1	4XPP		
2	45932C/Revised: December 2008		
3	TOBRAMYCIN		
4	FOR INJECTION, US	P	
5			
6	Rx only		
7			
8	This vial is intended i	for use by the hospital pharmacist in the extemporaneous	
9	preparation of IV sol	utions.	
10		PHARMACY BULK PACKAGE—	
11		NOT FOR DIRECT INFUSION	
12		NOT FOR DIRECT INFUSION	
13			
14	WARNINGS		
15	Patients treated with tobramycin injection and other aminoglycosides should be under		
16	close clinical observation, because these drugs have an inherent potential for causing		
17	ototoxicity and nephrotoxicity.		
18	Neurotoxicity, manifested as both auditory and vestibular ototoxicity, can occur.		
19	The auditory changes are irreversible, are usually bilateral, and may be partial or total.		
20	Eighth-nerve impairment and nephrotoxicity may develop, primarily in patients having		
21	preexisting renal dama	ge and in those with normal renal function to whom	
22	aminoglycosides are administered for longer periods or in higher doses than those		
23	recommended. Other manifestations of neurotoxicity may include numbness, skin		

tingling, muscle twitching, and convulsions. The risk of aminoglycoside-induced hearing loss increases with the degree of exposure to either high peak or high trough serum concentrations. Patients who develop cochlear damage may not have symptoms during therapy to warn them of eighth-nerve toxicity, and partial or total irreversible bilateral deafness may continue to develop after the drug has been discontinued.

Rarely, nephrotoxicity may not become apparent until the first few days after cessation of therapy. Aminoglycoside-induced nephrotoxicity usually is reversible.

Renal and eighth-nerve function should be closely monitored in patients with known or suspected renal impairment and also in those whose renal function is initially normal but who develop signs of renal dysfunction during therapy. Peak and trough serum concentrations of aminoglycosides should be monitored periodically during therapy to assure adequate levels and to avoid potentially toxic levels. Prolonged serum concentrations above 12 mcg/mL should be avoided. Rising trough levels (above 2 mcg/mL) may indicate tissue accumulation. Such accumulation, excessive peak concentrations, advanced age, and cumulative dose may contribute to ototoxicity and nephrotoxicity (see PRECAUTIONS). Urine should be examined for decreased specific gravity and increased excretion of protein, cells, and casts. Blood urea nitrogen, serum creatinine, and creatinine clearance should be measured periodically. When feasible, it is recommended that serial audiograms be obtained in patients old enough to be tested, particularly high-risk patients. Evidence of impairment of renal, vestibular, or auditory function requires discontinuation of the drug or dosage adjustment.

Tobramycin should be used with caution in premature and neonatal infants because of their renal immaturity and the resulting prolongation of serum half-life of the drug.

Concurrent and sequential use of other neurotoxic and/or nephrotoxic antibiotics, particularly other aminoglycosides (e.g., amikacin, streptomycin, neomycin, kanamycin, gentamicin, and paromomycin), cephaloridine, viomycin, polymyxin B, colistin, cisplatin, and vancomycin, should be avoided. Other factors that may increase patient risk are advanced age and dehydration.

Aminoglycosides should not be given concurrently with potent diuretics, such as ethacrynic acid and furosemide. Some diuretics themselves cause ototoxicity, and intravenously administered diuretics enhance aminoglycoside toxicity by altering antibiotic concentrations in serum and tissue.

Aminoglycosides can cause fetal harm when administered to a pregnant woman (see **PRECAUTIONS**).

DESCRIPTION:

Tobramycin sulfate, a water-soluble antibiotic of the aminoglycoside group, is derived from the actinomycete *Streptomyces tenebrarius*. Tobramycin for Injection, USP is supplied as a sterile powder and is intended for reconstitution with 30 mL of Sterile Water for Injection, USP. Each vial contains 1,200 mg of tobramycin activity. After dilution, the solution will contain 40 mg of tobramycin per mL. The product contains no preservative or sodium bisulfite.

Tobramycin sulfate is O-3-amino-3-deoxy- α -D-glucopyranosyl- $(1\rightarrow 4)$ -O-[2,6-

2 diamino-2,3,6-trideoxy- α -D-*ribo*-hexopyranosyl- $(1\rightarrow 6)$]-2-deoxy-L-streptamine, sulfate

3 (2:5)(salt) and has the molecular formula $(C_{18}H_{37}N_5O_9)_2 \bullet 5H_2SO_4$. The molecular weight

4 is 1425.45.

5 The structural formula for tobramycin is as follows:

The pharmacy bulk package of tobramycin is a container of a sterile preparation for parenteral use that contains multiple single doses. It is intended for use in a pharmacy admixture program. Package use is restricted to the preparation of admixtures for intravenous infusion or to the filling of empty sterile syringes for intravenous injection for patients with individualized dosing requirements.

CLINICAL PHARMACOLOGY:

Tobramycin is rapidly absorbed following intramuscular administration. Peak serum concentrations of tobramycin occur between 30 and 90 minutes after intramuscular administration. Following an intramuscular dose of 1 mg/kg of body weight, maximum serum concentrations reach about 4 mcg/mL, and measurable levels persist for as long as 8 hours. Therapeutic serum levels are generally considered to range from 4 to 6 mcg/mL.

- 1 When tobramycin is administered by intravenous infusion over a 1-hour period, the
- 2 serum concentrations are similar to those obtained by intramuscular administration.
- 3 Tobramycin is poorly absorbed from the gastrointestinal tract.
- 4 In patients with normal renal function, except neonates, tobramycin administered
- 5 every 8 hours does not accumulate in the serum. However, in those patients with reduced
- 6 renal function and in neonates, the serum concentration of the antibiotic is usually higher
- 7 and can be measured for longer periods of time than in normal adults. Dosage for such
- 8 patients must, therefore, be adjusted accordingly (see **DOSAGE AND**

9 **ADMINISTRATION**).

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occurs, and tobramycin is eliminated almost exclusively by glomerular filtration. Renal clearance is similar to that of endogenous creatinine. Ultrafiltration studies demonstrate that practically no serum protein binding occurs. In patients with normal renal

Following parenteral administration, little, if any, metabolic transformation

- function, up to 84% of the dose is recoverable from the urine in 8 hours and up to 93% in 24 hours.
 - Peak urine concentrations ranging from 75 to 100 mcg/mL have been observed following the intramuscular injection of a single dose of 1 mg/kg. After several days of treatment, the amount of tobramycin excreted in the urine approaches the daily dose administered. When renal function is impaired, excretion of tobramycin is slowed, and
- The serum half-life in normal individuals is 2 hours. An inverse relationship
 exists between serum half-life and creatinine clearance, and the dosage schedule should

accumulation of the drug may cause toxic blood levels.

1	be adjusted according to the degree of renal impairment (see DOSAGE AND		
2	ADMINISTRATION). In patients undergoing dialysis, 25% to 70% of the administered		
3	dose may be removed, depending on the duration and type of dialysis.		
4	Tobramycin can be detected in tissues and body fluids after parenteral		
5	administration. Concentrations in bile and stools ordinarily have been low, which		
6	suggests minimum biliary excretion. Tobramycin has appeared in low concentration in		
7	the cerebrospinal fluid following parenteral administration, and concentrations		
8	are dependent on dose, rate of penetration, and degree of meningeal inflammation. It has		
9	also been found in sputum, peritoneal fluid, synovial fluid, and abscess fluids,		
10	and it crosses the placental membranes. Concentrations in the renal cortex are several		
11	times higher than the usual serum levels.		
12	Probenecid does not affect the renal tubular transport of tobramycin.		
13			
14	Microbiology		
15	Tobramycin acts by inhibiting synthesis of protein in bacterial cells. In vitro tests		
16	demonstrate that tobramycin is bactericidal.		
17	Tobramycin has been shown to be active against most strains of the following		
18	organisms both in vitro and in clinical infections as described in INDICATIONS AND		
19	USAGE section:		
20	Aerobic Gram-positive microorganisms		
21	Staphylococcus aureus		
22	Aerobic Gram-negative microorganisms		
23	Citrobacter species		

1	Enterobacter species
2	Escherichia coli
3	Klebsiella species
4	Morganella morganii
5	Pseudomonas aeruginosa
6	Proteus mirabilis
7	Proteus vulgaris
8	Providencia species
9	Serratia species
10	
11	Aminoglycosides have a low order of activity against most gram-positive
12	organisms, including Streptococcus pyogenes, Streptococcus pneumoniae, and
13	enterococci.
14	Although most strains of enterococci demonstrate in vitro resistance, some strains
15	in this group are susceptible. In vitro studies have shown that an aminoglycoside
16	combined with an antibiotic that interferes with cell-wall synthesis affects some
17	enterococcal strains synergistically. The combination of penicillin G and tobramycin
18	results in a synergistic bactericidal effect in vitro against certain strains of Enterococcus
19	faecalis. However, this combination is not synergistic against other closely related
20	organisms, e.g., Enterococcus faecium. Speciation of enterococci alone cannot be used to
21	predict susceptibility. Susceptibility testing and tests for antibiotic synergism are
22	emphasized.
23	Cross-resistance between aminoglycosides may occur.

Susceptibility Tests

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Diffusion	Technio	ues

- 3 Quantitative methods that require measurement of zone diameters give the most precise
- 4 estimates of susceptibility of bacteria to antimicrobial agents. One such procedure
- 5 is the National Committee for Clinical Laboratory Standards (NCCLS)-approved
- 6 procedure. This method has been recommended for use with disks to test susceptibility
- 7 to tobramycin. Interpretation involves correlation of the diameters obtained in the disk
- 8 test with minimum inhibitory concentrations (MIC) for tobramycin.
- 9 Reports from the laboratory giving results of the standard single-disk
- susceptibility test with a 10 mcg tobramycin disk should be interpreted according to the
- 11 following criteria:

Zone Diameter (mm)	<u>Interpretation</u>
≥15	(S) Susceptible
13 to 14	(I) Intermediate
≤ 12	(R) Resistant

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generally achievable blood levels. A report of "Intermediate" suggests that the organism would be susceptible if high dosage is used or if the infection is confined to tissues and fluids in which high antimicrobial levels are obtained. A report of "Resistant" indicates that achievable concentrations are unlikely to be inhibitory and other therapy

A report of "Susceptible" indicates that the pathogen is likely to be inhibited by

should be selected.

Standardized procedures require the use of laboratory control organisms. The 10 mcg tobramycin disk should give the following zone diameters:

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1	Zone Organism Diameter (mm) E. coli ATCC 25922 18 to 26 P. aeruginosa ATCC 27853 19 to 25 S. aureus ATCC 25923 19 to 29			
2				
3	Dilution Techniques			
4	Broth and agar dilution methods, such as those recommended by the NCCLS ² , may be			
5	used to determine MICs of tobramycin. MIC test results should be interpreted			
6	according to the following criteria:			
7	MIC (mcg/mL) Interpretation ≤ 4 (S) Susceptible 8 (I) Intermediate ≥ 16 (R) Resistant			
8				
9				
10	laboratory control organisms. Tobramycin laboratory reagent should give the following			
11	MIC values:			
12 13	Organism (mcg/mL) E. faecalis ATCC 29212 8 to 32 E. coli ATCC 25922 0.25 to 1 P. aeruginosa ATCC 27853 0.25 to 1 S. aureus ATCC 29213 0.12 to 1			
14	INDICATIONS AND USAGE:			
15	Tobramycin for Injection, USP is indicated for the treatment of serious bacterial			
16	infections caused by susceptible strains of the designated microorganisms in the			
17	diseases listed below:			
18	Septicemia in the pediatric patient and adult caused by <i>P. aeruginosa</i> , <i>E. coli</i> , and			
19	Klebsiella spp.			

1 Lower respiratory tract infections caused by *P. aeruginosa*, *Klebsiella spp*, 2 Enterobacter spp, Serratia spp, E. coli, and S. aureus (penicillinase- and non-3 penicillinase-producing strains). 4 Serious central-nervous-system infections (meningitis) caused by susceptible 5 organisms. 6 Intra-abdominal infections, including peritonitis, caused by E. coli, Klebsiella spp, 7 and Enterobacter spp. 8 Skin, bone, and skin structure infections caused by P. aeruginosa, Proteus spp, 9 E. coli, Klebsiella spp, Enterobacter spp, and S. aureus. 10 Complicated and recurrent urinary tract infections caused by *P. aeruginosa*, 11 Proteus spp, (indole-positive and indole-negative), E. coli, Klebsiella spp, Enterobacter 12 spp, Serratia spp, S. aureus, Providencia spp, and Citrobacter spp. 13 Aminoglycosides, including tobramycin sulfate injection, USP are not indicated 14 in uncomplicated initial episodes of urinary tract infections unless the causative 15 organisms are not susceptible to antibiotics having less potential toxicity. Tobramycin for 16 Injection, USP may be considered in serious staphylococcal infections when penicillin or 17 other potentially less toxic drugs are contraindicated and when bacterial susceptibility 18 testing and clinical judgment indicate its use. 19 Bacterial cultures should be obtained prior to and during treatment to isolate and 20 identify etiologic organisms and to test their susceptibility to tobramycin. If susceptibility 21 tests show that the causative organisms are resistant to tobramycin, other appropriate 22 therapy should be instituted. In patients in whom a serious life-threatening

1 gram-negative infection is suspected, including those in whom concurrent therapy with a 2 penicillin or cephalosporin and an aminoglycoside may be indicated, treatment with 3 tobramycin may be initiated before the results of susceptibility studies are obtained. The 4 decision to continue therapy with tobramycin should be based on the results of 5 susceptibility studies, the severity of the infection, and the important additional concepts 6 discussed in the **WARNINGS** box above. 7 8 **CONTRAINDICATIONS:** 9 A hypersensitivity to any aminoglycoside is a contraindication to the use of tobramycin. 10 A history of hypersensitivity or serious toxic reactions to aminoglycosides may also 11 contraindicate the use of any other aminoglycoside because of the known cross-12 sensitivity of patients to drugs in this class. 13 14 **WARNINGS:** 15 See WARNINGS box above. 16 Serious allergic reactions including anaphylaxis and dermatologic reactions 17 including exfoliative dermatitis, toxic epidermal necrolysis, erythema multiforme, and 18 Stevens-Johnson Syndrome have been reported rarely in patients on tobramycin therapy. 19 Although rare, fatalities have been reported (see **CONTRAINDICATIONS**). 20 If an allergic reaction occurs, the drug should be discontinued and appropriate 21 therapy instituted. 22 Clostridium difficile associated diarrhea (CDAD) has been reported with use of 23 nearly all antibacterial agents, including Tobramycin for Injection, USP, and may range

1 in severity from mild diarrhea to fatal colitis. Treatment with antibacterial agents alters 2 the normal flora of the colon leading to overgrowth of *C. difficile*. 3 C. difficile produces toxins A and B which contribute to the development of 4 CDAD. Hypertoxin producing strains of C. difficile cause increased morbidity and 5 mortality, as these infections can be refractory to antimicrobial therapy and may require 6 colectomy. CDAD must be considered in all patients who present with diarrhea following 7 antibiotic use. Careful medical history is necessary since CDAD has been reported to 8 occur over two months after the administration of antibacterial agents. 9 If CDAD is suspected or confirmed, ongoing antibiotic use not directed against 10 C. difficile may need to be discontinued. Appropriate fluid and electrolyte management, 11 protein supplementation, antibiotic treatment of C. difficile, and surgical evaluation 12 should be instituted as clinically indicated. 13 14 **PRECAUTIONS:** 15 Serum and urine specimens for examination should be collected during therapy, as 16 recommended in the WARNINGS box. Serum calcium, magnesium, and sodium 17 should be monitored. 18 Peak and trough serum levels should be measured periodically during therapy. 19 Prolonged concentrations above 12 mcg/mL should be avoided. Rising trough levels 20 (above 2 mcg/mL) may indicate tissue accumulation. Such accumulation, advanced age, 21 and cumulative dosage may contribute to ototoxicity and nephrotoxicity. It is particularly 22 important to monitor serum levels closely in patients with known renal impairment.

A useful guideline would be to perform serum level assays after 2 or 3 doses, so
that the dosage could be adjusted if necessary, and at 3- to 4-day intervals during
therapy. In the event of changing renal function, more frequent serum levels should be
obtained and the dosage or dosage interval adjusted according to the guidelines
provided in **DOSAGE AND ADMINISTRATION**.

In order to measure the peak level, a serum sample should be drawn about 30

minutes following intravenous infusion or 1 hour after an intramuscular injection. Trough levels are measured by obtaining serum samples at 8 hours or just prior to the next dose of tobramycin. These suggested time intervals are intended only as guidelines and may vary according to institutional practices. It is important, however, that there be consistency within the individual patient program unless computerized pharmacokinetic dosing programs are available in the institution. These serum-level assays may be especially useful for monitoring the treatment of severely ill patients with changing renal function or of those infected with less susceptible organisms or those receiving maximum dosage.

Neuromuscular blockade and respiratory paralysis have been reported in cats receiving very high doses of tobramycin (40 mg/kg). The possibility of prolonged or secondary apnea should be considered if tobramycin is administered to anesthetized patients who are also receiving neuromuscular blocking agents, such as succinylcholine, tubocurarine, or decamethonium, or to patients receiving massive transfusions of citrated blood. If neuromuscular blockade occurs, it may be reversed by the administration of calcium salts.

Cross-allergenicity among aminoglycosides has been demonstrated.

1	In patients with extensive burns or cystic fibrosis, altered pharmacokinetics may
2	result in reduced serum concentrations of aminoglycosides. In such patients treated with
3	tobramycin, measurement of serum concentration is especially important as a basis for
4	determination of appropriate dosage.
5	Elderly patients may have reduced renal function that may not be evident in the
6	results of routine screening tests, such as BUN or serum creatinine. A creatinine
7	clearance determination may be more useful. Monitoring of renal function during
8	treatment with aminoglycosides is particularly important in such patients.
9	An increased incidence of nephrotoxicity has been reported following
10	concomitant administration of aminoglycoside antibiotics and cephalosporins.
11	Aminoglycosides should be used with caution in patients with muscular disorders,
12	such as myasthenia gravis or parkinsonism, since these drugs may aggravate muscle
13	weakness because of their potential curare-like effect on neuromuscular function.
14	Aminoglycosides may be absorbed in significant quantities from body surfaces
15	after local irrigation or application and may cause neurotoxicity and nephrotoxicity.
16	Aminoglycosides have not been approved for intraocular and/or subconjunctival
17	use. Physicians are advised that macular necrosis has been reported following
18	administration of aminoglycosides, including tobramycin, by these routes.
19	See WARNINGS box regarding concurrent use of potent diuretics and concurrent
20	and sequential use of other neurotoxic or nephrotoxic drugs.
21	The inactivation of tobramycin and other aminoglycosides by β-lactam-type
22	antibiotics (penicillins or cephalosporins) has been demonstrated in vitro and in

1 patients with severe renal impairment. Such inactivation has not been found in patients 2 with normal renal function who have been given the drugs by separate routes 3 of administration. 4 Therapy with tobramycin may result in overgrowth of nonsusceptible organisms. 5 If overgrowth of nonsusceptible organisms occurs, appropriate therapy should 6 be initiated. 7 8 Information for Patients 9 Diarrhea is a common problem caused by antibiotics which usually ends when the antibiotic is discontinued. Sometimes after starting treatment with antibiotics, patients 10 11 can develop watery and bloody stools (with or without stomach cramps and fever) even 12 as late as two or more months after having taken the last dose of the antibiotic. If this 13 occurs, patients should contact their physician as soon as possible. 14 15 Pregnancy Category D 16 Aminoglycosides can cause fetal harm when administered to a pregnant woman. 17 Aminoglycoside antibiotics cross the placenta, and there have been several reports of 18 total irreversible bilateral congenital deafness in children whose mothers received

tobramycin, she should be apprised of the potential hazard to the fetus.

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streptomycin during pregnancy. Serious side effects to mother, fetus, or newborn have

not been reported in the treatment of pregnant women with other aminoglycosides. If

tobramycin is used during pregnancy or if the patient becomes pregnant while taking

1	Pediatric Use
2	See INDICATIONS AND USAGE and DOSAGE AND ADMINISTRATION.
3	
4	Geriatric Use
5	Elderly patients may be at a higher risk of developing nephrotoxicity and ototoxicity
6	while receiving tobramycin (see WARNINGS, PRECAUTIONS, and
7	OVERDOSAGE). Other factors that may contribute to nephrotoxicity and ototoxicity
8	are rising trough levels, excessive peak concentrations, dehydration, concomitant
9	use of other neurotoxic or nephrotoxic drugs, and cumulative dose. Peak and trough
10	serum levels should be measured periodically during therapy to assure adequate
11	levels and to avoid potentially toxic levels (see WARNINGS and PRECAUTIONS).
12	Tobramycin is known to be substantially excreted by the kidney, and the risk of
13	toxic reactions to this drug may be greater in patients with impaired renal function. Dose
14	reduction is required for patients with impaired renal function (see DOSAGE AND
15	ADMINISTRATION). Elderly patients may have reduced renal function that may not
16	be evident in the results of routine screening tests, such as BUN or serum creatinine. A
17	creatinine clearance determination may be more useful. Monitoring of renal function
18	during treatment with aminoglycosides is particularly important in the elderly (see
19	PRECAUTIONS).
20	This product does not contain sodium.
21	
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23	

ADVERSE REACTIONS:

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1 2 **Neurotoxicity** 3 Adverse effects on both the vestibular and auditory branches of the eighth nerve have 4 been noted, especially in patients receiving high doses or prolonged therapy, in 5 those given previous courses of therapy with an ototoxin, and in cases of dehydration. 6 Symptoms include dizziness, vertigo, tinnitus, roaring in the ears, and hearing loss. 7 Hearing loss is usually irreversible and is manifested initially by diminution of high-tone 8 acuity. Tobramycin and gentamicin sulfates closely parallel each other in regard to 9 ototoxic potential. 10 11 **Nephrotoxicity** 12 Renal function changes, as shown by rising BUN, NPN, and serum creatinine and by 13 oliguria, cylindruria, and increased proteinuria, have been reported, especially in 14 patients with a history of renal impairment who are treated for longer periods or with 15 higher doses than those recommended. Adverse renal effects can occur in patients 16 with initially normal renal function. 17 Clinical studies and studies in experimental animals have been conducted to 18 compare the nephrotoxic potential of tobramycin and gentamicin. In some of the clinical 19 studies and in the animal studies, tobramycin caused nephrotoxicity significantly less 20 frequently than gentamicin. In some other clinical studies, no significant difference in the 21 incidence of nephrotoxicity between tobramycin and gentamicin was found. 22 Other reported adverse reactions possibly related to tobramycin include anemia,

granulocytopenia, and thrombocytopenia; and fever, rash, exfoliative dermatitis,

1 itching, urticaria, nausea, vomiting, diarrhea, headache, lethargy, pain at the injection

site, mental confusion, and disorientation. Laboratory abnormalities possibly related to

tobramycin include increased serum transaminases (AST, ALT); increased serum LDH

and bilirubin; decreased serum calcium, magnesium, sodium, and potassium; and

5 leukopenia, leukocytosis, and eosinophilia.

OVERDOSAGE:

Signs and Symptoms

The severity of the signs and symptoms following a tobramycin overdose are dependent on the dose administered, the patient's renal function, state of hydration, and age and whether or not other medications with similar toxicities are being administered concurrently. Toxicity may occur in patients treated more than 10 days, in adults given more than 5 mg/kg/day, in pediatric patients given more than 7.5 mg/kg/day, or in patients with reduced renal function where dose has not been appropriately adjusted.

Nephrotoxicity following the parenteral administration of an aminoglycoside is most closely related to the area under the curve of the serum concentration versus time graph. Nephrotoxicity is more likely if trough blood concentrations fail to fall below 2 mcg/mL and is also proportional to the average blood concentration. Patients who are elderly, have abnormal renal function, are receiving other nephrotoxic drugs, or are volume depleted are at greater risk for developing acute tubular necrosis. Auditory and vestibular toxicities have been associated with aminoglycoside overdose. These toxicities occur in patients treated longer than 10 days, in patients with abnormal renal function, in dehydrated patients, or in patients receiving medications with additive

- 1 auditory toxicities. These patients may not have signs or symptoms or may experience
- 2 dizziness, tinnitus, vertigo, and a loss of high-tone acuity as ototoxicity progresses.
- 3 Ototoxicity signs and symptoms may not begin to occur until long after the drug has been
- 4 discontinued.
- 5 Neuromuscular blockade or respiratory paralysis may occur following
- 6 administration of aminoglycosides. Neuromuscular blockade, respiratory failure, and
- 7 prolonged respiratory paralysis may occur more commonly in patients with myasthenia
- 8 gravis or Parkinson's disease. Prolonged respiratory paralysis may also occur in patients
- 9 receiving decamethonium, tubocurarine, or succinylcholine. If neuromuscular blockade
- occurs, it may be reversed by the administration of calcium salts but mechanical
- 11 assistance may be necessary.
- 12 If tobramycin were ingested, toxicity would be less likely because
- aminoglycosides are poorly absorbed from an intact gastrointestinal tract.

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Treatment

- 16 In all cases of suspected overdosage, call your Regional Poison Control Center to obtain
- 17 the most up-to-date information about the treatment of overdose. This recommendation
- is made because, in general, information regarding the treatment of overdosage may
- 19 change more rapidly than the package insert. In managing overdosage, consider the
- 20 possibility of multiple drug overdoses, interaction among drugs, and unusual drug
- 21 kinetics in your patient.
- The initial intervention in a tobramycin overdose is to establish an airway and
- 23 ensure oxygenation and ventilation. Resuscitative measures should be initiated

1	promptly if respiratory paralysis occurs.
2	Patients who have received an overdose of tobramycin and who have normal renal
3	function should be adequately hydrated to maintain a urine output of 3 to 5 mL/kg/hr.
4	Fluid balance, creatinine clearance, and tobramycin plasma levels should be carefully
5	monitored until the serum tobramycin level falls below 2 mcg/mL.
6	Patients in whom the elimination half-life is greater than 2 hours or whose renal
7	function is abnormal may require more aggressive therapy. In such patients, hemodialysis
8	may be beneficial.
9	
10	DOSAGE AND ADMINISTRATION:
11	The patient's pretreatment body weight should be obtained for calculation of correct
12	dosage. It is desirable to measure both peak and trough serum concentrations
13	(see WARNINGS box and PRECAUTIONS).
14	
15	Administration for Patients with Normal Renal Function
16	Adults with Serious Infections
17	3 mg/kg/day in 3 equal doses every 8 hours (see Table 1).
18	
19	Adults with Life-Threatening Infections
20	Up to 5 mg/kg/day may be administered in 3 or 4 equal doses (see Table 1). The dosage
21	should be reduced to 3 mg/kg/day as soon as clinically indicated. To prevent increased
22	toxicity due to excessive blood levels, dosage should not exceed 5 mg/kg/day unless
23	serum levels are monitored (see WARNINGSbox and PRECAUTIONS).

Table 1 Dosage schedule guide for adults with Normal Renal Function (Dosage at 8-Hour Intervals)

For		Usual Dose for Serious Infections	
. Weig	lent phing lb	1 mg/kg g8h	
kg	lb	(Total, 3 mg/kg/day)	
		mg/dose	mL/dose*
			q8h
120	264	120 mg	3 mL
115	253	115 mg	2.9 mL
110	242	110 mg	2.75 mL
105	231	105 mg	2.6 mL
100	220	100 mg	2.5 mL
95	209	95 mg	2.4 mL
90	198	90 mg	2.25 mL
85	187	85 mg	2.1 mL
80	176	80 mg	2 mL
75	165	75 mg	1.9 mL
70	154	70 mg	1.75 mL
65	143	65 mg	1.6 mL
60	132	60 mg	1.5 mL
55	121	55 mğ	1.4 mL
50 45	110	50 mg	1.25 mL
	99	45 mğ	1.1 mL
40	88	40 mg	1 mL

For Patient Weighing kg lb		Maximum Dose for Life- Threatening Infections (Reduce as soon as possible) 1.66 mg/kg q8h (Total, 5 mg/kg/day)	
	-	mg/dose	mL/dose* q8h
120	264	200 mg	5 mL
115	253	191 mg	4.75 mL
110	242	183 mg	4.5 mL
105	231	175 mg	4.4 mL
100	220	166 mg	4.2 mL
95	209	158 mg	4 mL
90 85	198 187	150 mg	3.75 mL 3.5 mL
80	176	141 mg 133 mg	3.3 mL
75	165	125 mg	3.1 mL
70	154	116 mg	2.9 mL
65	143	108 mg	2.7 mL
60	132	100 mg	2.5 mL
55	121	91 mg	2.25 mL
50	110	83 mg	2.1 mL
45	99	75 mg	1.9 mL
40	88	66 mg	1.6 mL

*Applicable to all product forms except the tobramycin injection, USP, (Pediatric).

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3 Pediatric patients (greater than 1 week of age)

- 4 6 to 7.5 mg/kg/day in 3 or 4 equally divided doses (2 to 2.5 mg/kg every 8 hours or 1.5 to
- 5 1.89 mg/kg every 6 hours).

1 Premature or full-term neonates 1 week of age or less 2 Up to 4 mg/kg/day may be administered in 2 equal doses every 12 hours. 3 It is desirable to limit treatment to a short term. The usual duration of treatment is 4 7 to 10 days. A longer course of therapy may be necessary in difficult and complicated 5 infections. In such cases, monitoring of renal, auditory, and vestibular functions is 6 advised, because neurotoxicity is more likely to occur when treatment is extended 7 longer than 10 days. 8 9 Dosage in Patients with Cystic Fibrosis 10 In patients with cystic fibrosis, altered pharmacokinetics may result in reduced serum 11 concentrations of aminoglycosides. Measurement of tobramycin serum concentration 12 during treatment is especially important as a basis for determining appropriate dose. In 13 patients with severe cystic fibrosis, an initial dosing regimen of 10 mg/kg/day in 4 14 equally divided doses is recommended. This dosing regimen is suggested only as a 15 guide. The serum levels of tobramycin should be measured directly during treatment due 16 to wide interpatient variability. 17 Administration for Patients with Impaired Renal Function 18 19 Whenever possible, serum tobramycin concentrations should be monitored during 20 therapy. 21 Following a loading dose of 1 mg/kg, subsequent dosage in these patients must be 22 adjusted, either with reduced doses administered at 8-hour intervals or with normal doses

given at prolonged intervals. Both of these methods are suggested as guides to be used

1 when serum levels of tobramycin cannot be measured directly. They are based on either 2 the creatinine clearance level or the serum creatinine level of the patient because 3 these values correlate with the half-life of tobramycin. The dosage schedule derived from 4 either method should be used in conjunction with careful clinical and laboratory 5 observations of the patient and should be modified as necessary. Neither method should 6 be used when dialysis is being performed. 7 8 Reduced dosage at 8-hour intervals 9 When the creatinine clearance rate is 70 mL or less per minute or when the serum 10 creatinine value is known, the amount of the reduced dose can be determined by 11 multiplying the normal dose from Table 1 by the percent of normal dose from the 12 accompanying nomogram. 13 An alternate rough guide for determining reduced dosage at 8-hour intervals (for 14 patients whose steadystate serum creatinine values are known) is to divide the normally 15 recommended dose by the patient's serum creatinine. 16 17 Normal dosage at prolonged intervals 18 If the creatinine clearance rate is not available and the patient's condition is stable, a 19 dosage frequency in hours for the dosage given in Table 1 can be determined by 20 multiplying the patient's serum creatinine by 6. 21 22 23

Dosage in Obese Patients

2 The appropriate dose may be calculated by using the patient's estimated lean body weight

plus 40% of the excess as the basic weight on which to figure mg/kg.

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Intravenous Administration

- 6 For intravenous administration, the usual volume of diluent (0.9% Sodium Chloride
- 7 Injection or 5% Dextrose Injection) is 50 to 100 mL for adult doses. For pediatric
- 8 patients, the volume of diluent should be proportionately less than that for adults. The
- 9 diluted solution usually should be infused over a period of 20 to 60 minutes.
- 10 Infusion periods of less than 20 minutes are not recommended because peak serum levels
- may exceed 12 mcg/mL (see **WARNINGS** box).
- Tobramycin should not be physically premixed with other drugs but should be
- administered separately according to the recommended dose and route.

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PREPARATION AND STORAGE:

- 16 Directions for Proper Use of Pharmacy Bulk Package-Not for direct infusion
- 17 The pharmacy bulk package is for use in the Hospital Pharmacy Admixture Service and
- only in a suitable work area, such as a laminar flow hood. Using aseptic technique, the
- 19 closure may be penetrated only 1 time after reconstitution using a suitable sterile transfer
- device or dispensing set, which allows measured dispensing of the contents. Use of a
- 21 syringe and needle is not recommended as it may cause leakage. After entry, entire
- contents of bulk vial should be dispensed within 24 hours.

1 Tobramycin for Injection, USP is supplied as a dry powder. The contents of the 2 vial should be diluted with 30 mL of Sterile Water for Injection, USP, to provide a 3 solution containing 40 mg of tobramycin per mL. Prior to reconstitution, the vial should 4 be stored at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature]. 5 After reconstitution, the solution should be kept in a refrigerator and used within 96 6 hours. If kept at room temperature, the solution must be used within 24 hours. 7 Prior to administration, parenteral drug products should be inspected visually for 8 particulate matter and discoloration whenever solution and container permit. 9 10 **HOW SUPPLIED:** 11 **Product NDC** 12 No No 13 300351 63323-303-51 Tobramycin for Injection, USP equivalent to 1.2 g tobramycin in a 50 mL pharmacy bulk package vial, 14 packaged in trays of 6. 15 16 17 Vial stoppers do not contain natural rubber latex. 18 19 **REFERENCES:** 20 1. National Committee for Clinical Laboratory Standards, Performance Standards for 21 Antimicrobial Disk Susceptibility Tests-Sixth Edition. Approved Standard NCCLS 22 Document M2-A6, Vol. 17, No. 1, NCCLS, Wayne, PA, 1997. 23 2. National Committee for Clinical Laboratory Standards, Methods for Dilution 24 Antimicrobial Susceptibility Tests for Bacteria that Grow Aerobically–Fourth Edition.

Approved Standard NCCLS Document M7-A4, Vol. 17, No. 2, NCCLS, Wayne, PA,

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