kidney,
368 intestinal epithelium and blood. Results from the human mass-balance study showed that
369 unchanged Decitabine in plasma accounted for approximately 2.4% of total radioactivity in
370 plasma. The major circulating metabolites are not believed to be pharmacologically active. The
371 presence of these metabolites in urine together with the high total body clearance and low
372 urinary excretion of unchanged Decitabine in the urine (~4% of the dose) indicate that
373 Decitabine is appreciably metabolized *in vivo*. *In vitro* studies show that Decitabine does not
374 inhibit nor induce CYP 450 enzymes up to more than 20-fold of the therapeutic maximum
375 observed plasma concentration (C_{max}). Thus; CYP-mediated metabolic drug interactions are
376 not anticipated, and Decitabine is unlikely to interact with agents metabolized through these
377 pathways. In addition, *in vitro* data show that Decitabine is a poor P-gp substrate.

378 Elimination

379 Mean plasma clearance following intravenous administration in cancer subjects was > 200 L/h
380 with moderate inter- subject variability (coefficient of variation [CV] is approximately 50%).
381 Excretion of unchanged drug appears to play only a minor role in the elimination of Decitabine.

382 Results from a mass balance study with radioactive ^{14}C -Decitabine in cancer patients showed
383 that 90% of the administered dose of Decitabine (4% unchanged drug) is excreted in the urine.

384 Additional information on special populations

385 The effects of renal or hepatic impairment, gender, age or race on the pharmacokinetics of
386 Decitabine have not been formally studied. Information on special populations was derived from
387 pharmacokinetic data from the 3 studies noted above, and from one Phase I study in MDS

388 subjects, (N = 14; 15 mg/m² x 3-hours q8h x 3 days)

389 *Elderly*

390 Population pharmacokinetic analysis showed that Decitabine pharmacokinetics are not
391 dependent on age (range studied 40 to 87 years; median 70 years).

392

393 *Paediatric population*

394 Population PK analysis of Decitabine showed that after accounting for body size, there is no
395 difference between Decitabine PK parameters in paediatric AML patients versus adults with
396 AML or MDS.

397 *Gender*

398 Population pharmacokinetic analysis of Decitabine did not show any clinically relevant
399 difference between men and women.

400 *Race*

401 Most of the patients studied were Caucasian. However, the population pharmacokinetic analysis
402 of Decitabine indicated that race had no apparent effect on the exposure to Decitabine.

403 *Hepatic impairment*

404 The PK of Decitabine have not been formally studied in patients with hepatic impairment.
405 Results from a human mass- balance study and *in vitro* experiments mentioned above
406 indicated that the CYP enzymes are unlikely to be involved in the metabolism of Decitabine. In
407 addition, the limited data from the population PK analysis indicated no significant PK
408 parameter dependencies on total bilirubin concentration despite a wide range of total bilirubin
409 levels. Thus, Decitabine exposure is not likely to be affected in patients with impaired hepatic
410 function.

411 *Renal impairment*

412 The PK of Decitabine have not been formally studied in patients with renal insufficiency. The
413 population PK analysis on the limited Decitabine data indicated no significant PK parameter
414 dependencies on normalized creatinine clearance, an indicator of renal function. Thus,
415 Decitabine exposure is not likely to be affected in patients with impaired renal function.

416 **5.3 Preclinical safety data**

417 Formal carcinogenicity studies have not been performed with Decitabine. Evidence from the

418 literature indicates that Decitabine has carcinogenic potential. The available data from *in vitro*
419 and *in vivo* studies provide sufficient evidence that Decitabine has genotoxic potential. Data
420 from the literature also indicate that Decitabine has adverse effects on all aspects of the
421 reproductive cycle, including fertility, embryo-foetal development and post-natal development.
422 Multi-cycle repeat-dose toxicity studies in rats and rabbits indicated that the primary toxicity
423 was myelosuppression, including effects on bone marrow, which was reversible on cessation
424 of treatment. Gastrointestinal toxicity was also observed and in males, testicular atrophy which
425 did not reverse over the scheduled recovery periods. Decitabine administration to
426 neonatal/juvenile rats showed a comparable general toxicity profile as in older rats.
427 Neurobehavioural development and reproductive capacity were unaffected when
428 neonatal/juvenile rats were treated at dose levels inducing myelosuppression. See section 4.2
429 for information on paediatric use.

430 **6. Pharmaceutical particulars**

431 **6.1 List of excipients**

432 Potassium dihydrogen phosphate

433 Sodium Hydroxide

434 Acetonitrile HP

435 Sodium hydroxide

436 Hydrochloric acid concentrated

437 Water for Injection

438 Nitrogen

439 **6.2 Incompatibilities**

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

442 **6.3 Shelf life**

443 Unopened vial

444 2 years.

445 Reconstituted and diluted solution

446 Within 15 minutes of reconstitution, the concentrate (in 10 ml of sterile water for injections)

447 must be further diluted with cold (2°C - 8°C) infusion fluids. This prepared diluted solution for
448 intravenous infusion can be stored at 2°C - 8°C for up to a maximum of 3 hours, followed by
449 up to 1 hour at room temperature (20°C - 25°C) before administration.

450 From a microbiological point of view, the product should be used within the time period
451 recommended above. It is the responsibility of the user to follow the recommended storage
452 times and conditions and ensure that reconstitution has taken place in aseptic conditions.

453 **6.4 Special precautions for storage**

454 Store below 30°C.

455 For storage conditions of the reconstituted and diluted medicinal product, see section 6.3.

456 **6.5 Nature and contents of container**

457 Tubular Type-I, clear glass vial (20R ISO Vial) stoppered with 20 mm grey bromobutyl rubber
458 stopper and sealed with 20 mm aluminum seal having Sky blue color polypropylene disc.

459 **6.6 Special precautions for disposal and other handling**

460 Recommendations for safe handling

461 Skin contact with the solution should be avoided and protective gloves must be worn. Standard
462 procedures for dealing with cytotoxic medicinal products should be adopted.

463 Reconstitution procedure

464 The powder should be aseptically reconstituted with 10 ml of water for injections. Upon
465 reconstitution, each ml contains approximately 5 mg of Decitabine at pH 6.7 to 7.3. Within 15
466 minutes of reconstitution, the solution must be further diluted with cold infusion fluids (sodium
467 chloride 9 mg/ml [0.9%] solution for injection or 5% glucose solution for injection) to a final
468 concentration of 0.15 to 1.0 mg/ml. For the shelf-life and the precaution for storage after
469 reconstitution, see section 6.3.

470 Decitabine should not be infused through the same intravenous access/line with other medicinal
471 products.

472 Disposal

473 This medicinal product is for single use only. Any unused medicinal product or waste material
474 should be disposed of in accordance with local requirements.

475 **7. Marketing Authorization Holder**

476 Imported by:

477 APL Pharma Thai Ltd

478 438 Phattanakarn 30, Phattanakarn Road,

479 Suanluang Subdistrict, Suanluang District,

480 Bangkok, Thailand 10250

481 Manufactured by:

482 Eugia Pharma Specialities Limited,

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

484 Shameerpet Mandal,

485 Medchal-Malkajgiri District,

486 Telangana, India.

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

488 **9. Date of authorization:**

489 **10. Date of revision of the text: March 1, 2023**