

PRODUCT MONOGRAPH

Pr CYTOSAR[®]

**Cytarabine for injection, USP
Lyophilized Powder for Injection
(100 mg, 500 mg, 1000 mg and 2000 mg)**

**Cytarabine Solution for Injection, House Std
Solution for Injection
(20 mg/mL and 100 mg/mL)**

Antileukemic Agent

Pfizer Canada Inc.
17,300 Trans-Canada Highway
Kirkland, Quebec, H9J 2M5

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PRODUCT MONOGRAPH

^{Pr}CYTOSAR[®] Cytarabine for injection USP

Antileukemic Agent

CAUTION: CYTOSAR (CYTARABINE) SHOULD BE USED ONLY BY PHYSICIANS EXPERIENCED WITH CANCER THERAPY DRUGS (SEE **WARNINGS AND PRECAUTIONS**). HEMATOLOGIC, RENAL, AND HEPATIC EVALUATIONS MUST BE DONE AT REGULAR INTERVALS.

ACTION AND CLINICAL PHARMACOLOGY

CYTOSAR (cytarabine) is metabolized by deoxycytidine kinase and other nucleotide kinases to the nucleotide triphosphate, an effective inhibitor of DNA polymerase; it is inactivated by pyrimidine nucleoside deaminase which converts it to the non-toxic uracil derivative. It appears that the balance of kinase and deaminase levels may be an important factor in determining sensitivity or resistance of the cell to cytarabine.

CYTOSAR is rapidly metabolized and is not effective orally; less than 20% of the orally administered dose is absorbed from the gastrointestinal tract.

Following rapid intravenous injection of CYTOSAR, the disappearance from plasma is biphasic. There is an initial distributive phase with a half-life of about 10 minutes, followed by a second elimination phase with a half-life of about 1 to 3 hours. After the distributive phase, over 80% of plasma radioactivity can be accounted for by the inactive metabolite 1- β -D-arabinofuranosyluracil (ara-U). Within 24 hours about 80% of the administered radioactivity can be recovered in the urine, approximately 90% of which is excreted as ara-U.

After subcutaneous or intramuscular administration of CYTOSAR, peak plasma levels of radioactivity are achieved about 20 to 60 minutes after injection and are considerably lower than those after intravenous administration.

Cerebrospinal fluid levels of cytarabine are low in comparison to plasma levels after single intravenous injection. However, in one patient in whom cerebrospinal levels were examined after 2 hours of constant intravenous infusion, levels approached 40% of the steady state plasma level. With intrathecal administration, levels of cytarabine in the cerebrospinal fluid declined with a first order half-life of about 2 hours. Because cerebrospinal fluid levels of deaminase are low, little conversion to ara-U was observed.

INDICATIONS AND CLINICAL USE

CYTOSAR (cytarabine) is indicated primarily for induction and maintenance of remission in acute leukemia in both adults and children.

It has been found useful in the treatment of acute myelocytic leukemia, chronic myelocytic leukemia (blast phase), acute lymphocytic leukemia and erythroleukemia. CYTOSAR may be used alone or in combination with other antineoplastic agents; the best results are obtained with combination therapy.

Children with non-Hodgkin's lymphoma have benefited from a combination drug program (LSA₂L₂) that included CYTOSAR.

CYTOSAR has been used intrathecally in newly diagnosed children with acute lymphocytic leukemia as well as in the treatment of meningeal leukemia.

CYTOSAR, in high dose 2-3 g/m² as an i.v. infusion over 1-3 hours given every 12 hours for 2-6 days with or without additional cancer chemotherapeutic agents, has been shown to be effective in the treatment of poor-risk leukemia, refractory leukemia, and relapsed acute leukemia.

Remissions induced by CYTOSAR not followed by maintenance treatment have been brief.

CONTRAINDICATIONS

CYTOSAR (cytarabine) is contraindicated in those patients who are hypersensitive to the drug.

WARNINGS AND PRECAUTIONS

General:

For induction therapy, patients should be treated in a facility with laboratory and supportive resources sufficient to monitor drug tolerance and protect and maintain a patient compromised by drug toxicity. The main toxic effect of CYTOSAR is bone marrow suppression with leukopenia, thrombocytopenia and anemia. Less serious toxicity includes nausea, vomiting, diarrhea and abdominal pain, oral ulceration, and hepatic dysfunction.

The physician must judge possible benefit to the patient against known toxic effects of this drug in considering the advisability of therapy with CYTOSAR. Before making this judgment or beginning treatment, the physician should be familiar with the following text.

When large intravenous doses are given quickly, patients are frequently nauseated and may vomit for several hours post injection. This problem tends to be less severe when the drug is infused.

Bacteriostatic water, one of the diluents recommended for reconstitution of CYTOSAR, contains benzyl alcohol (see PHARMACEUTICAL INFORMATION, Reconstitution of Lyophilized Powder). Benzyl alcohol has been reported to be associated with a fatal "Gaspings Syndrome" in pediatric patients. As premature and low birth weight infants may be at increased risk of developing this toxicity, they should not be given cytarabine reconstituted with a diluent containing benzyl alcohol (see WARNINGS AND PRECAUTIONS, Special Populations, Pediatrics).

If high dose therapy is used, do not use a diluent containing benzyl alcohol (see PHARMACEUTICAL INFORMATION, Reconstitution of Lyophilized Powder).

Do not use a diluent containing benzyl alcohol if using intrathecally (see PHARMACEUTICAL INFORMATION, Reconstitution of Lyophilized Powder).

Carcinogenesis, Mutagenesis, Impairment of Fertility:

Extensive chromosomal damage, including chromatoid breaks have been produced by cytarabine and malignant transformation of rodent cells in culture has been reported. Cytarabine is embryotoxic and teratogenic and produced peri- and postnatal toxicity in various species. Sperm head abnormalities were observed following cytarabine treatment in mice. (See TOXICOLOGY)

Cardiovascular:

High dose schedules: An increase in cardiomyopathy with subsequent death has been reported following experimental high dose CYTOSAR and cyclophosphamide therapy when used for bone marrow transplant preparation. This may be schedule dependent.

Gastrointestinal:

Abdominal tenderness (peritonitis) and Typhlitis with concurrent neutropenia and thrombocytopenia, have been reported in patients treated with conventional doses of cytarabine in combination with other drugs. Patients have responded to nonoperative medical management.

High dose schedule: Severe and at times fatal, GI toxicity (different from that seen with conventional therapy regimens of CYTOSAR) has been reported following high dose (2-3 g/m²) schedules of CYTOSAR). These reactions include severe gastrointestinal ulceration, including pneumatosis cystoides intestinalis, leading to peritonitis, bowel necrosis; and necrotizing colitis.

Genitourinary:

Tumor Lysis Syndrome: Like other cytotoxic drugs, CYTOSAR may induce hyperuricemia secondary to rapid lysis of neoplastic cells. The clinician should monitor the patient's blood uric acid level and be prepared to use such supportive and pharmacologic measurements as might be necessary to control this problem.

Hematologic Effects:

CYTOSAR (cytarabine) is a potent bone marrow suppressant; the severity depends on the dose of the drug and schedule of administration. Therapy should be started cautiously in patients with pre-existing drug-induced bone marrow suppression. Patients receiving this drug must be under close medical supervision and during induction therapy, should have leukocyte and platelet counts performed daily. Bone marrow examinations should be performed frequently after blasts

have disappeared from the peripheral blood. Facilities should be available for management of complications (possibly fatal) of bone marrow suppression (infection resulting from granulocytopenia and other impaired body defenses, and hemorrhage secondary to thrombocytopenia).

Hepatic/Biliary/Pancreatic and/or Renal Function:

The human liver apparently detoxifies a substantial fraction of an administered cytarabine dose. In particular, patients with renal or hepatic function impairment may have a higher likelihood of CNS toxicity after high-dose treatment with CYTOSAR. Use the drug with caution and at reduced dose in patients whose liver function is poor.

Periodic checks of bone marrow, liver and kidney function should be performed in patients receiving CYTOSAR.

Pancreatitis: Acute pancreatitis has been reported to occur in patients being treated with CYTOSAR in combination with other drugs.

High dose schedules: Other reactions have been reported following high dose (2-3 g/m² schedules of CYTOSAR) and include sepsis and liver abscess, and liver damage with increased hyperbilirubinemia.

Hypersensitivity Reactions:

Anaphylactic reactions have occurred with cytarabine treatment. Anaphylaxis that resulted in acute cardiopulmonary arrest and required resuscitation has been reported. This occurred immediately after the intravenous administration of CYTOSAR.

Immune:

Immunosuppressant Effects/Increased Susceptibility to Infections: Administration of live or live-attenuated vaccines in patients immunocompromised by chemotherapeutic agents including cytarabine, may result in serious or fatal infections. Vaccination with a live vaccine should be avoided in patients receiving cytarabine. Killed or inactivated vaccines may be administered; however, the response to such vaccines may be diminished.

Neurologic:

High dose schedules: Severe and at times fatal, CNS toxicity (different from that seen with conventional therapy regimens of CYTOSAR) has been reported following high dose (2-3 g/m² schedules of CYTOSAR). These reactions include cerebral and cerebellar dysfunction including personality changes, somnolence, convulsion and coma, usually reversible.

Delayed progressive ascending paralysis resulting in death has been reported in children with AML following intrathecal and intravenous cytarabine at conventional doses in combination with other drugs.

Cases of severe neurological adverse reactions that ranged from headache to paralysis, coma and stroke-like episodes have been reported mostly in pediatric patients given intravenous cytarabine in combination with intrathecal methotrexate.

Ophthalmologic:

High dose schedules: The following reactions have been reported following high dose (2-3 g/m²) schedules of CYTOSAR): reversible corneal toxicity and hemorrhagic conjunctivitis, which may be prevented or diminished by prophylaxis with a local corticosteroid eye drop.

Respiratory:

High dose schedules: Severe and sometimes fatal pulmonary toxicity, adult respiratory distress syndrome and pulmonary edema have occurred following high dose schedules with cytarabine therapy. A syndrome of sudden respiratory distress, rapidly progressing to pulmonary edema and radiographically pronounced cardiomegaly has been reported following experimental high dose CYTOSAR therapy used for the treatment of relapsed leukemia.

Skin:

Palmar plantar erythrodysesthesia: Palmar plantar erythrodysesthesia (PPE) has occurred with cytarabine treatment in adults and children. Severe cytarabine associated PPE that resulted in treatment discontinuation has been reported.

High dose schedules: Rarely, severe skin rash, leading to desquamation has been reported. Complete alopecia is more commonly seen with high dose therapy than with standard CYTOSAR treatment programs.

Special Populations**Pregnant Women:**

There are no studies on the use of cytarabine in pregnant women. CYTOSAR is known to be teratogenic in some animal species. Use of this drug in women who are or who may become pregnant should be undertaken only after due consideration of potential benefit and potential hazard to both mother and child. Women of childbearing potential should be advised to avoid becoming pregnant.

Normal infants have been born to mothers exposed to cytarabine during pregnancy (alone or in combination with other drugs); some of these infants were premature or of low birthweight. Some of the normal infants were followed up at ages ranging from six weeks to seven years following exposure, and showed no abnormalities. One apparently normal infant died at 80 days of gastroenteritis.

Congenital abnormalities have been reported, particularly when the fetus has been exposed to systemic therapy with cytarabine during the first trimester. These include upper and lower distal limb defects, and extremity and ear deformities.

Reports of pancytopenia, leucopenia, anemia, thrombocytopenia, electrolyte abnormalities, transient eosinophilia, increased IgM levels and hyperpyrexia, sepsis and death have occurred during the neonatal period to infants exposed to cytarabine in utero. Some of these infants were also premature.

Therapeutic abortions have been done in pregnant women on cytarabine. Normal fetuses have been reported while other reported fetal effects included enlarged spleen and Trisomy C chromosome abnormality in the chorionic tissue.

Because of the potential for abnormalities with cytotoxic therapy, particularly during the first trimester, a patient who is or who becomes pregnant while on CYTOSAR should be apprised of the potential risk to the fetus and the advisability of pregnancy continuation. There is a definite, but considerably reduced risk if therapy is initiated during the second or third trimester. Although normal infants have been delivered to patients treated in all three trimesters of pregnancy, follow-up of such infants would be advisable.

Bacteriostatic water, one of the diluents recommended for reconstitution of CYTOSAR, contains benzyl alcohol. Benzyl alcohol can cross the placenta (see WARNINGS AND PRECAUTIONS, Special Populations, Pediatrics).

Nursing Women:

It is not known whether this drug is excreted in human milk. Because many drugs are excreted in human milk and because of the potential for serious adverse reactions in nursing infants from cytarabine, a decision should be made whether to discontinue nursing or to discontinue the drug, taking into account the importance of the drug to the mother.

Pediatrics:

The safety of this drug for use in infants (under 1 year of age) is not established.

Gasping Syndrome: Cytarabine should not be given to premature and low birth weight infants when using a diluent that contains benzyl alcohol. The preservative benzyl alcohol has been associated with serious adverse events, including the “gasping syndrome”, and death in pediatric patients. Symptoms of gasping syndrome may include metabolic acidosis, seizure, bradycardia, gasping respiration and cardiovascular collapse. Although normal therapeutic doses of this product ordinarily deliver amounts of benzyl alcohol that are substantially lower than those reported in association with the “gasping syndrome”, the minimum amount of benzyl alcohol at which toxicity may occur is not known. The risk of benzyl alcohol toxicity depends on the quantity administered and the hepatic capacity to detoxify the chemical. Premature and low-birth weight infants may be more likely to develop toxicity. If cytarabine is used in high dose or intrathecal therapy, do not use a diluent containing benzyl alcohol. The preservative-free 0.9% sodium chloride can be used for reconstitution.

See also WARNING AND PRECAUTIONS, Neurologic.

Monitoring and Laboratory Tests:

Patients receiving CYTOSAR (cytarabine) must be monitored closely. Frequent platelet and leukocyte counts and bone marrow examinations are mandatory. Consider suspending or modifying therapy when drug-induced marrow depression has resulted in a platelet count under 50,000 or a polymorphonuclear granulocyte count under $1000/\text{mm}^3$. Counts of formed elements in the peripheral blood may continue to fall after the drug is stopped and reach lowest values after drug-free intervals of 12 of 24 days. When indicated, restart therapy when definite signs of

marrow recovery appear (on successive bone marrow studies). Patients whose drug is withheld until "normal" peripheral blood values are attained, may escape from control.

Interaction with other medicinal products:

Digoxin: Reversible decreases in steady-state plasma digoxin concentrations and renal glycoside excretion were observed in patients receiving beta-acetyldigoxin and chemotherapy regimens containing cyclophosphamide, vincristine and prednisone with or without cytarabine or procarbazine. Steady-state plasma digitoxin concentrations did not appear to change. Therefore, monitoring of plasma digoxin levels may be indicated in patients receiving similar combination chemotherapy regimens. The utilization of digitoxin for such patients may be considered as an alternative.

Gentamicin: An *in vitro* interaction study between gentamicin and cytarabine showed a cytarabine related antagonism for the susceptibility of *K. pneumoniae* strains. This study suggests that in patients on cytarabine being treated with gentamicin for a *K. pneumoniae* infection, the lack of a prompt therapeutic response may indicate the need for re-evaluation of antibacterial therapy.

Fluorocytosine: Clinical evidence showed possible inhibition of fluorocytosine efficacy therapy with cytarabine. This may be due to potential competitive inhibition of its uptake.

Methotrexate: Intravenous cytarabine given concomitantly with intrathecal methotrexate may increase the risk of severe neurological adverse reactions such as headache, paralysis, coma and stroke like episodes.

ADVERSE REACTIONS

Blood and Lymphatic System Disorders:

Because CYTOSAR (cytarabine) is a bone marrow suppressant, anemia, leukopenia, thrombocytopenia, megaloblastosis, and reduced reticulocytes can be expected as a result of its administration. The severity of these reactions is dose and schedule dependent. Cellular changes in the morphology of bone marrows and peripheral smears can be expected.

Following 5-day constant infusions or acute injections of 50 mg/m² to 600 mg/m², white cell depression follows a biphasic course. Regardless of initial count, dosage level, or schedule, there is an initial fall starting the first 24 hours with a nadir at days 7 to 9. This is followed by a brief rise which peaks around the twelfth day. A second and deeper fall reaches nadir at days 15 to 24. Then there is a rapid rise to above baseline in the next 10 days. Platelet depression is noticeable at 5 days with a peak depression occurring between days 12 to 15. Thereupon, a rapid rise to above baseline occurs in the next 10 days.

Infections and Infestations

Viral, bacterial, fungal, parasitic, or saprophytic infections, in any location on the body, may be associated with the use of CYTOSAR alone or in combination with other immunosuppressive agents following immunosuppressive doses that affect cellular or humoral immunity. These infections may be mild, but can be severe and at times fatal.

Musculoskeletal and connective tissue disorders

The Cytarabine Syndrome:

A cytarabine syndrome has been described by Castleberry. It is characterized by fever, myalgia, bone pain, occasionally chest pain, maculopapular rash, conjunctivitis and malaise. It usually occurs 6 to 12 hours following drug administration. Corticosteroids have been shown to be beneficial in treating or preventing this syndrome. If the symptoms of the syndrome are deemed treatable, corticosteroids should be contemplated as well as continuation of therapy with CYTOSAR.

Other adverse reactions:

Conventional Dose Therapy

Nausea and vomiting are most frequent following rapid intravenous injection.

Table 1 - Adverse Reactions (with conventional dose therapy) : The reported adverse reactions are listed below by MedDRA System Organ Class and by frequency. Frequencies are defined as: Very common (>10%), Common (>1%, ≤10%), Uncommon (>0.1%, ≤1%), Rare (>0.01%, ≤0.1%), and Frequency not known (cannot be estimated from available data).

| | |
|---|---|
| Blood and Lymphatic System Disorders: | |
| Very common | Bone marrow failure, thrombocytopenia, anaemia, anaemia megaloblastic, leukopenia, reticulocyte count decreased |
| Frequency not known | Bleeding (all sites) |
| Cardiac Disorders: | |
| Frequency not known | Pericarditis |
| Eye Disorders: | |
| Frequency not known | Conjunctivitis ^a |
| Gastrointestinal Disorders: | |
| Very common | Stomatitis, mouth ulceration, anal ulcer, anal inflammation, diarrhoea, vomiting, nausea, abdominal pain |
| Frequency not known | Bowel necrosis, Pancreatitis, oesophageal ulcer, oesophagitis |
| General Disorders and Administration Site Conditions: | |
| Very common | Pyrexia |
| Frequency not known | Chest pain, injection site reaction ^b |
| Hepatobiliary Disorders: | |
| Very common | Hepatic function abnormal |
| Frequency not known | Jaundice |
| Infections and Infestations: | |
| Very common | Sepsis, pneumonia, infection ^c |
| Frequency not known | Injection site cellulitis |
| Immune System Disorders: | |
| Frequency not known | Anaphylactic reaction, allergic oedema |
| Investigations: | |
| Very common | Biopsy bone marrow abnormal, blood smear test abnormal |
| Metabolism and Nutrition Disorders: | |
| Frequency not known | Decreased appetite |
| Musculoskeletal, Connective Tissue and Bone Disorders: | |
| Very common | Cytarabine syndrome |
| Nervous System Disorders: | |
| Frequency not known | Neurotoxicity, neuritis, dizziness, headache |
| Renal and Urinary Disorders: | |
| Frequency not known | Renal impairment, urinary retention |
| Respiratory, Thoracic and Mediastinal Disorders: | |
| Frequency not known | Dyspnoea, oropharyngeal pain |

| | |
|--|--|
| Skin and Subcutaneous Tissue Disorders: | |
| Very common | Alopecia, rash |
| Common | Skin ulcer |
| Frequency not known | Palmar-plantar erythrodysesthesia syndrome, urticaria, pruritus, freckling |
| Vascular Disorders: | |
| Frequency not known | Thrombophlebitis |

^amay occur with rash and may be hemorrhagic with high dose therapy

^bpain and inflammation at subcutaneous injection site

^cmay be mild, but can be severe and at times fatal

High Dose Therapy

Severe and at times fatal CNS, GI and pulmonary toxicity (different from that seen with conventional therapy regimens of CYTOSAR) has been reported following high dose schedules (2.0 g to 3.0 g/m² given every 12 hours for 12 doses).

Table 2: Adverse Reactions (with High Dose Therapy): The reported adverse reactions are listed below by MedDRA System Organ Class and by frequency. Frequencies are defined as: Very common (>10%), Common (>1%, ≤10%), Uncommon (>0.1%, ≤1%), Rare (>0.01%, ≤0.1%), and Frequency not known (cannot be estimated from available data).

| | |
|---|--|
| Cardiac Disorders: | |
| Frequency not known | Cardiomyopathy ^a |
| Eye Disorders: | |
| Very common | Corneal disorder |
| Frequency not known | hemorrhagic conjunctivitis ^b |
| Gastrointestinal Disorders: | |
| Common | Necrotising colitis |
| Frequency not known | Gastrointestinal necrosis, gastrointestinal ulcer, pneumatosis intestinalis, peritonitis |
| Hepatobiliary Disorders: | |
| Frequency not known | Liver injury, hyperbilirubinaemia |
| Infections and Infestations: | |
| Very common | Sepsis |
| Frequency not known | Liver abscess |
| Nervous System Disorders: | |
| Very common | Cerebral disorder, cerebellar disorder, somnolence |
| Frequency not known | Coma, convulsion, peripheral motor neuropathy, peripheral sensory neuropathy |
| Psychiatric Disorders: | |
| Frequency not known | Personality change ^c |
| Respiratory, Thoracic and Mediastinal Disorders: | |
| Very common | Acute respiratory distress syndrome, pulmonary oedema |
| Skin and Subcutaneous Tissue Disorders: | |
| Common | Skin exfoliation, |

^aWith subsequent death

^b may be prevented or diminished by prophylaxis with a local corticosteroid eyedrop

^c Personality change was reported in association with cerebral and cerebellar dysfunction.

Peripheral motor and sensory neuropathies after consolidation with high-dose CYTOSAR, daunorubicin, and asparaginase have occurred in adult patients with acute non lymphocytic leukemia. Patients treated with high-dose CYTOSAR should be observed for neuropathy since dose schedule alterations may be needed to avoid irreversible neurologic disorders.

Corneal toxicity consisting of ocular pain, tearing, foreign-body sensation, photophobia and blurred vision has been reported.

Rarely, severe skin rash, leading to desquamation has been reported. Complete alopecia is more commonly seen with high dose therapy than with standard CYTOSAR treatment programs.

If high dose therapy is used, do not use a diluent containing benzyl alcohol.

Intermediate dose therapy

A diffuse interstitial pneumonitis without clear cause that may have been related to Cytosar was reported in patients treated with experimental intermediate doses of CYTOSAR (1 gm/m²) with and without other chemotherapeutic agents (meta-AMSA, daunorubicin, VP-16).

Intrathecal therapy

CYTOSAR given intrathecally may cause systemic toxicity and careful monitoring of the hemopoietic system is indicated. Modification of other anti-leukemia therapy may be necessary. Major toxicity is rare. The most frequently reported reactions after intrathecal administration were nausea, vomiting and fever; these reactions are mild and self-limiting. Paraplegia has been reported. Necrotizing leukoencephalopathy with or without convulsion has been reported; in some cases, patients had also been treated with intrathecal methotrexate and/or hydrocortisone, as well as by central nervous system radiation. Isolated neurotoxicity has been reported. Blindness occurred in two patients in remission whose treatment had consisted of combination systemic chemotherapy, prophylactic central nervous system radiation and intrathecal CYTOSAR. When CYTOSAR is administered both intrathecally and intravenously within a few days, there is an increased risk of spinal cord toxicity, however, in serious life-threatening disease, concurrent use of intravenous and intrathecal CYTOSAR is left to the discretion of the treating physician.

SYMPTOMS AND TREATMENT OF OVERDOSAGE

There is no antidote for CYTOSAR (cytarabine) overdose.

Discontinuation of the drug and supportive therapy are of course indicated. Transfusions of platelets should be given if there is any sign of hemorrhage. Patients should be carefully observed for intercurrent infection and if such appears they should be rapidly and rigorously treated with appropriate antibiotic therapy.

Chronic overdose may cause serious bone marrow suppression. Daily hematological evaluation should be performed to prevent overdose. Nausea and vomiting, although a general side effect of the drug, may be an additional warning of overdose. Severe hemorrhage into the gastrointestinal tract may indicate overdose as may severe generalized infections.

Doses exceeding recommended dosage schedules have been used clinically and have been tolerated. The major toxicity with the use of 3 g/m² intravenous infusion over 1 hour every 12 hours for 12 doses and 3 g/m² continuous infusion for 4 days, other than reversible bone marrow suppression has been reversible corneal, cerebral and cerebellar dysfunction. Doses of 4.5 g/m² intravenous infusion over 1 hour every 12 hours for 12 doses has caused an unacceptable increase in irreversible CNS toxicity and death.

| |
|---|
| For management of a suspected drug overdose, contact your regional Poison Control Centre. |
|---|

DOSAGE AND ADMINISTRATION

CAUTION

The following precautionary measures are recommended in proceeding with the preparation and handling of cytotoxic agents such as CYTOSAR (cytarabine).

1. The procedure should be carried out in a vertical laminar flow hood (Biological Safety Cabinet - Class II).
2. Personnel should wear: PVC gloves, safety glasses, disposable gowns and masks.
3. All needles, syringes, vials, and other materials which have come in contact with CYTOSAR should be segregated and destroyed by incineration (sealed containers may explode). If incineration is not available, neutralization should be carried out using 5% sodium hypochlorite, or 5% sodium thiosulfate.
4. Personnel regularly involved in the preparation and handling of CYTOSAR should have bi-annual haematologic examinations.

CYTOSAR is not active orally. The schedule and method of administration varies with the program of therapy to be used. CYTOSAR may be given by intravenous infusion, injection/subcutaneously or intrathecally. When preparing cytarabine for intravenous high dose therapy or intrathecal use, do not use diluents containing benzyl alcohol (see PHARMACEUTICAL INFORMATION, Reconstitution of Lyophilized Powder and Handling of Solutions for Injection). It is recommended that CYTOSAR be reconstituted with preservative-free 0.9% sodium chloride for injection and used immediately.

Thrombophlebitis has occurred at the site of drug injection or infusion in some patients, and rarely patients have noted pain and inflammation at subcutaneous injection sites. In most instances, however, the drug has been well tolerated.

Patients can tolerate higher total doses when they receive the drug by rapid intravenous injection as compared with slow infusion. This phenomenon is related to the drug's rapid inactivation and brief exposure of susceptible normal and neoplastic cells to significant levels after rapid injection. Normal and neoplastic cells seem to respond to somewhat parallel fashion to these different modes of administration and no clear-cut clinical advantage has been demonstrated for either.

Clinical experience accumulated to date suggests that success with CYTOSAR is dependent more on adeptness in modifying day-to-day dosage to obtain maximum leukemic cell kill with tolerable toxicity than on the basic treatment schedule chosen at the outset of therapy. Toxicity necessitating dosage alteration almost always occurs.

Relatively constant plasma levels can be achieved by continuous intravenous infusion.

In many chemotherapeutic programs, CYTOSAR is used in combination with other cytotoxic drugs. The addition of these cytotoxic drugs has necessitated changes and dose alterations. The dosage schedules for combination therapy outlined below have been reported in the literature (see References).

DOSAGE SCHEDULES

Acute Myelocytic Leukemia - induction remission: adults

CYTOSAR 200 mg/m² daily by continuous infusion for 5 days (120 hours) - total dose 1000 mg/m². This course is repeated approximately every 2 weeks. Modifications must be made based on hematologic response.

Acute myelocytic leukemia - maintenance: adults

Maintenance programs are modifications of induction programs and, in general, use similar schedules of drug therapy as were used during induction. Most programs have a greater time spacing between courses of therapy during remission maintenance.

Acute myelocytic leukemia - induction and maintenance in children

Numerous studies have shown that childhood AML responds better than adult AML given similar regimens. Where the adult dosage is stated in terms of body weight or surface area, the children's dosage may be calculated on the same basis. When specified amounts of a drug are indicated for the adult dosage, these should be adjusted for children on the basis of such factors as age, body weight or body surface area.

Acute myelocytic leukemia – adults and children

The following tables outline the results of treatment with CYTOSAR alone and in combination with other chemotherapeutic agents, in the treatment of acute myelocytic leukemia in adults and children.

The treatment regimens outlined in the tables should not be compared for efficacy. These were independent studies with a number of variables involved, such as patient population, duration of disease, and previous treatment.

The responsiveness and course of childhood acute myelocytic leukemia (AML) appear to be different from that in adults. Numerous studies show response rates to be higher in children than in adults with similar treatment schedules. Experience indicates that at least with induction and initial drug responsiveness, childhood AML appears to be more similar to childhood acute lymphocytic leukemia (ALL) than to its adult variant.

TABLE I
Acute Myelocytic Leukemia - Remission Induction: Adults

| Drug Dosage Schedule* | | No. of Patients Evaluated | Complete Remissions | Investigators |
|--|--|---------------------------|---------------------|---------------------------------------|
| CYTOSAR Single-Dose Therapy | (Infusion) 10 mg/m ² 12 hrs/day | 12 | 2 (17%) | Ellison (1968) |
| | 30 mg/m ² 12 hrs/day | 41 | 10 (24%) | |
| | 10 mg/m ² 24 hrs/day | 9 | 2 (22%) | |
| | 30 mg/m ² 24 hrs/day | 36 | 2 (6%) | |
| | (Infusion) 200 mg/m ² 24 hrs/5 days | 36 | 9 (25%) | Bodey (1969) |
| | 10 mg/m ² i.v. injection initially, then infusions of 30 mg/m ² per 12 hrs or 60 mg/m ² /day for 4 days | 49 | 21 (43%) | Goodell (1970) |
| | (Infusion Therapy) 800 mg/m ² /2 days | 53 | 12 (23%) | Southwest Oncology Group (1974) |
| | 1000 mg/m ² /5 days | 60 | 24 (40%) | |
| 100 mg/m ² /day 1 hr infusion | 49 | 7 (14%) | Carey (1975) | |
| 5-12.5 mg/kg/12 hr infusion following i.v. synchronizing dose** | 5 | 5 (100%) | Lampkin (1976) | |
| Combined Therapy | CYTOSAR – doxorubicin | 41 | 30 (73%) | Preisler (1979) |
| | CYTOSAR - thioguanine daunorubicin | 28 | 22 (79%) | Gale (1977) |
| | CYTOSAR - doxorubicin vincristine – prednisolone | 35 | 23 (66%) | Weinstein (1980) |
| | CYTOSAR - daunorubicin thioguanine - prednisone vincristine | 139 | 84 (60%) | Glucksberg (1981) |
| | CYTOSAR – daunorubicin | 21 | 14 (67%) | Cassileth (1977) |

TABLE I (Cont'd)
Acute Myelocytic Leukemia - Remission Induction: Adults

| Drug Dosage Schedule* | | No. of Patients Evaluated | Complete Remissions | Investigator |
|--------------------------|------------------------|---------------------------|---------------------|-----------------|
| <i>High Dose Therapy</i> | CYTOSAR | 7 | 6 (86%) | Lister (1983) |
| | CYTOSAR | 21 | 12 (57%) | Herzig (1983) |
| | CYTOSAR | 11 | 8 (73%) | Preisler (1983) |
| | CYTOSAR - doxorubicin | 14 | 7 (50%) | Willemze (1982) |
| | CYTOSAR - asparaginase | 13 | 9 (69%) | Capizzi (1983) |

* Unless otherwise stated, all doses given until drug effect-modifications then based on hematologic reasons. See references.

** Highly experimental - requires ability to study mitotic indices.

TABLE II
Acute Myelocytic Leukemia - Remission Induction: Children (21 and under)

| Drug Therapy | No. of Patients Evaluated | Complete Remissions | Investigator |
|---|---------------------------|---------------------|------------------|
| CYTOSAR (5-12.5mg/kg following i.v. synchronizing dose**) | 16 | 12 (75%) | Lampkin (1976) |
| CYTOSAR, vincristine, doxorubicin, prednisolone | 48 | 35 (73%) | Weinstein (1980) |
| CYTOSAR, thioguanine, doxorubicin | 11 | 8 (72%) | Hagbin (1975) |
| CYTOSAR, thioguanine | 47 | 20 (43%) | Pizzo (1976) |
| CYTOSAR, cyclophosphamide | 12 | 7 (58%) | Pizzo (1976) |

** Highly experimental - requires ability to study mitotic indices.

Acute lymphocytic leukemia

In general, dosage schedules are similar to those used in acute myelocytic leukemia with some modification. CYTOSAR has been used in the treatment of acute lymphocytic leukemia in both adults and children. When CYTOSAR was used with other antineoplastic agents as part of a total therapy program, results were equal to or better than reported with such programs which did not include CYTOSAR. Used singly, or in combination with other agents, CYTOSAR has also been effective in treating patients who had relapsed on other therapy. Table III and IV summarize the results obtained in previously treated patients. Since these are independent studies with such variables as patient population, duration of disease and previous treatment, results shown should not be used for comparing the efficacy of the outlined treatment programs.

TABLE III
Acute Lymphocytic Leukemia - Remission Induction
Previously Treatment Patients
Adults and Children

| Drug Therapy | No. of Patients Evaluated | Complete Remissions | Response | Investigator |
|---|---------------------------|---------------------|----------|------------------|
| CYTOSAR 3-5 mg/kg/day (IV injection) | 43 | 2 (5%) | 15 (35%) | Howard (1968) |
| CYTOSAR - asparaginase | 9 | 8 (89%) | 8 (89%) | McElwain (1969) |
| CYTOSAR - cyclophosphamide | 11 | 7 (64%) | 9 (82%) | Bodey 1970 |
| CYTOSAR - prednisone | 83 | - | (49%) | Nesbitt (1970) |
| CYTOSAR 150-200 mg/m ² /5 days (infusion) | 34 | 1 (3%) | 4 (12%) | Wang (1970) |
| CYTOSAR - L - asparaginase - prednisone - vincristine - doxorubicin | 91 | 72 (79%) | - | Klemperer (1978) |
| CYTOSAR - L - asparaginase - prednisone - vincristine - doxorubicin | 55 | 42 (76%) | - | Klemperer (1978) |
| CYTOSAR - asparaginase | 22 | 13 (59%) | 15 (68%) | Ortaga (1972) |
| CYTOSAR - thioguanine | 19 | 9 (47%) | 9 (47%) | Bryan (1974) |

TABLE IV

| Drug Therapy | | No. of Patients Evaluated | Complete Remissions | Investigator |
|--------------------------|------------------------|---------------------------|---------------------|------------------|
| <i>High Dose Therapy</i> | CYTOSAR | 8 | 3 (38%) | Rohatinar (1983) |
| | CYTOSAR - doxorubicin | 3 | 2 (67%) | Willemze (1982) |
| | CYTOSAR - asparaginase | 10 | 3 (30%) | Capizzi (1983) |

Non-Hodgkin's lymphoma in children

CYTOSAR has been used as part of multi-drug program (LSA₂L₂) to treat non-Hodgkin's lymphoma in children. See Appendix A for complete dosage schedule.

High Dose Chemotherapy

Before instituting a program of high dose chemotherapy, the physician should be familiar with the literature, adverse reactions, precautions, contraindications, and warnings applicable to all the drugs involved in the program.

CYTOSAR

CYTOSAR: 2 g/m² infused over 3 hours every 12 hours x 12 doses (Days 1-6).

CYTOSAR

CYTOSAR: 3 g/m² infused over 1 hour every 12 hours x 12 doses (Days 1-6).

CYTOSAR

CYTOSAR: 3 g/m² infused over 75 minutes every 12 hours x 12 doses (Days 1-6).

CYTOSAR - doxorubicin

CYTOSAR: 3 g/m² infused over 2 hours every 12 hours x 12 doses (Days 1-6).

Doxorubicin: 30 mg/m² i.v. on Days 6-7.

CYTOSAR - asparaginase

CYTOSAR: 3 g/m² infused over 3 hours at 0 hours, 12 hours, 24 hours, and 36 hours. At 42 hours, 6000 units/m² of asparaginase i.m. (Days 1-2); repeat same schedule Days 8-9.

Combined Chemotherapy

Before instituting a program of combined chemotherapy, the physician should be familiar with the literature, adverse reactions, precautions, contraindications, and warnings applicable to all the drugs involved in the program.

CYTOSAR, doxorubicin

CYTOSAR: 100 mg/m²/day, continuous i.v. infusion (Days 1-10)

Doxorubicin: 30 mg/m²/day, i.v. infusion of 30 minutes (Days 1-3)

Additional (complete or modified) courses as necessary at 2-4 week intervals if leukemia is persistent.

CYTOSAR, thioguanine, daunorubicin

CYTOSAR: 100 mg/m², i.v. infusion over 30 minutes every 12 hours (Days 1-7).

Thioguanine: 100 mg/m², orally every 12 hours (Days 1-7).

Daunorubicin: 60 mg/m²/day, i.v. infusion (Days 5-7).

Additional (complete or modified) courses as necessary at 2-4 week intervals if leukemia is persistent.

CYTOSAR, doxorubicin, vincristine, prednisone

CYTOSAR: 100 mg/m²/day, continuous i.v. infusion (Days 1-7).

Doxorubicin: 30 mg/m²/day, i.v. infusion (Days 1-3).

Vincristine: 1.5 mg/m²/day, i.v. infusion (Days 1, 5)

Prednisolone: 40 mg/m²/day, i.v. infusion every 12 hours (Days 1-5).

Additional (complete or modified) courses as necessary at 2-4 week intervals if leukemia is persistent.

CYTOSAR, daunorubicin, thioguanine, prednisone, vincristine

CYTOSAR: 100 mg/m²/day, i.v. infusion (Days 1-10).

Daunorubicin: 70 mg/m²/day, i.v. infusion (Days 1-3).

Thioguanine: 100 mg/m², orally every 12 hours (Days 1-7).

Prednisone: 40 mg/m²/day, orally (Days 1-7).

Vincristine: 1 mg/m²/day, i.v. infusion (Days 1, 7)

Additional (complete or modified) courses as necessary at 2-4 week intervals if leukemia is persistent.

CYTOSAR, daunorubicin

CYTOSAR: 100 mg/m²/day, continuous i.v. infusion (Days 1-7).

Daunorubicin: 45 mg/m²/day, i.v. push (Days 1-3).

Additional (complete or modified) courses as necessary at 2-4 week intervals if leukemia is persistent.

Meningeal Leukemia - Intrathecal Use

CYTOSAR has been used intrathecally in acute leukemia in doses ranging from 5 mg/m² to 75 mg/m² of body surface area. The frequency of administration varied from once a day for 4 days to once every 4 days. The most frequently used dose was 30 mg/m² every 4 days until cerebrospinal fluid findings were normal, followed by one additional treatment. The dosage schedule is usually governed by the type and severity of central nervous system manifestations and the response to previous therapy.

CYTOSAR has been used intrathecally with SOLU-CORTEF Sterile Powder and methotrexate, both as prophylaxis in newly diagnosed children with acute lymphocytic leukemia, as well as in the treatment of meningeal leukemia. Sullivan has reported that prophylactic triple therapy has prevented late CNS disease and given overall cure and survival rates similar to those seen in patients in whom CNS radiation and intrathecal methotrexate was used as initial CNS prophylaxis. The dose of CYTOSAR was 30 mg/m², Solu-Cortef 15 mg/m², and methotrexate 15 mg/m². The physician should be familiar with this report before initiation of the regimen.

Prophylactic triple therapy following the successful treatment of the acute meningeal episode may be useful. The physician should familiarize himself with the current literature before instituting such a program.

CYTOSAR given intrathecally may cause systemic toxicity and careful monitoring of the hemopoietic system is indicated. Modification of the anti-leukemia therapy may be necessary. Major toxicity is rare. The most frequently reported reactions after intrathecal administration were nausea, vomiting and fever; these reactions are mild and self-limiting. Paraplegia has been reported. Necrotizing leukoencephalopathy occurred in 5 children; these patients had also been treated with intrathecal methotrexate and hydrocortisone, as well as by central nervous system radiation. Isolated neurotoxicity has been reported.

Blindness occurred in two patients in remission whose treatment had consisted of combination systemic chemotherapy, prophylactic central nervous system radiation and intrathecal CYTOSAR.

Focal leukemic involvement of the central nervous system may not respond to intrathecal CYTOSAR and may better be treated with radiotherapy.

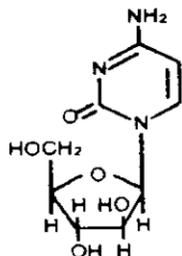
If used intrathecally, do not use a diluent containing benzyl alcohol. Reconstitute with preservative free saline and use immediately.

Dosage modification

The dosage of CYTOSAR must be modified or suspended when signs of serious hematologic depression appear. In general, consider discontinuing the drug if the patient has less than 50,000 platelets or 1000 polymorphonuclear granulocytes/mm³ in his peripheral blood. These guidelines may be modified depending on signs of toxicity in other systems and on the rapidity of fall in formed blood elements. Restart the drug when there are signs of marrow recovery and the above platelet and granulocyte levels have been attained. Withholding therapy until the patient's blood values are normal may result in escape of the patient's disease from control by the drug.

PHARMACEUTICAL INFORMATION

Trade name: CYTOSAR
Drug Substance:
Proper name: cytarabine USP
Chemical name: 4-amino-1-β-D-arabinofuranosyl-2(1H)-pyrimidinone
Structural formula:



Molecular formula: C₉H₁₃N₃O₅

Molecular weight: 243.22

Description:

- odourless, white to off-white crystalline powder
- melting point of 212° to 213°C
- pKa of 4.2
- partition coefficient (octanol-water) of 0.0071
- a synthetic nucleoside which differs from the normal nucleosides cytidine and deoxycytidine in that the sugar moiety is arabinose rather than ribose or deoxyribose
- pH of 7 for 10 mg/mL solution

Solubility:

- freely soluble in water
- slightly soluble in alcohol and chloroform

Composition

Each vial contains the labelled amount of cytarabine USP. Hydrochloric acid solution and/or sodium hydroxide solution is added to adjust the pH.

CYTOSAR Solution for Injection:

Solution of 20 mg/mL: Each mL contains 20 mg cytarabine, with sodium chloride, hydrochloride acid and/or sodium hydroxide to adjust pH and Water for Injection.

Solution of 100 mg/mL: Each mL contains 100 mg cytarabine, with hydrochloride acid and/or sodium hydroxide to adjust pH and Water for Injection.

Stability and Storage Recommendations

Sterile Lyophilized Powder: store at controlled room temperature (15°-30°C)

Solution for Injection: store at 15-25°C; protect from light

Characteristics of Reconstituted Solution from Lyophilized Powder:

- pH of reconstituted solution is approximately 5
- solutions reconstituted without a preservative should be used immediately
- solutions reconstituted with Bacteriostatic Water for Injection with Benzyl Alcohol 0.9% (for multi-dose use) may be stored at controlled room temperature (15°-30°C) for 48 hours
- discard any solution in which a slight haze develops

Reconstitution of Lyophilized Powder

CYTOSAR may be reconstituted with the diluents mentioned below and mixed with the compatible drugs mentioned in the **CHEMICAL STABILITY AND COMPATIBILITY** section. Compatibility must be assured before mixing with any other substance.

CYTOSAR may be reconstituted with the following diluents:

0.9% Sodium Chloride for Injection

Dextrose 5% in Water

Sterile Water for Injection

Bacteriostatic Water for Injection

When reconstituted with a diluent, the following concentrations result:

| Vial Size | Volume of Diluent to be added to Vial | Nominal Concentration |
|------------------|--|------------------------------|
| 100 mg | 5 mL | 20 mg/mL |
| 500 mg | 10 mL | 50 mg/mL |
| 1g | 10 mL | 100 mg/mL |
| 2g | 20 mL | 100 mg/mL |

Solutions reconstituted without a preservative should be used immediately. Solutions reconstituted with Bacteriostatic Water for Injection with Benzyl Alcohol 0.9% may be stored at controlled room temperature (15°-30°C) for 48 hours.

Handling of Solution for Injection

Single use only. Discard any unused portion. If a precipitate has formed as a result of exposure to low temperatures, redissolve by warming to 55°C for no longer than 30 minutes and then shake until the precipitate has dissolved. Allow to cool prior to use.

FOR INTRATHECAL USE: DO NOT USE DILUENT CONTAINING BENZYL ALCOHOL. RECONSTITUTE WITH PRESERVATIVE-FREE 0.9% SODIUM CHLORIDE FOR INJECTION. USE IMMEDIATELY.

FOR HIGH DOSE USE: DO NOT USE DILUENT CONTAINING BENZYL ALCOHOL.

CHEMICAL STABILITY AND COMPATIBILITY

Stability in Infusion Solutions

Lyophilized Powder: Chemical and physical stability studies of CYTOSAR have demonstrated that cytarabine is stable for seven days at room temperature when admixed at 0.5 mg/mL in glass i.v. bottles and plastic i.v. bags with: water for injection; 5% Dextrose injection; and 0.9% Sodium Chloride injection solutions. Also when similarly admixed at 8-32 mg/mL in glass i.v. bottles and plastic i.v. bags, cytarabine is stable for seven days at room temperature, -20°C, and 4°C in 5% Dextrose Injection; 5% Dextrose in 0.2% Sodium Chloride Injection; and, in 0.9% Sodium Chloride Injection Solutions.

Cytarabine is stable at room temperature at a concentration of 2 mg/mL in the presence of KCl equivalent to 50 meq/500 ml in Dextrose 5% in water and 0.9% Sodium Chloride for up to eight days.

CYTOSAR is compatible for 24 hours at 5°C with lactated Ringers, dextrose 5% in water, 0.9% sodium chloride, dextrose 5% in water in 0.9% sodium chloride.

Solution for Injection:

Dilutions of cytarabine should be made in Glucose 5% or Sodium Chloride 0.9% or Water for Injection intravenous infusions to concentrations as low as 0.1mg/mL (or up to the highest concentration required to provide the highest dose in the Dosage and Administration section) and are physically and chemically stable for up to 4 days at both 25°C/60% relative humidity, exposed to normal room lighting, and at 2-8°C, protected from light. However, as Cytarabine Solution for Injection and the infusion solutions prepared therefrom contain no antimicrobial agents, it is recommended that diluted solutions of Cytarabine Injection be used within 24 hours when stored at room temperature.

Drug Compatibilities

CYTOSAR 0.8 mg/mL and sodium cephalothin 1.0 mg/mL are chemically stable for 8 hours in dextrose 5% in water.

CYTOSAR 0.4 mg/mL and prednisolone sodium phosphate 0.2 mg/mL are compatible in dextrose 5% in water for 8 hours.

CYTOSAR 16 mcg/mL and vincristine sulfate 4 mcg/mL are compatible in dextrose 5% in water for 8 hours.

Drug incompatibilities

CYTOSAR has been known to be physically incompatible with heparin, insulin, 5-fluorouracil, penicillin G, and methylprednisolone sodium succinate.

AS WITH ALL INTRAVENOUS ADMIXTURES, DILUTION SHOULD BE MADE JUST PRIOR TO ADMINISTRATION AND THE RESULTING UNPRESERVED SOLUTION USED WITHIN 24 HOURS.

AVAILABILITY OF DOSAGE FORMS

CYTOSAR (cytarabine) is supplied as:

- Lyophilized sterile powder in vials containing 100 mg, 500mg, 1g and 2g of the drug.
- Solution for Injection (20 mg/mL) in polypropylene “cytosafe” vials of 100 mg/5 mL and 500 mg/25 mL
- Solution for Injection (100 mg/mL) in polypropylene “cytosafe” vials of 1000 mg/10 mL and 2000 mg/20 mL

PHARMACOLOGY

Cell Culture Studies

Cytarabine is cytotoxic to a wide variety of proliferating mammalian cells in culture. It exhibits cell phase specificity, primarily killing cells undergoing DNA synthesis (S-phase) and under certain conditions blocking the progression of cells from the G₁ phase to S-phase. Although the mechanism of action is not completely understood, it appears that cytarabine acts through the inhibition of DNA polymerase. A limited, but significant, incorporation of cytarabine into both DNA and RNA has also been reported. Extensive chromosomal damage, including chromatoid breaks has been produced by cytarabine and malignant transformation of rodent cells in culture has been reported. Deoxycytidine prevents or delays (but does not reverse) the cytotoxic activity.

Animal Studies

In experimental studies with mouse tumors, cytarabine was most effective in those tumors with a high growth fraction. The effect was dependent on the treatment schedule; optimal effects were achieved when the schedule (multiple closely spaced doses or constant infusion) ensured contact of the drug with the tumor cells when the maximum number of cells was in the susceptible S-phase. The best results were obtained when courses of therapy were separated by intervals sufficient to permit adequate host recovery.

Human Pharmacology

Cytarabine is capable of obliterating immune responses in man during administration. Suppression of antibody responses to E-coli-VI antigen and tetanus toxoid have been demonstrated. This suppression was obtained during both primary and secondary antibody responses.

Cytarabine also suppressed the development of cell-mediated immune responses such as delayed hypersensitivity skin reaction to dinitrochlorobenzene. However, it has no effect on already established delayed hypersensitivity reactions.

Following 5-day courses of intensive therapy with Cytarabine the immune response was suppressed, as indicated by the following parameters: macrophage ingress into skin windows; circulating antibody response following primary antigenic stimulation; lymphocyte blastogenesis with phytohemagglutinin. A few days after termination of therapy there was a rapid return to normal.

TOXICOLOGY

Animal Studies

Toxicity of cytarabine in experimental animals, as well as activity, is markedly influenced by the schedule of administration. For example, in mice, the LD₁₀ for single intraperitoneal administration is greater than 6000 mg/m². However, when administered in 8 doses, each separated by 3 hours, the LD₁₀ is less than 750 mg/m² total dose. Similarly, although a total dose of 1920 mg/m² administered as 12 injections at 6-hour intervals was lethal to beagle dogs (severe bone marrow hypoplasia with evidence of liver and kidney damage), dogs receiving the same total dose administered as 8 injections (again at 6-hour intervals) over a 48-hour period survived with minimal signs of toxicity.

The most consistent observation in surviving dogs was elevated transaminase levels. In all experimental species the primary limiting toxic effect is marrow suppression with leukopenia. In addition, cytarabine causes abnormal cerebellar development in the neonatal hamster and is teratogenic to the rat fetus.

The major dose-limiting toxicity of cytarabine observed in all tested species is myelosuppression, manifested by megaloblastosis, reticulocytopenia, leukopenia, and thrombocytopenia. Other target organs include liver, kidney, and brain. Extensive chromosomal damage, including chromatoid breaks have been produced by cytarabine and malignant transformation of rodent cells in culture has been reported. Cytarabine is embryotoxic and teratogenic and produced peri- and postnatal toxicity in various species. No formal fertility studies have been reported however sperm head abnormalities were observed following cytarabine treatment in mice.

APPENDIX A
LSA₂-L₂ Protocol

Woolner N, Burchenal JH, Lieberman PH, et al: Non-Hodgkin's Lymphoma in Children - A Comparative Study of Two Modalities of Therapy. *Cancer* 1976;37:123-134.

Induction Phase

Day 1. Cyclophosphamide 1,200 mg/m² single push injection.

Day 3 to 31. Prednisone 60 mg/m² po divided into three daily doses.

Day 3, 10, 17, 24. Vincristine 1.5 to 2.25 mg/m² intravenously.

Day 5, 27, 30. Spinal tap and intrathecal injection of Methotrexate 6.25 mg/m²

Day 12, 13. Daunomycin 60 mg/m² intravenously.

At the end of induction (last dose of intrathecal methotrexate) patient rests for 3-5 days before consolidation.

Consolidation Phase

Day 34 or 36, daily intravenous injections of cytosine arabinoside (Ara-C) 150 mg/m^2 for a total of 15 injections are given. (Injections are given from Monday through Friday.) Thioguanine 75 mg/m^2 is given orally, 8-12 hours after the injection of Ara-C. If the white blood count is 1500 or more and the platelet count 150,000 or more on the 5th day of Ara-C, the patient continues to receive the same dosage of thioguanine over the weekend. However, both are discontinued temporarily when there is evidence of marrow depression; this usually occurs after the initial seventh to tenth doses of the combination and ordinarily recovers within 7-10 days. Hence, the patients may receive more than 15 doses of thioguanine orally, but receive only 15 doses of i.v. cytosine arabinoside (Ara-C). This first phase of the consolidation takes an average of 30-35 days. The second phase of the consolidation should be started immediately after completion of the 15 doses of Ara-C; it entails daily i.v. administration of L-asparaginase, 60000 U/m^2 for a total of 12 injections, excluding weekends.

Two days after the last injection of the L-asparaginase, two more intrathecal (i.t.) injections of methotrexate are given 2 days apart. Three days after the last i.t. methotrexate, BCNU [1, 3-Bis (2 chloroethyl 1-1-nitrosourea)] 60 mg/m^2 is given i.v., which completes the consolidation. The average duration of the induction and consolidation is 85-100 days.

Maintenance Phase

The maintenance period consists of five cycles of 5 days each and is started 3-4 days after completion of consolidation.

Cycle I: Oral thioguanine 300 mg/m² for 4 consecutive days: i.v. cyclophosphamide 600 mg/m² on the 5th day.

Rest 7-10 days.

Cycle II: Oral hydroxyurea 2,400 mg/m² for 4 consecutive days: i.v. daunomycin 45 mg/m² on the 5th day.

Rest 7-10 days.

Cycle III: Oral methotrexate 10 mg/m² for 4 consecutive days: i.v. BCNU 60 mg/m² on the 5th day.

Rest 7-10 days.

Cycle IV: I.V. Ara-C 150 mg/m² for 4 consecutive days: i.v. vincristine 1.5 mg/m² on day 5.

Rest 7-10 days.

Cycle V: Two doses of i.t. methotrexate 6.25 mg/m² 2-3 days apart.

Rest 7-10 days and restart with Cycle I.

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PART III: CONSUMER INFORMATION

Pr **CYTOSAR**[®]
(cytarabine for injection)

This leaflet is part III of a three-part "Product Monograph" published when CYTOSAR was approved for sale in Canada and is designed specifically for Consumers. This leaflet is a summary and will not tell you everything about CYTOSAR. Contact a member of your healthcare team if you have any questions about the drug.

ABOUT THIS MEDICATION

What the medication is used for:

CYTOSAR is used alone or in combination with other anti-cancer medicines in the treatment of patients with certain types of leukaemia (cancer of the blood) and lymphomas (cancer of the lymph glands).

What it does:

Cytarabine is a cytotoxic drug that interferes with cell growth and causes cell death.

When it should not be used:

Do not take CYTOSAR:

- If you are allergic (hypersensitive) to cytarabine or any of the other ingredients in CYTOSAR

CYTOSAR should not be given to premature infants when using a diluent that contains benzyl alcohol.

What the medicinal ingredient is:

Cytarabine.

What the nonmedicinal ingredients are:

Hydrochloric acid solution and/or sodium hydroxide solution to adjust the pH.

What dosage forms it comes in:

CYTOSAR is supplied as:

- Freeze-dried sterile powder in vials containing 100 mg, 500mg, 1g and 2g of the drug
- Solution for Injection (20 mg/mL) in polypropylene "cytosafe" vials of 100 mg/5 mL and 500 mg/25 mL
- Solution for Injection (100 mg/mL) in polypropylene "cytosafe" vials of 1000 mg/10 mL and 2000 mg/20 mL

WARNINGS AND PRECAUTIONS

Serious Warnings and Precautions

CYTOSAR should be used only by doctors with experience in cancer medicines.

Serious side effects with CYTOSAR include:

- **Decreased production of blood cells (myelosuppression)**
- **Heart muscle disorders (cardiomyopathy).**
- **Anaphylactic reactions**
- **Tumour Lysis Syndrome (TLS)**
- **Secondary cancers (other cancers)**

In patients on a high dose schedule of CYTOSAR, severe gastrointestinal toxicity, central nervous system toxicity, pulmonary toxicity, at times fatal, and eye toxicity have been reported. Vaccination with a live vaccine should be avoided while taking CYTOSAR.

CYTOSAR may cause Tumour Lysis Syndrome [TLS]. TLS is a metabolic condition that results from dying cancer cells and involves changes in blood chemistry that can lead to kidney failure and abnormal heart rhythm, which may be fatal. Tell your doctor immediately if you have palpitations/irregular heartbeats; vomiting; fatigue/weakness; difficulty concentrating/trouble thinking; swelling, numbness or tingling in hands, face or feet; back pain; muscle cramps; fainting or trouble breathing.

Cases of acute pancreatitis, and cases of paralysis, at times fatal in children have been reported with the use of CYTOSAR in combination with other drugs.

Severe nervous system adverse reactions that ranged from headache to paralysis, coma and stroke-like episodes have been reported mostly in children (under 18 years of age) given intravenous (injected into the vein) cytarabine in combination with intrathecal (injected into the spinal cord) methotrexate.

Tell your doctor before taking CYTOSAR, if any of the following apply to you:

- Liver or kidney problems;
- Heart problems;
- Lung problems;
- Low blood cell counts;
- Skin problems;
- Pregnant or think you may be pregnant;
- Breast-feeding;
- Are male patient and plan to father a child.

The safety of CYTOSAR in infants (under 1 year of age) is not known.

Contraception

CYTOSAR may cause harm to an unborn child. Female patients who might get pregnant must use effective contraception during treatment with CYTOSAR. Since CYTOSAR may present in the semen, male patients who are not surgically sterile must agree to use effective contraception during treatment with CYTOSAR to prevent pregnancy in female partners. If pregnancy is suspected during treatment with CYTOSAR, inform your doctor immediately.

Driving and using machines:

If you experience (feel) dizziness, do not drive or use machinery.

INTERACTIONS WITH THIS MEDICATION

Tell your doctor or pharmacist if you are taking or have recently taken any other medicines, even those not prescribed, especially the following:

- 5-Fluorocytosine (a medicine used to treat fungal infections).
- Digoxin.
- Gentamicin (an antibiotic).
- Cyclophosphamide, vincristine and prednisone.
- Methotrexate (a medicine used to treat cancer).

PROPER USE OF THIS MEDICATION

Cytarabine will be given to you as an injection or an infusion into a vein (through a ‘drip’).

The dose of Cytarabine will be decided by your doctor based on your condition being treated for and your body surface area (your body weight and height will be used to calculate your body surface area).

Overdose:

In case of drug overdose, contact a healthcare practitioner, hospital emergency department or regional poison control centre, even if there are no symptoms.

SIDE EFFECTS AND WHAT TO DO ABOUT THEM

Like all medicines, CYTOSAR can have side effects.

Tell your doctor or nursing staff immediately, if you experience the following side effects:

- An allergic reaction such as sudden wheeziness, difficulty in breathing, swelling of eyelids, face or lips, rash or itching (especially affecting the whole body), hives.
- Feeling tired.
- Flu like symptoms e.g. raised temperature or fever and chills.
- Bruise more easily or bleed more than usual if you hurt yourself.

Other side effects include:

- Reactions at Injection site: inflammation to your veins (caused by a blood clot) and infection.
- Headaches or feeling dizzy, feeling of pins and needles, shaking and fits, drowsiness, experience problems in walking, speech problems, involuntary muscular movement, changes in your personality, tiredness, weakness, fainting.
- Hair loss, a skin rash or ulceration, peeling of the skin, itching or increased freckles.
- Infections
- Feeling sick, being sick, diarrhoea, loss of appetite,

abdominal pain. Abdominal swelling and bloody stool.

- Inflammation of the gullet, heartburn, sores and bleeding in the mouth, lips, or on the anus (back passage).
- Pancreatitis (pain in the upper abdomen) often accompanied by feeling sick or vomiting.
- Liver damage (seen as yellowing of the skin and whites of the eye).
- Difficulty or pain when passing urine. Blood in your urine and impaired kidney function.
- Feeling hot and feverish, conjunctivitis, and pain and numbness in joints, fingers, toes or face, swelling of the abdomen, legs, ankles and feet.
- May cause paralysis
- Shortness of breath, pneumonia, short or stabbing chest pain, build up of fluid in the lungs, sore throat.
- Muscle pain, bone pain.
- Fast heart beat
- Eye infection, irritation, pain and blurred vision. Visual loss. Intolerance to light.
- Cytarabine Syndrome can happen between 6 to 12 hours after receiving Cytarabine. The syndrome includes feeling generally unwell with a high temperature, pain in bone, muscle and sometimes the chest, blistery rash, sore eyes.
- Rash or blisters on the palms of the hands and soles of the feet

If any of the side effects get serious or if you notice any side effect not listed in this leaflet, please tell your doctor or nursing staff immediately.

SERIOUS SIDE EFFECTS, HOW OFTEN THEY HAPPEN AND WHAT TO DO ABOUT THEM

| Symptom / effect | Talk with your doctor or pharmacist | | Stop taking drug and call your doctor or pharmacist |
|--|-------------------------------------|--------------|---|
| | Only if severe | In all cases | |
| Pancreatitis (inflammation of the pancreas) with symptoms such as abdominal pain, fever, nausea, vomiting | | √ | |
| Decreased white blood cell and platelet counts with symptoms such as infection, fever, bleeding, bruising and rash | | √ | |

This is not a complete list of side effects. For any unexpected effects while taking CYTOSAR, contact your doctor or pharmacist.

HOW TO STORE IT

Keep out of reach and sight of children.

Sterile Freeze-dried Powder:

- Store at controlled room temperature (15-30°C).
- Solutions reconstituted without a preservative should be used immediately. Solutions reconstituted with Bacteriostatic Water for Injection with Benzyl Alcohol 0.9% may be stored at controlled room temperature (15-30°C) for 48 hours.

Sterile Solution for Injection:

- Store at 15-25°C; protect from light.
- Diluted solutions should be used within 24 hours when stored at room temperature (controlled temperature: not more than 25° C).

Medicines should not be disposed of via wastewater or household waste. Ask your pharmacist how to dispose of medicines no longer required. These measures will help to protect the environment.

REPORTING SUSPECTED SIDE EFFECTS

You can report any suspected adverse reactions associated with the use of health products to the Canada Vigilance Program by one of the following 3 ways:

Report online at www.healthcanada.gc.ca/medeffect

Call toll-free at 1-866-234-2345

Complete a Canada Vigilance Reporting Form and:

- Fax toll-free to 1-866-678-6789, or
- Mail to: Canada Vigilance Program
Health Canada
Postal Locator 0701C
Ottawa, ON K1A 0K9

Postage paid labels, Canada Vigilance Reporting Form and the adverse reaction reporting guidelines are available on the MedEffect™ Canada Web site at www.healthcanada.gc.ca/medeffect.

NOTE: Should you require information related to the management of side effects, contact your health professional. The Canada Vigilance Program does not provide medical advice.

MORE INFORMATION

This document plus the full Product Monograph, prepared for health professionals can be found at:
<http://www.Pfizer.ca> or by contacting Pfizer Canada Inc. Medical Information at 1-800-463-6001.

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