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**MYLERAN®** 

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4 Tablets

#### WARNING

MYLERAN is a potent drug. It should not be used unless a diagnosis of chronic myelogenous leukemia has been adequately established and the responsible physician is knowledgeable in assessing response to chemotherapy.

MYLERAN can induce severe bone marrow hypoplasia. Reduce or discontinue the dosage immediately at the first sign of any unusual depression of bone marrow function as reflected by an abnormal decrease in any of the formed elements of the blood. A bone marrow examination should be performed if the bone marrow status is uncertain.

SEE WARNINGS FOR INFORMATION REGARDING BUSULFAN-INDUCED

15 LEUKEMOGENESIS IN HUMANS.

#### DESCRIPTION

MYLERAN (busulfan) is a bifunctional alkylating agent. Busulfan is known chemically as 1,4-butanediol dimethanesulfonate and has the following structural formula:

CH<sub>3</sub>SO<sub>2</sub>O(CH<sub>2</sub>)<sub>4</sub>OSO<sub>2</sub>CH<sub>3</sub>

Busulfan is *not* a structural analog of the nitrogen mustards. MYLERAN is available in tablet form for oral administration. Each film-coated tablet contains 2 mg busulfan and the inactive ingredients hypromellose, lactose (anhydrous), magnesium stearate, pregelatinized starch, triacetin, and titanium dioxide.

The activity of busulfan in chronic myelogenous leukemia was first reported by D.A.G. Galton in 1953.

#### CLINICAL PHARMACOLOGY

31	Busulfan is a small, highly lipophilic molecule that easily crosses the blood brain barrier.
32	Following absorption, 32% and 47% of busulfan are bound to plasma proteins and red blood cells,
33	respectively.
34	Busulfan absorption from the gastrointestinal tract is essentially complete. This has been
35	demonstrated in radioactive studies after both intravenous and oral administration of <sup>35</sup> S-busulfan,
36	<sup>14</sup> C-busulfan, and <sup>3</sup> H-busulfan. Following intravenous administration of a single therapeutic dose of
37	<sup>35</sup> S-busulfan, there was rapid disappearance of radioactivity from the blood and 90% to 95% of the
38	<sup>35</sup> S-label disappeared within 3 to 5 minutes after injection. After either oral or intravenous
39	administration of <sup>35</sup> S-busulfan, 45% to 60% of the radioactivity was recovered in the urine in the
40	48 hours after administration; the majority of the total urinary excretion occurring in the first 24 hours.
41	Over 95% of the urinary <sup>35</sup> S-label occurs as <sup>35</sup> S-methanesulfonic acid. Oral and intravenous
42	administration of 1,4-14C-busulfan showed the same rapid initial disappearance of plasma
43	radioactivity as observed following the administration of <sup>35</sup> S-labeled drug. Cumulative radioactivity
44	in the urine after 48 hours was 25% to 30% of the administered dose (contrasting with 45% to 60%
45	for <sup>35</sup> S-busulfan), and suggests a slower excretion of the alkylating portion of the molecule and its
46	metabolites than for the sulfonoxymethyl moieties. Regardless of the route of administration,
47	1,4-14C-busulfan yielded a complex mixture of at least 12 radiolabeled metabolites in urine; the main
48	metabolite being 3-hydroxytetrahydrothiophene-1,1-dioxide. Pharmacokinetic studies employing <sup>3</sup> H-
49	busulfan labeled on the tetramethylene chain confirmed a rapid initial clearance of the radioactivity
50	from plasma, irrespective of whether the drug was given orally or intravenously.
51	A study compared a 2-mg single IV bolus injection to a single oral dose of a 2-mg tablet of
52	nonradioactive busulfan in 8 adult patients 13 to 60 years of age. The study demonstrated that the
53	mean? SD absolute bioavailability was 80%? 20% in adults. However, the absolute bioavailability
54	for 8 children 1.5 to 6 years of age was 68%? 31%.
55	In another study of 2, 4, and 6 mg of busulfan, given as a single oral dose on consecutive days
56	(starting with the lowest dose) in 5 adult patients, the mean dose-normalized (to 2 mg dose) area
57	under the plasma concentration-time curve (AUC) was about 130 ng?hr/mL, while the mean intra- and
58	inter-patient variability was about 16% and 21%, respectively. Busulfan was eliminated with a
59	plasma terminal elimination half-life $(t_{1/2})$ of about 2.6 hours, and demonstrated linear kinetics within
60	the range of 2 to 6 mg for both the maximum plasma concentration ( $C_{max}$ ) and AUC. The mean $C_{max}$ for
61	the 2-, 4-, and 6-mg doses (after dose normalization to 2 mg) was about 30 ng/mL. A recent study of 4

62	to 8 mg as single oral doses in 12 patients showed that the mean ? SD $C_{\text{max}}$ (after dose normalization
63	to 4 mg) was 68.2 ? 24.4 ng/mL, occurring at about 0.9 hours and the mean ? SD AUC (after dose
64	normalization to 4 mg) was 269 ? 62 ng?hr/mL. These results are consistent with previous results. In
65	addition, the mean? SD elimination half-life was 2.69? 0.49 hours.
66	The elimination of busulfan appears to be independent of renal function. This probably reflects the
67	extensive metabolism of the drug in the liver, since less than 2% of the administered dose is excreted
68	in the urine unchanged within 24 hours. The drug is metabolized by enzymatic activity to at least
69	12 metabolites, among which tetrahydrothiophene, tetrahydrothiophene 12-oxide, sulfolane, and
70	3-hydroxysulfolane were identified. These metabolites do not have cytotoxic activity.
71	There is no experience with the use of dialysis in an attempt to modify the clinical toxicity of
72	busulfan. One technical difficulty would derive from the extremely poor water solubility of busulfan.
73	Additionally, all studies of the metabolism of busulfan employing radiolabeled materials indicate
74	rapid chemical reactivity of the parent compound with prolonged retention of some of the metabolites
75	(particularly the metabolites arising from the "alkylating" portion of the molecule). The effectiveness
76	of dialysis at removing significant quantities of unreacted drug would be expected to be minimal in
77	such a situation.
78	Currently, there are no available data on the effect of food on busulfan bioavailability.
79	<b>Biochemical Pharmacology:</b> In aqueous media, busulfan undergoes a wide range of nucleophilic
80	substitution reactions. While this chemical reactivity is relatively non-specific, alkylation of the DNA
81	is felt to be an important biological mechanism for its cytotoxic effect. Coliphage T7 exposed to
82	busulfan was found to have the DNA crosslinked by intrastrand crosslinkages, but no interstrand
83	linkages were found.
84	The metabolic fate of busulfan has been studied in rats and humans using <sup>14</sup> C- and <sup>35</sup> S-labeled
85	materials. In humans, as in the rat, almost all of the radioactivity in <sup>35</sup> S-labeled busulfan is excreted in
86	the urine in the form of <sup>35</sup> S-methanesulfonic acid. Roberts and Warwick demonstrated that the
87	formation of methanesulfonic acid in vivo in the rat is not due to a simple hydrolysis of busulfan to
88	1,4-butanediol, since only about 4% of 2,3-14C-busulfan was excreted as carbon dioxide, whereas
89	2,3-14C-1,4-butanediol was converted almost exclusively to carbon dioxide. The predominant
90	reaction of busulfan in the rat is the alkylation of sulfhydryl groups (particularly cysteine and
91	cysteine-containing compounds) to produce a cyclic sulfonium compound which is the precursor of

the major urinary metabolite of the 4-carbon portion of the molecule, 3-hydroxytetrahydrothiophene-92 1,1-dioxide. This has been termed a "sulfur-stripping" action of busulfan and it may modify the 93 function of certain sulfur-containing amino acids, polypeptides, and proteins; whether this action 94 95 makes an important contribution to the cytotoxicity of busulfan is unknown. The biochemical basis for acquired resistance to busulfan is largely a matter of speculation. 96 97 Although altered transport of busulfan into the cell is one possibility, increased intracellular inactivation of the drug before it reaches the DNA is also possible. Experiments with other alkylating 98 agents have shown that resistance to this class of compounds may reflect an acquired ability of the 99 resistant cell to repair alkylation damage more effectively. 100 **Clinical Studies:** Although not curative, busulfan reduces the total granulocyte mass, relieves 101 102 symptoms of the disease, and improves the clinical state of the patient. Approximately 90% of adults with previously untreated chronic myelogenous leukemia will obtain hematologic remission with 103 regression or stabilization of organomegaly following the use of busulfan. It has been shown to be 104 superior to splenic irradiation with respect to survival times and maintenance of hemoglobin levels, 105 106 and to be equivalent to irradiation at controlling splenomegaly. 107 It is not clear whether busulfan unequivocally prolongs the survival of responding patients beyond the 31 months experienced by an untreated group of historical controls. Median survival figures of 31 108 to 42 months have been reported for several groups of patients treated with busulfan, but concurrent 109 control groups of comparable, untreated patients are not available. The median survival figures 110 reported from different studies will be influenced by the percentage of "poor risk" patients initially 111 entered into the particular study. Patients who are alive 2 years following the diagnosis of chronic 112 113 myelogenous leukemia, and who have been treated during that period with busulfan, are estimated to have a mean annual mortality rate during the second to fifth year which is approximately two thirds 114 that of patients who received either no treatment, conventional x-ray or <sup>32</sup>P-irradiation, or 115 116 chemotherapy with minimally active drugs. Busulfan is clearly less effective in patients with chronic myelogenous leukemia who lack the 117 Philadelphia (Ph<sup>1</sup>) chromosome. Also, the so-called "juvenile" type of chronic myelogenous 118 leukemia, typically occurring in young children and associated with the absence of a Philadelphia 119 chromosome, responds poorly to busulfan. The drug is of no benefit in patients whose chronic 120 121 myelogenous leukemia has entered a "blastic" phase.

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MYLERAN should not be used in patients whose chronic myelogenous leukemia has demonstrated

123	prior resistance to this drug.
124	MYLERAN is of no value in chronic lymphocytic leukemia, acute leukemia, or in the "blastic
125	crisis" of chronic myelogenous leukemia.
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127	INDICATIONS AND USAGE
128	MYLERAN (busulfan) is indicated for the palliative treatment of chronic myelogenous (myeloid,
129	myelocytic, granulocytic) leukemia.
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131	CONTRAINDICATIONS
132	MYLERAN is contraindicated in patients in whom a definitive diagnosis of chronic myelogenous
133	leukemia has not been firmly established.
134	MYLERAN is contraindicated in patients who have previously suffered a hypersensitivity reaction
135	to busulfan or any other component of the preparation.
136	
137	WARNINGS
138	The most frequent, serious side effect of treatment with busulfan is the induction of bone marrow
139	failure (which may or may not be anatomically hypoplastic) resulting in severe pancytopenia. The
140	pancytopenia caused by busulfan may be more prolonged than that induced with other alkylating
141	agents. It is generally felt that the usual cause of busulfan-induced pancytopenia is the failure to stop
142	administration of the drug soon enough; individual idiosyncrasy to the drug does not seem to be an
143	important factor. MYLERAN should be used with extreme caution and exceptional vigilance in
144	patients whose bone marrow reserve may have been compromised by prior irradiation or
145	chemotherapy, or whose marrow function is recovering from previous cytotoxic therapy. Although
146	recovery from busulfan-induced pancytopenia may take from 1 month to 2 years, this complication is
147	potentially reversible, and the patient should be vigorously supported through any period of severe
148	pancytopenia.
149	A rare, important complication of busulfan therapy is the development of bronchopulmonary
150	dysplasia with pulmonary fibrosis. Symptoms have been reported to occur within 8 months to
151	10 years after initiation of therapy—the average duration of therapy being 4 years. The histologic

findings associated with "busulfan lung" mimic those seen following pulmonary irradiation.
Clinically, patients have reported the insidious onset of cough, dyspnea, and low-grade fever. In some
cases, however, onset of symptoms may be acute. Pulmonary function studies have revealed
diminished diffusion capacity and decreased pulmonary compliance. It is important to exclude more
common conditions (such as opportunistic infections or leukemic infiltration of the lungs) with
appropriate diagnostic techniques. If measures such as sputum cultures, virologic studies, and
exfoliative cytology fail to establish an etiology for the pulmonary infiltrates, lung biopsy may be
necessary to establish the diagnosis. Treatment of established busulfan-induced pulmonary fibrosis is
unsatisfactory; in most cases the patients have died within 6 months after the diagnosis was
established. There is no specific therapy for this complication. MYLERAN should be discontinued if
this lung toxicity develops. The administration of corticosteroids has been suggested, but the results
have not been impressive or uniformly successful.
Busulfan may cause cellular dysplasia in many organs in addition to the lung. Cytologic
abnormalities characterized by giant, hyperchromatic nuclei have been reported in lymph nodes,
pancreas, thyroid, adrenal glands, liver, and bone marrow. This cytologic dysplasia may be severe
enough to cause difficulty in interpretation of exfoliative cytologic examinations from the lung,
bladder, breast, and the uterine cervix.
In addition to the widespread epithelial dysplasia that has been observed during busulfan therapy,
chromosome aberrations have been reported in cells from patients receiving busulfan.
Busulfan is mutagenic in mice and, possibly, in humans.
Malignant tumors and acute leukemias have been reported in patients who have received busulfan
therapy, and this drug may be a human carcinogen. The World Health Organization has concluded that
there is a causal relationship between busulfan exposure and the development of secondary
malignancies. Four cases of acute leukemia occurred among 243 patients treated with busulfan as
adjuvant chemotherapy following surgical resection of bronchogenic carcinoma. All 4 cases were
from a subgroup of 19 of these 243 patients who developed pancytopenia while taking busulfan 5 to 8
years before leukemia became clinically apparent. These findings suggest that busulfan is
leukemogenic, although its mode of action is uncertain.
Ovarian suppression and amenorrhea with menopausal symptoms commonly occur during busulfan
therapy in premenopausal patients. Busulfan has been associated with ovarian failure including
failure to achieve puberty in females. Busulfan interferes with spermatogenesis in experimental

183 animals, and there have been clinical reports of sterility, azoospermia, and testicular atrophy in male patients. 184 Hepatic veno-occlusive disease, which may be life threatening, has been reported in patients 185 186 receiving busulfan, usually in combination with cyclophosphamide or other chemotherapeutic agents prior to bone marrow transplantation. Possible risk factors for the development of hepatic 187 188 veno-occlusive disease include: total busulfan dose exceeding 16 mg/kg based on ideal body weight, and concurrent use of multiple alkylating agents (see CLINICAL PHARMACOLOGY and Drug 189 Interactions). 190 A clear cause-and-effect relationship with busulfan has not been demonstrated. Periodic 191 measurement of serum transaminases, alkaline phosphatase, and bilirubin is indicated for early 192 193 detection of hepatotoxicity. A reduced incidence of hepatic veno-occlusive disease and other regimen-related toxicities have been observed in patients treated with high-dose MYLERAN and 194 cyclophosphamide when the first dose of cyclophosphamide has been delayed for >24 hours after the 195 last dose of busulfan (see CLINICAL PHARMACOLOGY and Drug Interactions). 196 197 Cardiac tamponade has been reported in a small number of patients with thalassemia (2% in one series) who received busulfan and cyclophosphamide as the preparatory regimen for bone marrow 198 transplantation. In this series, the cardiac tamponade was often fatal. Abdominal pain and vomiting 199 200 preceded the tamponade in most patients. 201 **Pregnancy:** Pregnancy Category D. Busulfan may cause fetal harm when administered to a pregnant woman. Although there have been a number of cases reported where apparently normal children have 202 been born after busulfan treatment during pregnancy, one case has been cited where a malformed baby 203 204 was delivered by a mother treated with busulfan. During the pregnancy that resulted in the malformed infant, the mother received x-ray therapy early in the first trimester, mercaptopurine until the third 205 206 month, then busulfan until delivery. In pregnant rats, busulfan produces sterility in both male and female offspring due to the absence of germinal cells in testes and ovaries. Germinal cell aplasia or 207 sterility in offspring of mothers receiving busulfan during pregnancy has not been reported in humans. 208 209 There are no adequate and well-controlled studies in pregnant women. If this drug is used during pregnancy, or if the patient becomes pregnant while taking this drug, the patient should be apprised of 210 211 the potential hazard to the fetus. Women of childbearing potential should be advised to avoid 212 becoming pregnant.

PRECAUTIONS
<b>General:</b> The most consistent, dose-related toxicity is bone marrow suppression. This may be
manifest by anemia, leukopenia, thrombocytopenia, or any combination of these. It is imperative that
patients be instructed to report promptly the development of fever, sore throat, signs of local
infection, bleeding from any site, or symptoms suggestive of anemia. Any one of these findings may
indicate busulfan toxicity; however, they may also indicate transformation of the disease to an acute
"blastic" form. Since busulfan may have a delayed effect, it is important to withdraw the medication
temporarily at the first sign of an abnormally large or exceptionally rapid fall in any of the formed
elements of the blood. Patients should never be allowed to take the drug without close medical
supervision.
Seizures have been reported in patients receiving busulfan. As with any potentially epileptogenic
drug, caution should be exercised when administering busulfan to patients with a history of seizure
disorder, head trauma, or receiving other potentially epileptogenic drugs. Some investigators have
used prophylactic anticonvulsant therapy in this setting.
Information for Patients: Patients beginning therapy with busulfan should be informed of the
importance of having periodic blood counts and to immediately report any unusual fever or bleeding.
Aside from the major toxicity of myelosuppression, patients should be instructed to report any
difficulty in breathing, persistent cough, or congestion. They should be told that diffuse pulmonary
fibrosis is an infrequent, but serious and potentially life-threatening complication of long-term
busulfan therapy. Patients should be alerted to report any signs of abrupt weakness, unusual fatigue,
anorexia, weight loss, nausea and vomiting, and melanoderma that could be associated with a
syndrome resembling adrenal insufficiency. Patients should never be allowed to take the drug without
medical supervision and they should be informed that other encountered toxicities to busulfan include
infertility, amenorrhea, skin hyperpigmentation, drug hypersensitivity, dryness of the mucous
membranes, and rarely, cataract formation. Women of childbearing potential should be advised to
avoid becoming pregnant. The increased risk of a second malignancy should be explained to the
patient.
Laboratory Tests: It is recommended that evaluation of the hemoglobin or hematocrit, total white
blood cell count and differential count, and quantitative platelet count be obtained weekly while the
patient is on busulfan therapy. In cases where the cause of fluctuation in the formed elements of the

244	peripheral blood is obscure, bone marrow examination may be useful for evaluation of marrow status.
245	A decision to increase, decrease, continue, or discontinue a given dose of busulfan must be based not
246	only on the absolute hematologic values, but also on the rapidity with which changes are occurring.
247	The dosage of busulfan may need to be reduced if this agent is combined with other drugs whose
248	primary toxicity is myelosuppression. Occasional patients may be unusually sensitive to busulfan
249	administered at standard dosage and suffer neutropenia or thrombocytopenia after a relatively short
250	exposure to the drug. Busulfan should not be used where facilities for complete blood counts,
251	including quantitative platelet counts, are not available at weekly (or more frequent) intervals.
252	Drug Interactions: Busulfan may cause additive myelosuppression when used with other
253	myelosuppressive drugs.
254	In one study, 12 of approximately 330 patients receiving continuous busulfan and thioguanine
255	therapy for treatment of chronic myelogenous leukemia were found to have portal hypertension and
256	esophageal varices associated with abnormal liver function tests. Subsequent liver biopsies were
257	performed in 4 of these patients, all of which showed evidence of nodular regenerative hyperplasia.
258	Duration of combination therapy prior to the appearance of esophageal varices ranged from 6 to
259	45 months. With the present analysis of the data, no cases of hepatotoxicity have appeared in the
260	busulfan-alone arm of the study. Long-term continuous therapy with thioguanine and busulfan should
261	be used with caution.
262	Busulfan-induced pulmonary toxicity may be additive to the effects produced by other cytotoxic
263	agents.
264	The concomitant systemic administration of itraconazole to patients receiving high-dose
265	MYLERAN may result in reduced busulfan clearance (see CLINICAL PHARMACOLOGY). Patients
266	should be monitored for signs of busulfan toxicity when itraconazole is used concomitantly with
267	MYLERAN.
268	Busulfan clearance may be reduced in the presence of cyclophosphamide (see CLINICAL
269	PHARMACOLOGY).
270	Carcinogenesis, Mutagenesis, Impairment of Fertility: See WARNINGS section. The World
271	Health Organization has concluded that there is a causal relationship between busulfan exposure and
272	the development of secondary malignancies.
273	Pregnancy: Teratogenic Effects: Pregnancy Category D. See WARNINGS section.

274	Nonteratogenic Effects: There have been reports in the literature of small infants being born
275	after the mothers received busulfan during pregnancy, in particular, during the third trimester. One
276	case was reported where an infant had mild anemia and neutropenia at birth after busulfan was
277	administered to the mother from the eighth week of pregnancy to term.
278	Nursing Mothers: It is not known whether this drug is excreted in human milk. Because of the
279	potential for tumorigenicity shown for busulfan in animal and human studies, a decision should be
280	made whether to discontinue nursing or to discontinue the drug, taking into account the importance of
281	the drug to the mother.
282	Pediatric Use: See INDICATIONS AND USAGE and DOSAGE AND ADMINISTRATION
283	sections.
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285	ADVERSE REACTIONS
286	Hematological Effects: The most frequent, serious, toxic effect of busulfan is dose-related
287	myelosuppression resulting in leukopenia, thrombocytopenia, and anemia. Myelosuppression is most
288	frequently the result of a failure to discontinue dosage in the face of an undetected decrease in
289	leukocyte or platelet counts.
290	Aplastic anemia (sometimes irreversible) has been reported rarely, often following long-term
291	conventional doses and also high doses of MYLERAN.
292	Pulmonary: Interstitial pulmonary fibrosis has been reported rarely, but it is a clinically significant
293	adverse effect when observed and calls for immediate discontinuation of further administration of the
294	drug. The role of corticosteroids in arresting or reversing the fibrosis has been reported to be
295	beneficial in some cases and without effect in others.
296	Cardiac: Cardiac tamponade has been reported in a small number of patients with thalassemia who
297	received busulfan and cyclophosphamide as the preparatory regimen for bone marrow transplantation
298	(see WARNINGS).
299	One case of endocardial fibrosis has been reported in a 79-year-old woman who received a total
300	dose of 7,200 mg of busulfan over a period of 9 years for the management of chronic myelogenous
301	leukemia. At autopsy, she was found to have endocardial fibrosis of the left ventricle in addition to
302	interstitial pulmonary fibrosis.

303	<b>Ocular:</b> Busulfan is capable of inducing cataracts in rats and there have been several reports
304	indicating that this is a rare complication in humans.
305	<b>Dermatologic:</b> Hyperpigmentation is the most common adverse skin reaction and occurs in 5% to
306	10% of patients, particularly those with a dark complexion.
307	Metabolic: In a few cases, a clinical syndrome closely resembling adrenal insufficiency and
308	characterized by weakness, severe fatigue, anorexia, weight loss, nausea and vomiting, and
309	melanoderma has developed after prolonged busulfan therapy. The symptoms have sometimes been
310	reversible when busulfan was withdrawn. Adrenal responsiveness to exogenously administered
311	ACTH has usually been normal. However, pituitary function testing with metyrapone revealed a
312	blunted urinary 17-hydroxycorticosteroid excretion in 2 patients. Following the discontinuation of
313	busulfan (which was associated with clinical improvement), rechallenge with metyrapone revealed
314	normal pituitary-adrenal function.
315	Hyperuricemia and/or hyperuricosuria are not uncommon in patients with chronic myelogenous
316	leukemia. Additional rapid destruction of granulocytes may accompany the initiation of chemotherapy
317	and increase the urate pool. Adverse effects can be minimized by increased hydration, urine
318	alkalinization, and the prophylactic administration of a xanthine oxidase inhibitor such as allopurinol
319	<b>Hepatic Effects:</b> Esophageal varices have been reported in patients receiving continuous busulfan
320	and thioguanine therapy for treatment of chronic myelogenous leukemia (see PRECAUTIONS: Drug
321	Interactions). Hepatic veno-occlusive disease has been observed in patients receiving busulfan (see
322	WARNINGS).
323	Miscellaneous: Other reported adverse reactions include: urticaria, erythema multiforme, erythema
324	nodosum, alopecia, porphyria cutanea tarda, excessive dryness and fragility of the skin with
325	anhidrosis, dryness of the oral mucous membranes and cheilosis, gynecomastia, cholestatic jaundice,
326	and myasthenia gravis. Most of these are single case reports, and in many, a clear cause-and-effect
327	relationship with busulfan has not been demonstrated.
328	Seizures (see PRECAUTIONS: General) have been observed in patients receiving higher than
329	recommended doses of busulfan.
330	Observed During Clinical Practice: The following events have been identified during post-
331	approval use of busulfan. Because they are reported voluntarily from a population of unknown size,

332	estimates of frequency cannot be made. These events have been chosen for inclusion due to a
333	combination of their seriousness, frequency of reporting, or potential causal connection to busulfan.
334	Blood and Lymphatic: Aplastic anemia.
335	Eye: Cataracts, corneal thinning, lens changes.
336	Hepatobiliary Tract and Pancreas: Centrilobular sinusoidal fibrosis, hepatic veno-occlusive
337	disease, hepatocellular atrophy, hepatocellular necrosis, hyperbilirubinemia (see WARNINGS).
338	Non-site Specific: Infection, mucositis, sepsis.
339	Respiratory: Pneumonia.
340	Skin: Rash. An increased local cutaneous reaction has been observed in patients receiving
341	radiotherapy soon after busulfan.
342	
343	OVERDOSAGE
344	There is no known antidote to busulfan. The principal toxic effects are bone marrow depression
345	and pancytopenia. The hematologic status should be closely monitored and vigorous supportive
346	measures instituted if necessary. Induction of vomiting or gastric lavage followed by administration of
347	charcoal would be indicated if ingestion were recent. Dialysis may be considered in the management
348	of overdose as there is 1 report of successful dialysis of busulfan (see
349	CLINICALPHARMACOLOGY).
350	Gastrointestinal toxicity with mucositis, nausea, vomiting, and diarrhea has been observed when
351	MYLERAN was used in association with bone marrow transplantation.
352	Oral $LD_{50}$ single doses in mice are 120 mg/kg. Two distinct types of toxic response are seen at
353	median lethal doses given intraperitoneally. Within a matter of hours there are signs of stimulation of
354	the central nervous system with convulsions and death on the first day. Mice are more sensitive to this
355	effect than are rats. With doses at the $\mathrm{LD}_{50}$ there is also delayed death due to damage to the bone
356	marrow. At 3 times the $LD_{50}$ , atrophy of the mucosa of the large intestine is found after a week,
357	whereas that of the small intestine is little affected. After doses in the order of 10 times those used
358	therapeutically were added to the diet of rats, irreversible cataracts were produced after several
359	weeks. Small doses had no such effect.
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DOSAGE AND ADMINISTRATION

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393	Store at 25°C (77°F); excursions permitted to 15° to 30°C (59° to 86°F) (see USP Controlled
394	Room Temperature).
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