LEUKINE[®] SARGRAMOSTIM

Rx only

BERLEX



DESCRIPTION

LEUKINE® (sargramostim) is a recombinant human granulocyte-macrophage colony stimulating factor (rhu GM-CSF) produced by recombinant DNA technology in a yeast (S. cerevisiae) expression system. GM-CSF is a hematopoietic growth factor which stimulates proliferation and differentiation of hematopoietic progenitor cells. LEUKINE is a glycoprotein of 127 amino acids characterized by 3 primary molecular species having molecular masses of 19,500, 16,800 and 15,500 daltons. The amino acid sequence of LEUKINE differs from the natural human GM-CSF by a substitution of leucine at position 23, and the carbohydrate moiety may be different from the native protein. Sargramostim has been selected as the proper name for yeast-derived rhu GM-CSF.

The LEUKINE Liquid presentation is formulated as a sterile, preserved (11% benzyl alcohol), injectable solution (500 mcg/mL) in a vial. Lyophilized LEUKINE is a sterile, white, preservative-free powder (250 mcg) that requires reconstitution with 1 mL Sterile Water for Injection, USP or 1 mL Bacteriostatic Water for Injection, USP

LEUKINE Liquid and reconstituted lyophilized LEUKINE are clear, colorless liquids suitable for subcutaneous injection or intravenous infusion. LEUKINE Liquid contains 500 mcg (2.8 x 10⁶ IU/mL) sargramostim and 11% benzyl alcohol in a 1 mL solution. The vial of lyophilized LEUKINE contains 250 mcg (14 x 10⁶ IU/vial) sargramostim. The LEUKINE Liquid vial and reconstituted lyophilized LEUKINE vial also contain 40 mg/mL mannitol, USP; 10 mg/mL sucrose, NF; and 1.2 mg/mL tromethamine, USP, as excipients. Biological potency is expressed in International Units (IU) as tested against the WHO First International Reference Standard. The specific activity of LEUKINE is approximately 5.6 x 106 IU/mg.

CLINICAL PHARMACOLOGY

General GM-CSF belongs to a group of growth factors termed colony stimulating factors which support survival, clonal expansion, and differentiation of hematopoietic progenitor cells. GM-CSF induces partially committed progenitor cells to divide and differentiate in the granulocyte-macrophage pathways.

GM-CSF is also capable of activating mature granulocytes and macrophages. GM-CSF is a multilineage factor and, in addition to dose-dependent effects on the myelomonocytic lineage, can promote the proliferation of megakaryocytic and erythroid progenitors.¹ However, other factors are required to induce complete maturation in these two lineages. The various cellular responses (i.e., division, maturation, activation) are induced through GM-CSF binding to specific receptors expressed on the cell surface of target cells.2

In vitro Studies of LEUKINE in Human Cells The biological activity of GM-CSF is species-specific. Consequently, in vitro studies have been performed on human cells to characterize the pharmacological activity of LEUKINE. In vitro exposure of human bone marrow cells to LEUKINE at concentrations ranging from 1-100 ng/mL results in the proliferation of hematopoietic progenitors and in the formation of pure granulocyte, pure macrophage and mixed granulocyte-macrophage colonies.³ Chemotactic, anti-fungal and anti-parasitic⁴ activities of granulocytes and monocytes are increased by exposure to LEUKINE in vitro. LEUKINE increases the cytotoxicity of monocytes toward certain neoplastic cell lines³ and activates polymorphonuclear neutrophils to inhibit the growth of tumor cells.

In vivo Primate Studies of LEUKINE Pharmacology/toxicology studies of LEUKINE were performed in cynomology monkeys. An acute toxicity study revealed an absence of treatment-related toxicity following a single IV bolus injection at a dose of 300 mcg/kg. Two subacute studies were performed using IV injection (maximum dose 200 mco/kg/day x 14 days) and subcutaneous injection (maximum dose 200 mcg/kg/day x 28 days). No major visceral organ toxicity was documented. Notable histopathology findings included increased cellularity in hematologic organs and heart and lung tissues. A dose-dependent increase in leukocyte count, which consisted primarily of segmented neutrophils, occurred during the dosing period; increases in monocytes, basophils, eosinophils and lymphocytes were also noted. Leukocyte counts decreased to pretreatment values over a 1-2 week recovery period.

Pharmacokinetics Pharmacokinetic profiles have been analyzed in controlled studies of 24 normal male volunteers. Liquid and lyophilized LEUKINE, at the recommended dose of 250 mcg/m², have been determined to be bioequivalent based on the statistical evaluation of AUC.⁵

When LEUKINE (either liquid or lyophilized) was administered IV over 2 hours to normal volunteers, the mean beta half-life was approximately 60 minutes. Peak concentrations of GM-CSF were observed in blood samples obtained during or immediately after completion of LEUKINE infusion. For LEUKINE Liquid, the mean maximum concentration (Cmax) was 5.0 ng/mL, the mean clearance rate was approximately 420 mL/min/m² and the mean AUC (0-inf) was 640 ng/mL•min. Corresponding results for lyophilized LEUKINE in the same subjects were mean Cmax of 5.4 ng/mL, mean clearance rate of 431 mL/min/m², and mean AUC (0-inf) of 677 ng/mL+min, GM-CSF was last detected in blood samples obtained at 3 or 6 hours.

When LEUKINE (either liquid or lyophilized) was administered SC to normal volunteers, GM-CSF was detected in the serum at 15 minutes, the first sample point. The mean beta half-life was approximately 162 minutes. Peak levels occurred at 1 to 3 hours post injection, and LEUKINE remained detectable for up to 6 hours after injection. The mean Cmax was 1.5 ng/mL. For LEUKINE Liquid, the mean clearance was 549 mL/min/m² and the mean AUC (0-inf) was 549 ng/mL-min. For lyophilized LEUKINE, the mean clearance was 529 mL/min/m² and the mean AUC (0-inf) was 501 ng/mL•min.

Antibody Formation Serum samples collected before and after LEUKINE treatment from 214 patients with a variety of underlying diseases have been examined for the presence of antibodies. Neutralizing antibodies were detected in 5 of 214 patients (2.3%) after receiving LEUKINE by continuous IV infusion (3 patients) or subcutaneous injection (2 patients) for 28 to 84 days in multiple courses. All 5 patients had impaired hematopoiesis before the administration of LEUKINE and consequently the effect of the development of anti-GM-CSF antibodies on normal hematopoiesis could not be assessed. Drug-induced neutropenia, neutralization of endogenous GM-CSF activity and diminution of the therapeutic effect of LEUKINE secondary to formation of neutralizing antibody remain a theoretical possibility.

INDICATIONS AND USAGE

Use Following Induction Chemotherapy in Acute Myelogenous Leukemia LEUKINE is indicated for use following induction chemotherapy in older adult patients with acute myelogenous leukemia (AML) to shorten time to neutrophil recovery and to reduce the incidence of severe and life-threatening infections and infections resulting in death. The safety and efficacy of LEUKINE have not been assessed in patients with AML under 55 years of age.

The term acute myelogenous leukemia, also referred to as acute non-lymphocytic leukemia (ANLL), encompasses a beterogeneous group of leukemias arising from various non-lymphoid cell lines which have been defined morphologically by the French-American-British (FAB) system of classification.

Use in Mobilization and Following Transplantation of Autologous Peripheral Blood Progenitor Cells LEUKINE is indicated for the mobilization of hematopoietic progenitor cells into peripheral blood for collection by leukapheresis. Mobilization allows for the collection of increased numbers of progenitor cells capable of engraftment as compared with collection without mobilization. After myeloablative chemotherapy, the transplantation of an increased number of progenitor cells can lead to more rapid engraftment, which may result in a decreased need for supportive care. Myeloid reconstitution is further accelerated by administration of LEUKINE following peripheral blood progenitor cell transplantation

Use in Myeloid Reconstitution After Autologous Bone Marrow Transplantation LEUKINE is indicated for acceleration of myeloid recovery in patients with non-Hodgkin's lymphoma (NHL), acute lymphoblastic leukemia (ALL) and Hodgkin's disease undergoing autologous bone marrow transplantation (BMT). After autologous BMT in patients with NHL, ALL, or Hodgkin's disease, LEUKINE has been found to be safe and effective in accelerating myeloid engraftment, decreasing median duration of antibiotic administration, reducing the median duration of infectious episodes and shortening the median duration of hospitalization. Hematologic response to LEUKINE can be detected by complete blood count (CBC) with differential performed twice per week.

Use in Myeloid Reconstitution After Allogeneic Bone Marrow Transplantation LEUKINE is indicated for acceleration of myeloid recovery in patients undergoing allogeneic BMT from HLA-matched related donors. LEUKINE has been found to be safe and effective in accelerating myeloid engraftment, reducing the incidence of bacteremia and other culture positive infections, and shortening the median duration of hospitalization.

LEUKINE (sargramostim)

Use in Bone Marrow Transplantation Failure or Engraftment Delay LEUKINE is indicated in patients who have undergone allogeneic or autologous bone marrow transplantation (BMT) in whom engraftment is delayed or has failed. LEUKINE has been found to be safe and effective in prolonging survival of patients who are experiencing graft failure or engraftment delay, in the presence or absence of infection, following autologous or allogeneic BMT. Survival benefit may be relatively greater in those patients who demonstrate one or more of the following characteristics: autologous BMT failure or engraftment delay, no previous total body irradiation, malignancy other than leukemia or a multiple organ failure (MOF) score s 2 (See CLINICAL EXPERIENCE). Hematologic response to LEUKINE can be detected by complete blood count (CBC) with differential performed twice per week

CLINICAL EXPERIENCE

Acute Myelogenous Leukemia The safety and efficacy of sargramostim in patients with AML who are younger than 55 years of age have not been determined. Based on Phase II data suggesting the best therapeutic effects could be achieved in patients at highest risk for severe infections and mortality while neutropenic, the Phase III clinical trial was conducted in older patients. The safety and efficacy of LEUKINE in the treatment of AML were evaluated in a multi-center, randomized, doubleblind placebo-controlled trial of 99 newly diagnosed adult patients, 55-70 years of age, receiving induction with or without consolidation.⁶ A combination of standard doses of daunorubicin (days 1-3) and ara-C (days 1-7) was administered during induction and high dose ara-C was administered days 1-6 as a single course of consolidation, if given. Bone marrow evaluation was performed on day 10 following induction chemotherapy. If hypoplasia with <5% blasts was not achieved, patients immediately received a second cycle of induction chemotherapy. If the bone marrow was hypoplastic with <5% blasts on day 10 or 4 days following the second cycle of induction chemotherapy. LEUKINE (250 mcg/m²/day) or placebo was given IV over 4 hours each day starting 4 days after the completion of chemotherapy. Study drug was continued until an ANC >1500/mm³ for three consecutive days was attained or a maximum of 42 days. I FUKINE or placebo was also administered after the single course of consolidation chemotherapy if delivered (ara-C 3-6 weeks after induction following neutrophil recovery). Study drug was discontinued immediately if leukemic rearowth occurred. Hematological Recovery (in Days): Induction

Dataset

RRC

ANC>500/mm3

ANC>1000/mm3 b

PLT>20,000/mm^{3 c}

Patients with missing data censored

Generalized Wilcoxon

4 patients on placebo had missing values

LEUKINE (sargramostim) significantly shortened the median duration of ANC <500/mm³ by 4 days and <1000/mm³ by 7 days following induction (see table at right). 75% of patients receiving LEUKINE achieved ANC >500/mm³ by day 16, compared to day 25 for patients receiving placebo. The proportion of patients receiving 1 cycle (70%) or 2 cycles (30%) of induction was similar in both treatment groups; LEUKINE significantly shortened the median times to neutrophil recovery whether one cycle (12 versus 15 days) or two cycles (14 versus 23 days) of induction chemotherapy was administered. Median times to platelet (>20.000/mm³) and RBC transfusion independence were not significantly different between treatment groups.

During the consolidation phase of treatment, LEUKINE did not shorten the median time to recovery of ANC to 500/mm3 (13 days) or 1000/mm3 (14.5 days) compared to placebo. There were no significant differences in time to pl

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atelet and BBC transfusion independence	° 3 L
	** n_
The incidence of severe infections and deaths associated with infections	P-1

was significantly reduced in patients who received LEUKINE. During induction or consolidation, 27 of 52 patients receiving LEUKINE and 35 of 47 patients receiving placebo had at least one grade 3, 4 or 5 infection (p=0.02). Twenty-five patients receiving LEUKINE and 30 patients receiving placebo experienced severe and fatal infections during induction only. There were significantly fewer deaths from infectious causes in the sargramostim arm (3 versus 11, p=0.02). The majority of deaths in the placebo group were associated with fungal infections with pneumonia as the primary infection

Disease outcomes were not adversely affected by the use of LEUKINE. The proportion of patients achieving complete remission (CR) was higher in the LEUKINE group (69% as compared to 55% for the placebo group), but the difference was not significant (p=0.21). There was no significant difference in relapse rates; 12 of 36 patients who received LEUKINE and 5 of 26 patients who received placebo relapsed within 180 days of documented CR (p=0.26). The overall median survival was 378 days for patients receiving LEUKINE and 268 days for those on placebo (p=0.17). The study was not sized to assess the impact of LEUKINE treatment on response or survival.

Mobilization and Engraftment of PBPC A retrospective review was conducted of data from patients with cancer undergoing collection of peripheral blood progenitor cells (PBPC) at a single transplant center. Mobilization of PBPC and myeloid reconstitution post-transplant were compared between four groups of patients (n=196) receiving LEUKINE for mobilization and a historical control group who did not receive any mobilization treatment [progenitor cells collected by leukapheresis without mobilization (n=100)]. Sequential cohorts received LEUKINE. The cohorts differed by dose (125 or 250 mcg/m²/day), route (IV over 24 hours or SC) and use of LEUKINE post-transplant. Leukaphereses were initiated for all mobilization groups after the WBC reached 10,000/mm3. Leukaphereses continued until both a minimum number of mononucleated cells (MNC) were collected (6.5 or 8.0 x 10⁹/kg body weight) and a minimum number of phereses (5-8) were performed. Both minimum requirements varied by treatment cohort and planned conditioning regimen. If subjects failed to reach a WBC of 10,000 cells/mm³ by day 5, another cytokine was substituted for LEUKINE; these subjects were all successfully leukapheresed and transplanted. The most marked mobilization and post-transplant effects were seen in patients administered the higher dose of LEUKINE (250 mcg/m2) either IV (n=63) or SC (n=41).

PBPCs from patients treated at the 250 mcg/m²/day dose had significantly higher number of granulocyte-macrophage colony-forming units (CFU-GM) than those collected without mobilization. The mean value after thawing was 11.41 x 10⁴ CFU-GM/kg for all LEUKINE-mobilized patients, compared to 0.96 x 10⁴/kg for the nonmobilized group. A similar difference was observed in the mean number of erythrocyte burst-forming units (BFU-E) collected (23.96 x 104/kg for patients mobilized with 250 mc d patients).

J	mcg/m-	doses of	LEUKINE	administered	1.20	VS.	1.03 X	IU⁼/Kg	101 11011-1	nopilized
	After tran	nsplantatio	on, mobili	zed subjects	had	shor	ter tim	nes to	_	

myeloid engraftment and fewer days between transplantation and the	ANC and Platelet Recovery after PBPC Transplant						
last platelet transfusion compared to non-mobilized subjects. Neutrophil recovery (ANC >500/mm ³) was more rapid in patients		Route for	Post-transplant	ENGRA (median va	FTMENT Ilue in days)		
administered LEUKINE following PBPC transplantation with LEUKINE- mobilized cells (see table at right). Mobilized patients also had fewer		Mobilization	LEUKINE	ANC>500/mm3	Last platelet transfusion		
days to the last platelet transfusion and last RBC transfusion, and a shorter duration of boshitalization than did non-mobilized subjects	No Mobilization	_	по	29	28		
A second retrospective review of data from patients undergoing	LEUKINE	IV	no	21	24		
PBPC at another single transplant center was also conducted. LEUKINE	250 mcg/m ²	IV	yes	12	19		
was given SC at 250 mcg/m ² /day once a day (n=10) or twice a day		SC	yes	12	17		

sargramostim

n=52*

Median (25%, 75%)

13 (11, 16)

14 (12, 18)

11 (7, 14)

12 (9, 24)

2 patients on sargramostim and 4 patients on placebo had missing values.

patients on sargramostim and 3 patients on placebo had missing values.

atients on sargramostim and 4 patients on placebo had missing values.

Placebo

n=47

Median (25%, 75%)

17 (13, 25)

21 (13, 34)

12 (9, >42)

14 (9, 42)

p-value**

0.009

0.003

0.10

0.53

(n=21) until completion of the phereses. Phereses were begun on day 5 of LEUKINE administration and continued until the targeted MNC count of 9 x 10⁸/kg or CD34+ cell count of 1 x 10⁶/kg was reached. There was no difference in CD34+ cell count in patients receiving LEUKINE once or twice a day. The median time to ANC>500/mm³ was 12 days and to platelet recovery (>25,000/mm³ was 23 days

Survival studies comparing mobilized study patients to the non-mobilized patients and to an autologous historical bone marrow transplant group showed no differences in median survival time

Autologous Bone Marrow Transplantation⁷ Following a dose-ranging Phase I/II trial in patients undergoing autologous BMT for lymphoid malignancies,^{8,9} three single center, randomized, placebo-controlled and double-blinded studies were conducted to evaluate the safety and efficacy of LEUKINE for promoting hematopoietic reconstitution following autologous BMT. A total of 128 patients (65 LEUKINE, 63 placebo) were enrolled in these 3 studies. The majority of the patients had lymphoid malignancy (87 NHL, 17 ALL), 23 patients had Hodgkin's disease, and 1 patient had acute myeloblastic leukemia (AML). In 72 patients with NHL or ALL, the bone marrow harvest was purged prior to storage with one of several monoclonal antibodies. No chemical agent was used for in vitro treatment of the bone marrow. Preparative regimens in the 3 studies included cyclophosphamide (total dose 120-150 mg/kg) and total body irradiation (total dose 1,200-1,575 rads). Other regimens used in patients with Hodgkin's disease and NHL without radiotherapy consisted of 3 or more of the following in combination (expressed as total dose): cytosine arabinoside (400 mg/m²) and carmustine (300 mg/m²), cyclophosphamide (140-150 mg/kg), hydroxyurea (4.5 grams/m²) and etoposide (375-450 mg/m²).

Compared to placebo, administration of LEUKINE in 2 studies (n=44 and 47) significantly improved the following hematologic and clinical endpoints: time to neutrophil engraftment, duration of hospitalization and infection experience or antibacterial usage. In the third study (n=37) there was a positive trend toward earlier myeloid engraftment in favor of LEUKINE. This latter study differed from the other 2 in having enrolled a large number of patients with Hodgkin's disease who had also received extensive radiation and chemotherapy prior to harvest of autologous bone marrow. A subgroup analysis of the data from all 3 studies revealed that the median time to engraftment for patients with Hodgkin's disease, regardless of treatment, was 6 days longer when compared to patients with NHL and ALL, but that the overall beneficial LEUKINE treatment effect was the same. In the following combined analysis of the 3 studies, these 2 subgroups (NHL and ALL vs. Hodgkin's disease) are presented separately

Patients with Lymphoid Malignanc Lymphoma and Acute Lymphoblastic Le Mveloid engraftment (absolute neutrop) > 500 cells/mm³) in 54 natients receiv observed 6 days earlier than in 50 nati placebo (see table at right). Accelerate engraftment was associated with signif benefits. The median duration of hospi 6 days shorter for the LEUKINE group f placebo group. Median duration of infe (defined as fever and neutropenia: or 2 of the same organism: or fever >38°C a blood culture: or clinical evidence of inf receiving LEUKINE engrafted earlier than controls.

disease (GVHD) prophylaxis was cyclosporine A and a corticosteroid.

Accelerated myeloid engraftment was as improved the following: time to neutrophil duration of hospitalization, number of patier bacteremia and overall incidence of infection at right)

Median time to myeloid engraftment (A ≥ 500 cells/mm³) in 53 patients receiving (sargramostim) was 4 days less than in 56 treated with placebo (see table at right). Th patients with bacteremia and infection was lower in the LEUKINE group compared to the group (9/53 versus 19/56 and 30/53 versus statistical significance.

Three categories of patients were eligible for this study:

engraftment with ANC < 500 cells/mm³ for at least one week beyond day 21 post-transplantation). A total of 140 eligible patients from 35 institutions were treated with LEUKINE and evaluated in comparison to 103 historical control patients from a single institution. One hundred sixty-three patients had lymphoid or myeloid leukemia, 24 patients had non-Hodgkin's lymphoma, 19 patients had Hodgkin's disease and 37 patients had other diseases, such as aplastic anemia, myelodysplasia or non-hematologic malignancy. The majority of patients (223 out of 243) had received prior chemotherapy with or without radiotherapy and/or immunotherapy prior to preparation for transplantation

One hundred day survival was improved in favor of the patients treated with LEUKINE after graft failure following either autologous or allogeneic BMT. In addition, the median survival was improved by greater than 2-fold. The median survival of patients treated with LEUKINE after autologous failure was 474 days versus 161 days for the historical patients. Similarly, after allogeneic failure, the median survival was 97 days survival was better in patients with fewer impaired organs.

organ systems: cardiovascular, respiratory, gastrointestinal, hematologic, renal, hepatic and neurologic.¹⁰ Assessment of the MOF score is recommended as an additional method of determining the need to initiate treatment with LEUKINE in patients with graft failure or delay in engraftment following autologous or allogeneic BMT.

Factors that Contribute to Survival: The probability of survival was relatively greater for patients with any one of the following characteristics: autologous BMT failure or delay in engraftment, exclusion of total body irradiation from the preparative regimen, a non-leukemic malignancy or MOF score ≤ 2 (0, 1 or 2 dysfunctional organ systems). Leukemic subjects derived less benefit than other subjects.

CONTRAINDICATIONS

I FUKINE is contraindicated:

- 3) for concomitant use with chemotherapy and radiotherapy.

WARNINGS

Pediatric Use Benzyl alcohol is a constituent of LEUKINE Liquid and Bacteriostatic Water for Injection diluent. Benzyl alcohol has been reported to be associated with a fatal "Gasping Syndrome" in premature infants. Liquid solutions containing benzyl alcohol (including LEUKINE Liquid) or lyophilized LEUKINE reconstituted with Bacteriostatic Water for Injection, USP (0.9% benzyl alcohol) should not be administered to neonates (see PRECAUTIONS and DOSAGE AND ADMINISTRATION)

Fluid Retention Edema, capillary leak syndrome, pleural and/or pericardial effusion have been reported in patients after LEUKINE administration. In 156 patients enrolled in placebo-controlled studies using LEUKINE at a dose of 250 mcg/m²/day by 2-hour IV infusion, the reported incidences of fluid retention (LEUKINE vs. placebo) were as follows: peripheral edema, 11% vs. 7%; pleural effusion, 1% vs. 0%; and pericardial effusion, 4% vs. 1%. Capillary leak syndrome was not observed in this limited number of studies; based on other uncontrolled studies and reports from users of marketed LEUKINE, the incidence is estimated to be less than 1%. In patients with preexisting pleural and pericardial effusions, administration of LEUKINE may aggravate fluid retention; however, fluid retention associated with or worsened by LEUKINE has been reversible after interruption or dose reduction of LEUKINE with or without diuretic therapy. LEUKINE should be used with caution in patients with preexisting fluid retention, pulmonary infiltrates or congestive heart failure.

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y (Non-Hodgkin's eukemia): hil count [ANC] ving LELIKINE was	Autol	ogous BMT: Com of Res	bined Analysis sponses in Pat Median '	s from Placebo-C ients with NHL a Values (days)	Controlled Clin and ALL	nical Trials
ents treated with d mveloid		ANC ≥500/mm ³	ANC ≥1000/mm ³	Duration of Hospitalization	Duration of Infection	Duration of Antibacterial Therapy
icant clinical talization was	LEUKINE (n=54)	18*#	24*#	25*	1*	21*
han for the ctious episodes	Placebo (n=50)	24	32	31	4	25
positive cultures and 1 positive	* p <0.05 Wilcoxo Note: The single A	on or CMH ridit chi-si AML patient was not i	quared # p <0 ncluded.	0.05 Log rank		

3 days less in the group treated with LEUKINE. The median duration of antibacterial administration in the post-transplantation period was 4 days shorter for the patients treated with LEUKINE than for placebo-treated patients. The study was unable to detect a significant difference between the treatment groups in rate of disease relapse 24 months post-transplantation. As a group, leukemic subjects receiving LEUKINE derived less benefit than NHL subjects. However, both the leukemic and NHL groups

Patients with Hodgkin's Disease: If patients with Hodgkin's disease are analyzed separately, a trend toward earlier myeloid engraftment is noted. LEUKINE-treated patients engrafted earlier (by 5 days) than the placebo-treated patients (p=0.189, Wilcoxon) but the number of patients was small (n=22). Studies are in progress to confirm statistically the trend toward earlier engraftment with LEUKINE in patients with Hodgkin's disease.

Allogeneic Bone Marrow Transplantation A multi-center, randomized, placebo-controlled, and double-blinded study was conducted to evaluate the safety and efficacy of LEUKINE for promoting hematopoietic reconstitution following allogeneic BMT. A total of 109 patients (53 LEUKINE, 56 placebo) were enrolled in the study. Twenty-three patients (11 LEUKINE, 12 placebo) were 18 years old or younger. Sixty-seven patients had myeloid malignancies (33 AML, 34 CML), 17 had lymphoid malionancies (12 ALL, 5 NHL), 3 patients had Hodokin's disease, 6 had multiple myeloma, 9 had myelodysplastic disease, and 7 patients had aplastic anemia. In 22 patients at one of the seven study sites, bone marrow harvests were depleted of T cells. Preparative regimens included cvclophosphamide. husulfan. cvtosine arabinoside, etoposide, methotrexate, corticosteroids, and asparaginase. Some patients also received total body, splenic, or testicular irradiation. Primary graft-versus-host

ts with (see table		Allogeneic BN	IT: Analysis Median Val	of Data from Placebo ues (days or number of	-Controlled Clinical patients)	Trial
IC		ANC \geq 500/mm ³	ANC ≥ 1000/mm ³	Number of Patients with Infections	Number of Patients with Bacteremia	Days of Hospitalization
EUKINE patients	LEUKINE (n=53)	13*	14*	30*	9**	25*
e number of ignificantly	Placebo (n=56)	17	19	42	19	26

respectively). There were a number of secondary laboratory and clinical endpoints. Of these, only the incidence of severe (grade 3/4) mucositis was significantly improved in the LEUKINE group (4/53) compared to the placebo group (16/56) at p<0.05. LEUKINE-treated patients also had a shorter median duration of post-transplant IV antibiotic infusions, and shorter median number of days to last platelet and RBC transfusions compared to placebo patients, but none of these differences reached

Bone Marrow Transplantation Failure or Engraftment Delay A historically controlled study was conducted in patients experiencing graft failure following allogeneic or autologous BMT to determine whether LEUKINE improved survival after BMT failure.

1) patients displaying a delay in engraftment (ANC ≤ 100 cells/mm³ by day 28 post-transplantation);

2) patients displaying a delay in engraftment (ANC ≤ 100 cells/mm³ by day 21 post-transplantation) and who had evidence of an active infection; and

patients who lost their marrow graft after a transient engraftment (manifested by an average of ANC ≥ 500 cells/mm³ for at least one week followed by loss of

The MOF score is a simple clinical and laboratory assessment of 7 major

with LEUKINE treatment and 35 days for the historical controls. Improvement in								
Median Survival by Multiple Organ Failure (MOF) Category Median Survival (days)								
MOF (Composite MOF ≤ 2 Organs MOF > 2 Organs of Both Groups)								
utologous BMT								
LEUKINE	474 (n=58)	78.5 (n=10)	474 (n=68)					
Historical	165 (n=14)	39 (n=3)	161 (n=17)					
Allogeneic BMT	llogeneic BMT							
LEUKINE	174 (n=50)	27 (n=22)	97 (n=72)					
Historical	52.5 (n=60)	15.5 (n=26)	35 (n=86)					

1) in patients with excessive leukemic myeloid blasts in the bone marrow or peripheral blood (≥ 10%);

2) in patients with known hypersensitivity to GM-CSF, yeast-derived products or any component of the product;

Due to the potential sensitivity of rapidly dividing hematopoietic progenitor cells, LEUKINE should not be administered simultaneously with cytotoxic chemotherapy or radiotherapy or within 24 hours preceding or following chemotherapy or radiotherapy. In one controlled study, patients with small cell lung cancer received LEUKINE and concurrent thoracic radiotherapy and chemotherapy or the identical radiotherapy and chemotherapy without LEUKINE. The patients randomized to LEUKINE had significantly higher incidence of adverse events, including higher mortality and a higher incidence of grade 3 and 4 infections and grade 3 and 4 thrombocytopenia.1

LEUKINE (sargramostim)

WARNINGS (continued)

Respiratory Symptoms Sequestration of granulocytes in the pulmonary circulation has been documented following LEUKINE infusion,¹² and dyspnea has been reported occasionally in patients treated with LEUKINE. Special attention should be given to respiratory symptoms during or immediately following LEUKINE infusion, especially in patients with preexisting lung disease. In patients displaying dyspnea during LEUKINE administration, the rate of infusion should be reduced by half. If respiratory symptoms worsen despite infusion rate reduction, the infusion should be discontinued. Subsequent IV infusions may be administered following the standard dose schedule with careful monitoring. LEUKINE should be administered with caution in patients with hypoxia.

Cardiovascular Symptoms Occasional transient supraventricular arrhythmia has been reported in uncontrolled studies during LEUKINE administration, particularly in patients with a previous history of cardiac arrhythmia. However, these arrhythmias have been reversible after discontinuation of LEUKINE. LEUKINE should be used with caution in patients with preexisting cardiac disease.

Renal and Hepatic Dysfunction In some patients with preexisting renal or hepatic dysfunction enrolled in uncontrolled clinical trials, administration of LEUKINE has induced elevation of serum creatinine or bilirubin and hepatic enzymes. Dose reduction or interruption of LEUKINE administration has resulted in a decrease to pretreatment values. However, in controlled clinical trials the incidences of renal and hepatic dysfunction were comparable between LEUKINE (250 mcg/m²/day by 2-hour IV infusion) and placebo-treated patients. Monitoring of renal and hepatic function in patients displaying renal or hepatic dysfunction prior to initiation of treatment is recommended at least every other week during LEUKINE administration.

PRECAUTIONS

General Parenteral administration of recombinant proteins should be attended by appropriate precautions in case an allergic or untoward reaction occurs. Serious allergic or anaphylactic reactions have been reported. If any serious allergic or anaphylactic reaction occurs, LEUKINE therapy should immediately be discontinued and appropriate therapy initiated.

A syndrome characterized by respiratory distress, hypoxia, flushing, hypotension, syncope, and/or tachycardia has been reported following the first administration of LEUKINE (sargramostim) in a particular cycle. These signs have resolved with symptomatic treatment and usually do not recur with subsequent doses in the same cycle of treatment

Stimulation of marrow precursors with LEUKINE may result in a rapid rise in white blood cell (WBC) count. If the ANC exceeds 20,000 cells/mm3 or if the platelet count exceeds 500,000/mm3, LEUKINE administration should be interrupted or the dose reduced by half. The decision to reduce the dose or interrupt treatment should be based on the clinical condition of the patient. Excessive blood counts have returned to normal or baseline levels within 3 to 7 days following cessation of LEUKINE therapy. Twice weekly monitoring of CBC with differential (including examination for the presence of blast cells) should be performed to preclude development of

Growth Factor Potential IFUKINE is a growth factor that primarily stimulates normal myeloid precursors. However, the possibility that IFUKINE can act as a growth factor for any tumor type, particularly myeloid malignancies, cannot be excluded. Because of the possibility of tumor growth potentiation, precaution should be exercised when using this drug in any malignancy with myeloid characteristics.

Should disease progression be detected during LEUKINE treatment, LEUKINE therapy should be discontinued.

LEUKINE has been administered to patients with myelodysplastic syndromes (MDS) in uncontrolled studies without evidence of increased relapse rates. 13, 14, 15 Controlled studies have not been performed in patients with MDS.

Use in Patients Receiving Purged Bone Marrow LEUKINE is effective in accelerating myeloid recovery in patients receiving bone marrow purged by anti-B lymphocyte monoclonal antibodies. Data obtained from uncontrolled studies suggest that if in vitro marrow purging with chemical agents causes a significant decrease in the number of responsive hematopoietic progenitors, the patient may not respond to LEUKINE. When the bone marrow purging process preserves a sufficient number of progenitors (>1.2 x 104/kg), a beneficial effect of LEUKINE on myeloid engraftment has been reported.16

Use in Patients Previously Exposed to Intensive Chemotherapy/Radiotherapy In patients who before autologous BMT, have received extensive radiotherapy to hematopoietic sites for the treatment of primary disease in the abdomen or chest, or have been exposed to multiple myelotoxic agents (alkylating agents, anthracycline antibiotics and antimetabolites), the effect of LEUKINE on myeloid reconstitution may be limited.

Use in Patients with Malignancy Undergoing LEUKINE-Mobilized PBPC Collection When using LEUKINE to mobilize PBPC, the limited in vitro data suggest that tumor cells may be released and reinfused into the patient in the leukapheresis product. The effect of reinfusion of tumor cells has not been well studied and the data are inconclusive

Patient Monitoring I ELIKINE can induce variable increases in WBC and/or platelet counts. In order to avoid potential complications of excessive leukocytosis (WBC >50,000 cells/mm³; ANC >20,000 cells/mm³), a CBC is recommended twice per week during LEUKINE therapy. Monitoring of renal and hepatic function in patients displaying renal or hepatic dysfunction prior to initiation of treatment is recommended at least biweekly during LEUKINE administration. Body weight and hydration status should be carefully monitored during LEUKINE administration.

Drug Interaction Interactions between LEUKINE and other drugs have not been fully evaluated. Drugs which may potentiate the myeloproliferative effects of LEUKINE, such as lithium and corticosteroids, should be used with caution.

Carcinogenesis. Mutagenesis, Impairment of Fertility Animal studies have not been conducted with LEUKINE to evaluate the carcinogenic potential or the effect on fertility

Pregnancy (Category C) Animal reproduction studies have not been conducted with LEUKINE. It is not known whether LEUKINE can cause fetal harm when administered to a pregnant woman or can affect reproductive capability. LEUKINE should be given to a pregnant woman only if clearly needed.

Nursing Mothers It is not known whether LEUKINE is excreted in human milk. Because many drugs are excreted in human milk. LEUKINE should be administered to a nursing woman only if clearly needed.

Pediatric Use Safety and effectiveness in pediatric patients have not been established; however, available safety data indicate that LEUKINE does not exhibit any greater toxicity in pediatric patients than in adults. A total of 124 pediatric subjects between the ages of 4 months and 18 years have been treated with LEUKINE in clinical trials at doses ranging from 60-1,000 mcg/m²/day intravenously and 4-1,500 mcg/m²/day subcutaneously. In 53 pediatric patients enrolled in controlled studies at a dose of 250 mcg/m²/day by 2-hour IV infusion, the type and frequency of adverse events were comparable to those reported for the adult population. Liquid solutions containing benzyl alcohol (including LEUKINE Liquid) or lyophilized LEUKINE reconstituted with Bacteriostatic Water for Injection, USP (0.9% benzyl alcohol) should not be administered to neonates (see WARNINGS).

Geriatric Use In the clinical trials, experience in older patients (age ≥65 years), was limited to the acute myelogenous leukemia (AML) study. Of the 52 patients treated with LEUKINE in this randomized study, 22 patients were age 65-70 years and 30 patients were age 55-64 years. The number of placebo patients in each age group were 13 and 33 patients respectively. This was not an adequate database from which determination of differences in efficacy endpoints or safety assessments could be reliably made and this clinical study was not designed to evaluate difference between these two age groups. Analyses of general trends in safety and efficacy were undertaken and demonstrate similar patterns for older (65-70 yrs) vs younger patients (55-64 yrs). Greater sensitivity of some older individuals cannot be ruled out.

ADVERSE REACTIONS

Autologous and Allogeneic Bone Marrow
Transplantation LEUKINE is generally well
tolerated. In 3 placebo-controlled studies
enrolling a total of 156 patients after autologous
BMT or peripheral blood progenitor cell
transplantation events reported in at least

10% of patients who received IV LEUKINE or placebo were as reported at right: No significant differences were observed between LEUKINE and placebo-treated patients. in the type or frequency of laboratory abnormalities, including renal and hepatic parameters. In some patients with preexisting renal or hepatic dysfunction enrolled in uncontrolled clinical trials, administration of LEUKINE has induced elevation of serum creatinine or bilirubin and hepatic enzymes (see WARNINGS). In addition, there was no significant difference in relapse rate and 24 month survival between the LEUKINE and placebo-treated patients.

	Percent of AuBMT Patients Reporting Events									
	LEUKINE	Placebo		LEUKINE	Placebo					
Events by Body System	(n=79)	(n=77)	Events by Body System	(n=79)	(n=77)					
Body, General			Metabolic, Nutritional Disor	rder						
Fever	95	96	Edema	34	35					
Mucous membrane disorder	75	78	Peripheral edema	11	7					
Asthenia	66	51	Respiratory System							
Malaise	57	51	Dyspnea	28	31					
Sepsis	11	14	Lung disorder	20	23					
Digestive System			Hemic and Lymphatic Syste	m						
Nausea	90	96	Blood dyscrasia	25	27					
Diarrhea	89	82	Cardiovascular System							
Vomiting	85	90	Hemorrhage	23	30					
Anorexia	54	58	Urogenital System							
GI disorder	37	47	Urinary tract disorder	14	13					
GI hemorrhage	27	33	Kidney function abnormal	8	10					
Stomatitis	24	29	Nervous System							
Liver damage	13	14	CNS disorder	11	16					
Skin and Appendages										
Alopecia	73	74								
Rash	44	38								

In the placebo-controlled trial of 109 natients after allogeneic BMT events reported in at least 10% of patients who received IV LELIKINE or placebo were as reported at right-

There were no significant differences in the incidence or severity of GVHD, relapse rates and survival between the LEUKINE and placebotreated patients.

Adverse events observed for the patients treated with LEUKINE (sargramostim) in the historically controlled BMT failure study were similar to those reported in the placebocontrolled studies. In addition, headache (26%), pericardial effusion (25%), arthralgia (21%) and myalgia (18%) were also reported in patients treated with LEUKINE in the graft failure study.

In uncontrolled Phase I/II studies with LEUKINE in 215 patients, the most frequent adverse events were fever, asthenia, headache, bone pain, chills and myalgia. These systemi events were generally mild or moderate and were usually prevented or reversed by the administration of analgesics and antipyretics such as acetaminophen. In these uncontrolled trials, other infrequent events reported were dyspnea, peripheral edema, and rash.

Reports of events occurring with marketed LEUKINE include arrhythmia, fainting, eosinophilia, dizziness, hypotension, injection site reactions, pain (including abdominal, back, chest, and joint pain), tachycardia, thrombosis, and transient liver function abnormalities

In natients with preexisting edema capillary leak syndrome, pleural and/or nericardial effusion administration of LEUKINE may appravate fluid retention (see WARNINGS) Body weight and hydration status should be carefully monitored during LEUKINE administration.

Acute Myelogenous Leukemia Adverse events reported in at least 10% of patients who received LEUKINE or placebo were as reported at right:

Nearly all patients reported leukopenia. thrombocytopenia and anemia. The frequency and type of adverse events observed following induction were similar between LEUKINE and placebo groups. The only significant difference in the rates of these adverse events was an increase in skin associated events in the LEUKINE group (p=0.002). No significant differences were observed in laboratory results, renal or hepatic toxicity. No significant differences were observed between the LEUKINE- and placebo-treated patients for adverse events following consolidation. There was no significant difference in response rate or relanse rate.

In a historically controlled study of 86 patients with acute myelogenous leukemia (AML), the LEUKINE treated group exhibited an increased incidence of weight gain (p=0.007). low serum proteins and prolonged prothrombin time (p=0.02) when compared to the control group. Two I FUKINE treated patients had

progressive increase in circulating monocytes.

and promonocytes and blasts in the marrow which reversed when LEUKINE was discontinued. The historical control group exhibited an increased incidence of cardiac events (p=0.018), liver function abnormalities (p=0.008), and neurocortical hemorrhagic events (p=0.025).¹

Overdosage The maximum amount of LEUKINE that can be safely administered in single or multiple doses has not been determined. Doses up to 100 mcg/kg/day (4,000 mcg/m²/day or 16 times the recommended dose) were administered to 4 patients in a Phase I uncontrolled clinical study by continuous IV infusion for 7 to 18 days. Increases in WBC up to 200,000 cells/mm³ were observed. Adverse events reported were dyspnea, malaise, nausea, fever, rash, sinus tachycardia, headache and chills. All these events were reversible after discontinuation of LEUKINE

In case of overdosage, LEUKINE therapy should be discontinued and the patient carefully monitored for WBC increase and respiratory symptoms.

DOSAGE AND ADMINISTRATION

Neutrophil Recovery Following Chemotherapy in Acute Myelogenous Leukemia The recommended dose is 250 mcg/m²/day administered intravenously over a 4 hour period starting approximately on day 11 or 4 days following the completion of induction chemotherapy, if the day 10 bone marrow is hypoplastic with <5% blasts. If a second cycle of induction chemotherapy is necessary, LEUKINE should be administered approximately 4 days after the completion of chemotherapy if the bone marrow is hypoplastic with <5% blasts. LEUKINE should be continued until an ANC >1500 cells/mm³ for 3 consecutive days or a maximum of 42 days. LEUKINE should be discontinued immediately if leukemic regrowth occurs. If a severe adverse reaction occurs, the dose can be reduced by 50% or temporarily discontinued until the reaction abates

In order to avoid potential complications of excessive leukocytosis (WBC > 50,000 cells/mm³ or ANC > 20,000 cells/mm³) a CBC with differential is recommended twice per week during LEUKINE therapy. LEUKINE treatment should be interrupted or the dose reduced by half if the ANC exceeds 20,000 cells/mm³.

Mobilization of Perinheral Blood Progenitor Cells The recommended dose is 250 mcg/m²/day administered IV over 24 hours or SC once daily. Dosing should continue at the same dose through the period of PRPC collection. The optimal schedule for PRPC collection has not been established. In clinical studies, collection of PRPC was usually begun by day 5 and performed daily until protocol specified targets were achieved (see CLINICAL EXPERIENCE Mobilization and Engraftment of PBPC). If WBC > 50,000 cells/mm3, the LEUKINE dose should be reduced by 50%. If adequate numbers of progenitor cells are not collected, other mobilization therapy should be considered.

LEUKINE (sargramostim)

Parcent of Allogonaic RMT Datients Reporting Events									
I GIU		Disasha	rations neporting Events		Dlaasha				
Evente hy Rody Svetem	(n=53)	Place00 (n_56)	Events by Rody System	(n=53)	Placebo (n=56)				
	(11=33)	(11=30)		(11=33)	(11=30)				
Body, General	77	0.0	Metabolic/Nutritional Diso	raers	07				
Fever	//	80	Bilirubinemia	30	27				
Abdominal pain	38	23	Hyperglycemia	25	23				
Headache	36	36	Peripheral edema	15	21				
Chills	25	20	Increased creatinine	15	14				
Pain	17	36	Hypomagnesemia	15	9				
Asthenia	17	20	Increased SGPT	13	16				
Chest pain	15	9	Edema	13	11				
Back pain	9	18	Increased alk. phosphatase	8	14				
Digestive System			Respiratory System						
Diarrhea	81	66	Pharyngitis	23	13				
Nausea	70	66	Epistaxis	17	16				
Vomiting	70	57	Dyspnea	15	14				
Stomatitis	62	63	Rhinitis	11	14				
Anorexia	51	57	Hemic and Lymphatic Syst	em					
Dyspepsia	17	20	Thrombocytopenia	19	34				
Hematemesis	13	7	Leukopenia	17	29				
Dysphagia	11	7	Petechia	6	11				
GI hemorrhage	11	5	Agranulocytosis	6	11				
Constipation	8	11	Urogenital System						
Skin and Appendages			Hematuria	9	21				
Rash	70	73	Nervous System						
Alopecia	45	45	Paresthesia	11	13				
Pruritis	23	13	Insomnia	11	9				
Musculo-skeletal System			Anxiety	11	2				
Bone pain	21	5	Laboratory Abnormalities*						
Arthralgia	11	4	High glucose	41	49				
Special Senses			Low albumin	27	36				
Eve hemorrhade	11	0	High BUN	23	17				
Cardiovascular System			Low calcium	2	7				
Hypertension	34	32	High cholesterol	17	8				
Tachycardia	11	9	5		-				
*Crade 2 and 4 Jahoratory about	malition only	Donominator	n may yany duo to missing laborato	nu moneuromo	ato				

Adverse events observed in pediatric patients in controlled studies were comparable to those observed in adult patients.

Percent of AML Patients Reporting EventsLEUKINEPlaceboLEUKINEPlaceEvents by Body System(n=52)(n=47)Events by Body System(n=52)(n=47)Body, GeneralKetabolic/Nutritional DisorderKetabolic/Nutritional Disorder(n=52)(n=47)Fever (no infection)8174Metabolic/Nutritional Disorder49Infection6568Edema2523Weight loss3728Respiratory System4864Chills1926Hemic and Lymphatic System1921Allergy1215Coagulation1921Sweats613Cardiovascular System1926Digestive SystemHemorrhage2943Nausea5855Hypertension2532Liver7783Cardiac2332Diarrhea5253Hypotension1326Vomiting4634Urogenital System5057Anorexia1311Neuro-clinical4255Skin and Appendages43Neuro-clinical4252					
	LEUKINE	Placebo		LEUKINE	Placebo
Events by Body System	(n=52)	(n=47)	Events by Body System	(n=52)	(n=47)
Body, General			Metabolic/Nutritional Disc	order	
Fever (no infection)	81	74	Metabolic	58	49
Infection	65	68	Edema	25	23
Weight loss	37	28	Respiratory System		
Weight gain	8	21	Pulmonary	48	64
Chills	19	26	Hemic and Lymphatic Sys	tem	
Allergy	12	15	Coagulation	19	21
Sweats	6	13	Cardiovascular System		
Digestive System			Hemorrhage	29	43
Nausea	58	55	Hypertension	25	32
Liver	77	83	Cardiac	23	32
Diarrhea	52	53	Hypotension	13	26
Vomiting	46	34	Urogenital System		
Stomatitis	42	43	GU	50	57
Anorexia	13	11	Nervous System		
Abdominal distention	4	13	Neuro-clinical	42	53
Skin and Appendages			Neuro-motor	25	26
Skin	77	45	Neuro-psych	15	26
Alopecia	37	51	Neuro-sensory	6	11

Chloride Injection, USP to prepare IV infusion solutions HOW SUPPLIED

LEUKINE Liquid is available in vials containing 500 mcg/mL (2.8 x 10⁶ IU/mL) sargramostim. Lyophilized LEUKINE is available in vials containing 250 mcg (1.4 x 10⁶ IU/vial) saroramostim. Each dosage form is supplied as follows: Carton of 5 vials of Ivophilized LEUKINE 250 mcg (NDC 50419-002-33)

STORAGE

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immediately if blast cells appear or disease progression occurs.

should not be mixed together.

Preparation of LEUKINE

LEUKINE (sargramostim)

Post Peripheral Blood Progenitor Cell Transplantation The recommended dose is 250 mcg/m²/day administered IV over 24 hours or SC once daily beginning immediately following infusion of progenitor cells and continuing until an ANC>1500 cells/mm³ for 3 consecutive days is attained.

Myeloid Reconstitution After Autologous or Allogeneic Bone Marrow Transplantation The recommended dose is 250 mcg/m²/day administered IV over a 2-hour period beginning 2 to 4 hours after bone marrow infusion, and not less than 24 hours after the last dose of chemotherapy or radiotherapy. Patients should not receive LEUKINE until the post marrow infusion ANC is less than 500 cells/mm³. LEUKINE should be continued until an ANC >1500 cells/mm³ for 3 consecutive days is attained. If a severe adverse reaction occurs, the dose can be reduced by 50% or temporarily discontinued until the reaction abates. LEUKINE should be discontinued

In order to avoid potential complications of excessive leukocytosis (WBC > 50,000 cells/mm³, ANC > 20,000 cells/mm³) a CBC with differential is recommended twice per week during LEUKINE therapy. LEUKINE treatment should be interrupted or the dose reduced by 50% if the ANC exceeds 20,000 cells/mm³

Bone Marrow Transplantation Failure or Engraftment Delay The recommended dose is 250 mcg/m²/day for 14 days as a 2-hour IV infusion. The dose can be repeated after 7 days off therapy if engraftment has not occurred. If engraftment still has not occurred, a third course of 500 mcg/m²/day for 14 days may be tried after another 7 days off therapy. If there is still no improvement, it is unlikely that further dose escalation will be beneficial. If a severe adverse reaction occurs, the dose can be reduced by 50% or temporarily discontinued until the reaction abates. LEUKINE should be discontinued immediately if blast cells appear or disease progression occurs. In order to avoid potential complications of excessive leukocytosis (WBC > 50,000 cells/mm³, ANC > 20,000 cells/mm³) a CBC with differential is recommended twice per week during LEUKINE therapy. LEUKINE treatment should be interrupted or the dose reduced by half if the ANC exceeds 20,000 cells/mm³

1. LEUKINE Liguid is formulated as a sterile, preserved (11% benzyl alcohol), injectable solution (500 mcg/mL) in a vial. Lyophilized LEUKINE is a sterile, white, preservative-free powder (250 mcg) that requires reconstitution with 1 mL Sterile Water for Injection, USP, or 1 mL Bacteriostatic Water for Injection, USP. 2. LEUKINE Liquid may be stored for up to 20 days at 2-8°C once the vial has been entered. Discard any remaining solution after 20 day 3. Lyophilized LEUKINE (250 mcg) should be reconstituted aseptically with 10 mL of diluent (see below). The contents of vials reconstituted with different diluents

Sterile Water for Injection, USP (without preservative): Lyophilized LEUKINE vials contain no antibacterial preservative, and therefore solutions prepared with Sterile

Water for Injection, USP should be administered as soon as possible, and within 6 hours following reconstitution and/or dilution for IV infusion. The vial should not be re-entered or reused. Do not save any unused portion for administration more than 6 hours following reconstitution.

Bacteriostatic Water for Injection, USP (0.9% benzyl alcohol): Reconstituted solutions prepared with Bacteriostatic Water for Injection, USP (0.9% benzyl alcohol) may be stored for up to 20 days at 2-8°C prior to use. Discard reconstituted solution after 20 days. Previously reconstituted solutions mixed with freshly reconstituted solutions must be administered within 6 hours following mixing. Preparations containing benzyl alcohol (including LEUKINE Liquid and

lyophilized LEUKINE reconstituted with Bacteriostatic Water for Injection) should not be used in neonates (see WARNINGS).

4. During reconstitution of lyophilized LEUKINE the diluent should be directed at the side of the vial and the contents gently swirled to avoid foaming during dissolution. Avoid excessive or vigorous agitation; do not shake.

5. LEUKINE should be used for SC injection without further dilution. Dilution for IV infusion should be performed in 0.9% Sodium Chloride Injection, USP. If the final concentration of LEUKINE is below 10 mcg/mL, Albumin (Human) at a final concentration of 0.1% should be added to the saline prior to addition of LEUKINE to prevent adsorption to the components of the drug delivery system. To obtain a final concentration of 0.1% Albumin (Human), add 1 mg Albumin (Human) per 1 mL 0.9% Sodium Chloride Injection, USP (e.g., use 1 mL 5% Albumin [Human] in 50 mL 0.9% Sodium Chloride Injection, USP).

An in-line membrane filter should NOT be used for intravenous infusion of LEUKINE.

Store LEUKINE Liquid and reconstituted lyophilized LEUKINE solutions under refrigeration at 2-8°C (36-46°F); DO NOT FREEZE.

8. In the absence of compatibility and stability information, no other medication should be added to infusion solutions containing LEUKINE. Use only 0.9% Sodium

9. Aseptic technique should be employed in the preparation of all LEUKINE solutions. To assure correct concentration following reconstitution, care should be exercised to eliminate any air bubbles from the needle hub of the syringe used to prepare the diluent. Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration whenever solution and container permit.

Carton of 5 multiple-dose vials; each vial contains 1 mL of preserved 500 mcg/mL LEUKINE Liquid (NDC 50419-050-30).

LEUKINE should be refrigerated at 2-8°C (36-46°F). Do not freeze or shake. Do not use beyond the expiration date printed on the vial.

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Berlex Laboratories, Inc. U.S. Patent Nos. 5,391485; 5,393,870; and 5,229,496. Licensed under Research Corporation Technologies U.S. Patent No. 5,602,007, and under Novartis Corporation U.S. Patent Nos. 5,942,221; 5,908,763; 5,895,646; 5,891,429; and 5,720,952.