ſ

INTERNET ARCHIVE	DEC FEB
9 captures           6 Oct 03 - 20 Oct 08	2003 2005 2
<b>FDA</b> U.S. Food and Drug Administration of Health and Human Services	
CENTER FOR DRUG EVALUATION AND RESEARCH	i
FDA Home Page   CDER Home Page   CDER Site Info   Contact CDER   What's New @ CDER	_
CDER Home         About CDER         Drug Information         Regulatory Guidance         CDER Calendar         Specific Audiences         CDER Archives	-
Search GO powered Google"	
Methotrexate Preservative Free	
Company: Bigmar Application No.: 040266	
Approval Date: 02/26/1999	
• <u>Approval Letter(s)</u>	
• Printed Labeling	
• <u>Chemistry Review(s)</u>	
<ul> <li><u>Microbiology Review(s)</u></li> </ul>	
<ul> <li><u>Bioequivalence Review(s)</u></li> </ul>	
<ul> <li><u>Administrative Document(s)</u></li> </ul>	
• Correspondence	

Back to Drug Approval Page

Some documents are in **P**ortable **D**ocument **F**ormat (PDF) to retain the original format. To view or print these documents, you must use the Adobe Acrobat viewer. Acrobat is free and available directly from Adobe's website with full installation instructions.

Date created: July 31, 2003

CDER Home Page | CDER Site Info | Contact CDER | What's New @ CDER FDA Home Page | Search FDA Site | FDA A-Z Index | Contact FDA | HHS Home Page

FDA/Center for Drug Evaluation and Research



Find authenticated court documents without watermarks at <u>docketalarm.com</u>.

## CENTER FOR DRUG EVALUATION AND RESEARCH

# APPLICATION NUMBER: 40266

## **DRAFT FINAL PRINTED LABELING**

**DOCKET A L A R M** Find authenticated court documents without watermarks at <u>docketalarm.com</u>.



## METHOTREXATE FOR INJECTION USP

### WARNINGS

METHOTREXASE SHOULD BE USED ONLY BY PHYSI-CIANS WHOSE KNOMLEDGE AND EXPERIENCE INCLUDE THE USE OF ANTIMETABOLITE THERAPY.

BECAUSE OF THE POSSIBILITY OF SERIOUS TOXIC RE-ACTIONS (WHICH CAN BE FATAL):

METHOTREXATE SHOULD BE USED ONLY IN LIFE THREATENING "EOPLASTIC DISEASES, OR IN PATIENTS WITH PSORIASIS WITH SEVERE RECALCITRANT, DIS-ABLING DISEASE WHICH IS NOT ADEQUATELY RESPONSIVE TO OTHER FORMS OF THERAPY.

DEATHS HAVE BEEN REPORTED WITH THE USE OF METHOTREXATE IN THE TREATMENT OF MALIGNANCY AND PSORIASIS

PATIENTS SHOULD BE CLOSELY MONITORED FOR BONE MARROW, LIVER, LUNG AND KIDNEY TOXICITIES. (See PRECAUTIONS.)

PATIENTS SHOULD BE INFORMED BY THEIR PHYSICIAN OF THE RISKS INVOLVED AND BE UNDER A PHYSICIAN'S CARE THROUGHOUT THERAPY.

THE USE OF METHOTREXATE HIGH DOSE REGIMENS REC-COMMENDED FOR OSTEOSARCOMA RECURRES ME-TCULOUS CARE, USAN DOSAGE AND ADMINISTRATION I HIGH DOSE REGIMENS FOR OTHER NEOPLASTIC DIS-EASES ARE INVESTIGATIONAL AND A THERAPEUTIC ADVANTAGE HAS NOT BEEN ESTABLISHED.

METHOTREXATE FORMULATIONS AND DILUENTS CONTAINING PRESERVATIVES MUST NOT BE USED FOR INTRATHECAL OR HIGH DOSE METHOTREX-ATE THERAPY.

Methotrexate has been reported to cause fetal death and/or origental anomales. Therefore, it is not recommended for normen of childbearing potentia unless there is clear medical widence that the benefits can be expected to outweigh the providend disk. Pregnant women with postnasis should not sceive methotrexate. (See CONTRAINDICATIONS).

Methotrexate elimination is reduced in patients with im-paired renal function, ascites, or pleural effusions. Such patients require especiality careful monitoring for toxicity, and require dose reduction or, in some cases, discontinua-tion of methotrexate administration.

Unexpectedly severe (sometimes fatal) bone marrow sup-pression and gastrointestinal buictly have been reported with concomilant administration on methorescate (susually in high dosage) along with some nonsteroidal anti-inflammatory drugs (NSADS). (See PRECAUTIONS). Ding threactions).

4. Methotisexate causes hepatioxicity, through senatoxicity, through generality only after prolonged use. Acutely, fiver enzyme elevations are inclusion, and also do not appear predictive of subsequent hepatic disease. Liver biopy after sustained use often shows histologic changes, and librosis and cirrhosis have been reported; these latter lesions may not be praceded by symptoms or abnormal liver function less in the psortasis projulation. For this reason, periodic liver biops after long-term long-are the psortasis are usually recommended for psonatic parteents who are under long-term treatment (Se PRECAUTIONS, Organ System Toxicity, Hepatic.)

5. Methotrexate-induced lung disease is a potentially danger-ous lesion, which may occur acutely at any time during therapy and which has been reported at obses as low as 7.5 mg/week. It is not always fully reversible. Pulmonary symptoms (espe-cially a dir), nonproductive ocugi) may require interruption of treatment and ~ thui mvestigation.

Diarrhea and ulcerative stomatitis require interruption of therapy; otherwise, hernorrhagic enterlits and death from in-

÷ 1 1.00

testinal perforation may occur

7. Malignant lymphomas, which may regress following with-drawal of methotrexate, may occur in patients receiving low-doe methotrexate and, thus may not require cytotoxic treatment. Discontinue methotexate first and, if the lymphoma does not regress, appropriate treatment should be instituted.

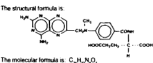
Like other cytotoxic drugs, methotrexate may induce "tu-mor lysis syndrome" in patients with rapidly growing tumors. Appropriate supportive and pharmacologic measures may prevent or alleviate this complication.

Severe, occasionally fatal, skin reactions have been reported following single or multiple doses of methotrexate. Reactions have occurred within days of oral, intramuscular, intravenous, or intrathecal methotrexate administration. Re-covery has been reported with discontinuation of therapy. (See PRECAUTIONS, Organ System Toxicity, skin.)

10. Potentially latal opportunistic infections, especially Pneumocystis carinii pneumonia may occur with methotrex ate therapy. DESCRIPTION

otrevate (formerly Amethopterin) is an antimetabolite used in eatment of certain neoplastic diseases and severe psoriasis.

Chemically methotrexate is N-[4-[[(2,4-diamino-6-pteridinyl) methyl]methylamino]benzoyl]-L-glutamic acid.



The molecular weight is: 454.45

Methotrexate for Injection is sterile and non-pyrogenic and may be given by the intramuscular, intravenous, intra-arterial, or intrathe-cal route. (See DOSAGE AND ADMIN(STRATION).

Each vial contains methotrexate sodium equivalent to 1 g methotr-exate. Contains no preservative. Sodium hydroxide and/or hydrochionc acid may be added to adjust the pH during manufac-ture to 8.5-8.7. The 1 g vial contains approximately 7mEq of sodium.

### CLINICAL PHARMACOLOGY

CLINICAL PHARMACCLOCY Methotexate inhibits dilyndroloa acid reductase. Dihydrololales must be induced to tetrahydrololales by this enzyme before they can be utilized as carriers of one-carbon groups in the synthesis of purine nucleototes and thymidylate. Therefore, methotexate inte-ners with DNA synthesis, repair and cellular replication. Actively proliferating tissues such as malignant cells, bone marrow, tetal cells, buccal and intestinal muccas, and cells of the urinary blad-der rare in gen-eral more sensitive to the effect of methotexate. When cellular proliferation in malignant tassues is greater than in most from all issues. Such or thore are used in pair inpair malignant growth without intervensible damage to normal lissues.

In psoriasis, the rate of production of epithelial cells in the skin is greatly increased over normal skin. This differential in profilera-tion rates is the basis for the use of methotrexate to control the psoriatic process.

Methotrerate in high doses, followed by leucovorin rescue, is used as a part of the treatment of patients with non-metastatic coleosa-rooma. The original retonale for high dose methotexate therapy was based on the concept of selective rescue of normal tissues by leucovorin. More recent evidence suggests that high dose meth-otexate may also overcome methotexate resistance caused by impaired active transport, decreased aftirity of dihydrofloic acid re-ductase for methotexate, increased levels of dihydrofloic acid re-ductase for methotexate, increased levels of dihydrofloic acid re-ductase for methotexate, increased levels of dihydrofloic acid or polygitamation of methotrexate. The actual mechanism of action is unknown. polygiutar

Two Pediatric Oncology Group studies (one randomized and one non-randomized) demonstrated a significant improvement in relapse-free survixa in patients with non-metastatic osteosa-rcoma, when high dose methotrexate with leucovorin rescue was used in combination with other chemotherapeutic agents following surgical resection of the primary tumor. These studies were not designed to demonstrate the specific contribution of high dose methotrexate/leucovorin rescue therapy to the efficacy of the combination. However, a contribution can be inferred from the reports of objective responses to this therapy

888) **9** Z in patients with metastatic osteosarcoma, and from reports of extensive tumor necrosis following preoperative administration of this therapy to patients with non-metastatic osteosarcoma.

Pharmacokinetics Absorption-Methotravate is generally completely absorbed from perenteral routes of injection. After inframusodia: injection, peak serum concentrations occur in 30 to 60 minutes.

Distribution-After intravenous and on the neuronal operation of the initial volume of distribution is approximately 0.16 Lkg (18% of body weight) and steady-state volume of distribution is approximately 0.4 to 0.8 Lkg (40% to 80% of body weight). Methotnexite competes with notwork balass for active transport across call membranes by means of a single carrier-mediated active transport process. At serum concen-tations greater than 100 micromolar, passive diflusion becomes a major pathway by which effective intraoetildar concentrations be achieved. Methotnexies in serum is approximately 50% protein bound. Laboratory studies demonstrate that it may be displaced from plasma adumin by vanious compounds including sufficientides, salicylates, tetracyclines, chloramphenicol, and phenytoin.

Methotrexate does not penetrate the blood-cerebrospinal fluid barrier in therapeutic amounts when given parenter-ally. High CSF concentrations of the drug may be attained by intrathecal administration.

By intraumedia administration. Metabolism - After absorption, metholtrexate undergoes hepätic and intracelidar metabolism to polyglutamated forms which can be converted back to metholtrexate by hydrolase enzymes. These polyglutamates act as inhibitors of dhydrolotate reduc-lase and thymidylate synthetase. Small amounts of these active metabolites vary among different cells, bissues and tumors. A small amount of metabolism to 7-bydroxymethorexate may occur at doess com-monly prescribed. Accumulation of this metabolite may become significant at the high doese used in osteogenic s arcoma. The aqueous solubility of 7-hydroxymethotrexate is 3 to 5 fold lower than the parent form after oral administration.

Half-Life - The terminal half-tile reported for methotrexate is ap proximately three to ten hours for patients receiving treatment los portiais or tow does entinecoptasic therapy (less than 30 mg/m<sup>2</sup>) For patients receiving high doses of methotrexate, the terminal half-tile is eight to 15 hours.

Excretion - Renal excretion is the primary route of elimination, and is dependent upon desage and route of administration. With N and ministration. 80% to 90% or the administered dose is excreted unchanged in the urine within .24 hours. There is limited bilary excretion amounting to 10% or lass of the administered dose. Enteroihepaic recirculation of methotrexate has been proposed.

Renal excretion occurs by giomenular fittration and active tubular secretion. Nonlinear elimination due to saturation of renal hubular reabsorption has been observed in psonatic patients at doses be-tween 7.5 and 30 mg. Impaired renal function, as well as concurrent use of drugs such as veak organic acids that also undergo tubular secretion, can markedly increase methortexate serum levels. Ex-cellent correlation has been reported between methotrexate clearance and endogenous creatinine clearance.

Methotrexate clearance rates vary widely and are generally de-created at higher does Delayed drug clearance has been identified as one of the major factors responsible for methotrexate toxicity it has been possibilized that the toxicity of methotrexate toxicity it drug ratter than the peak level is chieved. When a patient has de-layed drug alimination due to compromised renai function, a third space effusion, or other causes, methotrexate serum concentra-tions may remain elevated for prolonged periods.

The potential for taxicity from high dose regimens or delayed ex-cetion is reduced by the administration of leucovorin calcium during the final phase of methotexate plasma elimination. Pharmacoti-netic monitoring of methotexate sourun concentrations may help identify those patients at high risk for methotexate toxicity and ali in proper adjustment of leucovorin dosing. Quidelines for monitor-ing serum methotexate levels, and for adjustment of leucovorin dosing to reduce the risk of methotexate toxicity, are provided be-low in DOSAGE AND ADMINISTRATION.

Methotrexate has been detected in human breast milk. The highes breast milk to plasma concentration ratio reached was 0.081.

### INDICATIONS AND USAGE

Neoplastic Diseases Methotrexate is indicated in the treatment of gestational choriocar rinoma. chorioadenoma destruens and hydatidiform mole.

In acute lymphocytic leukemia, methotrexate is indicated in the p phytaxis of meningeal leukemia and is used in maintenance ther in combination with other chemotherapeutic agents. Methotrex is also indicated in the treatment of meningeal leukemia.

Methotrexate is used alone or in combination with other anticancer agents in the treatment of breast cancer, epidermoid cancers of the head and neck, advanced mycosis fungoides, and king cancer, par-ticularly squarous cell and small cell types. Methotrexate is also used in combination with other chemotherapeutic agents in the treat-ter of chemotherapeutic agents of the treatment of the second sec ment of advanced stage non-Hodgkin's lymphon

Methotrexate in high doses followed by leucovorin rescue in com-bination with other chemotherapeutic agents is effective in prolonging relapse-free survival in patients with non-meta-static osleosarcoma who have undergone surgical resection or ampu-tation for the primary tumor

Proclasis Methotics are is indicated in the symptomatic control of severe, re-calcitrant, disabling psoriasis that is not adequately responsive to other torms of therapy, but only when the diagnosis has been es-tablished, as by biopsy and/or after dermatologic consultation. It is important to ensure that a psoriasis thare is not due to an undiag-nosed concomitant disease affecting immune responses.

### CONTRAINDICATIONS

CONTRAINDICATIONS Methotrisate can cause tetal death or teratogenic effects when ad-ministered to a pregnant woman. Methotrozate is contraindicated in pregnant women with positiss and should be used in the treat-ment of neoplastic diseases only when the potential benefit outweights the risk to the fetus. Women of childbearing potential should not be started on methotrezate until pregnancy is excluded and should be hully counseled on the serious risk to the fetus (see PRECAUTIONS) should they become pregnant while undergoing treatment. Pregnancy should be avoided if either pathers is receiv-ing methotrezate, during and for at inimum of three months after therapy for male patients, and during and for at least one ovulatory cycle after therapy for lemale patients. (See Bored WARNINGS)

Because of the potential for serious adverse reactions from metho-trexale in breast fed inlants, it is contraindicated in nursing mothers.

Patients with psoriasis with alcoholism, alcoholic liver disease or other chronic liver disease should not receive methotrexate.

Patients with psoriasis who have overt or laboratory evidence of immunodeficiency syndromes should not receive methotrexate.

Patients with psoriasis who have preexisting blood dyscrasias, such as bone marrow hypoplasia, leukopenia, thrombocytopenia, or sig-nificant anemia, should not receive methotrexate.

Patients with a known hypersensitivity to methotrexate should not receive the drug.

### WARNINGS -SEE BOXED WARNINGS.

### PRECAUTIONS

Ceneral Metholirasate has the potential for serious toxicity (See Boxer WARNINGS) Toxic effects may be related in frequency and sever to to dose or frequency of administration but have been seen at al doses. Because they can occur at any time during therapy, it re-necessary to follow patients on methotrestate closely Most advers-reactions are reversible if detected early When such reactions do occur, the drug should be reduced in dosage or discontinued an appropriate corrective measures should be taken. If necessary, to could include the use of leucoronin calcium (See OVERDOSAGE if methotrestate therapy is reinstituted, it should be carred out and with increased alertness as to possible recurrence of toxicity.

The clinical pharmacology of methotrexate has not been well stur-red in older individuals. Due to diminished hepatic and renal function as well as decreased lotate stores in this population, relatively to doses should be considered, and these patients should be close monitored for early signs of toxicity.

Information for Patients Patients should be informed of the early signs and symptom of toxicity, of the need to see their physician promptly if the occur, and the need for close follow-up, including periodic lab-ratory tests to monitor toxicity.

Both the physician and pharmacist should emphasize to the p tient that the recommended dose is taken weekly in psonasis, at that mistaken daily use of the recommended dose has led to tai toxicity. Prescriptions should not be written or refilled on a PF

Patients should be informed of the potential benefit and risk in t use of methotrexate. The risk of effects on reproduction should

Find authenticated court documents without watermarks at docketalarm.com.

### https://web.archive.org/web/20050216215806/http://www.fda.gov/cder/foi/anda/99/40266\_Methotrexate\_Prntlbl.pdf

discussed with both male and female patients taking methotrexate

,

. . . . .

ARM

Patients undergoing methodrexate therapy should be closely moni-tored so that toxic effects are detected promptly Baseline assessment should neutode a complete blood count with differential and platiele counts, hapstec crywnes, renal innoction tests, and a chest X-ray. During therapy of psofasis, monitoring of these pa-meters is recommended, therafology at thesis monitoring to changing does or during antimecolastic therapy During insta-toring is usually indicated during antimecolastic therapy. During insta-toring is usually indicated during antimecolastic therapy. During insta-toring is usually indicated during antimecolastic therapy. During insta-toring may also be indicated. Transient time to the test test.

Transient liver function test abnormalities are observed frequently after methotexate administration and are usually not cause for modi-lication of methotexate therapy. Persistent liver function test abnormalities, and/or depression of serum atthumin may be indica-tors of serious liver forcifuly and require evaluation. (See PRECAUTIONS, Organ System Toxicity, Hepatic.)

A relationship between abnormal liver function tests and lib-rosis or cirrhosis of the liver has not been established for patients with psoriasis.

ulmonary function tests may be useful if methotrexate-induced ing disease is suspected, especially if baseline measurements re avaitable.

Drug Interactions Nonsteroidal anti-inflammatory drugs should not be administered prio to or concomitanity with the high doses of methodrexate used in the treatment of ostalosancoma. Concomitant administration of some NSAIDs with high dose methotrastate therapy has been re-ported to elevate and prolong serum methodrexate swets, resulting in deaths from severe hematologic and gastrointestinal toxicity.

Caution should be used when NSAIDs and salicylates are admini tered concombantly with lower doses of methotexate. These drup have been reported to reduce the bubutar secretion of methotre ate in an animal model and may enhence its toxicity.

Methotrexate is partially bound to serum albumin, and toxicity may be increased because of displacement by certain drugs, such as saic/ylates, phenyfbutazone, phenytoin, and autonamices. Renai bubular transport is also diminished by probeneoid, use of methotr-exate with this drug should be carefully monitored.

In the Ireatment of patients with osteosarcome, caution must be exercised if high-dose metholresate is administered in com-bination with a potentially nephrotoxic chemotherapeutic agent (eg. cispitatin)

## Oral antibodics such as tetracycline, chloramphenicol, and nonab-sorbable broad spectrum antibiotics, may decrease intestinal absorption of methodexate or interfore with the enterothepatic or-culation by inhibiting bowel flora and suppressing metabolism of the drug by bacteria.

Periciliins may reduce the renal clearance of methotrexate, in-creased serum concentrations of methotrexate with concornitant homatologic and gastrointestinal lossicity have been observed with high and low dose methotrexate. Use of methotrexate with penici-lins should be carriefully montored.

Patients receiving concomitant therapy with methotrexate and etrefinate or other retinoids should be monitored closely for pos-sible increased risk of hepatotoxicity.

Methotrexate may decrease the clearance of theophylline theophylline levels should be monitored when used con currently with methotrexate.

Vitamin preparations containing folic acid or its derivatives may decrease responses to systemically administered methotrer-ale. Preliminary animal and human studies have shown that small quantities of intravenously administered leucovorin en ter the CSF primarity as 5-methytietrahydrofolate and, in humans, remain 1-3 ordes or magnitude lower than the usual methotrevate concentrations following intrathecial administra-tion. However, high dosso al leucovorin may reduce the efficacy of intrathecally administered methotrexate.

## Folate deficiency states may increase methotrexate toxicity Timethoprim/suffamethoxazole has been reported rarely to increase

Carcinogenesis, Mutagenesis, Impairment of Fertility No controlled human data exist regarding the risk of neoplasia with methotrexate. Methotrexate has been evaluated in a number of animal studies for carcinogenic potential with inconclusive results.

Although there is evidence that methotrexate causes chromosomal damage to animal somatic cells and human bone memory cells, the chrical significance remains uncertain. Benefits should be weighed against the potential risks before using methotrexate alone or in combination with other drugs, espocially in pediatric patients or pediatric patients in the second s

Pregnancy: Teratogentic Effects, Pregnancy Category X See CONTRAIND/CATIONS.

## Nursing Mothers See CONTRAINDICATIONS.

ediatric Use alety and effectiveness in pediatric patients have not been estab hed, other than in cancer chemotherapy.

rgan System Toxicity astrainestrinal'It vorning, diarrhea, or stornalitis occur, which may suit in delyrdration, methotrexate should be discontinued und re-overy occurs. Methotrexate should be used with extreme caution it be presence of popic ulcor disease or ulcorative catifs.

atologic: Methotrexate can suppress hematopoiesis and cause via, leukopenia, and/or thrombocytopenia. In patients with ma-ncy and preveating hematopoietic impairment the drug should sed with caution, if at all.

In psoriasis, methotrexate should be stopped immediately if there is a significant drop in blood counts. In the treatment of neoplastic diseases, methotrexate should be continued only if the potential benefit warrants the risk of asvere myelosuppression. Patients with profound granulocytopenia and lever should be evaluated immediately and usually require parenteral broad-spectrum antibiotic therapy

Hepatic: Methotrexate has the potential for acute (elevated tran-saminases) and chronic (librosis and ciritosis) hepatotoxicity. Chronic loxicity is potentially Istal, it generally has occurred after protonged use (generally two years or more) and after a total does of at least 1.5 grams. In studies in psoriatic patients, hepatotoxicity appeared to be a function of total curuulative does and appeared to be enhanced by alcoholism, obesity, diabetes and advanced age. An accurate incidence rate has not been determined; the rate of progression and reversibility of lesions is not known. Special cau-tion is indicated in the presence of preexisting liver damage of impaired hepatic function.

In poniesis, liver function tests, including serum albumin, should be performed periodically prior to dosing but are often normal in the face of developing throats or princips. These leatons may be detectable only by biopsy. The usual rocommendation is to obtain a liver biopsy at 1) pretmerapy or shortly after initia-tion of threary (2 - 4 months), 2) a total curvulative dose of 1.5 grans, and 3) after each additional 1.0 to 1.5 grans. Moderate fibrosis or any cirrhosis normally saggests a repeat biopsy in 6 months. Milder histologic findings such as latty change and low grade portal inflammation, are relatively common pretherapy. Atthough these thread changes are usually not a ree-son to avoid or discontinue methotirexate therapy, the drug should be used with caution. uld be used with caution

Infection or Immunologic States: Methotrexate should be used infection or Immunologic States: Methotrexate should be used usually contraindicated in patients with over or laboratory evi-dence of immunodeficiency syndromes. Immunutation may be ineffective when given during methotrexate therapy. Immun-zation with ive virus vaccines is generally not recommended. There have been reports of disseminated vaccinia infections after smallpox immunizations in patients receiving methotiex-ate thetapy. Hypogammaglobulinemia has been reported rarely.

Potentially fatal opportunistic infections, especially *Pneumocyc* cariw pneumonia, may occur with methotrexate thorapy. Whe patient presents with putknonary symptoms, the possibility *Pneumocystis camii* pneumonia should be considered.

Prevince years cannot be entransing to the considered Neurologic: There have been reports of lexitoencephalopathy tolowing infravenous administration of methodrexate to patients who have had crainsignial intradiation. Sanous neurotoxicity fra-quently mandrested as generalized on focal seizures, has been reported with unexpectedly increased fraquency among pedi-atric patients with acute lymphoblastic leukenva who weter treated with intermediate-docs intravenous methodrexate (1 gm/ m<sup>3</sup>). Symptomatic patients were commonly noted to have leu-koncrephalopathy and/or microangiopathic calcifications on diagnostic imaging studies. Chronic leukencephalopathy has also been reported in patients who received repeated doses of high-dose methotrexate with leucovorin rescue even without

cranial irradation. Discontinuation of methotrexate does not al-ways result in complete recovery. A transient acute neurologic syndrome has been observed in pa-tients treated with high dose regimens. Manifestations of this stroke-like encephalopathy may include conflusion, hemiparesis, seizures and coma. The exact cause is unknown.

After the intrathecal use of methotrexate, the central nervous system toxicity which may occur can be classified as follows: acute chemical arachnoidits manifested by such symptoms as headach. Back pain, muchal rigidity, and fever; sub-acute myelopathy characterized by paraparesis/paraplegia associated with involvement with one or more spinal nerver roots; chronic leukoencephalopathy manifested by confusion, initability, som-nolence, ataxia, dementia, ascurues and coma. This condition can be progressive and even latal.

Pulmonary: Pulmonary symptoms (especially a dry nonproductive couplin or a non-specific pneumonitis occurring during methotres: ale therapy may be indicative of a potentially dangerous lesion and require interruption of treatment and careful investigation. Although chrically variable, the typical patient with methotresials indiced lung disease presents with lever, cough, dyspina. hypoxemia, and an infiftrate on chest x-ray, infection needs to be excluded. This lesion can occur at all dosages.

Renat: High doses of methotraxate used in the treatment of os-teosarcoma may cause renat damage leading to acute renat lature. Nephrotoxicity is due primarity to the precipitation of methotraxate and 7-hydroxymethotraxate in the renat lubules: Close attention to renat function including adequate hydration, urine alkalmization and measurement of serum methotrexate and creatinne levels are es-sential for safe administration.

Skin: Severa, occasionally fatal, dermatologic reactions, includ-ing toxic epidermal necrolysis, Stevens-Johnson syndrome, extolative dematitis, skin encrosis, acute y them multiorne, have been reported in children and acute y them days of oral, intranuscular, intravenous, or intrathecal metals admin-istration. Reactions were noted after single or multiple low, intermediate, or high doses of metholravate in patients with neoplestic and non-neoplestic diseases.

Other precautions: Methotrexate should be used with extreme cau-tion in the presence of debility.

Mothotrexate exits slowly from third space compartments (eg. pleural effusions or ascries). This results in a proionged ferminal plasma half-kile and unexpected loxocity. In patients with significant third space accumulations, it is advisable to evacuate the fluid before treatment and to monitor plasma methotrexate levels.

Lesions of psoniasis may be apgravated by concomitant exposure to ultraviolet radiation. Accitation dermatitis and sunburn may be recalled by the use of methotrexate.

ADVERSE REACTIONS IN GENERAL THE INCIDENCE AND SEVERITY OF ACUTE SIDE EFFECTS ARE RELATED TO DOSE AND FREQUENCY OF AD-MINISTRATIONS ARE UNSURSED ABOVE UNDER ORGAN SYSTEM TOXICITY IN THE PRECAUSE SCHON THAT SECTION SHOULD ALSO BE CONSULTED WHEN LOOKING FOR INFORMATION ABOUT ADVERSE REACTIONS WITH METHOTREKATE.

The most frequently reported adverse reactions include ulcarative stomatils, ieukopenia, nausea, and abdominal distines. Other fre-quently reported adverse effects are malaise, undue failque, chilis and lever, dizziness and decreased resistance to intection.

ADVERSE REACTIONS

neoplastic and non-neoplastic diseasus. Pneumocyctis carinii, monia was the most common inflection. Other reported infle-included nocedosis, histoplasmosis, cryptococoasis, Herpes z H. simplex hepatitis, and disseminated H. simplex.

Ophthalmic: conjunctivitis, serious visual changes of unknown etic

Pulmonary System: interstitial pneumonitis deaths have bee ported, and chronic interstitial obstructive pulmonary disease occasionally occurred.

Skin: erythematous rashes, pruritus, unicaria, photosen: ity, pigmentary changes, alopecia, ecchymosis, telangiect acne, turunciosis, erythema multiforme, toxic epider necrolysis, Stevens-Johnson syndrome, skin necrosis, effoliative dematitis.

Urogenital System: severe nephropathy or renal failure, azoter cystilis, hematuria; defective oogenesis or spemalogenesis, li serici dioposermia, menstruai dystancido, nu vaginal discharge, gynecomastia; infertiëty, abortion, letal defects.

Other rarer reactions related to or attributed to the use of mr ofrexate such as nodulosis, vascultis, arthratgia/myatgia, k of libido/impolence, diabetes, osteoporosis, sudden death, versible lymphomas, and tumor lysis syndrome. Anaphylact reactions have been reported.

Adverse Reactions in Psortasis: Dara no recent placebo-controlled trials in pasents with psor is: There are two literature reports (Roaniçk, 1969, and Mylo 1978) descriptions (Jarge series (n.204, 248) of psonasis patier realized and the methodreaste. Dosages ranged up to 25 mg per we and trials and the series of toru to four years. Whit the is option of biopcoa, photosensitivity and "burning of skin lessor (each 3% to thoose in the rheumatod arthritis situdies.

### OVERDOSAGE

OVEROCSAGE Leucovorin is indicated to diminish the toxicity and counteract the effect of inadvertently administered overdosages of methodraxat. Leu-covorin administration should begin as promptly as possible As the time interval between methodraxat administration and leu-covortin ini-liation increases, the effectiveness of ieu-covorin iounteracting toxicity decreases. Monitoring of the serum methot exate concentra-tion is essential in determining the optimal dos and duration of treatment with leucovorin.

In cases of massive overdosage, hydration and urnary alkaliniza tion may be necessary to prevent the precipitation of methodrezal and/or its metabolities in the renal tabules. Neither hemodialyais nor perior-neal dialysis has been shown to improve methodrezate elimination.

Accidental infrathecal overdosage may require intensive systemic support, high-dose systemic leucovorin, alkaline druresis and rapid CSF dramage and ventricololumbar perfusion.

## DOSAGE AND ADMINISTRATION

VOSAUC AND DAMINISTRATION Neoplastic Diseases Oral administration in tablet form is often preferred when low doses are being administered since absorption is rapid and ef-fective serum levels are obtained. Methotrexate for Injection may be given by the inframuscular, intravenous, intra-arterial, or in-frathecal route.

or in-trathecal route. Chorocarcinoma and similar trophobiastic diseases: Methotrexate is administered orally or intramuscularly in doses of 15 to 30 mg dayl for a five-day course. Such courses are usually repeated for 3 to 5 miss as required, with rest periods of more weeks interposed between courses. Unil any manifestimic more weeks subside: The efficiencess of interary is ontarily more units by thick should return to normal or less man 50 (U24 tr in unit) with the third or fourth course and usually be followed by usually after of method: Becale and the original of the discussion of the General establishing the each course of the discussion of the General establishing the each course of the discussion of the General establishing the each course of the discussion for the original session antilumor drugs has been reported as being useful. Sance hurtatisfism mole may smeetic chorocarcinoma. prophy-

Since hydatidiform mole may precede chonocarcinoma, prophy lactic chemotherapy with methotrexate has been recommended.

Chorioadenoma destruens is considered to be an invasive lorm of hydatid/orm mole. Methotrexate is administered in these disease states in doses similar to those recommended for choriocarcinoma.

Leukema: Acute lymphoblastic leukema in padiatric patients and young adolescents is the most responsive to present day chemo-therapy. In young aduits and older patients, clinical remission is more difficult to obtain and early relapse is more common.

Find authenticated court documents without watermarks at docketalarm.com.

DOCKET

ALARM

Other adverse reactions that have been reported with methotrex-ale are listed below by organ system. In the oncology setting, concomitant treatment and the underlying disease make specific attribution of a reaction to methotrezale difficult. timentary System: gingivitis, pharyngitis, stomatitis, anorexia, nau-ea, vomiting, diarrhea, hematemesis, melena, gastrointestinal /ceration and bieeding, enteniis, pancreatitis. Cardiovascular: pericarditis, pericardial effusion, hypotension, and thromboembolic events (including arterial thrombosis, carebral thrombosis, deep vein thrombosis, retinal vein thrombosis, throm-bophebits, and pulmonary embolus).

Central Mervous System: headaches, drowsiness, blurred vision. Aphasia, hemiparesis, paresis and convulsions have also occurred following administration of methodraxale. Following low does, there have been occasional reports of transmit suble cognitive dyslunc-tion, mood alteration or unusual cranial sensations.

Infection: There have been case reports of sometimes fatal oppor-tunistic infections in patients receiving methotrexate therapy for

## DOCKET A L A R M



# Explore Litigation Insights

Docket Alarm provides insights to develop a more informed litigation strategy and the peace of mind of knowing you're on top of things.

## **Real-Time Litigation Alerts**



Keep your litigation team up-to-date with **real-time alerts** and advanced team management tools built for the enterprise, all while greatly reducing PACER spend.

Our comprehensive service means we can handle Federal, State, and Administrative courts across the country.

## **Advanced Docket Research**



With over 230 million records, Docket Alarm's cloud-native docket research platform finds what other services can't. Coverage includes Federal, State, plus PTAB, TTAB, ITC and NLRB decisions, all in one place.

Identify arguments that have been successful in the past with full text, pinpoint searching. Link to case law cited within any court document via Fastcase.

## **Analytics At Your Fingertips**



Learn what happened the last time a particular judge, opposing counsel or company faced cases similar to yours.

Advanced out-of-the-box PTAB and TTAB analytics are always at your fingertips.

## API

Docket Alarm offers a powerful API (application programming interface) to developers that want to integrate case filings into their apps.

## LAW FIRMS

Build custom dashboards for your attorneys and clients with live data direct from the court.

Automate many repetitive legal tasks like conflict checks, document management, and marketing.

## FINANCIAL INSTITUTIONS

Litigation and bankruptcy checks for companies and debtors.

## E-DISCOVERY AND LEGAL VENDORS

Sync your system to PACER to automate legal marketing.