PAGE 1

- 1 **PRODUCT**
- 2 INFORMATION
- 3 INTRON[®] A
- 4 Interferon alfa-2b,
- 5 recombinant
- 6 For Injection
- 7

8 WARNING Alpha interferons, including INTRON[®] A, cause or aggravate fatal or
9 life-threatening neuropsychiatric, autoimmune, ischemic, and infectious
10 disorders. Patients should be monitored closely with periodic clinical and
11 laboratory evaluations. Patients with persistently severe or worsening signs or
12 symptoms of these conditions should be withdrawn from therapy. In many but
13 not all cases these disorders resolve after stopping INTRON A therapy. See
14 WARNINGS and ADVERSE REACTIONS.

15

16 **DESCRIPTION**

INTRON[®] A (Interferon alfa-2b) for intramuscular, subcutaneous, intralesional, or
 intravenous Injection is a purified sterile recombinant interferon product.

INTRON A recombinant for Injection has been classified as an alpha interferon 19 and is a water-soluble protein with a molecular weight of 19,271 daltons produced by 20 21 recombinant DNA techniques. It is obtained from the bacterial fermentation of a strain of Escherichia coli bearing a genetically engineered plasmid containing an interferon alfa-22 23 2b gene from human leukocytes. The fermentation is carried out in a defined nutrient 24 medium containing the antibiotic tetracycline hydrochloride at a concentration of 5 to 10 mg/L: the presence of this antibiotic is not detectable in the final product. The specific 25 activity of interferon alfa-2b, recombinant is approximately 2.6 x 10⁸ IU/mg protein as 26 27 measured by the HPLC assay. Powder for Injection

Vial Strength Million IU	mL Diluent	Final Concentration after Reconstitution million IU/mL*	mg INTRON A [†] per vial	Route of Administration
10	1	10	0.038	IM, SC, IV, IL
18	1	18	0.069	IM, SC, IV
50	1	50	0.192	IM, SC, IV

* Each mL also contains 20 mg glycine, 2.3 mg sodium phosphate dibasic, 0.55 mg sodium phosphate monobasic, and 1.0 mg human albumin.

[†] Based on the specific activity of approximately 2.6 x 10⁸ IU/mg protein, as measured by HPLC assay.

28 Prior to administration, the INTRON A Powder for Injection is to be reconstituted with

the provided Diluent for INTRON A (Sterile Water for Injection USP) (see **DOSAGE**

30 **AND ADMINISTRATION**). INTRON A Powder for Injection is a white to cream-colored

31 powder.

Solution Vials for Injection

Vial Strength	Concentration*	mg INTRON A [†] per vial	Route of Administration
18 [‡] MIU multidose	3 million IU/0.5 mL	0.088	IM, SC
25 [¶] MIU multidose	5 million IU/0.5 mL	0.123	IM, SC, IL

* Each mL contains 7.5 mg sodium chloride, 1.8 mg sodium phosphate dibasic,
 1.3 mg sodium phosphate monobasic, 0.1 mg edetate disodium, 0.1 mg polysorbate
 80, and 1.5 mg m-cresol as a preservative.

⁺ Based on the specific activity of approximately 2.6 x 10⁸ IU/mg protein as measured by HPLC assay.

[‡] This is a multidose vial which contains a total of 22.8 million IU of interferon alfa-2b, recombinant per 3.8 mL in order to provide the delivery of six 0.5-mL doses, each containing 3 million IU of INTRON A (for a label strength of 18 million IU).

34

Pen Strength	Concentration* million IU/1.5mL	INTRON A Dose Delivered (6 doses, 0.2 mL each)	mg INTRON A [†] per 1.5 mL	Route of Administration
3 MIU	22.5	3 MIU/0.2 mL	0.087	SC
5 MIU	37.5	5 MIU/0.2 mL	0.144	SC
10 MIU	75	10 MIU/0.2 mL	0.288	SC

Solution in Multidose Pens for Injection

* Each mL also contains 7.5 mg sodium chloride, 1.8 mg sodium phosphate dibasic, 1.3 mg sodium phosphate monobasic, 0.1 mg edetate disodium, 0.1 mg polysorbate 80, and 1.5 mg m-cresol as a preservative.

[†] Based on the specific activity of approximately 2.6 x 10⁸ IU/mg protein as measured by HPLC assay.

35 36

These packages do not require reconstitution prior to administration (see DOSAGE

37 AND ADMINISTRATION). INTRON A Solution for Injection is a clear, colorless solution.

38

39 CLINICAL PHARMACOLOGY

40 **General** The interferons are a family of naturally occurring small proteins and 41 glycoproteins with molecular weights of approximately 15,000 to 27,600 daltons 42 produced and secreted by cells in response to viral infections and to synthetic or 43 biological inducers.

44 **Preclinical Pharmacology** Interferons exert their cellular activities by binding to 45 specific membrane receptors on the cell surface. Once bound to the cell membrane, 46 interferons initiate a complex sequence of intracellular events. *In vitro* studies 47 demonstrated that these include the induction of certain enzymes, suppression of cell 48 proliferation, immunomodulating activities such as enhancement of the phagocytic 49 activity of macrophages and augmentation of the specific cytotoxicity of lymphocytes for 49 tornet cells.

50 target cells, and inhibition of virus replication in virus-infected cells.

¹ This is a multidose vial which contains a total of 32.0 million IU of interferon alfa-2b, recombinant per 3.2 mL in order to provide the delivery of five 0.5-mL doses, each containing 5 million IU of INTRON A (for a label strength of 25 million IU).

51 In a study using human hepatoblastoma cell line HB 611, the *in vitro* antiviral 52 activity of alpha interferon was demonstrated by its inhibition of hepatitis B virus (HBV) 53 replication.

54 The correlation between these *in vitro* data and the clinical results is unknown. 55 Any of these activities might contribute to interferon's therapeutic effects.

56 *Pharmacokinetics* The pharmacokinetics of INTRON[®] A were studied in 12 healthy 57 male volunteers following single doses of 5 million IU/m² administered intramuscularly, 58 subcutaneously, and as a 30-minute intravenous infusion in a crossover design.

59 The mean serum INTRON A concentrations following intramuscular and 60 subcutaneous injections were comparable. The maximum serum concentrations 61 obtained via these routes were approximately 18 to 116 IU/mL and occurred 3 to 62 12 hours after administration. The elimination half-life of INTRON A following both 63 intramuscular and subcutaneous injections was approximately 2 to 3 hours. Serum 64 concentrations were undetectable by 16 hours after the injections.

After intravenous administration, serum INTRON A concentrations peaked (135-273 IU/mL) by the end of the 30-minute infusion, then declined at a slightly more rapid rate than after intramuscular or subcutaneous drug administration, becoming undetectable 4 hours after the infusion. The elimination half-life was approximately 2 hours.

Urine INTRON A concentrations following a single dose (5 million IU/m²) were not detectable after any of the parenteral routes of administration. This result was expected since preliminary studies with isolated and perfused rabbit kidneys have shown that the kidney may be the main site of interferon catabolism.

There are no pharmacokinetic data available for the intralesional route of administration.

76 Serum Neutralizing Antibodies In INTRON A-treated patients tested for antibody 77 activity in clinical trials, serum anti-interferon neutralizing antibodies were detected in 78 0% (0/90) of patients with hairy cell leukemia, 0.8% (2/260) of patients treated 79 intralesionally for condylomata acuminata, and 4% (1/24) of patients with AIDS-Related 80 Kaposi's Sarcoma. Serum neutralizing antibodies have been detected in less than 3% of 81 patients treated with higher INTRON A doses in malignancies other than hairy cell 82 leukemia or AIDS-Related Kaposi's Sarcoma. The clinical significance of the 83 appearance of serum anti-interferon neutralizing activity in these indications is not 84 known.

85 Serum anti-interferon neutralizing antibodies were detected in 7% (12/168) of 86 patients either during treatment or after completing 12 to 48 weeks of treatment with 3 87 million IU TIW of INTRON A therapy for chronic hepatitis C and in 13% (6/48) of 88 patients who received INTRON A therapy for chronic hepatitis B at 5 million IU QD for 4 89 months, and in 3% (1/33) of patients treated at 10 million IU TIW. Serum anti-interferon 90 neutralizing antibodies were detected in 9% (5/53) of pediatric patients who received INTRON A therapy for chronic hepatitis B at 6 million IU/m² TIW. Among all chronic 91 92 hepatitis B or C patients, pediatrics and adults with detectable serum neutralizing 93 antibodies, the titers detected were low (22/24 with titers less than or equal to 1:40 and 94 2/24 with titers less than or equal to 1:160). The appearance of serum anti-interferon 95 neutralizing activity did not appear to affect safety or efficacy.

PAGE 4

97 Hairy Cell Leukemia In clinical trials in patients with hairy cell leukemia, there was 98 depression of hematopoiesis during the first 1 to 2 months of INTRON A treatment, 99 resulting in reduced numbers of circulating red and white blood cells, and platelets. 100 Subsequently, both splenectomized and nonsplenectomized patients achieved 101 substantial and sustained improvements in granulocytes, platelets, and hemoglobin 102 levels in 75% of treated patients and at least some improvement (minor responses) 103 INTRON A treatment resulted in a decrease in bone marrow occurred in 90%. 104 hypercellularity and hairy cell infiltrates. The hairy cell index (HCI), which represents 105 the percent of bone marrow cellularity times the percent of hairy cell infiltrate, was 106 greater than or equal to 50% at the beginning of the study in 87% of patients. The 107 percentage of patients with such an HCI decreased to 25% after 6 months and to 14% 108 after 1 year. These results indicate that even though hematologic improvement had 109 occurred earlier, prolonged INTRON A treatment may be required to obtain maximal 110 reduction in tumor cell infiltrates in the bone marrow.

111 The percentage of patients with hairy cell leukemia who required red blood cell or 112 platelet transfusions decreased significantly during treatment and the percentage of 113 patients with confirmed and serious infections declined as granulocyte counts improved. 114 Reversal of splenomegaly and of clinically significant hypersplenism was demonstrated 115 in some patients.

116 A study was conducted to assess the effects of extended INTRON A treatment 117 on duration of response for patients who responded to initial therapy. In this study, 126 118 responding patients were randomized to receive additional INTRON A treatment for 6 119 months or observation for a comparable period, after 12 months of initial INTRON A therapy. During this 6-month period, 3% (2/66) of INTRON A-treated patients relapsed 120 121 compared with 18% (11/60) who were not treated. This represents a significant 122 difference in time to relapse in favor of continued INTRON A treatment (P=0.006/0.01, 123 Log Rank/Wilcoxon). Since a small proportion of the total population had relapsed, 124 median time to relapse could not be estimated in either group. A similar pattern in 125 relapses was seen when all randomized treatment, including that beyond 6 months, and 126 available follow-up data were assessed. The 15% (10/66) relapses among INTRON A 127 patients occurred over a significantly longer period of time than the 40% (24/60) with 128 observation (P=0.0002/0.0001, Log Rank/Wilcoxon). Median time to relapse was 129 estimated, using the Kaplan-Meier method, to be 6.8 months in the observation group 130 but could not be estimated in the INTRON A group.

Subsequent follow-up with a median time of approximately 40 months
demonstrated an overall survival of 87.8%. In a comparable historical control group
followed for 24 months, overall median survival was approximately 40%.

134

135 **Malignant Melanoma** The safety and efficacy of INTRON A was evaluated as adjuvant 136 to surgical treatment in patients with melanoma who were free of disease (post surgery) 137 but at high risk for systemic recurrence. These included patients with lesions of Breslow 138 thickness greater than 4 mm, or patients with lesions of any Breslow thickness with 139 primary or recurrent nodal involvement. In a randomized, controlled trial in 280 patients, 143 patients received INTRON A therapy at 20 million IU/m² intravenously five times per 140 week for 4 weeks (induction phase) followed by 10 million IU/m² subcutaneously three 141 142 times per week for 48 weeks (maintenance phase). In the clinical trial, the median daily

INTRON A dose administered to patients was 19.1 million IU/m² during the induction
 phase and 9.1 million IU/m² during the maintenance phase. INTRON A therapy was
 begun less than or equal to 56 days after surgical resection. The remaining 137
 patients were observed.

147 INTRON A therapy produced a significant increase in relapse-free and overall 148 survival. Median time to relapse for the INTRON A-treated patients versus observation 149 patients was 1.72 years versus 0.98 years (P<0.01, stratified Log Rank). The estimated 150 5-year relapse-free survival rate, using the Kaplan-Meier method, was 37% for INTRON 151 A-treated patients versus 26% for observation patients. Median overall survival time for 152 INTRON A-treated patients versus observation patients was 3.82 years versus 2.78 153 years (P=0.047, stratified Log Rank). The estimated 5-year overall survival rate, using the Kaplan-Meier method, was 46% for INTRON A-treated patients versus 37% for 154 155 observation patients.

156 In a second study of 642 resected high-risk melanoma patients, subjects were 157 randomized equally to one of three groups; high-dose INTRON A therapy for 1 year 158 (same schedule as above), low-dose INTRON A therapy for 2 years (3 MU/d TIW SC), 159 and observation. Consistent with the earlier trial, high-dose INTRON A therapy 160 demonstrated an improvement in relapse-free survival (3-year estimated RFS 48% 161 versus 41%; median RFS 2.4 versus 1.6 years, P=not significant). Relapse-free 162 survival in the low-dose INTRON A arm was similar to that seen in the observation arm. 163 Neither high-dose nor low-dose INTRON A therapy showed a benefit in overall survival 164 as compared to observation in this study.

165

166 **Follicular Lymphoma** The safety and efficacy of INTRON A in conjunction with CHVP, 167 a combination chemotherapy regimen, was evaluated as initial treatment in patients with 168 clinically aggressive, large tumor burden, Stage III/IV follicular Non-Hodgkin's 169 Lymphoma. Large tumor burden was defined by the presence of any one of the 170 following: a nodal or extranodal tumor mass with a diameter of greater than 7 cm; 171 involvement of at least three nodal sites (each with a diameter of greater than 3 cm): systemic symptoms; splenomegaly; serous effusion, orbital or epidural involvement; 172 173 ureteral compression; or leukemia.

174 In a randomized, controlled trial, 130 patients received CHVP therapy and 175 135 patients received CHVP therapy plus INTRON A therapy at 5 million IU 176 subcutaneously three times weekly for the duration of 18 months. CHVP chemotherapy consisted of cyclophosphamide 600 mg/m², doxorubicin 25 mg/m², and teniposide (VM-177 178 26) 60 mg/m², administered intravenously on Day 1 and prednisone at a daily dose of 40 mg/m² given orally on Days 1 to 5. Treatment consisted of six CHVP cycles 179 180 administered monthly, followed by an additional six cycles administered every 2 months 181 for 1 year. Patients in both treatment groups received a total of 12 CHVP cycles over 182 18 months.

The group receiving the combination of INTRON A therapy plus CHVP had a significantly longer progression-free survival (2.9 years versus 1.5 years, P=0.0001, Log Rank test). After a median follow-up of 6.1 years, the median survival for patients treated with CHVP alone was 5.5 years while median survival for patients treated with CHVP plus INTRON A therapy had not been reached (P=0.004, Log Rank test). In three additional published, randomized, controlled studies of the addition of interferon

PAGE 6

alpha to anthracycline-containing combination chemotherapy regimens,¹⁻³ the addition
 of interferon alpha was associated with significantly prolonged progression-free survival.
 Differences in overall survival were not consistently observed.

192

193 **Condylomata Acuminata** Condylomata acuminata (venereal or genital warts) are 194 associated with infections of the human papilloma virus (HPV). The safety and efficacy 195 of INTRON A in the treatment of condylomata acuminata were evaluated in three 196 controlled double-blind clinical trials. In these studies, INTRON A doses of 1 million IU 197 per lesion were administered intralesionally three times a week (TIW), in less than or 198 equal to 5 lesions per patient for 3 weeks. The patients were observed for up to 16 199 weeks after completion of the full treatment course.

200 INTRON A treatment of condylomata was significantly more effective than 201 placebo, as measured by disappearance of lesions, decreases in lesion size, and by an 202 overall change in disease status. Of 192 INTRON A-treated patients and 206 placebo-203 treated patients who were evaluable for efficacy at the time of best response during the 204 course of the study, 42% of INTRON A patients versus 17% of placebo patients 205 experienced clearing of all treated lesions. Likewise, 24% of INTRON A patients versus 206 8% of placebo patients experienced marked (75% to less than 100%) reduction in lesion 207 size, 18% versus 9% experienced moderate (50% to 75%) reduction in lesion size, 10% 208 versus 42% had a slight (less than 50%) reduction in lesion size, 5% versus 24% had 209 no change in lesion size, and 0% versus 1% experienced exacerbation (P<0.001).

In one of these studies, 43% (54/125) of patients in whom multiple (less than or equal to 3) lesions were treated experienced complete clearing of all treated lesions during the course of the study. Of these patients, 81% remained cleared 16 weeks after treatment was initiated.

Patients who did not achieve total clearing of all their treated lesions had these same lesions treated with a second course of therapy. During this second course of treatment, 38% to 67% of patients had clearing of all treated lesions. The overall percentage of patients who had cleared all their treated lesions after two courses of treatment ranged from 57% to 85%.

219 INTRON A-treated lesions showed improvement within 2 to 4 weeks after the 220 start of treatment in the above study; maximal response to INTRON A therapy was 221 noted 4 to 8 weeks after initiation of treatment.

The response to INTRON A therapy was better in patients who had condylomata for shorter durations than in patients with lesions for a longer duration.

224 Another study involved 97 patients in whom three lesions were treated with either 225 an intralesional injection of 1.5 million IU of INTRON A per lesion followed by a topical 226 application of 25% podophyllin, or a topical application of 25% podophyllin alone. 227 Treatment was given once a week for 3 weeks. The combined treatment of INTRON A 228 and podophyllin was shown to be significantly more effective than podophyllin alone, as 229 determined by the number of patients whose lesions cleared. This significant difference 230 in response was evident after the second treatment (Week 3) and continued through 8 231 weeks post-treatment. At the time of the patient's best response, 67% (33/49) of the 232 INTRON A- and podophyllin-treated patients had all three treated lesions clear while 233 42% (20/48) of the podophyllin-treated patients had all three clear (P=0.003).

PAGE 7

AIDS-Related Kaposi's Sarcoma The safety and efficacy of INTRON A in the treatment of Kaposi's Sarcoma (KS), a common manifestation of the Acquired Immune Deficiency Syndrome (AIDS), were evaluated in clinical trials in 144 patients.

In one study, INTRON A doses of 30 million IU/m² were administered subcutaneously three times per week (TIW) to patients with AIDS-Related KS. Doses were adjusted for patient tolerance. The average weekly dose delivered in the first 4 weeks was 150 million IU; at the end of 12 weeks this averaged 110 million IU/week; and by 24 weeks averaged 75 million IU/week.

Forty-four percent of asymptomatic patients responded versus 7% of symptomatic patients. The median time to response was approximately 2 months and 1 month, respectively, for asymptomatic and symptomatic patients. The median duration of response was approximately 3 months and 1 month, respectively, for the asymptomatic and symptomatic patients. Baseline T4/T8 ratios were 0.46 for responders versus 0.33 for nonresponders.

In another study, INTRON A doses of 35 million IU were administered subcutaneously, daily (QD), for 12 weeks. Maintenance treatment, with every other day dosing (QOD), was continued for up to 1 year in patients achieving antitumor and antiviral responses. The median time to response was 2 months and the median duration of response was 5 months in the asymptomatic patients.

In all studies, the likelihood of response was greatest in patients with relatively intact immune systems as assessed by baseline CD4 counts (interchangeable with T4 counts). Results at doses of 30 million IU/m² TIW and 35 million IU/QD were subcutaneously similar and are provided together in TABLE 1. This table demonstrates the relationship of response to baseline CD4 count in both asymptomatic and symptomatic patients in the 30 million IU/m² TIW and the 35 million IU/QD treatment groups.

In the 30 million IU study group, 7% (5/72) of patients were complete responders
and 22% (16/72) of the patients were partial responders. The 35 million IU study had
13% (3/23 patients) complete responders and 17% (4/23) partial responders.

For patients who received 30 million IU TIW, the median survival time was longer in patients with CD4 greater than 200 (30.7 months) than in patients with CD4 less than or equal to 200 (8.9 months). Among responders, the median survival time was 22.6 months versus 9.7 months in nonresponders.

268

269 **Chronic Hepatitis C** The safety and efficacy of INTRON A in the treatment of chronic 270 hepatitis C was evaluated in 5 randomized clinical studies in which an INTRON A dose 271 of 3 million IU three times a week (TIW) was assessed. The initial three studies were 272 placebo-controlled trials that evaluated a 6-month (24-week) course of therapy. In each 273 of the three studies, INTRON A therapy resulted in a reduction in serum alanine 274 aminotransferase (ALT) in a greater proportion of patients versus control patients at the 275 end of 6 months of dosing. During the 6 months of follow-up, approximately 50% of the 276 patients who responded maintained their ALT response. A combined analysis 277 comparing pretreatment and post-treatment liver biopsies revealed histological 278 improvement in a statistically significantly greater proportion of INTRON A-treated 279 patients compared to controls.

280 Two additional studies have investigated longer treatment durations (up to 24 months).^{5,6} Patients in the two studies to evaluate longer duration of treatment had 281 hepatitis with or without cirrhosis in the absence of decompensated liver disease. 282 283 Complete response to treatment was defined as normalization of the final two serum 284 ALT levels during the treatment period. A sustained response was defined as a 285 complete response at the end of the treatment period, with sustained normal ALT 286 values lasting at least 6 months following discontinuation of therapy.

287 In Study 1, all patients were initially treated with INTRON A 3 million IU TIW 288 subcutaneously for 24 weeks (run-in-period). Patients who completed the initial 289 24-week treatment period were then randomly assigned to receive no further treatment, 290 or to receive 3 million IU TIW for an additional 48 weeks. In Study 2, patients who met 291 the entry criteria were randomly assigned to receive INTRON A 3 million IU TIW 292 subcutaneously for 24 weeks or to receive INTRON A 3 million IU TIW subcutaneously 293 for 96 weeks. In both studies, patient follow-up was variable and some data collection 294 was retrospective.

295 Results show that longer durations of INTRON A therapy improved the sustained 296 response rate (see TABLE 2). In patients with complete responses (CR) to INTRON A 297 therapy after 6 months of treatment (149/352 [42%]), responses were less often 298 sustained if drug was discontinued (21/70 [30%]) than if it was continued for 18 to 24 299 months (44/79 [56%]). Of all patients randomized, the sustained response rate in the 300 patients receiving 18 or 24 months of therapy was 22% and 26%, respectively, in the 301 two trials. In patients who did not have a CR by 6 months, additional therapy did not 302 result in significantly more responses, since almost all patients who responded to 303 therapy did so within the first 16 weeks of treatment.

A subset (less than 50%) of patients from the combined extended dosing studies 304 305 had liver biopsies performed both before and after INTRON A treatment. Improvement 306 in necroinflammatory activity as assessed retrospectively by the Knodell (Study 1) and 307 Scheuer (Study 2) Histology Activity Indices was observed in both studies. A higher 308 number of patients (58%, 45/78) improved with extended therapy than with shorter (6 months) therapy (38%, 34/89) in this subset. 309

Combination treatment with INTRON A and REBETOL[®] (ribavirin USP) provided 310 311 a significant reduction in virologic load and improved histologic response in adult 312 patients with compensated liver disease who were treatment-naïve or had relapsed 313 following therapy with alpha interferon alone; pediatric patients previously untreated with 314 alpha interferon experienced a sustained virologic response. See REBETOL prescribing 315 information for additional information.

316

317 Chronic Hepatitis B Adults The safety and efficacy of INTRON A in the treatment of 318 chronic hepatitis B were evaluated in three clinical trials in which INTRON A doses of 30 319 to 35 million IU per week were administered subcutaneously (SC), as either 5 million IU daily (QD), or 10 million IU three times a week (TIW) for 16 weeks versus no treatment. 320 321 All patients were 18 years of age or older with compensated liver disease, and had 322 chronic hepatitis B virus (HBV) infection (serum HBsAg positive for at least 6 months) and HBV replication (serum HBeAg positive). Patients were also serum HBV-DNA 323 positive, an additional indicator of HBV replication, as measured by a research assav.^{7,8} 324 325 All patients had elevated serum alanine aminotransferase (ALT) and liver biopsy

PAGE 9

findings compatible with the diagnosis of chronic hepatitis. Patients with the presence
 of antibody to human immunodeficiency virus (anti-HIV) or antibody to hepatitis delta
 virus (anti-HDV) in the serum were excluded from the studies.

Virologic response to treatment was defined in these studies as a loss of serum markers of HBV replication (HBeAg and HBV DNA). Secondary parameters of response included loss of serum HBsAg, decreases in serum ALT, and improvement in liver histology.

In each of two randomized controlled studies, a significantly greater proportion of INTRON A-treated patients exhibited a virologic response compared with untreated control patients (see TABLE 3). In a third study without a concurrent control group, a similar response rate to INTRON A therapy was observed. Pretreatment with prednisone, evaluated in two of the studies, did not improve the response rate and provided no additional benefit.

The response to INTRON A therapy was durable. No patient responding to INTRON A therapy at a dose of 5 million IU QD or 10 million IU TIW relapsed during the follow-up period, which ranged from 2 to 6 months after treatment ended. The loss of serum HBeAg and HBV DNA was maintained in 100% of 19 responding patients followed for 3.5 to 36 months after the end of therapy.

In a proportion of responding patients, loss of HBeAg was followed by the loss of HBsAg. HBsAg was lost in 27% (4/15) of patients who responded to INTRON A therapy at a dose of 5 million IU QD, and 35% (8/23) of patients who responded to 10 million IU TIW. No untreated control patient lost HBsAg in these studies.

In an ongoing study to assess the long-term durability of virologic response, 64 patients responding to INTRON A therapy have been followed for 1.1 to 6.6 years after treatment; 95% (61/64) remain serum HBeAg negative, and 49% (30/61) lost serum HBsAg.

INTRON A therapy resulted in normalization of serum ALT in a significantly
 greater proportion of treated patients compared to untreated patients in each of two
 controlled studies (see TABLE 4). In a third study without a concurrent control group,
 normalization of serum ALT was observed in 50% (12/24) of patients receiving INTRON
 A therapy.

Virologic response was associated with a reduction in serum ALT to normal or near normal (less than or equal to 1.5 x the upper limit of normal) in 87% (13/15) of patients responding to INTRON A therapy at 5 million IU QD, and 100% (23/23) of patients responding to 10 million IU TIW.

361 Improvement in liver histology was evaluated in Studies 1 and 3 by comparison 362 of pretreatment and 6-month post-treatment liver biopsies using the semiguantitative Knodell Histology Activity Index.⁹ No statistically significant difference in liver histology 363 was observed in treated patients compared to control patients in Study 1. Although 364 365 statistically significant histological improvement from baseline was observed in treated 366 patients in Study 3 ($P \le 0.01$), there was no control group for comparison. Of those 367 patients exhibiting a virologic response following treatment with 5 million IU QD or 10 368 million IU TIW, histological improvement was observed in 85% (17/20) compared to 369 36% (9/25) of patients who were not virologic responders. The histological 370 improvement was due primarily to decreases in severity of necrosis, degeneration, and 371 inflammation in the periportal, lobular, and portal regions of the liver (Knodell Categories

I + II + III). Continued histological improvement was observed in four responding
 patients who lost serum HBsAg and were followed 2 to 4 years after the end of INTRON
 A therapy.¹⁰

375

376 **Pediatrics** The safety and efficacy of INTRON A in the treatment of chronic hepatitis B 377 was evaluated in one randomized controlled trial of 149 patients ranging from 1 year to 378 17 years of age. Seventy-two patients were treated with 3 million IU/m² of INTRON A therapy administered subcutaneously three times a week (TIW) for 1 week; the dose 379 was then escalated to 6 million IU/m^2 TIW for a minimum of 16 weeks up to 24 weeks. 380 The maximum weekly dosage was 10 million IU TIW. Seventy-seven patients were 381 382 untreated controls. Study entry and response criteria were identical to those described 383 in the adult patient population.

384 Patients treated with INTRON A therapy had a better response (loss of HBV DNA 385 and HBeAg at 24 weeks of follow-up) compared to the untreated controls (24% [17/72] 386 versus 10% [8/77] P=0.05). Sixteen of the 17 responders treated with INTRON A therapy remained HBV DNA and HBeAg negative and had a normal serum ALT 12 to 387 388 24 months after completion of treatment. Serum HBsAg became negative in 7 out of 17 389 patients who responded to INTRON A therapy. None of the control patients who had an 390 HBV DNA and HBeAg response became HBsAg negative. At 24 weeks of follow-up, 391 normalization of serum ALT was similar in patients treated with INTRON A therapy 392 (17%, 12/72) and in untreated control patients (16%, 12/77). Patients with a baseline 393 HBV DNA less than 100 pg/mL were more likely to respond to INTRON A therapy than 394 were patients with a baseline HBV DNA greater than 100 pg/mL (35% versus 9%, 395 respectively). Patients who contracted hepatitis B through maternal vertical transmission had lower response rates than those who contracted the disease by other 396 397 means (5% versus 31%, respectively). There was no evidence that the effects on HBV 398 DNA and HBeAg were limited to specific subpopulations based on age, gender, or race. 399

400

	-	TABLE 1						
RESP	PONSE BY BASELINE CD4 (COUNT IN AIDS-RELATED	<u>KS PATIENTS</u>					
	<u>3</u>	<u>0 million IU/m² TIW, SC and</u>						
		<u>35 million IU QD, SC</u>						
	<u>Asymptomatic</u>		<u>Symptomatic</u>					
CD4<200	4/14 (29	9%) 0/19	(0%)					
200≤CD4≤400	6/12 (50	0%) 0/5	(0%)					
		} 58%						
CD4>400	5/7 (71	l%) 0/0	(0%)					
* Data for CD4, and asymptomatic and symptomatic classification were not available for all patients.								
			·					
		TABLE 2						
SUS	STAINED ALT RESPONSE R	ATE VERSUS DURATION C	F THERAPY					
	IN CHRONIC H	EPATITIS C PATIENTS						
	INTRON	A 3 Million IU TIW						
	Treat	ment Group [*] - Number of Pat	tients (%)					
			Difference					
Study	INTRON A 3 million IU	INTRON A 3 million IU	(Extended — 24					
Number	24 weeks of treatment	72 or 96 weeks of treatme	nt [†] weeks)					
			(95% CI) [‡]					

ALT respon	se at the	end of	follow-up
------------	-----------	--------	-----------

PAGE 11

1	12/101 (12%)	23/104 (22%)	10% (-3, 24)			
2	9/67 (13%)	21/80 (26%)	13% (-4, 30)			
Combined Studies	21/168 (12.5%)	44/184 (24%)	11.4% (2, 21)			
ALT response at the end of treatment						
	ALT response a	the end of treatment				
1	40/101 (40%)	51/104 (49%)				

* Intent-to-treat groups.

[†] Study 1: 72 weeks of treatment; Study 2: 96 weeks of treatment.

[‡] Confidence intervals adjusted for multiple comparisons due to 3 treatment arms in the study.

402 403

All Studies	15/38	(39%)	23/48	(48%)	6/91	(7%)	
3 ⁸			13/24 [§]	(54%)	2/27	(7%) [§]	NA [§]
2			10/24	(42%)	1/22	(5%)	0.005
1 ⁷	15/38	(39%)			3/42	(7%)	0.0009
Number	5 millio	n IU QD	10 million IU TIW		Controls		Value
Study	INTR	RON A	INTR	ON A	Untr	reated	P^{\ddagger}
		<u>Treatme</u>	nt Group [†] - N	lumber of Pa	ntients (%)		
			IA	DLE J			

* Loss of HBeAg and HBV DNA by 6 months post-therapy.

[†] Patients pretreated with prednisone not shown.

[‡] INTRON A treatment group versus untreated control.

[§] Untreated control patients evaluated after 24-week observation period. A subgroup subsequently received INTRON A therapy. A direct comparison is not applicable (NA).

TABLE 4

404

	ALT	<u> RESPONS</u>	<u>ES[*] IN CHR</u>	ONIC HEPA	TITIS B PAT	<u>IENTS</u>	
		Treatme	ent Group - N	lumber of Pa	tients (%)		
Study	INTF	RON A	INTR	ON A	Untr	eated	P^{\dagger}
Number	5 millio	n IU QD	10 million IU TIW		Controls		Value
1	16/38	(42%)			8/42	(19%)	0.03
2			10/24	(42%)	1/22	(5%)	0.0034
3			12/24 [‡]	(50%)	2/27	(7%) [‡]	NA [‡]
All Studies	16/38	(42%)	22/48	(46%)	11/91	(12%)	

* Reduction in serum ALT to normal by 6 months post-therapy.

[†] INTRON A treatment group versus untreated control.

[‡] Untreated control patients evaluated after 24-week observation period. A subgroup subsequently received INTRON A therapy. A direct comparison is not applicable (NA).

405

406 INDICATIONS AND USAGE

407 **Hairy Cell Leukemia** INTRON[®] A is indicated for the treatment of patients 18 years of 408 age or older with hairy cell leukemia.

409

410 **Malignant Melanoma** INTRON A is indicated as adjuvant to surgical treatment in 411 patients 18 years of age or older with malignant melanoma who are free of disease but 412 at high risk for systemic recurrence, within 56 days of surgery.

413

414 **Follicular Lymphoma** INTRON A is indicated for the initial treatment of clinically 415 aggressive (see **Clinical Pharmacology**) follicular Non-Hodgkin's Lymphoma in

PAGE 12

416 conjunction with anthracycline-containing combination chemotherapy in patients 18 417 years of age or older. Efficacy of INTRON A therapy in patients with low-grade, low-418 tumor burden follicular Non-Hodgkin's Lymphoma has not been demonstrated.

419

420 **Condvlomata Acuminata INTRON** A is indicated for intralesional treatment of selected 421 patients 18 years of age or older with condylomata acuminata involving external 422 surfaces of the genital and perianal areas (see **DOSAGE AND ADMINISTRATION**).

423 424

430

The use of this product in adolescents has not been studied.

425 AIDS-Related Kaposi's Sarcoma INTRON A is indicated for the treatment of selected 426 patients 18 years of age or older with AIDS-Related Kaposi's Sarcoma. The likelihood 427 of response to INTRON A therapy is greater in patients who are without systemic 428 symptoms, who have limited lymphadenopathy and who have a relatively intact immune 429 system as indicated by total CD4 count.

431 **Chronic Hepatitis C** INTRON A is indicated for the treatment of chronic hepatitis C in 432 patients 18 years of age or older with compensated liver disease who have a history of blood or blood-product exposure and/or are HCV antibody positive. Studies in these 433 434 patients demonstrated that INTRON A therapy can produce clinically meaningful effects 435 on this disease, manifested by normalization of serum alanine aminotransferase (ALT) 436 and reduction in liver necrosis and degeneration.

437 A liver biopsy should be performed to establish the diagnosis of chronic hepatitis. 438 Patients should be tested for the presence of antibody to HCV. Patients with other 439 causes of chronic hepatitis, including autoimmune hepatitis, should be excluded. Prior 440 to initiation of INTRON A therapy, the physician should establish that the patient has 441 compensated liver disease. The following patient entrance criteria for compensated liver 442 disease were used in the clinical studies and should be considered before INTRON A 443 treatment of patients with chronic hepatitis C:

444 445

446

- No history of hepatic encephalopathy, variceal bleeding, ascites, or other clinical signs of decompensation
- 447 Bilirubin Less than or equal to 2 mg/dL
- 448 Albumin Stable and within normal limits
- 449 Prothrombin Time Less than 3 seconds prolonged •
- WBC 450 Greater than or equal to 3000/mm³ •
- Greater than or equal to 70,000/mm³ 451 • Platelets
- 452
- 453 Serum creatinine should be normal or near normal.

454 Prior to initiation of INTRON A therapy, CBC and platelet counts should be 455 evaluated in order to establish baselines for monitoring potential toxicity. These tests 456 should be repeated at Weeks 1 and 2 following initiation of INTRON A therapy, and 457 monthly thereafter. Serum ALT should be evaluated at approximately 3-month intervals 458 to assess response to treatment (see DOSAGE AND ADMINISTRATION).

PAGE 13

Patients with preexisting thyroid abnormalities may be treated if thyroidstimulating hormone (TSH) levels can be maintained in the normal range by medication.
TSH levels must be within normal limits upon initiation of INTRON A treatment and TSH
testing should be repeated at 3 and 6 months (see **PRECAUTIONS, Laboratory Tests**).

INTRON A in combination with REBETOL[®] is indicated for the treatment of
 chronic hepatitis C in patients 3 years of age and older with compensated liver disease
 previously untreated with alpha interferon therapy and in patients 18 years of age and
 older who have relapsed following alpha interferon therapy. See REBETOL prescribing
 information for additional information.

469

470 Chronic Hepatitis B INTRON A is indicated for the treatment of chronic hepatitis B in 471 patients 1 year of age or older with compensated liver disease. Patients who have been 472 serum HBsAg positive for at least 6 months and have evidence of HBV replication 473 (serum HBeAg positive) with elevated serum ALT are candidates for treatment. Studies 474 in these patients demonstrated that INTRON A therapy can produce virologic remission 475 of this disease (loss of serum HBeAg) and normalization of serum aminotransferases. 476 INTRON A therapy resulted in the loss of serum HBsAg in some responding patients.

477 Prior to initiation of INTRON A therapy, it is recommended that a liver biopsy be 478 performed to establish the presence of chronic hepatitis and the extent of liver damage. 479 The physician should establish that the patient has compensated liver disease. The 480 following patient entrance criteria for compensated liver disease were used in the 481 clinical studies and should be considered before INTRON A treatment of patients with 482 chronic hepatitis B:

- 483
- No history of hepatic encephalopathy, variceal bleeding, ascites, or other signs of clinical decompensation
- 486 Bilirubin Normal
- 487 Albumin Stable and within normal limits
- Prothrombin Time *Adults* less than 3 seconds prolonged
- 489
 490 WBC
 Pediatrics less than or equal to 2 seconds prolonged
 Greater than or equal to 4000/mm³
- 491 Platelets Adults greater than or equal to 100,000/mm³
 - Pediatrics greater than or equal to 150,000/mm³

492 493

494 Patients with causes of chronic hepatitis other than chronic hepatitis B or chronic 495 hepatitis C should not be treated with INTRON A. CBC and platelet counts should be 496 evaluated prior to initiation of INTRON A therapy in order to establish baselines for 497 monitoring potential toxicity. These tests should be repeated at treatment Weeks 1, 2, 4, 8, 12, and 16. Liver function tests, including serum ALT, albumin, and bilirubin, 498 499 should be evaluated at treatment Weeks 1, 2, 4, 8, 12, and 16. HBeAg, HBsAg, and 500 ALT should be evaluated at the end of therapy, as well as 3- and 6-months post-501 therapy, since patients may become virologic responders during the 6-month period

502 following the end of treatment. In clinical studies in adults, 39% (15/38) of responding 503 patients lost HBeAg 1 to 6 months following the end of INTRON A therapy. Of 504 responding patients who lost HBsAg, 58% (7/12) did so 1 to 6 months post-treatment.

505 A transient increase in ALT greater than or equal to 2 times baseline value (flare) 506 can occur during INTRON A therapy for chronic hepatitis B. In clinical trials in adults and pediatrics, this flare generally occurred 8 to 12 weeks after initiation of therapy and 507 508 was more frequent in responders (adults 63%, 24/38; pediatrics 59%, 10/17) than in 509 nonresponders (adults 27%, 13/48; pediatrics 35%, 19/55). However, in adults and 510 pediatrics, elevations in bilirubin greater than or equal to 3 mg/dL (greater than or equal 511 to 2 times ULN) occurred infrequently (adults 2%, 2/86; pediatrics 3%, 2/72) during 512 therapy. When ALT flare occurs, in general, INTRON A therapy should be continued 513 unless signs and symptoms of liver failure are observed. During ALT flare, clinical 514 symptomatology and liver function tests including ALT, prothrombin time, alkaline 515 phosphatase, albumin, and bilirubin, should be monitored at approximately 2-week 516 intervals (see WARNINGS).

517

520

521

522

523

525

526

527

528

529

531

518 **CONTRAINDICATIONS**

519 INTRON[®] A is contraindicated in patients with:

- Hypersensitivity to interferon alpha or any component of the product
- Autoimmune hepatitis
- Decompensated liver disease

524 INTRON A and REBETOL[®] combination therapy is additionally contraindicated in:

- Patients with hypersensitivity to ribavirin or any other component of the product
- Women who are pregnant
- Men whose female partners are pregnant
- Patients with hemoglobinopathies (e.g., thalassemia major, sickle cell anemia)
- Patients with creatinine clearance less than 50 mL/min.
- 530 See REBETOL prescribing information for additional information.

532 WARNINGS

533 General Moderate to severe adverse experiences may require modification of the patient's dosage regimen, or in some cases termination of INTRON® A therapy. 534 535 Because of the fever and other "flu-like" symptoms associated with INTRON A 536 administration, it should be used cautiously in patients with debilitating medical 537 conditions, such as those with a history of pulmonary disease (e.g., chronic obstructive 538 pulmonary disease) or diabetes mellitus prone to ketoacidosis. Caution should also be 539 observed in patients with coagulation disorders (e.g., thrombophlebitis, pulmonary 540 embolism) or severe myelosuppression.

541

542 Cardiovascular Disorders

INTRON A therapy should be used cautiously in patients with a history of cardiovascular
disease. Those patients with a history of myocardial infarction and/or previous or
current arrhythmic disorder who require INTRON A therapy should be closely monitored
(see **PRECAUTIONS, Laboratory Tests**). Cardiovascular adverse experiences, which
include hypotension, arrhythmia, or tachycardia of 150 beats per minute or greater, and

548 rarely, cardiomyopathy and myocardial infarction have been observed in some INTRON 549 A-treated patients. Some patients with these adverse events had no history of 550 cardiovascular disease. Transient cardiomyopathy was reported in approximately 2% of 551 the AIDS-Related Kaposi's Sarcoma patients treated with INTRON A. Hypotension may 552 occur during INTRON A administration, or up to 2 days post-therapy, and may require 553 supportive therapy including fluid replacement to maintain intravascular volume.

554 Supraventricular arrhythmias occurred rarely and appeared to be correlated with 555 preexisting conditions and prior therapy with cardiotoxic agents. These adverse 556 experiences were controlled by modifying the dose or discontinuing treatment, but may 557 require specific additional therapy.

558

566

559 Cerebrovascular Disorders

560 Ischemic and hemorrhagic cerebrovascular events have been observed in patients 561 treated with interferon alpha-based therapies, including INTRON A. Events occurred in 562 patients with few or no reported risk factors for stroke, including patients less than 45 563 years of age. Because these are spontaneous reports, estimates of frequency cannot 564 be made and a causal relationship between interferon alpha-based therapies and these 565 events is difficult to establish.

567 Neuropsychiatric Disorders

568 DÉPRESSION AND SUICIDAL BEHAVIOR INCLUDING SUICIDAL IDEATION, 569 SUICIDAL ATTEMPTS, AND COMPLETED SUICIDES, HOMICIDAL IDEATION, AND 570 AGGRESSIVE BEHAVIOR SOMETIMES DIRECTED TOWARDS OTHERS, HAVE 571 BEEN REPORTED IN ASSOCIATION WITH TREATMENT WITH ALPHA 572 INTERFERONS, INCLUDING INTRON A THERAPY. If patients develop psychiatric 573 problems, including clinical depression, it is recommended that the patients be carefully 574 monitored during treatment and in the 6-month follow-up period.

575 INTRON A should be used with caution in patients with a history of psychiatric 576 disorders. INTRON A therapy should be discontinued for any patient developing severe 577 psychiatric disorder during treatment. Obtundation and coma have also been observed 578 in some patients, usually elderly, treated at higher doses. While these effects are 579 usually rapidly reversible upon discontinuation of therapy, full resolution of symptoms 580 has taken up to 3 weeks in a few severe episodes. If psychiatric symptoms persist or 581 worsen, or suicidal ideation or aggressive behavior towards others is identified, it is 582 recommended that treatment with INTRON A be discontinued and the patient followed, 583 with psychiatric intervention as appropriate. Narcotics, hypnotics, or sedatives may be 584 used concurrently with caution and patients should be closely monitored until the 585 adverse effects have resolved. Suicidal ideation or attempts occurred more frequently 586 among pediatric patients, primarily adolescents, compared to adult patients (2.4%) 587 versus 1%) during treatment and off-therapy follow-up. Cases of encephalopathy have 588 also been observed in some patients, usually elderly, treated with higher doses of 589 INTRON A.

590 Treatment with interferons may be associated with exacerbated symptoms of 591 psychiatric disorders in patients with co-occurring psychiatric and substance use 592 disorders. If treatment with interferons is initiated in patients with prior history or 593 existence of psychiatric condition or with a history of substance use disorders, treatment

PAGE 16

594 considerations should include the need for drug screening and periodic health 595 evaluation, including psychiatric symptom monitoring. Early intervention for re-596 emergence or development of neuropsychiatric symptoms and substance use is 597 recommended.

- 598
- 599

600 Bone Marrow Toxicity

601 INTRON A therapy suppresses bone marrow function and may result in severe 602 cytopenias including aplastic anemia. It is advised that complete blood counts (CBC) 603 be obtained pretreatment and monitored routinely during therapy (see **PRECAUTIONS**, 604 **Laboratory Tests**). INTRON A therapy should be discontinued in patients who develop 605 severe decreases in neutrophil (less than 0.5×10^9 /L) or platelet counts (less than 25×10^9 /L) (see **DOSAGE AND ADMINISTRATION**, Guidelines for Dose Modification). 607

608 **Ophthalmologic Disorders**

609 Decrease or loss of vision, retinopathy including macular edema, retinal artery or 610 vein thrombosis, retinal hemorrhages and cotton wool spots; optic neuritis, papilledema, 611 and serous retinal detachment may be induced or aggravated by treatment with 612 interferon alfa-2b or other alpha interferons. All patients should receive an eve 613 examination at baseline. Patients with preexisting ophthalmologic disorders (e.g., diabetic or hypertensive retinopathy) should receive periodic ophthalmologic exams 614 615 during interferon alpha treatment. Any patient who develops ocular symptoms should 616 receive a prompt and complete eye examination. Interferon alfa-2b treatment should be 617 discontinued in patients who develop new or worsening ophthalmologic disorders.

619 Endocrine Disorders

620 Infrequently, patients receiving INTRON A therapy developed thyroid 621 abnormalities, either hypothyroid or hyperthyroid. The mechanism by which INTRON A 622 may alter thyroid status is unknown. Patients with preexisting thyroid abnormalities whose thyroid function cannot be maintained in the normal range by medication should 623 624 not be treated with INTRON A. Prior to initiation of INTRON A therapy, serum TSH 625 should be evaluated. Patients developing symptoms consistent with possible thyroid 626 dysfunction during the course of INTRON A therapy should have their thyroid function 627 evaluated and appropriate treatment instituted. Therapy should be discontinued for 628 patients developing thyroid abnormalities during treatment whose thyroid function 629 cannot be normalized by medication. Discontinuation of INTRON A therapy has not 630 always reversed thyroid dysfunction occurring during treatment. Diabetes mellitus has 631 been observed in patients treated with alpha interferons. Patients with these conditions 632 who cannot be effectively treated by medication should not begin INTRON A therapy. 633 Patients who develop these conditions during treatment and cannot be controlled with 634 medication should not continue INTRON A therapy.

635

618

636 Gastrointestinal Disorders

637 Hepatotoxicity, including fatality, has been observed in interferon alpha-treated 638 patients, including those treated with INTRON A. Any patient developing liver function

PAGE 17

abnormalities during treatment should be monitored closely and if appropriate,
treatment should be discontinued.

642 **Pulmonary Disorders**

643 Dyspnea, pulmonary infiltrates, pneumonia, bronchiolitis obliterans, interstitial 644 pneumonitis, pulmonary hypertension, and sarcoidosis, some resulting in respiratory 645 failure and/or patient deaths, may be induced or aggravated by INTRON A or other 646 alpha interferons. Recurrence of respiratory failure has been observed with interferon 647 rechallenge. The etiologic explanation for these pulmonary findings has yet to be 648 established. Any patient developing fever, cough, dyspnea, or other respiratory 649 symptoms should have a chest X-ray taken. If the chest X-ray shows pulmonary 650 infiltrates or there is evidence of pulmonary function impairment, the patient should be 651 closely monitored, and, if appropriate, interferon alpha treatment should be 652 discontinued. While this has been reported more often in patients with chronic hepatitis 653 C treated with interferon alpha, it has also been reported in patients with oncologic 654 diseases treated with interferon alpha.

655

656 Autoimmune Disorders

657 Rare cases of autoimmune diseases including thrombocytopenia, vasculitis, 658 Ravnaud's phenomenon, rheumatoid arthritis, lupus erythematosus, and 659 rhabdomyolysis have been observed in patients treated with alpha interferons, including 660 patients treated with INTRON A. In very rare cases the event resulted in fatality. The 661 mechanism by which these events developed and their relationship to interferon alpha 662 therapy is not clear. Any patient developing an autoimmune disorder during treatment 663 should be closely monitored and, if appropriate, treatment should be discontinued.

665 Human Albumin

The powder formulations of this product contain albumin, a derivative of human
blood. Based on effective donor screening and product manufacturing processes, it
carries an extremely remote risk for transmission of viral diseases. A theoretical risk for
transmission of Creutzfeldt-Jakob disease (CJD) also is considered extremely remote.
No cases of transmission of viral diseases or CJD have ever been identified for albumin.

671

664

672 AIDS-Related Kaposi's Sarcoma INTRON A therapy should not be used for patients 673 with rapidly progressive visceral disease (see CLINICAL PHARMACOLOGY). Also of 674 note, there may be synergistic adverse effects between INTRON A and zidovudine. 675 Patients receiving concomitant zidovudine have had a higher incidence of neutropenia 676 than that expected with zidovudine alone. Careful monitoring of the WBC count is 677 indicated in all patients who are myelosuppressed and in all patients receiving other 678 myelosuppressive medications. The effects of INTRON A when combined with other 679 drugs used in the treatment of AIDS-related disease are unknown.

680

681 Chronic Hepatitis C and Chronic Hepatitis B Patients with decompensated liver
 682 disease, autoimmune hepatitis or a history of autoimmune disease, and patients who
 683 are immunosuppressed transplant recipients should not be treated with INTRON A.
 684 There are reports of worsening liver disease, including jaundice, hepatic

685 encephalopathy, hepatic failure, and death following INTRON A therapy in such 686 patients. Therapy should be discontinued for any patient developing signs and 687 symptoms of liver failure.

688 Chronic hepatitis B patients with evidence of decreasing hepatic synthetic 689 functions, such as decreasing albumin levels or prolongation of prothrombin time, who 690 nevertheless meet the entry criteria to start therapy, may be at increased risk of clinical 691 decompensation if a flare of aminotransferases occurs during INTRON A treatment. In such patients, if increases in ALT occur during INTRON A therapy for chronic hepatitis 692 693 B, they should be followed carefully, including close monitoring of clinical 694 symptomatology and liver function tests including ALT, prothrombin time, alkaline 695 phosphatase, albumin, and bilirubin. In considering these patients for INTRON A 696 therapy, the potential risks must be evaluated against the potential benefits of 697 treatment.

698

699 **Peripheral Neuropathy**

Peripheral neuropathy has been reported when alpha interferons were given in combination with telbivudine. In one clinical trial, an increased risk and severity of peripheral neuropathy was observed with the combination use of telbivudine and pegylated interferon alfa-2a as compared to telbivudine alone. The safety and efficacy of telbivudine in combination with interferons for the treatment of chronic hepatitis B has not been demonstrated.

706

Use with Ribavirin (see also REBETOL[®] prescribing information) REBETOL may
 cause birth defects and/or death of the unborn child. REBETOL therapy should not be
 started until a report of a negative pregnancy test has been obtained immediately prior
 to planned initiation of therapy. Patients should use at least two forms of contraception
 and have monthly pregnancy tests (see CONTRAINDICATIONS and PRECAUTIONS,
 Information for Patients).

713

Combination treatment with INTRON A and REBETOL was associated with hemolytic anemia. Hemoglobin less than 10 g/dL was observed in approximately 10% of adult and pediatric patients in clinical trials. Anemia occurred within 1 to 2 weeks of initiation of ribavirin therapy. Combination treatment with INTRON A and REBETOL should **not** be used in patients with creatinine clearance less than 50 mL/min. See REBETOL prescribing information for additional information.

720

721 **PRECAUTIONS**

General Acute serious hypersensitivity reactions (e.g., urticaria, angioedema, bronchoconstriction, anaphylaxis) have been observed rarely in INTRON[®] A-treated patients; if such an acute reaction develops, the drug should be discontinued immediately and appropriate medical therapy instituted. Transient rashes have occurred in some patients following injection, but have not necessitated treatment interruption.

728 While fever may be related to the flu-like syndrome reported commonly in 729 patients treated with interferon, other causes of persistent fever should be ruled out.

PAGE 19

There have been reports of interferon, including INTRON A, exacerbating
preexisting psoriasis and sarcoidosis as well as development of new sarcoidosis.
Therefore, INTRON A therapy should be used in these patients only if the potential
benefit justifies the potential risk.

Variations in dosage, routes of administration, and adverse reactions exist among different brands of interferon. Therefore, do not use different brands of interferon in any single treatment regimen.

737

Triglycerides Elevated triglyceride levels have been observed in patients treated with
interferons, including INTRON A therapy. Elevated triglyceride levels should be
managed as clinically appropriate. Hypertriglyceridemia may result in pancreatitis.
Discontinuation of INTRON A therapy should be considered for patients with
persistently elevated triglycerides (e.g., triglycerides greater than 1000 mg/dL)
associated with symptoms of potential pancreatitis, such as abdominal pain, nausea, or
vomiting.

746 **Drug Interactions** Interactions between INTRON A and other drugs have not been fully
747 evaluated. Caution should be exercised when administering INTRON A therapy in
748 combination with other potentially myelosuppressive agents such as zidovudine.
749 Concomitant use of alpha interferon and theophylline decreases theophylline clearance,
750 resulting in a 100% increase in serum theophylline levels.

751

745

Information for Patients Patients receiving INTRON A alone or in combination with
 REBETOL[®] should be informed of the risks and benefits associated with treatment and
 should be instructed on proper use of the product. To supplement your discussion with
 a patient, you may wish to provide patients with a copy of the MEDICATION GUIDE.

756 Patients should be informed of, and advised to seek medical attention for, 757 symptoms indicative of serious adverse reactions associated with this product. Such 758 adverse reactions may include depression (suicidal ideation), cardiovascular (chest 759 pain), ophthalmologic toxicity (decrease in/or loss of vision), pancreatitis or colitis 760 (severe abdominal pain), and cytopenias (high persistent fevers, bruising, dyspnea). 761 Patients should be advised that some side effects such as fatigue and decreased 762 concentration might interfere with the ability to perform certain tasks. Patients who are 763 taking INTRON A in combination with REBETOL must be thoroughly informed of the 764 risks to a fetus. Female patients and female partners of male patients must be told to 765 use two forms of birth control during treatment and for six months after therapy is 766 discontinued (see **MEDICATION GUIDE**).

Patients should be advised to remain well hydrated during the initial stages of
treatment and that use of an antipyretic may ameliorate some of the flu-like symptoms.

If a decision is made to allow a patient to self-administer INTRON A, they should be instructed, based on their treatment, if they should inject a dose of INTRON[®] A subcutaneously or intramuscularly. If it is too difficult for them to inject themselves, they should be instructed to ask someone who has been trained to give the injection to them. Patients should be instructed on the importance of site selection for self-administering the injection, as well as the importance on rotating the injection sites. A puncture

resistant container for the disposal of needles and syringes should be supplied.
Patients self-administering INTRON A should be instructed on the proper disposal of
needles and syringes and cautioned against reuse.

780 Dental and Periodontal Disorders Dental and periodontal disorders have been 781 reported in patients receiving ribavirin and interferon combination therapy. In addition, 782 dry mouth could have a damaging effect on teeth and mucous membranes of the mouth 783 during long-term treatment with the combination of REBETOL and interferon alfa-2b. 784 Patients should brush their teeth thoroughly twice daily and have regular dental 785 examinations. In addition, some patients may experience vomiting. If this reaction 786 occurs, they should be advised to rinse out their mouth thoroughly afterwards.

787

779

Laboratory Tests In addition to those tests normally required for monitoring patients,
 the following laboratory tests are recommended for all patients on INTRON A therapy,
 prior to beginning treatment and then periodically thereafter.

791 792

793

794

795

- Standard hematologic tests including hemoglobin, complete and differential white blood cell counts, and platelet count.
 - Blood chemistries electrolytes, liver function tests, and TSH.

Those patients who have preexisting cardiac abnormalities and/or are in advanced stages of cancer should have electrocardiograms taken prior to and during the course of treatment.

Mild-to-moderate leukopenia and elevated serum liver enzyme (SGOT) levels have been reported with intralesional administration of INTRON A (see **ADVERSE REACTIONS**); therefore, the monitoring of these laboratory parameters should be considered.

803 Baseline chest X-rays are suggested and should be repeated if clinically 804 indicated.

805 For malignant melanoma patients, differential WBC count and liver function tests 806 should be monitored weekly during the induction phase of therapy and monthly during 807 the maintenance phase of therapy.

808 For specific recommendations in chronic hepatitis C and chronic hepatitis B, see 809 **INDICATIONS AND USAGE**.

810

811 **Carcinogenesis, Mutagenesis, Impairment of Fertility** Studies with INTRON A have 812 not been performed to determine carcinogenicity.

813 Interferon may impair fertility. In studies of interferon administration in nonhuman 814 primates, menstrual cycle abnormalities have been observed. Decreases in serum 815 estradiol and progesterone concentrations have been reported in women treated with 816 human leukocyte interferon.¹² Therefore, fertile women should not receive INTRON A 817 therapy unless they are using effective contraception during the therapy period. 818 INTRON A therapy should be used with caution in fertile men.

819 Mutagenicity studies have demonstrated that INTRON A is not mutagenic.

820 Studies in mice (0.1, 1.0 million IU/day), rats (4, 20, 100 million IU/kg/day), and 821 cynomolgus monkeys (1.1 million IU/kg/day; 0.25, 0.75, 2.5 million IU/kg/day) injected

PAGE 21

with INTRON A for up to 9 days, 3 months, and 1 month, respectively, have revealed no
evidence of toxicity. However, in cynomolgus monkeys (4, 20, 100 million IU/kg/day)
injected daily for 3 months with INTRON A, toxicity was observed at the mid and high
doses and mortality was observed at the high dose.

826 However, due to the known species-specificity of interferon, the effects in 827 animals are unlikely to be predictive of those in man.

828 INTRON A in combination with REBETOL should be used with caution in fertile 829 men. See the REBETOL prescribing information for additional information.

Pregnancy Category C INTRON A has been shown to have abortifacient effects in Macaca mulatta (rhesus monkeys) at 15 and 30 million IU/kg (estimated human equivalent of 5 and 10 million IU/kg, based on body surface area adjustment for a 60-kg adult). There are no adequate and well-controlled studies in pregnant women. INTRON A therapy should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

838 Pregnancy Category X applies to combination treatment with INTRON A and 839 REBETOL (see CONTRAINDICATIONS). See REBETOL prescribing information for additional information. Significant teratogenic and/or embryocidal effects have been 841 demonstrated in all animal species exposed to ribavirin. REBETOL therapy is 842 contraindicated in women who are pregnant and in the male partners of women who are 843 pregnant. See CONTRAINDICATIONS and the REBETOL prescribing information.

844

830

837

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

850

Nursing Mothers It is not known whether this drug is excreted in human milk.
However, studies in mice have shown that mouse interferons are excreted into the milk.
Because of the potential for serious adverse reactions from the drug in nursing infants,
a decision should be made whether to discontinue nursing or to discontinue INTRON A
therapy, taking into account the importance of the drug to the mother.

856

857 Pediatric Use General Safety and effectiveness in pediatric patients have not been
 858 established for indications other than chronic hepatitis B and chronic hepatitis C.

Chronic Hepatitis B Safety and effectiveness in pediatric patients ranging in age from
 1 to 17 years have been established based upon one controlled clinical trial (see
 CLINICAL PHARMACOLOGY, INDICATIONS AND USAGE, and DOSAGE AND
 ADMINISTRATION, Chronic Hepatitis B Pediatrics).

Chronic Hepatitis C Safety and effectiveness in pediatric patients ranging in age from
 3 to 16 years have been established based upon clinical studies in 118 patients. See
 REBETOL prescribing information for additional information. Suicidal ideation or
 attempts occurred more frequently among pediatric patients compared to adult patients
 (2.4% versus 1%) during treatment and off-therapy follow-up (see WARNINGS,

PAGE 22

868 **Neuropsychiatric Disorders**). During a 48-week course of therapy there was a 869 decrease in the rate of linear growth (mean percentile assignment decrease of 7%) and 870 a decrease in the rate of weight gain (mean percentile assignment decrease of 9%). A 871 general reversal of these trends was noted during the 24-week post-treatment period.

Long-term data in a limited number of patients suggests that combination therapy
may induce a growth inhibition that results in reduced final adult height in some patients
(see ADVERSE REACTIONS, Chronic Hepatitis C Pediatrics).

875

894

Geriatric Use In all clinical studies of INTRON A, including studies as monotherapy and in combination with REBETOL (ribavirin USP) Capsules, only a small percentage of the subjects were aged 65 and over. These numbers were too few to determine if they respond differently from younger subjects except for the clinical trials of INTRON A in combination with REBETOL, where elderly subjects had a higher frequency of anemia (67%) than did younger patients (28%).

In a database consisting of clinical study and postmarketing reports for various
 indications, cardiovascular adverse events and confusion were reported more frequently
 in elderly patients receiving INTRON A therapy compared to younger patients.

885 In general, INTRON A therapy should be administered to elderly patients 886 cautiously, reflecting the greater frequency of decreased hepatic, renal, bone marrow, 887 and/or cardiac function and concomitant disease or other drug therapy. INTRON A is 888 known to be substantially excreted by the kidney, and the risk of adverse reactions to 889 INTRON A may be greater in patients with impaired renal function. Because elderly 890 patients often have decreased renal function, patients should be carefully monitored during treatment, and dose adjustments made based on symptoms and/or laboratory 891 CLINICAL PHARMACOLOGY DOSAGE 892 abnormalities (see and AND 893 ADMINISTRATION).

895 ADVERSE REACTIONS

General The adverse experiences listed below were reported to be possibly or probably related to INTRON[®] A therapy during clinical trials. Most of these adverse reactions were mild to moderate in severity and were manageable. Some were transient and most diminished with continued therapy.

The most frequently reported adverse reactions were "flu-like" symptoms, particularly fever, headache, chills, myalgia, and fatigue. More severe toxicities are observed generally at higher doses and may be difficult for patients to tolerate.

	TREA	ATMENT-RELA	TED ADVERSE EXPL	ERIENCES	S BY INC	DICATION			
			<u>Dosing Re</u>	<u>gimens</u>					
			Percentage (%)	of Patient	<u>s*</u>				
MALIGNANT MELANOMA	FOLLICULAR LYMPHOMA	HAIRY CELL LEUKEMIA	CONDYLOMATA ACUMINATA	AID RELA KAPC SARC	9S- ATED OSI'S OMA	CHRONIC HEPATITIS C		CHRO HEPATI	NIC TIS B
							Adı	ults	Pediatrics
<u>20 MIU/m²</u> <u>Induction (IV)</u> <u>10 MIU/m²</u> <u>Maintenance</u> (SC)	<u>5 MIU</u> <u>TIW/SC</u>	<u>2 MIU/m²</u> <u>TIW/SC</u>	<u>1</u> <u>MIU/lesion</u>	<u>30</u> <u>MIU/m</u> <u>TIW/S</u> <u>C</u>	<u>35</u> <u>MIU</u> <u>QD/S</u> <u>C</u>	<u>3</u> <u>MIU</u> <u>TIW</u>	<u>5</u> <u>MIU</u> QD	<u>10</u> <u>MIU</u> <u>TIW</u>	<u>6</u> <u>MIU/m²</u> <u>TIW</u>

PAGE 23

ADVERSE EXPERIENCE	N=143	N=135	N=145	N=352	N=74	N=29	N=183	N=101	N=78	N=116
Application-Site Disorders			20							
injection site inflammation		1					5	3		
other (≤5%)	burning, injection	on site bleeding, i	njection site pain,	injection site reac	tion (5% in	chronic he	patitis B pedi	atrics), itch	ing	
Blood Disorders (<5%)	anemia, anemia 14% in chronic thrombocytope	a hypochromic, gi hepatitis B pedia nia purpura	ranulocytopenia, h trics), thrombocyto	emolytic anemia, ppenia (10% in chi	leukopenia ronic hepati	, lymphocy tis C) (blee	tosis, neutro eding 8% in n	penia (9% i nalignant m	n chronic elanoma)	hepatitis C, ,
Body as a Whole	, , , , , , , , , , , , , , , , , , ,	• •								
facial edema		1		<1		10	<1	3	1	<1
weight decrease	3	13	<1	<1	5	3	10	2	5	3
other (≤5%)	allergic reaction nonspecific, lyn follicular lymph	n, cachexia, dehy nphadenitis, lymp oma), phlebitis su	dration, earache, l hadenopathy, mas perficial, scrotal/p	nernia, edema, hy stitis, periorbital eo enile edema, thirs	percalcemia dema, poor t, weakness	a, hypergly peripheral s, weight ir	cemia, hypo circulation, p crease	thermia, inf peripheral e	lammatio dema (6%	n 6 in
Cardiovascular System Disorders (<5%)	angina, arrhyth extrasystoles, h hypotension, p	mia, atrial fibrillati neart valve disord ulmonary embolis	on, bradycardia, c er, hematoma, hy m, Raynaud's dise	ardiac failure, car pertension (9% in ease, tachycardia,	diomegaly, chronic hep thrombosis	cardiomyo batitis C), h s, varicose	pathy, coron ypotension, vein	ary artery operations	lisorder, , phlebitis	s, postural
Endocrine System Disorders (<5%)	aggravation of	diabetes mellitus,	goiter, gynecoma	stia, hyperglycem	ia, hyperthy	vroidism, h	ypertriglyceri	demia, hyp	othyroidis	m, virilism
Flu-like Symptoms										
fever	81	56	68	56	47	55	34	66	86	94
headache	62	21	39	47	36	21	43	61	44	57
chills	54		46	45						
myalgia	75	16	39	44	34	28	43	59	40	27
fatigue	96	8	61	18	84	48	23	75	69	71
increased sweating	6	13	8	2	4	21	4	1	1	3
asthenia		63	7		11		40	5	15	5
rigors	2	7			30	14	16	38	42	30
arthralgia	6	8	8	9		3	16	19	8	15
dizziness	23		12	9	7	24	9	13	10	8
influenza-like symptoms	10	18	37		45	79	26	5		<1
back pain		15	19	6	1	3				
dry mouth	1	2	19		22	28	5	6	5	
chest pain	2	8	<1	<1	1	28	4	4		
malaise	6			14	5		13	9	6	3
pain	15	9	18	3	3	3				
(unspecified)	choct pain cub	stornal hyporthor	mia rhinitia rhinoi	rhoo						
	chest pain subs	sternal, hyperther	inia, minius, minu	IIIea						
System Disorders										
diarrhea	35	19	18	2	18	45	13	19	8	12
anorexia	69	21	19	1	38	41	14	43	53	43
nausea	66	24	21	17	28	21	19	50	33	18
taste alteration	24	2	13	<1	5	7	2	10		
abdominal pain	2	20	<5	1	5	21	16	5	4	23
loose stools		1		<1		10	2	2		2
vomiting	t	32	6	2	11	14	8	7	10	27
constipation	1	14	<1		1	10	4	5		2
gingivitis	2 [‡]	7 [‡]				14		1		
dyspepsia		2		2	4		7	3	8	3
other (<5%)	abdominal asci gastroenteritis, discoloration, g melena, mouth stomatitis ulcer	tes, abdominal dia gastrointestinal di ingival bleeding, ulceration, muco ative, taste loss, t	stension, colitis, d lisorder (7% in foll gum hyperplasia, l sitis, oral hemorrh ongue disorder, to	ysphagia, eructati icular lymphoma), halitosis, hemorrh age, oral leukopla ooth disorder	on, esophag gastrointes oids, increa kia, rectal b	gitis, flatule tinal hemo sed appeti leeding aff	ence, gallstor prrhage, gast te, increasec ter stool, rect	nes, gastric rointestinal I saliva, inte al hemorrh	ulcer, ga mucosal estinal dis age, stom	stritis, order, atitis,
Liver and Biliary System Disorders (<5%)	abnormal hepa (SGOT/SGPT) (15% in chronic	tic function tests, (elevated SGOT c hepatitis C), and	biliary pain, bilirub 63% in malignant I very rarely, hepa	inemia, hepatitis, melanoma and 24 tic encephalopath	increased I 1% in follicu y, hepatic fa	actate deh lar lympho ailure, and	ydrogenase, ma), jaundice death	increased e, right upp	transamir er quadra	ases nt pain
<u>Musculoskeletal</u> System Disorders										

PAGE 24

I REATMENT-RELATED A	ADVERSE EXPERIENCES B	

	Dosing Regimens									
				Percentage (%)	of Patient	<u>s*</u>				
	MALIGNANT MELANOMA	FOLLICULAR LYMPHOMA	HAIRY CELL LEUKEMIA	CONDYLOMATA ACUMINATA	AID RELA KAPC SARC	9S- TED 9SI'S OMA	CHRONIC HEPATITIS C [∥]		CHRO HEPATI	NIC TIS B
	221							Adı	ılts	Pediatrics
	<u>20 MIU/m</u> Induction (IV) <u>10 MIU/m²</u> <u>Maintenance</u> <u>(SC)</u>	<u>5 MIU</u> <u>TIW/SC</u>	<u>2 MIU/m²</u> <u>TIW/SC</u>	<u>1</u> <u>MIU/lesion</u>	<u>30</u> <u>MIU/m</u> <u>TIW/S</u> <u>C</u>	<u>35</u> <u>MIU</u> <u>QD/S</u> <u>C</u>	<u>3</u> <u>MIU</u> <u>TIW</u>	<u>5</u> <u>MIU</u> QD	<u>10</u> <u>MIU</u> <u>TIW</u>	<u>6</u> <u>MIU/m²</u> <u>TIW</u>
ADVERSE EXPERIENCE	N=143	N=135	N=145	N=352	N=74	N=29	N=183	N=101	N=78	N=116
musculoskeletal pain		18					21	9	1	10
other (<5%)	arteritis, arthritis atrophy, muscle	s, arthritis aggrava weakness, polya	ated, arthrosis, l arteritis nodosa,	oone disorder, bone p tendinitis, rheumatoio	ain, carpa d arthritis, s	l tunnel s spondylit	syndrome, hypo is	oreflexia, l	eg cramp	os, muscle
<u>Nervous System</u> <u>and Psychiatric</u> Disorders	1 27			;						
depression	40	9	6	3	9	28	19	17	6	4
paresthesia	13	13	6	1	3	21	5	6	3	<1
impaired		1		<1	3	14	3	8	5	3
amnesia	§	1	<5			14				
confusion	8	2	<5	1	12	10	1			2
bypoestbesia	0	1	<5	1	12	10				2
irritobility		1	<5	I		10		16	10	
initability	1	1					13 22¶	10	12	22
sommolence	1	2	<5	3	3		33	14	9	5
anxiety	1	9	Э	<1	1	3	5	2		3
insomnia	5	4		<1	3	3	12	11	6	8
nervousness	1	1		1		3	2	3		3
decreased libido	1	1	<5				1	5	1	
other (<5%)	abnormal coord (7% in chronic h delirium, dyspho flashes, hypere: manic depressio disorder, polyne vertigo (8% in fo	ination, abnormal hepatitis B pediatr onia, emotional la sthesia, hyperkine on, manic reactior europathy, psycho ollicular lymphoma	dreaming, abn ics), alcohol into bility, extrapyra esia, hypertonia n, migraine, neu sis, speech dis a)	ormal gait, abnormal i olerance, apathy, aph midal disorder, feeling , hypokinesia, impaire ıralgia, neuritis, neuro order, stroke, suicidal	thinking, a lasia, ataxi g of ebriety ed conscio pathy, neu l ideation, s	ggravate a, Bell's v, flushing usness, l urosis, pa suicide a	d depression, a palsy, CNS dys g, hearing disor labyrinthine dis aresis, paroniria ttempt, syncop	aggressive function, der, heari order, loss , parosmi e, tinnitus	e reaction coma, co ng impai s of cons a, person , tremor,	n, agitation prvulsions, rment, hot ciousness, nality twitching,
Reproduction System Disorders (<5%)	amenorrhea (12 penis disorder, s	2% in follicular lym sexual dysfunctio	nphoma), dysme n, uterine bleed	enorrhea, impotence, ing, vaginal dryness	leukorrhea	a, menor	rhagia, menstru	al irregula	arity, pelv	/ic pain,
<u>Resistance</u> <u>Mechanism</u> <u>Disorders</u>										
moniliasis		1		<1		17				
herpes simplex	1	2		1		3	1	5		
other (<5%)	abscess, conjur lymphoma), infe hepatitis C)	nctivitis, fungal inf action parasitic, ot	ection, hemoph itis media, seps	ilus, herpes zoster, in sis, stye, trichomonas,	fection, inf , upper res	ection ba	acterial, infectio tract infection, v	n nonspe viral infect	cific (7% ion (7% i	in follicular n chronic
<u>Respiratory</u> System Disorders										
dyspnea	15	14	<1		1	34	3	5		
coughing	6	13	<1			31	1	4		5
pharyngitis	2	8	<5	1	1	31	3	7	1	7
sinusitis	1	4				21	2			
nonproductive coughing	2	7				14	0	1		
nasal congestion	1	7		1		10	<1	4		
- other (≤5%)	asthma, bronch	itis (10% in follicu	lar lymphoma),	bronchospasm, cyan	osis, epist	axis (7%	in chronic hepa	atitis B pe	diatrics),	hemoptysis,

PAGE 25

TREATMENT-RELATED ADVERSE EXPERIENCES BY INDICATION

	Dosing Regimens									
				Percentage (%)	of Patient	<u>'S*</u>				
	MALIGNANT MELANOMA	FOLLICULAR LYMPHOMA	HAIRY CELL LEUKEMIA	CONDYLOMATA ACUMINATA	AID RELA KAPC SARC	OS- ATED OSI'S COMA	CHRONIC HEPATITIS C	I	CHRO HEPATI	NIC FIS B
								Adu	Ilts	Pediatrics
	<u>20 MIU/m²</u> <u>Induction (IV)</u> <u>10 MIU/m²</u> <u>Maintenance</u> <u>(SC)</u>	<u>5 MIU</u> <u>TIW/SC</u>	<u>2 MIU/m²</u> <u>TIW/SC</u>	<u>1</u> <u>MIU/lesion</u>	<u>30</u> <u>MIU/m</u> <u>TIW/S</u>	<u>35</u> <u>MIU</u> <u>QD/S</u> <u>C</u>	<u>3</u> <u>MIU</u> <u>TIW</u>	<u>5</u> <u>MIU</u> QD	<u>10</u> <u>MIU</u> <u>TIW</u>	<u>6</u> <u>MIU/m²</u> <u>TIW</u>
ADVERSE EXPERIENCE	N=143	N=135	N=145	N=352	<u>0</u> N=74	N=29	N=183	N=101	N=78	N=116
	hypoventilation, respiratory diso	, laryngitis, lung fil rder, respiratory i	orosis, pleural e nsufficiency, sn	ffusion, orthopnea, pl eezing, tonsillitis, trac	eural pain heitis, whe	, pneumo eezing	onia, pneumonit	is, pneum	othorax,	rales,
<u>Skin and</u> <u>Appendages</u> <u>Disorders</u>										
dermatitis	1		8				2	1		
alopecia	29	23	8		12	31	28	26	38	17
pruritus		10	11	1	7		9	6	4	3
rash	19	13	25		9	10	5	8	1	5
dry skin	1	3	9		9	10	4	3		<1
other (<5%) abnormal hair texture, acne, cellulitis, cyanosis of the hand, cold and clammy skin, dermatitis lichenoides, eczema, epidermal necrolysis, erythema, erythema nodosum, folliculitis, furunculosis, increased hair growth, lacrimal gland disorder, lacrimation, lipoma, maculopapular rash, melanosis, nail disorders, nonherpetic cold sores, pallor, peripheral ischemia, photosensitivity, pruritus genital, psoriasis, psoriasis aggravated, purpura (5% in chronic hepatitis C), rash erythematous, sebaceous cyst, skin depigmentation, skin discoloration, skin nodule, urticaria, vitiligo										
Urinary System Disorders (<5%)	albumin/protein polyuria (10% ir	in urine, cystitis, n follicular lympho	dysuria, hematu ma), renal insul	uria, incontinence, inc fficiency, urinary tract	reased BL infection (JN, mictu 5% in ch	rition disorder, ronic hepatitis (micturitior C)	n frequer	cy, nocturia,
Vision Disorders (<5%)	abnormal vision	n, blurred vision, d	iplopia, dry eye	s, eye pain, nystagmu	us, photop	hobia				
* Dash () indi	cates not reported									
† \/	where a standard state is a set									

[†] Vomiting was reported with nausea as a single term

[‡] Includes stomatitis/mucositis

§ Amnesia was reported with confusion as a single term

Percentages based upon a summary of all adverse events during 18 to 24 months of treatment

Predominantly lethargy

PAGE 26

Hairy Cell Leukemia The adverse reactions most frequently reported during clinical
trials in 145 patients with hairy cell leukemia were the "flu-like" symptoms of fever
(68%), fatigue (61%), and chills (46%).

907

908 Malignant Melanoma The INTRON A dose was modified because of adverse events in 909 65% (n=93) of the patients. INTRON A therapy was discontinued because of adverse 910 events in 8% of the patients during induction and 18% of the patients during 911 maintenance. The most frequently reported adverse reaction was fatigue, which was 912 observed in 96% of patients. Other adverse reactions that were recorded in greater 913 than 20% of INTRON A-treated patients included neutropenia (92%), fever (81%), 914 myalgia (75%), anorexia (69%), vomiting/nausea (66%), increased SGOT (63%), 915 headache (62%), chills (54%), depression (40%), diarrhea (35%), alopecia (29%), 916 altered taste sensation (24%), dizziness/vertigo (23%), and anemia (22%).

917 Adverse reactions classified as severe or life threatening (ECOG Toxicity Criteria 918 grade 3 or 4) were recorded in 66% and 14% of INTRON A-treated patients. 919 respectively. Severe adverse reactions recorded in greater than 10% of INTRON A-920 treated patients included neutropenia/leukopenia (26%), fatigue (23%), fever (18%), 921 myalgia (17%), headache (17%), chills (16%), and increased SGOT (14%). Grade 4 922 fatigue was recorded in 4% and grade 4 depression was recorded in 2% of INTRON A-923 treated patients. No other grade 4 AE was reported in more than 2 INTRON A-treated 924 patients. Lethal hepatotoxicity occurred in 2 INTRON A-treated patients early in the 925 clinical trial. No subsequent lethal hepatotoxicities were observed with adequate 926 monitoring of liver function tests (see PRECAUTIONS, Laboratory Tests).

927

928 Follicular Lymphoma Ninety-six percent of patients treated with CHVP plus INTRON A 929 therapy and 91% of patients treated with CHVP alone reported an adverse event of any 930 severity. Asthenia, fever, neutropenia, increased hepatic enzymes, alopecia, headache, 931 anorexia, "flu-like" symptoms, myalgia, dyspnea, thrombocytopenia, paresthesia, and 932 polyuria occurred more frequently in the CHVP plus INTRON A-treated patients than in patients treated with CHVP alone. Adverse reactions classified as severe or life 933 934 threatening (World Health Organization grade 3 or 4) recorded in greater than 5% of 935 CHVP plus INTRON A-treated patients included neutropenia (34%), asthenia (10%), 936 and vomiting (10%). The incidence of neutropenic infection was 6% in CHVP plus 937 INTRON A versus 2% in CHVP alone. One patient in each treatment group required 938 hospitalization.

939 Twenty-eight percent of CHVP plus INTRON A-treated patients had a temporary 940 modification/interruption of their INTRON A therapy, but only 13 patients (10%) 941 permanently stopped INTRON A therapy because of toxicity. There were four deaths 942 on study; two patients committed suicide in the CHVP plus INTRON A arm and two 943 patients in the CHVP arm had unwitnessed sudden death. Three patients with hepatitis 944 B (one of whom also had alcoholic cirrhosis) developed hepatotoxicity leading to 945 discontinuation of INTRON A. Other reasons for discontinuation included intolerable 946 asthenia (5/135), severe flu symptoms (2/135), and one patient each with exacerbation 947 of ankylosing spondylitis, psychosis, and decreased ejection fraction.

949 **Condylomata Acuminata** Eighty-eight percent (311/352) of patients treated with 950 INTRON A for condylomata acuminata who were evaluable for safety reported an 951 adverse reaction during treatment. The incidence of the adverse reactions reported 952 increased when the number of treated lesions increased from one to five. All 40 953 patients who had five warts treated reported some type of adverse reaction during 954 treatment.

Adverse reactions and abnormal laboratory test values reported by patients who were re-treated were qualitatively and quantitatively similar to those reported during the initial INTRON A treatment period.

958

AIDS-Related Kaposi's Sarcoma In patients with AIDS-Related Kaposi's Sarcoma,
 some type of adverse reaction occurred in 100% of the 74 patients treated with 30
 million IU/m² three times a week and in 97% of the 29 patients treated with 35 million IU
 per day.

963 Of these adverse reactions, those classified as severe (World Health 964 Organization grade 3 or 4) were reported in 27% to 55% of patients. Severe adverse reactions in the 30 million IU/m² TIW study included: fatigue (20%), influenza-like 965 966 symptoms (15%), anorexia (12%), dry mouth (4%), headache (4%), confusion (3%), 967 fever (3%), myalgia (3%), and nausea and vomiting (1% each). Severe adverse 968 reactions for patients who received the 35 million IU QD included: fever (24%), fatigue 969 (17%), influenza-like symptoms (14%), dyspnea (14%), headache (10%), pharyngitis 970 (7%), and ataxia, confusion, dysphagia, GI hemorrhage, abnormal hepatic function, 971 increased SGOT, myalgia, cardiomyopathy, face edema, depression, emotional lability, 972 suicide attempt, chest pain, and coughing (1 patient each). Overall, the incidence of 973 severe toxicity was higher among patients who received the 35 million IU per day dose.

974

975 Chronic Hepatitis C Adults Two studies of extended treatment (18-24 months) with 976 INTRON A show that approximately 95% of all patients treated experience some type of 977 adverse event and that patients treated for extended duration continue to experience 978 adverse events throughout treatment. Most adverse events reported are mild to 979 moderate in severity. However, 29/152 (19%) of patients treated for 18 to 24 months 980 experienced a serious adverse event compared to 11/163 (7%) of those treated for 6 981 months. Adverse events which occur or persist during extended treatment are similar in 982 type and severity to those occurring during short-course therapy.

983 Of the patients achieving a complete response after 6 months of therapy, 12/79 984 (15%) subsequently discontinued INTRON A treatment during extended therapy 985 because of adverse events, and 23/79 (29%) experienced severe adverse events 986 (WHO grade 3 or 4) during extended therapy.

In patients using combination treatment with INTRON A and REBETOL, the
 primary toxicity observed was hemolytic anemia. Reductions in hemoglobin levels
 occurred within the first 1 to 2 weeks of therapy. Cardiac and pulmonary events
 associated with anemia occurred in approximately 10% of patients treated with INTRON
 A/REBETOL therapy. See REBETOL prescribing information for additional information.

993 **Chronic Hepatitis C** *Pediatrics* In pediatric patients with chronic hepatitis C treated 994 with INTRON A 3 MIU/m² three times weekly and REBETOL 15 mg/kg per day, all

995 subjects (n=118) had at least one adverse event during 24-48 weeks of treatment, of 996 which 80% were considered to be mild or moderate in severity. Six percent discontinued 997 therapy due to adverse reactions and dose modifications were required in 30% of 998 subjects, most commonly for anemia and neutropenia. Adverse events occurring in 999 more than 50% of subjects included headache, fever, fatigue and anorexia. Adverse 1000 events occurring in 20-50% of subjects included influenza-like symptoms, abdominal 1001 pain, vomiting, nausea, myalgia, pharyngitis, diarrhea, viral infection, rigors, weight 1002 decrease, musculoskeletal pain, alopecia and dizziness. The most common laboratory 1003 test abnormalities were neutropenia (34%) and anemia (27%). Depression was reported 1004 in 13% (n=15) of children. Three of these subjects had suicidal ideation, and one 1005 attempted suicide. Weight loss and slowed growth are common in pediatric patients 1006 during combination therapy with INTRON A and REBETOL. Following treatment, 1007 rebound growth and weight gain occurred in most subjects. Long-term follow-up data in 1008 pediatric subjects, however, indicates that INTRON A in combination with REBETOL 1009 may induce a growth inhibition that results in reduced adult height in some patients (see 1010 PRECAUTIONS, Pediatric Use).

1011

1012 **Chronic Hepatitis B** *Adults* In patients with chronic hepatitis B, some type of adverse 1013 reaction occurred in 98% of the 101 patients treated at 5 million IU QD and 90% of the 1014 78 patients treated at 10 million IU TIW. Most of these adverse reactions were mild to 1015 moderate in severity, were manageable, and were reversible following the end of 1016 therapy.

Adverse reactions classified as severe (causing a significant interference with normal daily activities or clinical state) were reported in 21% to 44% of patients. The severe adverse reactions reported most frequently were the "flu-like" symptoms of fever (28%), fatigue (15%), headache (5%), myalgia (4%), rigors (4%), and other severe "flulike" symptoms, which occurred in 1% to 3% of patients. Other severe adverse reactions occurring in more than one patient were alopecia (8%), anorexia (6%), depression (3%), nausea (3%), and vomiting (2%).

1024 To manage side effects, the dose was reduced, or INTRON A therapy was 1025 interrupted in 25% to 38% of patients. Five percent of patients discontinued treatment 1026 due to adverse experiences. 1027

1028 **Chronic Hepatitis B** *Pediatrics* In pediatric patients with chronic hepatitis B (n=72) 1029 during 16-24 weeks of treatment, the most frequently reported adverse events were 1030 those commonly associated with interferon treatment: flu-like symptoms (100%), 1031 gastrointestinal system disorders (46%), and nausea and vomiting (40%). Neutropenia 1032 (13%) and thrombocytopenia (3%) were also reported. None of the adverse events was 1033 life threatening and most were moderate to severe and resolved upon dose reduction or 1034 drug discontinuation.

PAGE 29

ABNORMAL LABORATORY TEST VALUES BY INDICATION Dosing Regimens Percentage (%) of Patients

MALIGNANT FOLLICULAR HAIRY CELL CONDYLOMATA AIDS-RELATED CHRONIC CHRONIC MELANOMA LYMPHOMA LEUKEMIA ACUMINATA **KAPOSI'S SARCOMA HEPATITIS C HEPATITIS B** Adults Pediatrics 20 MIU/m² Induction (IV) <u>5 MIU</u> <u>2 MIU/m²</u> 1 <u>30 MIU/m²</u> <u>35</u> 3 5 10 <u>6</u> 10 MIU/m² TIW/SC TIW/SC MIU/lesion TIW/SC MIU <u>MIU</u> MIU MIU MIU/m² Maintenance QD/SC <u>TIW</u> QD <u>TIW</u> <u> TIW</u> (SC)Laboratory Tests N=143 N=135 N=352 N=69-73 N=26-28 N=75-103 N=113-115 N=145 N=140-171 N=96-101 26[¶] 32 23 17 Hemoglobin 22 8 NA ---1 15 26[†] 34[†] 9† White Blood Cell Count NA 17 22 68[†] ---10 15[‡] 12[‡] 5[‡] 1[‡] Platelet Count 15 13 NA 0 8 --Serum Creatinine 3 2 6 3 0 ---------0 3 13 8 0 Alkaline Phosphatase 4 ---4 ------------Lactate Dehydrogenase 1 0 -------------------Serum Urea Nitrogen 12 4 0 ------2 0 2 --SGOT 63 12 24 4 11 41 -----------SGPT 2 13 10 15 ----------------Granulocyte Count 92 36 45[§] 75[§] 61[§] 70[§] Total NA 39 ---31 ٠ 1000-<1500/mm³ 66 30 32 43 ---32 ٠ ------------750-<1000/mm³ 10 18 18 --21 24 • -----------500-<750/mm³ 25 17 9 7 ---1 ------------٠ <500/mm³ 1 13 2 2 2 ----------4 ٠

NA - Not Applicable - Patients' initial hematologic laboratory test values were abnormal due to their condition.

* Decrease of ≥2 g/dL

** Decrease of ≥2 g/dL; 14% 2-<3 g/dL; 3% ≥3 g/dL

[†] Decrease to <3000/mm³

[‡] Decrease to <70,000/mm³

§ Neutrophils plus bands

White Blood Cell Count was reported as neutropenia

[¶] Decrease of ≥2 g/dL; 20% 2-<3 g/dL; 6% ≥3 g/dL

PAGE 30

1035 Postmarketing Experience The following adverse reactions have been identified 1036 during postapproval use of INTRON A alone or in combination with REBETOL. Because 1037 these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to 1038 1039 drug exposure. 1040 1041 Blood and Lymphatic System Disorders 1042 pancytopenia (concurrent anemia, leukopenia, thrombocytopenia), aplastic 1043 anemia, pure red cell aplasia, thrombotic thrombocytopenic purpura, idiopathic 1044 thrombocytopenic purpura 1045 Ear and Labyrinth Disorders hearing loss 1046 1047 Endocrine Disorders 1048 hypopituitarism Eye Disorders 1049 Vogt-Koyanagi-Harada syndrome, serous retinal detachment 1050 1051 Gastrointestinal Disorders 1052 pancreatitis 1053 General Disorders and Administration Site Conditions 1054 asthenic conditions (including asthenia, malaise, fatigue) 1055 Immune System Disorders 1056 cases of acute hypersensitivity reactions, including anaphylaxis and angioedema, 1057 systemic lupus erythematosus, sarcoidosis or exacerbation of sarcoidosis 1058 Musculoskeletal and Connective Tissue Disorders 1059 mvositis 1060 Nervous System Disorders 1061 peripheral neuropathy Psychiatric Disorders 1062 1063 homicidal ideation, psychosis including hallucinations Renal and Urinary Disorders 1064 renal failure, renal insufficiency, nephrotic syndrome 1065 1066 Respiratory, Thoracic, and Mediastinal Disorders 1067 pulmonary hypertension 1068 Skin and Subcutaneous Tissue Disorders injection site necrosis, Stevens-Johnson syndrome, toxic epidermal necrolysis, 1069 1070 erythema multiforme, urticaria 1071 1072 **OVERDOSAGE** 1073 There is limited experience with overdosage. Postmarketing surveillance includes 1074 reports of patients receiving a single dose as great as 10 times the recommended dose. In general, the primary effects of an overdose are consistent with the effects seen with 1075 1076 therapeutic doses of interferon alfa-2b. Hepatic enzyme abnormalities, renal failure, hemorrhage, and myocardial infarction have been reported with single administration 1077 overdoses and/or with longer durations of treatment than prescribed (see ADVERSE 1078 1079 **REACTIONS**). Toxic effects after ingestion of interferon alfa-2b are not expected

PAGE 31

because interferons are poorly absorbed orally. Consultation with a poison center isrecommended.

1082

1083 **Treatment** There is no specific antidote for interferon alfa-2b. Hemodialysis and 1084 peritoneal dialysis are not considered effective for treatment of overdose.

1085

1086 DOSAGE AND ADMINISTRATION

1088 General

1089

1090 *IMPORTANT:* INTRON[®] A is supplied as 1) Powder for Injection/Reconstitution; 2) 1091 Solution for Injection in Vials; 3) Solution for Injection in Multidose Pens. Not all 1092 dosage forms and strengths are appropriate for some indications. It is important 1093 that you carefully read the instructions below for the indication you are treating to 1094 ensure you are using an appropriate dosage form and strength. 1095

1096 To enhance the tolerability of INTRON A, injections should be administered in the 1097 evening when possible.

1098

1099 To reduce the incidence of certain adverse reactions, acetaminophen may be 1100 administered at the time of injection.

- 1101
- 1102 1103

The solution should be allowed to come to room temperature before using.

1104 Hairy Cell Leukemia (see DOSAGE AND ADMINISTRATION, General) 1105

106 *Dose:* The recommended dose for the treatment of hairy cell leukemia is 2 million IU/m²
 107 administered intramuscularly or subcutaneously 3 times a week for up to 6 months.
 108 Patients with platelet counts of less than 50,000/mm³ should not be administered
 1109 INTRON A intramuscularly, but instead by subcutaneous administration. Patients who
 1109 are responding to therapy may benefit from continued treatment.

1111 1112

Dosage Forms for This Indication				
Dosage Form	Concentration	Route	Fixed Doses	
Powder 10 MIU (single dose)	10 MIU/mL	IM, SC	N/A	
Solution 18 MIU multidose	6 MIU/mL	IM, SC	N/A	
Solution 25 MIU multidose	10 MIU/mL	IM, SC	N/A	
Pen 3 MIU/dose multidose	15 MIU/mL	SC	1.5, 3.0, 4.5	
Pen 5 MIU/dose multidose	25 MIU/mL	SC	2.5, 5.0	

1113

1114 NOTE: INTRON A Powder for Injection does not contain a preservative. The vial 1115 must be discarded after reconstitution and withdrawal of a single dose.

- 1116
- 1117 Dose Adjustment:
- 1118
- If severe adverse reactions develop, the dosage should be modified (50% reduction)
 or therapy should be temporarily withheld until the adverse reactions abate and then
 resume at 50% (1 MIU/m² TIW).

PAGE 32

- If severe adverse reactions persist or recur following dosage adjustment, INTRON A should be permanently discontinued.
- INTRON A should be discontinued for progressive disease or failure to respond after six months of treatment.
- 1126
- 1127
- 1128

7 Malignant Melanoma (see DOSAGE AND ADMINISTRATION, General)

1129 INTRON A adjuvant treatment of malignant melanoma is given in two phases, induction
1130 and maintenance.
1131

Induction Recommended Dose: The recommended daily dose of INTRON A in induction is 20 million IU/m² as an intravenous infusion, over 20 minutes, 5 consecutive days per week, for 4 weeks (see **Dose Adjustment** below).

1135 1136

Dosage Forms for This Indication					
Dosage Form	Concentration	Route			
Powder 10 MIU	10 MIU/mL	IV			
Powder 18 MIU	18 MIU/mL	IV			
Powder 50 MIU	50 MIU/mL	IV			

1137

1138 NOTE: INTRON A Solution for Injection in vials or Multidose Pens is NOT 1139 recommended for intravenous administration and should not be used for the 1140 induction phase of malignant melanoma.

1142NOTE: INTRON A Powder for Injection does not contain a preservative. The vial1143must be discarded after reconstitution and withdrawal of a single dose.

1144

1141

1145 Dose Adjustment: NOTE: Regular laboratory testing should be performed to monitor
 1146 laboratory abnormalities for the purpose of dose modifications (see PRECAUTIONS,
 1147 Laboratory Tests).

- 1148
- INTRON A should be withheld for severe adverse reactions, including granulocyte counts greater than 250/mm³ but less than 500/mm³ or SGPT/SGOT greater than 5-10x upper limit of normal, until adverse reactions abate. INTRON A treatment should be restarted at 50% of the previous dose.
- 1153 INTRON A should be permanently discontinued for:
 - Toxicity that does not abate after withholding INTRON A
 - Severe adverse reactions which recur in patients receiving reduced doses of INTRON A
- Granulocyte count less than 250/mm³ or SGPT/SGOT of greater than 10x upper limit of normal
- 1159

1154 1155

1156

1160 *Maintenance Recommended Dose:* The recommended dose of INTRON A for 1161 maintenance is 10 million IU/m² as a subcutaneous injection three times per week for 1162 48 weeks (see *Dose Adjustment* below).

1163 1164

Dosage Forms for This Indication

PAGE 33

	Dosage Form	Concentration	Route	Fixed Doses			
	Powder 10 MIU (single dose)*	10 MIU/mL	SC	N/A			
	Powder 18 MIU (single dose)**	18 MIU/mL	SC	N/A			
	Solution 18 MIU multidose	6 MIU/mL	SC	N/A			
	Solution 25 MIU multidose	10 MIU/mL	SC	N/A			
	Pen 3 MIU/dose multidose*	15 MIU/mL	SC	1.5, 3.0, 4.5, 6.0			
	Pen 5 MIU/dose multidose	25 MIU/mL	SC	7.5, 10.0			
4405	Pen 10 MIU/dose multidose	50 MIU/mL	SC	10.0, 15.0, 20.0			
1165	*Patients receiving 50% dose redu	ction only					
1166	**Patients receiving full dose only						
1167							
1168	NOTE: INTRON A Powder f	or Injection doe	es not contain a pre	servative. The vial			
1169	must be discarded after red	constitution and	d withdrawal of a si	ngle dose.			
1170	Dose Adjustment: NOTE: R	Regular laborator	v testing should be p	erformed to monitor			
1171	laboratory abnormalities for t	he nurnose of de	se modifications (se				
1170	Laboratory Tosts)						
1172	Laboratory resis).						
1173							
1174	 INTRON A should be wire 	thheld for sever	e adverse reactions,	including granulocyte			
1175	counts greater than 250/r	ກm³ but less tha	n 500/mm³ or SGPT/	SGOT greater than 5-			
1176	10x upper limit of norm	al, until adverse	e reactions abate.	INTRON A treatment			
1177	should be restarted at 50°	% of the previous	s dose.				
1178		, b					
1170		monontly diagon	tiousd for				
11/9	INTRON A should be per	manently discon					
1180	 I oxicity that does i 	not abate after w	ithnolding IN I RON F	A			
1181	 Severe adverse re 	actions which re	ecur in patients receit	ving reduced doses of			
1182	INTRON A						
1183	 Granulocyte count 	less than 250/m	m ³ or SGPT/SGOT c	of greater than 10x			
1184	upper limit of norm	al		0			
1185							
1186	Follicular Lymphoma (sool			Conoral)			
1100	Folicular Lympholia (see		ADMINISTRATION, V	Selleral)			
1187	a -						
1188	Dose: The recommended do	se of INTRON A	for the treatment of	follicular lymphoma is			
1189	5 million IU subcutaneously t	hree times per w	eek for up to 18 mor	oths in conjunction			
1190	with anthracycline-containing	chemotherapy i	egimen and following	g completion of the			
1191	chemotherapy regimen.		-				
1192							
1193		Dosage Forms for	This Indication				
	Dosage Form	Concentration	Route	Fixed Doses			
	Powder 10 MILL (single dose)	10 MILI/mI	SC	N/A			

Dosage Form	Concentration	Route	Fixed Doses
Powder 10 MIU (single dose)	10 MIU/mL	SC	N/A
Solution 18 MIU multidose	6 MIU/mL	SC	N/A
Solution 25 MIU multidose	10 MIU/mL	SC	N/A
Pen 5 MIU/dose multidose	25 MIU/mL	SC	2.5, 5.0
Pen 10 MIU/dose multidose	50 MIU/mL	SC	5.0

1194

1195

1196

NOTE: INTRON A Powder for Injection does not contain a preservative. The vial must be discarded after reconstitution and withdrawal of a single dose.

1197

1198 Dose Adjustment:

PAGE 34

- Doses of myelosuppressive drugs were reduced by 25% from a full-dose CHOP
 regimen, and cycle length increased by 33% (e.g., from 21 to 28 days) when alpha
 interferon was added to the regimen.
- Delay chemotherapy cycle if neutrophil count was less than 1500/mm³ or platelet count was less than 75,000/mm³.
- INTRON A should be permanently discontinued if SGOT exceeds greater than 5x
 the upper limit of normal or serum creatinine greater than 2.0 mg/dL (see
 WARNINGS).
- Administration of INTRON A therapy should be withheld for a neutrophil count less than 1000/mm³, or a platelet count less than 50,000/mm³.
- INTRON A dose should be reduced by 50% (2.5 MIU TIW) for a neutrophil count greater than 1000/mm³, but less than 1500/mm³. The INTRON A dose may be re-escalated to the starting dose (5 million IU TIW) after resolution of hematologic toxicity (ANC greater than 1500/mm³).

1215 Condylomata Acuminata (see DOSAGE AND ADMINISTRATION, General)

1217 Dose: The recommended dose is 1.0 million IU per lesion in a maximum of 5 lesions in
1218 a single course. The lesions should be injected three times weekly on alternate days for
1219 3 weeks. An additional course may be administered at 12 to 16 weeks.

1220 1221

1216

Dosage Forms for This Indication				
Dosage Form Concentration Route				
Powder 10 MIU (single dose)	10 MIU/mL	IL		
Solution 25 MIU multidose	10 MIU/mL	IL		

1222

1223 **NOTE: INTRON A Powder for Injection does not contain a preservative. The vial** 1224 **must be discarded after reconstitution and withdrawal of a single dose.**

1225

1227

1228

1226 **NOTE: Do not use the following formulations for this indication:**

- the 18 million or 50 million IU Powder for Injection
- the 18 million IU multidose INTRON A Solution for Injection
- the Multidose Pens
- 1229 1230 1231

Dose Adjustment: None

1232

1233 **Technique for Injection**: The injection should be administered intralesionally using a 1234 Tuberculin or similar syringe and a 25- to 30-gauge needle. The needle should be 1235 directed at the center of the base of the wart and at an angle almost parallel to the plane 1236 of the skin (approximately that in the commonly used PPD test). This will deliver the 1237 interferon to the dermal core of the lesion, infiltrating the lesion and causing a small 1238 wheal. Care should be taken not to go beneath the lesion too deeply; subcutaneous 1239 injection should be avoided, since this area is below the base of the lesion. Do not 1240 inject too superficially since this will result in possible leakage, infiltrating only the 1241 keratinized layer and not the dermal core.

1242

1243 AIDS-Related Kaposi's Sarcoma (see DOSAGE AND ADMINISTRATION, General)

PAGE 35

1244

1245 **Dose**: The recommended dose of INTRON A for Kaposi's Sarcoma is 30 million
 1246 IU/m²/dose administered subcutaneously or intramuscularly three times a week until
 1247 disease progression or maximal response has been achieved after 16 weeks of

1248 treatment. Dose reduction is frequently required (see **Dose Adjustment** below).

- 1249
- 1250

1251

1254

Dosa	age Forms	for This	Indication

Dosage Form	Concentration	Route
Powder 50 MIU	50 MIU/mL	IM, SC

1252 NOTE: INTRON A Solution for Injection either in vials or in Multidose Pens should 1253 NOT be used for AIDS-Related Kaposi's Sarcoma.

1255 NOTE: INTRON A Powder for Injection does not contain a preservative. The vial
 1256 must be discarded after reconstitution and withdrawal of a single dose.

1258 Dose Adjustment:

1259

1257

- INTRON A dose should be reduced by 50% or withheld for severe adverse reactions.
- INTRON A may be resumed at a reduced dose if severe adverse reactions abate with interruption of dosing.
- INTRON A should be permanently discontinued if severe adverse reactions persist or if they recur in patients receiving a reduced dose.

1267 Chronic Hepatitis C (see DOSAGE AND ADMINISTRATION, General)

1268

1269 **Dose**: The recommended dose of INTRON A for the treatment of chronic hepatitis C is 3 million IU three times a week (TIW) administered subcutaneously or intramuscularly. 1270 In patients tolerating therapy with normalization of ALT at 16 weeks of treatment, 1271 INTRON A therapy should be extended to 18 to 24 months (72 to 96 weeks) at 3 million 1272 1273 IU TIW to improve the sustained response rate (see CLINICAL PHARMACOLOGY, 1274 **Chronic Hepatitis C**). Patients who do not normalize their ALTs or have persistently high levels of HCV RNA after 16 weeks of therapy rarely achieve a sustained response 1275 1276 with extension of treatment. Consideration should be given to discontinuing these 1277 patients from therapy.

1278 When INTRON A is administered in combination with REBETOL[®], patients with 1279 impaired renal function and/or those over the age of 50 should be carefully monitored 1280 with respect to the development of anemia. See REBETOL prescribing information for 1281 dosing when used in combination with REBETOL for adults and pediatric patients.

1282 1283

1284

Dosage Form	Concentration	Route	Fixed Doses	
Solution 18 MIU multidose	6 MIU/mL	IM, SC	N/A	
Pen 3 MIU/dose multidose	15 MIU/mL	SC	1.5, 3.0	

Dosage Forms for This Indication

PAGE 36

1286

1287 **Dose Adjustment:** If severe adverse reactions develop during INTRON A treatment. the dose should be modified (50% reduction) or therapy should be temporarily 1288 discontinued until the adverse reactions abate. If intolerance persists after dose 1289 1290 adjustment, INTRON A therapy should be discontinued.

1291 1292

1293

Chronic Hepatitis B Adults (see DOSAGE AND ADMINISTRATION, General)

1294 **Dose**: The recommended dose of INTRON A for the treatment of chronic hepatitis B is 30 to 35 million IU per week, administered subcutaneously or intramuscularly, either as 1295 1296 5 million IU daily (QD) or as 10 million IU three times a week (TIW) for 16 weeks.

- 1297
- 1298

Dosage Form	Concentration	Route	Fixed Doses		
Powder 10 MIU (single dose)	10 MIU/mL	IM, SC	N/A		
Solution 25 MIU multidose	10 MIU/mL	IM, SC	N/A		
Pen 5 MIU/dose multidose	25 MIU/mL	SC	2.5, 5.0, 10.0		
Pen 10 MIU/dose multidose	50 MIU/mL	SC	5.0, 10.0		

Dosage Forms for This Indication

1299

1300 NOTE: INTRON A Powder for Injection does not contain a preservative. The vial must be discarded after reconstitution and withdrawal of a single dose. 1301

- 1302
- 1303
- 1304

Chronic Hepatitis B Pediatrics (see DOSAGE AND ADMINISTRATION, General)

1305 Dose: The recommended dose of INTRON A for the treatment of chronic hepatitis B is 3 million IU/m² three times a week (TIW) for the first week of therapy followed by dose 1306 escalation to 6 million IU/m² TIW (maximum of 10 million IU TIW) administered 1307 subcutaneously for a total duration of 16 to 24 weeks. 1308

- 1309
- 1310

Dosage Forms for This Indication				
Dosage Form	Concentration	Route	Fixed Doses	
Powder 10 MIU (single dose)	10 MIU/mL	SC	N/A	
Solution 25 MIU multidose	10 MIU/mL	SC	N/A	
Pen 3 MIU/dose multidose	15 MIU/mL	SC	1.5, 3.0, 4.5, 6.0	
Pen 5 MIU/dose multidose	25 MIU/mL	SC	2.5, 5.0, 7.5, 10.0	
Pen 10 MIU/dose multidose	50 MIU/mL	SC	5.0, 10.0, 15.0, 20.0	

1311

1312 NOTE: INTRON A Powder for Injection does not contain a preservative. The vial 1313 must be discarded after reconstitution and withdrawal of a single dose.

1314

1315 **Dose Adjustment:** If severe adverse reactions or laboratory abnormalities develop 1316 during INTRON A therapy, the dose should be modified (50% reduction) or discontinued if appropriate, until the adverse reactions abate. If intolerance persists after dose 1317 1318 adjustment, INTRON A therapy should be discontinued.

- 1319
- 1320 For patients with decreases in white blood cell, granulocyte or platelet counts, the 1321 following guidelines for dose modification should be followed:
- 1322

INTRON A Dose	White Blood Cell Count	Granulocyte Count	Platelet Count
Reduce 50%	<1.5 x 10 ⁹ /L	<0.75 x 10 ⁹ /L	<50 x 10 ⁹ /L
Permanently Discontinue	<1.0 x 10 ⁹ /L	<0.5 x 10 ⁹ /L	<25 x 10 ⁹ /L

- 1323
- 1324 1325

INTRON A therapy was resumed at up to 100% of the initial dose when white blood cell, granulocyte, and/or platelet counts returned to normal or baseline values.

1326 1327

PREPARATION AND ADMINISTRATION Reconstitution of INTRON[®] A Powder for 1328 1329 Injection The reconstituted solution is clear and colorless to light yellow. The INTRON 1330 A powder reconstituted with Sterile Water for Injection USP is a single-use vial and does not contain a preservative. DO NOT RE-ENTER VIAL AFTER WITHDRAWING THE 1331 1332 DISCARD UNUSED PORTION (see DOSAGE AND ADMINISTRATION). DOSE. 1333 Once the dose from the single-dose vial has been withdrawn, the sterility of any 1334 remaining product can no longer be guaranteed. Pooling of unused portions of some 1335 medications has been linked to bacterial contamination and morbidity.

1336

1337 • Intramuscular, Subcutaneous, or Intralesional Administration

Inject 1 mL Diluent (Sterile Water for Injection USP) for INTRON A into the INTRON A
Swirl gently to hasten complete dissolution of the powder. The appropriate
INTRON A dose should then be withdrawn and injected intramuscularly,
subcutaneously, or intralesionally (see **MEDICATION GUIDE** for detailed instructions).

Please refer to the **MEDICATION GUIDE** for detailed, step-by-step instructions on how to inject the INTRON A dose. After preparation and administration of the INTRON A injection, it is essential to follow the procedure for proper disposal of syringes and needles (see **MEDICATION GUIDE** for detailed instructions).

1346 Parenteral drug products should be inspected visually for particulate matter and 1347 discoloration prior to administration.

1348

1349 • Intravenous Infusion

The infusion solution should be prepared immediately prior to use. Based on the desired dose, the appropriate vial strength(s) of INTRON A should be reconstituted with the diluent provided. Inject 1 mL Diluent (Sterile Water for Injection USP) for INTRON A into the INTRON A vial. Swirl gently to hasten complete dissolution of the powder. The appropriate INTRON A dose should then be withdrawn and injected into a 100-mL bag of 0.9% Sodium Chloride Injection USP. The final concentration of INTRON A should not be less than 10 million IU/100 mL.

Please refer to the MEDICATION GUIDE for detailed, step-by-step instructions on
how to inject the INTRON A dose. After preparation and administration of INTRON A, it
is essential to follow the procedure for proper disposal of syringes and needles.

1360

1361 INTRON A Solution for Injection in Vials INTRON A Solution for Injection is supplied
 1362 in two multidose vials. The solutions for injection do not require reconstitution prior to
 1363 administration; the solution is clear and colorless.

PAGE 38

1365 The appropriate dose should be withdrawn from the vial and injected 1366 intramuscularly, subcutaneously, or intralesionally.

1368INTRON A Solution for Injection is not recommended for intravenous1369administration.

1370

1367

Solution for Injection in Multidose Pens The INTRON A Solution for Injection Multidose Pens are designed to deliver 3 to 12 doses, depending on the individual dose, using a simple dial mechanism, and are for subcutaneous injections only. Only the needles provided in the packaging should be used for the INTRON A Solution for Injection Multidose Pen. A new needle is to be used each time a dose is delivered using the pen. To avoid the possible transmission of disease, each INTRON A Solution for Injection Multidose Pen is for single patient use only.

1378

1382

1384

Please refer to the MEDICATION GUIDE for detailed, step-by-step instructions
on how to inject the INTRON A dose. After preparation and administration of INTRON
A, it is essential to follow the procedure for proper disposal of syringes and needles.

1383 HOW SUPPLIED

1385 **INTRON[®] A Powder for Injection**

1386 INTRON A Powder for Injection, 10 million IU per vial and Diluent for INTRON A 1387 (Sterile Water for Injection USP) 1 mL per vial; boxes containing 1 INTRON A vial and 1 1388 vial of INTRON A Diluent (NDC 0085-0571-02).

1389 INTRON A Powder for Injection, 18 million IU per vial and Diluent for INTRON A
1390 (Sterile Water for Injection USP) 1 mL per vial; boxes containing 1 vial of INTRON A
1391 and 1 vial of INTRON A Diluent (NDC 0085-1110-01).

1392 INTRON A Powder for Injection, 50 million IU per vial and Diluent for INTRON A
1393 (Sterile Water for Injection USP) 1 mL per vial; boxes containing 1 INTRON A vial and 1
1394 vial of INTRON A Diluent (NDC 0085-0539-01).

1395

1396INTRON A Solution for Injection in Multidose Pens

INTRON A Solution for Injection, 6 doses of 3 million IU (18 million IU)
Multidose Pen (22.5 million IU per 1.5 mL per pen); boxes containing 1 INTRON A
Multidose Pen, six disposable needles and alcohol swabs (NDC 0085-1242-01).

INTRON A Solution for Injection, 6 doses of 5 million IU (30 million IU) Multidose
Pen (37.5 million IU per 1.5 mL per pen); boxes containing 1 INTRON A Multidose Pen,
six disposable needles and alcohol swabs (NDC 0085-1235-01).

INTRON A Solution for Injection, 6 doses of 10 million IU (60 million IU)
Multidose Pen (75 million IU per 1.5 mL per pen); boxes containing 1 INTRON A
Multidose Pen, six disposable needles and alcohol swabs (NDC 0085-1254-01).

1406

1407 **INTRON A Solution for Injection in Vials**

INTRON A Solution for Injection, 18 million IU multidose vial (22.8 million IU per
3.8 mL per vial); boxes containing 1 vial of INTRON A Solution for Injection (NDC 00851168-01).

PAGE 39

1411 INTRON A Solution for Injection, 25 million IU multidose vial (32 million IU per 1412 3.2 mL per vial); boxes containing 1 vial of INTRON A Solution for Injection (NDC 0085-1413 1133-01). 1414 1415 Storage 1416 1417 **INTRON A Powder for Injection/Reconstitution** 1418 INTRON A Powder for Injection should be stored in the refrigerator at 2° to 8°C (36°-1419 46°F). After reconstitution, the solution should be used immediately, but may be 1420 stored up to 24 hours at 2° to 8°C (36°-46°F). Throw away any medicine left in the 1421 vial after you withdraw 1 dose. 1422 INTRON A Solution for Injection in Vials 1423 INTRON A Solution for Injection in vials should be stored in the refrigerator at 2° to 1424 8°C (36°-46°F). 1425 INTRON A Solution for Injection in Multidose Pens 1426 INTRON A Solution for Injection in Multidose Pens should be stored in the 1427 refrigerator at 2° to 8°C (36°-46°F). 1428 INTRON A Solution for Injection and INTRON A Solution for Injection in the • 1429 **Multidose Pens** 1430 INTRON A Solution for Injection and INTRON A Solution for Injection in the 1431 Multidose Pens should not be frozen and should be kept away from heat. Throw 1432 away any unused INTRON A Multidose Pen remaining after 4 weeks. Throw away 1433 any unused INTRON A Solution for Injection remaining in the vial after one month. 1434 1435 **References:** 1436 1. Smalley R, et al. N Engl J Med. 1992;327:1336-1341. 1437 2. Aviles A, et al. Leukemia and Lymphoma. 1996;20:495-499. 3. 1438 Unterhalt M, et al. Blood. 1996;88(10 Suppl 1):1744A. 1439 4. Schiller J, et al. J Biol Response Mod. 1989;8:252-261. 1440 5. Poynard T, et al. N Engl J Med. 1995;332(22)1457-1462. 1441 Lin R, et al. J Hepatol. 1995;23:487-496. 6. 1442 7. Perrillo R, et al. N Engl J Med. 1990;323:295-301. 1443 8. Perez V, et al. J Hepatol. 1990;11:S113-S117. 1444 9. Knodell R, et al. *Hepatology*. 1981;1:431-435. 1445 10. Perrillo R, et al. Ann Intern Med. 1991;115:113-115. 1446 11. Kauppila A, et al. Int J Cancer. 1982;29:291-294.

PAGE 40



Whitehouse Station, NJ 08889 USA.

1448 1449

1450

1451 Copyright[©] 1986, 2011 Schering Corporation, a subsidiary of **Merck & Co., Inc.** All

1452 rights reserved.

1453 U.S. Patent Nos. 5,935,566 and 6,610,830.

1454

1455 Rev. 11/2013

1456

MEDICATION GUIDE

INTRON[®] A (In-tron-aye)

(Interferon alfa-2b, recombinant)

Read this Medication Guide before you start taking INTRON A, and each time you get a refill. There may be new information. This information does not take the place of talking with your healthcare provider about your medical condition or your treatment.

If you are taking INTRON A with REBETOL, also read the Medication Guide for REBETOL[®] (ribavirin) Capsules and Oral Solution.

INTRON A alone is a treatment for certain types of cancers and hepatitis B virus. INTRON A by itself or with REBETOL is a treatment for some people infected with hepatitis C virus.

What is the most important information I should know about INTRON A?

INTRON A can cause serious side effects that:

- may cause death, or
- may worsen certain serious diseases that you may already have.

Tell your healthcare provider right away if you have any of the symptoms listed below while taking INTRON A. If symptoms get worse, or become severe and continue, your healthcare provider may tell you to stop taking INTRON A permanently. In many, but not all people, these symptoms go away after they stop taking INTRON A.

1. Heart problems. Some people who take INTRON A may develop heart problems, including:

- low blood pressure
- fast heart rate or abnormal heart beats
- trouble breathing or chest pain
- heart attacks or heart muscle problems (cardiomyopathy)
- 2. Stroke or symptoms of a stroke. Symptoms may include weakness, loss of coordination, and numbness. Stroke or symptoms of a stroke may happen in people who have some risk factors or no known risk factors for a stroke.
- **3. Mental health problems and suicide.** INTRON A may cause you to develop mood or behavior problems that may get worse during treatment with INTRON A or after your last dose, including:
 - irritability (getting upset easily)
 - depression (feeling low, feeling bad about yourself, or feeling hopeless)
 - aggressive behavior
 - thoughts of hurting yourself or others, or suicide
 - former drug addicts may fall back into drug addiction or overdose

If you have these symptoms, your healthcare provider should carefully monitor you during treatment with INTRON A and for 6 months after your last dose.

4. New or worsening autoimmune disease. Some people taking INTRON A develop autoimmune diseases (a condition where the body's immune cells attack other cells or organs in the body), including rheumatoid arthritis, systemic lupus erythematosus, sarcoidosis, and psoriasis. In some people who already have an autoimmune disease, the disease may get worse while on INTRON A.

5. Infections. Some people who take INTRON A may get an infection. Symptoms may include:

- fever
- chills
- bloody diarrhea
- burning or pain with urination
- urinating often
- coughing up mucus (phlegm) that is discolored (for example yellow or pink)

While taking INTRON A, you should see a healthcare provider regularly for check-ups and blood tests to make sure that your treatment is working and to check for side effects.

What is INTRON A?

INTRON A is a prescription medicine that is used:

- to treat adults with a blood cancer called hairy cell leukemia
- to treat certain adults with a type of skin cancer called malignant melanoma
- to treat adults with some types of Follicular Non-Hodgkin's Lymphoma along with certain chemotherapy medicines
- to treat certain adults with genital warts (condylomata acuminate), by injecting the medicine directly into the warts
- to treat certain adults with a type of cancer caused by AIDS, called AIDS-related Kaposi's Sarcoma
- alone to treat adults with chronic (lasting a long time) hepatitis C infection with stable liver problems
- with REBETOL to treat chronic (lasting a long time) hepatitis C infection in people 3 years and older with stable liver problems
- to treat chronic (lasting a long time) hepatitis B infection in people 1 year and older with stable liver problems

Who should not take INTRON A?

Do not take INTRON A if you:

- had a serious allergic reaction to another alpha interferon product or are allergic to any of the ingredients in INTRON A. See the end of this Medication Guide for a complete list of ingredients. Ask your healthcare provider if you are not sure.
- have certain types of hepatitis (autoimmune hepatitis)
- have certain other liver problems

Talk to your healthcare provider before taking INTRON A if you have any of these conditions.

What should I tell my healthcare provider before taking INTRON A?

Before you take INTRON A, tell your healthcare provider if you:

- See "What is the most important information I should know about INTRON A?"
- have or ever had any problems with your heart, including heart attack or have high blood pressure
- have or ever had bleeding problems or blood clots
- are being treated for a mental illness or had treatment in the past for any mental illness, including depression and suicidal behavior
- have any kind of autoimmune disease (where the body's immune system attacks the body's own cells), such as psoriasis, systemic lupus erythematosus, rheumatoid arthritis
- have or ever had low blood cell counts
- have ever been addicted to drugs or alcohol
- have liver problems (other than hepatitis B or C)
- have or had lung problems, such as chronic obstructive pulmonary disease (COPD)
- have diabetes
- have colitis (inflammation of your intestine)
- have a condition that suppresses your immune system, such as cancer
- have hepatitis B or C infection
- have HIV infection (the virus that causes AIDS)
- have kidney problems
- have high blood triglyceride levels (fat in your blood)
- have an organ transplant and are taking medicine that keeps your body from rejecting your transplant (suppresses your immune system)
- have any other medical conditions
- are pregnant or plan to become pregnant. It is not known if INTRON A will harm your unborn baby. You should use effective birth control during treatment with INTRON A. Talk to your healthcare provider about birth control choices for you during treatment with INTRON A. Tell your healthcare provider if you become pregnant during treatment with INTRON A.
- are breast-feeding or plan to breast-feed. It is not known if INTRON A passes into your breast milk. You and your healthcare provider should decide if you will use INTRON A or breast-feed. You should not do both.

Tell your healthcare provider about all the medicines you take, including prescription and non-prescription medicines, vitamins, and herbal supplements. INTRON A and certain other medicines may affect each other and cause side effects.

Especially tell your healthcare provider if you take:

- the anti-hepatitis B medicine telbivudine (Tyzeka)
- the anti-HIV medicine zidovudine (Retrovir)
- theophylline (Theo-24, Elixophyllin, Uniphyl, Theolair). Your healthcare provider may need to monitor the amount of theophylline in your body and make changes to your theophylline dose.

Know the medicines you take. Keep a list of them and show it to your healthcare provider and pharmacist when you get a new medicine.

How should I take INTRON A?

- INTRON A is given as an injection under the skin (subcutaneous) or into a muscle (intramuscular), into genital lesions, or as an injection into a vein (intravenous), depending on the condition that is being treated.
- Your healthcare provider will decide your dose of INTRON A and how often you will take it.
- If your healthcare provider decides that you can inject INTRON A for your condition, inject it exactly as prescribed, under your skin (subcutaneous injection) or into your muscle (intramuscular injection). Do not change your dose or how you inject INTRON A unless your healthcare provider tells you to.
- Do not take more than your prescribed dose.
- Your healthcare provider should show you how to prepare and measure your dose of INTRON A and how to inject yourself before you use INTRON A for the first time.
- You should not inject INTRON A until your healthcare provider has shown you how to use INTRON A the right way.
- INTRON A comes as:
 - a powder for injection in a vial that is used only 1 time (single-use vial). The powder must be mixed with water for injection (a diluent) before you inject it.
 - a solution for injection in a multi-dose vial
 - a solution for injection in a pen that is used more than 1 time (multidose pen)
- See the attached Instructions for Use for detailed instructions for preparing and injecting a dose of INTRON A.

- If you miss a dose of INTRON A, take the missed dose as soon as possible during the same day or the next day, then continue on your regular dosing schedule. If several days go by after you miss a dose, check with your healthcare provider to see what to do.
- Do not inject more than 1 dose or take more than your prescribed dose without talking to your healthcare provider.
- If you take too much INTRON A, call your healthcare provider right away. Your healthcare provider may examine you more closely, and do blood tests.
- Your healthcare provider should do regular blood tests before you start INTRON A, and during your treatment to see how well the treatment is working and to check for side effects.

What are the possible side effects of INTRON A?

INTRON A may cause serious side effects including:

- See "What is the most important information I should know about INTRON A?"
- **Blood problems.** INTRON A can affect your bone marrow and cause low white blood cell and platelet counts. In some people, these blood counts may fall to dangerously low levels. If your blood cell counts become very low, you can get infections or have bleeding problems.
- Serious eye problems. INTRON A may cause eye problems that may lead to vision loss or blindness. You should have an eye exam before you start taking INTRON A. If you have eye problems or have had them in the past, you may need eye exams while taking INTRON A. Tell your healthcare provider or eye doctor right away if you have any vision changes while taking INTRON A.
- **Thyroid problems.** Some people develop changes in the function of their thyroid. Symptoms of thyroid problems include:
 - o problems concentrating
 - o feeling cold or hot all the time
 - o changes in your weight
 - o skin changes
- **Blood sugar problems**. Some people may develop high blood sugar or diabetes. If you have high blood sugar or diabetes before starting INTRON A, talk to your healthcare provider before you take INTRON A. If you develop high blood sugar or diabetes while taking INTRON A, your healthcare provider may tell you to stop INTRON A and prescribe a different medicine for you. Symptoms of high blood sugar or diabetes may include:
 - increased thirst
 - o tiredness
 - o urinating more often than normal
 - o increased appetite
 - o weight loss
 - o your breath smells like fruit

Lung problems including:

- trouble breathing
- o pneumonia
- o inflammation of lung tissue
- new or worse high blood pressure of the lungs (pulmonary hypertension). This can be severe and may lead to death.

You may need to have a chest X-ray or other tests if you develop fever, cough, shortness of breath, or other symptoms of a lung problem during treatment with INTRON A.

- Severe liver problems, or worsening of liver problems including liver failure and • death. Symptoms may include:
 - o nausea
 - o loss of appetite
 - o tiredness
 - o diarrhea
 - yellowing of your skin or the white part of your eyes 0
 - bleeding more easily than normal 0
 - o swelling of your stomach area (abdomen)
 - o confusion
 - o sleepiness
 - o you cannot be awakened (coma)

Serious allergic reactions and skin reactions. Symptoms may include: .

o itching

0

o swelling of your face, eyes,

- o chest pain
- o feeling faint

lips, tongue, or throat

o skin rash, hives, sores in your

• trouble breathing anxiousness

- mouth, or your skin blisters and peels
- Swelling of your pancreas (pancreatitis) and intestines (colitis). Symptoms may • include:
 - severe stomach area (abdomen) pain 0
 - severe back pain
 - o nausea
 - o vomiting
 - fever 0
- New or worsening autoimmune disease. Some patients taking INTRON A develop • autoimmune diseases (a condition where the body's immune cells attack other cells or organs in the body), including rheumatoid arthritis, systemic lupus erythematosus, sarcoidosis, and psoriasis. In some patients who already have an autoimmune disease, the disease may worsen while on INTRON A.
- Nerve problems. People who take INTRON A or other alpha interferon products with • telbivudine (Tyzeka) can develop nerve problems such as continuing numbness, tingling, or burning sensation in the arms or legs (peripheral neuropathy). Call your healthcare provider if you have any of these symptoms.

- **Growth problems in children.** Weight loss and slowed growth are common in children during combination treatment with INTRON A and REBETOL. Most children will go through a growth spurt and gain weight after treatment stops. Some children may not reach the height that they were expected to have before treatment. Talk to your healthcare provider if you are concerned about your child's growth during treatment with INTRON A and REBETOL.
- Dental and gum problems.

Tell your healthcare provider right away if you have any of the symptoms listed above.

The most common side effects of INTRON A include:

- **Flu-like symptoms**. Symptoms may include: headache, muscle aches, tiredness, and fever. Some of these symptoms may be decreased by injecting your INTRON A dose in the evening. Talk to your healthcare provider about which over-the-counter medicines you can take to help prevent or decrease some of the symptoms.
- Tiredness. Many people become very tired during treatment with INTRON A.
- Appetite problems. Nausea, loss of appetite, and weight loss can happen with INTRON A.
- Skin reactions. Redness, swelling, and itching are common at the injection site.
- Hair thinning.

Tell your healthcare provider if you have any side effect that bothers you or that does not go away.

These are not all the side effects of INTRON A. For more information, ask your healthcare provider or pharmacist.

Call your doctor for medical advice about side effects. You may report side effects to the FDA at 1–800–FDA–1088.

How should I store INTRON A?

INTRON A Solution for Injection and INTRON A Solution for Injection in the Multidose Pens:

- Store in the refrigerator between $36^{\circ}F$ to $46^{\circ}F$ ($2^{\circ}C$ to $8^{\circ}C$).
- INTRON A Solution for Injection in Multidose vials for injection and INTRON A Solution for Injection in the Multidose Pens may be used to give more than 1 injection of medicine.
- Do not freeze.
- o Throw away any unused INTRON A Multidose Pen remaining after 4 weeks.
- Throw away any unused INTRON A Solution for Injection remaining in the vial after one month.

INTRON A Powder for Injection:

Before mixing, store in the refrigerator between 36°F to 46°F (2°C to 8°C).

- After mixing the INTRON A Powder for Injection, use the solution right away or store the solution in the refrigerator for up to 24 hours between $36^{\circ}F$ to $46^{\circ}F$ ($2^{\circ}C$ to $8^{\circ}C$).
- Throw away any medicine left in the vial after you withdraw 1 dose.
- o Do not freeze.

Keep INTRON A and all medicines out of the reach of children.

General Information about INTRON A

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. Do not use INTRON A for a condition for which it was not prescribed. Do not give INTRON A to other people, even if they have the same symptoms that you have. It may harm them.

This Medication Guide summarizes the most important information about INTRON A. If you would like more information, ask your healthcare provider. You can ask your healthcare provider or pharmacist for information about INTRON A that was written for health care professionals.

• For more information, go to www.IntronA.com or call 1-800-622-4477.

What are the ingredients in INTRON A?

Active ingredient: interferon alfa-2b

Inactive ingredients:

- **Powder for injection contains**: glycine, sodium phosphate dibasic, sodium phosphate monobasic, human albumin. Sterile water for injection is provided as a diluent.
- **Solution Multidose vials for injection contain**: sodium chloride, sodium phosphate dibasic, sodium phosphate monobasic, edetate disodium, polysorbate 80, and m-cresol as a preservative.
- Solution in Multidose Pens for injection contain: sodium chloride, sodium phosphate dibasic, sodium phosphate monobasic, edetate disodium, polysorbate 80, and m-cresol as a preservative.

The Medication Guide has been approved by the U.S. Food and Drug Administration.

Manufactured by: Schering Corporation, a subsidiary of **Merck & Co., Inc.,** Whitehouse Station, NJ 08889 USA

Copyright © 1996, 2011 Schering Corporation, a subsidiary of Merck & Co., Inc. All rights reserved.

Revised: 11/2013

mg-mk2958-mtl-XXXXrXXX