
HIGHLIGHTS OF PRESCRIBING INFORMATION These highlights do not include all the information needed to use ARSENIC TRIOXIDE INJECTION safely and effectively. See full prescribing information for ARSENIC TRIOXIDE INJECTION.

ARSENIC TRIOXIDE injection, for intravenous use Initial U.S. Approval: 2000

WARNING: DIFFERENTIATION SYNDROME, CARDIAC CONDUCTION ABNORMALITIES, and ENCEPHALOPATHY INCLUDING WERNICKE'S

See full prescribing information for complete boxed warning.

- Patients with acute promyelocytic leukemia (APL) treated with arsenic trioxide have experienced symptoms of differentiation syndrome, which may be life-threatening or fatal. If differentiation syndrome is suspected, immediately initiate high-dose corticosteroids and hemodynamic monitoring until resolution. Temporarily withhold arsenic trioxide. (2.3, 5.1)
- Arsenic trioxide can cause QTc interval prolongation, complete atrioventricular block and torsade de pointes, which can be fatal. Before administering arsenic trioxide, assess the QTc interval, correct electrolyte abnormalities, and consider discontinuing drugs known to prolong QTc interval. Do not administer arsenic trioxide to patients with ventricular arrhythmia or prolonged QTc interval. Withhold arsenic trioxide until resolution and resume at reduced dose for QTc prolongation. (2.3, 5.2)
- Serious encephalopathy, including Wernicke's, has occurred with arsenic trioxide. If Wernicke's encephalopathy is suspected, immediately interrupt arsenic trioxide and initiate parenteral thiamine. Monitor until symptoms resolve or improve and thiamine levels normalize. (5.3)

INDICATIONS AND USAGE

Arsenic trioxide injection is an arsenical indicated:

• For induction of remission and consolidation in patients with APL who are refractory to, or have relapsed from, retinoid and anthracycline chemotherapy, and whose APL is characterized by the presence of the t(15;17) translocation or PML/RAR-alpha gene expression. (1.2)

DOSAGE AND ADMINISTRATION
Relapsed or refractory APL:
• Induction: Administer 0.15 mg/kg/day intravenously daily until bone marrow remission. Do not exceed

- Induction: Administer 0.15 mg/kg/day intravenously daily until bone marrow remission. Do not exceed 60 days. (2.2)
- *Consolidation:* Administer 0.15 mg/kg/day intravenously daily for 25 doses over a period up to 5 weeks. (2.2)

CONTRAINDICATIONS				
Hypersensitivity to arsenic. (4)				

------ WARNINGS AND PRECAUTIONS ------

- <u>Hepatotoxicity</u>: Monitor hepatic function tests at least twice weekly during induction and at least once weekly during consolidation. Withhold arsenic trioxide for certain elevations in AST, alkaline phosphatase and bilirubin and resume at reduced dose upon resolution. (2.3, 5.4)
- <u>Carcinogenesis</u>: Arsenic trioxide is a human carcinogen. Monitor patients for the development of second primary malignancies. (5.5)
- Embryo-Fetal Toxicity: Can cause fetal harm. Advise of potential risk to a fetus and use of effective

ADVERSE REACTIONS The most common adverse reactions (> 30%) are nausea, cough, fatigue, pyrexia, headache, abdominal pain, vomiting, tachycardia, diarrhea, dyspnea, hypokalemia, leukocytosis, hyperglycemia, hypomagnesemia, insomnia, dermatitis, edema, QTc prolongation, rigors, sore throat, arthralgia, paresthesia, and pruritus. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Amneal Pharmaceuticals at 1-877-835-5472 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

······ USE IN SPECIFIC POPULATIONS ······

- Lactation: Advise not to breastfeed. (8.2)
- <u>Renal Impairment:</u> Monitor patients with severe renal impairment (creatinine clearance less than 30 mL/min) for toxicity when treated with arsenic trioxide; dose reduction may be warranted. (8.6)
- <u>Hepatic Impairment</u>: Monitor patients with severe hepatic impairment (Child-Pugh Class C) for toxicity when treated with arsenic trioxide. (8.7)

See 17 for PATIENT COUNSELING INFORMATION.

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FULL PRESCRIBING INFORMATION

WARNING: DIFFERENTIATION SYNDROME, CARDIAC CONDUCTION ABNORMALITIES AND ENCEPHALOPATHY INCLUDING WERNICKE'S

Differentiation Syndrome: Patients with acute promyelocytic leukemia (APL) treated with arsenic trioxide have experienced differentiation syndrome, which may be life-threatening or fatal. Signs and symptoms may include unexplained fever, dyspnea, hypoxia, acute respiratory distress, pulmonary infiltrates, pleural or pericardial effusions, weight gain, peripheral edema, hypotension, renal insufficiency, hepatopathy, and multi-organ dysfunction, in the presence or absence of leukocytosis. If differentiation syndrome is suspected, immediately initiate high-dose corticosteroids and hemodynamic monitoring until resolution. Temporarily withhold arsenic trioxide [see Dosage and Administration (2.3), Warnings and Precautions (5.1)].

Cardiac Conduction Abnormalities: Arsenic trioxide can cause QTc interval prolongation, complete atrioventricular block and torsade de pointes, which can be fatal. Before administering arsenic trioxide, assess the QTc interval, correct electrolyte abnormalities, and consider discontinuing drugs known to prolong QTc interval. Do not administer arsenic trioxide to patients with a ventricular arrhythmia or prolonged QTc interval. Withhold arsenic trioxide until resolution and resume at reduced dose for QTc prolongation [see Dosage and Administration (2.3), Warnings and Precautions (5.2)].

Encephalopathy: Serious encephalopathy, including Wernicke's, has occurred with arsenic trioxide. Wernicke's is a neurologic emergency. Consider testing thiamine levels in patients at risk for thiamine deficiency. Administer parenteral thiamine in patients with or at risk for thiamine deficiency. Monitor patients for neurological symptoms and nutritional status while receiving arsenic trioxide. If Wernicke's encephalopathy is suspected, immediately interrupt arsenic trioxide and initiate parenteral thiamine. Monitor until symptoms resolve or improve and thiamine levels normalize [see Warnings and Precautions (5.3)].

1 INDICATIONS AND USAGE

1.2 Relapsed or Refractory APL

Arsenic trioxide injection is indicated for induction of remission and consolidation in patients with APL who are refractory to, or have relapsed from, retinoid and anthracycline chemotherapy, and whose APL is characterized by the presence of the t(15;17) translocation or PML/RAR-alpha gene expression.

2 DOSAGE AND ADMINISTRATION

2.2 Recommended Dosage for Relapsed or Refractory APL

A treatment course for patients with relapsed or refractory APL consists of 1 induction

cycle and 1 consolidation cycle [see Clinical Studies (14.2)].

- For the induction cycle, the recommended dosage of arsenic trioxide injection is 0.15 mg/kg/day intravenously daily until bone marrow remission or up to a maximum of 60 days.
- For the consolidation cycle, the recommended dosage of arsenic trioxide injection is 0.15 mg/kg/day intravenously daily for 25 doses over a period of up to 5 weeks. Begin consolidation 3 to 6 weeks after completion of induction cycle.

2.3 Monitoring and Dosage Modifications for Adverse Reactions

During induction, monitor coagulation studies, blood counts, and chemistries at least 2 to 3 times per week through recovery. During consolidation, monitor coagulation studies, blood counts, and chemistries at least weekly.

Table 2 shows the dosage modifications for adverse reactions due to arsenic trioxide injection when used alone.

Table 2: Dosage Modifications for Adverse Reactions of Arsenic Trioxide Injection

Adverse Reaction	Dosage Modification		
Differentiation syndrome, defined by the presence of 2 or more of the following: - Unexplained fever - Dyspnea - Pleural and/or pericardial effusion - Pulmonary infiltrates - Renal failure - Hypotension - Weight gain greater than 5 kg [see Warnings and Precautions (5.1)]	 Temporarily withhold arsenic trioxide injection. Administer dexamethasone 10 mg intravenously every 12 hours until the resolution of signs and symptoms for a minimum of 3 days. Resume treatment when the clinical condition improves and reduce the dose of the withheld drug(s) by 50%. Increase the dose of the withheld drug(s) to the recommended dosage after one week in the absence of recurrence of symptoms of differentiation syndrome. If symptoms re-appear, decrease arsenic trioxide injection to the previous dose. 		
QTc (Framingham formula) Prolongation greater than 450 msec for men or greater than 460 msec for women <i>[see Warnings and Precautions (5.2)]</i>	 Withhold arsenic trioxide injection and any medication known to prolong the QTc interval. Correct electrolyte abnormalities. After the QTc normalizes and electrolyte abnormalities are corrected, resume treatment with arsenic trioxide injection at a 50% reduced dose (0.075 mg/kg/day daily) for one week after resolution. If the 50% reduced dose is tolerated for one week (in the absence of QTc prolongation), increase the dose of arsenic trioxide injection to 0.11 mg/kg/day daily for 		

	 the next week. The dose of arsenic trioxide injection can be increased to 0.15 mg/kg/day in the absence of QTc prolongation during that 14-day dose-escalation period.
Hepatotoxicity, defined by 1 or more of the following: - Total bilirubin (TB) greater than 3 times the upper limit of normal (ULN) - Aspartate aminotransferase (AST) greater than 5 times the ULN - Alkaline phosphatase (AP) greater than 5 times the ULN <i>[see Warnings and</i> <i>Precautions (5.4)]</i>	 Withhold arsenic trioxide injection. Resume treatment at a 50% reduced dose of the withheld drug(s) when TB is less than 1.5 times the ULN and AP/AST are less than 3 times the ULN. Increase the dose of the withheld drug(s) back to the recommended dosage after one week on the reduced dose in the absence of worsening of hepatotoxicity. Discontinue the withheld drug(s) permanently if hepatotoxicity recurs.
Other severe or life-threatening (grade 3 to 4) nonhematologic reactions [see Adverse Reactions (6)]	 Temporarily withhold arsenic trioxide injection. When the adverse reaction resolves to no more than mild (grade 1), resume arsenic trioxide injection reduced by 2 dose levels (see Table 3 below).
Moderate (grade 2) nonhematologic reactions [see Adverse Reactions (6)]	 Reduce the dose of arsenic trioxide injection by 1 dose level (see Table 3 below).
Leukocytosis (WBC count greater than 10 Gi/L) [<i>see Adverse Reactions (6.1)</i>]	 Administer hydroxyurea. Hydroxyurea may be discontinued when the WBC declines below 10 Gi/L.
Myelosuppression, defined by 1 or more of the following: - absolute neutrophil count less than 1 Gi/L - platelets less than 50 Gi/L lasting more than 5 weeks <i>[see Adverse Reactions (6)]</i>	 Consider reducing the dose of arsenic trioxide injection by 1 dose level (see Table 3 below). If myelosuppression lasts ≥ 50 days or occurs on 2 consecutive cycles, assess a marrow aspirate for remission status. In the case of molecular remission, resume arsenic trioxide injection at 1 dose level lower (see Table 3 below).

Table 3: Dose Reduction Levels for Hematologic and Nonhematologic Toxicities

	Arsenic Trioxide Injection
Dose Level	mg/kg intravenously
	once daily

Starting level	0.15	
-1	0.11	
-2	0.10	
-3	0.075	

2.4 Preparation and Administration

Reconstitution

Dilute arsenic trioxide injection with 100 mL to 250 mL 5% Dextrose Injection, USP or 0.9% Sodium Chloride Injection, USP, using proper aseptic technique, immediately after withdrawal from the vial. Do not save any unused portions for later administration.

After dilution, store arsenic trioxide injection for no more than 24 hours at room temperature and 48 hours when refrigerated.

<u>Administration</u>

Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit.

Administer arsenic trioxide injection as an intravenous infusion over 2 hours. The infusion duration may be extended up to 4 hours if acute vasomotor reactions are observed. A central venous catheter is not required.

The arsenic trioxide injection vial is single-dose and does not contain any preservatives. Discard unused portions of each vial properly. Do not mix arsenic trioxide injection with other medications.

Safe Handling Procedures

Arsenic trioxide injection is a hazardous drug. Follow applicable special handling and disposal procedures¹.

3 DOSAGE FORMS AND STRENGTHS

Injection: 10 mg/10 mL (1 mg/mL) arsenic trioxide clear solution in a single-dose vial.

Injection: 12 mg/6 mL (2 mg/mL) arsenic trioxide clear solution in a single-dose vial.

4 CONTRAINDICATIONS

Arsenic trioxide injection is contraindicated in patients with hypersensitivity to arsenic.

5 WARNINGS AND PRECAUTIONS

5.1 Differentiation Syndrome

Differentiation syndrome, which may be life-threatening or fatal, has been observed in patients with acute promyelocytic leukemia (APL) treated with arsenic trioxide. In clinical trials, 16% to 23% of patients treated with arsenic trioxide for APL developed differentiation syndrome. Signs and symptoms include unexplained fever, dyspnea, hypoxia, acute respiratory distress, pulmonary infiltrates, pleural or pericardial effusion,

weight gain, peripheral edema, hypotension, renal insufficiency, hepatopathy and multiorgan dysfunction. Differentiation syndrome has been observed with and without concomitant leukocytosis, and it has occurred as early as day 1 of induction to as late as the second month induction therapy.

If differentiation syndrome is suspected, temporarily withhold arsenic trioxide and immediately initiate dexamethasone 10 mg intravenously every 12 hours and hemodynamic monitoring until resolution of signs and symptoms for a minimum of 3 days [see Dosage and Administration (2.3)].

5.2 Cardiac Conduction Abnormalities

Patients treated with arsenic trioxide can develop QTc prolongation, torsade de pointes, and complete atrioventricular block. In the clinical trial of patients with relapsed or refractory APL treated with arsenic trioxide monotherapy, 40% had at least one ECG tracing with a QTc interval greater than 500 msec. A prolonged QTc was observed between 1 and 5 weeks after start of arsenic trioxide infusion, and it usually resolved by 8 weeks after arsenic trioxide infusion. There are no data on the effect of arsenic trioxide on the QTc interval during the infusion of the drug.

The risk of torsade de pointes is related to the extent of QTc prolongation, concomitant administration of QTc prolonging drugs, a history of torsade de pointes, pre-existing QTc interval prolongation, congestive heart failure, administration of potassium-wasting diuretics, or other conditions that result in hypokalemia or hypomagnesemia. The risk may be increased when arsenic trioxide is co-administered with medications that can lead to electrolyte abnormalities (such as diuretics or amphotericin B) [see Drug Interactions (7)].

Prior to initiating therapy with arsenic trioxide, assess the QTc interval by electrocardiogram, correct pre-existing electrolyte abnormalities, and consider discontinuing drugs known to prolong QTc interval. Do not administer arsenic trioxide to patients with a ventricular arrhythmia or prolonged QTc. If possible, discontinue drugs that are known to prolong the QTc interval. If it is not possible to discontinue the interacting drug, perform cardiac monitoring frequently [see Drug Interactions (7)]. During arsenic trioxide therapy, maintain potassium concentrations above 4 mEq/L and magnesium concentrations above 1.8 mg/dL. Monitor ECG weekly and more frequently for clinically unstable patients.

For patients who develop a QTc Framingham greater than 450 msec for men or greater than 460 msec for women, withhold arsenic trioxide and any medication known to prolong the QTc interval. Correct electrolyte abnormalities. When the QTc normalizes and electrolyte abnormalities are corrected, resume arsenic trioxide at a reduced dose [see Dosage and Administration (2.3)].

5.3 Encephalopathy

Serious encephalopathies were reported in patients receiving arsenic trioxide. Monitor patients for neurological symptoms, such as confusion, decreased level of consciousness, seizures, cognitive deficits, ataxia, visual symptoms and ocular motor dysfunction. Advise patients and caregivers of the need for close observation.

Wernicke's Encephalopathy

Wernicke's encephalopathy occurred in patients receiving arsenic trioxide. Wernicke's

encephalopathy is a neurologic emergency that can be prevented and treated with thiamine. Consider testing thiamine levels in patients at risk for thiamine deficiency (e.g., chronic alcohol use, malabsorption, nutritional deficiency, concomitant use of furosemide). Administer parenteral thiamine in patients with or at risk for thiamine deficiency. Monitor patients for neurological symptoms and nutritional status while receiving arsenic trioxide. If Wernicke's encephalopathy is suspected, immediately interrupt arsenic trioxide and initiate parenteral thiamine. Monitor until symptoms resolve or improve and thiamine levels normalize.

5.4 Hepatotoxicity

Long-term liver abnormalities can occur in patients with APL treated with arsenic trioxide.

During treatment with arsenic trioxide, monitor hepatic function tests at least twice weekly during induction and at least once weekly during consolidation. Withhold arsenic trioxide if elevations in AST or alkaline phosphatase occur to greater than 5 times the upper limit of normal and/or elevation in serum total bilirubin occurs to greater than 3 times the upper limit of normal and resume at reduced dose upon resolution [see Dosage and Administration (2.3)].

5.5 Carcinogenesis

The active ingredient of arsenic trioxide injection, arsenic trioxide, is a human carcinogen. Monitor patients for the development of second primary malignancies.

5.6 Embryo-Fetal Toxicity

Arsenic trioxide can cause fetal harm when administered to a pregnant woman. Arsenic trioxide was embryolethal and teratogenic in rats when administered on gestation day 9 at a dose approximately 10 times the recommended human daily dose on a mg/m² basis. A related trivalent arsenic, sodium arsenite, produced teratogenicity when administered during gestation in mice at a dose approximately 5 times the projected human dose on a mg/m² basis and in hamsters at an intravenous dose approximately equivalent to the projected human daily dose on a mg/m² basis.

Advise pregnant women of the potential risk to a fetus. Advise females of reproductive potential to use effective contraception during treatment with arsenic trioxide and for 6 months after the last dose. Advise males with female partners of reproductive potential to use effective contraception during treatment with arsenic trioxide and for 3 months after the last dose [see Use in Specific Populations (8.1, 8.3)].

6 ADVERSE REACTIONS

The following clinically significant adverse reactions are described elsewhere in the labeling:

- Differentiation Syndrome [see Warnings and Precautions (5.1)]
- Cardiac Conduction Abnormalities [see Warnings and Precautions (5.2)]
- Encephalopathy [see Warnings and Precautions (5.3)]
- Hepatotoxicity [see Warnings and Precautions (5.4)]
- Carcinogenesis [see Warnings and Precautions (5.5)]

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

Relapsed or Refractory APL

Safety information was available for 52 patients with relapsed or refractory APL who participated in clinical trials of arsenic trioxide. Forty patients in the Study PLRXAS01 received the recommended dose of 0.15 mg/kg, of whom 28 completed both induction and consolidation cycles. An additional 12 patients with relapsed or refractory APL received doses generally similar to the recommended dose.

Serious adverse reactions observed in the 40 patients with refractory or relapsed APL enrolled in Study PLRXAS01 included differentiation syndrome (n=3), hyperleukocytosis (n=3), QTc interval \geq 500 msec (n=16, 1 with torsade de pointes), atrial dysrhythmias (n=2), and hyperglycemia (n=2).

The most common adverse reactions (> 30%) were nausea, cough, fatigue, pyrexia, headache, abdominal pain, vomiting, tachycardia, diarrhea, dyspnea, hypokalemia, leukocytosis, hyperglycemia, hypomagnesemia, insomnia, dermatitis, edema, QTc prolongation, rigors, sore throat, arthralgia, paresthesia, and pruritus.

Table 5 describes the adverse reactions in patients aged 5 to 73 years with APL who received arsenic trioxide at the recommended dose. Similar adverse reactions profiles were seen in the other patient populations who received arsenic trioxide.

Table 5: Adverse Reactions (≥ 5%) in Patients with Relapsed or Refractory APL Who Received Arsenic Trioxide in Study PLRXAS01

Body SystemAdverse	Any GradeAdverse Reactions		Grade ≥ 3Adverse Reactions		
reaction	n	%	n		%
Gastrointestinal	disorde	ers			
Nausea	30	75			
Abdominal pain (lower & upper)	23	58	4		10
Vomiting	23	58			
Diarrhea	21	53			
Sore throat	14	35			
Constipation	11	28	1	1	3
Anorexia	9	23			
Appetite decreased	6	15			
Loose stools	4	10			
Dyspepsia	4	10			
Oral blistering	3	8			
Fecal incontinence	3	8			
Gastrointestinal hemorrhage	3	8			

Dry mouth	3	8		
Abdominal	h	0		
tenderness	5	ð		
Diarrhea	5	0		
hemorrhagic	3	8		
Abdominal	_	0		
distension	3	Ø		
Respiratory	1			
Cough	26	65		
Dyspnea	21	53	4	10
Epistaxis	10	25		
Hypoxia	9	23	4	10
Pleural effusion	8	20	1	3
Post nasal drin	5	13	-	
Wheezing	5	13		
Decreased breath		1.5		
sounds	4	10		
Crenitations	4	10		
Rales	4	10		
Hemontysis	3	8		
Tachynnea	3	0 8		
Rhonchi	3	8		
General disorder	p reand a	o dminist	ration site condit	ions
Fatique	25	63	nation site condit	5
Pyrovia (fovor)	25	63	2	<u>5</u> Б
Edema - non-	2.5	05	۷	5
specific	16	40		
Rigors	15	38		
Chest nain	10	25	2	5
Injection site nain	8	20	۷	5
Pain - non-specific	6	20 15	1	3
Injection site	0	15	<u>ــــــــــــــــــــــــــــــــــــ</u>	5
arythoma	5	13		
Weight gain	5	13		
Injection site	5	13		
edema	4	10		
Weakness	4	10	2	5
Hemorrhage	<u>ד</u> ר	8	۷	5
Weight loss	3	8		
Drug	5	0		
hypersensitivity	2	5	1	3
Nervous system	disorde	ars		
Headache	24	60	1	3
Insomnia	17	43	<u>-</u> 1	<u>~</u>
Parosthosia	13	22	<u>-</u> 2	۲ ۲
nizzinese			۷	
(excluding vertige)	9	23		
Tremor	5	13		
		<u>1</u>		

Convulsion	3	8	2	5		
Somnolence	3	8				
Coma	2	5	2	5		
Cardiac disorders						
Tachycardia	22	55				
ECG OT corrected						
interval prolonged	16	40				
> 500 msec						
Palpitations	4	10				
ECG abnormal						
other than OT	~					
interval	3	8				
prolongation						
Metabolism and	nutritio	n disoro	ders			
Hypokalemia	20	50	5	13		
Hypomagnesemia	18	45	5	13		
Hyperalycemia	18	45	5	13		
	2		2 2	5		
Hyporkalomia	0	20 10	2	5		
	/ E	10	2			
ASTINCIERSED	C ₄	13	1	3		
Нуросаксетка	4	10				
Hypoglycemia	3	8				
Acidosis	2	5				
Hematologic diso	rders	1				
Leukocytosis	20	50	1	3		
Anemia	8	20	2	5		
Thrombocytopenia	7	18	5	13		
Febrile neutropenia	5	13	3	8		
Neutropenia	4	10	4	10		
Disseminated						
intravascular	3	8	3	8		
coagulation						
Lymphadenopathy	3	8				
Skin and subcuta	aneous	tissue (disorders			
Dermatitis	17	43				
Pruritus	13	33	1	3		
Ecchymosis	8	20				
Dry skin	6	15				
Erythema - non-	_					
specific	5	13				
Increased						
sweating	5	13				
Facial edema	3	8				
Night sweats	3	8				
Potochiao	ך א	8				
Hyperniamontation	ך ג	Q Q				
Non crocific claim	ر ا	υ				
	3	8				
lesions						

Urticaria	3	8				
Local exfoliation	2	5				
Evelid edema	2	5				
Musculoskeletal,	conne	ctive tis	sue, and bone di	sorders		
, Arthralgia	13	33	3	8		
Myalgia	10	25	2	5		
Bone pain	9	23	4	10		
Back pain	7	18	1	3		
Neck pain	5	13				
Pain in limb	5	13	2	5		
Psychiatric disor	ders					
Anxiety	12	30				
Depression	8	20				
Agitation	2	5				
Confusion	2	5				
Vascular disorde	rs					
Hypotension	10	25	2	5		
Flushing	4	10				
Hypertension	4	10				
Pallor	4	10				
Infections and in	festatio	ons				
Sinusitis	8	20				
Herpes simplex	5	13				
Upper respiratory	5	13	1	3		
tract infection	-		-			
Bacterial infection -	3	8	1	3		
non-specific	2	0				
Herpes zoster	3	8 r				
Nasopharyngitis	2	5 F				
Oral candidiasis	2	5 F	n	Г.		
Sepsis	<u>k</u> ama di	D Condona	Ζ	D		
	stem a	sorders				
Vayınai bomorrhado	5	13				
Intermenstrual						
hleeding	3	8				
Ocular disorders						
Eve irritation	4	10				
Blurred vision	4	10				
Drv eve	3	8				
Painful red eve	2	5				
Renal and urinary	v disoro	lers				
Renal failure	3	8	1	3		
Renal impairment	3	8				
Oliguria	2	5				
Incontinence	2	5				
Ear disorders						
Earache	3	8				

Tinnitus	2	5		
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Other Clinically Relevant Adverse Reactions

Leukocytosis

Arsenic trioxide can induce proliferation of leukemic promyelocytes resulting in a rapid increase in white blood cell count. Leukocytosis greater than 10 Gi/L developed during induction therapy in 50% of patients receiving arsenic trioxide monotherapy for relapsed/refractory APL. In the relapsed/refractory setting, a relationship did not exist between baseline WBC counts and development of hyperleukocytosis nor baseline WBC counts and peak WBC counts.

6.2 Postmarketing Experience

The following adverse reactions have been identified during postapproval use of arsenic trioxide. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate the frequency or establish a causal relationship to drug exposure.

Cardiac disorders: Ventricular extrasystoles in association with QT prolongation, ventricular tachycardia in association with QT prolongation, including torsade de pointes, atrioventricular block, and congestive heart failure

Ear and labyrinth disorders: Deafness

Hematologic disorders: Pancytopenia, bone marrow necrosis

Infections: Herpes zoster

Investigations: Gamma-glutamyltransferase increased

Musculoskeletal and connective tissue disorders: Bone pain, myalgia, rhabdomyolysis

Neoplasms benign, malignant and unspecified: Melanoma, pancreatic cancer, squamous cell carcinoma

Nervous system disorders: Peripheral neuropathy, paresis, seizures, confusion, encephalopathy, Wernicke's encephalopathy, posterior reversible encephalopathy syndrome

Skin and subcutaneous tissue disorders: Toxic epidermal necrolysis

7 DRUG INTERACTIONS

Drugs That Can Prolong the QT/QTc Interval

Concomitant use of these drugs and arsenic trioxide may increase the risk of serious QT/QTc interval prolongation [see Warnings and Precautions (5.1)]. Discontinue or replace with an alternative drug that does not prolong the QT/QTc interval while the patient is using arsenic trioxide. Monitor ECGs more frequently in patients when it is not feasible to avoid concomitant use.

Drugs That Can Lead to Electrolyte Abnormalities

Electrolyte abnormalities increase the risk of serious QT/QTc interval prolongation [see

Warnings and Precautions (5.1)]. Avoid concomitant use of drugs that can lead to electrolyte abnormalities. Monitor electrolytes more frequently in patients who must receive concomitant use of these drugs and arsenic trioxide.

Drugs That Can Lead to Hepatotoxicity

Use of these drugs and arsenic trioxide may increase the risk of serious hepatotoxicity *[see Warnings and Precautions (5.4)]*. Discontinue or replace with an alternative drug that does not cause hepatotoxicity while the patient is using arsenic trioxide. Monitor liver function tests more frequently in patients when it is not feasible to avoid concomitant use.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

<u>Risk Summary</u>

Based on the mechanism of action [see Clinical Pharmacology (12.1)] and findings in animal studies, arsenic trioxide can cause fetal harm when administered to a pregnant woman. Arsenic trioxide was embryolethal and teratogenic in rats when administered on gestation day 9 at a dose approximately 10 times the recommended human daily dose on a mg/m² basis (see Data). A related trivalent arsenic, sodium arsenite, produced teratogenicity when administered during gestation in mice at a dose approximately 5 times the projected human dose on a mg/m² basis and in hamsters at an intravenous dose approximately equivalent to the projected human daily dose on a mg/m² basis. There are no studies with the use of arsenic trioxide in pregnant women, and limited published data on arsenic trioxide use during pregnancy are insufficient to inform a drug-associated risk of major birth defects and miscarriage. Advise pregnant women of the potential risk to a fetus.

The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2% to 4% and 15% to 20%, respectively.

<u>Data</u>

Human Data

One patient was reported to deliver a live infant with no reported congenital anomalies after receiving arsenic trioxide during the first five months of pregnancy. A second patient became pregnant three months after discontinuing arsenic trioxide and was reported to have a normal pregnancy outcome. A third patient was a pregnant healthcare provider who experienced dermal contact with liquid arsenic trioxide and had a normal pregnancy outcome after treatment and monitoring. A fourth patient who became pregnant while receiving arsenic trioxide had a miscarriage.

Animal Data

Studies in pregnant mice, rats, hamsters, and primates have shown that inorganic arsenicals cross the placental barrier when given orally or by injection. An increase in resorptions, neural-tube defects, anophthalmia and microphthalmia were observed in

rats administered 10 mg/kg of arsenic trioxide on gestation day 9 (approximately 10 times the recommended human daily dose on a mg/m² basis). Similar findings occurred in mice administered a 10 mg/kg dose of a related trivalent arsenic, sodium arsenite (approximately 5 times the projected human dose on a mg/m² basis), on gestation days 6, 7, 8, or 9. Intravenous injection of 2 mg/kg sodium arsenite (approximately equivalent to the projected human daily dose on a mg/m² basis) on gestation day 7 (the lowest dose tested) resulted in neural-tube defects in hamsters.

8.2 Lactation

<u>Risk Summary</u>

Arsenic trioxide is excreted in human milk. There are no data on the effects of arsenic trioxide on the breastfed child or on milk production. Because of the potential for serious adverse reactions in a breastfed child, advise women not to breastfeed during treatment with arsenic trioxide and for 2 weeks after the final dose.

8.3 Females and Males of Reproductive Potential

Arsenic trioxide can cause fetal harm when administered to a pregnant woman [see Use in Specific Populations (8.1)].

Pregnancy Testing

Conduct pregnancy testing in females of reproductive potential prior to initiation of arsenic trioxide.

Contraception

Females

Advise females of reproductive potential to use effective contraception during treatment with arsenic trioxide and for 6 months after the final dose.

Males

Advise males with female partners of reproductive potential to use effective contraception during treatment with arsenic trioxide and for 3 months after the final dose.

Infertility

Males

Based on testicular toxicities including decreased testicular weight and impaired spermatogenesis observed in animal studies, arsenic trioxide may impair fertility in males of reproductive potential [see Nonclinical Toxicology (13.1)].

8.4 Pediatric Use

The safety and efficacy of arsenic trioxide as a single agent for treatment of pediatric patients with relapsed or refractory APL is supported by the pivotal phase 2 study in 40 patients with relapsed or refractory APL. Five patients below the age of 18 years (age range: 5 to 16 years) were treated with arsenic trioxide at the recommended dose of 0.15 mg/kg/day. A literature review included an additional 17 patients treated with arsenic trioxide for relapsed or refractory APL, with ages ranging from 4 to 21 years. No differences in efficacy and safety were observed by age.

8.5 Geriatric Use

Use of arsenic trioxide as monotherapy in patients with relapsed or refractory APL is supported by the open-label, single-arm trial that included 6 patients aged 65 and older (range: 65 to 73 years). A literature review included an additional 4 patients aged 69 to 72 years who were treated with arsenic trioxide for relapsed or refractory APL. No overall differences in safety or effectiveness were observed between these patients and younger patients.

8.6 Renal Impairment

Exposure of arsenic trioxide may be higher in patients with severe renal impairment [see Clinical Pharmacology (12.3)]. Monitor patients with severe renal impairment (creatinine clearance [CLcr] less than 30 mL/min) frequently for toxicity; a dose reduction may be warranted.

The use of arsenic trioxide in patients on dialysis has not been studied.

8.7 Hepatic Impairment

Since limited data are available across all hepatic impairment groups, caution is advised in the use of arsenic trioxide in patients with hepatic impairment [*see Clinical Pharmacology* (12.3)]. Monitor patients with severe hepatic impairment (Child-Pugh Class C) frequently for toxicity.

10 OVERDOSAGE

Manifestations

Manifestations of arsenic trioxide overdosage include convulsions, muscle weakness, and confusion.

<u>Management</u>

For symptoms of arsenic trioxide overdosage, immediately discontinue arsenic trioxide and consider chelation therapy.

A conventional protocol for acute arsenic intoxication includes dimercaprol administered at a dose of 3 mg/kg intramuscularly every 4 hours until immediate life-threatening toxicity has subsided. Thereafter, penicillamine at a dose of 250 mg orally, up to a maximum frequency of four times per day (\leq 1 g per day), may be given.

11 DESCRIPTION

Arsenic trioxide injection is a sterile injectable solution of arsenic trioxide. The molecular formula of arsenic trioxide in the solid state is As_2O_3 , with a molecular weight of 197.84 g/mol and the following structural formula:



Arsenic trioxide is a white to off-white powder. It is practically insoluble to sparingly soluble in water. It dissolves in solutions of alkali hydroxides (NaOH 1 M). It is practically insoluble in ethanol, chloroform and ethyl ether.

Arsenic trioxide injection is available in 10 mL or 6 mL, single-dose vials containing 10 mg or 12 mg of arsenic trioxide respectively. Arsenic trioxide injection is formulated as a sterile, nonpyrogenic, clear solution of arsenic trioxide in water for injection using sodium hydroxide and dilute hydrochloric acid to adjust to pH 8. Arsenic trioxide injection is preservative-free. Arsenic trioxide, the active ingredient, is present at a concentration of 1 mg/mL in 10 mL vial and 2 mg/mL in 6 mL vial. Inactive ingredients and their respective approximate concentrations are sodium hydroxide (1.2 mg/mL) for solubilization, and sodium hydroxide and hydrochloric acid for pH adjustment to pH 8.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

The mechanism of action of arsenic trioxide is not completely understood. Arsenic trioxide causes morphological changes and DNA fragmentation characteristic of apoptosis in NB4 human promyelocytic leukemia cells *in vitro*. Arsenic trioxide also causes damage or degradation of the fusion protein promyelocytic leukemia (PML)-retinoic acid receptor (RAR)-alpha.

12.2 Pharmacodynamics

Cardiac Electrophysiology

In a single-arm trial of arsenic trioxide (0.15 mg/kg daily), 16 of 40 patients (40%) had a QTc interval greater than 500 msec. Prolongation of the QTc was observed between 1 and 5 weeks after arsenic trioxide infusion, and then returned towards baseline by the end of 8 weeks after arsenic trioxide infusion.

12.3 Pharmacokinetics

The inorganic, lyophilized form of arsenic trioxide, when placed into solution, immediately forms the hydrolysis product arsenious acid (As^{III}). As^{III} is the pharmacologically active species of arsenic trioxide. Monomethylarsonic acid (MMA^V), and dimethylarsinic acid (DMA^V) are the main pentavalent metabolites formed during metabolism, in addition to arsenic acid (As^V) a product of As^{III} oxidation.

The pharmacokinetics of arsenical species ([As^{III}], [As^V], [MMA^V], [DMA^V]) were determined in 6 APL patients following once-daily doses of 0.15 mg/kg for 5 days per week. Over the total single-dose range of 7 mg to 32 mg (administered as 0.15 mg/kg),

systemic exposure (AUC) appears to be linear.

Peak plasma concentrations of arsenious acid (As^{III}), the primary active arsenical species were reached at the end of infusion (2 hours). Plasma concentration of As^{III} declined in a biphasic manner with a mean elimination half-life of 10 to 14 hours and is characterized by an initial rapid distribution phase followed by a slower terminal elimination phase. The daily exposure to As^{III} (mean AUC_{0-24h}) was 194 ng·hr/mL (n=5) on Day 1 of Cycle 1 and 332 ng·hr/mL (n=6) on Day 25 of Cycle 1, which represents an approximate 2-fold accumulation.

The primary pentavalent metabolites, MMA^V and DMA^V , are slow to appear in plasma (approximately 10 to 24 hours after first administration of arsenic trioxide), but, due to their longer half-life, accumulate more upon multiple dosing than does As^{III} . The mean estimated terminal elimination half-lives of the metabolites MMA^V and DMA^V are 32 hours and 72 hours, respectively. Approximate accumulation ranged from 1.4- to 8-fold following multiple dosing as compared to single-dose administration. As^V is present in plasma only at relatively low levels.

Distribution

The volume of distribution (V_{ss}) for As^{|||} is large (mean 562 L, N=10) indicating that As^{|||} is widely distributed throughout body tissues. V_{ss} is also dependent on body weight and increases as body weight increases.

<u>Elimination</u>

Metabolism

Much of the As^{III} is distributed to the tissues where it is methylated to the less cytotoxic metabolites, monomethylarsonic acid (MMA^V) and dimethylarsinic acid (DMA^V) by methyltransferases primarily in the liver. The metabolism of arsenic trioxide also involves oxidation of As^{III} to As^V, which may occur in numerous tissues via enzymatic or nonenzymatic processes. As^V is present in plasma only at relatively low levels following administration of arsenic trioxide.

Excretion

Approximately 15% of the administered arsenic trioxide dose is excreted in the urine as unchanged As^{III}. The methylated metabolites of As^{III} (MMA^V, DMA^V) are primarily excreted in the urine. The total clearance of As^{III} is 49 L/h and the renal clearance is 9 L/h. Clearance is not dependent on body weight or dose administered over the range of 7 mg to 32 mg.

Specific Populations

Patients with Renal Impairment

The effect of renal impairment on the pharmacokinetics of As^{III}, As^V, and the pentavalent metabolites MMA^V and DMA^V was evaluated in 20 patients with advanced malignancies. Patients were classified as having normal renal function (creatinine clearance [CLcr] > 80 mL/min, n=6), mild renal impairment (CLcr 50 to 80 mL/min, n=5), moderate renal impairment (CLcr 30 to 49 mL/min, n=6), or severe renal impairment (CLcr < 30 mL/min, n=3). Following twice-weekly administration of 0.15 mg/kg over a 2-hour infusion, the mean AUC_{0-INF} for As^{III} was comparable among the normal, mild and moderate renal impairment groups. However, in the **severe** renal impairment group, the mean AUC_{0-INF} for As^{III} was approximately 48% higher than that in the normal group.

Systemic exposure to MMA^V and DMA^V tended to be larger in patients with renal impairment; however, the clinical consequences of this increased exposure are not known. As^V plasma levels were generally below the limit of assay quantitation in patients with impaired renal function [see Use in Specific Populations (8.6)]. The use of arsenic trioxide in patients on dialysis has not been studied.

Patients with Hepatic Impairment

The effect of pharmacokinetics of As^{III}, As^V, and the pentavalent metabolites MMA^V and DMA^V was evaluated following administration of 0.25 to 0.50 mg/kg of arsenic trioxide in patients with hepatocellular carcinoma. Patients were classified as having normal hepatic function (n=4), mild hepatic impairment (Child-Pugh class A, n=12), moderate hepatic impairment (Child-Pugh class B, n=3), or severe hepatic impairment (Child-Pugh class C, n=1). No clear trend toward an increase in systemic exposure to As^{III}, As^V, MMA^V or DMA^V was observed with decreasing level of hepatic function as assessed by dosenormalized (per mg dose) AUC in the mild and moderate hepatic impairment groups. However, the one patient with severe hepatic impairment had mean dose-normalized AUC_{0-24h} and C_{max} values 40% and 70% higher, respectively, than those patients with normal hepatic function. The mean dose-normalized trough plasma levels for both MMA^V and DMA^V in this severely hepatically impaired patient were 2.2-fold and 4.7-fold higher, respectively, than those in the patients with normal hepatic function [see Use in Specific *Populations (8.7)*].

Pediatric Patients

Following intravenous administration of 0.15 mg/kg/day of arsenic trioxide in 10 APL patients (median age = 13.5 years, range 4 to 20 years), the daily exposure to As^{III} (mean AUC_{0-24h}) was 317 ng·hr/mL on Day 1 of Cycle 1 [see Use in Specific Populations (8.4)].

Drug Interaction Studies

No formal assessments of pharmacokinetic drug-drug interactions between arsenic trioxide and other drugs have been conducted. The methyltransferases responsible for metabolizing arsenic trioxide are not members of the cytochrome P450 family of isoenzymes. *In vitro* incubation of arsenic trioxide with human liver microsomes showed no inhibitory activity on substrates of the major cytochrome P450 (CYP) enzymes such as 1A2, 2A6, 2B6, 2C8, 2C9, 2C19, 2D6, 2E1, 3A4/5, and 4A9/11. The pharmacokinetics of drugs that are substrates for these CYP enzymes are not expected to be affected by concomitant treatment with arsenic trioxide.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenicity studies have not been conducted with arsenic trioxide [see Warnings and Precautions (5.6)].

Arsenic trioxide and trivalent arsenite salts have not been demonstrated to be mutagenic to bacteria, yeast, or mammalian cells. Arsenite salts are clastogenic *in vitro* (human fibroblast, human lymphocytes, Chinese hamster ovary cells, Chinese hamster V79 lung cells). Trivalent arsenic was genotoxic in the chromosome aberrations assay and micronucleus bone marrow assay in mice.

The effect of arsenic on fertility has not been adequately studied in humans. Decreased testicular weight and impaired spermatogenesis have been reported in animal studies. Male Wistar rat pups were administered 1.5 mg/kg sodium arsenite solution via the intraperitoneal route from postnatal days 1 to 14 and testes were collected for evaluation on postnatal days 15, 21, and 50. Results of this study revealed an altered morphology of the seminiferous tubules along with degeneration of spermatogenic cells, increased number of sperm with abnormal morphology, and decreased sperm counts. In beagle dogs administered intravenous arsenic trioxide for 90 days, reduced inner cell layers within seminiferous tubules and significantly decreased numbers of spermatozytes, spermatozoa, and sperm cells were observed at doses of 1 mg/kg/day and higher. The 1 mg/kg/day dose is approximately 3 times the recommended human daily dose on a mg/m² basis.

14 CLINICAL STUDIES

14.2 Relapsed or Refractory APL

Arsenic trioxide was investigated in Study PLRXAS01, an open-label, single-arm trial in 40 patients with relapsed or refractory APL who were previously treated with an anthracycline and a retinoid regimen. Patients received arsenic trioxide 0.15 mg/kg/day intravenously over 1 to 2 hours until the bone marrow was cleared of leukemic cells or for a maximum of 60 days. The CR (absence of visible leukemic cells in bone marrow and peripheral recovery of platelets and white blood cells with a confirmatory bone marrow \geq 30 days later) rate in this population of previously treated patients was 28 of 40 (70%). Among the 22 patients who had relapsed less than one year after treatment with tretinoin, there were 18 complete responders (82%). Of the 18 patients receiving arsenic trioxide \geq one year from tretinoin treatment, there were 10 complete responders (55%). The median time to bone marrow remission was 44 days and to onset of CR was 53 days. Three of 5 children, 5 years or older, achieved CR. No children less than 5 years old were treated.

Three to six weeks following bone marrow remission, 31 patients received consolidation therapy with arsenic trioxide, at the same dose, for 25 additional days over a period up to 5 weeks. In follow-up treatment, 18 patients received further arsenic trioxide as a maintenance course. Fifteen patients had bone marrow transplants. At last follow-up, 27 of 40 patients were alive with a median follow-up time of 484 days (range 280 to 755) and 23 of 40 patients remained in complete response with a median follow-up time of 483 days (range 280 to 755).

Cytogenetic conversion to no detection of the APL chromosome rearrangement was observed in 24 of 28 (86%) patients who met the response criteria defined above, in 5 of 5 (100%) patients who met some, but not all, of the response criteria, and 3 of 7 (43%) of patients who did not respond. RT-PCR conversions to no detection of the APL gene rearrangement were demonstrated in 22 of 28 (79%) of patients who met the response criteria, in 3 of 5 (60%) of patients who met some, but not all, of the response criteria, in 2 of 7 (29%) of patients who did not respond.

Responses were seen across all age groups tested, ranging from 6 to 72 years. The ability to achieve a CR was similar for both sexes. There were insufficient patients of Black, Hispanic, or Asian ancestry to estimate relative response rates in these groups,

but responses were seen in each group.

15 REFERENCES

1. "OSHA Hazardous Drugs." OSHA. http://www.osha.gov/SLTC/hazardousdrugs/index.html

16 HOW SUPPLIED/STORAGE AND HANDLING

How Supplied

Arsenic trioxide injection is supplied as a sterile, clear, colorless solution in single-dose vials.

It is available as follows:

10 mg/10 mL (1 mg/mL):

10 mL Single-Dose Vial:	NDC 70121-1483-1
10 Single-Dose Vials in 1 Carton:	NDC 70121-1483-7
<u>12 mg/6 mL (2 mg/mL):</u>	
6 mL Single-Dose Vial:	NDC 70121-1658-6
10 Single-Dose Vials in 1 Carton:	NDC 70121-1658-1

Storage and Handling

Store at 20° to 25°C (68° to 77°F); excursions permitted between 15° to 30°C (59° to 86°F) [see USP Controlled Room Temperature]. Do not freeze.

Arsenic trioxide injection is a hazardous drug. Follow applicable special handling and disposal procedures¹.

17 PATIENT COUNSELING INFORMATION

Differentiation Syndrome

Advise patients that symptoms of APL differentiation syndrome include fever, sudden weight gain, dizziness/lightheadedness, labored breathing, and accumulation of fluid in the lungs, heart, and chest. This syndrome is managed by immediate treatment with high-dose corticosteroids. Advise patients to immediately report any of these symptoms *[see Warnings and Precautions (5.1)]*.

Cardiac Conduction Abnormalities

Advise patients that arsenic trioxide may cause ECG abnormalities, including QT prolongation. If extreme, this prolongation has the potential to cause fainting, irregular heartbeat, or more serious side effects. Advise patients to immediately report any of these symptoms. Advise patients to provide a complete list of current medications as caution should be taken when arsenic trioxide is co-administered with other medications that can cause QT prolongation or lead to electrolyte abnormalities [see Warnings and Precautions (5.2) and Drug Interactions (7)].

Encephalopathy and Wernicke's Encephalopathy (WE)

Advise patients that symptoms of encephalopathies include neurological symptoms such as confusion, decreased level of consciousness, seizures, cognitive deficits, ataxia, visual symptoms and ocular motor dysfunction. Advise patients and caregivers to closely monitor for neurological symptoms and immediately report them to their healthcare provider [see Warnings and Precautions (5.3)].

Advise patients at risk for thiamine deficiency (e.g., chronic alcohol use, malabsorption, nutritional deficiency, concomitant use of furosemide) that Wernicke's encephalopathy is a neurologic emergency that can be prevented and treated with thiamine supplementation, and to immediately report any neurological symptoms to their healthcare provider [see Warnings and Precautions (5.3)].

Embryo-Fetal Toxicity

Advise pregnant women of the potential risk to a fetus. Advise females of reproductive potential to inform their healthcare provider of a known or suspected pregnancy [see Warnings and Precautions (5.5) and Use in Specific Populations (8.1)].

Advise females of reproductive potential to use effective contraception during treatment with arsenic trioxide and for 6 months after the last dose [see Use in Specific Populations (8.3)].

Advise males with female partners of reproductive potential to use effective contraception during treatment with arsenic trioxide for 3 months after the last dose *[see Use in Specific Populations (8.3)]*.

Lactation

Advise women not to breastfeed during treatment with arsenic trioxide and for 2 weeks after the last dose [see Use in Specific Populations (8.2)].

Infertility

Advise males of reproductive potential that arsenic trioxide may impair fertility [see Use in Specific Population (8.3)].

Other Adverse Reactions

Advise patients of the expected adverse reactions of arsenic trioxide. Most patients in clinical trials experienced some drug-related toxicity, most commonly leukocytosis, gastrointestinal symptoms (nausea, vomiting, diarrhea, and abdominal pain), fatigue, edema, hyperglycemia, dyspnea, cough, rash or itching, headaches, and dizziness. Advise patients to call their healthcare provider at the onset of any adverse reactions *[see Adverse Reactions (6.1)]*.

Manufactured by: **Amneal Oncology Pvt. Ltd.** Telangana 509301, INDIA

Distributed by: **Amneal Pharmaceuticals LLC** Bridgewater, NJ 08807

Rev. 07-2021-02

PRINCIPAL DISPLAY PANEL

NDC 70121-1483-1 Arsenic Trioxide Injection 10 mg/10 mL (1 mg/1 mL) Rx Only Vial Label Amneal Pharmaceuticals LL



NDC: 70121-1483-7 Arsenic Trioxide Injection 10 mg/10 mL (1 mg/1 mL) Rx Only Carton Label

Amneal Pharmaceuticals LLC



NDC 70121-1658-6 Arsenic Trioxide Injection 12 mg/6 mL (2 mg/1 mL) Rx Only Vial Label Amneal Pharmaceuticals LLC



NDC: 70121-1658-1 Arsenic Trioxide Injection 10 mg/10 mL (1 mg/1 mL) Rx Only Carton Label Amneal Pharmaceuticals LLC



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WATER (UNII: 059QF0K00R) Packaging Item Code Package Description Marketing Start Date Marketing End Date Item Schole Item Schole Item Schole Marketing End Date Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Schole Item Sch	SC	DIUM HYDRO	(IDE (UNII: 55X04	QC32I)					
Item Code Package Description Marketing Start Date Marketing End Date NDC:70121- 1483-7 10 in 1 CARTON 01/25/2021	W	ATER (UNII: 059	QF0KO0R)						
Package DescriptionMarketing Start DateMarketing End Date1NDC:70121- 1483-710 in 1 CARTON01/25/2021									
#Item CodePackage DescriptionMarketing Start DateMarketing End Date1NDC:70121- 1483-710 in 1 CARTON01/25/2021	Packaging								
NDC:70121- 1483-7 10 in 1 CARTON 01/25/2021	#	Item Code	Pa	ackage Description	M	ark	eting Start Date	Marketing End Date	
	1	NDC:70121- 1483-7	10 in 1 CARTON		01/2	25/2	2021		

1	NDC:70121- 1483-1	10 mL in 1 VIAL, SINGLE-DOSE; Type 0: Not a Combination Product							
M	Marketing Information								
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date					
AN	IDA	ANDA210739	01/25/2021						

ARSENIC TRIOXIDE

arsenic trioxide injection, solution

Product Information							
Product Type			HUMAN PRESCRIPTION DRUG	Item Code (Source)		NDC:70121-1658	
R	oute of Admir	nistration	INTRAVENOUS				
A	ctive Ingred	lient/Active	Moiety				
	Ingredient Name				Basis of Strength Stren		
ARSENIC TRIOXIDE (UNII: S7V92P67HO) (ARSENIC CATION (3+) - UNII:C96613F5AV) ARSENIC CATION (3+) -			ARSENIC TRIOXIDE 2 mg in 1 m		2 mg in 1 mL		
In	active Ingr	edients					
	Ingredient Name Strength						rength
H١	DROCHLORIC	ACID (UNII: QTT)	.7582CB)				
sc	DIUM HYDRO	(IDE (UNII: 55X04	QC32I)				
WATER (UNII: 059QF0KO0R)							
Pa	ackaging						
#	Item Code	Pa	ackage Description	Marketing Star Date		: Marketing End Date	
1	NDC:70121- 1658-1	10 in 1 CARTON		08/20/2021			
1	NDC:70121- 1658-6	6 mL in 1 VIAL, 9 Combination Pro	SINGLE-DOSE; Type 0: Not a oduct				
Marketing Information							
	Marketing	Applicat	tion Number or Monograph	Marke	ting Start	Mark	ceting End
	Category		Citation	00/20/20	Date		Date
AN	IDA	ANDA210739	9	08/20/20	21		

Revised: 7/2021

Amneal Pharmaceuticals LLC