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# PEGASYS®

## 3

Rx only

## (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 Therapy should be withdrawn in patients with persistently severe or worsening signs or symptoms of these conditions.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

11 and ADVERSE REACTIONS).

Use with Ribavirin. Ribavirin, including COPEGUS®, may cause birth defects 12 13

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.

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The anemia associated with ribavirin therapy may result in a worsening of cardiac 16

disease. Ribavirin is genotoxic and mutagenic and should be considered a potential

carcinogen (see COPEGUS Package Insert for additional information and other

WARNINGS). 18

### **DESCRIPTION**

- 20 PEGASYS, peginterferon alfa-2a, is a covalent conjugate of recombinant alfa-2a
- 21 interferon (approximate molecular weight [MW] 20,000 daltons) with a single branched
- 22 bis-monomethoxy polyethylene glycol (PEG) chain (approximate MW 40,000 daltons).
- 23 The PEG moiety is linked at a single site to the interferon alfa moiety via a stable amide 24
- bond to lysine. Peginterferon alfa-2a has an approximate molecular weight of 60,000 25 daltons. Interferon alfa-2a is produced using recombinant DNA technology in which a
- 26 cloned human leukocyte interferon gene is inserted into and expressed in Escherichia
- 27 coli.
- 28 PEGASYS is supplied as an injectable solution in vials and prefilled syringes.
- 29 180 µg/1.0 mL Vial: A vial contains approximately 1.2 mL of solution to deliver 1.0 mL
- 30 of drug product. Subcutaneous (sc) administration of 1.0 mL delivers 180 µg of drug
- 31 product (expressed as the amount of interferon alfa-2a), 8.0 mg sodium chloride, 0.05 mg
- 32 polysorbate 80, 10.0 mg benzyl alcohol, 2.62 mg sodium acetate trihydrate, and 0.05 mg
- 33 acetic acid. The solution is colorless to light yellow and the pH is  $6.0 \pm 0.5$ .
- 34 180 μg/0.5 mL Prefilled Syringe: Each syringe contains 0.6 mL of solution to deliver
- 35 0.5 mL of drug product. Subcutaneous (sc) administration of 0.5 mL delivers 180 µg of
- 36 drug product (expressed as the amount of interferon alfa-2a), 4.0 mg sodium chloride.
- 37 0.025 mg polysorbate 80, 5.0 mg benzyl alcohol, 1.3085 mg sodium acetate trihydrate,
- 38 and 0.0231 mg acetic acid. The solution is colorless to light yellow and the pH is
- 39  $6.0 \pm 0.5$ .

### 40 CLINICAL PHARMACOLOGY

### 41 Pharmacodynamics

- 42 Interferons bind to specific receptors on the cell surface initiating intracellular signaling
- via a complex cascade of protein-protein interactions leading to rapid activation of gene
- 44 transcription. Interferon-stimulated genes modulate many biological effects including the
- 45 inhibition of viral replication in infected cells, inhibition of cell proliferation and
- immunomodulation. The clinical relevance of these in vitro activities is not known.
- 47 PEGASYS stimulates the production of effector proteins such as serum neopterin and 2',
- 48 5'-oligoadenylate synthetase.

### 49 Pharmacokinetics

- 50 Maximal serum concentrations (C<sub>max</sub>) and AUC increased in a nonlinear dose related
- 51 manner following administration of 90 to 270 µg of PEGASYS. Maximal serum
- 52 concentrations (C<sub>max</sub>) occur between 72 to 96 hours post-dose.
- Week 48 mean trough concentrations (16 ng/mL; range 4 to 28) at 168 hours post-dose
- are approximately 2-fold higher than week 1 mean trough concentrations (9 ng/mL; range
- 55 0 to 15). Steady-state serum levels are reached within 5 to 8 weeks of once weekly
- dosing. The peak to trough ratio at week 48 is approximately 2. The mean systemic
- 57 clearance in healthy subjects given PEGASYS was 94 mL/h, which is approximately
- 58 100-fold lower than that for interferon alfa-2a (ROFERON®-A). The mean terminal half-
- 59 life after sc dosing in patients with chronic hepatitis C was 160 hours (range 84 to 353
- 60 hours) compared to 5 hours (range 3.7 to 8.5 hours) for ROFERON-A.

## 61 Special Populations

- 62 Gender and Age
- 63 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
- 9 years taking 180 μg PEGASYS, but peak concentrations were similar (9 vs. 10 ng/mL) in
- those older and younger than 62 years.

### 67 Pediatric Patients

- In a population pharmacokinetics study, 14 children 2 to 8 years of age with CHC
- 69 received PEGASYS based on their body surface area (BSA of the child x
- 70 180 µg/1.73m<sup>2</sup>). The clearance of PEGASYS in children was nearly 4-fold lower
- 71 compared to the clearance reported in adults.
- 72 Steady-state trough levels in children with the BSA-adjusted dosing were similar to
- 73 trough levels observed in adults with 180 µg fixed dosing. Time to reach the steady state
- in children is approximately 12 weeks, whereas in adults, steady state is reached within 5 to 8 weeks. In these children receiving the BSA adjusted dose, the mean exposure (AUC)
- to 8 weeks. In these children receiving the BSA adjusted dose, the mean exposure (AUC) during the dosing interval is predicted to be 25% to 70% higher than that observed in
- adults receiving 180 µg fixed dosing. The safety and effectiveness of PEGASYS in
- 78 patients below the age of 18 years have not been established (see PRECAUTIONS:
- 79 Pediatric Use).

- 80 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).
- 83 The pharmacokinetics of ribavirin following administration of COPEGUS have not been
- 84 studied in patients with renal impairment and there are limited data from clinical trials on
- 85 administration of COPEGUS in patients with creatinine clearance <50 mL/min.
- 86 Therefore, patients with creatinine clearance <50 mL/min should not be treated with
- 87 COPEGUS (see WARNINGS and DOSAGE AND ADMINISTRATION).
- 88 Effect of Food on Absorption of Ribavirin
- 89 Bioavailability of a single oral dose of ribavirin was increased by co-administration with
- a high-fat meal. The absorption was slowed (T<sub>max</sub> was doubled) and the AUC<sub>0-192h</sub> and
- 91 C<sub>max</sub> increased by 42% and 66%, respectively, when COPEGUS was taken with a high-
- 92 fat meal compared with fasting conditions (see **DOSAGE AND ADMINISTRATION**).

## 93 Drug Interactions

- 94 Nucleoside Analogues
- 95 In vitro data indicate ribavirin reduces phosphorylation of lamivudine, stavudine, and
- 96 zidovudine. However, no pharmacokinetic (e.g., plasma concentrations or intracellular
- 97 triphosphorylated active metabolite concentrations) or pharmacodynamic (e.g., loss of
- 98 HIV/HCV virologic suppression) interaction was observed when ribavirin and
- 99 lamivudine (n=18), stavudine (n=10), or zidovudine (n=6) were co-administered as part
- of a multi-drug regimen to HCV/HIV coinfected patients (see PRECAUTIONS: Drug
- 101 Interactions).
- 102 In vitro, didanosine or its active metabolite (dideoxyadenosine 5'-triphosphate) is
- increased when didanosine is co-administered with ribavirin (see PRECAUTIONS:
- 104 **Drug Interactions**).
- 105 Drugs Metabolized by Cytochrome P450
- There was no effect on the pharmacokinetics of representative drugs metabolized by CYP
- 107 2C9, CYP 2C19, CYP 2D6 or CYP 3A4.
- 108 Treatment with PEGASYS once weekly for 4 weeks in healthy subjects was associated
- with an inhibition of P450 1A2 and a 25% increase in theophylline AUC (see
- 110 PRECAUTIONS: Drug Interactions).
- 111 Methadone
- 112 The pharmacokinetics of concomitant administration of methadone and PEGASYS were
- evaluated in 24 PEGASYS naive chronic hepatitis C (CHC) patients (15 male, 9 female)
- who received 180 μg PEGASYS subcutaneously weekly. All patients were on stable
- methadone maintenance therapy (median dose 95 mg, range 30 mg to 150 mg) prior to
- receiving PEGASYS. Mean methodone PK parameters were 10% to 15% higher after 4
- weeks of PEGASYS treatment as compared to baseline (see PRECAUTIONS: Drug
- 118 Interactions). Methadone did not significantly alter the PK of PEGASYS as compared to
- a PK study of 6 chronic hepatitis C patients not receiving methadone.

### 120 CLINICAL STUDIES

### 121 Chronic Hepatitis C Studies 1, 2, and 3: PEGASYS Monotherapy

- 122 The safety and effectiveness of PEGASYS for the treatment of hepatitis C virus infection
- 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.
- 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
- 128 20% of subjects had cirrhosis or bridging fibrosis. Study 3 enrolled patients with a
- histological diagnosis of cirrhosis (78%) or bridging fibrosis (22%).
- 130 In Study 1 (n=630), patients received either ROFERON-A (interferon alfa-2a) 3 MIU
- three times/week (tiw), PEGASYS 135 µg once each week (qw) or PEGASYS 180 µg
- qw. In Study 2 (n=526), patients received either ROFERON-A 6 MIU tiw for 12 weeks
- 133 followed by 3 MIU tiw for 36 weeks or PEGASYS 180 µg qw. In Study 3 (n=269),
- patients received ROFERON-A 3 MIU tiw, PEGASYS 90 µg qw or PEGASYS 180 µg
- once each week.
- 136 In all three studies, treatment with PEGASYS 180 µg resulted in significantly more
- patients who experienced a sustained response (defined as undetectable HCV RNA [<50]
- 138 IU/mL] using the COBAS AMPLICOR® HCV Test, version 2.0 and normalization of
- 139 ALT on or after study week 68) compared to treatment with ROFERON-A. In Study 1,
- response to PEGASYS 135 µg was not different from response to 180 µg. In Study 3,
- 141 response to PEGASYS 90 µg was intermediate between PEGASYS 180 µg and
- 142 ROFERON-A.

## 143 Table 1 Sustained Response to Monotherapy Treatment

	Study 1				Study 2		Study 3		
	ROFERON-A 3 MIU (N=207)	PEGASYS 180 μg (N=208)	DIFF* (95% CI)	ROFERON-A 6/3 MIU (N=261)	PEGASYS 180 μg (N=265)	DIFF* (95% CI)	ROFERON-A 3 MIU (N=86)	PEGASYS 180 μg (N=87)	DIFF* (95% CI)
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)

\*Percent difference between PEGASYS and ROFERON-A treatment.

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.

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149 Of the patients who did not demonstrate either undetectable HCV RNA or at least a

2log<sub>10</sub> drop in HCV RNA titer from baseline by 12 weeks of PEGASYS 180 μg therapy,

- 151 2% (3/156) achieved a sustained virologic response (see DOSAGE AND
- 152 **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.

# Chronic Hepatitis C Studies 4 and 5: PEGASYS/COPEGUS Combination

### 157 Therapy

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- 158 The safety and effectiveness of PEGASYS in combination with COPEGUS for the
- 159 treatment of hepatitis C virus infection were assessed in two randomized controlled
- 160 clinical trials. All patients were adults, had compensated liver disease, detectable hepatitis
- 161 C virus, liver biopsy diagnosis of chronic hepatitis, and were previously untreated with
- interferon. Approximately 20% of patients in both studies had compensated cirrhosis
- 163 (Child-Pugh class A). Patients coinfected with HIV were excluded from these studies.
- 164 In Study 4, patients were randomized to receive either PEGASYS 180 μg sc once weekly
- 165 (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<sup>®</sup> (interferon alfa-2b
- 167 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
- 169 treatment assignment was blinded. Sustained virological response was defined as
- undetectable (<50 IU/mL) HCV RNA on or after study week 68. PEGASYS in
- combination with COPEGUS resulted in a higher SVR compared to PEGASYS alone or
- interferon alfa-2b and ribavirin (**Table 2**). In all treatment arms, patients with viral
- genotype 1, regardless of viral load, had a lower response rate.

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

	Interferon alfa-2b +	PEGASYS +	PEGASYS +
·	Ribavirin 1000 mg or 1200 mg	Placebo	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%)

\*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 (see **Table 3**), all patients received PEGASYS 180  $\mu$ 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 10<sup>6</sup> HCV RNA copies/mL serum) were preferentially assigned to treatment for 48 weeks.

- 185 **HCV** Genotypes
- 186 HCV 1 and 4 - Irrespective of baseline viral titer, treatment for 48 weeks with
- 187 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) 188
- 189 compared to shorter treatment (24 weeks) and/or 800 mg COPEGUS.
- 190 HCV 2 and 3 - Irrespective of baseline viral titer, treatment for 24 weeks with
- 191 PEGASYS and 800 mg of COPEGUS resulted in a similar SVR compared to longer
- 192 treatment (48 weeks) and/or 1000 mg or 1200 mg of COPEGUS (see **Table 3**).
- 193 The numbers of patients with genotype 5 and 6 were too few to allow for meaningful
- 194 assessment.

195 Table 3 Sustained Virologic Response as a Function of Genotype 196 (Study 5)

24 Weel	ks Treatment	48 Weeks Treatment		
PEGASYS + COPEGUS	PEGASYS + COPEGUS	PEGASYS + COPEGUS	PEGASYS + COPEGUS	
800 mg (N=207)	1000 mg or 1200 mg*	800 mg (N=361)	1000 mg or 1200 mg*	
29/101 (29%)	48/118 (41%)	99/250 (40%)	138/271 (51%)	
79/96 (82%)	116/144 (81%)	75/99 (76%)	117/153 (76%)	
0/5 (0%)	7/12 (58%)	5/8 (63%)	9/11 (82%)	
	PEGASYS + COPEGUS 800 mg (N=207) 29/101 (29%) 79/96 (82%)	COPEGUS  800 mg 1000 mg or 1200 mg* (N=207) (N=280)  29/101 (29%) 48/118 (41%)  79/96 (82%) 116/144 (81%)	PEGASYS +         PEGASYS +         PEGASYS +           COPEGUS         COPEGUS         COPEGUS           800 mg         1000 mg or 1200 mg*         800 mg           (N=207)         (N=280)         (N=361)           29/101 (29%)         48/118 (41%)         99/250 (40%)           79/96 (82%)         116/144 (81%)         75/99 (76%)	

- 197 \*1000 mg for body weight <75 kg; 1200 mg for body weight ≥75 kg.
- 198 Other Treatment Response Predictors
- 199 Treatment response rates are lower in patients with poor prognostic factors receiving
- 200 pegylated interferon alpha therapy. In studies 4 and 5, treatment response rates were
- 201 lower in patients older than 40 years (50% vs. 66%), in patients with cirrhosis (47% vs.
- 202
- 59%), in patients weighing over 85 kg (49% vs. 60%), and in patients with genotype 1
- 203 with high vs. low viral load (43% vs. 56%). African-American patients had lower
- 204 response rates compared to Caucasians.
- 205 Paired liver biopsies were performed on approximately 20% of patients in studies 4 and
- 206 5. Modest reductions in inflammation compared to baseline were seen in all treatment
- 207 groups.
- 208 In studies 4 and 5, lack of early virologic response by 12 weeks (defined as HCV RNA
- 209 undetectable or >2log<sub>10</sub> lower than baseline) was grounds for discontinuation of
- 210 treatment. Of patients who lacked an early viral response by 12 weeks and completed a
- 211 recommended course of therapy despite a protocol-defined option to discontinue therapy,
- 212 5/39 (13%) achieved an SVR. Of patients who lacked an early viral response by 24
- 213 weeks, 19 completed a full course of therapy and none achieved an SVR.

### 214 Chronic Hepatitis C and Coinfection with HIV (CHC/HIV) Study 6: PEGASYS 215 Monotherapy and PEGASYS/COPEGUS Combination Therapy

216 In Study 6, patients with CHC/HIV were randomized to receive either PEGASYS 180 µg 217 sc once weekly (qw) plus an oral placebo, PEGASYS 180 µg qw plus COPEGUS 800 mg po daily or ROFERON-A (interferon alfa-2a), 3 MIU sc tiw plus COPEGUS 800 218 219 mg po daily. All patients received 48 weeks of therapy and sustained virologic response 220 (SVR) was assessed at 24 weeks of treatment-free follow-up. COPEGUS or placebo 221 treatment assignment was blinded in the PEGASYS treatment arms. All patients were 222 adults, had compensated liver disease, detectable hepatitis C virus, liver biopsy diagnosis 223 of chronic hepatitis C, and were previously untreated with interferon. Patients also had 224 CD4+ cell count ≥200 cells/μL or CD4+ cell count ≥100 cells/μL but <200 cells/μL and 225 HIV-1 RNA <5000 copies/mL, and stable status of HIV. Approximately 15% of patients 226 in the study had cirrhosis. Results are shown in Table 4.

Table 4 Sustained Virologic Response in Patients with Chronic **Hepatitis C Coinfected with HIV (Study 6)** 

	ROFERON-A + COPEGUS 800 mg	PEGASYS + Placebo	PEGASYS + COPEGUS 800 mg
	(N=289)	(N=289)	(N=290)
All patients	33 (11%)*	58 (20%)*	116 (40%)
Genotype 1	12/171 (7%)	24/175 (14%)	51/176 (29%)
Genotypes 2, 3	18/89 (20%)	32/90 (36%)	59/95 (62%)
		1	

\*PEGASYS + COPEGUS vs. PEGASYS; PEGASYS + COPEGUS vs. ROFERON-A + COPEGUS pvalue < 0.0001 (Cochran-Mantel-Haenszel).

232 Treatment response rates are lower in CHC/HIV patients with poor prognostic factors (including HCV genotype 1, HCV RNA >800,000 IU/mL, and cirrhosis) receiving 233 234 pegylated interferon alpha therapy. Geographic region is not a prognostic factor for 235 response. However, poor prognostic factors occur more frequently in the US population than in the non-US population.

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- 237 Of the patients who did not demonstrate either undetectable HCV RNA or at least a 2log<sub>10</sub> reduction from baseline in HCV RNA titer by 12 weeks of PEGASYS and 238
- 239 COPEGUS combination therapy, 2% (2/85) achieved an SVR.
- In CHC patients with HIV coinfection who received 48 weeks of PEGASYS alone or in 240 241 combination with COPEGUS treatment, mean and median HIV RNA titers did not 242 increase above baseline during treatment or 24 weeks post-treatment.

## Chronic Hepatitis B Studies 7 and 8: PEGASYS Monotherapy

244 The safety and effectiveness of PEGASYS for the treatment of chronic hepatitis B were 245 assessed in controlled clinical trials in HBeAg positive (Study 7) and HBeAg negative 246 (Study 8) patients with chronic hepatitis B.

- Patients were randomized to PEGASYS 180 μg sc once weekly (qw), PEGASYS 180 μg
- sc qw combined with lamivudine 100 mg once daily po or lamivudine 100 mg once daily
- po. All patients received 48 weeks of their assigned therapy followed by 24 weeks of
- 250 treatment-free follow-up. Assignment to receipt of PEGASYS or no PEGASYS was not
- 251 masked.
- 252 All patients were adults with compensated liver disease, had chronic hepatitis B virus
- 253 (HBV) infection, and evidence of HBV replication (serum HBV >500,000 copies/mL for
- 254 Study 7 and >100,000 copies/mL for Study 8) as measured by PCR (COBAS
- 255 AMPLICOR® HBV Assay). All patients had serum alanine aminotransferase (ALT)
- between 1 and 10 times the upper limit of normal (ULN) and liver biopsy findings
- compatible with the diagnosis of chronic hepatitis.
- 258 The results observed in the PEGASYS and lamivudine monotherapy groups are shown in
- 259 **Table 5**.

Table 5 Percentage of Patients with Serological, Virological,
 Biochemical, and Histological Response

	]	Study 7 HBeAg posi		Study 8 HBeAg negative			
	Lami	vudine	PEGASYS	Lami	vudine	PEGASYS	
	24111	Vacino	Legis	Luiii	v dame	LONGIS	
	N =	= 272	N = 271	N = 181		N = 177	
	ĘOT <sup>1</sup>	EOF <sup>2</sup>	EOF <sup>2</sup>	EOT <sup>I</sup>	EOF <sup>2</sup>	EOF <sup>2</sup>	
HBeAg Seroconversion (%)	20	19*	32*	NA	NA	NA	
HBV DNA Response (%) <sup>3</sup>	62	22***	32***	85	29**	43**	
ALT Normalization (%)	62	28	41	73	44**	59**	
HBsAg Seroconversion (%)	0	0	3	1	. 0	3	
	N =	= 184	N = 207	N = 125		N = 143	
Histological Improvement (%) <sup>4</sup>	ND .	40	41	ND	41	48	
Changes in Ishak fibrosis score compared to baseline (%):						, , , , , ,	
- Improved <sup>5</sup> - Unchanged - Worsened <sup>5</sup>	ND	32 20 16	25 25 26	ND	31 23 15	32 30 19	

<sup>262</sup> End of Treatment (week 48)

<sup>263 &</sup>lt;sup>2</sup> End of follow-up – 24 weeks post-treatment (week 72)

- <sup>3</sup><100,000 copies/mL for HBeAg positive and <20,000 copies/mL for HBeAg negative patients
- 265 <sup>4</sup>≥2 point decrease in Ishak necro-inflammatory score from baseline with no worsening of the Ishak fibrosis
- score. Not all patients provided both initial and end of follow-up biopsies (missing biopsy rates: 19% to
- 267 24% in the PEGASYS and 31% to 32% in the Lamivudine arms)
- <sup>5</sup>Change of 1 point or more in Ishak fibrosis score
- 269 \*p<0.001; \*\*p<0.01; \*\*\*p=0.012 (primary efficacy endpoints Cochran-Mantel-Haenszel test comparisons
- of PEGASYS to Lamivudine)

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- 272 PEGASYS co-administered with lamivudine did not result in any additional sustained
- 273 response when compared to PEGASYS monotherapy.
- 274 Conclusions regarding comparative efficacy of PEGASYS and lamivudine treatment
- based upon the end of follow-up results are limited by the different mechanisms of action
- of the two compounds. Most treatment effects of lamivudine are unlikely to persist 24
- weeks after therapy is withdrawn.

### 278 INDICATIONS AND USAGE

- 279 PEGASYS, peginterferon alfa-2a, alone or in combination with COPEGUS, is indicated
- 280 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. Patients in whom
- 282 efficacy was demonstrated included patients with compensated liver disease and
- 283 histological evidence of cirrhosis (Child-Pugh class A) and patients with HIV disease that
- 284 is clinically stable (e.g., antiretroviral therapy not required or receiving stable
- antiretroviral therapy).
- 286 PEGASYS is indicated for the treatment of adult patients with HBeAg positive and
- 287 HBeAg negative chronic hepatitis B who have compensated liver disease and evidence of
- 288 viral replication and liver inflammation.

### 289 **CONTRAINDICATIONS**

- 290 PEGASYS is contraindicated in patients with:
- Hypersensitivity to PEGASYS or any of its components
- Autoimmune hepatitis
- Hepatic decompensation (Child-Pugh score greater than 6 [class B and C]) in cirrhotic patients before or during treatment
- Hepatic decompensation with Child-Pugh score greater than or equal to 6 in cirrhotic
   CHC patients coinfected with HIV before or during treatment
- 297 PEGASYS is contraindicated in neonates and infants because it contains benzyl alcohol.
- 298 Benzyl alcohol is associated with an increased incidence of neurologic and other
- complications in neonates and infants, which are sometimes fatal.
- 300 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 (e.g., thalassemia major, sickle-cell anemia)

### 305 WARNINGS

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

### 310 Neuropsychiatric

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

### 323 Infections

- While fever may be associated with the flu-like syndrome reported commonly during
- interferon therapy, other causes of high or persistent fever must be ruled out, particularly
- in patients with neutropenia. Serious and severe infections (bacterial, viral, fungal), some
- fatal, have been reported during treatment with alpha interferons including PEGASYS.
- 328 Appropriate anti-infective therapy should be started immediately and discontinuation of
- 329 therapy should be considered.

### 330 Bone Marrow Toxicity

- 331 PEGASYS suppresses bone marrow function and may result in severe cytopenias.
- Ribavirin may potentiate the neutropenia and lymphopenia induced by alpha interferons
- 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
- monitored routinely during therapy (see PRECAUTIONS: Laboratory Tests).
- 336 PEGASYS and COPEGUS should be used with caution in patients with baseline
- neutrophil counts <1500 cells/mm<sup>3</sup>, with baseline platelet counts <90,000 cells/mm<sup>3</sup> or
- 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
- 340 (see DOSAGE AND ADMINISTRATION: Dose Modifications).

- 341 Severe neutropenia and thrombocytopenia occur with a greater incidence in HIV
- 342 coinfected patients than monoinfected patients and may result in serious infections or
- 343 bleeding (see ADVERSE REACTIONS).

#### 344 Cardiovascular Disorders

- 345 Hypertension, supraventricular arrhythmias, chest pain, and myocardial infarction have
- 346 been observed in patients treated with PEGASYS.
- 347 PEGASYS should be administered with caution to patients with pre-existing cardiac
- 348 disease. Because cardiac disease may be worsened by ribavirin-induced anemia, patients
- 349 with a history of significant or unstable cardiac disease should not use COPEGUS (see
- 350 WARNINGS: Anemia and COPEGUS Package Insert).

#### 351 **Cerebrovascular Disorders**

- 352 Ischemic and hemorrhagic cerebrovascular events have been observed in patients treated
- 353 with interferon alfa-based therapies, including PEGASYS. Events occurred in patients
- 354 with few or no reported risk factors for stroke, including patients less than 45 years of
- 355 age. Because these are spontaneous reports, estimates of frequency cannot be made and a
- 356 causal relationship between interferon alfa-based therapies and these events is difficult to
- 357 establish.

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### **Hepatic Failure and Hepatitis Exacerbations**

- 359 Chronic hepatitis C (CHC) patients with cirrhosis may be at risk of hepatic
- 360 decompensation and death when treated with alpha interferons, including PEGASYS.
- 361 Cirrhotic CHC patients coinfected with HIV receiving highly active antiretroviral therapy
- 362 (HAART) and interferon alfa-2a with or without ribavirin appear to be at increased risk
- 363 for the development of hepatic decompensation compared to patients not receiving
- 364 HAART. In Study 6, among 129 CHC/HIV cirrhotic patients receiving HAART, 14
- 365 (11%) of these patients across all treatment arms developed hepatic decompensation
- 366 resulting in 6 deaths. All 14 patients were on NRTIs, including stavudine, didanosine,
- 367 abacavir, zidovudine, and lamivudine. These small numbers of patients do not permit
- 368 discrimination between specific NRTIs for the associated risk. During treatment,
- 369 patients' clinical status and hepatic function should be closely monitored, and PEGASYS
- 370 treatment should be immediately discontinued if decompensation (Child-Pugh score ≥6)
- 371 is observed (see **CONTRAINDICATIONS**).
- 372 Exacerbations of hepatitis during hepatitis B therapy are not uncommon and are
- 373 characterized by transient and potentially severe increases in serum ALT. Chronic
- 374 hepatitis B patients experienced transient acute exacerbations (flares) of hepatitis B (ALT
- 375 elevation >10-fold higher than the upper limit of normal) during PEGASYS treatment
- 376 (12% and 18%) and post-treatment (7% and 12%) in HBeAg negative and HBeAg
- 377 positive patients, respectively. Marked transaminase flares while on PEGASYS therapy
- 378 have been accompanied by other liver test abnormalities. Patients experiencing ALT
- 379 flares should receive more frequent monitoring of liver function. PEGASYS dose
- 380 reduction should be considered in patients experiencing transaminase flares. If ALT
- 381 increases are progressive despite reduction of PEGASYS dose or are accompanied by
- 382 increased bilirubin or evidence of hepatic decompensation, PEGASYS should be

- immediately discontinued (see ADVERSE REACTIONS: Chronic Hepatitis B and
- 384 DOSAGE AND ADMINISTRATION: Dose Modifications).

## 385 Hypersensitivity

- 386 Severe acute hypersensitivity reactions (e.g., urticaria, angioedema, bronchoconstriction,
- and anaphylaxis) have been rarely observed during alpha interferon and ribavirin therapy.
- 388 If such reaction occurs, therapy with PEGASYS and COPEGUS should be discontinued
- and appropriate medical therapy immediately instituted. Serious skin reactions including
- 390 vesiculobullous eruptions, reactions in the spectrum of Stevens Johnson Syndrome
- 391 (erythema multiforme major) with varying degrees of skin and mucosal involvement and
- 392 exfoliative dermatitis (erythroderma) have been rarely reported in patients receiving
- 393 PEGASYS with and without ribavirin. Patients developing signs or symptoms of severe
- 394 skin reactions must discontinue therapy (see ADVERSE REACTIONS: Postmarketing
- 395 Experience).

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### **Endocrine Disorders**

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

### 403 Autoimmune Disorders

- 404 Development or exacerbation of autoimmune disorders including myositis, hepatitis,
- 405 thrombotic thrombocytopenic purpura, idiopathic thrombocytopenic purpura, psoriasis,
- 406 rheumatoid arthritis, interstitial nephritis, thyroiditis, and systemic lupus erythematosus
- 407 have been reported in patients receiving alpha interferon. PEGASYS should be used with
- 408 caution in patients with autoimmune disorders.

### Pulmonary Disorders

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

### 415 Colitis

- 416 Ulcerative and hemorrhagic/ischemic colitis, sometimes fatal, have been observed within
- 417 12 weeks of starting alpha interferon treatment. Abdominal pain, bloody diarrhea, and
- 418 fever are the typical manifestations of colitis. PEGASYS should be discontinued
- 419 immediately if these symptoms develop. The colitis usually resolves within 1 to 3 weeks
- 420 of discontinuation of alpha interferon.

### **Pancreatitis**

- 422 Pancreatitis, sometimes fatal, has occurred during alpha interferon and ribavirin
- 423 treatment. PEGASYS and COPEGUS should be suspended if symptoms or signs

- 424 suggestive of pancreatitis are observed. PEGASYS and COPEGUS should be
- discontinued in patients diagnosed with pancreatitis.

## 426 Ophthalmologic Disorders

- 427 Decrease or loss of vision, retinopathy including macular edema, retinal artery or vein
- 428 thrombosis, retinal hemorrhages and cotton wool spots, optic neuritis, papilledema and
- 429 serous retinal detachment are induced or aggravated by treatment with PEGASYS or
- 430 other alpha interferons. All patients should receive an eye examination at baseline.
- 431 Patients with pre-existing ophthalmologic disorders (e.g., diabetic or hypertensive
- 432 retinopathy) should receive periodic ophthalmologic exams during interferon alpha
- 433 treatment. Any patient who develops ocular symptoms should receive a prompt and
- complete eye examination. PEGASYS treatment should be discontinued in patients who
- develop new or worsening ophthalmologic disorders.

### 436 Pregnancy: Use with Ribavirin (also, see COPEGUS Package Insert)

- Ribavirin may cause birth defects and/or death of the exposed fetus. Extreme care
- must be taken to avoid pregnancy in female patients and in female partners of male
- 439 patients taking PEGASYS and COPEGUS combination therapy. COPEGUS
- 440 THERAPY SHOULD NOT BE STARTED UNLESS A REPORT OF A
- 441 NEGATIVE PREGNANCY TEST HAS BEEN OBTAINED IMMEDIATELY
- PRIOR TO INITIATION OF THERAPY. Women of childbearing potential and
- 443 men must use two forms of effective contraception during treatment and for at least
- 6 months after treatment has concluded. Routine monthly pregnancy tests must be
- performed during this time (see BOXED WARNING, CONTRAINDICATIONS,
- 446 PRECAUTIONS: Information for Patients, and COPEGUS Package Insert).

### 447 Anemia

- 448 The primary toxicity of ribavirin is hemolytic anemia. Hemoglobin <10 g/dL was
- observed in approximately 13% of COPEGUS and PEGASYS treated patients in chronic
- 450 hepatitis C clinical trials (see PRECAUTIONS: Laboratory Tests). The anemia
- associated with COPEGUS occurs within 1 to 2 weeks of initiation of therapy with
- 452 maximum drop in hemoglobin observed during the first eight weeks. BECAUSE THE
- 453 INITIAL DROP IN HEMOGLOBIN MAY BE SIGNIFICANT, IT IS ADVISED THAT
- 454 HEMOGLOBIN OR HEMATOCRIT BE OBTAINED PRE-TREATMENT AND AT
- 455 WEEK 2 AND WEEK 4 OF THERAPY OR MORE FREQUENTLY IF CLINICALLY
- 456 INDICATED. Patients should then be followed as clinically appropriate.
- 457 Fatal and nonfatal myocardial infarctions have been reported in patients with anemia
- 458 caused by ribavirin. Patients should be assessed for underlying cardiac disease before
- 459 initiation of ribavirin therapy. Patients with pre-existing cardiac disease should have
- 460 electrocardiograms administered before treatment, and should be appropriately monitored
- during therapy. If there is any deterioration of cardiovascular status, therapy should be
- suspended or discontinued (see DOSAGE AND ADMINISTRATION: COPEGUS
- 463 **Dosage Modification Guidelines**). Because cardiac disease may be worsened by drug-
- 464 induced anemia, patients with a history of significant or unstable cardiac disease should
- not use COPEGUS (see COPEGUS Package Insert).

- 466 Renal
- 467 It is recommended that renal function be evaluated in all patients started on COPEGUS.
- 468 COPEGUS should not be administered to patients with creatinine clearance <50 mL/min
- 469 (see CLINICAL PHARMACOLOGY: Special Populations).

### 470 PRECAUTIONS

471 General

479

- The safety and efficacy of PEGASYS alone or in combination with COPEGUS have not
- 473 been established in:
- Patients who have failed alpha interferon treatment with or without ribavirin
- Liver or other organ transplant recipients
- Hepatitis B patients coinfected with HCV or HIV
- Hepatitis C patients coinfected with HBV or coinfected with HIV with a CD4+ cell
   count <100 cells/μL</li>
- Caution should be exercised in initiating treatment in any patient with baseline risk of severe anemia (e.g., spherocytosis, history of GI bleeding).
- 482 Renal Impairment
- 483 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
- 485 toxicity should be closely monitored. Doses of PEGASYS should be adjusted
- 486 accordingly, PEGASYS should be used with caution in patients with creatinine clearance
- 487 <50 mL/min (see **DOSAGE AND ADMINISTRATION: Dose Modifications**).
- 488 COPEGUS should not be used in patients with creatinine clearance <50 mL/min (see
- 489 **COPEGUS Package Insert**).
- 490 Information for Patients
- 491 Patients receiving PEGASYS alone or in combination with COPEGUS should be
- 492 directed in its appropriate use, informed of the benefits and risks associated with
- 493 treatment, and referred to the PEGASYS and, if applicable, COPEGUS (ribavirin)
- 494 MEDICATION GUIDES.
- 495 PEGASYS and COPEGUS combination therapy must not be used by women who are
- 496 pregnant or by men whose female partners are pregnant. COPEGUS therapy should not
- 497 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
- 499 female partners of childbearing potential must be advised of the teratogenic/embryocidal
- 500 risks and must be instructed to practice effective contraception during COPEGUS therapy
- and for 6 months post-therapy. Patients should be advised to notify the healthcare
- 502 provider immediately in the event of a pregnancy (see CONTRAINDICATIONS and
- 503 WARNINGS).

- Women of childbearing potential and men must use two forms of effective contraception
- during treatment and during the 6 months after treatment has been stopped; routine
- 506 monthly pregnancy tests must be performed during this time (see
- 507 CONTRAINDICATIONS and COPEGUS Package Insert).
- To monitor maternal and fetal outcomes of pregnant women exposed to COPEGUS, the
- 509 Ribavirin Pregnancy Registry has been established. Patients should be encouraged to
- 510 register by calling 1-800-593-2214.
- Patients should be advised that laboratory evaluations are required before starting therapy
- and periodically thereafter (see Laboratory Tests). Patients should be instructed to
- 513 remain well hydrated, especially during the initial stages of treatment. Patients should be
- advised to take COPEGUS with food.
- Patients should be informed that it is not known if therapy with PEGASYS alone or in
- 516 combination with COPEGUS will prevent transmission of HCV or HBV infection to
- others or prevent cirrhosis, liver failure or liver cancer that might result from HCV or
- 518 HBV infection. Patients who develop dizziness, confusion, somnolence, and fatigue
- should be cautioned to avoid driving or operating machinery.
- 520 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
- 523 syringes. The full container should be disposed of according to the directions provided by
- 524 the physician (see **MEDICATION GUIDE**).

### 525 Laboratory Tests

- 526 Before beginning PEGASYS or PEGASYS and COPEGUS combination therapy,
- 527 standard hematological and biochemical laboratory tests are recommended for all
- patients. Pregnancy screening for women of childbearing potential must be performed.
- 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 weeks, and then
- every 4 to 6 weeks or more frequently if abnormalities were found. Thyroid stimulating
- hormone (TSH) was measured every 12 weeks. Monthly pregnancy testing should be
- performed during combination therapy and for 6 months after discontinuing therapy.
- The entrance criteria used for the clinical studies of PEGASYS may be considered as a
- guideline to acceptable baseline values for initiation of treatment:
- Platelet count ≥90,000 cells/mm³ (as low as 75,000 cells/mm³ in HCV patients with
- cirrhosis or 70,000 cells/mm<sup>3</sup> in patients with CHC and HIV)
- Absolute neutrophil count (ANC) ≥1500 cells/mm<sup>3</sup>
- Serum creatinine concentration <1.5 x upper limit of normal

- TSH and T<sub>4</sub> within normal limits or adequately controlled thyroid function
- CD4+ cell count ≥200 cells/μL or CD4+ cell count ≥100 cells/μL but <200 cells/μL and HIV-1 RNA <5000 copies/mL in patients coinfected with HIV
- Hemoglobin ≥12 g/dL for women and ≥13 g/dL for men in CHC monoinfected patients
- Hemoglobin ≥11 g/dL for women and ≥12 g/dL for men in patients with CHC and HIV
- 550 PEGASYS treatment was associated with decreases in WBC, ANC, lymphocytes, and
- platelet counts often starting within the first 2 weeks of treatment (see ADVERSE
- 552 **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
- 555 must be ruled out, particularly in patients with neutropenia (see WARNINGS:
- 556 Infections).
- In chronic hepatitis C, transient elevations in ALT (2-fold to 5-fold above baseline) were
- observed in some 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 is accompanied by increased bilirubin, PEGASYS therapy
- 561 should be discontinued (see DOSAGE AND ADMINISTRATION: Dose
- 562 Modifications).
- Unlike hepatitis C, during hepatitis B therapy and follow up, transient elevations in ALT
- of 5 to 10 x ULN were observed in 25% and 27% and of >10 x ULN were observed in
- 565 12% and 18%, of HBeAg negative and HBeAg positive patients, respectively. These
- 566 ALT elevations have been accompanied by other liver test abnormalities (see
- 567 WARNINGS: Hepatic Failure and Hepatitis Exacerbations and DOSAGE AND
- 568 ADMINISTRATION: Dose Modifications).

### 569 **Drug Interactions**

- 570 Theophylline
- 571 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
- 573 serum levels should be monitored and appropriate dose adjustments considered for
- patients given both theophylline and PEGASYS (see CLINICAL PHARMACOLOGY:
- 575 **Drug Interactions**).
- 576 Methadone
- 577 In a PK study of HCV patients concomitantly receiving methodone, treatment with
- 578 PEGASYS once weekly for 4 weeks was associated with methadone levels that were
- 579 10% to 15% higher than at baseline (see CLINICAL PHARMACOLOGY: Drug
- Interactions). The clinical significance of this finding is unknown; however, patients
- should be monitored for the signs and symptoms of methadone toxicity.

- 582 Nucleoside Analogues
- 583 NRTIs
- In Study 6 among the CHC/HIV coinfected cirrhotic patients receiving NRTIs cases of
- hepatic decompensation (some fatal) were observed (see WARNINGS: Hepatic Failure
- and Hepatitis Exacerbations).
- Patients receiving PEGASYS/COPEGUS and NRTIs should be closely monitored for
- treatment associated toxicities. Physicians should refer to prescribing information for the
- 589 respective NRTIs for guidance regarding toxicity management. In addition, dose
- reduction or discontinuation of PEGASYS, COPEGUS or both should also be considered
- if worsening toxicities are observed (see WARNINGS, PRECAUTIONS, DOSAGE
- 592 AND ADMINISTRATION: Dose Modifications).
- 593 Didanosine
- 594 Co-administration of COPEGUS and didanosine is not recommended. Reports of fatal
- 595 hepatic failure, as well as peripheral neuropathy, pancreatitis, and symptomatic
- 596 hyperlactatemia/lactic acidosis have been reported in clinical trials (see CLINICAL
- 597 PHARMACOLOGY: Drug Interactions).
- 598 Zidovudine
- 599 In Study 6, patients who were administered zidovudine in combination with
- 600 PEGASYS/COPEGUS developed severe neutropenia (ANC <500) and severe anemia
- 601 (hemoglobin <8 g/dL) more frequently than similar patients not receiving zidovudine
- 602 (neutropenia 15% vs. 9%) (anemia 5% vs. 1%). Discontinuation of zidovudine should be
- 603 considered as medically appropriate. Dose reduction or discontinuation of PEGASYS,
- 604 COPEGUS or both should also be considered if worsening clinical toxicities are
- observed, including hepatic decompensation (e.g., Childs Pugh > 6).
- 606 Lamivudine, Stavudine, and Zidovudine
- 607 In vitro studies have shown ribavirin can reduce the phosphorylation of pyrimidine
- 608 nucleoside analogs such as lamivudine, stavudine, and zidovudine. No evidence of a
- 609 pharmacokinetic or pharmacodynamic interaction was seen when ribavirin was co-
- administered with lamivudine, stavudine, and/or zidovudine in HIV/HCV coinfected
- patients (see CLINICAL PHARMACOLOGY: Drug Interactions).
- 612 Carcinogenesis, Mutagenesis, Impairment of Fertility
- 613 Carcinogenesis
- PEGASYS has not been tested for its carcinogenic potential.
- 615 Mutagenesis
- 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
- the presence or absence of metabolic activation.

- 619 Use with Ribavirin
- 620 Ribavirin is genotoxic and mutagenic. The carcinogenic potential of ribavirin has not
- 621 been fully determined. In a p53 (+/-) mouse carcinogenicity study at doses up to the
- 622 maximum tolerated dose of 100 mg/kg/day ribavirin was not oncogenic. However, on a
- 623 body surface area basis, this dose was 0.5 times maximum recommended human 24-hour
- 624 dose of ribavirin. A study in rats to assess the carcinogenic potential of ribavirin is
- 625 ongoing (see COPEGUS Package Insert).
- 626 Impairment of Fertility
- 627 PEGASYS may impair fertility in women. Prolonged menstrual cycles and/or
- 628 amenorrhea were observed in female cynomolgus monkeys given sc injections of
- 629 600 µg/kg/dose (7200 µg/m<sup>2</sup>/dose) of PEGASYS every other day for one month, at
- approximately 180 times the recommended weekly human dose for a 60 kg person (based 630
- 631 on body surface area). Menstrual cycle irregularities were accompanied by both a
- 632 decrease and delay in the peak 17\beta-estradiol and progesterone levels following
- 633 administration of PEGASYS to female monkeys. A return to normal menstrual rhythm
- 634 followed cessation of treatment. Every other day dosing with 100 µg/kg (1200 µg/m<sup>2</sup>)
- 635 PEGASYS (equivalent to approximately 30 times the recommended human dose) had no
- 636 effects on cycle duration or reproductive hormone status.
- 637 The effects of PEGASYS on male fertility have not been studied. However, no adverse
- 638 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 10<sup>6</sup> IU/kg/day. 639
- 640 Use with Ribavirin
- 641 Ribavirin has shown reversible toxicity in animal studies of male fertility (see
- 642 **COPEGUS Package Insert).**
- 643 Pregnancy
- 644 Pregnancy: Category C
- 645 PEGASYS has not been studied for its teratogenic effect. Non-pegylated interferon alfa-
- 646 2a treatment of pregnant Rhesus monkeys at approximately 20 to 500 times the human
- 647 weekly dose resulted in a statistically significant increase in abortions. No teratogenic
- 648 effects were seen in the offspring delivered at term. PEGASYS should be assumed to
- 649 have abortifacient potential. There are no adequate and well-controlled studies of
- 650 PEGASYS in pregnant women. PEGASYS is to be used during pregnancy only if the
- 651
- potential benefit justifies the potential risk to the fetus. PEGASYS is recommended for
- 652 use in women of childbearing potential only when they are using effective contraception
- 653 during therapy.
- 654 **Pregnancy: Category X: Use With Ribavirin (see CONTRAINDICATIONS)**
- 655 Significant teratogenic and/or embryocidal effects have been demonstrated in all
- 656 animal species exposed to ribavirin. COPEGUS therapy is contraindicated in
- 657 women who are pregnant and in the male partners of women who are pregnant (see
- 658 CONTRAINDICATIONS, WARNINGS, and COPEGUS Package Insert).

- 659 Ribavirin Pregnancy Registry
- A Ribavirin Pregnancy Registry has been established to monitor maternal and fetal
- outcomes of pregnancies of female patients and female partners of male patients exposed
- 662 to ribavirin during treatment and for 6 months following cessation of treatment.
- Healthcare providers and patients are encouraged to report such cases by calling 1-800-
- 664 593-2214.

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### **Nursing Mothers**

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

### 671 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.
- 674 PEGASYS contains benzyl alcohol. Benzyl alcohol has been reported to be associated
- with an increased incidence of neurological and other complications in neonates and
- infants, which are sometimes fatal (see **CONTRAINDICATIONS**).

### 677 Geriatric Use

- Younger patients have higher virologic response rates than older patients. Clinical studies
- 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
- of vounger subjects. Adverse reactions related to alpha interferons, such as CNS, cardiac,
- and systemic (e.g., 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
- 687 monitor renal function. PEGASYS should be used with caution in patients with creatinine
- 688 clearance <50 mL/min and COPEGUS should not be administered to patients with
- 689 creatinine clearance <50 mL/min.

### ADVERSE REACTIONS

- 691 PEGASYS alone or in combination with COPEGUS causes a broad variety of serious
- adverse reactions (see **BOXED WARNING** and **WARNINGS**). The most common life-
- 693 threatening or fatal events induced or aggravated by PEGASYS and COPEGUS were
- 694 depression, suicide, relapse of drug abuse/overdose, and bacterial infections, each
- occurring at a frequency of <1%. Hepatic decompensation occurred in 2% (10/574) of
- 696 CHC/HIV patients (see WARNINGS: Hepatic Failure and Hepatitis Exacerbations).
- In all hepatitis C studies, one or more serious adverse reactions occurred in 10% of CHC
- monoinfected patients and in 19% of CHC/HIV patients receiving PEGASYS alone or in
- 699 combination with COPEGUS. The most common serious adverse event (3% in CHC and

- 700 5% in CHC/HIV) was bacterial infection (e.g., sepsis, osteomyelitis, endocarditis,
- 701 pyelonephritis, pneumonia). Other SAEs occurred at a frequency of <1% and included:
- suicide, suicidal ideation, psychosis, aggression, anxiety, drug abuse and drug overdose,
- angina, hepatic dysfunction, fatty liver, cholangitis, arrhythmia, diabetes mellitus,
- autoimmune phenomena (e.g., hyperthyroidism, hypothyroidism, sarcoidosis, systemic
- lupus erythematosus, rheumatoid arthritis), peripheral neuropathy, aplastic anemia, peptic
- 706 ulcer, gastrointestinal bleeding, pancreatitis, colitis, corneal ulcer, pulmonary embolism.
- 707 coma, myositis, cerebral hemorrhage, thrombotic thrombocytopenic purpura, psychotic
- disorder, and hallucination.
- Nearly all patients in clinical trials experienced one or more adverse events. For hepatitis
- 710 C patients, the most commonly reported adverse reactions were psychiatric reactions.
- 711 including depression, insomnia, irritability, anxiety, and flu-like symptoms such as
- fatigue, pyrexia, myalgia, headache, and rigors. Other common reactions were anorexia,
- 713 nausea and vomiting, diarrhea, arthralgias, injection site reactions, alopecia, and pruritus.
- 714 Overall 11% of CHC monoinfected patients receiving 48 weeks of therapy with
- 715 PEGASYS either alone or in combination with COPEGUS discontinued therapy; 16% of
- 716 CHC/HIV coinfected patients discontinued therapy. The most common reasons for
- 717 discontinuation of therapy were psychiatric, flu-like syndrome (e.g., lethargy, fatigue,
- 718 headache), dermatologic, and gastrointestinal disorders and laboratory abnormalities
- 719 (thrombocytopenia, neutropenia, and anemia).
- 720 Overall 39% of patients with CHC or CHC/HIV required modification of PEGASYS
- 721 and/or COPEGUS therapy. The most common reason for dose modification of
- 722 PEGASYS in CHC and CHC/HIV patients was for laboratory abnormalities, neutropenia
- 723 (20% and 27%, respectively) and thrombocytopenia (4% and 6%, respectively). The most
- 724 common reason for dose modification of COPEGUS in CHC and CHC/HIV patients was
- anemia (22% and 16%, respectively).
- 726 PEGASYS dose was reduced in 12% of patients receiving 1000 mg to 1200 mg
- 727 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
- 729 COPEGUS for 48 weeks and in 12% of patients receiving 800 mg COPEGUS for 24
- 730 weeks.
- 731 Chronic hepatitis C monoinfected patients treated for 24 weeks with PEGASYS and 800
- mg COPEGUS were observed to have lower incidence of serious adverse events (3% vs.
- 733 10%), Hgb <10 g/dL (3% vs. 15%), dose modification of PEGASYS (30% vs. 36%) and
- 734 COPEGUS (19% vs. 38%) and of withdrawal from treatment (5% vs. 15%) compared to
- patients treated for 48 weeks with PEGASYS and 1000 mg or 1200 mg COPEGUS. On
- patients treated for 10 weeks with 1 Edito 15 and 1000 ing of 1200 ing CO1 Edob. On
- 736 the other hand the overall incidence of adverse events appeared to be similar in the two
- 737 treatment groups.
- 738 Because clinical trials are conducted under widely varying and controlled
- 739 conditions, adverse reaction rates observed in clinical trials of a drug cannot be
- 740 directly compared to rates in the clinical trials of another drug. Also, the adverse
- event rates listed here may not predict the rates observed in a broader patient
- 742 population in clinical practice.

Table 6 Adverse Reactions Occurring in ≥5% of Patients in Chronic Hepatitis C Clinical Trials (Pooled Studies 1, 2, 3, and Study 4)

		herapy (Pooled lies 1-3)		nation Therapy idy 4
Body System	PEGASYS 180 μg 48 week†	ROFERON-A*†	PEGASYS 180 µg + 1000 mg or 1200 mg COPEGUS 48 week**	Intron <sup>®</sup> A + 1000 mg or 1200 mg REBETOL <sup>®</sup> 48 week**
	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
Diarrhea	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
Thrombocytopenia	5	2	5	<1
Metabolic and Nutritional				
Anorexia	17	17	24	26
Weight decrease	4	3	10	10

		therapy (Pooled lies 1-3)	CHC Combination Therapy Study 4		
Body System	PEGASYS 180 μg 48 week†	ROFERON-A*†	PEGASYS 180 μg + 1000 mg or 1200 mg COPEGUS 48 week**	Intron® A + 1000 mg or 1200 mg REBETOL® 48 week**	
	N=559	N=554	N=451	N=443	
	%	%	%	%	
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	
Resistance Mechanism Disorders					
Overall	10	6	12	10	
Psychiatric	***************************************				
Irritability/Anxiety/	19	22	33	38	
Nervousness					
Insomnia	19	23	30	37	
Depression	18	19	20	28	
Concentration impairment	8	10	10	13	
Mood alteration	3	2	5	6	
Respiratory, Thoracic and Mediastinal					
Dyspnea	4	2	13	14	
Cough	4	3	10	7	
Dyspnea exertional	<1	<1	4	7	

		therapy (Pooled lies 1-3)	CHC Combination Therap Study 4		
Body System	PEGASYS 180 µg 48 week†	ROFERON-A*†	PEGASYS 180 µg + 1000 mg or 1200 mg COPEGUS 48 week**	Intron® A + 1000 mg or 1200 mg REBETOL® 48 week**	
	N=559	N=554	N=451	N=443	
	%	%	%	%	
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	

<sup>746 †</sup> Pooled studies 1, 2, and 3

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### **CHC With HIV Coinfection**

The adverse event profile of coinfected patients treated with PEGASYS and COPEGUS in Study 6 was generally similar to that shown for monoinfected patients in Study 4 (**Table 6**). Events occurring more frequently in coinfected patients were neutropenia (40%), anemia (14%), thrombocytopenia (8%), weight decrease (16%), and mood alteration (9%).

## Chronic Hepatitis B

In clinical trials of 48 week treatment duration, the adverse event profile of PEGASYS in chronic hepatitis B was similar to that seen in chronic hepatitis C PEGASYS monotherapy use, except for exacerbations of hepatitis (see WARNINGS: Hepatic Failure and Hepatitis Exacerbations). Six percent of PEGASYS treated patients in the hepatitis B studies experienced one or more serious adverse events.

<sup>\*</sup> Either 3 MIU or 6/3 MIU of ROFERON-A

<sup>748 \*\*</sup>Study 4

<sup>‡</sup> Severe hematologic abnormalities (lymphocyte <0.5 x  $10^9$ /L; hemoglobin <10 g/dL; neutrophil <0.75 x  $10^9$ /L; platelet <50 x  $10^9$ /L).

- 764 The most common or important serious adverse events in the hepatitis B studies were
- 765 infections (sepsis, appendicitis, tuberculosis, influenza), hepatitis B flares, anaphylactic
- 766 shock, thrombotic thrombocytopenic purpura.
- 767 The most commonly observed adverse reactions were pyrexia (54% vs. 4%), headache
- 768 (27% vs. 9%), fatigue (24% vs. 10%), myalgia (26% vs. 4%), alopecia (18% vs. 2%), and
- 769 anorexia (16% vs. 3%) in the PEGASYS and lamivudine groups respectively.
- 770 Overall 5% of hepatitis B patients discontinued PEGASYS therapy and 40% of patients
- 771 required modification of PEGASYS dose. The most common reason for dose
- 772 modification in patients receiving PEGASYS therapy was for laboratory abnormalities
- 773 including neutropenia (20%), thrombocytopenia (13%), and ALT disorders (11%).

### 774 **Laboratory Test Values**

- 775 The laboratory test values observed in the hepatitis B trials (except where noted below)
- 776 were similar to those seen in the PEGASYS monotherapy hepatitis C trials.

#### 777 Neutrophils

- 778 In the hepatitis C studies, decreases in neutrophil count below normal were observed in
- 779 95% of all patients treated with PEGASYS either alone or in combination with
- 780 COPEGUS. Severe potentially life-threatening neutropenia (ANC <  $0.5 \times 10^9$ /L) occurred
- 781 in 5% of CHC patients and 12% of CHC/HIV patients receiving PEGASYS either alone
- 782 or in combination with COPEGUS. Modification of PEGASYS dose for neutropenia
- 783 occurred in 17% of patients receiving PEGASYS monotherapy and 22% of patients
- 784 receiving PEGASYS/COPEGUS combination therapy. In the CHC/HIV patients 27%
- 785 required modification of interferon dosage for neutropenia. Two percent of patients with
- 786 CHC and 10% of patients with CHC/HIV required permanent reductions of PEGASYS
- 787 dosage and <1% required permanent discontinuation. Median neutrophil counts return to
- 788 pre-treatment levels 4 weeks after cessation of therapy (see DOSAGE AND
- 789 **ADMINISTRATION: Dose Modifications).**

#### 790 Lymphocytes

- 791 Decreases in lymphocyte count are induced by interferon alpha therapy. PEGASYS plus
- 792 COPEGUS combination therapy induced decreases in median total lymphocyte counts
- 793 (56% in CHC and 40% in CHC/HIV, with median decrease of 1170 cells/mm<sup>3</sup> in CHC
- and 800 cells/mm<sup>3</sup> in CHC/HIV). In the hepatitis C studies, lymphopenia was observed 794
- 795 during both monotherapy (81%) and combination therapy with PEGASYS and
- 796 COPEGUS (91%). Severe lymphopenia (<0.5 x 10<sup>9</sup>/L) occurred in approximately 5% of
- 797 all monotherapy patients and 14% of all combination PEGASYS and COPEGUS therapy
- 798
- recipients. Dose adjustments were not required by protocol. The clinical significance of
- 799 the lymphopenia is not known.
- 800 In CHC with HIV coinfection, CD4 counts decreased by 29% from baseline (median
- decrease of 137 cells/mm<sup>3</sup>) and CD8 counts decreased by 44% from baseline (median 801
- decrease of 389 cells/mm<sup>3</sup>) in the PEGASYS plus COPEGUS combination therapy arm. 802
- 803 Median lymphocyte CD4 and CD8 counts return to pre-treatment levels after 4 to 12
- 804 weeks of the cessation of therapy. CD4% did not decrease during treatment.

- 805 Platelets
- In the hepatitis C studies, platelet counts decreased in 52% of CHC patients and 51% of
- 807 CHC/HIV patients treated with PEGASYS alone (respectively median decrease of 41%
- and 35% from baseline), and in 33% of CHC patients and 47% of CHC/HIV patients
- receiving combination therapy with COPEGUS (median decrease of 30% from baseline).
- Moderate to severe thrombocytopenia (<50,000/mm<sup>3</sup>) was observed in 4% of CHC and
- 811 8% of CHC/HIV patients. Median platelet counts return to pre-treatment levels 4 weeks
- after the cessation of therapy.
- 813 Hemoglobin
- In the hepatitis C studies, the hemoglobin concentration decreased below 12 g/dL in 17%
- 815 (median Hgb reduction of 2.2 g/dL) of monotherapy and 52% (median Hgb reduction of
- 816 3.7 g/dL) of combination therapy patients. Severe anemia (Hgb <10 g/dL) was
- encountered in 13% of all patients receiving combination therapy and in 2% of CHC
- 818 patients and 8% of CHC/HIV patients receiving PEGASYS monotherapy. Dose
- modification for anemia in COPEGUS recipients treated for 48 weeks occurred in 22% of
- 820 CHC patients and 16% of CHC/HIV patients (see DOSAGE AND
- 821 **ADMINISTRATION: Dose Modifications).**
- 822 Triglycerides
- 823 Triglyceride levels are elevated in patients receiving alfa interferon therapy and were
- 824 elevated in the majority of patients participating in clinical studies receiving either
- 825 PEGASYS alone or in combination with COPEGUS. Random levels ≥400 mg/dL were
- 826 observed in about 20% of CHC patients. Severe elevations of triglycerides (>1000)
- mg/dL) occurred in 2% of CHC monoinfected patients.
- 828 In HCV/HIV coinfected patients, fasting levels ≥400 mg/dL were observed in up to 36%
- 829 of patients receiving either PEGASYS alone or in combination with COPEGUS. Severe
- elevations of triglycerides (>1000 mg/dL) occurred in 7% of coinfected patients.
- 831 ALT Elevations
- 832 Chronic Hepatitis C
- One percent of patients in the hepatitis C trials experienced marked elevations (5- to 10-
- fold above the upper limit of normal) in ALT levels during treatment and follow-up.
- These transaminase elevations were on occasion associated with hyperbilirubinemia and
- were managed by dose reduction or discontinuation of study treatment. Liver function
- 837 test abnormalities were generally transient. One case was attributed to autoimmune
- hepatitis, which persisted beyond study medication discontinuation (see **DOSAGE AND**
- 839 **ADMINISTRATION: Dose Modifications**).
- 840 Chronic Hepatitis B
- 841 Transient ALT elevations are common during hepatitis B therapy with PEGASYS.
- Wenty-five percent and 27% of patients experienced elevations of 5 to 10 x ULN and
- 843 12% and 18% had elevations of >10 x ULN during treatment of HBeAg negative and
- HBeAg positive disease, respectively. Flares have been accompanied by elevations of
- total bilirubin and alkaline phosphatase and less commonly with prolongation of PT and
- reduced albumin levels. Eleven percent of patients had dose modifications due to ALT

- flares and <1% of patients were withdrawn from treatment (see WARNINGS: Hepatic
- 848 Failure and Hepatitis Exacerbations and DOSAGE AND ADMINISTRATION:
- **Dose Modifications**).
- ALT flares of 5 to 10 x ULN occurred in 13% and 16% of patients, while ALT flares of
- >10 x ULN occurred in 7% and 12% of patients in HBeAg negative and HBeAg positive
- disease, respectively, after discontinuation of PEGASYS therapy.
- 853 Thyroid Function
- 854 PEGASYS alone or in combination with COPEGUS was associated with the
- development of abnormalities in thyroid laboratory values, some with associated clinical
- 856 manifestations. In the hepatitis C studies, hypothyroidism or hyperthyroidism requiring
- treatment, dose modification or discontinuation occurred in 4% and 1% of PEGASYS
- 858 treated patients and 4% and 2% of PEGASYS and COPEGUS treated patients,
- 859 respectively. Approximately half of the patients, who developed thyroid abnormalities
- 860 during PEGASYS treatment, still had abnormalities during the follow-up period (see
- 861 **PRECAUTIONS:** Laboratory Tests).
- 862 Immunogenicity
- 863 Chronic Hepatitis C
- 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
- percent of patients (25/835) receiving PEGASYS with or without COPEGUS, developed
- low-titer neutralizing antibodies (using an assay with a sensitivity of 100 INU/mL).
- 868 Chronic Hepatitis B
- Wenty-nine percent (42/143) of hepatitis B patients treated with PEGASYS for 24
- weeks developed binding antibodies to interferon alfa-2a, as assessed by an ELISA assay.
- 871 Thirteen percent of patients (19/143) receiving PEGASYS developed low-titer
- neutralizing antibodies (using an assay with a sensitivity of 100 INU/mL).
- 873 The clinical and pathological significance of the appearance of serum neutralizing
- antibodies is unknown. No apparent correlation of antibody development to clinical
- response or adverse events was observed. The percentage of patients whose test results
- were considered positive for antibodies is highly dependent on the sensitivity and
- specificity of the assays.
- Additionally, the observed incidence of antibody positivity in these assays may be
- 879 influenced by several factors including sample timing and handling, concomitant
- medications, and underlying disease. For these reasons, comparison of the incidence of
- antibodies to PEGASYS with the incidence of antibodies to other products may be
- misleading.
- 883 Postmarketing Experience
- The following adverse reactions have been identified and reported during post-approval
- 885 use of PEGASYS therapy: dehydration, hearing impairment, hearing loss, and serious
- 886 skin reactions (see WARNINGS: Hypersensitivity). Because these reactions are
- reported voluntarily from a population of uncertain size, it is not always possible to

- 888 reliably estimate their frequency or establish a causal relationship to drug exposure.
- Decisions to include these reactions in labeling are typically based on one or more of the
- 890 following factors: (1) seriousness of the reaction, (2) frequency of reporting or (3)
- strength of causal connection to PEGASYS.

### 892 **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
- 897 enzymes, neutropenia, and thrombocytopenia. There is no specific antidote for
- 898 PEGASYS. Hemodialysis and peritoneal dialysis are not effective.

### 899 DOSAGE AND ADMINISTRATION

- There are no safety and efficacy data on treatment of chronic hepatitis C or hepatitis B for
- 901 longer than 48 weeks. For patients with hepatitis C, consideration should be given to
- 902 discontinuing therapy after 12 to 24 weeks of therapy if the patient has failed to
- 903 demonstrate an early virologic response defined as undetectable HCV RNA or at least a
- 904 2log<sub>10</sub> reduction from baseline in HCV RNA titer by 12 weeks of therapy (see
- 905 CLINICAL STUDIES).
- 906 A patient should self-inject PEGASYS only if the physician determines that it is
- 907 appropriate and the patient agrees to medical follow-up as necessary and training in
- 908 proper injection technique has been provided to him/her (see illustrated PEGASYS
- 909 MEDICATION GUIDE for directions on injection site preparation and injection
- 910 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
- and prefilled syringes with particulate matter or discoloration should be returned to the
- 914 pharmacist.

### 915 Chronic Hepatitis C

### 916 **PEGASYS Monotherapy**

- 917 The recommended dose of PEGASYS monotherapy for chronic hepatitis C is 180 μg (1.0
- 918 mL vial or 0.5 mL prefilled syringe) once weekly for 48 weeks by subcutaneous
- administration in the abdomen or thigh.

### 920 PEGASYS and COPEGUS Combination Therapy

- 921 The recommended dose of PEGASYS when used in combination with ribavirin for
- 922 chronic hepatitis C is 180 µg (1.0 mL vial or 0.5 mL prefilled syringe) once weekly. The
- 923 recommended dose of COPEGUS and duration for PEGASYS/COPEGUS therapy is
- based on viral genotype (see **Table 7**).
- 925 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 (e.g., genotype), response to therapy, and tolerability of the regimen.

- 928 Since COPEGUS absorption increases when administered with a meal, patients are
- 929 advised to take COPEGUS with food.

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### Table 7 PEGASYS and COPEGUS Dosing Recommendations

	_	
PEGASYS Dose	COPEGUS Dose	Duration
100	<75 kg = 1000 mg	48 weeks
180 μg	≥75 kg = 1200 mg	48 weeks
180 μg	800 mg	24 weeks
	180 μg	<75  kg = 1000  mg $≥75  kg = 1200  mg$

- 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.

### 934 CHC with HIV Coinfection

### 935 **PEGASYS Monotherapy**

- 936 The recommended dose of PEGASYS monotherapy for chronic hepatitis C in patients
- coinfected with HIV is 180 µg (1.0 mL vial or 0.5 mL prefilled syringe) once weekly for
- 938 48 weeks by subcutaneous administration in the abdomen or thigh.

### 939 **PEGASYS/COPEGUS Combination Therapy**

- The recommended dose when used in combination with ribavirin is PEGASYS 180 µg sc
- once weekly and COPEGUS 800 mg po daily given in two divided doses for a total of 48
- weeks, regardless of genotype.
- 943 Since COPEGUS absorption increases when administered with a meal, patients are
- advised to take COPEGUS with food.

### 945 Chronic Hepatitis B

### 946 **PEGASYS Monotherapy**

- The recommended dose of PEGASYS monotherapy for hepatitis B is 180 µg (1.0 mL
- 948 vial or 0.5 mL prefilled syringe) once weekly for 48 weeks by subcutaneous
- administration in the abdomen or thigh.

### 950 **Dose Modifications**

- 951 If severe adverse reactions or laboratory abnormalities develop during combination
- 952 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.

### 955 **PEGASYS**

- 956 General
- When dose modification is required for moderate to severe adverse reactions (clinical
- and/or laboratory), initial dose reduction to 135 µg (which is 0.75 mL for the vials or
- adjustment to the corresponding graduation mark for the syringes) is generally adequate.
- However, in some cases, dose reduction to 90 µg (which is 0.5 mL for the vials or

adjustment to the corresponding graduation mark for the syringes) may be needed.

962 Following improvement of the adverse reaction, re-escalation of the dose may be

onsidered (see WARNINGS, PRECAUTIONS, and ADVERSE REACTIONS).

### 964 Hematological

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## Table 8 PEGASYS Hematological Dose Modification Guidelines

Laboratory Values	Reduce PEGASYS Dose to:	Discontinue PEGASYS if:
ANC $\geq$ 750/mm <sup>3</sup> ANC $\leq$ 750/mm <sup>3</sup>	Maintain 180 μg Reduce to 135 μg	ANC <500/mm <sup>3</sup> , treatment should be suspended until ANC values return to more than 1000/mm <sup>3</sup>
		Reinstitute at 90 µg and monitor ANC
Platelet ≥50,000/mm <sup>3</sup>	Maintain 180 μg	Platelet count <25,000/mm <sup>3</sup>
Platelet <50,000/mm <sup>3</sup>	Reduce to 90 µg	

Psychiatric: Depression

Table 9

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

<b>Depression</b> Severity	Initial Management (4-8 weeks)		Depression		
	Dose modification	Visit schedule	Remains stable	Improves	Worsens
Mild	No change	weekly by visit	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)	1	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	Psychiatric therapy necessary		

consultation	

### 969 Renal Function

- 970 In patients with end-stage renal disease requiring hemodialysis, dose reduction to 135 μg
- 971 PEGASYS is recommended. Signs and symptoms of interferon toxicity should be closely
- 972 monitored.
- 973 Liver Function
- 974 If ALT increases are progressive despite dose reduction or accompanied by increased
- 975 bilirubin or evidence of hepatic decompensation, therapy should be immediately
- 976 discontinued.
- In chronic hepatitis C patients with progressive ALT increases above baseline values, the
- dose of PEGASYS should be reduced to 135 µg and more frequent monitoring of liver
- 979 function should be performed. After PEGASYS dose reduction or withholding, therapy
- 980 can be resumed after ALT flares subside.
- 981 In chronic hepatitis B patients with elevations in ALT (>5 x ULN), more frequent
- 982 monitoring of liver function should be performed and consideration should be given to
- 983 either reducing the dose of PEGASYS to 135 μg or temporarily discontinuing treatment.
- 984 After PEGASYS dose reduction or withholding, therapy can be resumed after ALT flares
- 985 subside.
- In patients with persistent, severe (ALT >10 times above the upper limit of normal)
- hepatitis B flares, consideration should be given to discontinuation of treatment.

### 988 COPEGUS

## Table 10 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	

<sup>\*</sup> One 200 mg tablet in the morning and two 200 mg tablets in the evening.

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Once COPEGUS has been withheld due to a laboratory abnormality or clinical 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 1200 mg).

997	Renal Impairment
998 999	COPEGUS should not be used in patients with creatinine clearance <50 mL/min (see CLINICAL PHARMACOLOGY, WARNINGS and COPEGUS Package Insert).
1000	HOW SUPPLIED
1001	Single Dose Vial
1002 1003 1004	Each PEGASYS (peginterferon alfa-2a) 180 μg single use, clear glass vial provides 1.0 mL containing 180 μg peginterferon alfa-2a for sc injection. Each package contains 1 vial (NDC 0004-0350-09).
1005	Prefilled Syringes Monthly Convenience Pack
1006 1007 1008 1009	Four prefilled syringes of PEGASYS (peginterferon alfa-2a), 180 µg single use, graduated, clear glass prefilled syringes, in a box with 4 needles and 4 alcohol swabs (NDC 0004-0352-39). Each syringe is a 0.5 mL (½ cc) volume syringe supplied with a 27-gauge, ½-inch needle with needle-stick protection device.
1010	Storage
1011 1012 1013	Store in the refrigerator at 2°C to 8°C (36°F to 46°F). Do not freeze or shake. Protect from light. Vials and prefilled syringes are for single use only. Discard any unused portion.
1014 1015	REBETRON <sup>®</sup> , REBETROL <sup>®</sup> , and INTRON <sup>®</sup> are registered trademarks of Schering Corporation.
1016	PI Revised: April 2009
1017	MEDICATION GUIDE
1018	PEGASYS®
1019	(peginterferon alfa-2a)
1020 1021 1022 1023 1024 1025	Before you start taking PEGASYS (PEG-ah-sis), alone or in combination with COPEGUS® (Co-PEG-UHS), please read this Medication Guide carefully. Read this Medication Guide each time you refill your prescription in case new information has been added and make sure the pharmacist has given you the medicine your healthcare provider prescribed for you. Reading the information in this Medication Guide does not take the place of talking with your healthcare provider.
1026 1027	If you are taking PEGASYS in combination with COPEGUS, you should also read the Medication Guide for COPEGUS (ribavirin, USP) Tablets.
1028 1029	What is the most important information I should know about PEGASYS therapy?
1030 1031 1032 1033 1034	PEGASYS, taken alone or in combination with COPEGUS, is a treatment for some people who are infected with hepatitis C virus. PEGASYS taken alone is a treatment for some people who are infected with the hepatitis B virus. However, PEGASYS and COPEGUS can have serious side effects that may cause death in rare cases. Before starting PEGASYS therapy, you should talk with your healthcare provider about the

- possible benefits and the possible side effects of treatment, to decide if either of these
- treatments is right for you. If you begin treatment you will need to see your healthcare
- provider regularly for examinations and blood tests to make sure your treatment is
- working and to check for side effects.
- The most serious possible side effects of PEGASYS taken alone or in combination with
- 1040 COPEGUS include:

### 1041 Problems with Pregnancy:

- 1042 Taking PEGASYS in combination with COPEGUS tablets can cause death, serious
- birth defects or other harm to your unborn child. Therefore, if you are pregnant or
- 1044 your partner is pregnant or plans to become pregnant, do not take
- 1045 PEGASYS/COPEGUS combination therapy. Female patients and female partners
- of male patients being treated with PEGASYS/COPEGUS combination therapy
- must not become pregnant during treatment and for 6 months after treatment has
- stopped. During this time, you must have pregnancy tests that show you are not
- pregnant. You must also use two effective forms of birth control during therapy and
- for 6 months after stopping therapy. Male patients should use a condom with
- spermicide as one of the two forms. You must use birth control even if you believe that
- you are not fertile or that your fertility is low. You should talk to your healthcare provider
- about birth control for you and your partner.
- 1054 If you are pregnant, you or your male partner must not take PEGASYS/COPEGUS
- 1055 combination therapy. If you or your partner are being treated and you become
- pregnant either during treatment or within 6 months of stopping treatment, call
- 1057 your healthcare provider right away.
- 1058 If you or a female sexual partner becomes pregnant, you should tell your healthcare
- provider. There is a Ribavirin Pregnancy Registry that collects information about
- pregnancy outcomes of female patients and female partners of male patients exposed to
- ribavirin. You or your healthcare provider are encouraged to contact the Registry at 1-
- 1062 800-593-2214.

1063

### Mental health problems and suicide:

- 1064 PEGASYS and PEGASYS/COPEGUS combination therapy may cause some patients to
- develop mood or behavioral problems. Signs of these problems include irritability
- 1066 (getting easily upset), depression (feeling low, feeling bad about yourself or feeling
- hopeless), and anxiety. Some patients may have aggressive behavior. Former drug addicts
- may fall back into drug addiction or overdosage. Some patients think about hurting or
- killing themselves or other people and some have killed (suicide) or hurt themselves or
- hurt other people. You must tell your healthcare provider if you are being treated for a
- nate of the people. For most ten your nearlifeare provider it you are being treated for a
- 1071 mental illness or have a history of mental illness, including depression and suicidal
- behavior or if you are or have ever been addicted to drugs or alcohol. Call your
- 1073 healthcare provider immediately if you develop any of these problems while on
- 1074 PEGASYS treatment.

### 1075 Heart problems:

- 1076 Some patients taking PEGASYS or PEGASYS/COPEGUS therapies may develop
- problems with their heart, including low blood pressure, fast heart rate, and very rarely,
- heart attacks. Tell your healthcare provider if you have had any heart problems in the
- 1079 past.

### 1080 Blood problems:

- Many patients taking PEGASYS have had a drop in the number of their white blood cells
- and their platelets. If the numbers of these blood cells are too low, you could be at risk for
- serious infections or bleeding.
- 1084 COPEGUS causes a decrease in the number of your red blood cells (anemia). This can be
- dangerous, especially for patients who already have heart or circulatory (cardiovascular)
- problems. If you have or have ever had any cardiovascular problems, talk with your
- healthcare provider before taking the combination of PEGASYS and COPEGUS.

### 1088 Liver problems:

- 1089 Infrequently, some patients with hepatitis C and liver scarring can develop sudden severe
- worsening (failure) of their liver disease while taking PEGASYS. Patients infected with
- both the hepatitis C virus and HIV can have an increased chance of having liver failure
- during PEGASYS treatment.
- Some patients taking PEGASYS for hepatitis B have had a rise in a blood test that
- measures liver inflammation. If you have a rise in this blood test, your liver may need to
- be watched more closely with additional blood tests.

### 1096 Infections:

- Some patients taking interferon have had serious infections. Sometimes these infections
- have been fatal. If you develop a fever that does not go away or gets higher, call your
- healthcare provider right away. Your healthcare provider will need to examine you to rule
- out your having a serious infection.

### 1101 Eye problems:

- 1102 Changes in vision such as a decrease or loss of vision (blindness) may happen in some
- patients. You should have an eye exam before you take PEGASYS. If you have eye
- problems or have had them in the past you may need eye exams while you are taking
- PEGASYS. Tell your healthcare provider or eye doctor immediately if you have changes
- in your vision.

### 1107 Body organ problems:

- 1108 Some patients may experience lung problems (such as difficulty breathing or
- pneumonia). Certain symptoms like severe stomach pain may mean that your internal
- organs are being damaged. Cases of weakness, loss of coordination and numbness due to
- stroke have been reported in patients taking PEGASYS/COPEGUS, including patients
- with few or no reported risk factors for stroke.

- 1113 Call your healthcare provider immediately if you develop any of these 1114 conditions: 1115 • You become very depressed, think about suicide or injuring/killing another 1116 1117 You have severe chest pain 1118 You have trouble breathing 1119 You have a change in your vision 1120 You become pregnant 1121 You notice unusual bleeding or bruising 1122 You have psoriasis (a skin disease) and it gets worse while taking PEGASYS 1123 You have weakness, loss of coordination, numbness or difficulty speaking
- 1124 High fever or a fever that does not go away 1125
  - You have severe stomach pain or lower back pain
- 1126 Bloody diarrhea

1131

1127 Skin rash can occur in patients taking PEGASYS. In some patients a rash 1128 can be serious. If you develop a rash with fever, blisters, or sores in your mouth, nose or eyes or conjunctivitis (red or inflamed eyes, like "pink eye"), 1129 1130 stop using PEGASYS and call your doctor right away

1132 For more information on possible side effects with PEGASYS therapy, alone or in 1133 combination with COPEGUS, please read the section on "What are the possible side 1134 effects of PEGASYS, and PEGASYS taken with COPEGUS?" in this Medication 1135 Guide. You should also read the Medication Guide for COPEGUS tablets if you are 1136 taking that medicine with PEGASYS.

#### 1137 What is PEGASYS?

- PEGASYS is a drug used to treat adults who have a lasting (chronic) infection with 1138
- 1139 hepatitis C virus or hepatitis B virus and who show signs that the virus is damaging the
- 1140 liver. Patients with hepatitis have the virus in their blood and in their liver. PEGASYS 1141 reduces the amount of hepatitis C virus in the body and helps the body's immune system
- fight the virus. The drug COPEGUS are tablets that may be taken with PEGASYS to help 1142
- fight the virus infection. Do not take COPEGUS by itself. 1143
- 1144 In some patients that have received PEGASYS treatment for approximately one year to
- 1145 treat hepatitis C, the amount of the hepatitis virus in the body was decreased to a level so
- 1146 low that it could not be measured by blood tests. After 3 months of therapy, your
- 1147 healthcare provider may ask you to have a blood test to help determine how you are
- 1148 responding to your treatment.
- 1149 It is not known if PEGASYS, used alone or in combination with COPEGUS, can cure
- 1150 hepatitis (permanently eliminate the virus) or if it can prevent liver failure or liver cancer
- 1151 that is caused by hepatitis infection.
- It is also not known if PEGASYS, alone or in combination with COPEGUS, will prevent 1152
- 1153 one infected person from infecting another person with hepatitis.

### Who should not take PEGASYS, or PEGASYS with COPEGUS?

- Do not take PEGASYS or PEGASYS/COPEGUS therapy if you:
- are pregnant, planning to get pregnant during treatment or during the 6 months after treatment or breast-feeding
- are a male patient with a female sexual partner who is pregnant or plans to become pregnant at any time while you are being treated with COPEGUS or during the 6 months after your treatment has ended
- have hepatitis caused by your immune system attacking your liver (autoimmune hepatitis)
- have unstable or severe liver disease
- had an allergic reaction to another alpha interferon or are allergic to any of the
   ingredients in PEGASYS or COPEGUS tablets
- Do not take PEGASYS, alone or in combination with COPEGUS, if you have abnormal red blood cells such as sickle-cell anemia or thalassemia major.

# lf you have ever had any of the following conditions or serious medical problems, tell your healthcare provider before you start taking PEGASYS:

- History of or current severe mental illness (such as depression or anxiety)
- History of drug or alcohol addiction or abuse
- History of heart disease or previous heart attack
- 1174 History of cancer
- Autoimmune disease (where the body's immune system attacks the body's own cells), such as psoriasis (a skin disease), systemic lupus erythematosus, rheumatoid arthritis
- 1178 Kidney problems
- Blood disorders (bleeding problems)
- 1180 Diabetes (high blood sugar)
- Problems with the thyroid gland
- Liver problems, other than hepatitis C or hepatitis B
- Colitis (an inflammation of the bowels)
- Eye problems
- 1185 Sleep problems
- 1186 HIV infection
- Organ transplant and are taking medicine that keeps your body from rejecting your transplant (suppresses your immune system)
- You should tell your healthcare provider if you are taking or planning to take other prescription or nonprescription medicines or vitamin and mineral supplements or herbal medicines.
- Also tell your healthcare provider if you are taking any of the following medicines:
- Theophylline: Your healthcare provider may need to monitor the amount of theophylline in your body and make changes to your theophylline dose.

- HIV medications called nucleoside reverse transcriptase inhibitors (abacavir, didanosine, emtricitabine, lamivudine, tenofovir, stavudine or zidovudine). Some patients developed serious liver problems including death.
- Didanosine: Do not take COPEGUS and didanosine.

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1201 If you have any questions about your health condition or about taking PEGASYS alone or in combination with COPEGUS, you should talk to your healthcare provider.

### 1203 How should I take PEGASYS, or PEGASYS with COPEGUS?

- 1204 PEGASYS is given by injection under the skin (subcutaneous injection). PEGASYS 1205 comes in two different forms (a liquid in a single use vial and a liquid in a prefilled 1206 syringe). Your healthcare provider will determine which is best for you. Your healthcare 1207 provider will also decide whether you will take PEGASYS alone or with COPEGUS. 1208 Your dose of PEGASYS is given as a single injection once per week. At some point, your 1209 healthcare provider may change your dose of PEGASYS or COPEGUS. Do not change 1210 your dose unless your healthcare provider tells you to change it. It is important that you 1211 take PEGASYS and COPEGUS exactly as your healthcare provider tells you. Once you 1212 start treatment with PEGASYS, do not switch to another brand of interferon without 1213 talking to your healthcare provider. Other interferons may not have the same effect on the
- Take your prescribed dose of PEGASYS once a week, on the same day of each week and at approximately the same time. Your total dose of COPEGUS tablets should be divided so you take it twice a day with food (breakfast and dinner). Taking half your dose of

treatment of your disease. Switching brands will also require a change in your dose.

- 1218 COPEGUS in the morning and the other half at night will keep the medicine in your body
- at a steady level. Do not take more than your prescribed dose of PEGASYS or
- 1220 COPEGUS. Be sure to read the Medication Guide for COPEGUS (ribavirin, USP)
- 1221 for complete instructions on how to take the COPEGUS tablets.
- 1222 Your healthcare provider will train you and/or the person that will be giving you the
- 1223 PEGASYS injections on the proper way to give injections. Whether you give yourself the
- injection or another person gives the injection to you, it is important that you are
- comfortable with preparing and injecting a dose of PEGASYS, and you understand the
- instructions in "How do I inject PEGASYS?" At the end of this guide there are detailed instructions on how to prepare and give yourself an injection of PEGASYS
- using the form your healthcare provider has prescribed for you.
- 1229 If you miss a dose and you remember within 2 days of when you should have taken
- 1230 PEGASYS, give yourself an injection of PEGASYS as soon as you remember. Take your
- next dose on the day you would usually take it. If **more than 2 days** have passed, ask your healthcare provider what you should do. If you miss a dose of COPEGUS, take the
- missed dose as soon as you remember during the same day. Do not take 2 doses too close
- together in time. If it is late in the day, wait until the next day and go back on schedule.
- 1235 Do not double the next dose.
- 1236 If you take more than the prescribed amount of PEGASYS, call your healthcare provider
- 1237 right away. Your healthcare provider may want to examine you and take blood for
- 1238 testing.

- You must get regular blood tests to help your healthcare provider check how the treatment is working and to check for side effects.
- 1241 What should I avoid while taking PEGASYS, or PEGASYS with COPEGUS?
- If you are pregnant do not start taking or continue taking COPEGUS in combination with PEGASYS. (See "What is the most important information I should know about PEGASYS therapy? Problems with Pregnancy".)
- Avoid becoming pregnant while taking PEGASYS, alone or in combination with COPEGUS. PEGASYS, alone or in combination with COPEGUS, may harm your unborn child (death or serious birth defects) or cause you to lose your baby (miscarry). (See "What is the most important information I should know about PEGASYS therapy? Problems with Pregnancy".)
- Do not breast-feed your baby while on PEGASYS, alone or in combination with COPEGUS.
- **Drinking alcohol**, including beer, wine and liquor. This may make your liver disease worse.
- **Taking other medicines**. Take only medicines prescribed or approved by your healthcare provider. These include prescription and nonprescription medicines and herbal supplements.
- What are the possible side effects of PEGASYS, and PEGASYS taken with COPEGUS?
- 1259 Also see "What is the most important information I should know about PEGASYS therapy?" in this Medication Guide.
- The possible serious side effects of PEGASYS and PEGASYS/COPEGUS combination therapy are:
- Mental health problems including depression and suicidal thoughts
- **Blood problems including anemia.** Anemia is a reduction in the number of red blood cells you have which can be dangerous, especially if you have heart or breathing problems. Tell your health care provider right away if you feel tired, have chest pain or shortness of breath. These may be signs of low red blood cell counts.
- 1268 Serious infections
- **Stroke.** Some patients may experience weakness, loss of coordination, and numbness due to stroke.
- Autoimmune problems: Some patients may develop a disease where the body's own immune system begins to attack itself (autoimmune disease) while on PEGASYS therapy. These diseases can include rheumatoid arthritis, systemic lupus erythematosus, psoriasis or thyroid problems. In some patients who already have an autoimmune disease, the disease may worsen while on PEGASYS therapy.
- **Heart problems:** PEGASYS may cause some patients to experience chest pain, and very rarely a heart attack. Patients who already have heart disease could be at greatest risk. Tell your healthcare provider if you have or have had a heart problem in the past.
- **Liver problems:** Some patients may develop worsening of liver function. Some of the symptoms may include stomach bloating, confusion, brown urine, and yellow eyes. Tell your healthcare provider immediately if any of these symptoms occur.

- Eye problems including changes in vision.
- **Harm to unborn children**. PEGASYS and PEGASYS/COPEGUS may cause birth defects or death of an unborn child. For more details, see "What is the most important information I should know about PEGASYS therapy?" in this Medication Guide.

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- 1287 Common, but less serious, side effects include:
- Flu-like symptoms: Most patients who take PEGASYS have flu-like symptoms that usually lessen after the first few weeks of treatment. Flu-like symptoms may include fever, chills, muscle aches, joint pain, and headaches. Taking pain and fever reducers such as acetaminophen or ibuprofen before you take PEGASYS can help with these symptoms. You can also try taking PEGASYS at night. You may be able to sleep through the symptoms.
- Extreme fatigue (tiredness): Many patients may become extremely tired while on PEGASYS therapy.
- **Upset stomach:** Nausea, taste changes, diarrhea, and loss of appetite occur commonly.
- **Blood sugar problems:** Some patients may develop a problem with the way their body controls their blood sugar and may develop diabetes.
- **Thyroid problems**: Some patients develop changes in the function of their thyroid. Symptoms of thyroid changes include the inability to concentrate, feeling cold or hot all the time, a change in your weight, and changes to your skin.
- Skin reactions: Some patients may develop redness, swelling, dry or itchy skin at the site of injection. If after several days these symptoms do not disappear, contact your health care provider. You may get a rash during therapy. If this occurs, your health care provider may recommend medicine to treat the rash.
- **Hair thinning:** Temporary hair loss is not uncommon during treatment with PEGASYS.
- 1309 Trouble sleeping
- These are not all of the side effects of PEGASYS, and PEGASYS taken with COPEGUS.
- 1311 Your healthcare provider or pharmacist can give you a more complete list. Call your
- doctor for medical advice about side effects. Call your doctor for medical advice about
- side effects. You may report side effects to FDA at 1-800-FDA-1088. You may also
- report side effects to Roche at 1-800-526-6367.
- Talk to your healthcare provider if you are worried about side effects or find them very
- bothersome.

#### 1317 General advice about prescription medicines

- 1318 Medicines are sometimes prescribed for purposes other than those listed in a Medication
- Guide. If you have any concerns or questions about PEGASYS, contact your healthcare
- provider. Do not use PEGASYS for a condition or person other than that for which it is
- prescribed. If you want to know more about PEGASYS, your healthcare provider or
- pharmacist will be able to provide you with detailed information that is written for health-
- 1323 care providers.

1324	the Medication Guide supplied with that medicine.
1326	Keep this and all drugs out of the reach of children.
1327	This Medication Guide has been approved by the US Food and Drug Administration.
1328	MG Revised: April 2009
1329 1330	Medication Guide Appendix: Instructions for Preparing and Giving a Dose with a PEGASYS® Prefilled Syringe
1331	How should I store PEGASYS Prefilled Syringes?
1332 1333 1334 1335	PEGASYS must be stored in the refrigerator at a temperature of 2°C to 8°C (36°F to 46°F). Do not leave PEGASYS outside of the refrigerator for more than 24 hours. Do not freeze PEGASYS. Keeping PEGASYS at temperatures outside the recommended range can destroy the medicine.
1336	Each PEGASYS prefilled syringe can only be used once. Discard after use.
1337	To avoid product foaming, do not shake the prefilled syringe of PEGASYS.
1338	Protect PEGASYS from light during storage.
1339	Keep this and all other medicines out of the reach of children.
1340 1341 1342 1343 1344	How do I prepare and inject PEGASYS?  You should read through all of these directions and ask your healthcare provider for help if you have any questions before trying to give yourself an injection. It is important to follow these directions carefully. Talk to your healthcare provider if you have any questions about PEGASYS.
1345 1346 1347 1348	Your healthcare provider may not want you to take all the medicine that comes in the prefilled syringe. To appropriately administer the dose that your healthcare provider tells you to take, you may have to get rid of some of the medicine before injecting the medicine.
1349 1350 1351 1352 1353 1354	If you ever switch between using prefilled syringes and vials, talk to your healthcare provider about how much PEGASYS to use. Equal volumes of liquid from the prefilled syringes and the vials DO NOT contain the same amount of PEGASYS. If you switch between prefilled syringes and vials, you will have to adjust the volume of liquid that you use to give your injection. If you do not adjust this, you could accidentally take too much or too little of your medicine.
1355 1356	If you are giving this injection to someone else, a healthcare provider must teach you how to avoid needle sticks. Being stuck by a used needle can pass diseases on to you.
1357 1358	The prefilled syringes are used for injecting PEGASYS under the surface of the skin (subcutaneous).
1359 1360	

- 1. Collect all the materials you will need before you start to give the injection: 1361 One PEGASYS prefilled syringe Monthly Convenience Pack containing an 1362 inner carton holding the PEGASYS prefilled syringe 1363 A puncture-resistant container for cleaning up when you are finished 1364 1365 1366 2. Open the convenience pack and look at the contents. Each convenience pack has everything you need for the PEGASYS injection. 1367 4 single use syringes filled with medicine (should be colorless to light 1368 1369 yellow) four 27-gauge, ½-inch needles with needle-stick protection device 1370 4 alcohol swabs 1371 3. Take the syringe out of the refrigerator. If there is foam in the solution, put it back in 1372 the refrigerator for use at a later time and use another syringe. 1373 4. Lay the syringe on a flat clean surface and wait a few minutes until it reaches room 1374 temperature. If you notice condensation water on the outside of the syringe, wait 1375 1376 another few minutes until it disappears. Wash your hands with soap and warm water to prevent infection. 1377 6. After the syringe has warmed up, pick it up by the glass barrel and look at it carefully. 1378 Do not use PEGASYS if: 1379 1380 the medicine is cloudy the medicine has particles floating in it 1381 the medicine is any color besides colorless to light yellow 1382 the expiration date has passed 1383 Glass Barrel
  - 1384

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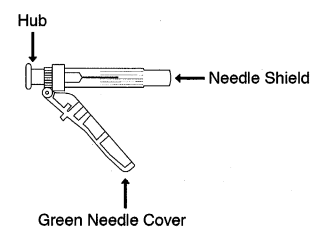
1385 7. Attachment of the needle to the PEGASYS prefilled syringe:

Plunger Rod

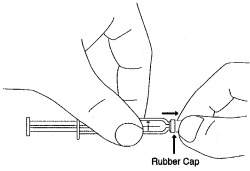
• Remove the needle from its package. Do not remove the needle shield yet. Keep the needle covered until just before you give the injection.

Stopper

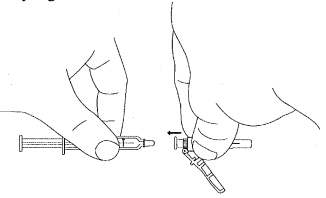
Rubber Cap



 • Remove and discard the rubber cap from the tip of the syringe barrel.



- Hold the needle close to the hub where the green needle cover connects.
- Put the needle onto the syringe by using an easy twisting motion to tighten the needle onto the syringe.

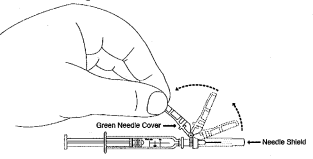


1396 1397 Here is a picture of the assembled syringe: 1398 Hub Glass Barrel Needle Shield Plunger Rod Stopper Green Needle Cover 1399 1400 Keep the syringe in a horizontal position until ready for use. 1401 If you need to set the syringe down, make sure the plastic needle shield covers 1402 the needle. Never let the needle touch any surface. 1403 8. Decide where you will give the injection. 1404 Pick a place on your stomach or thigh (see the picture below). Avoid your 1405 navel and waistline. You should use a different place each time you give 1406 yourself an injection. 1407 1408 1409 9. Prepare your skin for the injection. 1410 To minimize the discomfort from injections, you may want to gently tap the 1411 area where you plan to give yourself an injection. 1412 Clean the area using the alcohol pad. Let the skin dry for 10 seconds. 1413

1414 10. Prepare the syringe for injection.

1415 • Pull the green needle cov

Pull the green needle cover back from the needle toward the syringe barrel. The green needle cover will remain in the position you set, do not remove it. This is the needle-stick protection device.



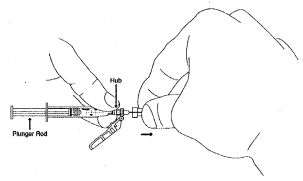
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• Hold the syringe-needle assembly tightly at the hub.

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• Remove the clear plastic needle shield covering the needle by pulling it straight off.

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11. Remove air bubbles from the syringe.

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• Hold the syringe with the needle pointing up to the ceiling.

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• If you see little bubbles, pull down slightly on the plunger rod.

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• Using your thumb and finger, gently tap the syringe to bring air bubbles to the top (small air bubbles may remain on the glass surface).

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• Press the plunger in slightly to push air bubbles out of the syringe.

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• If you find that there is still small air bubbles on glass surface after you push the air bubbles out, you can still give yourself the injection, the small air bubbles will not hurt you.

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12. Dose adjustment.

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• Your healthcare provider may not want you to take all the medicine that comes in the prefilled syringe.

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 To appropriately administer the dose that your healthcare provider tells you to take, you may have to get rid of some of the medicine before injecting the medicine.

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The syringe has markings for 180 mcg, 135 mcg, and 90 mcg. Your healthcare 1442 provider will tell you which mark to use. 1443 1444 1445 1446 1447 Once you know which mark to use, slowly and carefully press on the plunger 1448 rod of the syringe to push out medicine from the syringe. Keep pressing until the edge of the plunger stopper reaches the right mark on the side of the 1449 1450 svringe. Do not decrease or increase your dose of PEGASYS unless your healthcare 1451 1452 provider tells you to. 1453 1454 13. Give the injection of PEGASYS. Position the point of the needle (the bevel) so it is facing up. 1455 1456 1457 Pinch a fold of skin on your stomach or thigh firmly with your thumb and 1458 1459 forefinger. 1460 Hold the syringe like a pencil at a 45° to 90° angle to your skin. In one quick 1461 motion, insert the needle as far as it will go into the pinched area of skin. Pull 1462 the plunger of the syringe back very slightly. If blood comes into the syringe, 1463 1464 the needle has entered a blood vessel. Do not inject. Withdraw the needle

and discard the syringe as outlined in step 11. Repeat the above steps

with a new prefilled syringe and prepare a new site.

1467 1468 • If no blood appears, release your skin and slowly push the plunger all the way down so that you get all of your medicine.



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• Pull out the needle at same angle you put it in.

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• Wipe the area with an alcohol swab.

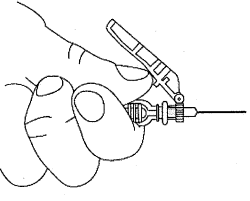
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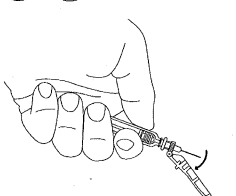
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14. For safety reasons, before you dispose of the syringe and needle, push the green needle cover toward the needle. Then place the free end of the green cap on a flat surface and push down on it until it clicks and covers over the needle. Always place used syringes and needles in a puncture-resistant container immediately after use and never reuse them. Keep your disposal container out of the reach of children.



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### How should I dispose of materials used to inject PEGASYS?

There may be special state and local laws for disposal of used needles and syringes. Your healthcare provider or pharmacist should provide you with instructions on how to properly dispose of your used syringes and needles. Always follow these instructions.

1483 The instructions below should be used as a general guide for proper disposal:

• The needles and syringes should never be reused.

- Place all used needles and syringes in a puncture-proof disposable container that is available through your pharmacy or healthcare provider (Sharp's container).
   DO NOT use glass or clear plastic containers for disposal of needles and syringes.
- Dispose of the full container as instructed by your healthcare provider or pharmacist.

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- DO NOT throw the container in your household trash. DO NOT recycle. Keep the container out of the reach of children.
- 1492 MG Appendix: Prefilled Syringe revision date: April 2009
- Medication Guide Appendix: Instructions for Preparing and Giving a Dose with a PEGASYS® Vial
- 1495 How should I store PEGASYS vials?
- 1496 PEGASYS must be stored in the refrigerator at a temperature of 2°C to 8°C (36°F to
- 1497 46°F). Do not leave PEGASYS outside of the refrigerator for more than 24 hours. Do not
- 1498 freeze PEGASYS. Keeping PEGASYS at temperatures outside the recommended range
- can destroy the medicine.
- 1500 Each PEGASYS vial can only be used once. Discard after use.
- Do not shake the vial of PEGASYS. If PEGASYS is shaken too hard, it will not work
- 1502 properly.
- 1503 Protect PEGASYS from light during storage.
- 1504 Keep this and all other medicines out of the reach of children.
- 1505 How do I inject PEGASYS?
- 1506 The following instructions will help you learn how to measure your dose and give
- yourself an injection of PEGASYS. You should read through all of these directions and ask your healthcare provider for help if you have any questions before trying to give
- yourself an injection. It is important to follow these directions carefully. Talk to your
- healthcare provider if you have any questions about PEGASYS.
- 1511 If you are giving an injection to someone else, a healthcare provider must teach you how
- to avoid needle sticks. Being stuck by a used needle can pass diseases on to you.
- 1513 1. Collect all the materials you will need before you start to give the injection:
- One vial of PEGASYS
- One syringe and needle
- Several alcohol pads
- A puncture-resistant container to dispose of the needle and syringe when you are finished
- 1519 2. Check the date on the carton the PEGASYS comes in and make sure the expiration date has not passed, then remove a vial from the package and look at the medicine.
- Do not use PEGASYS if:
- 1522 the medicine is cloudy
- 1523 the medicine has particles floating in it

1524 the medicine is any color besides colorless to light yellow 1525 the expiration date has passed 1526 3. Warm the refrigerated medicine by gently rolling it in the palms of your hands for 1527 about one minute. Do not shake. 1528 4. Wash your hands with soap and warm water to prevent infection. 5. Take the vial of PEGASYS and flip off the plastic top covering the vial opening, and 1529 1530 clean the rubber stopper on the top of the vial with a different alcohol pad. 1531 1532 If you are not sure how much medicine to use or which mark to use, STOP and call 1533 your healthcare provider right away. 1534 6. Remove the needle and syringe from their packaging and attach the needle to the end 1535 of the syringe. 1536 Pull the plunger back so the end of it is to the mark on the syringe barrel that 1537 matches the dose prescribed for you by your healthcare provider. This will pull air 1538 into the syringe barrel. 1539 Push the needle through the center of the stopper on the vial. 1540 1541 Slowly inject all the air from the syringe into the air space above the solution. Do 1542 not inject air into the fluid. 1543 1544 Keep the needle inside the vial and turn both upside down. Hold the vial and 1545 syringe straight up. Slowly pull back on the plunger until the medicine is in the 1546 syringe up to the mark that matches your dose. Make sure the needle tip always

stays in the medicine (not in the air space above it).

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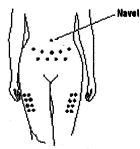


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- When the medicine is up to the right mark on the syringe barrel, take the syringe and needle out of the rubber stopper on the vial.
- Keep the syringe pointing up until you are ready to use it.
- If you need to set the syringe down, make sure that you never let the needle touch any surface.
- 7. Remove air bubbles from the syringe.
- Hold the syringe with the needle pointing up to the ceiling.
- Using your thumb and finger, tap the syringe to bring air bubbles to the top.
- Press the plunger in slightly to push air bubbles out of the syringe.
- 1558 8. Decide where you will give the injection.
  - Pick a place on your stomach or thigh (see the picture below). Avoid your navel and waistline. You should use a different place each time you give yourself an injection.



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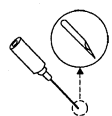
- 9. Prepare your skin for the injection.
  - To minimize the discomfort from injections, you may want to gently tap the area where you plan to give yourself an injection.
    - Clean the area using an alcohol pad. Let the skin dry for 10 seconds.

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- 10. Give the injection of PEGASYS.
- Position the point of the needle (the bevel) so it is facing up.

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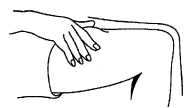


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• Pinch a fold of skin on your stomach or thigh firmly between your thumb and forefinger.

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- Hold the syringe like a pencil at a 45° to 90° angle to your skin. In one quick motion, insert the needle as far as it will go into the pinched area of skin. Pull the plunger of the syringe back very slightly. If blood comes into the syringe, the needle has entered a blood vessel. Do not inject. Withdraw the needle and discard the syringe as outlined in step 11. Repeat the above steps with a new vial and syringe and prepare a new site.
  - If no blood appears, release your skin and slowly push the plunger all the way down so that you get all of your medicine.

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- Pull out the needle at same angle you put it in. Wipe the area with an alcohol pad.
- 1587 11. For safety reasons, always place used syringes and needles in a puncture-resistant container immediately after use and never reuse them.
  - If you are using a syringe with a needle-stick protection device, before you dispose of the syringe and needle, place the free end of the green cap on a flat surface and push down on it until it clicks and covers over the needle.

#### How should I dispose of materials used to inject PEGASYS?

- There may be special state and local laws for disposal of used needles and syringes. Your healthcare provider or pharmacist should provide you with instructions on how to properly dispose of your used syringes and needles. Always follow these instructions.
- 1596 The instructions below should be used as a general guide for proper disposal:
- The needles and syringes should never be reused.
- Place all used needles and syringes in a puncture-proof disposable container that is available through your pharmacy or healthcare provider (Sharp's container).
  - DO NOT use glass or clear plastic containers for disposal of needles and syringes.
- Dispose of the full container as instructed by your healthcare provider or pharmacist.

DO NOT throw the container in your household trash. DO NOT recycle. Keep the container out of the reach of children.

1605 MG Appendix: Vial revision date: October 2008

Roche

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