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3 Product Information
4 PegIntron™
5 (Peginterferon alfa-2b)
6 Powder for Injection
7

Alpha interferons, including PegIntron™, 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. Patients with persistently severe or worsening signs or symptoms of these conditions should be withdrawn from therapy. In many but not all cases these disorders resolve after stopping PegIntron™ therapy. See WARNINGS, ADVERSE REACTIONS.

Use with Ribavirin. Ribavirin may cause birth defects and/or death of the unborn child. Extreme care must be taken to avoid pregnancy in female patients and in female partners of male patients. Ribavirin causes hemolytic anemia. The anemia associated with REBETOL therapy may result in a worsening of cardiac disease. Ribavirin is genotoxic and mutagenic and should be considered a potential carcinogen. (See REBETOL package insert for additional information and other warnings.)

8 DESCRIPTION

9 PegIntron™, peginterferon alfa-2b, Powder for Injection is a covalent conjugate of
10 recombinant alfa-2b interferon with monomethoxy polyethylene glycol (PEG). The
11 average molecular weight of the PEG portion of the molecule is 12,000 daltons. The
12 average molecular weight of the PegIntron molecule is approximately 31,000
13 daltons. The specific activity of peginterferon alfa-2b is approximately 0.7×10^8
14 IU/mg protein.

15 Interferon alfa-2b, is a water-soluble protein with a molecular weight of 19,271
16 daltons produced by recombinant DNA techniques. It is obtained from the bacterial



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17 fermentation of a strain of *Escherichia coli* bearing a genetically engineered plasmid
18 containing an interferon gene from human leukocytes.

19 **PegIntron is supplied in both vials and the Redipen® for subcutaneous use.**

20 **Vials**

21 Each vial contains either 74 mcg, 118.4 mcg, 177.6 mcg, or 222 mcg of PegIntron as
22 a white to off-white tablet-like solid, that is whole/in pieces or as a loose powder, and
23 1.11 mg dibasic sodium phosphate anhydrous, 1.11 mg monobasic sodium
24 phosphate dihydrate, 59.2 mg sucrose and 0.074 mg polysorbate 80. Following
25 reconstitution with 0.7 mL of the supplied Sterile Water for Injection, USP, each vial
26 contains PegIntron at strengths of either 50 mcg per 0.5 mL, 80 mcg per 0.5 mL, 120
27 mcg per 0.5 mL, or 150 mcg per 0.5 mL.

28 **Redipen®**

29 Redipen® is a dual-chamber glass cartridge containing lyophilized PegIntron as a
30 white to off-white tablet or powder that is whole or in pieces in the sterile active
31 chamber and a second chamber containing Sterile Water for Injection, USP. Each
32 PegIntron Redipen® contains either 67.5 mcg, 108 mcg, 162 mcg, or 202.5 mcg of
33 PegIntron, and 1.013 mg dibasic sodium phosphate anhydrous, 1.013 mg
34 monobasic sodium phosphate dihydrate, 54 mg sucrose and 0.0675 mg polysorbate
35 80. Each cartridge is reconstituted to allow for the administration of up to 0.5 mL of
36 solution. Following reconstitution, each Redipen® contains PegIntron at strengths of
37 either 50 mcg per 0.5 mL, 80 mcg per 0.5 mL, 120 mcg per 0.5 mL, or 150 mcg per
38 0.5 mL for a single use. Because a small volume of reconstituted solution is lost
39 during preparation of PegIntron, each Redipen® contains an excess amount of
40 PegIntron powder and diluent to ensure delivery of the labeled dose.

41 **CLINICAL PHARMACOLOGY**

42 **General:** The biological activity of PegIntron is derived from its interferon alfa-2b
43 moiety. Interferons exert their cellular activities by binding to specific membrane
44 receptors on the cell surface and initiate a complex sequence of intracellular events.



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45 These include the induction of certain enzymes, suppression of cell proliferation,
46 immunomodulating activities such as enhancement of the phagocytic activity of
47 macrophages and augmentation of the specific cytotoxicity of lymphocytes for target
48 cells, and inhibition of virus replication in virus-infected cells. Interferon alfa
49 upregulates the Th1 T-helper cell subset in *in vitro* studies. The clinical relevance of
50 these findings is not known.

51 **Pharmacodynamics:** PegIntron raises concentrations of effector proteins such as
52 serum neopterin and 2'5' oligoadenylate synthetase, raises body temperature, and
53 causes reversible decreases in leukocyte and platelet counts. The correlation
54 between the *in vitro* and *in vivo* pharmacologic and pharmacodynamic and clinical
55 effects is unknown.

56 **Pharmacokinetics:** Following a single subcutaneous (SC) dose of PegIntron, the
57 mean absorption half-life ($t_{1/2 k_a}$) was 4.6 hours. Maximal serum concentrations
58 (C_{max}) occur between 15-44 hours post-dose, and are sustained for up to 48-72
59 hours. The C_{max} and AUC measurements of PegIntron increase in a dose-related
60 manner. After multiple dosing, there is an increase in bioavailability of PegIntron.
61 Week 48 mean trough concentrations (320 pg/mL; range 0, 2960) are approximately
62 3-fold higher than Week 4 mean trough concentrations (94 pg/mL; range 0, 416).
63 The mean PegIntron elimination half-life is approximately 40 hours (range 22 to 60
64 hours) in patients with HCV infection. The apparent clearance of PegIntron is
65 estimated to be approximately 22.0 mL/hr·kg. Renal elimination accounts for 30% of
66 the clearance.

67 Pegylation of interferon alfa-2b produces a product (PegIntron) whose clearance is
68 lower than that of non-pegylated interferon alfa-2b. When compared to INTRON A,
69 PegIntron (1 mcg/kg) has approximately a 7-fold lower mean apparent clearance
70 and a 5-fold greater mean half-life permitting a reduced dosing frequency. At
71 effective therapeutic doses, PegIntron has approximately 10-fold greater C_{max} and
72 50-fold greater AUC than interferon alfa-2b.

73 **Special Populations**

74 **Renal Dysfunction**



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75 Following multiple dosing of PegIntron (1 mcg/kg SC given every week for four
76 weeks) the clearance of PegIntron is reduced by a mean of 17% in patients with
77 moderate renal impairment (creatinine clearance 30-49 mL/min) and by a mean of
78 44% in patients with severe renal impairment (creatinine clearance 10-29 mL/min)
79 compared to subjects with normal renal function. Clearance was similar in patients
80 with severe renal impairment not on dialysis and patients who are receiving
81 hemodialysis. The dose of PegIntron for monotherapy should be reduced in patients
82 with moderate or severe renal impairment (See **DOSAGE AND ADMINISTRATION:**
83 **DOSE REDUCTION**). REBETOL should not be used in patients with creatinine
84 clearance < 50 mL/min (See **REBETOL Package Insert, WARNINGS**).

85 **Gender**

86 During the 48 week treatment period with PegIntron, no differences in the
87 pharmacokinetic profiles were observed between male and female patients with
88 chronic hepatitis C infection.

89 **Geriatric Patients**

90 The pharmacokinetics of geriatric subjects (> 65 years of age) treated with a single
91 subcutaneous dose of 1 mcg/kg of PegIntron were similar in C_{max} , AUC, clearance,
92 or elimination half-life as compared to younger subjects (28 to 44 years of age).

93 **Effect of Food on Absorption of Ribavirin** Both AUC_{if} and C_{max} increased by
94 70% when REBETOL Capsules were administered with a high-fat meal (841 kcal,
95 53.8 g fat, 31.6 g protein, and 57.4 g carbohydrate) in a single-dose pharmacokinetic
96 study. (See **DOSAGE AND ADMINISTRATION**.)

97 **Drug Interactions**

98 **Drugs Metabolized by Cytochrome P-450**

99 The pharmacokinetics of representative drugs metabolized by CYP1A2 (caffeine),
100 CYP2C8/9 (tolbutamide), CYP2D6 (dextromethorphan), CYP3A4 (midazolam) and
101 N-acetyltransferase (dapson) were studied in 22 patients with chronic hepatitis C
102 who received PegIntron (1.5 mcg/kg) once weekly for 4 weeks. PegIntron treatment



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103 resulted in a 28% (mean) increase in a measure of CYP2C8/9 activity. PegIntron
104 treatment also resulted in a 66% (mean) increase in a measure of CYP2D6 activity;
105 however, the effect was variable as 13 patients had an increase, 5 patients had a
106 decrease, and 4 patients had no significant change (see **PRECAUTIONS: Drug**
107 **Interactions**).

108 No significant effect was observed on the pharmacokinetics of representative drugs
109 metabolized by CYP1A2, CYP3A4, or N-acetyltransferase. The effects of PegIntron
110 on CYP2C19 activity were not assessed.

111 **Methadone**

112 The pharmacokinetics of concomitant administration of methadone and PegIntron
113 were evaluated in 18 PegIntron naïve chronic hepatitis C patients receiving 1.5
114 mcg/kg/week PegIntron SC weekly. All patients were on stable methadone
115 maintenance therapy receiving ≥ 40 mg/day prior to initiating PegIntron. Mean
116 methadone AUC was approximately 16% higher after 4 weeks of PegIntron
117 treatment as compared to baseline. In 2 patients, methadone AUC was
118 approximately double after 4 weeks of PegIntron treatment as compared to baseline
119 (see **PRECAUTIONS: Drug Interactions**).

120 **Use with Ribavirin:**

121 Ribavirin has been shown *in vitro* to inhibit phosphorylation of zidovudine,
122 lamivudine, and stavudine. However, in a study with another pegylated interferon in
123 combination with ribavirin, no pharmacokinetic (e.g., plasma concentrations or
124 intracellular triphosphorylated active metabolite concentrations) or
125 pharmacodynamic (e.g., loss of HIV/HCV virologic suppression) interaction was
126 observed when ribavirin and lamivudine (n=18); stavudine (n=10); or zidovudine
127 (n=6) were co-administered as part of a multi-drug regimen to HIV/HCV co-infected
128 patients. Exposure to didanosine or its active metabolite (dideoxyadenosine 5'-
129 triphosphate) is increased when didanosine is co-administered with ribavirin, which
130 could cause or worsen clinical toxicities (see **PRECAUTIONS: Drug Interactions**).



131 **CLINICAL STUDIES**132 **PegIntron Monotherapy-Study 1**

133 A randomized study compared treatment with PegIntron (0.5, 1, or 1.5 mcg/kg once
 134 weekly SC) to treatment with INTRON A (3 million units three times weekly SC) in
 135 1219 adults with chronic hepatitis from HCV infection. The patients were not
 136 previously treated with interferon alfa, had compensated liver disease, detectable
 137 HCV RNA, elevated ALT, and liver histopathology consistent with chronic hepatitis.
 138 Patients were treated for 48 weeks and were followed for 24 weeks post-treatment.
 139 Seventy percent of all patients were infected with HCV genotype 1, and 74 percent
 140 of all patients had high baseline levels of HCV RNA (more than 2 million copies per
 141 mL of serum), two factors known to predict poor response to treatment.
 142 Response to treatment was defined as undetectable HCV RNA and normalization of
 143 ALT at 24 weeks post-treatment. The response rates to the 1 and 1.5 mcg/kg
 144 PegIntron doses were similar (approximately 24%) to each other and were both
 145 higher than the response rate to INTRON A (12%). (See **Table 1.**)

146

Table 1. Rates of Response to Treatment-Study 1

	A PegIntron 0.5 mcg/kg (N=315)	B PegIntron 1 mcg/kg (N=298)	C INTRON A 3 MIU TIW (N=307)	B - C (95% CI) Difference between PegIntron 1 mcg/kg and INTRON A
Treatment Response (Combined Virologic Response and ALT Normalization)	17%	24%	12%	11 (5, 18)
Virologic Response ^a	18%	25%	12%	12 (6,19)
ALT Normalization	24%	29%	18%	11 (5,18)

147 ^aSerum HCV is measured by a research-based quantitative polymerase chain reaction assay by a central
 148 laboratory.

149

150 Patients with both viral genotype 1 and high serum levels of HCV RNA at
 151 baseline were less likely to respond to treatment with PegIntron. Among patients
 152 with the two unfavorable prognostic variables, 8% (12/157) responded to PegIntron
 153 treatment and 2% (4/169) responded to INTRON A. Doses of PegIntron higher than
 154 the recommended dose did not result in higher response rates in these patients.



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155 Patients receiving PegIntron with viral genotype 1 had a response rate of 14%
156 (28/199) while patients with other viral genotypes had a 45% (43/96) response rate.

157 Ninety-six percent of the responders in the PegIntron groups and 100% of
158 responders in the INTRON A group first cleared their viral RNA by week 24 of
159 treatment. (See **DOSAGE AND ADMINISTRATION.**)

160 The treatment response rates were similar in men and women. Response
161 rates were lower in African American and Hispanic patients and higher in Asians
162 compared to Caucasians. Although African Americans had a higher proportion of
163 poor prognostic factors compared to Caucasians the number of non-Caucasians
164 studied (9% of the total) was insufficient to allow meaningful conclusions about
165 differences in response rates after adjusting for prognostic factors.

166

167 Liver biopsies were obtained before and after treatment in 60% of patients. A
168 modest reduction in inflammation compared to baseline that was similar in all four
169 treatment groups was observed.

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171 **PegIntron/REBETOL Combination Therapy-Study 2**

172 A randomized study compared treatment with two PegIntron/REBETOL® (ribavirin,
173 USP) regimens [PegIntron 1.5 mcg/kg SC once weekly (QW)/REBETOL 800 mg PO
174 daily (in divided doses); PegIntron 1.5 mcg/kg SC QW for 4 weeks then 0.5 mcg/kg
175 SC QW for 44 weeks/REBETOL 1000/1200 mg PO daily (in divided doses)] with
176 INTRON A [3 MIU SC thrice weekly (TIW)/REBETOL 1000/1200 mg PO daily (in
177 divided doses)] in 1530 adults with chronic hepatitis C. Interferon naïve patients
178 were treated for 48 weeks and followed for 24 weeks posttreatment. Eligible patients
179 had compensated liver disease, detectable HCV RNA, elevated ALT, and liver
180 histopathology consistent with chronic hepatitis.

181 Response to treatment was defined as undetectable HCV RNA at 24 weeks
182 posttreatment. The response rate to the PegIntron 1.5 mcg/kg plus ribavirin 800 mg
183 dose was higher than the response rate to INTRON A/REBETOL (see **Table 2**).



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184 The response rate to PegIntron 1.5→0.5 mcg/kg/REBETOL was essentially the
 185 same as the response to INTRON A/REBETOL (data not shown).

	PegIntron 1.5 mcg/kg QW REBETOL 800 mg QD	INTRON A 3 MIU TIW REBETOL 1000/1200 mg QD
Overall response ^{1,2}	52% (264/511)	46% (231/505)
Genotype 1	41% (141/348)	33% (112/343)
Genotype 2-6	75% (123/163)	73% (119/162)

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¹ Serum HCV RNA is measured with a research-based quantitative polymerase chain reaction assay by a central laboratory.

² Difference in overall treatment response (PegIntron/REBETOL vs. INTRON A/REBETOL) is 6% with 95% confidence interval of (0.18, 11.63) adjusted for viral genotype and presence of cirrhosis at baseline. Response to treatment was defined as undetectable HCV RNA at 24 weeks posttreatment.

194 Patients with viral genotype 1, regardless of viral load, had a lower response
 195 rate to PegIntron (1.5 mcg/kg)/REBETOL (800 mg) compared to patients with other
 196 viral genotypes. Patients with both poor prognostic factors (genotype 1 and high viral
 197 load) had a response rate of 30% (78/256) compared to a response rate of 29%
 198 (71/247) with INTRON A/REBETOL.

199 Patients with lower body weight tended to have higher adverse event rates
 200 (see **ADVERSE REACTIONS**) and higher response rates than patients with higher
 201 body weights. Differences in response rates between treatment arms did not
 202 substantially vary with body weight.

203 Treatment response rates with PegIntron/REBETOL were 49% in men and
 204 56% in women. Response rates were lower in African American and Hispanic
 205 patients and higher in Asians compared to Caucasians. Although African Americans
 206 had a higher proportion of poor prognostic factors compared to Caucasians, the
 207 number of non-Caucasians studied (11% of the total) was insufficient to allow
 208 meaningful conclusions about differences in response rates after adjusting for
 209 prognostic factors in this study.



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210 Liver biopsies were obtained before and after treatment in 68% of patients.
 211 Compared to baseline approximately 2/3 of patients in all treatment groups were
 212 observed to have a modest reduction in inflammation.

213 **PegIntron/REBETOL Combination Therapy-Study 3**

214 In a large United States community-based study (Study 3), 4913 patients with
 215 chronic hepatitis C were randomized to receive PegIntron 1.5 mcg/kg SC once
 216 weekly (QW) in combination with a REBETOL dose of 800-1400 mg (weight-based
 217 dosing- [WBD]) or 800 mg (Flat) PO daily (in divided doses) for 24 or 48 weeks
 218 based on genotype. Response to treatment was defined as undetectable HCV RNA
 219 (based on an assay with a lower limit of detection of 125 IU/mL) at 24 weeks
 220 posttreatment.

221 Treatment with PegIntron 1.5 mcg/kg and REBETOL 800-1400 mg resulted in a
 222 higher sustained virologic response compared to PegIntron in combination with a flat
 223 800 mg daily dose of REBETOL. Subjects weighing >105 kg obtained the greatest
 224 benefit with WBD, although a modest benefit was also observed in subjects
 225 weighing >85-105 kg (Table 3). The benefit of WBD in subjects weighing >85 kg was
 226 observed with HCV genotypes 1-3. Insufficient data were available to reach
 227 conclusions regarding other genotypes. Use of WBD resulted in an increased
 228 incidence of anemia (see ADVERSE REACTIONS and Laboratory Values).

229

230 **Table 3. SVR Rate by Treatment and Baseline Weight- Study 3**

Treatment Group	Subject Baseline Weight			
	<65 kg (<143 lb)	65-85 kg (143-188 lb)	>85-105 kg (>188-231 lb)	>105 kg (>231 lb)
WBD*	50% (173/348)	45% (449/994)	42% (351/835)	47% (138/292)
Flat	51% (173/342)	44% (443/1011)	39% (318/819)	33% (91/272)

231 * p=0.01, primary efficacy comparison (based on data from subjects weighing 65 kg or higher at
 232 baseline and utilizing a logistic regression analysis that includes treatment [WBD or Flat], genotype
 233 and presence/absence of advanced fibrosis, in the model).

234



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235 A total of 1552 subjects weighing >65 kg in Study 3 had genotype 2 or 3 and
236 were randomized to 24 or 48 weeks of therapy. No additional benefit was observed
237 with the longer treatment duration.

238 INDICATIONS AND USAGE

239 PegIntron is indicated for use alone or in combination with REBETOL for the
240 treatment of chronic hepatitis C in patients with compensated liver disease who have
241 not been previously treated with interferon alpha and are at least 18 years of age.

242

243 CONTRAINDICATIONS

244 **PegIntron is contraindicated in patients with:**

- 245 • hypersensitivity to PegIntron or any other component of the product
- 246 • autoimmune hepatitis
- 247 • hepatic decompensation (Child-Pugh score >6 [class B and C]) in cirrhotic CHC
248 patients before or during treatment

249 PegIntron/REBETOL combination therapy is additionally contraindicated in:

- 250 • patients with hypersensitivity to ribavirin or any other component of the
251 product
- 252 • women who are pregnant
- 253 • men whose female partners are pregnant
- 254 • patients with hemoglobinopathies (e.g., thalassemia major, sickle-cell
255 anemia)
- 256 • patients with creatinine clearance < 50 mL/min

257 WARNINGS

258 Patients should be monitored for the following serious conditions, some of which
259 may become life threatening. Patients with persistently severe or worsening signs or
260 symptoms should be withdrawn from therapy.



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261 **Neuropsychiatric events**

262 Life-threatening or fatal neuropsychiatric events, including suicide, suicidal and
263 homicidal ideation, depression, relapse of drug addiction/overdose, and aggressive
264 behavior sometimes directed towards others have occurred in patients with and
265 without a previous psychiatric disorder during PegIntron treatment and follow-up.
266 Psychoses, hallucinations, bipolar disorders, and mania have been observed in
267 patients treated with alpha interferons. PegIntron should be used with extreme
268 caution in patients with a history of psychiatric disorders. Patients should be advised
269 to report immediately any symptoms of depression and/or suicidal ideation to their
270 prescribing physicians. Physicians should monitor all patients for evidence of
271 depression and other psychiatric symptoms. If patients develop psychiatric
272 problems, including clinical depression, it is recommended that the patients be
273 carefully monitored during treatment and in the 6-month follow-up period. If
274 psychiatric symptoms persist or worsen, or suicidal ideation or aggressive behavior
275 towards others is identified, it is recommended that treatment with PegIntron be
276 discontinued, and the patient followed, with psychiatric intervention as appropriate. In
277 severe cases, PegIntron should be stopped immediately and psychiatric intervention
278 instituted. (See **DOSAGE AND ADMINISTRATION: Dose Reduction**). Cases of
279 encephalopathy have been observed in some patients, usually elderly, treated with
280 higher doses of PegIntron.

281 **Bone marrow toxicity**

282 PegIntron suppresses bone marrow function, sometimes resulting in severe
283 cytopenias. PegIntron should be discontinued in patients who develop severe
284 decreases in neutrophil or platelet counts (see **DOSAGE AND ADMINISTRATION:**
285 **Dose Reduction**). Ribavirin may potentiate the neutropenia induced by interferon
286 alpha. Very rarely alpha interferons may be associated with aplastic anemia.

287 **Hepatic Failure**

288 Chronic hepatitis C (CHC) patients with cirrhosis may be at risk of hepatic
289 decompensation and death when treated with alpha interferons, including PegIntron.



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290 Cirrhotic CHC patients co-infected with HIV receiving highly active antiretroviral
291 therapy (HAART) and alpha interferons with or without ribavirin appear to be at
292 increased risk for the development of hepatic decompensation compared to patients
293 not receiving HAART. During treatment, patients' clinical status and hepatic function
294 should be closely monitored, and PegIntron treatment should be immediately
295 discontinued if decompensation (Child-Pugh score >6) is observed (See
296 **CONTRAINDICATIONS**).

297 **Endocrine disorders**

298 PegIntron causes or aggravates hypothyroidism and hyperthyroidism.
299 Hyperglycemia has been observed in patients treated with PegIntron. Diabetes
300 mellitus has been observed in patients treated with alpha interferons. Patients with
301 these conditions who cannot be effectively treated by medication should not begin
302 PegIntron therapy. Patients who develop these conditions during treatment and
303 cannot be controlled with medication should not continue PegIntron therapy.

304 **Cardiovascular events**

305 Cardiovascular events, which include hypotension, arrhythmia, tachycardia,
306 cardiomyopathy, angina pectoris, and myocardial infarction, have been observed in
307 patients treated with PegIntron. PegIntron should be used cautiously in patients with
308 cardiovascular disease. Patients with a history of myocardial infarction and
309 arrhythmic disorder who require PegIntron therapy should be closely monitored (see
310 **Laboratory Tests**). Patients with a history of significant or unstable cardiac disease
311 should not be treated with PegIntron/REBETOL combination therapy. (see
312 **REBETOL package insert**.)

313 **Cerebrovascular disorders**

314 Ischemic and hemorrhagic cerebrovascular events have been observed in patients
315 treated with interferon alfa-based therapies, including PegIntron. Events occurred in
316 patients with few or no reported risk factors for stroke, including patients less than 45
317 years of age. Because these are spontaneous reports, estimates of frequency



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318 cannot be made and a causal relationship between interferon alfa-based therapies
319 and these events is difficult to establish.

320

321 **Pulmonary disorders**

322 Dyspnea, pulmonary infiltrates, pneumonia, bronchiolitis obliterans, interstitial
323 pneumonitis and sarcoidosis, some resulting in respiratory failure and/or patient
324 deaths, may be induced or aggravated by PegIntron or alpha interferon therapy.
325 Recurrence of respiratory failure has been observed with interferon rechallenge.
326 PegIntron combination treatment should be suspended in patients who develop
327 pulmonary infiltrates or pulmonary function impairment. Patients who resume
328 interferon treatment should be closely monitored.

329 **Colitis**

330 Fatal and nonfatal ulcerative or hemorrhagic/ischemic colitis have been observed
331 within 12 weeks of the start of alpha interferon treatment. Abdominal pain, bloody
332 diarrhea, and fever are the typical manifestations. PegIntron treatment should be
333 discontinued immediately in patients who develop these symptoms and signs. The
334 colitis usually resolves within 1-3 weeks of discontinuation of alpha interferons.

335 **Pancreatitis**

336 Fatal and nonfatal pancreatitis have been observed in patients treated with alpha
337 interferon. PegIntron therapy should be suspended in patients with signs and
338 symptoms suggestive of pancreatitis and discontinued in patients diagnosed with
339 pancreatitis.

340 **Autoimmune disorders**

341 Development or exacerbation of autoimmune disorders (e.g., thyroiditis, thrombotic
342 thrombocytopenic purpura, idiopathic thrombocytopenic purpura, rheumatoid
343 arthritis, interstitial nephritis, systemic lupus erythematosus, and psoriasis) have
344 been observed in patients receiving PegIntron. PegIntron should be used with
345 caution in patients with autoimmune disorders.



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346 **Ophthalmologic disorders**

347 Decrease or loss of vision, retinopathy including macular edema, retinal artery or
348 vein thrombosis, retinal hemorrhages and cotton wool spots, optic neuritis, and
349 papilledema may be induced or aggravated by treatment with peginterferon alfa-2b
350 or other alpha interferons. All patients should receive an eye examination at
351 baseline. Patients with preexisting ophthalmologic disorders (e.g., diabetic or
352 hypertensive retinopathy) should receive periodic ophthalmologic exams during
353 interferon alpha treatment. Any patient who develops ocular symptoms should
354 receive a prompt and complete eye examination. Peginterferon alfa-2b treatment
355 should be discontinued in patients who develop new or worsening ophthalmologic
356 disorders.

357 **Hypersensitivity**

358 Serious, acute hypersensitivity reactions (e.g., urticaria, angioedema,
359 bronchoconstriction, anaphylaxis) and cutaneous eruptions (Stevens Johnson
360 syndrome, toxic epidermal necrolysis) have been rarely observed during alpha
361 interferon therapy. If such a reaction develops during treatment with PegIntron,
362 discontinue treatment and institute appropriate medical therapy immediately.
363 Transient rashes do not necessitate interruption of treatment.

364 **Use with Ribavirin—(See also REBETOL Package Insert.)**

365 **REBETOL may cause birth defects and/or death of the unborn child. REBETOL**
366 **therapy should not be started until a report of a negative pregnancy test has**
367 **been obtained immediately prior to planned initiation of therapy. Patients**
368 **should use at least two forms of contraception and have monthly pregnancy**
369 **tests (see BOXED WARNING, CONTRAINDICATIONS and PRECAUTIONS:**
370 **Information for Patients and REBETOL package insert).**

371 **Anemia**

372 Ribavirin caused hemolytic anemia in 10% of PegIntron/REBETOL-treated patients
373 within 1-4 weeks of initiation of therapy. Complete blood counts should be obtained



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374 pretreatment and at week 2 and week 4 of therapy or more frequently if clinically
375 indicated. Anemia associated with REBETOL therapy may result in a worsening of
376 cardiac disease. Decrease in dosage or discontinuation of REBETOL may be
377 necessary. (See **DOSAGE AND ADMINISTRATION: Dose Reduction.**)

378 **PRECAUTIONS**

- 379 • PegIntron alone or in combination with REBETOL has not been studied in
380 patients who have failed other alpha interferon treatments.
- 381 • The safety and efficacy of PegIntron alone or in combination with REBETOL for
382 the treatment of hepatitis C in liver or other organ transplant recipients have not
383 been studied. In a small (n=16) single-center, uncontrolled case experience,
384 renal failure in renal allograft recipients receiving interferon alpha and ribavirin
385 combination therapy was more frequent than expected from the center's previous
386 experience with renal allograft recipients not receiving combination therapy. The
387 relationship of the renal failure to renal allograft rejection is not clear.
- 388 • The safety and efficacy of PegIntron/REBETOL for the treatment of patients with
389 HCV co-infected with HIV or HBV have not been established.

390 **Triglycerides:**

391 Elevated triglyceride levels have been observed in patients treated with interferon
392 alfa including PegIntron therapy. Hypertriglyceridemia may result in pancreatitis (see
393 **WARNINGS: Pancreatitis**). Elevated triglyceride levels should be managed as
394 clinically appropriate. Discontinuation of PegIntron therapy should be considered for
395 patients with symptoms of potential pancreatitis, such as abdominal pain, nausea, or
396 vomiting and persistently elevated triglycerides (e.g., triglycerides >1000 mg/dL).

397 **Patients with renal insufficiency**

398 Increases in serum creatinine levels have been observed in patients with renal
399 insufficiency receiving interferon alfa products, including PegIntron. Patients with
400 impaired renal function should be closely monitored for signs and symptoms of
401 interferon toxicity, including increases in serum creatinine, and PegIntron dosing



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402 should be adjusted accordingly or discontinued (see **CLINICAL PHARMACOLOGY:**
403 **Pharmacokinetics and DOSAGE AND ADMINISTRATION: Dose Reduction**).
404 PegIntron monotherapy should be used with caution in patients with creatinine
405 clearance < 50 mL/min; the potential risks should be weighed against the potential
406 benefits in these patients. Combination therapy with REBETOL must not be used in
407 patients with creatinine clearance < 50 mL/min (see **REBETOL Package Insert**
408 **WARNINGS**).

409 **Information for Patients:** Patients receiving PegIntron alone or in combination with
410 REBETOL should be directed in its appropriate use, informed of the benefits and
411 risks associated with treatment, and referred to the MEDICATION GUIDES for
412 PegIntron and, if applicable, REBETOL.

413 Patients must be informed that REBETOL may cause birth defects and/or
414 death of the unborn child. Extreme care must be taken to avoid pregnancy in female
415 patients and in female partners of male patients during treatment with combination
416 PegIntron/REBETOL therapy and for 6 months post-therapy. Combination
417 PegIntron/REBETOL therapy should not be initiated until a report of a negative
418 pregnancy test has been obtained immediately prior to initiation of therapy. It is
419 recommended that patients undergo monthly pregnancy tests during therapy and for
420 6 months post-therapy. (See **CONTRAINdicATIONS and REBETOL package**
421 **insert**.)

422 Patients should be informed that there are no data regarding whether
423 PegIntron therapy will prevent transmission of HCV infection to others. Also, it is not
424 known if treatment with PegIntron will cure hepatitis C or prevent cirrhosis, liver
425 failure, or liver cancer that may be the result of infection with the hepatitis C virus.

426 Patients should be advised that laboratory evaluations are required before
427 starting therapy and periodically thereafter (see **Laboratory Tests**). It is advised that
428 patients be well hydrated, especially during the initial stages of treatment. "Flu-like"
429 symptoms associated with administration of PegIntron may be minimized by bedtime
430 administration of PegIntron or by use of antipyretics.



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431 Patients should be advised to use a puncture-resistant container for the
432 disposal of used syringes, needles, and the Redipen®. The full container should be
433 disposed of in accordance with state and local laws. Patients should be thoroughly
434 instructed in the importance of proper disposal. Patients should also be cautioned
435 against reusing or sharing needles, syringes, or the Redipen®.

436 **Dental and periodontal disorders:** Dental and periodontal disorders have been
437 reported in patients receiving PegIntron/REBETOL combination therapy. In addition,
438 dry mouth could have a damaging effect on teeth and mucous membranes of the
439 mouth during long-term treatment with the combination of REBETOL and PegIntron.
440 Patients should brush their teeth thoroughly twice daily and have regular dental
441 examinations. If vomiting occurs, patients should be advised to rinse out their mouth
442 thoroughly afterwards.

443 **Laboratory Tests:** PegIntron alone or in combination with ribavirin may cause
444 severe decreases in neutrophil and platelet counts, and hematologic, endocrine
445 (e.g., TSH), and hepatic abnormalities. Transient elevations in ALT (2-5 fold above
446 baseline) were observed in 10% of patients treated with PegIntron, and was not
447 associated with deterioration of other liver functions. Triglyceride levels are
448 frequently elevated in patients receiving alpha interferon therapy including PegIntron
449 and should be periodically monitored.

450 Patients on PegIntron or PegIntron/REBETOL combination therapy should
451 have hematology and blood chemistry testing before the start of treatment and then
452 periodically thereafter. In the clinical trial CBC (including hemoglobin, neutrophil and
453 platelet counts) and chemistries (including AST, ALT, bilirubin, and uric acid) were
454 measured during the treatment period at weeks 2, 4, 8, 12, and then at 6-week
455 intervals or more frequently if abnormalities developed. TSH levels were measured
456 every 12 weeks during the treatment period. HCV RNA should be measured at 6
457 months of treatment. PegIntron or PegIntron/REBETOL combination therapy should
458 be discontinued in patients with persistent high viral levels.



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459 Patients who have pre-existing cardiac abnormalities should have
460 electrocardiograms administered before treatment with PegIntron/REBETOL.

461 **Drug Interactions**

462 Caution should be used when administering PegIntron with medications metabolized
463 by CYP2C8/9 (e.g., warfarin and phenytoin) or CYP2D6 (e.g., flecainide) (see
464 **CLINICAL PHARMACOLOGY; Drug Interactions**).

465 **Methadone**

466 In a pharmacokinetic study of 18 chronic hepatitis C patients concomitantly receiving
467 methadone, treatment with PegIntron once weekly for 4 weeks was associated with
468 a mean increase of 16% in methadone AUC; in 2 out of 18 patients, methadone
469 AUC doubled (see **CLINICAL PHARMACOLOGY: Drug Interactions**). The clinical
470 significance of this finding is unknown; however, patients should be monitored for
471 the signs and symptoms of increased narcotic effect.

472 **Use with Ribavirin:**

473 **Nucleoside Analogues**

474 Hepatic decompensation (some fatal) has occurred in cirrhotic HIV/HCV co-infected
475 patients receiving combination antiretroviral therapy for HIV and interferon alfa and
476 ribavirin. Adding treatment with alfa interferons alone or in combination with ribavirin
477 may increase the risk in this patient subset. Patients receiving interferon with
478 ribavirin and Nucleoside Reverse Transcriptase Inhibitors (NRTIs) should be closely
479 monitored for treatment-associated toxicities, especially hepatic decompensation
480 and anemia. Discontinuation of NRTIs should be considered as medically
481 appropriate (see **Individual NRTI Product Information**). Dose reduction or
482 discontinuation of interferon, ribavirin or both should also be considered if worsening
483 clinical toxicities are observed, including hepatic decompensation (e.g., Child-Pugh >
484 6).

485 **Stavudine, Lamivudine, and Zidovudine:** *In vitro* studies have shown ribavirin can
486 reduce the phosphorylation of pyrimidine nucleoside analogues such as stavudine,



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487 lamivudine, and zidovudine. In a study with another pegylated interferon alfa, no
488 evidence of a pharmacokinetic or pharmacodynamic (e.g., loss of HIV/HCV virologic
489 suppression) interaction was seen when ribavirin was co-administered with
490 zidovudine, lamivudine, or stavudine in HIV/HCV co-infected patients (see
491 **CLINICAL PHARMACOLOGY: Drug Interactions**).

492 Although there was no evidence of loss of HIV/HCV virologic suppression
493 when ribavirin was co-administered with zidovudine, HIV/HCV co-infected patients
494 who were administered zidovudine in combination with pegylated interferon alfa and
495 ribavirin developed severe neutropenia (ANC <500) and severe anemia (hemoglobin
496 <8 g/dL) more frequently than similar patients not receiving zidovudine.

497 **Didanosine:** Co-administration of REBETOL Capsules or Oral Solution and
498 didanosine is not recommended. Reports of fatal hepatic failure, as well as
499 peripheral neuropathy, pancreatitis, and symptomatic hyperlactatemia/lactic
500 acidosis have been reported in clinical trials (see **CLINICAL PHARMACOLOGY:**
501 **Drug Interactions**).

502 **Carcinogenesis, Mutagenesis, and Impairment of Fertility**

503 **Carcinogenesis and Mutagenesis:** PegIntron has not been tested for its
504 carcinogenic potential. Neither PegIntron, nor its components interferon or
505 methoxypolyethylene glycol caused damage to DNA when tested in the standard
506 battery of mutagenesis assays, in the presence and absence of metabolic activation.

507 **Use with Ribavirin:** Ribavirin is genotoxic and mutagenic and should be considered
508 a potential carcinogen. See REBETOL package insert for additional warnings
509 relevant to PegIntron therapy in combination with ribavirin.

510 **Impairment of Fertility:** PegIntron may impair human fertility. Irregular menstrual
511 cycles were observed in female cynomolgus monkeys given subcutaneous injections
512 of 4239 mcg/m² PegIntron alone every other day for 1 month (approximately 345
513 times the recommended weekly human dose based upon body surface area). These
514 effects included transiently decreased serum levels of estradiol and progesterone,



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515 suggestive of anovulation. Normal menstrual cycles and serum hormone levels
516 resumed in these animals 2 to 3 months following cessation of PegIntron treatment.
517 Every other day dosing with 262 mcg/m² (approximately 21 times the weekly human
518 dose) had no effects on cycle duration or reproductive hormone status. The effects
519 of PegIntron on male fertility have not been studied.

520 **Pregnancy Category C: PegIntron monotherapy:** Non-pegylated interferon alfa-
521 2b, has been shown to have abortifacient effects in *Macaca mulatta* (rhesus
522 monkeys) at 15 and 30 million IU/kg (estimated human equivalent of 5 and 10 million
523 IU/kg, based on body surface area adjustment for a 60 kg adult). PegIntron should
524 be assumed to also have abortifacient potential. There are no adequate and well-
525 controlled studies in pregnant women. PegIntron therapy is to be used during
526 pregnancy only if the potential benefit justifies the potential risk to the fetus.
527 Therefore, PegIntron is recommended for use in fertile women only when they are
528 using effective contraception during the treatment period.

529 **Pregnancy Category X: Use with Ribavirin**

530 **Significant teratogenic and/or embryocidal effects have been demonstrated in**
531 **all animal species exposed to ribavirin. REBETOL therapy is contraindicated**
532 **in women who are pregnant and in the male partners of women who are**
533 **pregnant. (See CONTRAINDICATIONS and the REBETOL Package Insert.)**

534 **Ribavirin Pregnancy Registry: A Ribavirin Pregnancy Registry has been**
535 **established to monitor maternal-fetal outcomes of pregnancies in female**
536 **patients and female partners of male patients exposed to ribavirin during**
537 **treatment and for 6 months following cessation of treatment. Physicians and**
538 **patients are encouraged to report such cases by calling 1-800-593-2214.**

539 **Nursing Mothers:** It is not known whether the components of PegIntron and/or
540 REBETOL are excreted in human milk. Studies in mice have shown that mouse
541 interferons are excreted in breast milk. Because of the potential for adverse
542 reactions from the drug in nursing infants, a decision must be made whether to



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543 discontinue nursing or discontinue the PegIntron and REBETOL treatment, taking
544 into account the importance of the therapy to the mother.

545 **Pediatric:** Safety and effectiveness in pediatric patients below the age of 18 years
546 have not been established.

547 **Geriatric:** In general, younger patients tend to respond better than older patients to
548 interferon-based therapies. Clinical studies of PegIntron alone or in combination with
549 REBETOL did not include sufficient numbers of subjects aged 65 and over,
550 however, to determine whether they respond differently than younger subjects.
551 Treatment with alpha interferons, including PegIntron, is associated with
552 neuropsychiatric, cardiac, pulmonary, GI and systemic (flu-like) adverse effects.
553 Because these adverse reactions may be more severe in the elderly, caution should
554 be exercised in the use of PegIntron in this population. This drug is known to be
555 substantially excreted by the kidney. Because elderly patients are more likely to
556 have decreased renal function, the risk of toxic reactions to this drug may be greater
557 in patients with impaired renal function (see **CLINICAL PHARMACOLOGY: Special**
558 **Populations: Renal dysfunction**). REBETOL should not be used in patients with
559 creatinine clearance <50 mL/min. When using PegIntron/REBETOL therapy, refer
560 also to the REBETOL Package Insert.

561 **ADVERSE REACTIONS**

562 Study 1 compared PegIntron monotherapy with INTRON A monotherapy.
563 Study 2 compared combination therapy of PegIntron/REBETOL with combination
564 therapy with INTRON A/REBETOL. In these studies, nearly all study patients in
565 clinical trials experienced one or more adverse events.

566 Adverse events that occurred in Studies 1 and 2 at >5% incidence are provided in
567 **Table 4** by treatment group. Due to potential differences in ascertainment
568 procedures, adverse event rate comparisons across studies should not be made.

569 **Table 4. Adverse Events Occurring in > 5% of Patients**



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Adverse Events	Percentage of Patients Reporting Adverse Events*			
	Study 1		Study 2	
	PegIntron1 mcg/kg (n=297)	INTRON A 3 MIU (n=303)	PegIntron 1.5 mcg/kg/ REBETOL (n=511)	INTRON A/ REBETOL (n=505)
Application Site				
Injection Site Inflammation/Reaction	47	20	75	49
Autonomic Nervous Sys.				
Mouth Dry	6	7	12	8
Sweating Increased	6	7	11	7
Flushing	6	3	4	3
Body as a Whole				
Fatigue/Asthenia	52	54	66	63
Headache	56	52	62	58
Rigors	23	19	48	41
Fever	22	12	46	33
Weight Decrease	11	13	29	20
RUQ Pain	8	8	12	6
Chest Pain	6	4	8	7
Malaise	7	6	4	6
Central/Periph. Nerv. Sys.				
Dizziness	12	10	21	17
Endocrine				
Hypothyroidism	5	3	5	4
Gastrointestinal				
Nausea	26	20	43	33
Anorexia	20	17	32	27
Diarrhea	18	16	22	17
Vomiting	7	6	14	12
Abdominal Pain	15	11	13	13
Dyspepsia	6	7	9	8
Constipation	1	3	5	5
Hematologic Disorders				
Neutropenia	6	2	26	14
Anemia	0	0	12	17
Leukopenia	<1	0	6	5
Thrombocytopenia	7	<1	5	2
Liver and Biliary System				
Hepatomegaly	6	5	4	4
Musculoskeletal				
Myalgia	54	53	56	50
Arthralgia	23	27	34	28
Musculoskeletal Pain	28	22	21	19
Psychiatric				



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Adverse Events	Percentage of Patients Reporting Adverse Events*			
	Study 1		Study 2	
	PegIntron1 mcg/kg (n=297)	INTRON A 3 MIU (n=303)	PegIntron 1.5 mcg/kg/ REBETOL (n=511)	INTRON A/ REBETOL (n=505)
Insomnia	23	23	40	41
Depression	29	25	31	34
Anxiety/Emotional Lability/Irritability	28	34	47	47
Concentration Impaired	10	8	17	21
Agitation	2	2	8	5
Nervousness	4	3	6	6
Reproductive, Female				
Menstrual Disorder	4	3	7	6
Resistance Mechanism				
Infection Viral	11	10	12	12
Infection Fungal	<1	3	6	1
Respiratory System				
Dyspnea	4	2	26	24
Coughing	8	5	23	16
Pharyngitis	10	7	12	13
Rhinitis	2	2	8	6
Sinusitis	7	7	6	5
Skin and Appendages				
Alopecia	22	22	36	32
Pruritus	12	8	29	28
Rash	6	7	24	23
Skin Dry	11	9	24	23
Special Senses, Other				
Taste Perversion	<1	2	9	4
Vision Disorders				
Vision Blurred	2	3	5	6
Conjunctivitis	4	2	4	5

570 *Patients reporting one or more adverse events. A patient may have reported more than one
571 adverse event within a body system/organ class category.

572 The adverse event profile in Study 3, which compared PegIntron/weight-
573 based REBETOL combination to a PegIntron/flat dose REBETOL regimen, revealed
574 an increased rate of anemia with weight-based dosing (29% vs. 19% for weight-
575 based vs. flat dose regimens, respectively). However, the majority of cases of
576 anemia were mild and responded to dose reductions.



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577 The incidence of serious adverse events was comparable in all studies. In the
578 PEG monotherapy trial (Study 1) the incidence of serious adverse events was
579 similar—(about 12%) in all treatment groups. In Study 2, the incidence of serious
580 adverse events was 17% in the PegIntron/REBETOL groups compared to 14% in
581 the INTRON A/REBETOL group. In Study 3, there was a similar incidence of serious
582 adverse events reported for the weight-based REBETOL group (12%) and with the
583 flat dose REBETOL regimen.

584 In many but not all cases, adverse events resolved after dose reduction or
585 discontinuation of therapy. Some patients experienced ongoing or new serious
586 adverse events during the 6-month follow-up period. There have been 19 patient
587 deaths which occurred during treatment or during follow-up in these clinical trials. In
588 Study 1, there was one suicide in a patient receiving PegIntron monotherapy and
589 two deaths among patients receiving INTRON A monotherapy (1 murder/suicide and
590 1 sudden death). In Study 2, there was one suicide in a patient receiving
591 PegIntron/REBETOL combination therapy; and 1 patient death in the INTRON
592 A/REBETOL group (motor vehicle accident). In Study 3, there were 14 deaths, 2 of
593 which were probable suicides and one was an unexplained death in a person with a
594 relevant medical history of depression.

595 In Studies 1 and 2, 10-14% of patients receiving PegIntron, alone or in
596 combination with REBETOL, discontinued therapy compared with 6% treated with
597 INTRON A alone and 13% treated with INTRON A in combination with REBETOL.
598 Similarly in Study 3, 15% of patients receiving PegIntron in combination with weight-
599 based REBETOL and 14% of patients receiving PegIntron and flat dose REBETOL
600 discontinued therapy due to an adverse event. The most common reasons for
601 discontinuation of therapy were related to known interferon effects of psychiatric,
602 systemic (e.g., fatigue, headache), or gastrointestinal adverse events.

603 In Study 2, dose reductions due to adverse reactions occurred in 42% of
604 patients receiving PegIntron (1.5 mcg/kg)/REBETOL and in 34% of those receiving
605 INTRON A/REBETOL. The majority of patients (57%) weighing 60 kg or less
606 receiving PegIntron (1.5 mcg/kg)/REBETOL required dose reduction. Reduction of



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607 interferon was dose related (PegIntron 1.5 mcg/kg > PegIntron 0.5 mcg/kg or
608 INTRON A), 40%, 27%, 28%, respectively. Dose reduction for REBETOL was
609 similar across all three groups, 33-35%. The most common reasons for dose
610 modifications were neutropenia (18%), or anemia (9%) (see **Laboratory Values**).
611 Other common reasons included depression, fatigue, nausea, and
612 thrombocytopenia. In Study 3, dose modifications due to adverse events occurred
613 more frequently with WBD compared to flat dosing (29 and 23%, respectively).

614 In the PegIntron/REBETOL combination trials the most common adverse
615 events were psychiatric which occurred among 77% of patients in Study 2 and 68-
616 69% of patients in Study 3. These psychiatric adverse events included most
617 commonly depression, irritability, and insomnia, each reported by approximately 30-
618 40% of subjects in all treatment groups. Suicidal behavior (ideation, attempts, and
619 suicides) occurred in 2% of all patients during treatment or during follow-up after
620 treatment cessation (see **WARNINGS**).

621 PegIntron induced fatigue or headache in approximately two-thirds of
622 patients, with fever or rigors in approximately half of the patients. The severity of
623 some of these systemic symptoms (e.g., fever and headache) tends to decrease as
624 treatment continues. In Studies 1 and 2, application site inflammation and reaction
625 (e.g., bruise, itchiness, and irritation) occurred at approximately twice the incidence
626 with PegIntron therapies (in up to 75% of patients) compared with INTRON A.
627 However, injection site pain was infrequent (2-3%) in all groups. In Study 3 there
628 was a 23-24% incidence overall for injection site reactions or inflammation.

629 In Study 2, many patients continued to experience adverse events several
630 months after discontinuation of therapy. By the end of the 6-month follow-up period,
631 the incidence of ongoing adverse events by body class in the PegIntron
632 1.5/REBETOL group was 33% (psychiatric), 20% (musculoskeletal), and 10% (for
633 endocrine and for GI). In approximately 10-15% of patients weight loss, fatigue, and
634 headache had not resolved.

635 Individual serious adverse events occurred at a frequency $\leq 1\%$ and included
636 suicide attempt, suicidal ideation, severe depression; psychosis, aggressive



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637 reaction, relapse of drug addiction/overdose; nerve palsy (facial, oculomotor);
638 cardiomyopathy, myocardial infarction, angina, pericardial effusion, retinal ischemia,
639 retinal artery or vein thrombosis, blindness, decreased visual acuity, optic neuritis,
640 transient ischemic attack, supraventricular arrhythmias, loss of consciousness;
641 neutropenia, infection (sepsis, pneumonia, abscess, cellulitis); emphysema,
642 bronchiolitis obliterans, pleural effusion, gastroenteritis, pancreatitis, gout,
643 hyperglycemia, hyperthyroidism and hypothyroidism, autoimmune thrombocytopenia
644 with or without purpura, rheumatoid arthritis, interstitial nephritis, lupus-like
645 syndrome, sarcoidosis, aggravated psoriasis; urticaria, injection-site necrosis,
646 vasculitis, and phototoxicity.

647 **Laboratory Values**

648 Changes in selected laboratory values during treatment with PegIntron alone or in
649 combination with REBETOL treatment are described below. **Decreases in**
650 **hemoglobin, neutrophils, and platelets may require dose reduction or**
651 **permanent discontinuation from therapy. (See DOSAGE AND**
652 **ADMINISTRATION: Dose Reduction.)**

653 **Hemoglobin.** Hemoglobin levels decreased to <11 g/dL in about 30% of patients in
654 Study 2. In Study 3, 47% of patients receiving WBD REBETOL and 33% on flat dose
655 REBETOL had decreases in hemoglobin levels <11 g/dl. Reductions in hemoglobin
656 to <9 g/dL occurred more frequently in patients receiving WBD compared to flat
657 dosing (4% and 2%, respectively). In Study 2, dose modification was required in 9%
658 and 13% of patients in the PegIntron/REBETOL and INTRON A/REBETOL groups.
659 Hemoglobin levels become stable by treatment week 4-6 on average. The typical
660 pattern observed was a decrease in hemoglobin levels by treatment week 4 followed
661 by stabilization and a plateau, which was maintained to the end of treatment. In the
662 PegIntron monotherapy trial hemoglobin decreases were generally mild and dose
663 modifications were rarely necessary. (See **DOSAGE AND ADMINISTRATION:**
664 **Dose Reduction.**)

665 **Neutrophils.** Decreases in neutrophil counts were observed in a majority of patients
666 treated with PegIntron alone (70%) or as combination therapy with REBETOL in



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667 Study 2 (85%) and INTRON A/REBETOL (60%). Severe potentially life-threatening
668 neutropenia ($<0.5 \times 10^9/L$) occurred in 1% of patients treated with PegIntron
669 monotherapy, 2% of patients treated with INTRON A/REBETOL and in
670 approximately 4% of patients treated with PegIntron/REBETOL. Two percent of
671 patients receiving PegIntron monotherapy and 18% of patients receiving
672 PegIntron/REBETOL in Study 2 required modification of interferon dosage. Few
673 patients ($< 1\%$) required permanent discontinuation of treatment. Neutrophil counts
674 generally return to pre-treatment levels 4 weeks after cessation of therapy. (See
675 **DOSAGE AND ADMINISTRATION: Dose Reduction.**)

676 **Platelets.** Platelet counts decreased to $<100,000/mm^3$ in approximately 20% of
677 patients treated with PegIntron alone or with REBETOL and in 6% of patients treated
678 with INTRON A/REBETOL. Severe decreases in platelet counts ($<50,000/mm^3$)
679 occur in $<4\%$ of patients. Patients may require discontinuation or dose modification
680 as a result of platelet decreases. (See **DOSAGE AND ADMINISTRATION: Dose**
681 **Reduction.**) In Study 2, 1% or 3% of patients required dose modification of INTRON
682 A or PegIntron, respectively. Platelet counts generally returned to pretreatment
683 levels 4 weeks after the cessation of therapy.

684 **Triglycerides.** Elevated triglyceride levels have been observed in patients treated
685 with interferon alphas including PegIntron.

686 **Thyroid Function.** Development of TSH abnormalities, with and without clinical
687 manifestations, are associated with interferon therapies. In Study 2, clinically
688 apparent thyroid disorders occur among patients treated with either INTRON A or
689 PegIntron (with or without REBETOL) at a similar incidence (5% for hypothyroidism
690 and 3% for hyperthyroidism). Subjects developed new onset TSH abnormalities
691 while on treatment and during the follow-up period. At the end of the follow-up period
692 7% of subjects still had abnormal TSH values.

693 **Bilirubin and uric acid.** In Study 2, 10-14% of patients developed
694 hyperbilirubinemia and 33-38% developed hyperuricemia in association with
695 hemolysis. Six patients developed mild to moderate gout.



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696 **Postmarketing Experience**

697 The following adverse reactions have been identified and reported during post-
698 approval use of PegIntron therapy: aphthous stomatitis, erythema multiforme,
699 hearing impairment, hearing loss, memory loss, migraine headache, myositis,
700 peripheral neuropathy, renal insufficiency, renal failure, rhabdomyolysis, seizures,
701 Stevens Johnson syndrome, thrombotic thrombocytopenic purpura, toxic epidermal
702 necrolysis, vertigo, and pure red cell aplasia. Because the reports of these reactions
703 are voluntary and the population of uncertain size, it is not always possible to reliably
704 estimate the frequency of the reaction or establish a causal relationship to drug
705 exposure.

706 **Immunogenicity:** Approximately 2% of patients receiving PegIntron (32/1759) or
707 INTRON A (11/728) with or without REBETOL developed low-titer (≤ 160)
708 neutralizing antibodies to PegIntron or INTRON A. The clinical and pathological
709 significance of the appearance of serum neutralizing antibodies is unknown. No
710 apparent correlation of antibody development to clinical response or adverse events
711 was observed. The incidence of posttreatment binding antibody ranged from 8 to 15
712 percent. The data reflect the percentage of patients whose test results were
713 considered positive for antibodies to PegIntron in a Biacore assay that is used to
714 measure binding antibodies, and in an antiviral neutralization assay, which
715 measures serum-neutralizing antibodies. The percentage of patients whose test
716 results were considered positive for antibodies is highly dependent on the sensitivity
717 and specificity of the assays. Additionally the observed incidence of antibody
718 positivity in these assays may be influenced by several factors including sample
719 timing and handling, concomitant medications, and underlying disease. For these
720 reasons, comparison of the incidence of antibodies to PegIntron with the incidence
721 of antibodies to other products may be misleading.

722 **OVERDOSAGE**

723 There is limited experience with overdosage. In the clinical studies, a few patients
724 accidentally received a dose greater than that prescribed. There were no instances



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725 in which a participant in the monotherapy or combination therapy trials received
 726 more than 10.5 times the intended dose of PegIntron. The maximum dose received
 727 by any patient was 3.45 mcg/kg weekly over a period of approximately 12 weeks.
 728 The maximum known overdosage of REBETOL was an intentional ingestion of 10 g
 729 (fifty 200-mg capsules). There were no serious reactions attributed to these
 730 overdosages. In cases of overdosing, symptomatic treatment and close observation
 731 of the patient are recommended.

732 **DOSAGE AND ADMINISTRATION**

733 There are no safety and efficacy data on treatment for longer than 1 year. A patient
 734 should self-inject PegIntron only if it has been determined that it is appropriate and
 735 the patient agrees to medical follow-up as necessary and training in proper injection
 736 technique has been given to him/her.

737 It is recommended that patients receiving PegIntron, alone or in combination
 738 with ribavirin, be discontinued from therapy if HCV viral levels remain high after 6
 739 months of therapy.

740 **PegIntron Monotherapy**

741 The recommended dose of PegIntron regimen is 1 mcg/kg/week subcutaneously for
 742 1 year. The dose should be administered on the same day of the week.

743 The volume of PegIntron to be injected depends on patient weight (see **Table 5**
 744 below).

745 **Table 5. Recommended PegIntron Monotherapy Dosing**

Body weight kg (lbs)	PegIntron Redipen® or Vial Strength to Use	Amount of PegIntron (mcg) To Administer	Volume (mL) of PegIntron to Administer
≤45 (≤100)	50 mcg per 0.5 mL	40	0.4
46 – 56 (101 – 124)		50	0.5



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57 - 72 (125 - 159)	80 mcg per 0.5 mL	64	0.4
73 - 88 (160 - 195)		80	0.5
89 - 106 (196 - 234)	120 mcg per 0.5 mL	96	0.4
107 - 136 (235 - 300)		120	0.5
137 - 160 (301 - 353)	150 mcg per 0.5 mL	150	0.5

*When reconstituted as directed

746
747

748 PegIntron/REBETOL Combination Therapy

749 The recommended dose of PegIntron is 1.5 mcg/kg/week in combination with 800-
750 1400 mg of REBETOL based on patient body weight. The volume of PegIntron to be
751 injected depends on the strength of PegIntron and patient's body weight. (See **Table**
752 **6.**)

753 The treatment duration for patients with genotype 1 is 48 weeks. Patients with
754 genotype 2 and 3 should be treated for 24 weeks.

755

756 **TABLE 6. Recommended PegIntron Combination Therapy Dosing**

757

Body weight kg (lbs)	PegIntron Redipen® or Vial Strength to Use	Amount of PegIntron (mcg) To Administer	Volume (mL)*of PegIntron to Administer	REBETOL Daily Dose	REBETOL Number of Capsules
<40 (<87)	50 mcg per 0.5 mL	50	0.5	800 mg/day	2 x 200 mg capsules A.M. 2 x 200 mg capsules P.M.
40-50 (88-111)	80 mcg per 0.5 mL	64	0.4	800 mg/day	2 x 200 mg capsules A.M. 2 x 200 mg capsules P.M.



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51-60 (112-133)		80	0.5	800 mg/day	2 x 200 mg capsules A.M. 2 x 200 mg capsules P.M.
61-65 (134-144)	120 mcg per 0.5 mL	96	0.4	800 mg/day	2 x 200 mg capsules A.M. 2 x 200 mg capsules P.M.
66-75 (145-166)		96	0.4	1000 mg/day	2 x 200 mg capsules A.M. 3 x 200 mg capsules P.M.
76-85 (167-188)		120	0.5	1000 mg/day	2 x 200 mg capsules A.M. 3 x 200 mg capsules P.M.
86-105 (189-231)	150 mcg per 0.5 mL	150	0.5	1200 mg/day	3 x 200 mg capsules A.M. 3 x 200 mg capsules P.M.
>105 (>231)				1400 mg/day	3 x 200 mg capsules A.M. 4 x 200 mg capsules P.M.

758 * When reconstituted as directed.

759

760 REBETOL should be taken with food. REBETOL should not be used in patients with
761 creatinine clearance <50 mL/min.

762 Dose Reduction

763 If a serious adverse reaction develops during the course of treatment (see
764 **WARNINGS**) discontinue or modify the dosage of PegIntron and/or REBETOL until
765 the adverse event abates or decreases in severity. If persistent or recurrent serious
766 adverse events develop despite adequate dosage adjustment, discontinue
767 treatment. For guidelines for dose modifications and discontinuation based on
768 laboratory parameters, see **Tables 7** and **8**. Dose reduction of PegIntron may be
769 accomplished by utilizing a lower dose strength as shown in **Table 9** or **10**. For vials,
770 50% dose reduction may also be accomplished by reducing the volume
771 administered by one-half without changing the dose strength.

772 In the combination therapy trial dose reductions occurred among 42% of patients
773 receiving PegIntron 1.5 mcg/kg/REBETOL 800 mg daily including 57% of those
774 patients weighing 60 kg or less (see **ADVERSE REACTIONS**).

775 Table 7. Guidelines for Modification or Discontinuation of PegIntron or 776 PegIntron/REBETOL and for Scheduling Visits for Patients with Depression

Depression Severity ¹	Initial Management (4-8 wks)			Depression	
	Dose modification	Visit schedule	Remains stable	Improves	Worsens



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Mild	No change	Evaluate once weekly by visit and/or phone.	Continue weekly visit schedule.	Resume normal visit schedule.	(See moderate or severe depression)
Moderate	Decrease IFN dose 50%	Evaluate once weekly (office visit at least every other week).	Consider psychiatric consultation. Continue reduced dosing.	If symptoms improve and are stable for 4 wks, may resume normal visit schedule. Continue reduced dosing or return to normal dose.	(See severe depression)
Severe	Discontinue IFN/R permanently.	Obtain immediate psychiatric consultation.	Psychiatric therapy as necessary		

⁷⁷⁷
⁷⁷⁸ See DSM-IV for definitions.

⁷⁷⁹
⁷⁸⁰ **Table 8. Guidelines for Dose Modification and Discontinuation of PegIntron or PegIntron/REBETOL for Hematologic Toxicity**

Laboratory Values	PegIntron	REBETOL
Hgb* <10.0 g/dL <8.5 g/dL	----- Permanently discontinue	Decrease by 200 mg/day Permanently discontinue
WBC <1.5 x10 ⁹ /L <1.0 x10 ⁹ /L	Reduce dose by 50% Permanently discontinue	----- Permanently discontinue
Neutrophil <0.75 x10 ⁹ /L <0.5 x10 ⁹ /L	Reduce dose by 50% Permanently discontinue	----- Permanently discontinue
Platelets <80 x10 ⁹ /L <50 x10 ⁹ /L	Reduce dose by 50% Permanently discontinue	----- Permanently discontinue

⁷⁸¹
⁷⁸² * For patients with a history of stable cardiac disease receiving PegIntron in combination with ribavirin,
⁷⁸³ the PegIntron dose should be reduced by half and the ribavirin dose by 200 mg/day if a > 2 g/dL decrease in
⁷⁸⁴ hemoglobin is observed during any 4-week period. Both PegIntron and ribavirin should be permanently
⁷⁸⁵ discontinued if patients have hemoglobin levels <12 g/dL after this ribavirin dose reduction.

⁷⁸⁶
⁷⁸⁷ **Table 9. Reduced PegIntron Dose (0.5 mcg/kg) for (1 mcg/kg) Monotherapy**

Body weight kg (lbs)	PegIntron Redipen [®] /Vial Strength to Use	Amount of PegIntron(mcg) To Administer	Volume (mL)** of PegIntron to Administer
-------------------------	--	--	---



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≤45 (≤100)	50 mcg per 0.5 mL*	20	0.2
46 - 56 (101-124)		25	0.25
57 - 72 (125-159)	50 mcg per 0.5 mL	30	0.3
73 - 88 (160-195)		40	0.4
89-106 (196-234)	50 mcg per 0.5 mL	50	0.5
107-136 (235-300)	80 mcg per 0.5 mL	64	0.4
137-160 (301-353)		80	0.5

788
789
790

* Must use vial. Minimum delivery for Redipen® 0.3 mL

** When reconstituted as directed



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791 **TABLE 10. Reduced PegIntron Dose (0.75 mcg/kg) for (1.5 mcg/kg) Combination**
 792 **Therapy**

793

Body weight (kg)(lbs)	PegIntron Redipen®/Vial Strength to Use	Amount of PegIntron(mcg) to Administer	Volume (mL)** of PegIntron to Administer
<40 (<87)	50 mcg per 0.5 mL*	25	0.25
40-50 (88-111)	50 mcg per 0.5 mL	30	0.3
51-60 (112-133)		40	0.4
61-75 (134-166)	50 mcg per 0.5 mL	50	0.5
76-85 (167-188)	80 mcg per 0.5 mL	64	0.4
>85 (>188)		80	0.5

794 * Must use vial. Minimum delivery for Redipen® 0.3 mL

795 ** When reconstituted as directed

796 **Renal Function**

797 In patients with moderate renal dysfunction (creatinine clearance 30-50 mL/min), the
 798 PegIntron dose should be reduced by 25%. Patients with severe renal dysfunction
 799 (creatinine clearance 10-29 mL/min) including those on hemodialysis, should have
 800 the PegIntron dose reduced by 50%. If renal function decreases during treatment,
 801 PegIntron therapy should be discontinued. When PegIntron is administered in
 802 combination with REBETOL, subjects with impaired renal function and/or those over
 803 the age of 50 should be more carefully monitored with respect to the development of
 804 anemia.

805 **Preparation and Administration**806 **PegIntron Redipen®**

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807 PegIntron Redipen® consists of a dual-chamber glass cartridge with sterile,
808 lyophilized peginterferon alfa-2b in the active chamber and Sterile Water for
809 Injection, USP in the diluent chamber. The PegIntron in the glass cartridge should
810 appear as a white to off-white tablet-shaped solid that is whole or in pieces, or
811 powder. To reconstitute the lyophilized peginterferon alfa-2b in the Redipen®, hold
812 the Redipen® upright (dose button down) and press the two halves of the pen
813 together until there is an audible click. Gently invert the pen to mix the solution. **DO**
814 **NOT SHAKE**. The reconstituted solution has a concentration of either 50 mcg per
815 0.5 mL, 80 mcg per 0.5 mL, 120 mcg per 0.5 mL, or 150 mcg per 0.5 mL for a single
816 subcutaneous injection. Visually inspect the solution for particulate matter and
817 discoloration prior to administration. The reconstituted solution should be clear and
818 colorless. Do not use the solution if it is discolored, or not clear, or if particulates are
819 present.

820 Keeping the pen upright, attach the supplied needle and select the
821 appropriate PegIntron dose by pulling back on the dosing button until the dark bands
822 are visible and turning the button until the dark band is aligned with the correct dose.
823 The prepared PegIntron solution is to be injected subcutaneously.

824 The PegIntron Redipen® is a single-use pen and does not contain a
825 preservative. The reconstituted solution should be used immediately and cannot be
826 stored for more than 24 hours at 2°-8° C (see **Storage**). **DO NOT REUSE THE**
827 **REDIPEN®**. The sterility of any remaining product can no longer be guaranteed.
828 **DISCARD THE UNUSED PORTION.** Pooling of unused portions of some
829 medications has been linked to bacterial contamination and morbidity.

830 **PegIntron Vials**

831 Two B-D® Safety Lok™ syringes are provided in the package; one syringe is for the
832 reconstitution steps and one for the patient injection. There is a plastic safety sleeve
833 to be pulled over the needle after use. The syringe locks with an audible click when
834 the green stripe on the safety sleeve covers the red stripe on the needle. Instructions
835 for the preparation and administration of PegIntron Powder for Injection are provided
836 below.



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837 **Reconstitute the PegIntron lyophilized product with only 0.7 mL of 1.25**
838 **mL of supplied diluent (Sterile Water for Injection, USP). The diluent vial is for**
839 **single use only. The remaining diluent should be discarded.** No other
840 medications should be added to solutions containing PegIntron, and PegIntron
841 should not be reconstituted with other diluents. Swirl gently to hasten complete
842 dissolution of the powder. The reconstituted solution should be clear and colorless.
843 Visually inspect the solution for particulate matter and discoloration prior to
844 administration. The solution should not be used if discolored or cloudy, or if
845 particulates are present.

846 The appropriate PegIntron dose should be withdrawn and injected
847 subcutaneously. PegIntron vials are for single use only and do not contain a
848 preservative. The reconstituted solution should be used immediately and cannot be
849 stored for more than 24 hours at 2°-8° C (see **Storage**). **DO NOT REUSE THE**
850 **VIAL.** The sterility of any remaining product can no longer be guaranteed. **DISCARD**
851 **THE UNUSED PORTION.** Pooling of unused portions of some medications has
852 been linked to bacterial contamination and morbidity.

853 After preparation and administration of the PegIntron for injection, it is
854 essential to follow the state and/or local procedures for proper disposal of syringes,
855 needles, and the Redipen®. A puncture-resistant container should be used for
856 disposal. Patients should be instructed in how to properly dispose of used syringes,
857 needles, or the Redipen® and be cautioned against the reuse of these items.

858 **Storage**

859 **PegIntron Redipen®**

860 PegIntron Redipen® should be stored at 2° - 8°C (36°-46°F).

861 After reconstitution, the solution should be used immediately, but may be stored up
862 to 24 hours at 2° - 8°C (36° - 46°F). The reconstituted solution contains no
863 preservative, and is clear and colorless. **DO NOT FREEZE.**



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864 **PegIntron Vials**

865 PegIntron should be stored at 25°C (77°F); excursions permitted to 15°-30°C (59°-
 866 86°F) [see USP Controlled Room Temperature]. After reconstitution with supplied
 867 Diluent the solution should be used immediately, but may be stored up to 24 hours
 868 at 2°-8°C (36°-46°F). The reconstituted solution contains no preservative, is clear
 869 and colorless. **DO NOT FREEZE.**

870 **HOW SUPPLIED**871 **PegIntron Redipen®**

Each PegIntron Redipen® Package Contains:	
A box containing one 50 mcg per 0.5 mL PegIntron Redipen® and 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1323-01)
A box containing one 80 mcg per 0.5 mL PegIntron Redipen® and 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1316-01)
A box containing one 120 mcg per 0.5 mL PegIntron Redipen® and 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1297-01)
A box containing one 150 mcg per 0.5 mL PegIntron Redipen® and 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1370-01)

872

Each PegIntron Redipen® PAK 4 Contains:	
A box containing four 50 mcg per 0.5 mL PegIntron Redipen® Units, each containing 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1323-02)
A box containing four 80 mcg per 0.5 mL PegIntron Redipen® Units, each containing 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1316-02)
A box containing four 120 mcg per 0.5 mL PegIntron Redipen® Units, each containing 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1297-02)
A box containing four 150 mcg per 0.5 mL PegIntron Redipen® Units, each containing 1 B-D® needle and 2 alcohol swabs.	(NDC 0085-1370-02)

873

874 **PegIntron Vials**

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Each PegIntron Package Contains:	
A box containing one 50 mcg per 0.5 mL vial of PegIntron Powder for Injection and one 1.25 mL vial of Diluent (Sterile Water for Injection, USP), 2 B-D® Safety Lok™ syringes with a safety sleeve and 2 alcohol swabs.	(NDC 0085-1368-01)
A box containing one 80 mcg per 0.5 mL vial of PegIntron Powder for Injection and one 1.25 mL vial of Diluent (Sterile Water for Injection, USP), 2 B-D® Safety Lok™ syringes with a safety sleeve and 2 alcohol swabs.	(NDC 0085-1291-01)
A box containing one 120 mcg per 0.5 mL vial of PegIntron Powder for Injection and one 1.25 mL vial of Diluent (Sterile Water for Injection, USP), 2 B-D® Safety Lok™ syringes with a safety sleeve and 2 alcohol swabs.	(NDC 0085-1304-01)
A box containing one 150 mcg per 0.5 mL vial of PegIntron Powder for Injection and one 1.25 mL vial of Diluent (Sterile Water for Injection, USP), 2 B-D® Safety Lok™ syringes with a safety sleeve and 2 alcohol swabs.	(NDC 0085-1279-01)

875 Schering Corporation

876 Kenilworth, NJ 07033 USA

877 **REV. 3/24/08**

878 **XXXXXXXXXT** U.S. Patent Nos. 5,908,621; 5,951,974; 6,042,822; 6,177,074;

879 6,180,096; 6,250,469; 6,482,613; 6,524,570; and 6,610,830.

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881 B-D is a registered trademark of Becton Dickinson and Company.

882 Safety-Lok is a trademark of Becton Dickinson and Company.



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1 032408.final draft Med Guide powder for injection

2 **MEDICATION GUIDE**

3 **PegIntron®**

4 **Peginterferon alfa-2b**

5
6 **Including appendix with instructions for using PegIntron® Powder for Injection**

7
8 Read this Medication Guide carefully before you start taking PegIntron (**Peg In-tron**) or
9 PegIntron/REBETOL (**REB-eh-tole**) combination therapy. Read the Medication Guide each
10 time you refill your prescription because there may be new information. The information in
11 this Medication Guide does not take the place of talking with your health care provider
12 (doctor, nurse, nurse practitioner, or physician's assistant).

13
14 **If you are taking PegIntron/REBETOL combination therapy, also read the Medication**
15 **Guide for REBETOL (ribavirin, USP) Capsules.**

16
17 **What is the most important information I should know about PegIntron and**
18 **PegIntron/REBETOL combination therapy?**

19
20 PegIntron (peginterferon) is a treatment for some people who are infected with hepatitis C
21 virus. However, PegIntron and PegIntron/REBETOL combination therapy can have serious
22 side effects that may cause death in rare cases. Before you decide to start treatment, you
23 should talk to your health care provider about the possible benefits and side effects of
24 PegIntron or PegIntron/REBETOL combination therapy. If you begin treatment you will
25 need to see your health care provider regularly for medical examinations and lab tests to
26 make sure your treatment is working and to check for side effects.

27
28 **REBETOL capsules may cause birth defects and/or death of an unborn child. If you**
29 **are pregnant, you or your male partner must not take PegIntron/REBETOL**
30 **combination therapy. You must not become pregnant while either you or your partner**
31 **are being treated with the combination PegIntron/REBETOL therapy, or for 6 months**
32 **after stopping therapy. Men and women should use birth control while taking the**
33 **combination therapy and for 6 months afterwards. If you or your partner are being**
34 **treated and you become pregnant either during treatment or within 6 months of**
35 **stopping treatment, call your health care provider right away. There is a Ribavirin**
36 **Pregnancy Registry that collects information about pregnancy outcomes in female**
37 **patients and female partners of male patients exposed to ribavirin. You or your**
38 **healthcare provider are encouraged to contact the Registry at 1-800-593-2214.**

39
40 **If you are taking PegIntron or PegIntron/REBETOL therapy you should call your**
41 **health care provider immediately if you develop any of these symptoms:**

42
43 **New or worsening mental health problems such as thoughts about killing or hurting**
44 **yourself or others, trouble breathing, chest pain, severe stomach or lower back pain,**
45 **bloody diarrhea or bloody bowel movements, high fever, bruising, bleeding, or**
46 **decreased vision.**



47
48 The most serious possible side effects of PegIntron and PegIntron/REBETOL therapy
49 include:

50
51 **Problems with Pregnancy.** Combination PegIntron/REBETOL therapy can cause
52 death, serious birth defects, or other harm to your unborn child. If you are a woman of
53 childbearing age you must not become pregnant during treatment and for 6 months
54 after you have stopped therapy. You must have a negative pregnancy test immediately
55 before beginning treatment, during treatment and for 6 months after you have stopped
56 therapy. Both male and female patients must use effective forms of birth control during
57 treatment and for the 6 months after treatment is completed. Male patients should use
58 a condom. If you are a female, you must use birth control even if you believe that you are
59 not fertile or that your fertility is low. You should talk to your health care provider about
60 birth control for you and your partner.

61
62 **Mental health problems and suicide.** PegIntron and PegIntron/REBETOL therapies may
63 cause patients to develop mood or behavioral problems. These can include irritability
64 (getting easily upset) and depression (feeling low, feeling bad about yourself, or feeling
65 hopeless). Some patients may have aggressive behavior. Former drug addicts may fall back
66 into drug addiction or overdose. Some patients think about hurting or killing themselves or
67 other people and some have killed (suicide) or hurt themselves or others. You must tell your
68 health care provider if you are being treated for a mental illness or had treatment in the past
69 for any mental illness, including depression and suicidal behavior. You should tell your
70 health care provider if you have ever been addicted to drugs or alcohol.

71
72 **Heart problems.** Some patients taking PegIntron or PegIntron/REBETOL therapy may
73 develop problems with their heart, including low blood pressure, fast heart rate, and very
74 rarely, heart attacks. Tell your health care provider if you have had any heart problems in the
75 past.

76
77 **Blood problems.** PegIntron and PegIntron/REBETOL therapies commonly lower two types
78 of blood cells (white blood cells and platelets). In some patients, these blood counts may fall
79 to dangerously low levels. If your blood counts become very low, this could lead to
80 infections or bleeding.

81
82 REBETOL therapy causes a decrease in the number of red blood cells you have (anemia).
83 This can be dangerous, especially for patients who already have heart or circulatory
84 (cardiovascular) problems. Talk with your health care provider before taking combination
85 PegIntron/REBETOL therapy if you have or have ever had any cardiovascular problems.

86
87 **Body organ problems.** Certain symptoms like severe stomach pain may mean that your
88 internal organs are being damaged. Cases of weakness, loss of coordination, and numbness
89 due to stroke have been reported in patients taking PegIntron, including patients with few or
90 no reported risk factors for stroke.



93 *For other possible side effects, see "What are the possible side effects of PegIntron and*
 94 *PegIntron/REBETOL" in this Medication Guide.*

95

96 **What is PegIntron and PegIntron/REBETOL combination therapy?**

97 The PegIntron product is a drug used to treat adults who have a lasting (chronic) infection
 98 with hepatitis C virus and who show signs that the virus is damaging the liver.

99 PegIntron/REBETOL combination therapy consists of two medications also used to treat
 100 hepatitis C infection. Patients with hepatitis C have the virus in their blood and in their liver.
 101 PegIntron reduces the amount of virus in the body and helps the body's immune system fight
 102 the virus. REBETOL (ribavirin) is a drug that helps to fight the viral infection but does not
 103 work when used by itself to treat chronic hepatitis C.

104

105 It is not known if PegIntron or PegIntron/REBETOL therapies can cure hepatitis C
 106 (permanently eliminate the virus) or if it can prevent liver failure or liver cancer that is
 107 caused by hepatitis C infection.

108

109 It is also not known if PegIntron or PegIntron/REBETOL combination therapy will prevent
 110 one infected person from infecting another person with hepatitis C.

111

112 **Who should not take PegIntron or PegIntron/REBETOL therapy?**

113 Do not take PegIntron or PegIntron/REBETOL therapy if you:

114 • are pregnant, planning to get pregnant during treatment or during the 6 months after
 115 treatment, or breast-feeding

116

117 • are a male patient with a female sexual partner who is pregnant or plans to become
 118 pregnant at any time while you are being treated with REBETOL or during the 6
 119 months after your treatment has ended

120

121 • have hepatitis caused by your immune system attacking your liver (autoimmune
 122 hepatitis) or unstable liver disease

123

124 • had an allergic reaction to another alpha interferon or are allergic to any of the
 125 ingredients in PegIntron or REBETOL Capsules. If you have any doubts, ask your
 126 health care provider

127

128 • Do not take PegIntron/REBETOL combination therapy if you have abnormal red
 129 blood cells such as sickle-cell anemia or thalassemia major

130

131 **If you have any of the following conditions or serious medical problems, discuss them**
 132 **with your health care provider before taking PegIntron or PegIntron/REBETOL**
 133 **therapy:**

134 • depression or anxiety

135 • sleep problems

136 • high blood pressure

137 • previous heart attack, or other heart problems

138 • liver problems (other than hepatitis C infection)



- 139 • any kind of autoimmune disease (where the body's immune system attacks the body's
- 140 own cells), such as psoriasis, systemic lupus erythematosus, rheumatoid arthritis
- 141 • thyroid problems
- 142 • diabetes
- 143 • colitis (inflammation of the bowels)
- 144 • cancer
- 145 • hepatitis B infection
- 146 • HIV infection
- 147 • kidney problems
- 148 • bleeding problems
- 149 • alcoholism
- 150 • drug abuse or addiction
- 151 • body organ transplant and are taking medicine that keeps your body from rejecting your
- 152 transplant (suppresses your immune system)

153

154

155 **How should I take PegIntron or PegIntron/REBETOL?**

156 Your health care provider will decide whether you will take PegIntron therapy alone or the
 157 combination of PegIntron/REBETOL, as well as the correct dose (based on your weight).
 158 PegIntron and PegIntron/REBETOL are given for one year. Take your prescribed dose of
 159 PegIntron once a week, on the same day of each week and at approximately the same time.
 160 Take the medicine for the full year and do not take more than the prescribed dose.
 161 REBETOL Capsules should be taken with food. When you take REBETOL with food, more
 162 of the medicine (70% more on average) is taken up by your body. You should take
 163 REBETOL the same way every day (twice a day with food) to keep the medicine in your
 164 body at a steady level. This will help your health care provider to decide how your treatment
 165 is working and how to change the number of REBETOL capsules you take if you have side
 166 effects from REBETOL. **Be sure to read the Medication Guide for REBETOL**
 167 **(ribavirin, USP) for complete instructions on how to take the REBETOL capsules.**

168

169 You should be completely comfortable with how to prepare PegIntron; how to set the dose
 170 you take, and how to inject yourself before you use PegIntron for the first time. PegIntron
 171 comes in two different forms, a powder in a single-use vial and a Redipen[®] single-use
 172 delivery system. See the attached appendix for detailed instructions for preparing and giving
 173 a dose of PegIntron.

174

175 If you miss a dose of the PegIntron product, take the missed dose as soon as possible during
 176 the same day or the next day, then continue on your regular dosing schedule. If several days
 177 go by after you miss a dose, check with your health care provider about what to do. Do not
 178 double the next dose or take more than one dose a week without talking to your health care
 179 provider. Call your health care provider right away if you take more than your prescribed
 180 PegIntron dose. Your health care provider may wish to examine you more closely, and take
 181 blood for testing.

182



183 If you miss a dose of REBETOL capsules, take the missed dose as soon as possible during
 184 the same day. If an entire day has gone by, check with your health care provider about what
 185 to do. Do not double the next dose.

186

187 You must get regular blood tests to help your health care provider check how the treatment is
 188 working and to check for side effects.

189

190 Tell your health care provider if you are taking or planning to take other prescription or non-
 191 prescription medicines, including vitamin and mineral supplements and herbal medicines.

192

193 **What should I avoid while taking PegIntron or PegIntron/REBETOL therapies?**

194 • If you are pregnant do not start taking PegIntron/REBETOL combination therapy.

195 • Avoid becoming pregnant while taking PegIntron or PegIntron/REBETOL.

196 PegIntron and PegIntron/REBETOL may harm your unborn child (death or serious birth
 197 defects) or cause you to lose your baby (miscarry). **If you or your partner become**
 198 **pregnant during treatment or during the 6 months after treatment with**
 199 **PegIntron/REBETOL combination therapy, immediately report the pregnancy to your**
 200 **health care provider. You or your health care provider should call 1-800-593-2214.** By
 201 calling this number, information about you and/or your partner will be added to a pregnancy
 202 registry that will be used to help you and your health care provider make decisions about
 203 your treatment for hepatitis in the future. You, your partner and/or your health care provider
 204 will be asked to provide follow-up information on the outcome of the pregnancy.

205

206 • Do not breast-feed your baby while taking PegIntron.

207

208 **What are the possible side effects of PegIntron and PegIntron/REBETOL combination**
 209 **therapy?**

210

211 **Possible, serious side effects include:**

212 • **Mental health problems including suicide, blood problems, heart problems, body**
 213 **organ problems.** See "What is the most important information I should know about
 214 PegIntron and PegIntron/REBETOL combination therapy?"

215

216 • **Other body organ problems.** A few patients have lung problems (such as
 217 pneumonia or inflammation of the lung tissue), inflammation of the kidney, and eye
 218 disorders.

219

220 • **New or worsening autoimmune disease.** Some patients taking PegIntron or
 221 PegIntron/REBETOL develop autoimmune diseases (a condition where the body's
 222 immune cells attack other cells or organs in the body), including rheumatoid arthritis,
 223 systemic lupus erythematosus, and psoriasis. In some patients who already have an
 224 autoimmune disease, the disease worsens on PegIntron and PegIntron/REBETOL
 225 combination therapy.

226

227 **Common but less serious side effects include:**

228

- 229 • **Flu-like symptoms.** Most patients who take PegIntron or PegIntron/REBETOL
 230 therapy have "flu-like" symptoms (headache, muscle aches, tiredness and fever).
 231 Some of these symptoms (fever, headache) usually lessen after the first few weeks of
 232 therapy. You can reduce some of these symptoms by injecting your PegIntron dose at
 233 bedtime. Over-the-counter pain and fever reducers, such as acetaminophen or
 234 ibuprofen, can be used to prevent or reduce the fever and headache.
 235
- 236 • **Extreme fatigue (tiredness).** Many patients become extremely tired while on
 237 PegIntron or PegIntron/REBETOL combination therapy.
 238
- 239 • **Appetite problems.** Nausea, loss of appetite, and weight loss occur commonly.
 240
- 241 • **Thyroid problems.** Some patients develop changes in the function of their thyroid.
 242 Symptoms of thyroid changes include the inability to concentrate, feeling cold or hot
 243 all the time, a change in your weight and changes to your skin.
 244
- 245 • **Blood sugar problems.** Some patients develop problems with the way their body
 246 controls their blood sugar and may develop high blood sugar or diabetes.
 247
- 248 • **Skin reactions.** Redness, swelling, and itching are common at the site of injection.
 249 If after several days these symptoms do not disappear contact your health care
 250 provider. You may get a rash during therapy. If this occurs, your health care
 251 provider may recommend medicine to treat the rash.
 252
- 253 • **Hair thinning.** Hair thinning is common during PegIntron and PegIntron/REBETOL
 254 treatment. Hair loss stops and hair growth returns after therapy is stopped.
 255

256 These are not all of the side effects of PegIntron or PegIntron/REBETOL combination
 257 therapy. Your health care provider or pharmacist can give you a more complete list.
 258

259 **Call your doctor for medical advice about side effects. You may report side effects to FDA at**
 260 **1-800-FDA-1088.**
 261

262 **General advice about prescription medicines:**

263 Medicines are sometimes prescribed for purposes other than those listed in a Medication
 264 Guide. If you have any concerns about PegIntron, ask your health care provider. Your health
 265 care provider or pharmacist can give you information about PegIntron that was written for
 266 health care professionals. Do not use PegIntron for a condition for which it was not
 267 prescribed. Do not share this medication with other people.
 268

269 **If you are taking PegIntron/REBETOL combination therapy, also read the Medication**
 270 **Guide for REBETOL (ribavirin, USP) Capsules.**
 271

272 *This Medication Guide has been approved by the U.S. Food and Drug Administration.*

273 **Revised May 2007**
 274



275 **How do I prepare and inject the PegIntron Dose?**

276 Before you inject PegIntron, the powder must be mixed with **0.7 mL** of the supplied
 277 DILUENT for PegIntron, Sterile Water for Injection (diluent). You should carefully
 278 follow the directions given to you by your health care provider.

279
 280 The vial of mixed PegIntron should be used immediately. **DO NOT** prepare more than one
 281 vial at a time. If you don't use the vial of the prepared solution right away, it must be
 282 stored in a refrigerator and used within 24 hours.

283
 284 **Storing PegIntron**

285 PegIntron Powder should be stored at room temperature (25 ° C, 77°F); avoid exposure to
 286 heat. After mixing, the PegIntron solution should be used immediately but may be stored
 287 in the refrigerator up to 24 hours. The solution contains no preservatives. **DO NOT**
 288 FREEZE.

289 **Preparing the PegIntron solution:**

- 290
 291 1. Find a clean, well-lit, non-slip flat working surface and assemble all of the supplies
 292 you will need for an injection. All of the supplies you will need for an injection are
 293 in the PegIntron Powder for Injection package. The package contains:
 294 ■ a vial of PegIntron powder
 295
 296 ■ a 1.25 mL vial of DILUENT
 297
 298 ■ 2 disposable syringes, and
 299
 300 ■ alcohol swabs
 301
 302 2. Check the date printed on the PegIntron carton to make sure that the expiration date
 303 has not passed. Remove one vial and look at the contents. The PegIntron in the vial
 304 should appear as a white to off-white tablet-like solid that is whole/in pieces or as a
 305 loose powder.

306
 307 If you have already mixed the PegIntron solution and it has been stored properly in the
 308 refrigerator, take it out of the refrigerator and allow the solution to come to room
 309 temperature.

- 310
 311 3. Wash your hands thoroughly with soap and water, rinse and towel dry. It is important
 312 to keep your work area, your hands and injection site clean to minimize the risk of
 313 infection.

314
 315 The disposable syringes have needles that are already attached and cannot be removed.
 316 Each syringe has a clear plastic safety sleeve that is pulled over the needle for disposal
 317 after use. The safety sleeve should remain tight against the flange while using the
 318 syringe and moved over the needle only when ready for disposal. **Figure A.**

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The syringes and needles are for single use only.

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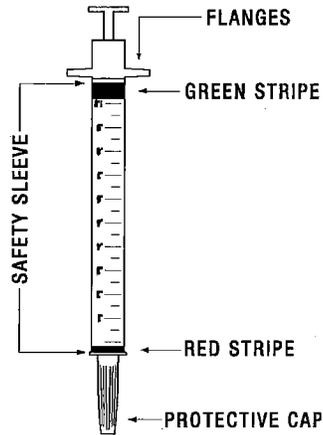


Figure A

4. Remove the protective wrapper from ONE of the syringes provided and use for the following steps 5-7. Make sure that the syringe safety sleeve is sitting against the flange. (see **Figure A**).
5. Remove the protective plastic cap from the tops of both the supplied DILUENT and the PegIntron vials. Clean the rubber stopper on the top of both vials with an alcohol swab.
6. Carefully remove the protective cap straight off of the needle to avoid damaging the needle point. Fill the syringe with air by pulling the plunger to 0.7 mL (**Figure B**). Hold the DILUENT vial upright. Do not touch the cleaned top of the vial with your hands (**Figure C**). Insert the needle through the center of the rubber stopper of the DILUENT vial, and inject the air from the syringe into the vial (**Figure D**). Turn the vial upside down and make sure the tip of the needle is in the liquid. Withdraw only 0.7 mL of DILUENT by pulling the plunger back to the 0.7 mL mark on the side of the syringe (**Figure E**). Remove the needle from the vial (**Figure F**). **Discard the remaining DILUENT.**

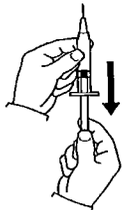


Figure B

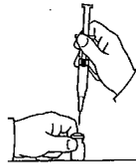


Figure C

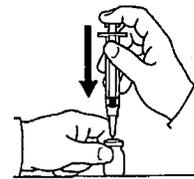


Figure D

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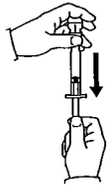


Figure E

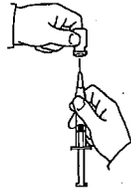


Figure F

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7. Insert the needle through the center of the rubber stopper of the PegIntron vial, and place the needle tip against the glass wall of the vial (Figure G). SLOWLY inject the 0.7 mL DILUENT so that the stream of DILUENT runs down the side of the vial. To prevent bubbles from forming, DO NOT AIM THE STREAM of diluent directly on the tablet-like SOLID or POWDER in the bottom of the vial. Remove the needle from the vial.

Firmly grasp the safety sleeve and pull it over the exposed needle until you hear a click. The green stripe on the safety sleeve will completely cover the red stripe on the needle. (See Figure O in the section: "Injecting the PegIntron dose.") Discard the syringe and needle in the puncture proof container.

8. GENTLY swirl the vial in a gentle circular motion (Figure H), until the PegIntron is completely dissolved. DO NOT SHAKE the vial. If any powder remains undissolved in the vial, gently turn the vial upside down until all of the powder is dissolved. It is not unusual for the solution to appear cloudy or bubbly for a few minutes. If air bubbles do form, wait until the solution has settled and all bubbles have risen to the top before withdrawing your dose from the vial.



Figure G



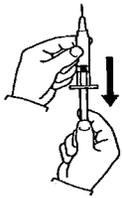
Figure H

9. After the solution has settled and is completely dissolved it should be clear, colorless and without particles, but there may be a ring of foam or bubbles on the surface, this is normal. Do not use it if you see particles or the color is not correct.

10. After the PegIntron powder is dissolved but before you withdraw your dose, clean the rubber stopper again with an alcohol swab.

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11. Unwrap the second syringe provided. You will use it to give yourself the injection. Carefully remove the protective cap from the needle and fill the syringe with air by pulling the plunger to the number on the side of the syringe (mL) that corresponds to your prescribed dose (**Figure J**). Hold the PegIntron vial upright. **DO NOT** touch the cleaned top of the vial with your hands (**Figure K**). Insert the needle into the vial containing the PegIntron solution and inject the air into the center of the vial (**Figure L**).



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Figure J

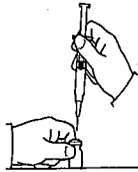


Figure K

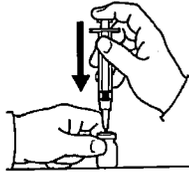
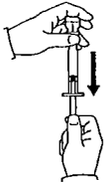


Figure L

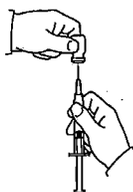
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12. Turn the PegIntron vial upside down. Be sure the tip of the needle is in the PegIntron solution. While holding the vial and syringe with one hand slowly pull the plunger back to withdraw the exact amount of PegIntron into the syringe your health care provider told you to use (**Figure M**).



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421
422
Figure M

13. Remove the needle from the vial (**Figure N**) and check for air bubbles in the syringe. If you see any bubbles, hold the syringe with the needle pointing up and gently tap the syringe until the bubbles rise. Then push the plunger in slowly until the bubbles disappear.



423 Figure N

424

425

426 **Injecting the PegIntron Dose**427 **Selecting the Site for Injection.**

428 The best sites for giving yourself an injection are those areas with a layer of fat between
 429 the skin and muscle, like your thigh, the outer surface of your upper arm, and abdomen.
 430 Do not inject yourself in the area near your navel or waistline. If you are very thin, you
 431 should only use the thigh or outer surface of the arm for injection.

432

433 You should use a different site each time you inject PegIntron to avoid soreness at any one
 434 site. Do not inject PegIntron solution into an area where the skin is irritated, red, bruised,
 435 infected or has scars, stretch marks or lumps.

436

437 14. Clean the skin where the injection is to be given with an alcohol swab, and wait for
 438 the area to dry. Remove the protective cap from the needle. Make sure the safety sleeve of
 439 the syringe is pushed firmly against the syringe flange so that the needle is fully exposed
 440 (see **Figure A**).

441

442 15. With one hand, pinch a 2-inch fold of loose skin. With your other hand, pick up the
 443 syringe and hold it like a pencil. Position the bevel of the needle facing up and insert the
 444 needle approximately $\frac{1}{4}$ inch into the pinched skin at approximately a 45 to 90 degree
 445 angle with a quick dart-like thrust. After the needle is in, remove the hand that you used to
 446 pinch your skin and use it to hold the syringe barrel. Pull the plunger of the syringe back
 447 very slightly. If blood comes into the syringe, the needle has entered a blood vessel. **Do**
 448 **not inject.** Withdraw the needle and discard the syringe as outlined in step 17. Repeat the
 449 above steps with a new vial to prepare a new syringe and inject the medicine at a new site.
 450 If no blood is present in the syringe, inject the medicine by gently pressing the plunger all
 451 the way down the syringe barrel.

452

453 16. Hold an alcohol swab near the needle and pull the needle straight out of the skin.
 454 Press the alcohol swab over the injection site for several seconds. Do not massage the
 455 injection site. If there is bleeding, cover it with a bandage.

456

457 17. After injecting your dose, firmly grasp the safety sleeve and pull it over the exposed
 458 needle until you hear a click, and the green stripe on the safety sleeve covers the red stripe
 459 on the needle (**Figure O**). Discard the syringe and needle in the Sharp's container supplied
 460 to you.

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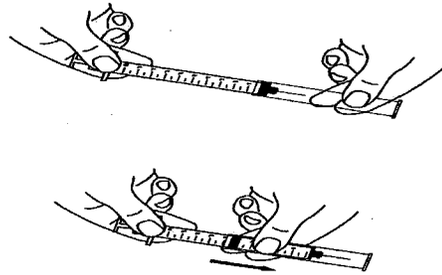


Figure O

18. After 2 hours, check the injection site for redness, swelling, or tenderness. If you have a skin reaction and it doesn't clear up in a few days, contact your health care provider or nurse.

How do I dispose of the used syringes and needles?

Discard used safety lock syringes and needles in a Sharp's container or other puncture proof container like a coffee can. DO NOT USE glass or clear plastic containers. Your health care provider or nurse will tell you how to dispose a full container. Always keep the container out of reach of children.

Manufactured by: Schering Corporation, Kenilworth, NJ 07033 USA

DATE 3/08

1 **032408.final draft Med Guide Redipen**
2 **MEDICATION GUIDE**
3 **PegIntron™ Redipen® Single-dose Delivery System**
4 **(Peginterferon alfa-2b)**

5
6 **Including appendix with instructions for using PegIntron™ Redipen® Single-dose**
7 **Delivery System**

8
9 Read this Medication Guide carefully before you start taking PegIntron™ (**Peg In-tron**) or
10 PegIntron™/REBETOL® (**REB-eh-tole**) combination therapy. Read the Medication Guide
11 each time you refill your prescription because there may be new information. The
12 information in this Medication Guide does not take the place of talking with your health care
13 provider (doctor, nurse, nurse practitioner, or physician's assistant).

14
15 If you are taking PegIntron/REBETOL combination therapy, also read the Medication Guide
16 for REBETOL (ribavirin, USP) Capsules.

17
18 **What is the most important information I should know about PegIntron and**
19 **PegIntron/REBETOL combination therapy?**

20
21 PegIntron (peginterferon) is a treatment for some people who are infected with hepatitis C
22 virus. However, PegIntron and PegIntron/REBETOL combination therapy can have serious
23 side effects that may cause death in rare cases. Before you decide to start treatment, you
24 should talk to your health care provider about the possible benefits and side effects of
25 PegIntron or PegIntron/REBETOL combination therapy. If you begin treatment you will
26 need to see your health care provider regularly for medical examinations and lab tests to
27 make sure your treatment is working and to check for side effects.

28
29 **REBETOL capsules may cause birth defects and/or death of an unborn child. If you are**
30 **pregnant, you or your male partner must not take PegIntron/REBETOL combination**
31 **therapy. You must not become pregnant while either you or your partner are being**
32 **treated with the combination PegIntron/REBETOL therapy, or for 6 months after**
33 **stopping therapy. Men and women should use birth control while taking the**
34 **combination therapy and for 6 months afterwards. If you or your partner are being**
35 **treated and you become pregnant, either during treatment or within 6 months of**
36 **stopping treatment, call your health care provider right away. There is a Ribavirin**
37 **Pregnancy Registry that collects information about pregnancy outcomes of female**
38 **patients and female partners of male patients exposed to ribavirin. You or your**
39 **healthcare provider are encouraged to contact the Registry at 1-800-593-2214.**

40
41 If you are taking PegIntron or PegIntron/REBETOL therapy you should call your
42 health care provider immediately if you develop any of these symptoms:

43
44 New or worsening mental health problems, such as thoughts about killing or hurting
45 yourself or others, trouble breathing, chest pain, severe stomach or lower back pain,

46 **bloody diarrhea or bloody bowel movements, high fever, bruising, bleeding, or**
47 **decreased vision.**

48

49 The most serious possible side effects of PegIntron and PegIntron/REBETOL therapy
50 include:

51

52 **Problems with Pregnancy. Combination PegIntron/REBETOL therapy can cause**
53 **death, serious birth defects, or other harm to your unborn child. If you are a woman of**
54 **childbearing age, you must not become pregnant during treatment and for 6 months**
55 **after you have stopped therapy. You must have a negative pregnancy test immediately**
56 **before beginning treatment, during treatment, and for 6 months after you have stopped**
57 **therapy. Both males and female patients must use effective forms of birth control**
58 **during treatment and for the 6 months after treatment is completed. Male patients**
59 **should use a condom.** If you are a female, you must use birth control even if you believe
60 that you are not fertile or that your fertility is low. You should talk to your health care
61 provider about birth control for you and your partner.

62

63 **Mental health problems and suicide.** PegIntron and PegIntron/REBETOL therapies may
64 cause patients to develop mood or behavioral problems. These can include irritability (getting
65 easily upset) and depression (feeling low, feeling bad about yourself, or feeling hopeless).
66 Some patients may have aggressive behavior. Former drug addicts may fall back into drug
67 addiction or overdose. Some patients think about hurting or killing themselves or other
68 people and some have killed (suicide) or hurt themselves or others. You must tell your
69 health care provider if you are being treated for a mental illness or had treatment in the past
70 for any mental illness, including depression and suicidal behavior. You should tell your
71 health care provider if you have ever been addicted to drugs or alcohol.

72

73 **Heart problems.** Some patients taking PegIntron or PegIntron/REBETOL therapy may
74 develop problems with their heart, including low blood pressure, fast heart rate, and very
75 rarely, heart attacks. Tell your health care provider if you have had any heart problems in the
76 past.

77

78 **Blood problems.** PegIntron and PegIntron/REBETOL therapies commonly lower two types
79 of blood cells (white blood cells and platelets). In some patients, these blood counts may fall
80 to dangerously low levels. If your blood counts become very low, this could lead to
81 infections or bleeding.

82

83 REBETOL therapy causes a decrease in the number of red blood cells you have (anemia).
84 This can be dangerous, especially for patients who already have heart or circulatory
85 (cardiovascular) problems. Talk with your health care provider before taking combination
86 PegIntron/REBETOL therapy if you have, or have ever had any cardiovascular problems.

87

88 **Body organ problems.** Certain symptoms like severe stomach pain may mean that your
89 internal organs are being damaged. Cases of weakness, loss of coordination, and numbness
90 due to stroke have been reported in patients taking PegIntron, including patients with few or
91 no reported risk factors for stroke.

92

93 *For other possible side effects, see "What are the possible side effects of PegIntron and*
94 *PegIntron/REBETOL" in this Medication Guide.*

95

96 **What is PegIntron and PegIntron/REBETOL combination therapy?**

97 The PegIntron product is a drug used to treat adults who have a lasting (chronic) infection
98 with hepatitis C virus and who show signs that the virus is damaging the liver.

99 PegIntron/REBETOL combination therapy consists of two medications also used to treat
100 hepatitis C infection. Patients with hepatitis C have the virus in their blood and in their liver.
101 PegIntron reduces the amount of virus in the body and helps the body's immune system fight
102 the virus. REBETOL (ribavirin) is a drug that helps to fight the viral infection, but does not
103 work when used by itself to treat chronic hepatitis C.

104

105 It is not known if PegIntron or PegIntron/REBETOL therapies can cure hepatitis C
106 (permanently eliminate the virus), or if it can prevent liver failure or liver cancer that is
107 caused by hepatitis C infection.

108

109 It is also not known if PegIntron or PegIntron/REBETOL combination therapy will prevent
110 one infected person from infecting another person with hepatitis C.

111

112 **Who should not take PegIntron or PegIntron/REBETOL therapy?**

113 Do not take PegIntron or PegIntron/REBETOL therapy if you:

114

- are pregnant, planning to get pregnant during treatment or during the 6 months after
115 treatment, or breast-feeding

116

- are a male patient with a female sexual partner who is pregnant, or plans to become
117 pregnant at any time while you are being treated with REBETOL, or during the 6
118 months after your treatment has ended

119

- have hepatitis caused by your immune system attacking your liver (autoimmune
120 hepatitis) or unstable liver disease

121

- had an allergic reaction to another alpha interferon or are allergic to any of the
122 ingredients in PegIntron or REBETOL Capsules. If you have any doubts, ask your
123 health care provider.

124

- Do not take PegIntron/REBETOL combination therapy if you have abnormal red
125 blood cells such as sickle-cell anemia or thalassemia major.

126

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137

**If you have any of the following conditions or serious medical problems, discuss them
with your health care provider before taking PegIntron or PegIntron/REBETOL
therapy:**

- depression or anxiety
- sleep problems
- high blood pressure
- previous heart attack, or other heart problems

- 138 • liver problems (other than hepatitis C infection)
- 139 • any kind of autoimmune disease (where the body's immune system attacks the body's
- 140 own cells), such as psoriasis, systemic lupus erythematosus, rheumatoid arthritis
- 141 • thyroid problems
- 142 • diabetes
- 143 • colitis (inflammation of the bowels)
- 144 • cancer
- 145 • hepatitis B infection
- 146 • HIV infection
- 147 • kidney problems
- 148 • bleeding problems
- 149 • alcoholism
- 150 • drug abuse or addiction
- 151 • body organ transplant and are taking medicine that keeps your body from rejecting your
- 152 transplant (suppresses your immune system)

153

154

155 **How should I take PegIntron or PegIntron/REBETOL?**

156 Your health care provider will decide whether you will take PegIntron therapy alone or the
157 combination of PegIntron/REBETOL, as well as the correct dose (based on your weight).
158 PegIntron and PegIntron/REBETOL are given for one year. Take your prescribed dose of
159 PegIntron ONCE A WEEK, on the same day of each week and at approximately the same
160 time. Take the medicine for the full year and do not take more than the prescribed dose.
161 REBETOL Capsules should be taken with food. When you take REBETOL with food, more
162 of the medicine (70% more on average) is taken up by your body. You should take
163 REBETOL the same way every day (twice a day with food) to keep the medicine in your
164 body at a steady level. This will help your health care provider to decide how your treatment
165 is working and how to change the number of REBETOL capsules you take if you have side
166 effects from REBETOL. **Be sure to read the Medication Guide for REBETOL**
167 **(ribavirin, USP) for complete instructions on how to take the REBETOL capsules.**

168

169 You should be completely comfortable with how to prepare PegIntron, how to set the dose
170 you take, and how to inject yourself before you use PegIntron for the first time. PegIntron
171 comes in two different forms, a powder in a single-use vial and a Redipen single-use delivery
172 system. See the attached appendix for detailed instructions for preparing and giving a dose of
173 PegIntron.

174

175 If you miss a dose of the PegIntron product, take the missed dose as soon as possible during
176 the same day or the next day, then continue on your regular dosing schedule. If several days
177 go by after you miss a dose, check with your health care provider about what to do. Do not
178 double the next dose or take more than one dose a week without talking to your health care
179 provider. Call your health care provider right away if you take more than your prescribed
180 PegIntron dose. Your health care provider may wish to examine you more closely, and take
181 blood for testing.

182

183 If you miss a dose of REBETOL capsules, take the missed dose as soon as possible during
184 the same day. If an entire day has gone by, check with your health care provider about what
185 to do. Do not double the next dose.

186

187 You must get regular blood tests to help your health care provider check how the treatment is
188 working and to check for side effects.

189

190 Tell your health care provider if you are taking or planning to take other prescription or non-
191 prescription medicines, including vitamin and mineral supplements and herbal medicines.

192

193 **What should I avoid while taking PegIntron or PegIntron/REBETOL therapies?**

194 • If you are pregnant do not start taking PegIntron/REBETOL combination therapy.

195 • Avoid becoming pregnant while taking PegIntron or PegIntron/REBETOL.

196 PegIntron and PegIntron/REBETOL may harm your unborn child (death or serious birth
197 defects) or cause you to lose your baby (miscarry). **If you or your partner becomes**
198 **pregnant during treatment or during the 6 months after treatment with**
199 **PegIntron/REBETOL combination therapy, immediately report the pregnancy to your**
200 **health care provider. You or your health care provider should call 1-800-593-2214.** By
201 calling this number, information about you and/or your partner will be added to a pregnancy
202 registry that will be used to help you and your health care provider make decisions about
203 your treatment for hepatitis in the future. You, your partner, and/or your health care provider
204 will be asked to provide follow-up information on the outcome of the pregnancy.

205

206 • Do not breast-feed your baby while taking PegIntron.

207

208 **What are the possible side effects of PegIntron and PegIntron/REBETOL combination**
209 **therapy?**

210

211 **Possible, serious side effects include:**

212 **Mental health problems including suicide, blood problems, heart problems, body organ**
213 **problems.** See "What is the most important information I should know about PegIntron and
214 PegIntron/REBETOL combination therapy?"

215

216 **Other body organ problems.** A few patients have lung problems (such as pneumonia or
217 inflammation of the lung tissue), inflammation of the kidney, and eye disorders.

218

219 **New or worsening autoimmune disease.** Some patients taking PegIntron or
220 PegIntron/REBETOL develop autoimmune diseases (a condition where the body's immune
221 cells attack other cells or organs in the body), including rheumatoid arthritis, systemic lupus
222 erythematosus, and psoriasis. In some patients who already have an autoimmune disease, the
223 disease worsens on PegIntron and PegIntron/REBETOL combination therapy.

224

225 Common but less serious side effects include:

226

227 **Flu-like symptoms.** Most patients who take PegIntron or PegIntron/REBETOL therapy have
228 "flu-like" symptoms (headache, muscle aches, tiredness, and fever). Some of these symptoms

229 (fever, headache) usually lessen after the first few weeks of therapy. You can reduce some of
230 these symptoms by injecting your PegIntron dose at bedtime. Over-the-counter pain and
231 fever reducers, such as acetaminophen or ibuprofen, can be used to prevent or reduce the
232 fever and headache.

233

234 **Extreme fatigue (tiredness).** Many patients become extremely tired while on PegIntron or
235 PegIntron/REBETOL combination therapy.

236

237 **Appetite problems.** Nausea, loss of appetite, and weight loss, occur commonly.

238

239 **Thyroid problems.** Some patients develop changes in the function of their thyroid.
240 Symptoms of thyroid changes include the inability to concentrate, feeling cold or hot all the
241 time, a change in your weight, and changes to your skin.

242

243 **Blood sugar problems.** Some patients develop problems with the way their body controls
244 their blood sugar, and may develop high blood sugar or diabetes.

245

246 **Skin reactions.** Redness, swelling, and itching are common at the site of injection. If after
247 several days these symptoms do not disappear contact your health care provider. You may
248 get a rash during therapy. If this occurs, your health care provider may recommend medicine
249 to treat the rash.

250

251 **Hair thinning.** Hair thinning is common during PegIntron and PegIntron/REBETOL
252 treatment. Hair loss stops and hair growth returns after therapy is stopped.

253

254 These are not all of the side effects of PegIntron or PegIntron/REBETOL combination
255 therapy. Your health care provider or pharmacist can give you a more complete list.

256

257 Call your doctor for medical advice about side effects. You may report side effects to FDA at
258 1-800-FDA-1088.

259

260 **General advice about prescription medicines:**

261 Medicines are sometimes prescribed for purposes other than those listed in a Medication
262 Guide. If you have any concerns about PegIntron, ask your health care provider. Your health
263 care provider or pharmacist can give you information about PegIntron that was written for
264 health care professionals. Do not use PegIntron for a condition for which it was not
265 prescribed. Do not share this medication with other people.

266

267 If you are taking PegIntron/REBETOL combination therapy, also read the Medication Guide
268 for REBETOL (ribavirin, USP) Capsules.

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270 *This Medication Guide has been approved by the U.S. Food and Drug Administration.*

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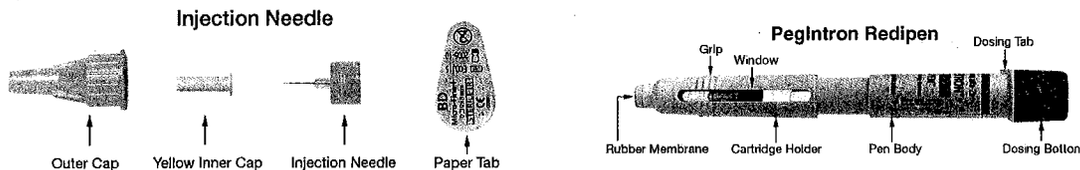
273 **How do I prepare and inject the PegIntron Redipen Dose?**

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 275 The PegIntron Redipen system is for a single use, by one person only, ONCE A WEEK. The
 276 Redipen must not be shared. Use only the injection needle provided in the packaging for the
 277 PegIntron Redipen system. If you have problems with the Redipen system or the PegIntron
 278 solution, you should contact your health care provider or pharmacist.

279
 280 The following instructions explain how to prepare and inject yourself with the PegIntron
 281 Redipen system. Please read the instructions carefully and follow them step by step. Your
 282 health care provider will instruct you on how to self-inject with the PegIntron Redipen. Do
 283 not attempt to inject yourself unless you are sure you understand the procedure and
 284 requirements for self-injection.

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How to Use the PegIntron Redipen Single-dose Delivery System.



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Storing PegIntron

PegIntron Redipen should be stored in the refrigerator at 2°C to 8°C (36°F to 46°F); avoid exposure to heat. After mixing, the PegIntron solution should be used immediately but may be stored in the refrigerator up to 24 hours at 2°C to 8°C (36°F - 46°F). The solution contains no preservatives. DO NOT FREEZE.

Preparation

1. Find a clean, well-lit, non-slip flat working surface and assemble all of the supplies you will need for an injection. All of the supplies you will need are in the PegIntron Redipen package. The package contains:
 - a PegIntron Redipen single-dose delivery system
 - one disposable needle
 - two alcohol swabs, and
 - Dosing tray; (The dosing tray is the bottom half of the Redipen package.)
2. Take the PegIntron Redipen out of the refrigerator and allow the medicine to come to room temperature. Before removing the Redipen from the carton, check the expiration date printed on the PegIntron Redipen carton to make sure that the expiration date has not passed. Do not use if the expiration date has passed.

- 313 3. After taking the PegIntron Redipen out of the carton, look in the window of the Redipen
314 and make sure the PegIntron in the cartridge holder window is a white, to off white tablet
315 that is whole, or in pieces, or powdered.
316 4. Wash your hands thoroughly with soap and water, rinse, and towel dry. It is important
317 to keep your work area, your hands, and the injection site clean to minimize the risk of
318 infection.

319

320 **1. Mix the Drug**

321

322 **Key points:**

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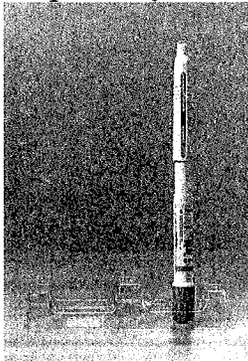
324 **Before you mix the PegIntron, make sure it is at room temperature. It is important that**
325 **you keep the PegIntron Redipen UPRIGHT (Dosing Button down) as shown in Figure**
326 **1.**

327

328 a. Hold the PegIntron Redipen **UPRIGHT (Figure 1a)** in the dosing tray on a hard, flat,
329 non-slip surface with the dosing button **down**. You may want to hold the Redipen using
330 the grip.

331

332 b. To mix the powder and the liquid, keep the Redipen upright in the dosing tray and
333 press the top half of the Redipen downward toward the hard, flat, non-slip surface **until**
334 **you hear the click** (Figure 1b). Once you've heard the click, you will notice in the
335 window that both dark stoppers are now touching. The dosing button should be flush with
336 the pen body.

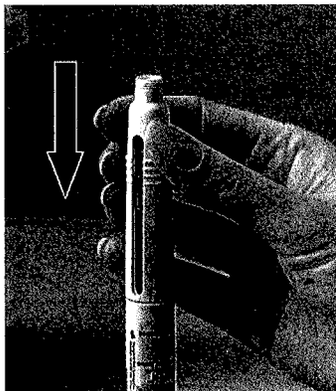


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339 **Figure 1a**

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341 **Figure 1b**

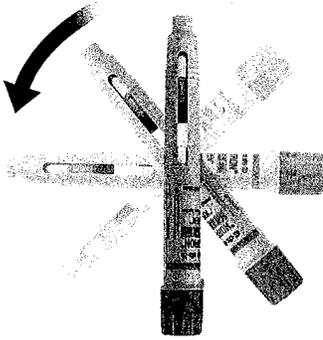
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343 c. Wait several seconds for the powder to completely dissolve.

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345

346 **d. Gently turn the PegIntron Redipen upside down twice (Figure 2). To avoid**
347 **excessive foaming, DO NOT SHAKE.**



348

349 **Figure 2**

350

351 e. Keep the PegIntron™ Redipen® **UPRIGHT**, with the dosing button down. Then, look
352 through the Redipen® window to see that the mixed PegIntron™ solution is completely
353 dissolved. The solution should be clear and colorless **before use**. Before attaching the
354 needle, it is normal to see some small bubbles in the Redipen® window, near the top of
355 the solution. Do not use the solution if it is discolored, or not clear, or if particulates are
356 present.

357

358 **f. Place the PegIntron Redipen back into the dosing tray provided in the packaging**
359 **(Figure 3). The dosing button will be on the bottom.**

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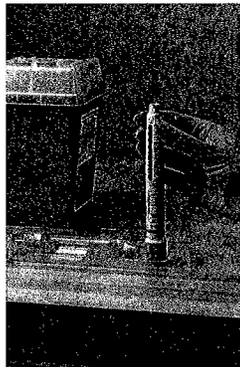


Figure 3

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2. Attach the Needle

- a. Wipe the rubber membrane of the PegIntron Redipen with one alcohol swab.
- b. Remove the protective paper tab from the injection needle, but do NOT remove either the outer cap or the yellow inner cap from the injection needle. Keeping the PegIntron Redipen UPRIGHT in the dosing tray, FIRMLY push the injection needle straight into the Redipen rubber membrane, and screw it firmly in place, in a clockwise direction (Figure 4). Remember to leave the needle caps in place when you attach the needle to the Redipen. Pushing the needle through the rubber membrane, “primes” the needle and allows the extra liquid and air in the pen to be removed.

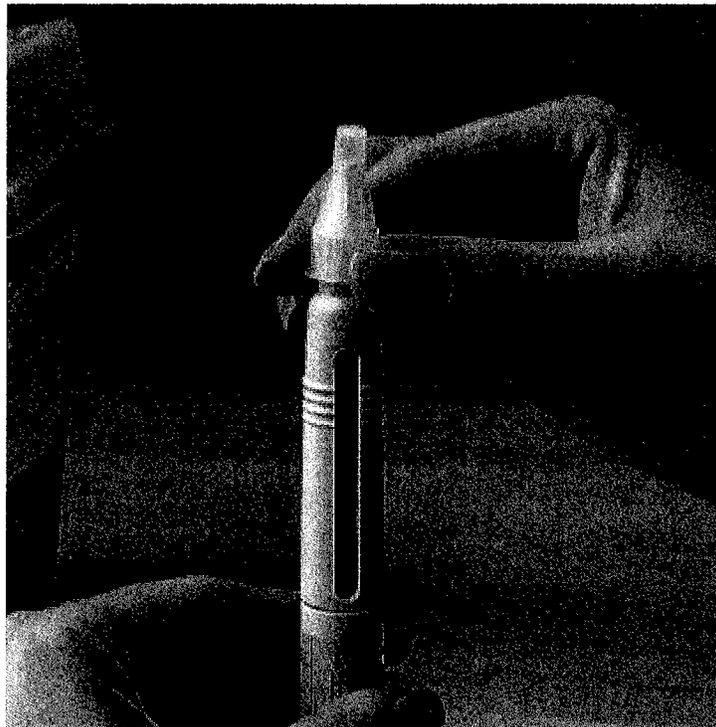


Figure 4

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NOTE: Some fluid will trickle out. This is **normal**. The dark stoppers move up and you will no longer see the fluid in the window once the needle is successfully primed.

3. Dialing the Dose

- a. **Remove the PegIntron Redipen from the dosing tray (Figure 5a).**
Holding the PegIntron Redipen firmly, pull the dosing button out as far as it will go. You will see a dark band-

Do not push the dosing button in until you are ready to self-inject the PegIntron dose.

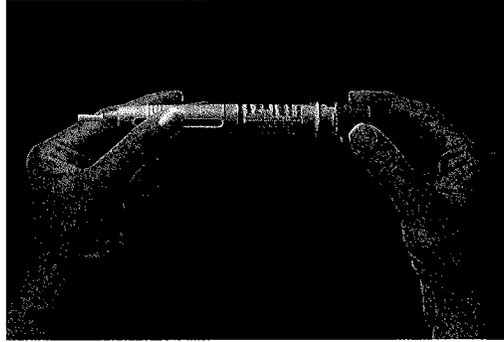


Figure 5a

b. Turn the dosing button until your prescribed dose is lined up with the dosing tab (**Figure 5b**). The dosing button will turn freely. If you have trouble dialing your dose, check to make sure the dosing button has been pulled out as far as it will go (**Figure 5c**).

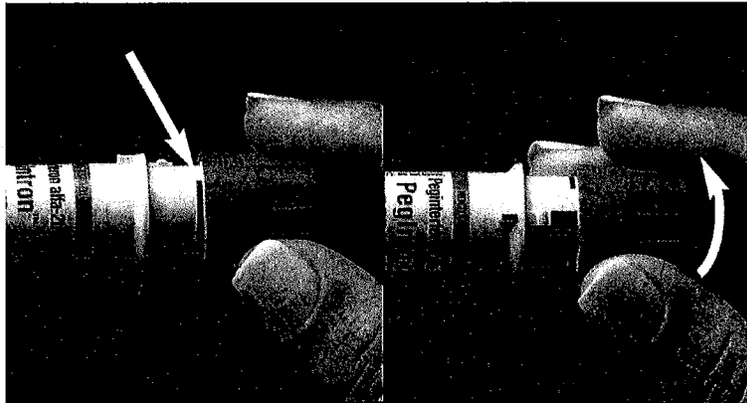


Figure 5b

Figure 5c

c. Carefully lay the PegIntron Redipen down on a hard, flat, non-slip surface. Do NOT remove either of the needle caps and do NOT push the dosing button in until you are ready to self-inject the PegIntron dose.

4. Injecting the PegIntron Dose

Choosing an Injection Site

The best sites for giving yourself an injection are those areas with a layer of fat between the skin and muscle, like your thigh, the outer surface of your upper arm, and abdomen. Do not inject yourself in the area near your navel or waistline. If you are very thin, you should only use the thigh or outer surface of the arm for injection.

You should use a different site each time you inject PegIntron to avoid soreness at any one site. Do not inject PegIntron into an area where the skin is irritated, red, bruised, infected, or has scars, stretch marks, or lumps.

- 420 a. Clean the skin where the injection is to be given with the second alcohol swab provided,
421 and wait for the area to dry.
422 b. Remove the **outer** cap from the needle (**Figure 6a**). There may be some liquid around the
423 yellow inner needle cap (**Figure 6b**). This is normal.
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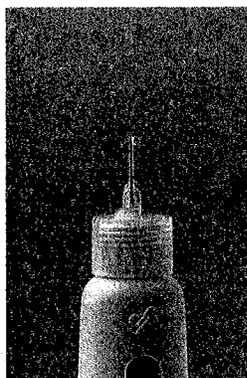


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428 **Figure 6a****Figure 6b**

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- 430 c. Once the injection site is dry, remove the **yellow** inner needle cap (**Figure 6c**). You are
431 now ready to inject.
432



433

434 **Figure 6c**

435

- 436 **d. Hold the PegIntron Redipen with your fingers wrapped around the pen body barrel**
437 **and your thumb on the dosing button (Figure 7).**

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- With your other hand, pinch the skin in the area you have cleaned for injection.
- Insert the needle into the pinched skin at an angle of 45° to 90°.
- Press the dosing button down slowly and firmly until you can't push it any further.
- Keep your thumb pressed down on the dosing button for an additional 5 seconds to ensure that you get the complete dose.
- Remove the needle from your skin.