1 CLOLAR[™] FOR INTRAVENOUS INFUSION

2 (clofarabine)

3 DESCRIPTION

- 4 CLOLAR[™] For Intravenous Infusion (CLOLAR[™]; clofarabine) contains clofarabine, a
- 5 purine nucleoside anti-metabolite. CLOLAR[™] (1 mg/mL) is supplied in a 20 mL, single-use
- 6 vial. The 20 mL vial contains 20 mg clofarabine formulated in 20 mL unbuffered normal
- 7 saline (comprised of Water for Injection, USP, and Sodium Chloride USP). The pH range of
- 8 the solution is 4.5 to 7.5. The solution is clear and practically colorless, and free from
- 9 foreign matter.

10

- 11 The chemical structure of clofarabine is 2-chloro-9-(2-deoxy-2-fluoro- β -D-
- 12 arabinofuranosyl)-9H-purin-6-amine. The molecular formula of clofarabine is
- 13 $C_{10}H_{11}ClFN_5O_3$ with a molecular weight of 303.68.

14



Clofarabine

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18 CLINICAL PHARMACOLOGY

19 Mechanism of Action: Clofarabine is sequentially metabolized intracellularly to the 5'-20 monophosphate metabolite by deoxycytidine kinase and mono- and di-phosphokinases to the 21 active 5'-triphosphate metabolite. Clofarabine has high affinity for the activating 22 phosphorylating enzyme, deoxycytidine kinase, equal to or greater than that of the natural 23 substrate, deoxycytidine. Clofarabine inhibits DNA synthesis by decreasing cellular 24 deoxynucleotide triphosphate pools through an inhibitory action on ribonucleotide reductase. 25 and by terminating DNA chain elongation and inhibiting repair through incorporation into 26 the DNA chain by competitive inhibition of DNA polymerases. The affinity of clofarabine 27 triphosphate for these enzymes is similar to or greater than that of deoxyadenosine 28 triphosphate. In preclinical models, clofarabine has demonstrated the ability to inhibit DNA 29 repair by incorporation into the DNA chain during the repair process. Clofarabine 5'-30 triphosphate also disrupts the integrity of mitochondrial membrane, leading to the release of 31 the pro-apoptotic mitochondrial proteins, cytochrome C and apoptosis-inducing factor, 32 leading to programmed cell death.

33

34 Clofarabine is cytotoxic to rapidly proliferating and quiescent cancer cell types *in vitro*.

35

36 Human Pharmacokinetics: The population pharmacokinetics of CLOLAR[™] were studied in 37 40 pediatric patients aged 2 to 19 years (21 males/19 females) with relapsed or refractory ALL or AML. At the given 52 mg/m^2 dose, similar concentrations were obtained over a 38 39 wide range of BSAs. Clofarabine was 47% bound to plasma proteins, predominantly to 40 albumin. Based on non-compartmental analysis, systemic clearance and volume of distribution at steady-state were estimated to be 28.8 $L/h/m^2$ and 172 L/m^2 , respectively. The 41 42 terminal half-life was estimated to be 5.2 hours. No apparent difference in pharmacokinetics 43 was observed between patients with ALL and AML or between males and females.

response was found in this population.

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Based on 24-hour urine collections in the pediatric studies, 49-60% of the dose is excreted in
the urine unchanged. *In vitro* studies using isolated human hepatocytes indicate very limited
metabolism (0.2%), therefore the pathways of non-renal elimination remain unknown.

No relationship between clofarabine or clofarabine triphosphate exposure and toxicity or

51

Although no clinical drug-drug interaction studies have been conducted to date, on the basis of the *in vitro* studies, cytochrome p450 inhibitors and inducers are unlikely to affect the metabolism of clofarabine. The effect of clofarabine on the metabolism of cytochrome p450 substrates has not been studied. The pharmacokinetics of clofarabine have not been evaluated in patients with renal or hepatic dysfunction.

57

58 CLINICAL STUDIES

Sixty-six (66) pediatric ALL patients were exposed to CLOLAR[™]. Fifty-eight (58) of the
patients received the recommended pediatric dose of CLOLAR[™] 52 mg/m² daily × 5 as an
intravenous infusion (IVI).

62

The safety and efficacy of CLOLAR[™] were evaluated in pediatric patients with refractory or relapsed hematologic malignancies in an open-label, dose-escalation, noncomparative study. The starting dose of CLOLAR[™] was 11.25 mg/m²/day IVI daily × 5 and escalated to 70 mg/m²/day IVI daily × 5. This dosing schedule was repeated every 2 to 6 weeks depending on toxicity and response. Nine of 17 ALL patients were treated with CLOLAR[™] 52 mg/m² daily × 5. In the 17 ALL patients there were 2 complete remissions (12.5%) and 2 partial remissions (12.5%) at varying doses. Dose-limiting toxicities (DLTs) in this study were

reversible hyperbilirubinemia and elevated transaminase levels and skin rash, experienced at 70 mg/m². As a result of this study, the recommended dose for subsequent study in pediatric 72 patients was determined to be $52 \text{ mg/m}^2/\text{day}$ for 5 days.

73

74 Single Arm Study in Pediatric ALL

75 A single arm study was conducted in relapsed/refractory pediatric patients with ALL at a single dose. All patients had disease that had relapsed after and/or was refractory to two or 76 77 more prior therapies. Most patients, 46/49 (93.8%), had received 2 to 4 prior regimens and 15/49 (30.6%) of the patients had undergone at least 1 prior transplant. The median age of 78 79 the treated patients was 12 years. There were more males, 29/49 (59.2%), than females, 80 20/49 (40.8%). Most of the patients were either Caucasian (n=20, 40.8%) or Hispanic (n=20, 40.8%), with 12.2% African-American (n=6), and 6.1% Other race (n=3). All patients 81 received a dose of 52 mg/m² daily \times 5 IVI. There was no dose modification during the 82 83 remission induction phase of treatment (maximum of 2 cycles). Doses could be modified (reduced/delayed) during the post-induction phase. There was no dose escalation. The 84 85 planned study endpoint was the rate of Complete Remission (CR), defined as no evidence of circulating blasts or extramedullary disease, an M1 bone marrow (<5% blasts), and recovery 86 of peripheral counts (platelets $> 100 \times 10^9$ L and absolute neutrophil count (ANC) $> 1.0 \times 10^9$ L and absolute neutrophil count (ANC) 87 10⁹ L) and Complete Remission in the Absence of Total Platelet Recovery (CRp), defined as 88 meeting all criteria for CR except for recovery of platelet counts to $> 100 \times 10^9$ L. Partial 89 90 Response (PR) was also determined, defined as complete disappearance of circulating blasts, 91 an M2 bone marrow (> 5% and < 25% blasts), and appearance of normal progenitor cells or 92 an M1 marrow that did not qualify for CR or CRp. Transplantation rate was not a study 93 endpoint.

94

Response rates for these studies were determined by an unblinded Independent ResponseReview Panel (IRRP).

- 98 Table 1 summarizes results for the pediatric ALL study. Responses were seen in both pre-B
- 99 and T-cell immunophenotypes of ALL. The median cumulative dose was 540 mg (range 29-
- 100 1905 mg) in 1 (42.9%), 2 (38.8%) or 3 or more (18.4%) cycles.

101

102

Table 1:	Results	in	Pediatric	ALL	Study
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n=49					
Responses	n	%	95% CI		
CR	6	12.2	4.6 to 24.8		
CRp	4	8.2	2.3 to 19.6		
PR	5	10.2	3.4 to 22.2		

103

104 Of the 15 responding pediatric ALL patients, 6 had post-clofarabine bone marrow

105 transplantation, so that duration of response could not be determined. In the 9 responding

106 patients who were not transplanted, the response durations for CR were 43, 50, 82, 93+, and

107 160+ days; for CRp the response duration was 32 days; and for PR the response durations

108 were 7, 16, and 21 days.

109

110 INDICATIONS AND USAGE

111 CLOLAR[™] is indicated for the treatment of pediatric patients 1 to 21 years old with relapsed

112 or refractory acute lymphoblastic leukemia after at least two prior regimens. This use is

113 based on the induction of complete responses. Randomized trials demonstrating increased

114 survival or other clinical benefit have not been conducted.

116 CONTRAINDICATIONS

- 117 None
- 118

119 WARNINGS

120 CLOLAR[™] should be administered under the supervision of a qualified physician 121 experienced in the use of antineoplastic therapy. Suppression of bone marrow function 122 should be anticipated. This is usually reversible and appears to be dose dependent. The use of CLOLAR[™] is likely to increase the risk of infection, including severe sepsis, as a result of 123 bone marrow suppression. Administration of CLOLAR[™] results in a rapid reduction in 124 125 peripheral leukemia cells. For this reason, patients undergoing treatment with CLOLAR[™] 126 should be evaluated and monitored for signs and symptoms of tumor lysis syndrome, as well 127 as signs and symptoms of cytokine release (eg. tachypnea, tachycardia, hypotension, 128 pulmonary edema) that could develop into systemic inflammatory response syndrome 129 (SIRS)/capillary leak syndrome, and organ dysfunction. Physicians are encouraged to give 130 continuous IV fluids throughout the five days of CLOLAR[™] administration to reduce the 131 effects of tumor lysis and other adverse events. Allopurinol should be administered if hyperuricemia is expected. CLOLAR[™] should be discontinued immediately in the event of 132 133 clinically significant signs or symptoms of SIRS or capillary leak syndrome, either of which can be fatal, and use of steroids, diuretics, and albumin considered. CLOLAR™ can be re-134 135 instituted when the patient is stable, generally at a lower dose.

136

137 Severe bone marrow suppression, including neutropenia, anemia, and thrombocytopenia, has

138 been observed in patients treated with CLOLAR[™]. At initiation of treatment, most patients

139 in the clinical studies had hematological impairment as a manifestation of leukemia. Because

- 140 of the pre-existing immunocompromised condition of these patients and prolonged
- 141 neutropenia that can result from treatment with CLOLAR[™], patients are at increased risk for

142	severe opportunistic infections.	Careful hematological	monitoring	during f	herany is
144	severe opportunistic infections.	Calciul liciliatological	monitoring	uuring i	nerapy is

- 143 important, and hepatic and renal function should be assessed prior to and during treatment
- 144 with CLOLAR[™] because of CLOLAR[™]'s predominantly renal excretion and because the
- 145 liver is a target organ for CLOLARTM toxicity. The respiratory status and blood pressure
- should be closely monitored during infusion of CLOLAR[™].
- 147

148 Hepatic and Renal Impairment

149 CLOLAR[™] has not been studied in patients with hepatic or renal dysfunction. Its use in

150 such patients should be undertaken only with the greatest caution.

151

152 Pregnancy – Teratogenic Effects: Pregnancy Category D

153 CLOLAR[™] (clofarabine) may cause fetal harm when administered to a pregnant woman.

154 Clofarabine was teratogenic in rats and rabbits. Developmental toxicity (reduced fetal body

155 weight and increased post-implantation loss) and increased incidences of malformations and

156 variations (gross external, soft tissue, skeletal and retarded ossification) were observed in rats

157 receiving 54 mg/m²/day (approximately equivalent to the recommended clinical dose on a

 158 mg/m^2 basis), and in rabbits receiving 12 mg/m^2 /day (approximately 23% of the

- 159 recommended clinical dose on a mg/m^2 basis).
- 160

161 There are no adequate and well-controlled studies in pregnant women using clofarabine. If

162 this drug is used during pregnancy, or if the patient becomes pregnant while taking this drug,

163 the patient should be apprised of the potential hazard to the fetus.

164

165 Women of childbearing potential should be advised to avoid becoming pregnant while166 receiving treatment with clofarabine.

168 **PRECAUTIONS**

169 Information for Patients and Caregivers

170 Physicians are advised to discuss the following with patients to whom CLOLAR[™] will be

administered and patient caregivers, as appropriate.

172

173 Dehydration/Hypotension

Patients receiving CLOLAR[™] may experience vomiting and diarrhea; they should therefore
be advised regarding appropriate measures to avoid dehydration. Patients should be
instructed to seek medical advice if they experience symptoms of dizziness, lightheadedness,
fainting spells, or decreased urine output. CLOLAR[™] administration should be stopped if
the patient develops hypotension for any reason during the 5 days of administration. If
hypotension is transient and resolves without pharmacological intervention, CLOLAR[™]
treatment can be re-instituted, generally at a lower dose.

181

182 Concomitant Medications

183 Since CLOLAR[™] is excreted primarily by the kidneys, drugs with known renal toxicity

should be avoided during the 5 days of CLOLAR[™] administration. In addition, since the

185 liver is a known target organ for CLOLAR[™] toxicity, concomitant use of medications known

186 to induce hepatic toxicity should also be avoided. Patients taking medications known to

affect blood pressure or cardiac function should be closely monitored during administration
of CLOLAR[™].

190 Pregnancy/Nursing

- 191 All patients should be advised to use effective contraceptive measures to prevent pregnancy.
- 192 Female patients should be advised to avoid breast feeding during treatment with CLOLAR[™].
- 193

194 Laboratory Tests

- 195 Complete blood counts and platelet counts should be obtained at regular intervals during
- 196 CLOLAR[™] therapy, and more frequently in patients who develop cytopenias. In addition
- 197 liver and kidney function should be monitored frequently during the 5 days of CLOLAR™
- 198 administration.

199

200 **Drug Interactions**

201 Although no clinical drug-drug interaction studies have been conducted to date, on the basis

- 202 of the *in vitro* studies, cytochrome p450 inhibitors and inducers are unlikely to affect the
- 203 metabolism of clofarabine. The effect of clofarabine on the metabolism of cytochrome p450
- 204 substrates has not been studied.

205

206 Drug/Laboratory Tests Interactions

There are no known clinically significant interactions of CLOLAR[™] with other medications
or laboratory tests. No formal drug/laboratory test interaction studies have been conducted
with CLOLAR[™].

211 Carcinogenesis, Mutagenesis, Impairment of Fertility

212 Carcinogenesis

213 Clofarabine has not been tested for carcinogenic potential.

214

215 Mutagenesis

- 216 Clofarabine showed clastogenic activity in the *in vitro* mammalian cell chromosome
- aberration assay (CHO cells) and in the *in vivo* rat micronucleus assay. It did not show
- 218 evidence of mutagenic activity in the bacterial mutation assay (Ames test).

219

220 Impairment of Fertility

221 Studies in mice, rats, and dogs have demonstrated dose-related adverse effects on male reproductive organs. Seminiferous tubule and testicular degeneration and atrophy were 222 reported in male mice receiving IP doses of 3 mg/kg/day (9 mg/m²/day, approximately 17% 223 of clinical recommended dose on a mg/m^2 basis). The testes of rats receiving 25 mg/kg/day224 $(150 \text{ mg/m}^2/\text{day}, \text{ approximately 3 times the recommended clinical dose on a mg/m}^2 \text{ basis})$ in 225 226 a 6-month IV study had bilateral degeneration of the seminiferous epithelium with retained 227 spermatids and atrophy of interstitial cells. In a 6-month IV dog study, cell degeneration of 228 the epididymis and degeneration of the seminiferous epithelium in the testes were observed in dogs receiving 0.375 mg/kg/day (7.5 mg/m²/day, approximately 14% of the clinical 229 recommended dose on a mg/m^2 basis). Ovarian atrophy or degeneration and uterine mucosal 230 apoptosis were observed in female mice at 75 mg/kg/day (225 mg/m²/day, approximately 231 4 fold of recommended human dose on a mg/m^2 basis), the only dose administered to female 232 mice. The effect on human fertility is unknown. 233 234

235	Pregnancy
236	Teratogenic Effects: Pregnancy Category D
237	See WARNINGS.
238	
239	Nursing Mothers
240	It is not known whether clofarabine or its metabolites are excreted in human milk. Because
241	of the potential for tumorigenicity shown for clofarabine in animal studies and the potential
242	for serious adverse reactions, women treated with clofarabine should not nurse.
243	
244	Other Special Population: Adults
245	Safety and efficacy have not been established in adults. One study was performed in highly
246	refractory and/or relapsed adult patients with hematologic malignancies. The Phase 2 dose of
247	CLOLAR [™] was determined to be 40 mg/m ² /day administered as a 1- to 2-hour IVI daily × 5
248	every 28 days.
249	
250	ADVERSE REACTIONS
251	One hundred thirteen (113) pediatric patients with ALL (67) or AML (46) were exposed to
252	CLOLAR™. Ninety six (96) of the pediatric patients treated in clinical trials received the
253	recommended dose of CLOLAR TM 52 mg/m ² daily \times 5.
254	
255	The most common adverse effects after CLOLAR™ treatment, regardless of causality, were

256 gastrointestinal tract symptoms, including vomiting, nausea, and diarrhea; hematologic

- 257 effects, including anemia, leukopenia, thrombocytopenia, neutropenia, and febrile
- 258 neutropenia; and infection.

259

- 260 Table 2 lists adverse events by System Organ Class regardless of causality, including severe
- or life threatening events (NCI CTC grade 3 or grade 4), reported in $\geq 10\%$ of the 96 patients
- in the 52 mg/m²/day dose group. More detailed information and follow-up of certain events
 is given below.
- 264

	52 mg/m ² (N=96)					
System Organ Class	Т	otal	1	ade 3	Grade 4	
Adverse Event ¹	Ν	%	n	%	n	%
Blood and Lymphatic System Disorders		-	-		-	-
Febrile neutropenia	55	57	51	53	3	3
Neutropenia	10	10	3	3	7	7
Transfusion reaction	10	10	3	3		
Cardiac Disorders			-			
Tachycardia NOS	33	34	6	6		
Gastrointestinal Disorders			-			
Abdominal pain NOS	35	36	7	7		
Constipation	20	21				
Diarrhea NOS	51	53	10	10		
Gingival bleeding	14	15	7	7	1	1
Nausea	72	75	14	15	1	1
Sore throat NOS	13	14			•	
Vomiting NOS	80	83	8	8	1	1
General Disorders and Administration Site Condition	ns		•			
Edema NOS	19	20	1	1	2	2
Fatigue	35	36	3	3	1	1
Injection site pain	13	14	1	1		
Lethargy	11	11			•	
Mucosal inflammation NOS	17	18	3	3	•	
Pain NOS	18	19	6	6	1	1
Pyrexia	39	41	15	16		
Rigors	36	38	3	3		
Hepato-Biliary Disorders						
Hepatomegaly	14	15	8	8		
Jaundice NOS	14	15	2	2		
Infections and Infestations	<u>,</u>		•			
Bacteremia	10	10	10	10		
Cellulitis	11	11	9	9		
Herpes simplex	11	11	6	6		
Oral candidiasis	12	13	2	2		
Pneumonia NOS	10	10	5	5	2	2
Sepsis NOS	14	15	7	7	7	7
Staphylococcal infection NOS	12	13	10	10		
Investigations	· · · · ·		•			
Weight decreased	10	10				

Table 2:	Most Commonly Reported (>=10% Overall) Adverse Events
	by System Organ Class (N=96)

ſ

Table 2:Most Commonly Reported (>=System Organ Class (N=9)			Adve	rse Evo	ents b	у
	52 mg/m ² (N=96)					
System Organ Class	Total			ade 3		de 4
Adverse Event ¹	n	%	n	%	n	%
Metabolism and Nutrition Disorders	<u> </u>				-	
Anorexia	30	31	5	5	7	7
Appetite decreased NOS	11	11				
Musculoskeletal, Connective Tissue and Bone Disorders			•		•	
Arthralgia	11	11	3	3		
Back pain	12	13	3	3		
Myalgia	13	14				
Pain in limb	28	29	5	5		
Nervous System Disorders						
Dizziness (exc vertigo)	15	16				
Headache NOS	44	46	4	4		
Somnolence	10	10	1	1		
Tremor NEC	10	10				
Psychiatric Disorders			•		•	
Anxiety NEC	21	22	2	2		
Depression NEC	11	11	1	1		
Irritability	11	11	1	1		•
Renal and Urinary Disorders						
Hematuria	16	17	2	2		
Respiratory, Thoracic and Mediastinal Disorders						
Cough	18	19				
Dyspnea NOS	12	13	4	4	2	2
Epistaxis	30	31	14	15		
Pleural effusion	10	10	3	3	2	2
Respiratory distress	13	14	6	6	5	5
Skin and Subcutaneous Tissue Disorders						
Contusion	11	11	1	1		
Dermatitis NOS	39	41	7	7		
Dry skin	10	10	1	1		
Erythema NEC	17	18				
Palmar-plantar erythrodysesthesia syndrome		13	4	4		
Petechiae	28	29	7	7		
Pruritus NOS	45	47	1	1		
Vascular Disorders			T			
Flushing	17	18				
Hypertension NOS	11	11	4	4		
Hypotension NOS		29	12	13	7	7

¹ Patients with more than one occurrence of the same preferred term are counted only once. Grade 4 includes deaths (Grade 5).

268 Cardiovascular

- 269 The most frequently reported cardiac disorder was tachycardia (34%), which was however,
- already present in 27.4% of patients at study entry. Most of the cardiac adverse events were
- 271 reported in the first 2 cycles.

272

Pericardial effusion was a frequent finding in these patients on post-treatment studies, [19/55
(35%)]. The effusion was almost always minimal to small and in no cases had hemodynamic
significance.

276

277 Left ventricular systolic dysfunction (LVSD) was also noted. Fifteen out of fifty-five

278 patients [15/55 (27%)] had some evidence of LVSD after study entry. In most cases where

subsequent follow-up data were available, the LVSD appeared to be transient. The exact

etiology for the LVSD is unclear because of previous therapy or serious concurrent illness.

281

282 Hepatic

283 Hepato-biliary toxicities were frequently observed in pediatric patients during treatment with

284 CLOLAR[™]. Grade 3 or 4 elevated AST occurred in 38% of patients and grade 3 or 4

- elevated ALT occurred in 44% of patients. Grade 3 or 4 elevated bilirubin occurred in 15%
- of patients, with 2 cases of grade 4 hyperbilirubinemia resulting in treatment discontinuation.

- For patients with follow-up data, elevations in AST and ALT were transient and typically of
- 289 <2 weeks duration. The majority of AST and ALT elevations occurred within 1 week of
- 290 CLOLARTM administration and returned to baseline or \leq grade 2 within several days.
- 291 Although less common, elevations in bilirubin appeared to be more persistent. Where

follow-up data are available, the median time to recovery from grade 3 and grade 4

elevations in bilirubin to \leq grade 2 was 6 days.

294

295 Infection

At baseline 47% of the patients had 1 or more concurrent infections. A total of 85% of
patients experienced at least 1 infection after CLOLAR[™] treatment, including fungal, viral
and bacterial infections.

299

300 Renal

The most prevalent renal toxicity was elevated creatinine. Grade 3 or 4 elevated creatinine
occurred in 6% of patients. Nephrotoxic medications, tumor lysis, and tumor lysis with
hyperuricemia may contribute to renal toxicity.

304

305 Systemic Inflammatory Response Syndrome (SIRS)/Capillary Leak Syndrome

306 Capillary leak syndrome or SIRS (signs and symptoms of cytokine release, e.g., tachypnea, 307 tachycardia, hypotension, pulmonary edema) occurred in 4 pediatric patients overall (3 ALL, 1 AML). Several patients developed rapid onset of respiratory distress, hypotension, 308 309 capillary leak (pleural and pericardial effusions), and multi-organ failure. Close monitoring for this syndrome and early intervention are recommended. The use of prophylactic steroids 310 (eg, 100 mg/m² hydrocortisone on Days 1 through 3) may be of benefit in preventing signs or 311 symptoms of SIRS or capillary leak. Physicians should be alert to early indications of this 312 syndrome and should immediately discontinue CLOLAR[™] administration if they occur and 313 provide appropriate supportive measures. After the patient is stabilized and organ function 314 has returned to baseline, re-treatment with CLOLAR[™] can be considered at a lower dose. 315

317 Overdosage

- 318 There were no known overdoses of CLOLAR[™]. The highest daily dose administered to a
- human to date (on a mg/m² basis) has been 70 mg/m²/day \times 5 days (2 pediatric ALL
- 320 patients). The toxicities included in these 2 patients included grade 4 hyperbilirubinemia,
- 321 grade 2 and 3 vomiting, and grade 3 maculopapular rash.

322

323 DOSAGE AND ADMINISTRATION

324 **Recommended Dose**

325 CLOLAR[™] should be diluted per instructions below with 5% dextrose injection, USP or

326 0.9% sodium chloride injection, USP prior to intravenous infusion (IVI).

327

The recommended pediatric dose and schedule is 52 mg/m² administered by intravenous infusion (IVI) over 2 hours daily for 5 consecutive days. Treatment cycles are repeated

following recovery or return to baseline organ function, approximately every 2 to 6 weeks.

The dosage is based on the patient's body surface area (BSA), calculated using the actual

height and weight before the start of each cycle. To prevent drug incompatibilities, no other

333 medications should be administered through the same intravenous line.

334

335 CLOLAR[™] has not been studied in patients with hepatic or renal dysfunction. Its use in
336 such patients should be undertaken only with the greatest caution.

337

338 Physicians are encouraged to give continuous IV fluids throughout the 5 days of CLOLAR™

administration to reduce the effects of tumor lysis and other adverse events. The use of

prophylactic steroids (e.g., 100 mg/m² hydrocortisone on Days 1 through 3) may be of

341 benefit in preventing signs or symptoms of SIRS or capillary leak (e.g., hypotension). If 342 patients show early signs or symptoms of SIRS or capillary leak (e.g., hypotension), the 343 physician should immediately discontinue CLOLAR[™] administration and provide 344 appropriate supportive measures. Close monitoring of renal and hepatic function during the 5 days of CLOLAR[™] administration is advised. If substantial increases in creatinine or 345 346 bilirubin are noted, physicians should immediately discontinue administration of 347 CLOLAR[™]. CLOLAR[™] should be re-instituted when the patient is stable and organ function has returned to baseline, possibly at a lower dose. If hyperuricemia is anticipated 348 349 (tumor lysis), patients should prophylactically receive allopurinol. 350

351

352 STORAGE AND HANDLING

Vials containing undiluted CLOLAR[™] should be stored at 25°C (77°F); excursions permitted
to 15-30°C (59-86°F).

355

356 CLOLAR[™] should be filtered through a sterile 0.2 µm syringe filter and then further diluted
357 with 5% dextrose injection USP or 0.9% sodium chloride injection USP prior to intravenous
358 infusion (IVI). The resulting admixture may be stored at room temperature, but must be used
359 within 24 hours of preparation.

360

361 HOW SUPPLIED

362 CLOLAR[™] is formulated at a concentration of 1 mg/mL in sodium chloride (9 mg/mL),

363 USP, and water for injection, USP, quantity sufficient (qs) to 1 mL. CLOLAR™ is supplied

in 20 mL flint vials in a box of 4 (NDC 58468-0100-2). The 20 mL flint vials contain 20 mL

365 (20 mg) of solution. The pH range of the solution is 4.5 to 7.5. The solution is clear and

366 practically colorless, is preservative free, and is free from foreign matter.

367		
368	Rx only	
369	U.S. Patents: 4,751,	221; 4, 918,179; 5,384,310; 5,661,136, 6,680,382 B2.
370	Other patents pending	g.
371		
372	NAME AND ADDR	ESS OF MANUFACTURER
373	Manufactured by:	AAI Development Services
374		Charleston, SC 29405
375	Manufactured for:	Genzyme Corporation
376		4545 Horizon Hill Blvd
377		San Antonio, TX 78229
378	Distributed by:	Genzyme Corporation
379		500 Kendall Street
380		Cambridge, MA 02142
381		