

2	PEGASYS â
3	(peginterferon alfa-2a

Alpha interferons, including PEGASYS (peginterferon alfa-2a), may cause or aggravate fatal or life-threatening neuropsychiatric, autoimmune, ischemic, and infectious disorders. Patients should be monitored closely with periodic clinical and laboratory evaluations. Therapy should be withdrawn in patients with persistently severe or worsening signs or symptoms of these conditions. In many, but not all cases, these disorders resolve after stopping PEGASYS therapy (see WARNINGS and ADVERSE REACTIONS).

Use with Ribavirin. Ribavirin, including COPEGUS, may cause birth defects and/or death of the fetus. Extreme care must be taken to avoid pregnancy in female patients and in female partners of male patients. Ribavirin causes hemolytic anemia. The anemia associated with ribavirin therapy may result in a worsening of cardiac disease. Ribavirin is genotoxic and mutagenic and should be considered a potential carcinogen (see COPEGUS Package Insert for additional information and other WARNINGS).

DESCRIPTION

- 19 PEGASYS, peginterferon alfa-2a, is a covalent conjugate of recombinant alfa-2a
- 20 interferon (approximate molecular weight [MW] 20,000 daltons) with a single branched
- bis-monomethoxy polyethylene glycol (PEG) chain (approximate MW 40,000 daltons).
- The PEG moiety is linked at a single site to the interferon alfa moiety via a stable amide
- bond to lysine. Peginterferon alfa-2a has an approximate molecular weight of 60,000
- daltons. Interferon alfa-2a is produced using recombinant DNA technology in which a
- 25 cloned human leukocyte interferon gene is inserted into and expressed in Escherichia
- 26 *coli*.

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- Each vial contains approximately 1.2 mL of solution to deliver 1.0 mL of drug product.
- 28 Subcutaneous (sc) administration of 1.0 mL delivers 180 µg of drug product (expressed
- as the amount of interferon alfa-2a), 8.0 mg sodium chloride, 0.05 mg polysorbate 80,
- 30 10.0 mg benzyl alcohol, 2.62 mg sodium acetate trihydrate, and 0.05 mg acetic acid. The
- 31 solution is colorless to light yellow and the pH is 6.0 ± 0.01 .

CLINICAL PHARMACOLOGY

Pharmacodynamics

- 34 Interferons bind to specific receptors on the cell surface initiating intracellular signaling
- 35 via a complex cascade of protein-protein interactions leading to rapid activation of gene
- 36 transcription. Interferon-stimulated genes modulate many biological effects including the
- 37 inhibition of viral replication in infected cells, inhibition of cell proliferation, and
- immunomodulation. The clinical relevance of these in vitro activities is not known.

- 39 PEGASYS stimulates the production of effector proteins such as serum neopterin and 2',
- 40 5'-oligoadenylate synthetase.

41 Pharmacokinetics

- 42 Maximal serum concentrations (C_{max}) occur between 72 to 96 hours post dose. The C_{max}
- and AUC measurements of PEGASYS increase in a dose-related manner. Week 48 mean
- 44 trough concentrations (16 ng/mL; range 4 to 28) at 168 hours post dose are approximately
- 45 2-fold higher than week 1 mean trough concentrations (8 ng/mL; range 0 to 15). Steady-
- state serum levels are reached within 5 to 8 weeks of once weekly dosing. The peak to
- 47 trough ratio at week 48 is approximately 2.0.
- The mean systemic clearance in healthy subjects given PEGASYS was 94 mL/h, which is
- 49 approximately 100-fold lower than that for interferon alfa-2a (ROFERON®-A). The
- mean terminal half-life after sc dosing in patients with chronic hepatitis C was 80 hours
- 51 (range 50 to 140 hours) compared to 5.1 hours (range 3.7 to 8.5 hours) for
- 52 ROFERON®-A.

53 **Special Populations**

- 54 Gender and Age
- 55 PEGASYS administration yielded similar pharmacokinetics in male and female healthy
- subjects. The AUC was increased from 1295 to 1663 ng·h/mL in subjects older than 62
- 57 years taking 180 μg PEGASYS, but peak concentrations were similar (9 vs 10 ng/mL) in
- those older and younger than 62 years.
- 59 Pediatric Patients
- The pharmacokinetics of PEGASYS have not been adequately studied in pediatric
- 61 patients.
- 62 Renal Dysfunction
- In patients with end stage renal disease undergoing hemodialysis, there is a 25% to 45%
- reduction in PEGASYS clearance (see **PRECAUTIONS: Renal Impairment**).
- The pharmacokinetics of ribavirin following administration of COPEGUS have not been
- studied in patients with renal impairment and there are limited data from clinical trials on
- 67 administration of COPEGUS in patients with creatinine clearance <50 mL/min.
- Therefore, patients with creatinine clearance <50 mL/min should not be treated with
- 69 COPEGUS (see WARNINGS and DOSAGE AND ADMINISTRATION).
- 70 Effect of Food on Absorption of Ribavirin
- Bioavailability of a single oral dose of ribavirin was increased by co-administration with
- 72 a high-fat meal. The absorption was slowed (T_{max} was doubled) and the AUC_{0-192h} and
- 73 C_{max} increased by 42% and 66%, respectively, when COPEGUS was taken with a high-
- fat meal compared with fasting conditions (see **DOSAGE AND ADMINISTRATION**).

75 **Drug Interactions**

- 76 Nucleoside Analogues
- Ribavirin has been shown in vitro to inhibit phosphorylation of zidovudine and stavudine
- 78 which could lead to decreased anti-retroviral activity. Exposure to didanosine or its active
- 79 metabolite (dideoxyadenosine 5'-triphosphate) is increased when didanosine is co-
- administered with ribavirin (see **PRECAUTIONS: Drug Interactions**).

81 **CLINICAL STUDIES**

82 **PEGASYS Monotherapy (Studies 1, 2, and 3)**

- 83 The safety and effectiveness of PEGASYS for the treatment of hepatitis C virus infection
- 84 were assessed in three randomized, open-label, active-controlled clinical studies. All
- patients were adults, had compensated liver disease, detectable hepatitis C virus (HCV),
- liver biopsy diagnosis of chronic hepatitis, and were previously untreated with interferon.
- 87 All patients received therapy by sc injection for 48 weeks, and were followed for an
- additional 24 weeks to assess the durability of response. In studies 1 and 2, approximately
- 89 20% of subjects had cirrhosis or bridging fibrosis. Study 3 enrolled patients with a
- 90 histological diagnosis of cirrhosis (78%) or bridging fibrosis (22%).
- 91 In study 1 (n=630), patients received either ROFERON-A (interferon alfa-2a) 3MIU
- 92 three times/week (tiw), PEGASYS 135 µg once each week (qw) or PEGASYS 180 µg
- 93 qw. In study 2 (n=526), patients received either ROFERON-A 6 MIU tiw for 12 weeks
- 94 followed by 3 MIU tiw for 36 weeks or PEGASYS 180 µg qw. In study 3 (n=269),
- 95 patients received ROFERON-A 3 MIU tiw, PEGASYS 90 µg qw or PEGASYS 180 µg
- 96 once each week.
- In all three studies, treatment with PEGASYS 180 µg resulted in significantly more
- 98 patients who experienced a sustained response (defined as undetectable HCV RNA and
- 99 normalization of ALT on or after study week 68) compared to treatment with
- 100 ROFERON-A. In study 1, response to PEGASYS 135 µg was not different from response
- 101 to 180 μg. In study 3, response to PEGASYS 90 μg was intermediate between PEGASYS
- 102 180 μg and ROFERON-A.

Table 1 Sustained Response to Monotherapy Treatment

	Study 1				Study 2			Study 3		
	ROFERON-A 3 MIU	PEGASYS 180 mg	DIFF* (95% CI)	ROFERON-A 6/3 MIU	PEGASYS 180 mg	DIFF* (95% CI)	ROFERON-A 3 MIU	PEGASYS 180 mg	DIFF* (95% CI)	
	(N=207)	(N=208)		(N=261)	(N=265)		(N=86)	(N=87)		
Combined Virologic and Biologic Sustained Response	11%	24%	13 (6, 20)	17%	35%	18 (11, 25)	7%	23%	16 (6, 26)	
Sustained Virologic Response**	11%	26%	15 (8, 23)	19%	38%	19 (11, 26)	8%	30%	22 (11, 33)	

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Matched pre- and post-treatment liver biopsies were obtained in approximately 70% of patients. Similar modest reductions in inflammation compared to baseline were observed in all treatment groups.

- Of the patients who did not demonstrate either undetectable HCV RNA or at least a
- 113 2-log10 drop in HCV RNA titer from baseline by 12 weeks of PEGASYS 180 μg
- therapy, 2% (3/156) achieved a sustained virologic response (see **DOSAGE AND**
- 115 **ADMINISTRATION**).
- Averaged over study 1, study 2, and study 3, response rates to PEGASYS were 23%
- among patients with viral genotype 1 and 48% in patients with other viral genotypes. The
- treatment response rates were similar in men and women.

PEGASYS/COPEGUS Combination Therapy (Studies 4 and 5)

- 120 The safety and effectiveness of PEGASYS in combination with COPEGUS for the
- treatment of hepatitis C virus infection were assessed in two randomized controlled
- 122 clinical trials. All patients were adults, had compensated liver disease, detectable hepatitis
- 123 C virus, liver biopsy diagnosis of chronic hepatitis, and were previously untreated with
- interferon. Approximately 20% of patients in both studies had compensated cirrhosis
- 125 (Child-Pugh class A).
- In study 4, patients were randomized to receive either PEGASYS 180 µg sc once weekly
- 127 (qw) with an oral placebo, PEGASYS 180 µg qw with COPEGUS 1000 mg po (body
- weight <75 kg) or 1200 mg po (body weight ≥75 kg) or REBETRON™ (interferon alfa-
- 2b 3 MIU sc tiw plus ribavirin 1000 mg or 1200 mg po). All patients received 48 weeks
- of therapy followed by 24 weeks of treatment-free follow-up. COPEGUS or placebo
- treatment assignment was blinded. PEGASYS in combination with COPEGUS resulted
- in a higher SVR (defined as undetectable HCV RNA at the end of the 24-week treatment-
- free follow-up period) compared to PEGASYS alone or interferon alfa-2b and ribavirin

^{*}Percent difference between PEGASYS and Roferon-A treatment

^{**}COBAS AMPLICOR® HCV Test, version 2.0

134 (Table 2). In all treatment arms, patients with viral genotype 1 regardless of viral load,

had a lower response rate compared to patients with other viral genotypes.

Table 2 Sustained Virologic Response to Combination Therapy (Study 4)

	Interferon alfa-2b+ Ribavirin 1000 mg or 1200 mg	PEGASYS + placebo	PEGASYS + COPEGUS 1000 mg or 1200 mg
All patients	197/444 (44%)	65/224 (29%)	241/453 (53%)
Genotype 1	103/285 (36%)	29/145 (20%)	132/298 (44%)
Genotypes 2-6	94/159 (59%)	36/79 (46%)	109/155 (70%)

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Difference in overall treatment response (PEGASYS/COPEGUS – Interferon alfa -2b/ribavirin) was 9% (95% CI 2.3, 15.3).

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In study 5, all patients received PEGASYS 180 μ g sc qw and were randomized to treatment for either 24 or 48 weeks and to a COPEGUS dose of either 800 mg or 1000 mg/1200 mg (for body weight <75 kg / \geq 75 kg). Assignment to the four treatment arms was stratified by viral genotype and baseline HCV viral titer. Patients with genotype 1 and high viral titer (defined as >2 x 10⁶ HCV RNA copies/mL serum) were preferentially assigned to treatment for 48 weeks.

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148 Genotype 1

149 Irrespective of baseline viral titer, treatment for 48 weeks with PEGASYS and 1000 mg

or 1200 mg of COPEGUS resulted in higher SVR (defined as undetectable HCV RNA at

the end of the 24-week treatment-free follow-up period) compared to shorter treatment

152 (24 weeks) and/or 800 mg COPEGUS.

153 Genotype non-1

154 Irrespective of baseline viral titer, treatment for 24 weeks with PEGASYS and 800 mg of

155 COPEGUS resulted in a similar SVR compared to longer treatment (48 weeks) and/or

156 1000 mg or 1200 mg of COPEGUS (see Table 3).

Table 3 Sustained Virologic Response as a Function of Genotype (Study 5)

24 Weeks	s Treatment	48 Weeks Treatment		
PEGASYS +	PEGASYS +	PEGASYS +	PEGASYS +	
COPEGUS	COPEGUS	COPEGUS	COPEGUS	
800 mg	1000 mg or 1200 mg*	800 mg	1000 mg or 1200 mg*	
(N=207)	(N=280)	(N=361)	(N=436)	

Genotype 1	29/101 (29%)	48/118 (41%)	99/250 (40%)	138/271 (51%)
Genotype 2-3	79/96 (82%)	116/144 (81%)	75/ 99 (76%)	117/153 (76%)

159 *1000 mg for body weight <75 kg; 1200 mg for body weight ≥75 kg.

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Among the 36 patients with genotype 4, response rates were similar to those observed in patients with genotype 1 (data not shown). The numbers of patients with genotype 5 and 6 were too few to allow for meaningful assessment.

164 Treatment Response in Patient Subgroups

- 165 Treatment response rates are lower in patients with poor prognostic factors receiving
- pegylated interferon alpha therapy. In studies 4 and 5, treatment response rates were
- lower in patients older than 40 years (50% vs 66%), in patients with cirrhosis (47% vs
- 168 59%), in patients weighing over 85 kg (49% vs 60%), and in patients with genotype 1
- with high vs low viral load (43% vs 56%). African American patients had lower response
- 170 rates compared to Caucasians.
- Paired liver biopsies were performed on approximately 20% of patients in Studies 4 and
- 5. Modest reductions in inflammation compared to baseline were seen in all treatment
- 173 groups.
- 174 In studies 4 and 5, lack of early virologic response at 12 weeks (defined as HCV RNA
- 175 undetectable or >2log10 lower than baseline) was grounds for discontinuation of
- 176 treatment. Of patients who lacked an early viral response at 12 weeks and completed a
- 177 recommended course of therapy despite a protocol-defined option to discontinue therapy,
- 178 5/39 (13%) achieved an SVR. Of patients who lacked an early viral response at 24 weeks,
- 179 nineteen completed a full course of therapy and none achieved an SVR.

180 INDICATIONS AND USAGE

- PEGASYS, peginterferon alfa-2a, alone or in combination with COPEGUS, is indicated
- for the treatment of adults with chronic hepatitis C virus infection who have compensated
- liver disease and have not been previously treated with interferon alpha.

184 CONTRAINDICATIONS

- 185 PEGASYS is contraindicated in patients with:
- hypersensitivity to PEGASYS or any of its components
- autoimmune hepatitis
- hepatic decompensation (Child-Pugh class B and C) before or during treatment
- 189 PEGASYS is contraindicated in neonates and infants because it contains benzyl alcohol.
- Benzyl alcohol is associated with an increased incidence of neurologic and other
- complications in neonates and infants, which are sometimes fatal.
- 192 PEGASYS and COPEGUS combination therapy is additionally contraindicated in:

- Patients with known hypersensitivity to COPEGUS or to any component of the tablet.
- Women who are pregnant.
- Men whose female partners are pregnant.
- Patients with hemoglobinopathies (eg, thalassemia major, sickle-cell anemia).

197 WARNINGS

198 General

- 199 Patients should be monitored for the following serious conditions, some of which may
- 200 become life threatening. Patients with persistently severe or worsening signs or
- symptoms should have their therapy withdrawn (see **BOXED WARNING**).

202 **Neuropsychiatric**

- 203 Life-threatening or fatal neuropsychiatric reactions may manifest in patients receiving
- 204 therapy with PEGASYS and include suicide, suicidal ideation, depression, relapse of
- 205 drug addiction and drug overdose. These reactions may occur in patients with and
- without previous psychiatric illness.
- 207 PEGASYS should be used with extreme caution in patients who report a history of
- depression. Neuropsychiatric adverse events observed with alpha interferon treatment
- 209 include aggressive behavior, psychoses, hallucinations, bipolar disorders and mania.
- 210 Physicians should monitor all patients for evidence of depression and other psychiatric
- 211 symptoms. Patients should be advised to report any sign or symptom of depression or
- suicidal ideation to their prescribing physicians. In severe cases, therapy should be
- 213 stopped immediately and psychiatric intervention instituted (see ADVERSE
- 214 **REACTIONS** and **DOSAGE AND ADMINISTRATION**).

215 Infections

- 216 Serious and severe bacterial infections, some fatal, have been observed in patients treated
- 217 with alpha interferons including PEGASYS. Some of the infections have been associated
- 218 with neutropenia. PEGASYS should be discontinued in patients who develop severe
- 219 infections and appropriate antibiotic therapy instituted.

220 Bone Marrow Toxicity

- 221 PEGASYS suppresses bone marrow function and may result in severe cytopenias.
- Ribayirin may potentiate the neutropenia and lymphopenia induced by alpha interferons
- 223 including PEGASYS. Very rarely alpha interferons may be associated with aplastic
- anemia. It is advised that complete blood counts (CBC) be obtained pre-treatment and
- 225 monitored routinely during therapy (see **PRECAUTIONS: Laboratory Tests**).
- 226 PEGASYS and COPEGUS should be used with caution in patients with baseline
- neutrophil counts <1500 cells/mm³, with baseline platelet counts <90,000 cells/mm³ or
- 228 baseline hemoglobin <10 g/dL. PEGASYS therapy should be discontinued, at least

- temporarily, in patients who develop severe decreases in neutrophil and/or platelet counts
- 230 (see **DOSAGE AND ADMINISTRATION: Dose Modifications**).

231 Cardiovascular Disorders

- 232 Hypertension, supraventricular arrhythmias, chest pain, and myocardial infarction have
- been observed in patients treated with PEGASYS.
- 234 PEGASYS should be administered with caution to patients with preexisting cardiac
- disease. Because cardiac disease may be worsened by ribavirin-induced anemia, patients
- with a history of significant or unstable cardiac disease should not use COPEGUS (see
- 237 WARNING: Anemia and COPEGUS Package Insert).

238 **Hypersensitivity**

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- 239 Severe acute hypersensitivity reactions (eg, urticaria, angioedema, bronchoconstriction,
- anaphylaxis) have been rarely observed during alpha interferon and ribavirin therapy. If
- such reaction occurs, therapy with PEGASYS and COPEGUS should be discontinued
- and appropriate medical therapy immediately instituted.

Endocrine Disorders

- 244 PEGASYS causes or aggravates hypothyroidism and hyperthyroidism. Hyperglycemia,
- 245 hypoglycemia, and diabetes mellitus have been observed to develop in patients treated
- 246 with PEGASYS. Patients with these conditions at baseline who cannot be effectively
- treated by medication should not begin PEGASYS therapy. Patients who develop these
- 248 conditions during treatment and cannot be controlled with medication may require
- 249 discontinuation of PEGASYS therapy.

250 Autoimmune Disorders

- 251 Development or exacerbation of autoimmune disorders including myositis, hepatitis, ITP,
- 252 psoriasis, rheumatoid arthritis, interstitial nephritis, thyroiditis, and systemic lupus
- 253 erythematosus have been reported in patients receiving alpha interferon. PEGASYS
- should be used with caution in patients with autoimmune disorders.

255 **Pulmonary Disorders**

- 256 Dyspnea, pulmonary infiltrates, pneumonia, bronchiolitis obliterans, interstitial
- 257 pneumonitis and sarcoidosis, some resulting in respiratory failure and/or patient deaths,
- 258 may be induced or aggravated by PEGASYS or alpha interferon therapy. Patients who
- 259 develop persistent or unexplained pulmonary infiltrates or pulmonary function
- 260 impairment should discontinue treatment with PEGASYS.

261 Colitis

- 262 Ulcerative, and hemorrhagic/ischemic colitis, sometimes fatal, have been observed within
- 263 12 weeks of starting alpha interferon treatment. Abdominal pain, bloody diarrhea, and
- 264 fever are the typical manifestations of colitis. PEGASYS should be discontinued

- immediately if these symptoms develop. The colitis usually resolves within 1 to 3 weeks
- of discontinuation of alpha interferon.

267 Pancreatitis

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- 268 Pancreatitis, sometimes fatal, has occurred during alpha interferon and ribavirin
- 269 treatment. PEGASYS and COPEGUS should be suspended if symptoms or signs
- 270 suggestive of pancreatitis are observed. PEGASYS and COPEGUS should be
- 271 discontinued in patients diagnosed with pancreatitis.

Ophthalmologic Disorders

- 273 Decrease or loss of vision, retinopathy including macular edema, retinal artery or vein
- 274 thrombosis, retinal hemorrhages and cotton wool spots, optic neuritis, and papilledema
- are induced or aggravated by treatment with PEGASYS or other alpha interferons. All
- 276 patients should receive an eye examination at baseline. Patients with preexisting
- 277 ophthalmologic disorders (eg, diabetic or hypertensive retinopathy) should receive
- 278 periodic ophthalmologic exams during interferon alpha treatment. Any patient who
- 279 develops ocular symptoms should receive a prompt and complete eye examination.
- 280 PEGASYS treatment should be discontinued in patients who develop new or worsening
- 281 ophthalmologic disorders.
- 282 Use With Ribavirin (Also, see COPEGUS Package Insert.)
- 283 Ribavirin may cause birth defects and/or death of the exposed fetus.
- 284 Extreme care must be taken to avoid pregnancy in female patients and in
- 285 female partners of male patients taking PEGASYS and COPEGUS
- 286 combination therapy. COPEGUS THERAPY SHOULD NOT BE STARTED
- 287 UNLESS A REPORT OF A NEGATIVE PREGNANCY TEST HAS BEEN
- 288 OBTAINED IMMEDIATELY PRIOR TO INITIATION OF THERAPY. Women of
- 289 childbearing potential and men must use two forms of effective
- 290 contraception during treatment and for at least six months after treatment
- 291 has concluded. Routine monthly pregnancy tests must be performed
- 292 during this time (see BOXED WARNING, CONTRAINDICATIONS,
- 293 PRECAUTIONS: Information for Patients, and COPEGUS Package Insert).

294 Anemia

- 295 The primary toxicity of ribavirin is hemolytic anemia. Hemoglobin <10 g/dL was
- observed in approximately 13% of COPEGUS and PEGASYS treated patients in clinical
- 297 trials (see PRECAUTIONS: Laboratory Tests). The anemia associated with
- 298 COPEGUS occurs within 1 to 2 weeks of initiation of therapy with maximum drop in
- 299 hemoglobin observed during the first eight weeks. BECAUSE THE INITIAL DROP IN
- 300 HEMOGLOBIN MAY BE SIGNIFICANT, IT IS ADVISED THAT HEMOGLOBIN
- 301 OR HEMATOCRIT BE OBTAINED PRETREATMENT AND AT WEEK 2 AND
- 302 WEEK 4 OF THERAPY OR MORE FREQUENTLY IF CLINICALLY INDICATED.
- Patients should then be followed as clinically appropriate.

- 304 Fatal and nonfatal myocardial infarctions have been reported in patients with anemia
- 305 caused by ribavirin. Patients should be assessed for underlying cardiac disease before
- 306 initiation of ribavirin therapy. Patients with pre-existing cardiac disease should have
- 307 electrocardiograms administered before treatment, and should be appropriately monitored
- during therapy. If there is any deterioration of cardiovascular status, therapy should be
- 309 suspended or discontinued (see **DOSAGE AND ADMINISTRATION: COPEGUS**
- 310 **Dose Modification Guidelines**). Because cardiac disease may be worsened by drug-
- induced anemia, patients with a history of significant or unstable cardiac disease should
- 312 not use COPEGUS (see **COPEGUS Package Insert**).

313 Renal

- 314 It is recommended that renal function be evaluated in all patients started on COPEGUS.
- 315 COPEGUS should not be administered to patients with creatinine clearance
- 316 <50 mL/minute (see CLINICAL PHARMACOLOGY: Special Populations).

317 **PRECAUTIONS**

318 General

- 319 The safety and efficacy of PEGASYS alone or in combination with COPEGUS for the
- 320 treatment of hepatitis C have not been established in
- Patients who have failed other alpha interferon treatments
- Liver or other organ transplant recipients
- Patients co-infected with human immunodeficiency virus (HIV) or hepatitis B virus
- 324 (HBV)

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325 Renal Impairment

- 326 A 25% to 45% higher exposure to PEGASYS is seen in subjects undergoing
- hemodialysis. In patients with impaired renal function, signs and symptoms of interferon
- 328 toxicity should be closely monitored. Doses of PEGASYS should be adjusted
- accordingly. PEGASYS should be used with caution in patients with creatinine clearance
- 330 <50 mL/min (see **DOSAGE AND ADMINISTRATION: Dose Modifications**).

Information for Patients

- 332 Patients receiving PEGASYS alone or in combination with COPEGUS should be
- directed in its appropriate use, informed of the benefits and risks associated with
- treatment, and referred to the PEGASYS and, if applicable, COPEGUS (ribavirin)
- 335 MEDICATION GUIDES.
- PEGASYS and COPEGUS combination therapy must not be used by women who are
- pregnant or by men whose female partners are pregnant. COPEGUS therapy should not
- be initiated until a report of a negative pregnancy test has been obtained immediately
- before starting therapy. Female patients of childbearing potential and male patients with
- female partners of childbearing potential must be advised of the teratogenic/embryocidal

- risks and must be instructed to practice effective contraception during COPEGUS therapy
- and for 6 months posttherapy. Patients should be advised to notify the physician
- 343 immediately in the event of a pregnancy (see CONTRAINDICATIONS and
- 344 **WARNINGS**).
- Women of childbearing potential and men must use two forms of effective contraception
- during treatment and during the 6 months after treatment has concluded; routine monthly
- pregnancy tests must be performed during this time (see **CONTRAINDICATIONS** and
- 348 **COPEGUS Package Insert**).
- 349 If pregnancy does occur during treatment or during 6 months post-therapy, the patient
- must be advised of the significant teratogenic risk of COPEGUS therapy to the fetus. To
- 351 monitor maternal-fetal outcomes of pregnant women exposed to COPEGUS, the
- 352 COPEGUS Pregnancy Registry has been established. Physicians and patients are strongly
- encouraged to register by calling 1-800-526-6367.
- Patients should be advised that laboratory evaluations are required before starting therapy
- and periodically thereafter (see Laboratory Tests). Patients should be instructed to
- 356 remain well hydrated, especially during the initial stages of treatment. Patients should be
- advised to take COPEGUS with food.
- 358 Patients should be informed that it is not known if therapy with PEGASYS alone or in
- 359 combination with COPEGUS will prevent transmission of HCV infection to others or
- 360 prevent cirrhosis, liver failure or liver cancer that might result from HCV infection.
- Patients who develop dizziness, confusion, somnolence, and fatigue should be cautioned
- 362 to avoid driving or operating machinery.
- 363 If home use is prescribed, a puncture-resistant container for the disposal of used needles
- and syringes should be supplied to the patients. Patients should be thoroughly instructed
- in the importance of proper disposal and cautioned against any reuse of any needles and
- syringes. The full container should be disposed of according to the directions provided by
- the physician (see **MEDICATION GUIDE**).

Laboratory Tests

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- 369 Before beginning PEGASYS or PEGASYS and COPEGUS combination therapy,
- 370 standard hematological and biochemical laboratory tests are recommended for all
- patients. Pregnancy screening for women of childbearing potential must be performed.
- 372 After initiation of therapy, hematological tests should be performed at 2 weeks and 4
- weeks and biochemical tests should be performed at 4 weeks. Additional testing should
- be performed periodically during therapy. In the clinical studies, the CBC (including
- hemoglobin level and white blood cell and platelet counts) and chemistries (including
- liver function tests and uric acid) were measured at 1, 2, 4, 6, and 8, and then every 4
- weeks or more frequently if abnormalities were found. Thyroid stimulating hormone
- 378 (TSH) was measured every 12 weeks. Monthly pregnancy testing should be performed
- during combination therapy and for 6 months after discontinuing therapy.
- 380 The entrance criteria used for the clinical studies of PEGASYS may be considered as a
- 381 guideline to acceptable baseline values for initiation of treatment:

- Platelet count ≥90,000 cells/mm³ (as low as 75,000 cells/mm³ in patients with cirrhosis)
- Caution should be exercised in initiating treatment in any patient with baseline risk of severe anemia (eg spherocytosis, history of GI bleeding).
- Absolute neutrophil count (ANC) ≥1500 cells/mm³
- Serum creatinine concentration <1.5 x upper limit of normal
- TSH and T₄ within normal limits or adequately controlled thyroid function
- 389 PEGASYS treatment was associated with decreases in WBC, ANC, lymphocytes and
- 390 platelet counts often starting within the first 2 weeks of treatment (see ADVERSE
- 391 **REACTIONS**). Dose reduction is recommended in patients with hematologic
- abnormalities (see **DOSAGE AND ADMINISTRATION: Dose Modifications**).
- While fever is commonly caused by PEGASYS therapy, other causes of persistent fever
- 394 must be ruled out, particularly in patients with neutropenia (see WARNINGS:
- 395 **Infections**).
- 396 Transient elevations in ALT (2-fold to 5-fold above baseline) were observed in some
- 397 patients receiving PEGASYS, and were not associated with deterioration of other liver
- function tests. When the increase in ALT levels is progressive despite dose reduction or
- 399 is accompanied by increased bilirubin, PEGASYS therapy should be discontinued (see
- 400 **DOSAGE AND ADMINISTRATION: Dose Modifications**).

401 **Drug Interactions**

- 402 Treatment with PEGASYS once weekly for 4 weeks in healthy subjects was associated
- with an inhibition of P450 1A2 and a 25% increase in the ophylline AUC. The ophylline
- 404 serum levels should be monitored and appropriate dose adjustments considered for
- patients given both theophylline and PEGASYS (see **PRECAUTIONS**). There was no
- 406 effect on the pharmacokinetics of representative drugs metabolized by CYP 2C9, CYP
- 407 2C19, CYP 2D6 or CYP 3A4. In patients with chronic hepatitis C treated with
- 408 PEGASYS in combination with COPEGUS, PEGASYS treatment did not affect ribavirin
- 409 distribution or clearance.

410 Nucleoside Analogues

- 411 Didanosine
- 412 Co-administration of COPEGUS and didanosine is not recommended. Reports of fatal
- 413 hepatic failure, as well as peripheral neuropathy, pancreatitis, and symptomatic
- 414 hyperlactatemia/lactic acidosis have been reported in clinical trials (see **CLINICAL**
- 415 **PHARMACOLOGY: Drug Interactions**).

- 416 Stavudine and Zidovudine
- 417 Ribavirin can antagonize the in vitro antiviral activity of stavudine and zidovudine
- against HIV. Therefore, concomitant use of ribavirin with either of these drugs should be
- 419 avoided.

420 Carcinogenesis, Mutagenesis, Impairment of Fertility

- 421 Carcinogenesis
- 422 PEGASYS has not been tested for its carcinogenic potential.
- 423 Mutagenesis
- 424 PEGASYS did not cause DNA damage when tested in the Ames bacterial mutagenicity
- assay and in the in vitro chromosomal aberration assay in human lymphocytes, either in
- 426 the presence or absence of metabolic activation.
- 427 Use With Ribavirin
- 428 Ribavirin is genotoxic and mutagenic. The carcinogenic potential of ribavirin has not
- been fully determined. In a p53 (+/-) mouse carcinogenicity study at doses up to the
- 430 maximum tolerated dose of 100 mg/kg/day ribavirin was not oncogenic. However, on a
- body surface area basis, this dose was 0.5 times maximum recommended human 24-hour
- dose of ribavirin. A study in rats to assess the carcinogenic potential of ribavirin is
- 433 ongoing.
- 434 Mutagenesis (see **COPEGUS Package Insert**)
- 435 Impairment of Fertility
- 436 PEGASYS may impair fertility in women. Prolonged menstrual cycles and/or
- 437 amenorrhea were observed in female cynomolgus monkeys given SC injections of
- 438 600 ug/kg/dose (7200 ug/m²/dose) of PEGASYS every other day for one month, at
- approximately 180 times the recommended weekly human dose for a 60 kg person (based
- on body surface area). Menstrual cycle irregularities were accompanied by both a
- 441 decrease and delay in the peak 17β -estradiol and progesterone levels following
- administration of PEGASYS to female monkeys. A return to normal menstrual rhythm
- followed cessation of treatment. Every other day dosing with 100 µg/kg (1200 µg/m²)
- PEGASYS (equivalent to approximately 30 times the recommended human dose) had no
- effects on cycle duration or reproductive hormone status.
- The effects of PEGASYS on male fertility have not been studied. However, no adverse
- effects on fertility were observed in male Rhesus monkeys treated with non-pegylated
- interferon alfa-2a for 5 months at doses up to 25 x 10⁶ IU/kg/day (see **COPEGUS**
- 449 **Package Insert**).

450 Pregnancy

- 451 Pregnancy: Category C
- 452 PEGASYS has not been studied for its teratogenic effect. Non-pegylated interferon alfa-
- 453 2a treatment of pregnant Rhesus monkeys at approximately 20 to 500 times the human
- 454 weekly dose resulted in a statistically significant increase in abortions. No teratogenic
- effects were seen in the offspring delivered at term. PEGASYS should be assumed to
- 456 have abortifacient potential. There are no adequate and well-controlled studies of
- 457 PEGASYS in pregnant women. PEGASYS is to be used during pregnancy only if the
- potential benefit justifies the potential risk to the fetus. PEGASYS is recommended for
- 459 use in women of childbearing potential only when they are using effective contraception
- 460 during therapy.

461 Pregnancy: Category X: Use With Ribavirin (see CONTRAINDICATIONS)

- 462 Significant teratogenic and/or embryocidal effects have been demonstrated in all
- animal species exposed to ribavirin. COPEGUS therapy is contraindicated in
- 464 women who are pregnant and in the male partners of women who are pregnant (see
- 465 CONTRAINDICATIONS, WARNINGS, and COPEGUS Package Insert).
- 466 If pregnancy occurs in a patient or partner of a patient during treatment or during the 6
- 467 months after treatment cessation, such cases should be reported to the COPEGUS
- 468 Pregnancy Registry at 1-800-526-6367.

Nursing Mothers

469

- 470 It is not known whether peginterferon or ribavirin or its components are excreted in
- 471 human milk. The effect of orally ingested peginterferon or ribavirin from breast milk on
- 472 the nursing infant has not been evaluated. Because of the potential for adverse reactions
- 473 from the drugs in nursing infants, a decision must be made whether to discontinue
- 474 nursing or discontinue PEGASYS and COPEGUS treatment.

475 **Pediatric Use**

- The safety and effectiveness of PEGASYS, alone or in combination with COPEGUS in
- patients below the age of 18 years have not been established.
- 478 PEGASYS contains benzyl alcohol. Benzyl alcohol has been reported to be associated
- 479 with an increased incidence of neurological and other complications in neonates and
- infants, which are sometimes fatal (see **CONTRAINDICATIONS**).

481 Geriatric Use

- 482 Younger patients have higher virologic response rates than older patients. Clinical studies
- 483 of PEGASYS alone or in combination with COPEGUS did not include sufficient
- numbers of subjects aged 65 or over to determine whether they respond differently from
- 485 younger subjects. Adverse reactions related to alpha interferons, such as CNS, cardiac,
- and systemic (eg, flu-like) effects may be more severe in the elderly and caution should

- be exercised in the use of PEGASYS in this population. PEGASYS and COPEGUS are
- excreted by the kidney, and the risk of toxic reactions to this therapy may be greater in
- patients with impaired renal function. Because elderly patients are more likely to have
- decreased renal function, care should be taken in dose selection and it may be useful to
- 491 monitor renal function. PEGASYS should be used with caution in patients with creatinine
- 492 clearance <50 mL/min and COPEGUS should not be administered to patients with
- 493 creatinine clearance <50 mL/min.

ADVERSE REACTIONS

- 495 PEGASYS alone or in combination with COPEGUS causes a broad variety of serious
- adverse reactions (see **BOXED WARNING** and **WARNINGS**). In all studies, one or
- 497 more serious adverse reactions occurred in 10% of patients receiving PEGASYS alone or
- in combination with COPEGUS.
- The most common life-threatening or fatal events induced or aggravated by PEGASYS
- and COPEGUS were depression, suicide, relapse of drug abuse/overdose, and bacterial
- infections; each occurred at a frequency of <1%.
- Nearly all patients in clinical trials experienced one or more adverse events. The most
- 503 commonly reported adverse reactions were psychiatric reactions, including depression,
- 504 irritability, anxiety, and flu-like symptoms such as fatigue, pyrexia, myalgia, headache
- and rigors.

494

- Overall 11% of patients receiving 48 weeks of therapy with PEGASYS either alone (7%)
- or in combination with COPEGUS (10%) discontinued therapy. The most common
- reasons for discontinuation of therapy were psychiatric, flu-like syndrome (eg, lethargy,
- fatigue, headache), dermatologic and gastrointestinal disorders.
- 510 The most common reason for dose modification in patients receiving combination
- therapy was for laboratory abnormalities; neutropenia (20%) and thrombocytopenia (4%)
- for PEGASYS and anemia (22%) for COPEGUS.
- 513 PEGASYS dose was reduced in 12% of patients receiving 1000 mg to 1200 mg
- 514 COPEGUS for 48 weeks and in 7% of patients receiving 800 mg COPEGUS for 24
- weeks. COPEGUS dose was reduced in 21% of patients receiving 1000 mg to 1200 mg
- 516 COPEGUS for 48 weeks and 12% in patients receiving 800 mg COPEGUS for 24 weeks.
- 517 Because clinical trials are conducted under widely varying and controlled
- 518 conditions, adverse reaction rates observed in clinical trials of a drug cannot be
- 519 directly compared to rates in the clinical trials of another drug. Also, the adverse
- 520 event rates listed here may not predict the rates observed in a broader patient
- 521 population in clinical practice.

Table 4 Adverse Reactions Occurring in 35% of Patients in Hepatitis C Clinical Trials (Pooled Studies 1, 2, 3, and Study 4)

	ιω ς τη	1 -		T
Body System	PEGASYS 180 mg 48 wk†	ROFERON-A*†	PEGASYS 180 mg + 1000 mg or 1200 mg COPEGUS 48 wk **	Intron A + 1000 mg or 1200 mg REBETOL ^a 48 wk**
	N=559	N=554	N=451	N=443
	%	%	%	%
Application Site Disorders				
Injection site reaction	22	18	23	16
Endocrine Disorders				
Hypothyroidism	3	2	4	5
Flu-like symptoms and signs				
Fatigue/Asthenia	56	57	65	68
Pyrexia	37	41	41	55
Rigors	35	44	25	37
Pain	11	12	10	9
Gastrointestinal				
Nausea/vomiting	24	33	25	29
Diamhea	16	16	11	10
Abdominal pain	15	15	8	9
Dry mouth	6	3	4	7
Dyspepsia	<1	1	6	5
Hematologic‡				
Lymphopenia	3	5	14	12
Anemia	2	1	11	11
Neutropenia	21	8	27	8

Body System	PEGASYS 180 mg 48 wk†	ROFERON-A*†	PEGASYS 180 mg + 1000 mg or 1200 mg COPEGUS 48 wk **	Intron A + 1000 mg or 1200 mg REBETOL ^â 48 wk**
	N=559	N=554	N=451	N=443
	%	%	%	%
Thrombocytopenia	5	2	5	<1
Metabolic and Nutritional				
Anorexia	17	17	24	26
Weight decrease	4	3	10	10
Musculoskeletal, Connective Tissue and Bone				
Myalgia	37	38	40	49
Arthralgia	28	29	22	23
Back pain	9	10	5	5
Neurological				
Headache	54	58	43	49
Dizziness (excluding vertigo)	16	12	14	14
Memory impairment	5	4	6	5
Psychiatric				
Irritability/Anxiety/Nervo usness	19	22	33	38
Insomnia	19	23	30	37
Depression	18	19	20	28
Concentration impairment	8	10	10	13
Mood alteration	3	2	5	6

Body System	PEGASYS 180 mg 48 wk†	ROFERON-A*†	PEGASYS 180 mg + 1000 mg or 1200 mg COPEGUS 48 wk **	Intron A + 1000 mg or 1200 mg REBETOL ^a 48 wk**
	N=559	N=554	N=451	N=443
	%	%	%	%
Resistance Mechanism Disorders				
Overall	10	6	12	10
Respiratory, Thoracic and Mediastinal				
Dyspnea	4	2	13	14
Cough	4	3	10	7
Dyspnea exertional	<1	<1	4	7
Skin and Subcutaneous Tissue				
Alopecia	23	30	28	33
Pruritus	12	8	19	18
Dermatitis	8	3	16	13
Dry Skin	4	3	10	13
Rash	5	4	8	5
Sweating Increased	6	7	6	5
Eczema	1	1	5	4
Visual Disorders				
Vision Blurred	4	2	5	2

⁵²⁵ † Pooled studies 1, 2, and 3 526

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529

Patients treated for 24 weeks with PEGASYS and 800 mg COPEGUS were observed to have lower incidence of serious adverse events (3% vs 10%), Hgb <10g/dL (3% vs 15%), dose modification of PEGASYS (30% vs 36%) and COPEGUS (19% vs 38%) and of

^{*} Either 3 MIU or 6/3 MIU of ROFERON-A

⁵²⁷ **Study 4

⁵²⁸ ‡ Severe hematologic abnormalities

- withdrawal from treatment (5% vs 15%) compared to patients treated for 48 weeks with
- PEGASYS and 1000 mg or 1200 mg COPEGUS. On the other hand the overall incidence
- of adverse events appeared to be similar in the two treatment groups.
- 536 The most common serious adverse event (3%) was bacterial infection (eg, sepsis,
- osteomyelitis, endocarditis, pyelonephritis, pneumonia). Other SAEs occurred at a
- frequency of <1% and included: suicide, suicidal ideation, psychosis, aggression, anxiety,
- 539 drug abuse and drug overdose, angina, hepatic dysfunction, fatty liver, cholangitis,
- 540 arrhythmia, diabetes mellitus, autoimmune phenomena (eg, hyperthyroidism,
- 541 hypothyroidism, sarcoidosis, systemic lupus erythematosis, rheumatoid arthritis)
- 542 peripheral neuropathy, aplastic anemia, peptic ulcer, gastrointestinal bleeding,
- 543 pancreatitis, colitis, corneal ulcer, pulmonary embolism, coma, myositis, and cerebral
- hemorrhage.

545

Laboratory Test Values

- 546 Hemoglobin
- 547 The hemoglobin concentration decreased below 12g/dL in 17% (median Hgb
- drop = 2.2 g/dL) of monotherapy and 52% (median Hgb drop = 3.7 g/dL) of combination
- 549 therapy patients. Severe anemia (Hgb <10 g/dL) was encountered in 13% of patients
- receiving combination therapy and 2% of monotherapy recipients. Dose modification for
- anemia was required in 22% of ribavirin recipients treated for 48 weeks. Hemoglobin
- decreases in PEGASYS monotherapy were generally mild and did not require dose
- modification (see **DOSAGE AND ADMINISTRATION: Dose Modifications**).
- 554 Neutrophils
- Decreases in neutrophil count below normal were observed in 95% of patients treated
- with PEGASYS either alone or in combination with COPEGUS. Severe potentially life-
- threatening neutropenia (ANC $<0.5\times10^9/L$) occurred in approximately 5% of patients
- receiving PEGASYS either alone or in combination with COPEGUS. Seventeen percent
- of patients receiving PEGASYS monotherapy and 20% to 24% of patients receiving
- 560 PEGASYS/COPEGUS combination therapy required modification of interferon dosage
- for neutropenia. Two percent of patients required permanent reductions of PEGASYS
- dosage and <1% required permanent discontinuation. Median neutrophil counts return to
- pre-treatment levels 4 weeks after cessation of therapy (see **DOSAGE AND**
- **ADMINISTRATION: Dose Modifications**).
- 565 Lymphocytes
- Decreases in lymphocyte count are induced by interferon alpha therapy. Lymphopenia
- was observed during both monotherapy (86%) and combination therapy with PEGASYS
- and COPEGUS (94%). Severe lymphopenia (<0.5x10⁹/L) occurred in approximately 5%
- of monotherapy patients and 14% of combination PEGASYS AND COPEGUS therapy
- recipients. Dose adjustments were not required by protocol. Median lymphocyte counts
- return to pre-treatment levels after 4 to 12 weeks of the cessation of therapy. The clinical
- significance of the lymphopenia is not known.

- 573 Platelets
- Platelet counts decreased in 52% of patients treated with PEGASYS alone (median drop
- 575 45% from baseline), 33% of patients receiving combination with COPEGUS (median
- drop 30% from baseline). Median platelet counts return to pretreatment levels 4 weeks
- after the cessation of therapy.
- 578 Triglycerides
- 579 Triglyceride levels are elevated in patients receiving alfa interferon therapy and were
- 580 elevated in the majority of patients participating in clinical studies receiving either
- 581 PEGASYS alone or in combination with COPEGUS. Random levels higher ≥400 mg/dL
- were observed in about 20% of patients.
- 583 ALT Elevations
- Less than 1% of patients experienced marked elevations (5- to 10-fold above baseline) in
- 585 ALT levels during treatment. These transaminase elevations were on occasion associated
- with hyperbilirubinemia and were managed by dose reduction or discontinuation of study
- 587 treatment. Liver function test abnormalities were generally transient. One case was
- 588 attributed to autoimmune hepatitis, which persisted beyond study medication
- discontinuation (see **DOSAGE AND ADMINISTRATION: Dose Modifications**).
- 590 Thyroid function
- 591 PEGASYS alone or in combination with COPEGUS was associated with the
- development of abnormalities in thyroid laboratory values, some with associated clinical
- 593 manifestations. Hypothyroidism or hyperthyroidism requiring treatment, dose
- modification or discontinuation occurred in 4% and 1% of PEGASYS treated patients
- 595 and 4% and 2% of PEGASYS and COPEGUS treated patients, respectively.
- 596 Approximately half of the patients, who developed thyroid abnormalities during
- 597 PEGASYS treatment, still had abnormalities during the follow-up period (see
- 598 **PRECAUTIONS: Laboratory Tests**).
- 599 Immunogenicity
- Nine percent (71/834) of patients treated with PEGASYS with or without COPEGUS
- developed binding antibodies to interferon alfa-2a, as assessed by an ELISA assay. Three
- 602 percent of patients (25/835) receiving PEGASYS with or without COPEGUS, developed
- low-titer neutralizing antibodies (using an assay of a sensitivity of 100 INU/mL).
- The clinical and pathological significance of the appearance of serum neutralizing
- antibodies is unknown. No apparent correlation of antibody development to clinical
- 606 response or adverse events was observed. The percentage of patients whose test results
- 607 were considered positive for antibodies is highly dependent on the sensitivity and
- specificity of the assays.
- 609 Additionally the observed incidence of antibody positivity in these assays may be
- 610 influenced by several factors including sample timing and handling, concomitant
- 611 medications, and underlying disease. For these reasons, comparison of the incidence of

- antibodies to PEGASYS with the incidence of antibodies to these products may be
- 613 misleading.

614 **OVERDOSAGE**

- There is limited experience with overdosage. The maximum dose received by any patient
- was 7 times the intended dose of PEGASYS (180 µg/day for 7 days). There were no
- serious reactions attributed to overdosages. Weekly doses of up to 630 µg have been
- administered to patients with cancer. Dose-limiting toxicities were fatigue, elevated liver
- 619 enzymes, neutropenia, and thrombocytopenia. There is no specific antidote for
- 620 PEGASYS. Hemodialysis and peritoneal dialysis are not effective.

DOSAGE AND ADMINISTRATION

- There are no safety and efficacy data on treatment for longer than 48 weeks.
- 623 Consideration should be given to discontinuing therapy after 12-24 weeks of therapy if
- 624 the patient has failed to demonstrate an early virologic response (see CLINICAL
- 625 **STUDIES**).

621

641

626 **PEGASYS**

- The recommended dose of PEGASYS monotherapy is 180 µg (1.0 mL) once weekly for
- 48 weeks by subcutaneous administration in the abdomen or thigh.

629 PEGASYS and COPEGUS COMBINATION

- The recommended dose of PEGASYS when used in combination with ribavirin is 180 µg
- 631 (1.0 mL) once weekly. The recommended dose of COPEGUS and duration for
- PEGASYS/COPEGUS therapy is based on viral genotype (see Table 5).
- The daily dose of COPEGUS is 800 mg to 1200 mg administered orally in two divided
- doses. The dose should be individualized to the patient depending on baseline disease
- characteristics (eg, genotype), response to therapy, and tolerability of the regimen.
- 636 Since COPEGUS absorption increases when administered with a meal, patients are
- advised to take COPEGUS with food.

Table 5 PEGASYS and COPEGUS Dosing Recommendations

Genotype	PEGASYS Dose	COPEGUS Dose	Duration
Genotype 1, 4	180 µg	<75 kg = 1000 mg	48 weeks
Genotype 1, 4	160 μg	≥75 kg = 1200 mg	48 weeks
Genotype 2, 3	180 μg	800 mg	24 weeks

Genotypes 2 and 3 showed no increased response to treatment beyond 24 weeks (see Table 3).

Data on genotypes 5 and 6 are insufficient for dosing recommendations.

A patient should self-inject PEGASYS only if the physician determines that it is appropriate and the patient agrees to medical follow-up as necessary and training in

- proper injection technique has been provided to him/her (see illustrated PEGASYS
- 645 **MEDICATION GUIDE** for directions on injection site preparation and injection
- 646 instructions).
- PEGASYS should be inspected visually for particulate matter and discoloration before
- administration, and not used if particulate matter is visible or product is discolored. Vials
- with particulate matter or discoloration should be returned to the pharmacist.

650 **Dose Modifications**

- 651 If severe adverse reactions or laboratory abnormalities develop during combination
- 652 COPEGUS/PEGASYS therapy, the dose should be modified or discontinued, if
- appropriate, until the adverse reactions abate. If intolerance persists after dose
- adjustment, COPEGUS/PEGASYS therapy should be discontinued.

655 **PEGASYS**

- 656 General
- When dose modification is required for moderate to severe adverse reactions (clinical
- and/or laboratory), initial dose reduction to 135 µg (0.75 mL) is generally adequate.
- However, in some cases, dose reduction to 90 µg (0.5 mL) may be needed. Following
- improvement of the adverse reaction, re-escalation of the dose may be considered (see
- WARNINGS, PRECAUTIONS, and ADVERSE REACTIONS).

662 Hematological

Table 6 PEGASYS Hematological Dose Modification Guidelines

Laboratory Values	PEGASYS Dose Reduction	Discontinue PEGASYS if:
ANC <750/mm ³	135 μg	ANC <500/mm³, treatment should be suspended until ANC values return to more than 1000/mm³. Reinstitute at 90 µg and monitor ANC
Platelet <50,000/mm ³	90 μg	Platelet count <25,000/mm ³

664 Psychiatric: Depression

665

666

Table 7 Guidelines for Modification or Discontinuation of PEGASYS and for Scheduling Visits for Patients with Depression

Depression Severity		nagement veeks)	Depression		
	Dose modification	Visit schedule	Remains stable	Improves	Worsens

Mild	No change	Evaluate once weekly by visit and/or phone	Continue weekly visit schedule	Resume normal visit schedule	(See moderate or severe depression)
Moderate	Decrease PEGASYS dose to 135 µg (in some cases dose reduction to 90 µg may be needed)	Evaluate once weekly (office visit at least every other week)	Consider psychiatric consultation. Continue reduced dosing	If symptoms improve and are stable for 4 weeks, may resume normal visit schedule. Continue reduced dosing or return to normal dose	(See severe depression)
Severe	Discontinue PEGASYS permanently	Obtain immediate psychiatric consultation	Psychiatric ther	apy necessary	

667 Renal Function

- In patients with end-stage renal disease requiring hemodialysis, dose reduction to 135 µg
- 669 PEGASYS is recommended. Signs and symptoms of interferon toxicity should be closely
- 670 monitored.

671 Liver Function

- 672 In patients with progressive ALT increases above baseline values, the dose of PEGASYS
- should be reduced to 135 µg. If ALT increases are progressive despite dose reduction or
- 674 accompanied by increased bilirubin or evidence of hepatic decompensation, therapy
- should be immediately discontinued.

676 **COPEGUS**

677 Table 8 COPEGUS Dosage Modification Guidelines

Laboratory Values	Reduce Only COPEGUS Dose to 600 mg/day* if:	Discontinue COPEGUS if:
Hemoglobin in patients with no cardiac disease	<10 g/dL	<8.5 g/dL
Hemoglobin in patients with history of stable cardiac disease	≥2 g/dL decrease in hemoglobin during any 4 week period treatment	<12 g/dL despite 4 weeks at reduced dose

- * One 200 mg tablet in the morning and two 200 mg tablets in the evening.
- Once COPEGUS has been withheld due to a laboratory abnormality or clinical
- 680 manifestation, an attempt may be made to restart COPEGUS at 600 mg daily and further
- increase the dose to 800 mg daily depending upon the physician's judgment. However, it
- is not recommended that COPEGUS be increased to the original dose (1000 mg or
- 683 1200 mg).
- 684 Renal Impairment
- 685 COPEGUS should not be used in patients with creatinine clearance <50 mL/min (see
- 686 WARNINGS and COPEGUS Package Insert).

687 **HOW SUPPLIED**

688 **Single Dose Vial**

- Each PEGASYS (peginterferon alfa-2a) 180 µg single use, clear glass vial provides
- 690 1.0 mL containing 180 μg peginterferon alfa-2a for SC injection. Each package contains
- 691 1 vial (NDC 0004-0350-09).

692 Monthly Convenience Pack

- 693 Four vials of PEGASYS (peginterferon alfa-2a), 180 μg single use, in a box with 4
- 694 syringes and 8 alcohol swabs (NDC 0004-0350-39). Each syringe is a 1 mL (1 cc)
- of volume syringe supplied with a 27 gauge, ½ inch needle with needle-stick protection
- 696 device.

697 **Storage**

- 698 Store in the refrigerator at 36° to 46°F (2° to 8°C). Do not freeze or shake. Protect from
- 699 light. Vials are for single use only. Discard any unused portion.
- 700 REBETRONTM is a trademark of Schering Corporation.
- 701 Rx only

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