

PEXARITE 100 mg, 500 mg powder for concentrate for solution for infusion

Summary of Product Characteristics Updated 6-July-2022 | APL Pharma Thai Company

1. Name of the medicinal product

PEXARITE 100 mg powder for concentrate for solution for infusion

PEXARITE 500 mg powder for concentrate for solution for infusion

2. Qualitative and quantitative composition

PEXARITE 100 mg powder for concentrate for solution for infusion

Each vial contains 100 mg of pemetrexed (as pemetrexed disodium).

Excipient with known effect

Each vial contains approximately 11 mg sodium.

PEXARITE 500 mg powder for concentrate for solution for infusion

Each vial contains 500 mg of pemetrexed (as pemetrexed disodium).

Excipient with known effect

Each vial contains approximately 54 mg sodium.

After reconstitution (see section 6.6), each vial contains 25 mg/ml of pemetrexed.

For the full list of excipients see section 6.1.

3. Pharmaceutical form

Powder for concentrate for solution for infusion.

White to either light yellow or green-yellow lyophilised powder.

4. Clinical particulars

4.1 Therapeutic indications

Malignant pleural mesothelioma

PEXARITE in combination with cisplatin is indicated for the treatment of chemotherapy naïve patients with unresectable malignant pleural mesothelioma.

Non-small cell lung cancer

PEXARITE in combination with cisplatin is indicated for the first line treatment of patients with locally advanced or metastatic non-small cell lung cancer other than predominantly squamous cell histology (see section 5.1).

PEXARITE is indicated as monotherapy for the maintenance treatment of locally advanced or metastatic non-small cell lung cancer other than predominantly squamous cell histology in patients whose disease has not progressed immediately following platinum-based chemotherapy (see section 5.1).

PEXARITE is indicated as monotherapy for the second line treatment of patients with locally advanced or metastatic non-small cell lung cancer other than predominantly squamous cell histology (see section 5.1).

4.2 Posology and method of administration

Posology

PEXARITE must only be administered under the supervision of a physician qualified in the use of anti-cancer chemotherapy.

PEXARITE in combination with cisplatin

The recommended dose of PEXARITE is 500 mg/m² of body surface area (BSA) administered as

42 an intravenous infusion over 10 minutes on the first day of each 21-day cycle. The recommended
 43 dose of cisplatin is 75 mg/m² BSA infused over two hours approximately 30 minutes after
 44 completion of the Pemetrexed infusion on the first day of each 21-day cycle. Patients must
 45 receive adequate anti-emetic treatment and appropriate hydration prior to and/or after receiving
 46 cisplatin

47 (see also cisplatin Summary of Product Characteristics for specific dosing advice).

48 *PEXARITE as single agent*

49 In patients treated for non-small cell lung cancer after prior chemotherapy, the recommended
 50 dose of PEXARITE is 500 mg/m² BSA administered as an intravenous infusion over 10 minutes
 51 on the first day of each 21-day cycle.

52 *Pre-medication regimen*

53 To reduce the incidence and severity of skin reactions, a corticosteroid should be given the day
 54 prior to, on the day of, and the day after Pemetrexed administration. The corticosteroid should be
 55 equivalent to 4 mg of dexamethasone administered orally twice a day (see section 4.4).

56 To reduce toxicity, patients treated with pemetrexed must also receive vitamin supplementation
 57 (see section 4.4). Patients must take oral folic acid or a multivitamin containing folic acid (350 to
 58 1000 micrograms) on a daily basis. At least five doses of folic acid must be taken during the
 59 seven days preceding the first dose of pemetrexed, and dosing must continue during the full
 60 course of therapy and for 21 days after the last dose of pemetrexed. Patients must also receive
 61 an intramuscular injection of vitamin B₁₂ (1000 micrograms) in the week preceding the first dose
 62 of pemetrexed and once every three cycles thereafter. Subsequent vitamin B₁₂ injections may be
 63 given on the same day as pemetrexed.

64 *Monitoring*

65 Patients receiving pemetrexed should be monitored before each dose with a complete blood
 66 count, including a differential white cell count (WCC) and platelet count. Prior to each
 67 chemotherapy administration blood chemistry tests should be collected to evaluate renal and
 68 hepatic function. Before the start of any cycle of chemotherapy, patients are required to have the
 69 following: absolute neutrophil count (ANC) should be ≥ 1500 cells/mm³ and platelets should be ≥
 70 100,000 cells/mm³.

71 Creatinine clearance should be ≥ 45 ml/min.

72 The total bilirubin should be ≤ 1.5 times upper limit of normal. Alkaline phosphatase (AP),
 73 aspartate aminotransferase (AST or SGOT) and alanine aminotransferase (ALT or SGPT) should
 74 be ≤ 3 times upper limit of normal. Alkaline phosphatase, AST and ALT ≤ 5 times upper limit of
 75 normal is acceptable if liver has tumour involvement.

76 *Dose adjustments*

77 Dose adjustments at the start of a subsequent cycle should be based on nadir haematologic
 78 counts or maximum non-haematologic toxicity from the preceding cycle of therapy. Treatment
 79 may be delayed to allow sufficient time for recovery. Upon recovery patients should be retreated
 80 using the guidelines in Tables 1, 2 and 3, which are applicable for PEXARITE used as a single
 81 agent or in combination with cisplatin.

Table 1 - Dose modification table for PEXARITE (as single agent or in combination) and cisplatin – Haematologic toxicities	
Nadir ANC < 500 /mm ³ and nadir platelets ≥ 50,000 /mm ³	75 % of previous dose (both PEXARITE and cisplatin)
Nadir platelets < 50,000 /mm ³ regardless of nadir ANC	75 % of previous dose (both PEXARITE and cisplatin)
Nadir platelets < 50,000/mm ³ with bleeding ^a , regardless of nadir ANC	50% of previous dose (both PEXARITE and cisplatin)

82 ^a These criteria meet the National Cancer Institute Common Toxicity Criteria (CTC v2.0; NCI
83 1998) definition of \geq CTC Grade 2 bleeding

84 If patients develop non-haematologic toxicities \geq Grade 3 (excluding neurotoxicity), PEXARITE
85 should be withheld until resolution to less than or equal to the patient's pre-therapy value.
86 Treatment should be resumed according to the guidelines in Table 2.

Table 2: Dose modification table for PEXARITE (as single agent or in combination) and cisplatin– Non-haematologic toxicities ^{a, b}		
	Dose of pemetrexed (mg/m²)	Dose for cisplatin (mg/m²)
Any Grade 3 or 4 toxicities except mucositis	75 % of previous dose	75 % of previous dose
Any diarrhoea requiring hospitalisation (irrespective of grade) or grade 3 or 4 diarrhoea.	75 % of previous dose	75 % of previous dose
Grade 3 or 4 mucositis	50 % of previous dose	100 % of previous dose

87

88 ^a National Cancer Institute Common Toxicity Criteria (CTC v2.0; NCI 1998) ^b Excluding
89 neurotoxicity

90 In the event of neurotoxicity, the recommended dose adjustment for PEXARITE and cisplatin is
91 documented in Table 3. Patients should discontinue therapy if Grade 3 or 4 neurotoxicity is
92 observed.

Table 3 - Dose modification table for PEXARITE (as single agent or in combination) and cisplatin – Neurotoxicity		
CTC ^a Grade	Dose of pemetrexed (mg/m²)	Dose for cisplatin (mg/m²)
0 – 1	100 % of previous dose	100 % of previous dose
2	100 % of previous dose	50 % of previous dose

93

94 ^a National Cancer Institute Common Toxicity Criteria (CTC v2.0; NCI 1998)

95 Treatment with PEXARITE should be discontinued if a patient experiences any haematologic or
96 non-haematologic Grade 3 or 4 toxicity after 2 dose reductions or immediately if Grade 3 or 4
97 neurotoxicity is observed.

98 Special populations

99 *Elderly*

100 In clinical studies, there has been no indication that patients 65 years of age or older are at
101 increased risk of adverse events compared to patients younger than 65 years old. No dose
102 reductions other than those recommended for all patients are necessary.

103 *Paediatric population*

104 There is no relevant use of PEXARITE in the paediatric population in malignant pleural
105 mesothelioma and non-small cell lung cancer.

106 *Patients with renal impairment* (standard cockcroft and gault formula or glomerular filtration rate
107 measured Tc99m-DPTA serum clearance method)

108 Pemetrexed is primarily eliminated unchanged by renal excretion. In clinical studies, patients with
109 creatinine clearance of \geq 45 ml/min required no dose adjustments other than those
110 recommended for all patients. There are insufficient data on the use of pemetrexed in patients
111 with creatinine clearance below 45 ml/min; therefore the use of pemetrexed is not recommended

112 (see section 4.4).

113 *Patients with hepatic impairment*

114 No relationships between AST (SGOT), ALT (SGPT), or total bilirubin and pemetrexed
115 pharmacokinetics were identified. However, patients with hepatic impairment such as bilirubin >
116 1.5 times the upper limit of normal and/or aminotransferase > 3.0 times the upper limit of normal
117 (hepatic metastases absent) or > 5.0 times the upper limit of normal (hepatic metastases
118 present) have not been specifically studied.

119 Method of administration

120 PEXARITE is for intravenous use. PEXARITE should be administered as an intravenous infusion
121 over 10 minutes on the first day of each 21-day cycle.

122 For precautions to be taken before handling or administering PEXARITE and for instructions on
123 reconstitution and dilution of PEXARITE before administration, see section 6.6.

124 **4.3 Contraindications**

125 Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

126 Breast-feeding (see section 4.6).

127 Concomitant yellow fever vaccine (see section 4.5).

128 **4.4 Special warnings and precautions for use**

129 Pemetrexed can suppress bone marrow function as manifested by neutropenia,
130 thrombocytopenia and anaemia (or pancytopenia) (see section 4.8). Myelosuppression is usually
131 the dose-limiting toxicity. Patients should be monitored for myelosuppression during therapy and
132 pemetrexed should not be given to patients until absolute neutrophil count (ANC) returns to \geq
133 1500 cells/mm³ and platelet count returns to \geq 100,000 cells/mm³. Dose reductions for
134 subsequent cycles are based on nadir ANC, platelet count and maximum non-haematologic
135 toxicity seen from the previous cycle (see section 4.2).

136 Less toxicity and reduction in Grade 3/4 haematologic and non-haematologic toxicities such as
137 neutropenia, febrile neutropenia and infection with Grade 3/4 neutropenia were reported when
138 pre-treatment with folic acid and vitamin B₁₂ was administered. Therefore, all patients treated
139 with pemetrexed must be instructed to take folic acid and vitamin B₁₂ as a prophylactic measure
140 to reduce treatment-related toxicity (see section 4.2).

141 Skin reactions have been reported in patients not pre-treated with a corticosteroid. Pre-treatment
142 with dexamethasone (or equivalent) can reduce the incidence and severity of skin reactions (see
143 section 4.2).

144 An insufficient number of patients has been studied with creatinine clearance of below 45 ml/min.
145 Therefore, the use of pemetrexed in patients with creatinine clearance of < 45 ml/min is not
146 recommended (see section 4.2).

147 Patients with mild to moderate renal insufficiency (creatinine clearance from 45 to 79 ml/min)
148 should avoid taking non-steroidal anti-inflammatory drugs (NSAIDs) such as ibuprofen, and
149 acetylsalicylic acid (> 1.3 g daily) for 2 days before, on the day of, and 2 days following
150 pemetrexed administration (see section 4.5).

151 In patients with mild to moderate renal insufficiency eligible for pemetrexed therapy NSAIDs with
152 long elimination half-lives should be interrupted for at least 5 days prior to, on the day of, and at
153 least 2 days following pemetrexed administration (see section 4.5).

154 Serious renal events, including acute renal failure, have been reported with pemetrexed alone or
155 in association with other chemotherapeutic agents. Many of the patients in whom these occurred
156 had underlying risk factors for the development of renal events including dehydration or pre-
157 existing hypertension or diabetes. Nephrogenic diabetes insipidus and renal tubular necrosis
158 were also reported in post marketing setting with pemetrexed alone or with other
159 chemotherapeutic agents. Most of these events resolved after pemetrexed withdrawal. Patients

160 should be regularly monitored for acute tubular necrosis, decreased renal function and signs and
161 symptoms of nephrogenic diabetes insipidus (e.g. hypernatraemia).

162 The effect of third space fluid, such as pleural effusion or ascites, on pemetrexed is not fully
163 defined. A phase 2 study of pemetrexed in 31 solid tumour patients with stable third space fluid
164 demonstrated no difference in pemetrexed dose normalized plasma concentrations or clearance
165 compared to patients without third space fluid collections. Thus, drainage of third space fluid
166 collection prior to pemetrexed treatment should be considered but may not be necessary.

167 Due to the gastrointestinal toxicity of pemetrexed given in combination with cisplatin, severe
168 dehydration has been observed. Therefore, patients should receive adequate antiemetic
169 treatment and appropriate hydration prior to and/or after receiving treatment.

170 Serious cardiovascular events, including myocardial infarction and cerebrovascular events have
171 been uncommonly reported during clinical studies with pemetrexed, usually when given in
172 combination with another cytotoxic agent. Most of the patients in whom these events have been
173 observed had pre-existing cardiovascular risk factors (see section 4.8).

174 Immunodepressed status is common in cancer patients. As a result, concomitant use of live
175 attenuated vaccines is not recommended (see section 4.3 and 4.5).

176 Pemetrexed can have genetically damaging effects. Sexually mature males are advised not to
177 father a child during the treatment and up to 6 months thereafter. Contraceptive measures or
178 abstinence are recommended. Owing to the possibility of pemetrexed treatment causing
179 irreversible infertility, men are advised to seek counselling on sperm storage before starting
180 treatment.

181 Women of childbearing potential must use effective contraception during treatment with
182 pemetrexed (see section 4.6).

183 Cases of radiation pneumonitis have been reported in patients treated with radiation either prior,
184 during or subsequent to their pemetrexed therapy. Particular attention should be paid to these
185 patients and caution exercised with use of other radiosensitising agents.

186 Cases of radiation recall have been reported in patients who received radiotherapy weeks or
187 years previously.

188 Excipients

189 PEXARITE 100 mg powder for concentrate for solution for infusion

190 This medicinal product contains less than 1 mmol sodium (23 mg) per vial, that is to say
191 essentially 'sodium-free'.

192 PEXARITE 500 mg powder for concentrate for solution for infusion

193 This medicinal product contains 54 mg of sodium per vial, equivalent to 2.7% of the WHO
194 recommended maximum daily intake of 2 g sodium for an adult.

195 **4.5 Interaction with other medicinal products and other forms of interaction**

196 Pemetrexed is mainly eliminated unchanged renally by tubular secretion and to a lesser extent by
197 glomerular filtration. Concomitant administration of nephrotoxic drugs (e.g. aminoglycoside, loop
198 diuretics, platinum compounds, cyclosporin) could potentially result in delayed clearance of
199 pemetrexed. This combination should be used with caution. If necessary, creatinine clearance
200 should be closely monitored.

201 Concomitant administration of substances that are also tubularly secreted (e.g. probenecid,
202 penicillin) could potentially result in delayed clearance of pemetrexed. Caution should be made
203 when these drugs are combined with pemetrexed. If necessary, creatinine clearance should be
204 closely monitored.

205 In patients with normal renal function (creatinine clearance \geq 80 ml/min), high doses of non-
206 steroidal anti-inflammatory drugs (NSAIDs, such as ibuprofen > 1600 mg/day) and acetylsalicylic
207 acid at higher dose (\geq 1.3 g daily) may decrease pemetrexed elimination and, consequently,

208 increase the occurrence of pemetrexed adverse events. Therefore, caution should be made
209 when administering higher doses of NSAIDs or acetylsalicylic acid, concurrently with
210 pemetrexed to patients with normal function (creatinine clearance \geq 80 ml/min).

211 In patients with mild to moderate renal insufficiency (creatinine clearance from 45 to 79 ml/min),
212 the concomitant administration of pemetrexed with NSAIDs (e.g. ibuprofen) or acetylsalicylic
213 acid at higher dose should be avoided for 2 days before, on the day of, and 2 days following
214 pemetrexed administration (see section 4.4).

215 In the absence of data regarding potential interaction with NSAIDs having longer half-lives such
216 as piroxicam or rofecoxib, the concomitant administration with pemetrexed in patients with mild
217 to moderate renal insufficiency should be interrupted for at least 5 days prior to, on the day of,
218 and at least 2 days following pemetrexed administration (see section 4.4). If concomitant
219 administration of NSAIDs is necessary, patients should be monitored closely for toxicity,
220 especially myelosuppression and gastrointestinal toxicity.

221 Pemetrexed undergoes limited hepatic metabolism. Results from *in vitro* studies with human liver
222 microsomes indicated that pemetrexed would not be predicted to cause clinically significant
223 inhibition of the metabolic clearance of drugs metabolised by CYP3A, CYP2D6, CYP2C9, and
224 CYP1A2.

225 Interactions common to all cytotoxics

226 Due to the increased thrombotic risk in patients with cancer, the use of anticoagulation treatment
227 is frequent. The high intra-individual variability of the coagulation status during diseases and the
228 possibility of interaction between oral anticoagulants and anticancer chemotherapy require
229 increased frequency of INR (International Normalised Ratio) monitoring, if it is decided to treat
230 the patient with oral anticoagulants.

231 Concomitant use contraindicated: Yellow fever vaccine: risk of fatal generalised vaccinale
232 disease (see section 4.3).

233 Concomitant use not recommended: Live attenuated vaccines (except yellow fever, for which
234 concomitant use is contraindicated): risk of systemic, possibly fatal, disease. The risk is
235 increased in subjects who are already immunosuppressed by their underlying disease. Use an
236 inactivated vaccine where it exists (poliomyelitis) (see section 4.4).

237 **4.6 Fertility, pregnancy and lactation**

238 Women of childbearing potential / Contraception in males and females

239 Women of childbearing potential must use effective contraception during treatment with
240 pemetrexed.

241 Pemetrexed can have genetically damaging effects. Sexually mature males are advised not to
242 father a child during the treatment and up to 6 months thereafter. Contraceptive measures or
243 abstinence are recommended.

244 Pregnancy

245 There are no data from the use of pemetrexed in pregnant women but pemetrexed, like other
246 anti-metabolites, is suspected to cause serious birth defects when administered during
247 pregnancy. Animal studies have shown reproductive toxicity (see section 5.3). Pemetrexed
248 should not be used during pregnancy unless clearly necessary, after a careful consideration of
249 the needs of the mother and the risk for the foetus (see section 4.4).

250 Breast-feeding

251 It is unknown whether pemetrexed is excreted in human milk and adverse reactions on the
252 breast-feeding child cannot be excluded. Breast-feeding must be discontinued during
253 pemetrexed therapy (see section 4.3).

254 Fertility

255 Owing to the possibility of pemetrexed treatment causing irreversible infertility, men are advised

256 to seek counselling on sperm storage before starting treatment.

257 4.7 Effects on ability to drive and use machines

258 No studies on the effects on the ability to drive and use machines have been performed.

259 However, it has been reported that pemetrexed may cause fatigue. Therefore, patients should
260 be cautioned against driving or operating machines if this event occurs.

261 4.8 Undesirable effects

262 Summary of the safety profile

263 The most commonly reported undesirable effects related to pemetrexed, whether used as
264 monotherapy or in combination, are bone marrow suppression manifested as anaemia,
265 neutropenia, leukopenia, thrombocytopenia; and gastrointestinal toxicities, manifested as
266 anorexia, nausea, vomiting, diarrhoea, constipation, pharyngitis, mucositis, and stomatitis. Other
267 undesirable effects include renal toxicities, increased aminotransferases, alopecia, fatigue,
268 dehydration, rash, infection/sepsis and neuropathy. Rarely seen events include Stevens-Johnson
269 syndrome and toxic epidermal necrolysis.

270 Tabulated list of adverse reactions

271 The table 4 lists the adverse drug events regardless of causality associated with pemetrexed
272 used either as a monotherapy treatment or in combination with cisplatin from the pivotal
273 registration studies (JMCH, JMEI, JMBD, JMEN and PARAMOUNT) and from the post marketing
274 period.

275 ADRs are listed by MedDRA body system organ class. The following convention has been used
276 for classification of frequency: very common: $\geq 1/10$; common: $\geq 1/100$ to $< 1/10$; uncommon: \geq
277 $1/1,000$ to $< 1/100$; rare: $\geq 1/10,000$ to $< 1/1,000$; very rare: $< 1/10,000$ and not known (cannot be
278 estimated from available data).

279 **Table 4. Frequencies of all grades adverse drug events regardless of causality from the**
280 **pivotal registration studies: JMEI (ALIMTA vs Docetaxel), JMDB (ALIMTA and Cisplatin**
281 **versus GEMZAR and Cisplatin, JMCH (ALIMTA plus Cisplatin versus Cisplatin), JMEN**
282 **and PARAMOUNT (Pemetrexed plus Best Supportive Care versus Placebo plus Best**
283 **Supportive Care) and from post-marketing period.**

284

285

System Organ Class (MedDRA)	Very common	Common	Uncommon	Rare	Very rare	Not known
Infections and infestations	Infection ^a Pharyngitis	Sepsis ^b			Dermo-hypodermatitis	
Blood and lymphatic system disorders	Neutropenia Leukopenia Haemoglobin decreased	Febrile neutropenia Platelet count decreased	Pancytopenia	Autoimmune haemolytic anaemia		
Immune System disorders		Hypersensitivity		Anaphylactic shock		
Metabolism and nutrition		Dehydration				

disorders						
Nervous system disorders		Taste disorder Peripheral motor neuropathy Peripheral sensory neuropathy Dizziness	Cerebrovascular accident Ischaemic stroke Haemorrhage intracranial			
Eye disorders		Conjunctivitis Dry eye Lacrimation increased Keratoconjunctivitis sicca Eyelid oedema Ocular surface disease				
Cardiac disorders		Cardiac failure Arrhythmia	Angina Myocardial infarction Coronary artery disease Arrhythmia supra-ventricular			
Vascular disorders			Peripheral ischaemia ^c			
Respiratory, thoracic and mediastinal disorders			Pulmonary embolism Interstitial pneumonitis ^{bd}			
Gastrointestinal disorders	Stomatitis Anorexia Vomiting Diarrhoea Nausea	Dyspepsia Constipation Abdominal pain	Rectal haemorrhage Gastrointestinal haemorrhage Intestinal perforation Oesophagitis			

			Colitis ^e			
Hepatobiliary disorders		Alanine aminotransferase increased Aspartate aminotransferase increased		Hepatitis		
Skin and subcutaneous tissue disorders	Rash Skin exfoliation	Hyperpigmentation Pruritus Erythema multiforme Alopecia Urticaria		Erythema	Stevens Johnson syndrome ^b Toxic epidermal necrolysis ^b Pemphigoid Dermatitis bullous Acquired epidermolysis bullosa Erythematous oedema ^f Pseudocellulitis Dermatitis Eczema Prurigo	
Renal and urinary disorders	Creatinine clearance decreased Blood creatinine increased ^e	Renal failure Glomerular filtration rate decreased				Nephrogenic diabetes insipidus Renal tubular necrosis
General disorders and administration site conditions	Fatigue	Pyrexia Pain Oedema Chest pain Mucosal inflammation				
Investigations		Gamma-glutamyl-transferase				

		increased				
Injury, poisoning and procedural complications			Radiation oesophagitis Radiation pneumonitis	Recall phenomenon		

286 ^a with and without neutropenia

287 ^b in some cases fatal

288 ^c sometimes leading to extremity necrosis

289 ^d with respiratory insufficiency

290 ^e seen only in combination with cisplatin

291 ^f mainly of the lower limbs

292 Reporting of suspected adverse reactions

293 Reporting suspected adverse reactions after authorisation of the medicinal product is important. It
294 allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare
295 professionals are asked to report any suspected adverse reactions to
296 pharmacovigilance@aurobindo.com.

297 **4.9 Overdose**

298 Reported symptoms of overdose include neutropenia, anaemia, thrombocytopenia, mucositis,
299 sensory polyneuropathy and rash. Anticipated complications of overdose include bone marrow
300 suppression as manifested by neutropenia, thrombocytopenia and anaemia. In addition,
301 infection with or without fever, diarrhoea, and/or mucositis may be seen. In the event of
302 suspected overdose, patients should be monitored with blood counts and should receive
303 supportive therapy as necessary. The use of calcium folinate / folic acid in the management of
304 pemetrexed overdose should be considered.

305 **5. Pharmacological properties**

306 **5.1 Pharmacodynamic properties**

307 Pharmacotherapeutic group: Folic acid analogues, ATC code: L01BA04

308 PEXARITE (pemetrexed) is a multi-targeted anti-cancer antifolate agent that exerts its action by
309 disrupting crucial folate-dependent metabolic processes essential for cell replication.

310 *In vitro* studies have shown that pemetrexed behaves as a multitargeted antifolate by inhibiting
311 thymidylate synthase (TS), dihydrofolate reductase (DHFR), and glycinamide ribonucleotide
312 formyltransferase (GARFT), which are key folate-dependent enzymes for the *de novo*
313 biosynthesis of thymidine and purine nucleotides. Pemetrexed is transported into cells by both
314 the reduced folate carrier and membrane folate binding protein transport systems. Once in the
315 cell, pemetrexed is rapidly and efficiently converted to polyglutamate forms by the enzyme
316 folylpolyglutamate synthetase. The polyglutamate forms are retained in cells and are even more
317 potent inhibitors of TS and GARFT. Polyglutamation is a time- and concentration-dependent
318 process that occurs in tumour cells and, to a lesser extent, in normal tissues. Polyglutamated
319 metabolites have an increased intracellular half-life resulting in prolonged drug action in
320 malignant cells.

321 Clinical efficacy

322 *Mesothelioma*

323 EMPHACIS, a multicentre, randomised, single-blind phase 3 study of pemetrexed plus cisplatin
324 versus cisplatin in chemo-naïve patients with malignant pleural mesothelioma, has shown that
325 patients treated with pemetrexed and cisplatin had a clinically meaningful 2.8-month median

326 survival advantage over patients receiving cisplatin alone.

327 During the study, low-dose folic acid and vitamin B₁₂ supplementation was introduced to patients'
328 therapy to reduce toxicity. The primary analysis of this study was performed on the population of
329 all patients randomly assigned to a treatment arm who received study drug (randomised and
330 treated). A subgroup analysis was performed on patients who received folic acid and vitamin B₁₂
331 supplementation during the entire course of study therapy (fully supplemented). The results of
332 these analyses of efficacy are summarised in the table below:

333

334 **Table 5. Efficacy of pemetrexed plus cisplatin vs. cisplatin in malignant pleural**
335 **mesothelioma**

Efficacy parameter	Randomized and treated patients		Fully supplemented patients	
	Pemetrexed/ cisplatin	Cisplatin	Pemetrexed/ cisplatin	Cisplatin
	(N = 226)	(N = 222)	(N = 168)	(N = 163)
Median overall survival (months) (95 % CI)	12.1 (10.0 - 14.4)	9.3 (7.8 - 10.7)	13.3 (11.4 - 14.9)	10.0 (8.4 - 11.9)
Log Rank p-value*	0.020		0.051	
Median time to tumour (months) (95 % CI)	5.7 (4.9 - 6.5)	3.9 (2.8 - 4.4)	6.1 (5.3 - 7.0)	3.9 (2.8 - 4.5)
Log Rank p-value*	0.001		0.008	
Time to treatment failure (95 % CI)	4.5 (3.9 - 4.9)	2.7 (2.1 - 2.9)	4.7 (4.3 - 5.6)	2.7 (2.2 - 3.1)
Log Rank p-value*	0.001		0.001	
Overall response rate** (95 % CI)	41.3 % (34.8 - 48.1)	16.7 % (12.0 - 22.2)	45.5 % (37.8 - 53.4)	19.6 % (13.8 - 26.6)
Fisher's exact p-value*	< 0.001		< 0.001	

336 Abbreviation: CI = confidence interval

337 * p-value refers to comparison between arms.

338 ** In the pemetrexed/cisplatin arm, randomized and treated (N = 225) and fully supplemented (N
339 = 167)

340 A statistically significant improvement of the clinically relevant symptoms (pain and dyspnoea)
341 associated with malignant pleural mesothelioma in the pemetrexed/cisplatin arm (212 patients)
342 versus the cisplatin arm alone (218 patients) was demonstrated using the Lung Cancer Symptom
343 Scale. Statistically significant differences in pulmonary function tests were also observed. The
344 separation between the treatment arms was achieved by improvement in lung function in the
345 pemetrexed/cisplatin arm and deterioration of lung function over time in the control arm.

346 There are limited data in patients with malignant pleural mesothelioma treated with pemetrexed
347 alone. Pemetrexed at a dose of 500 mg/m² was studied as a single-agent in 64 chemo-naïve
348 patients with malignant pleural mesothelioma. The overall response rate was 14.1 %.

349 *NSCLC, second-line treatment*

350 A multicentre, randomised, open label phase 3 study of pemetrexed versus docetaxel in patients
351 with locally advanced or metastatic NSCLC after prior chemotherapy has shown median survival
352 times of 8.3 months for patients treated with pemetrexed (Intent To Treat population n = 283) and
353 7.9 months for patients treated with docetaxel (ITT n = 288). Prior chemotherapy did not include
354 pemetrexed. An analysis of the impact of NSCLC histology on the treatment effect on overall
355 survival was in favour of pemetrexed versus docetaxel for other than predominantly squamous
356 histologies (n = 399, 9.3 versus 8.0 months, adjusted HR = 0.78; 95% CI = 0.61-1.00, p = 0.047)

357 and was in favour of docetaxel for squamous cell carcinoma histology (n = 172, 6.2 versus 7.4
 358 months, adjusted HR = 1.56; 95% CI = 1.08-2.26, p = 0.018). There were no clinically relevant
 359 differences observed for the safety profile of pemetrexed within the histology subgroups.

360 Limited clinical data from a separate randomized, Phase 3, controlled trial, suggest that efficacy
 361 data (overall survival, progression free survival) for pemetrexed are similar between patients
 362 previously pre treated with docetaxel (n = 41) and patients who did not receive previous
 363 docetaxel treatment (n = 540).

364

365 **Table 6. Efficacy of pemetrexed vs docetaxel in NSCLC - ITT population**

	Pemetrexed	Docetaxel
Survival Time (months)	(n = 283)	(n = 288)
▪ Median (m)	8.3	7.9
▪ 95 % CI for median	(7.0 - 9.4)	(6.3 - 9.2)
▪ HR	0.99	
▪ 95 % CI for HR	(.82 - 1.20)	
▪ Non-inferiority p-value (HR)	.226	
Progression free survival (months)	(n = 283)	(n = 288)
▪ Median	2.9	2.9
▪ HR (95 % CI)	0.97 (.82 – 1.16)	
Time to treatment failure (TTTF)	(n = 283)	(n = 288)
▪ Median	2.3	2.1
▪ HR (95 % CI)	0.84 (.71 - .997)	
Response (n: qualified for response)	(n = 264)	(n = 274)
▪ Response rate (%) (95 % CI)	9.1 (5.9 - 13.2)	8.8 (5.7 - 12.8)
▪ Stable disease (%)	45.8	46.4

366 Abbreviations: CI = confidence interval; HR = hazard ratio; ITT = intent to treat; n = total
 367 population size.

368 *NSCLC, first-line treatment*

369 A multicentre, randomised, open-label, Phase 3 study of pemetrexed plus cisplatin versus
 370 gemcitabine plus cisplatin in chemo-naïve patients with locally advanced or metastatic (Stage IIIb
 371 or IV) non-small cell lung cancer (NSCLC) showed that pemetrexed plus cisplatin (Intent-To-
 372 Treat [ITT] population n = 862) met its primary endpoint and showed similar clinical efficacy as
 373 gemcitabine plus cisplatin (ITT n = 863) in overall survival (adjusted hazard ratio 0.94; 95% CI =
 374 0.84-1.05). All patients included in this study had an ECOG performance status 0 or 1.

375 The primary efficacy analysis was based on the ITT population. Sensitivity analyses of main
 376 efficacy endpoints were also assessed on the Protocol Qualified (PQ) population. The efficacy
 377 analyses using PQ population are consistent with the analyses for the ITT population and support
 378 the non-inferiority of AC versus GC.

379 Progression free survival (PFS) and overall response rate were similar between treatment arms:
 380 median PFS was 4.8 months for pemetrexed plus cisplatin versus 5.1 months for gemcitabine
 381 plus cisplatin (adjusted hazard ratio 1.04; 95% CI = 0.94-1.15), and overall response rate was
 382 30.6% (95% CI = 27.3-33.9) for pemetrexed plus cisplatin versus 28.2% (95% CI = 25.0-31.4) for
 383 gemcitabine plus cisplatin. PFS data were partially confirmed by an independent review
 384 (400/1725 patients were randomly selected for review).

385 The analysis of the impact of NSCLC histology on overall survival demonstrated clinically
 386 relevant differences in survival according to histology, see table below.

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Table 7. Efficacy of pemetrexed + cisplatin vs. gemcitabine + cisplatin in first-line non-small cell lung cancer – ITT population and histology subgroups.

ITT population and histology subgroups	Median overall survival in months (95% CI)				Adjusted hazard ratio (HR) (95% CI)	Superiority p-value
	Pemetrexed + cisplatin		Gemcitabine + cisplatin			
ITT population (N = 1725)	10.3 (9.8 – 11.2)	N=862	10.3 (9.6 – 10.9)	N=863	0.94 ^a (0.84 – 1.05)	0.259
Adenocarcinoma (N=847)	12.6 (10.7 – 14.6)	N=436	10.9 (10.2 – 11.6)	N=411	0.84 (0.71–0.99)	0.033
Large cell (N=153)	10.4 (8.6 – 14.1)	N=76	6.7 (5.5 – 9.0)	N=77	0.67 (0.48–0.96)	0.027
Other (N=252)	8.6 (6.8 – 10.2)	N=106	9.2 (8.1 – 10.6)	N=146	1.08 (0.81–1.45)	0.586
Squamous cell (N=473)	9.4 (8.4 – 10.2)	N=244	10.8 (9.5 – 12.1)	N=229	1.23 (1.00–1.51)	0.050

390 Abbreviations: CI = confidence interval; ITT = intent-to-treat; N = total population size.

391 ^a Statistically significant for noninferiority, with the entire confidence interval for HR well below the
392 1.17645 noninferiority margin (p < 0.001).

393 **Kaplan Meier plots of overall survival by histology**

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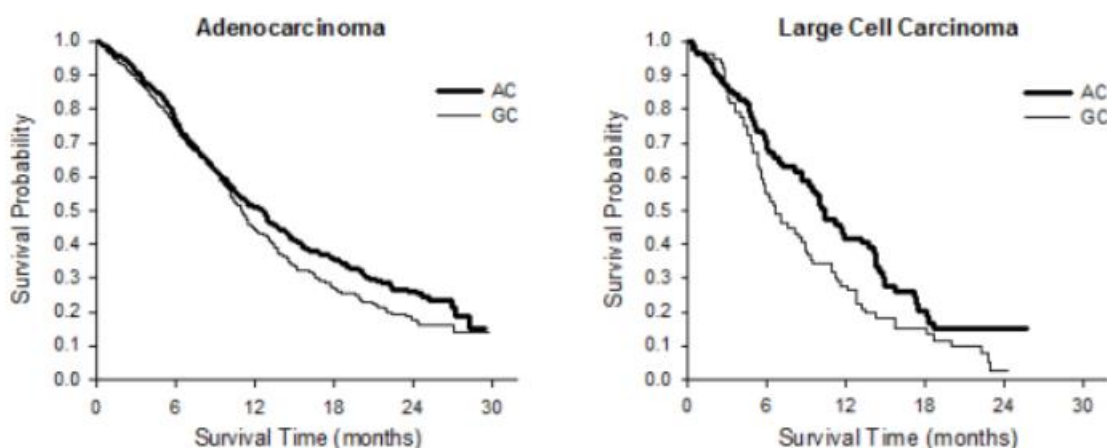
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405 There were no clinically relevant differences observed for the safety profile of pemetrexed plus
406 cisplatin within the histology subgroups.

407 Patients treated with pemetrexed and cisplatin required fewer transfusions (16.4% versus 28.9%,
408 p<0.001), red blood cell transfusions (16.1% versus 27.3%, p<0.001) and platelet transfusions
409 (1.8% versus 4.5%, p=0.002). Patients also required lower administration of
410 erythropoietin/darbopoietin (10.4% versus 18.1%, p<0.001), G-CSF/GM-CSF (3.1% versus 6.1%,
411 p=0.004), and iron preparations (4.3% versus 7.0%, p=0.021).

412 *NSCLC, maintenance treatment*

413 **JMEN**

414 A multicentre, randomised, double-blind, placebo-controlled Phase 3 study (JMEN), compared
415 the efficacy and safety of maintenance treatment with pemetrexed plus best supportive care
416 (BSC) (n = 441) with that of placebo plus BSC (n = 222) in patients with locally advanced (Stage
417 IIIB) or metastatic (Stage IV) Non Small Cell Lung Cancer (NSCLC) who did not progress after 4
418 cycles of first line doublet therapy containing Cisplatin or Carboplatin in combination with

419 Gemcitabine, Paclitaxel, or Docetaxel. First line doublet therapy containing pemetrexed was not
420 included. All patients included in this study had an ECOG performance status 0 or 1. Patients
421 received maintenance treatment until disease progression. Efficacy and safety were measured
422 from the time of randomisation after completion of first line (induction) therapy. Patients received
423 a median of 5 cycles of maintenance treatment with pemetrexed and 3.5 cycles of placebo. A
424 total of 213 patients (48.3%) completed ≥ 6 cycles and a total of 103 patients (23.4%) completed
425 ≥ 10 cycles of treatment with pemetrexed.

426 The study met its primary endpoint and showed a statistically significant improvement in PFS in
427 the pemetrexed arm over the placebo arm (n = 581, independently reviewed population; median
428 of 4.0 months and 2.0 months, respectively) (hazard ratio = 0.60, 95% CI = 0.49-0.73, p <
429 0.00001). The independent review of patient scans confirmed the findings of the investigator
430 assessment of PFS. The median OS for the overall population (n = 663) was 13.4 months for the
431 pemetrexed arm and 10.6 months for the placebo arm, hazard ratio = 0.79 (95% CI = 0.65-0.95,
432 p = 0.01192).

433 Consistent with other pemetrexed studies, a difference in efficacy according to NSCLC histology
434 was observed in JMEN. For patients with NSCLC other than predominantly squamous cell
435 histology (n = 430, independently reviewed population) median PFS was 4.4 months for the
436 pemetrexed arm and 1.8 months for the placebo arm, hazard ratio = 0.47 (95% CI = 0.37-0.60, p
437 = 0.00001). The median OS for patients with NSCLC other than predominantly squamous cell
438 histology (n = 481) was 15.5 months for the pemetrexed arm and 10.3 months for the placebo
439 arm, hazard ratio = 0.70 (95% CI = 0.56-0.88, p = 0.002). Including the induction phase the
440 median OS for patients with NSCLC other than predominantly squamous cell histology was 18.6
441 months for the pemetrexed arm and 13.6 months for the placebo arm, hazard ratio = 0.71 (95%
442 CI = 0.56-0.88, p = 0.002).

443 The PFS and OS results in patients with squamous cell histology suggested no advantage for
444 pemetrexed over placebo.

445 There were no clinically relevant differences observed for the safety profile of pemetrexed within
446 the histology subgroups.

447 **JMEN: Kaplan Meier plots of progression-free survival (PFS) and overall survival**
448 **pemetrexed versus placebo in patients with NSCLC other than predominantly squamous**
449 **cell histology:**

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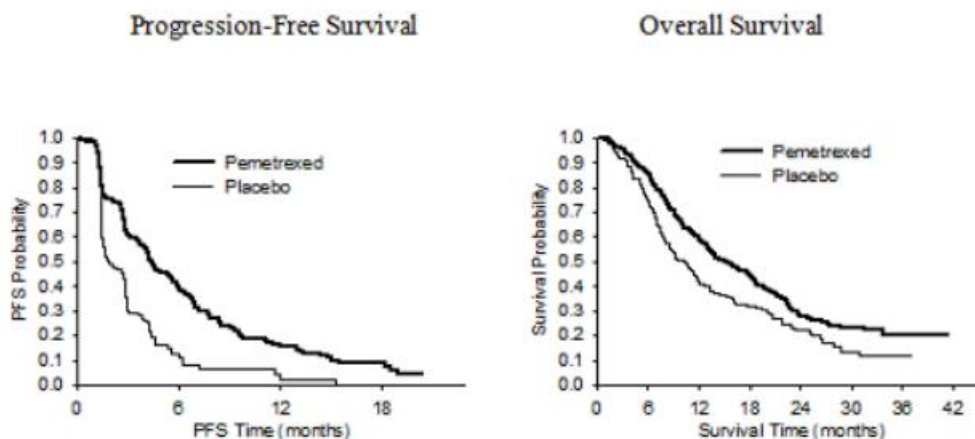
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463 PARAMOUNT

464 A multicentre, randomised, double-blind, placebo-controlled Phase 3 study (PARAMOUNT),

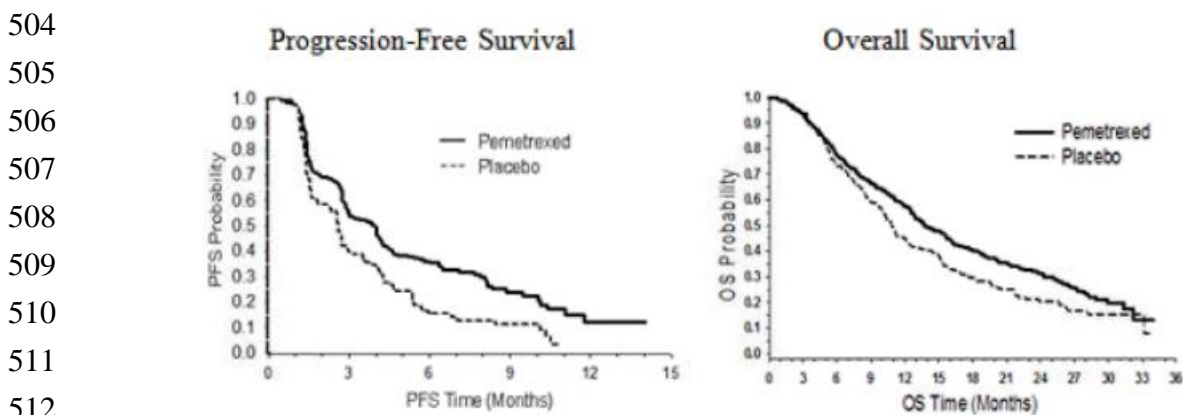


465 compared the efficacy and safety of continuation maintenance treatment with pemetrexed plus
 466 BSC (n = 359) with that of placebo plus BSC (n = 180) in patients with locally advanced (Stage
 467 IIIB) or metastatic (Stage IV) NSCLC other than predominantly squamous cell histology who did
 468 not progress after 4 cycles of first line doublet therapy of pemetrexed in combination with
 469 cisplatin. Of the 939 patients treated with pemetrexed plus cisplatin induction, 539 patients were
 470 randomised to maintenance treatment with pemetrexed or placebo. Of the randomised patients,
 471 44.9% had a complete/partial response and 51.9% had a response of stable disease to
 472 pemetrexed plus cisplatin induction. Patients randomised to maintenance treatment were
 473 required to have an ECOG performance status 0 or 1. The median time from the start of
 474 pemetrexed plus cisplatin induction therapy to the start of maintenance treatment was 2.96
 475 months on both the pemetrexed arm and the placebo arm. Randomised patients received
 476 maintenance treatment until disease progression. Efficacy and safety were measured from the
 477 time of randomisation after completion of first line (induction) therapy. Patients received a median
 478 of 4 cycles of maintenance treatment with pemetrexed and 4 cycles of placebo. A total of 169
 479 patients (47.1%) completed ≥ 6 cycles maintenance treatment with pemetrexed, representing at
 480 least 10 total cycles of pemetrexed.

481 The study met its primary endpoint and showed a statistically significant improvement in PFS in
 482 the pemetrexed arm over the placebo arm (n = 472, independently reviewed population; median
 483 of 3.9 months and 2.6 months, respectively) (hazard ratio = 0.64, 95% CI = 0.51-0.81, p =
 484 0.0002). The independent review of patient scans confirmed the findings of the investigator
 485 assessment of PFS. For randomised patients, as measured from the start of pemetrexed plus
 486 cisplatin first line induction treatment, the median investigator-assessed PFS was 6.9 months for
 487 the pemetrexed arm and 5.6 months for the placebo arm (hazard ratio = 0.59 95% CI = 0.47-
 488 0.74).

489 Following pemetrexed plus cisplatin induction (4 cycles), treatment with pemetrexed was
 490 statistically superior to placebo for OS (median 13.9 months versus 11.0 months, hazard ratio =
 491 0.78, 95%CI=0.64-0.96, p=0.0195). At the time of this final survival analysis, 28.7% of patients
 492 were alive or lost to follow up on the pemetrexed arm versus 21.7% on the placebo arm. The
 493 relative treatment effect of pemetrexed was internally consistent across subgroups (including
 494 disease stage, induction response, ECOG PS, smoking status, gender, histology and age) and
 495 similar to that observed in the unadjusted OS and PFS analyses. The 1 year and 2 year survival
 496 rates for patients on pemetrexed were 58% and 32% respectively, compared to 45% and 21% for
 497 patients on placebo. From the start of pemetrexed plus cisplatin first line induction treatment, the
 498 median OS of patients was 16.9 months for the pemetrexed arm and 14.0 months for the placebo
 499 arm (hazard ratio= 0.78, 95% CI= 0.64-0.96). The percentage of patients that received post
 500 study treatment was 64.3% for pemetrexed and 71.7% for placebo.

501 **PARAMOUNT:Kaplan Meier plot of progression-free survival (PFS) and Overall Survival**
 502 **(OS) for continuation pemetrexed maintenance versus placebo in patients with NSCLC**
 503 **other than predominantly squamous cell histology (measured from randomisation)**



514 The pemetrexed maintenance safety profiles from the two studies JMEN and PARAMOUNT were
515 similar.

516 **5.2 Pharmacokinetic properties**

517 The pharmacokinetic properties of pemetrexed following single-agent administration have been
518 evaluated in 426 cancer patients with a variety of solid tumours at doses ranging from 0.2 to 838
519 mg/m² infused over a 10-minute period.

520 Pemetrexed has a steady-state volume of distribution of 9 l/m². *In vitro* studies indicate that
521 pemetrexed is approximately 81 % bound to plasma proteins. Binding was not notably affected
522 by varying degrees of renal impairment. Pemetrexed undergoes limited hepatic metabolism.
523 Pemetrexed is primarily eliminated in the urine, with 70 % to 90 % of the administered dose
524 being recovered unchanged in urine within the first 24 hours following administration. *In vitro*
525 studies indicate that pemetrexed is actively secreted by OAT3 (organic anion transporter).
526 Pemetrexed total systemic clearance is 91.8 ml/min and the elimination half-life from plasma is
527 3.5 hours in patients with normal renal function (creatinine clearance of 90 ml/min). Between
528 patient variability in clearance is moderate at 19.3 %. Pemetrexed total systemic exposure (AUC)
529 and maximum plasma concentration increase proportionally with dose. The pharmacokinetics of
530 pemetrexed are consistent over multiple treatment cycles.

531 The pharmacokinetic properties of pemetrexed are not influenced by concurrently administered
532 cisplatin. Oral folic acid and intramuscular vitamin B₁₂ supplementation do not affect the
533 pharmacokinetics of pemetrexed.

534 **5.3 Preclinical safety data**

535 Administration of pemetrexed to pregnant mice resulted in decreased foetal viability, decreased
536 foetal weight, incomplete ossification of some skeletal structures and cleft palate.

537 Administration of pemetrexed to male mice resulted in reproductive toxicity characterised by
538 reduced fertility rates and testicular atrophy. In a study conducted in beagle dog by intravenous
539 bolus injection for 9 months, testicular findings (degeneration/necrosis of the seminiferous
540 epithelium) have been observed. This suggests that pemetrexed may impair male fertility.
541 Female fertility was not investigated.

542 Pemetrexed was not mutagenic in either the *in vitro* chromosome aberration test in Chinese
543 hamster ovary cells, or the Ames test. Pemetrexed has been shown to be clastogenic in the *in*
544 *vivo* micronucleus test in the mouse.

545 Studies to assess the carcinogenic potential of pemetrexed have not been conducted.

546 **6. Pharmaceutical particulars**

547 **6.1 List of excipients**

548 Mannitol

549 Hydrochloric acid

550 Sodium hydroxide

551 **6.2 Incompatibilities**

552 Pemetrexed is physically incompatible with diluents containing calcium, including lactated
553 Ringer's injection and Ringer's injection. In the absence of other compatibility studies this
554 medicinal product must not be mixed with other medicinal products.

555 **6.3 Shelf life**

556 Unopened vial

557 2 years.

558 Reconstituted and infusion solutions

559 When prepared as directed, reconstituted and infusion solutions of PEXARITE contain no

560 antimicrobial preservatives. Chemical and physical in-use stability of reconstituted and infusion
561 solutions of pemetrexed were demonstrated for 24 hours at refrigerated temperature. From a
562 microbiological point of view, the product should be used immediately. If not used immediately,
563 in-use storage times and conditions prior to use are the responsibility of the user and would not
564 be longer than 24 hours at 2°C to 8°C.

565 **6.4 Special precautions for storage**

566 Unopened vial

567 Store below 30°C.

568 This medicinal product does not require any special storage conditions

569 For storage conditions after reconstitution of the medicinal product, see section 6.3.

570 **6.5 Nature and contents of container**

571 PEXARITE 100 mg powder for concentrate for solution for infusion

572 Type I glass vial with rubber stopper containing 100 mg of pemetrexed.

573 Pack of 1 vial.

574 PEXARITE 500 mg powder for concentrate for solution for infusion

575 Type I glass vial with rubber stopper containing 500 mg of pemetrexed.

576 Pack of 1 vial.

577 **6.6 Special precautions for disposal and other handling**

578 1. Use aseptic technique during the reconstitution and further dilution of pemetrexed for
579 intravenous infusion administration.

580 2. Calculate the dose and the number of PEXARITE vials needed. Each vial contains an excess
581 of pemetrexed to facilitate delivery of label amount.

582 3. PEXARITE 100 mg

583 Reconstitute 100-mg vials with 4.2 ml of sodium chloride 9 mg/ml (0.9 %) solution for injection,
584 without preservative, resulting in a solution containing 25 mg/ml pemetrexed.

585 Pemetrexed 500 mg

586 Reconstitute 500-mg vials with 20 ml of sodium chloride 9 mg/ml (0.9 %) solution for injection,
587 without preservative, resulting in a solution containing 25 mg/ml pemetrexed.

588 Gently swirl each vial until the powder is completely dissolved. The resulting solution is clear and
589 ranges in colour from colourless to yellow or green-yellow without adversely affecting product
590 quality. The pH of the reconstituted solution is between 6.6 and 7.8. **Further dilution is**
591 **required.**

592 4. The appropriate volume of reconstituted pemetrexed solution must be further diluted to 100 ml
593 with sodium chloride 9 mg/ml (0.9 %) solution for injection, without preservative, and
594 administered as an intravenous infusion over 10 minutes.

595 5. Pemetrexed infusion solutions prepared as directed above are compatible with polyvinyl
596 chloride lined administration sets and infusion bags.

597 6. Parenteral medicinal products must be inspected visually for particulate matter and
598 discolouration prior to administration. If particulate matter is observed, do not administer.

599 7. Pemetrexed solutions are for single use only. Any unused medicinal product or waste material
600 must be disposed of in accordance with local requirements.

601 Preparation and administration precautions

602 As with other potentially toxic anticancer agents, care should be exercised in the handling and

603 preparation of pemetrexed infusion solutions. The use of gloves is recommended. If a
604 pemetrexed solution contacts the skin, wash the skin immediately and thoroughly with soap and
605 water. If pemetrexed solutions contact the mucous membranes, flush thoroughly with water.
606 Pemetrexed is not a vesicant. There is not a specific antidote for extravasation of pemetrexed.
607 There have been few reported cases of pemetrexed extravasation, which were not assessed as
608 serious by the investigator. Extravasation should be managed by local standard practice as with
609 other non-vesicants.

610 **7. Marketing authorization holder**

611 Imported by:

612 APL Pharma Thai Ltd

613 438 Phattanakarn 30, Phattanakarn Road,

614 Suanluang Subdistrict, Suanluang District,

615 Bangkok, Thailand 10250

616 Manufactured by:

617 Eugia Pharma Specialities Limited,

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

619 Shameerpet Mandal,

620 Medchal-Malkajgiri District,

621 Telangana, India.

622 **8. Marketing authorization Number(s)**

623 1C...../.....(NG)

624 **9. Date of authorization:**

625 **10. Date of revision of the text:** July 6, 2022