1	ROCHE
2	ROCEPHIN®
3	(ceftriaxone sodium)
4	FOR INJECTION
5	Rx only
6 7 8	To reduce the development of drug-resistant bacteria and maintain the effectiveness of Rocephin and other antibacterial drugs, Rocephin should be used only to treat or prevent infections that are proven or strongly suspected to be caused by bacteria.
9 10 11 12 13	DESCRIPTION: Rocephin is a sterile, semisynthetic, broad-spectrum cephalosporin antibiotic for intravenous or intramuscular administration. Ceftriaxone sodium is $(6R,7R)$ -7-[2-(2-Amino-4-thiazolyl)glyoxylamido]-8-oxo-3-[[(1,2,5,6-tetrahydro-2-methyl-5,6-dioxo- as -triazin-3-yl)thio]methyl]-5-thia-1-azabicyclo[4.2.0]oct-2-ene-2-carboxylic acid, 7^2 -(Z)-(O -methyloxime), disodium salt, sesquaterhydrate.
14 15	The chemical formula of ceftriaxone sodium is $C_{18}H_{16}N_8Na_2O_7S_3 \cdot 3.5H_2O$. It has a calculated molecular weight of 661.59 and the following structural formula:
	NH ₂ CCONH H S CH ₃ CH ₃ NO - Na+ CH ₃ OCH ₃ OCH ₃ O - Na+
16	COO - Na+
17 18 19 20	Rocephin is a white to yellowish-orange crystalline powder which is readily soluble in water, sparingly soluble in methanol and very slightly soluble in ethanol. The pH of a 1% aqueous solution is approximately 6.7. The color of Rocephin solutions ranges from light yellow to amber, depending on the length of storage, concentration and diluent used.
21 22	Rocephin contains approximately 83 mg (3.6 mEq) of sodium per gram of ceftriaxone activity.
23 24 25 26	CLINICAL PHARMACOLOGY: Average plasma concentrations of ceftriaxone following a single 30-minute intravenous (IV) infusion of a 0.5, 1 or 2 gm dose and intramuscular (IM) administration of a single 0.5 (250 mg/mL or 350 mg/mL concentrations) or 1 gm dose in healthy subjects are presented in Table 1.

Table 1 Ceftriaxone Plasma Concentrations After Single Dose Administration

Dose/Route			Average Plasma Concentrations (μg/mL)						
	0.5 hr	1 hr	2 hr	4 hr	6 hr	8 hr	12 hr	16 hr	24 hr
0.5 gm IV*	82	59	48	37	29	23	15	10	5
0.5 gm IM				•					
[.] 250 mg/mL	22	33	38	35	30	26	16	ND	5
0.5 gm IM									
350 mg/mL	20	32	38	34	31	24	16	ND	5
1 gm IV*	151	111	88	67	53	43	28	18	9
1 gm IM	40	68	76	68	56	44	29	ND	ND
2 gm IV*	257	192	154	117	89	74	46	31	15

29 *IV doses were infused at a constant rate over 30 minutes.

30 ND = Not determined.

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31 Ceftriaxone was completely absorbed following IM administration with mean maximum

32 plasma concentrations occurring between 2 and 3 hours postdosing. Multiple IV or IM

doses ranging from 0.5 to 2 gm at 12- to 24-hour intervals resulted in 15% to 36%

34 accumulation of ceftriaxone above single dose values.

35 Ceftriaxone concentrations in urine are high, as shown in Table 2.

Table 2 Urinary Concentrations of Ceftriaxone After Single Dose
 Administration

Dose/Route	Average Urinary Concentrations (µg/mL)					
	0-2 hr	2-4 hr	4-8 hr	8-12 hr	12-24 hr	24-48 hr
0.5 gm IV	526	366	142	87	70	15
0.5 gm IM	115	425	308	127	96	28
1 gm IV	995	855	293	147	132	32
1 gm IM	504	628	418	237	ND	ND
2 gm IV	2692	1976	757	274	198	40

38 ND = Not determined.

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Thirty-three percent to 67% of a ceftriaxone dose was excreted in the urine as unchanged drug and the remainder was secreted in the bile and ultimately found in the feces as microbiologically inactive compounds. After a 1 gm IV dose, average concentrations of

42 ceftriaxone, determined from 1 to 3 hours after dosing, were 581 μg/mL in the

43 gallbladder bile, 788 μ g/mL in the common duct bile, 898 μ g/mL in the cystic duct bile,

44 78.2 μg/gm in the gallbladder wall and 62.1 μg/mL in the concurrent plasma.

45 Over a 0.15 to 3 gm dose range in healthy adult subjects, the values of elimination half-

46 life ranged from 5.8 to 8.7 hours; apparent volume of distribution from 5.78 to 13.5 L;

plasma clearance from 0.58 to 1.45 L/hour; and renal clearance from 0.32 to 0.73 L/hour.

48 Ceftriaxone is reversibly bound to human plasma proteins, and the binding decreased

from a value of 95% bound at plasma concentrations of <25 µg/mL to a value of 85%

bound at 300 µg/mL. Ceftriaxone crosses the blood placenta barrier.

- 51 The average values of maximum plasma concentration, elimination half-life, plasma
- 52 clearance and volume of distribution after a 50 mg/kg IV dose and after a 75 mg/kg IV
- dose in pediatric patients suffering from bacterial meningitis are shown in Table 3.
- 54 Ceftriaxone penetrated the inflamed meninges of infants and pediatric patients; CSF
- 55 concentrations after a 50 mg/kg IV dose and after a 75 mg/kg IV dose are also shown in
- 56 Table 3.

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57 Table 3 Average Pharmacokinetic Parameters of Ceftriaxone in Pediatric Patients With Meningitis

	50 mg/kg IV	75 mg/kg IV
Maximum Plasma Concentrations (μg/mL)	216	275
Elimination Half-life (hr)	4.6	4.3
Plasma Clearance (mL/hr/kg)	49	60
Volume of Distribution (mL/kg)	338	373
CSF Concentration—inflamed meninges (µg/mL)	5.6	6.4
Range (µg/mL)	1.3-18.5	1.3-44
Time after dose (hr)	3.7 (± 1.6)	3.3 (± 1.4)

Compared to that in healthy adult subjects, the pharmacokinetics of ceftriaxone were only minimally altered in elderly subjects and in patients with renal impairment or hepatic dysfunction (Table 4); therefore, dosage adjustments are not necessary for these patients with ceftriaxone dosages up to 2 gm per day. Ceftriaxone was not removed to any significant extent from the plasma by hemodialysis. In 6 of 26 dialysis patients, the elimination rate of ceftriaxone was markedly reduced, suggesting that plasma concentrations of ceftriaxone should be monitored in these patients to determine if dosage adjustments are necessary.

Table 4 Average Pharmacokinetic Parameters of Ceftriaxone in Humans

Subject Group	Elimination	Plasma	Volume of
	Half-Life	Clearance	Distribution
	(hr)	(L/hr)	(L)
Healthy Subjects	5.8-8.7	0.58-1.45	5.8-13.5
Elderly Subjects (mean age, 70.5 yr)	8.9	0.83	10.7
Patients With Renal Impairment			
Hemodialysis Patients (0-5 mL/min)*	14.7	0.65	13.7
Severe (5-15 mL/min)	15.7	0.56	12.5
Moderate (16-30 mL/min)	11.4	0.72	11.8
Mild (31-60 mL/min)	12.4	0.70	13.3
Patients With Liver Disease	8.8	1.1	13.6

*Creatinine clearance.

Pharmacokinetics in the Middle Ear Fluid: In one study, total ceftriaxone concentrations (bound and unbound) were measured in middle ear fluid obtained during the insertion of tympanostomy tubes in 42 pediatric patients with otitis media. Sampling times were from 1 to 50 hours after a single intramuscular injection of 50 mg/kg of

- 74 ceftriaxone. Mean (\pm SD) ceftriaxone levels in the middle ear reached a peak of 35 (\pm 12)
- 75 μg/mL at 24 hours, and remained at 19 (± 7) μg/mL at 48 hours. Based on middle ear
- fluid ceftriaxone concentrations in the 23 to 25 hour and the 46 to 50 hour sampling time
- 77 intervals, a half-life of 25 hours was calculated. Ceftriaxone is highly bound to plasma
- 78 proteins. The extent of binding to proteins in the middle ear fluid is unknown.
- 79 Microbiology: The bactericidal activity of ceftriaxone results from inhibition of cell wall
- 80 synthesis. Ceftriaxone has a high degree of stability in the presence of beta-lactamases,
- both penicillinases and cephalosporinases, of gram-negative and gram-positive bacteria.
- 82 Ceftriaxone has been shown to be active against most strains of the following
- 83 microorganisms, both in vitro and in clinical infections described in the INDICATIONS
- 84 AND USAGE section.
- 85 Aerobic gram-negative microorganisms:
- 86 Acinetobacter calcoaceticus
- 87 Enterobacter aerogenes
- 88 Enterobacter cloacae
- 89 Escherichia coli
- 90 Haemophilus influenzae (including ampicillin-resistant and beta-lactamase producing
- 91 strains
- 92 Haemophilus parainfluenzae
- 93 Klebsiella oxytoca
- 94 Klebsiella pneumoniae
- 95 Moraxella catarrhalis (including beta-lactamase producing strains)
- 96 Morganella morganii
- 97 Neisseria gonorrhoeae (including penicillinase- and nonpenicillinase-producing strains)
- 98 Neisseria meningitidis
- 99 Proteus mirabilis
- 100 Proteus vulgaris
- 101 Serratia marcescens
- 102 Ceftriaxone is also active against many strains of *Pseudomonas aeruginosa*.
- 103 NOTE: Many strains of the above organisms that are multiply resistant to other
- 104 antibiotics, eg, penicillins, cephalosporins, and aminoglycosides, are susceptible to
- 105 ceftriaxone.
- 106 Aerobic gram-positive microorganisms:
- 107 Staphylococcus aureus (including penicillinase-producing strains)
- 108 Staphylococcus epidermidis
- 109 Streptococcus pneumoniae
- 110 Streptococcus pyogenes
- 111 Viridans group streptococci
- 112 NOTE: Methicillin-resistant staphylococci are resistant to cephalosporins, including
- 113 ceftriaxone. Most strains of Group D streptococci and enterococci, eg, Enterococcus
- 114 (Streptococcus) faecalis, are resistant.

115 116 117 118	Anaerobic microorganisms: Bacteroides fragilis Clostridium species Peptostreptococcus species	
119	NOTE: Most strains of Clostridium difficile are	e resistant.
120 121 122 123 124	The following in vitro data are available, <u>bu</u> Ceftriaxone exhibits in vitro minimal inhibite less against most strains of the following reffectiveness of ceftriaxone in treating clinic have not been established in adequate and well	ry concentrations (MICs) of ≤8 µg/mL or nicroorganisms, however, the safety and al infections due to these microorganisms
125 126 127 128 129 130	Aerobic gram-negative microorganisms: Citrobacter diversus Citrobacter freundii Providencia species (including Providencia re Salmonella species (including Salmonella typh Shigella species	
131 132	Aerobic gram-positive microorganisms: Streptococcus agalactiae	
133 134 135	Anaerobic microorganisms: Prevotella (Bacteroides) bivius Porphyromonas (Bacteroides) melaninogenicu	s
136 137 138 139 140 141 142 143 144	Susceptibility Tests: Dilution Techniques: Quantitative methods as inhibitory concentrations (MICs). These MIC bacteria to antimicrobial compounds. The standardized procedure. Standardized procedure or agar) or equivalent with standardized in concentrations of ceftriaxone powder. The MIC the following criteria for aerobic organism gonorrhoeae, and Streptococcus spp, including	s provide estimates of the susceptibility of MICs should be determined using a ares are based on a dilution method (broth noculum concentrations and standardized C values should be interpreted according to so ther than <i>Haemophilus</i> spp, <i>Neisseria</i>
	MIC (μg/mL)	<u>Interpretation</u>
145 146	≤8 16-32 ≥64 The following interpretive criteria ² should be using Haemophilus Test Media (HTM).	(S) Susceptible (I) Intermediate (R) Resistant e used when testing Haemophilus species
	MIC (µg/mL)	<u>Interpretation</u>
	≤2	(S) Susceptible
	•	

- 147 The absence of resistant strains precludes defining any categories other than
- 148 "Susceptible". Strains vielding results suggestive of a "Nonsusceptible" category should
- 149 be submitted to a reference laboratory for further testing.
- The following interpretive criteria² should be used when testing Neisseria gonorrhoeae 150
- when using GC agar base and 1% defined growth supplement. 151

MIC (µg/mL) Interpretation ≤0.25 (S) Susceptible

- 152 The absence of resistant strains precludes defining any categories other than
- "Susceptible". Strains yielding results suggestive of a "Nonsusceptible" category should 153
- 154 be submitted to a reference laboratory for further testing.
- The following interpretive criteria² should be used when testing Streptococcus spp 155
- 156 including Streptococcus pneumoniae using cation-adjusted Mueller-Hinton broth with 2
- 157 to 5% lysed horse blood.

MIC (μg/mL)	<u>Interpretation</u>
≤0.5	(S) Susceptible
1	(I) Intermediate
≥2	(R) Resistant

- A report of "Susceptible" indicates that the pathogen is likely to be inhibited if the 158 159 antimicrobial compound in the blood reaches the concentrations usually achievable. A 160 report of "Intermediate" indicates that the results should be considered equivocal, and if
- 161 the microorganism is not fully susceptible to alternative, clinically feasible drugs, the test
- 162 should be repeated. This category implies possible clinical applicability in body sites
- 163 where the drug is physiologically concentrated or in situations where high dosage of the
- 164 drug can be used. This category also provides a buffer zone which prevents small
- 165 uncontrolled technical factors from causing major discrepancies in interpretation. A
- 166
- report of "Resistant" indicates that the pathogen is not likely to be inhibited if the
- 167 antimicrobial compound in the blood reaches the concentrations usually achievable; other
- 168 therapy should be selected.

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- 169 Standardized susceptibility test procedures require the use of laboratory control
- 170 microorganisms to control the technical aspects of the laboratory procedures.
- 171 Standardized ceftriaxone powder should provide the following MIC values:²

Microorganism	ATCC®#	MIC (μg/mL)
Escherichia coli	25922	0.03 - 0.12
Staphylococcus aureus	29213	1 - 8*
Pseudomonas aeruginosa	27853	8 - 32
Haemophilus influenzae	49247	0.06 - 0.25
Neisseria gonorrhoeae	49226	0.004 - 0.015
Streptococcus pneumoniae	49619	0.03 - 0.12

^{*} A bimodal distribution of MICs results at the extremes of the acceptable range should be suspect and control validity should be verified with data from other control strains.

Diffusion Techniques: Quantitative methods that require measurement of zone 174 diameters also provide reproducible estimates of the susceptibility of bacteria to 175 antimicrobial compounds. One such standardized procedure³ requires the use of 176 standardized inoculum concentrations. This procedure uses paper discs impregnated with 177 30 µg of ceftriaxone to test the susceptibility of microorganisms to ceftriaxone. 178 179 Reports from the laboratory providing results of the standard single-disc susceptibility test with a 30 ug ceftriaxone disc should be interpreted according to the following criteria 180 for aerobic organisms other than Haemophilus spp, Neisseria gonorrhoeae, and 181 182 Streptococcus spp: Zone Diameter (mm) Interpretation (S) Susceptible ≥21 (I) Intermediate 14-20 ≤13 (R) Resistant The following interpretive criteria³ should be used when testing *Haemophilus* species 183 when using Haemophilus Test Media (HTM). 184 Zone Diameter (mm) Interpretation ≥26 (S) Susceptible The absence of resistant strains precludes defining any categories other than 185 "Susceptible". Strains yielding results suggestive of a "Nonsusceptible" category should 186 be submitted to a reference laboratory for further testing. 187 The following interpretive criteria³ should be used when testing Neisseria gonorrhoeae 188 when using GC agar base and 1% defined growth supplement. 189 Zone Diameter (mm) Interpretation (S) Susceptible ≥35 190 The absence of resistant strains precludes defining any categories other than "Susceptible". Strains yielding results suggestive of a "Nonsusceptible" category should 191 be submitted to a reference laboratory for further testing. 192 193 The following interpretive criteria³ should be used when testing Streptococcus spp other 194 than Streptococcus pneumoniae when using Mueller-Hinton agar supplemented with 5% 195 sheep blood incubated in 5% CO₂. Zone Diameter (mm) Interpretation (S) Susceptible ≥27 25-26 (I) Intermediate (R) Resistant ≤24 196 Interpretation should be as stated above for results using dilution techniques. Interpretation involves correlation of the diameter obtained in the disc test with the MIC 197 198 for ceftriaxone. 199 Disc diffusion interpretive criteria for ceftriaxone discs against Streptococcus 200 pneumoniae are not available, however, isolates of pneumococci with oxacillin zone

- 201 diameters of >20 mm are susceptible (MIC \leq 0.06 $\mu g/mL$) to penicillin and can be
- 202 considered susceptible to ceftriaxone. Streptococcus pneumoniae isolates should not be
- 203 reported as penicillin (ceftriaxone) resistant or intermediate based solely on an oxacillin
- zone diameter of \leq 19 mm. The ceftriaxone MIC should be determined for those isolates
- 205 with oxacillin zone diameters ≤19 mm.
- 206 As with standardized dilution techniques, diffusion methods require the use of laboratory
- 207 control microorganisms that are used to control the technical aspects of the laboratory
- 208 procedures. For the diffusion technique, the 30 µg ceftriaxone disc should provide the
- 209 following zone diameters in these laboratory test quality control strains:³

<u>Microorganism</u>	ATCC®#	Zone Diameter Ranges (mm)
Escherichia coli	25922	29 - 35
Staphylococcus aureus	25923	22 - 28
Pseudomonas aeruginosa	27853	17 - 23
Haemophilus influenzae	49247	31 - 39
Neisseria gonorrhoeae	49226	39 - 51
Streptococcus pneumoniae	49619	30 - 35

- 210 Anaerobic Techniques: For anaerobic bacteria, the susceptibility to ceftriaxone as MICs
- 211 can be determined by standardized test methods.⁴ The MIC values obtained should be
- 212 interpreted according to the following criteria:

MIC (μg/mL)	Interpretation
≤16	(S) Susceptible
32	(I) Intermediate
>64	(R) Resistant

- 213 As with other susceptibility techniques, the use of laboratory control microorganisms is
- 214 required to control the technical aspects of the laboratory standardized procedures.
- 215 Standardized ceftriaxone powder should provide the following MIC values for the
- 216 indicated standardized anaerobic dilution⁴ testing method:

<u>Method</u>	Microorganism	ATCC®#	$MIC (\mu g/mL)$
Agar	Bacteroides fragilis	25285	32 - 128
	Bacteroides thetaiotaomicron	29741	64 - 256
Broth	Bacteroides thetaiotaomicron	29741	32 - 128

- 217 ATCC® is a registered trademark of the American Type Culture Collection.
- 218 INDICATIONS AND USAGE: Before instituting treatment with Rocephin, appropriate
- 219 specimens should be obtained for isolation of the causative organism and for
- 220 determination of its susceptibility to the drug. Therapy may be instituted prior to
- 221 obtaining results of susceptibility testing.
- 222 To reduce the development of drug-resistant bacteria and maintain the effectiveness of
- 223 Rocephin and other antibacterial drugs, Rocephin should be used only to treat or prevent
- 224 infections that are proven or strongly suspected to be caused by susceptible bacteria.
- 225 When culture and susceptibility information are available, they should be considered in

- 226 selecting or modifying antibacterial therapy. In the absence of such data, local
- 227 epidemiology and susceptibility patterns may contribute to the empiric selection of
- 228 therapy.
- 229 Rocephin is indicated for the treatment of the following infections when caused by
- 230 susceptible organisms:
- 231 LOWER RESPIRATORY TRACT INFECTIONS caused by Streptococcus pneumoniae,
- 232 Staphylococcus aureus, Haemophilus influenzae, Haemophilus parainfluenzae,
- 233 Klebsiella pneumoniae, Escherichia coli, Enterobacter aerogenes, Proteus mirabilis or
- 234 Serratia marcescens.
- 235 ACUTE BACTERIAL OTITIS MEDIA caused by Streptococcus pneumoniae,
- 236 Haemophilus influenzae (including beta-lactamase producing strains) or Moraxella
- 237 catarrhalis (including beta-lactamase producing strains).
- 238 NOTE: In one study lower clinical cure rates were observed with a single dose of
- 239 Rocephin compared to 10 days of oral therapy. In a second study comparable cure rates
- 240 were observed between single dose Rocephin and the comparator. The potentially lower
- 241 clinical cure rate of Rocephin should be balanced against the potential advantages of
- 242 parenteral therapy (see CLINICAL STUDIES).
- 243 SKIN AND SKIN STRUCTURE INFECTIONS caused by Staphylococcus aureus,
- 244 Staphylococcus epidermidis, Streptococcus pyogenes, Viridans group streptococci,
- 245 Escherichia coli, Enterobacter cloacae, Klebsiella oxytoca, Klebsiella pneumoniae,
- 246 Proteus mirabilis, Morganella morganii,* Pseudomonas aeruginosa, Serratia
- 247 marcescens, Acinetobacter calcoaceticus, Bacteroides fragilis* or Peptostreptococcus
- 248 species.
- 249 URINARY TRACT INFECTIONS (complicated and uncomplicated) caused by
- 250 Escherichia coli, Proteus mirabilis, Proteus vulgaris, Morganella morganii or Klebsiella
- 251 pneumoniae.
- 252 UNCOMPLICATED GONORRHEA (cervical/urethral and rectal) caused by Neisseria
- 253 gonorrhoeae, including both penicillinase- and nonpenicillinase-producing strains, and
- 254 pharyngeal gonorrhea caused by nonpenicillinase-producing strains of Neisseria
- 255 gonorrhoeae.
- 256 PELVIC INFLAMMATORY DISEASE caused by Neisseria gonorrhoeae. Rocephin, like
- 257 other cephalosporins, has no activity against Chlamydia trachomatis. Therefore, when
- cephalosporins are used in the treatment of patients with pelvic inflammatory disease and
- 259 Chlamydia trachomatis is one of the suspected pathogens, appropriate antichlamydial
- 260 coverage should be added.
- 261 BACTERIAL SEPTICEMIA caused by Staphylococcus aureus, Streptococcus
- 262 pneumoniae, Escherichia coli, Haemophilus influenzae or Klebsiella pneumoniae.
- 263 BONE AND JOINT INFECTIONS caused by Staphylococcus aureus, Streptococcus
- 264 pneumoniae, Escherichia coli, Proteus mirabilis, Klebsiella pneumoniae or Enterobacter
- 265 species.

- 266 INTRA-ABDOMINAL INFECTIONS caused by Escherichia coli, Klebsiella pneumoniae,
- 267 Bacteroides fragilis, Clostridium species (Note: most strains of Clostridium difficile are
- 268 resistant) or Peptostreptococcus species.
- 269 MENINGITIS caused by Haemophilus influenzae, Neisseria meningitidis or
- 270 Streptococcus pneumoniae. Rocephin has also been used successfully in a limited number
- 271 of cases of meningitis and shunt infection caused by Staphylococcus epidermidis * and
- 272 Escherichia coli.*
- 273 *Efficacy for this organism in this organ system was studied in fewer than ten infections.
- 274 SURGICAL PROPHYLAXIS: The preoperative administration of a single 1 gm dose of
- 275 Rocephin may reduce the incidence of postoperative infections in patients undergoing
- 276 surgical procedures classified as contaminated or potentially contaminated (eg. vaginal or
- 277 abdominal hysterectomy or cholecystectomy for chronic calculous cholecystitis in high-
- 278 risk patients, such as those over 70 years of age, with acute cholecystitis not requiring
- 279 therapeutic antimicrobials, obstructive jaundice or common duct bile stones) and in
- surgical patients for whom infection at the operative site would present serious risk (eg.
- 281 during coronary artery bypass surgery). Although Rocephin has been shown to have been
- as effective as cefazolin in the prevention of infection following coronary artery bypass
- 283 surgery, no placebo-controlled trials have been conducted to evaluate any cephalosporin
- 284 antibiotic in the prevention of infection following coronary artery bypass surgery.
- 285 When administered prior to surgical procedures for which it is indicated, a single 1 gm
- dose of Rocephin provides protection from most infections due to susceptible organisms
- 287 throughout the course of the procedure.
- 288 CONTRAINDICATIONS: Rocephin is contraindicated in patients with known allergy
- 289 to the cephalosporin class of antibiotics.
- 290 Hyperbilirubinemic neonates, especially prematures, should not be treated with
- 291 Rocephin. In vitro studies have shown that ceftriaxone can displace bilirubin from its
- 292 binding to serum albumin and bilirubin encephalopathy can possibly develop in these
- 293 patients.
- 294 Rocephin should not be administered concurrently with calcium treatment in newborns
- because of the risk of precipitation of ceftriaxone-calcium salt (see WARNINGS).
- 296 WARNINGS: BEFORE THERAPY WITH ROCEPHIN IS INSTITUTED, CAREFUL
- 297 INQUIRY SHOULD BE MADE TO DETERMINE WHETHER THE PATIENT HAS
- 298 HAD PREVIOUS HYPERSENSITIVITY REACTIONS TO CEPHALOSPORINS,
- 299 PENICILLINS OR OTHER DRUGS. THIS PRODUCT SHOULD BE GIVEN
- 300 CAUTIOUSLY TO PENICILLIN-SENSITIVE PATIENTS. ANTIBIOTICS SHOULD
- 301 BE ADMINISTERED WITH CAUTION TO ANY PATIENT WHO HAS
- 302 DEMONSTRATED SOME FORM OF ALLERGY, PARTICULARLY TO DRUGS.
- 303 SERIOUS ACUTE HYPERSENSITIVITY REACTIONS MAY REQUIRE THE USE
- 304 OF SUBCUTANEOUS EPINEPHRINE AND OTHER EMERGENCY MEASURES.
- 305 Clostridium difficile associated diarrhea (CDAD) has been reported with use of
- 306 nearly all antibacterial agents, including Rocephin, and may range in severity from

- 307 mild diarrhea to fatal colitis. Treatment with antibacterial agents alters the normal
- 308 flora of the colon leading to overgrowth of C. difficile.
- 309 C. difficile produces toxins A and B which contribute to the development of CDAD.
- 310 Hypertoxin producing strains of C. difficile cause increased morbidity and mortality, as
- 311 these infections can be refractory to antimicrobial therapy and may require colectomy.
- 312 CDAD must be considered in all patients who present with diarrhea following antibiotic
- 313 use. Careful medical history is necessary since CDAD has been reported to occur over
- 314 two months after the administration of antibacterial agents.
- 315 If CDAD is suspected or confirmed, ongoing antibiotic use not directed against C.
- 316 difficile may need to be discontinued. Appropriate fluid and electrolyte management,
- 317 protein supplementation, antibiotic treatment C. difficile, and surgical evaluation should
- 318 be instituted as clinically indicated.
- 319 Ceftriaxone must not be mixed or administered simultaneously with calcium-containing
- 320 solutions or products, even via different infusion lines.
- 321 Calcium-containing solutions or products must not be administered within 48 hours of
- 322 last administration of ceftriaxone.
- 323 Cases of fatal reactions with calcium-ceftriaxone precipitates in lung and kidneys in
- 324 neonates and prematures have been described. In some cases the infusion lines and times
- 325 of administration of ceftriaxone and calcium-containing solutions differed (see
- 326 CONTRAINDICATIONS and ADVERSE REACTIONS).
- 327 **PRECAUTIONS:** General: Prescribing Rocephin in the absence of a proven or strongly
- 328 suspected bacterial infection or a prophylactic indication is unlikely to provide benefit to
- 329 the patient and increases the risk of the development of drug-resistant bacteria.
- 330 Although transient elevations of BUN and serum creatinine have been observed, at the
- 331 recommended dosages, the nephrotoxic potential of Rocephin is similar to that of other
- 332 cephalosporins.
- 333 Ceftriaxone is excreted via both biliary and renal excretion (see CLINICAL
- 334 PHARMACOLOGY). Therefore, patients with renal failure normally require no
- adjustment in dosage when usual doses of Rocephin are administered, but concentrations
- 336 of drug in the serum should be monitored periodically. If evidence of accumulation
- exists, dosage should be decreased accordingly.
- 338 Dosage adjustments should not be necessary in patients with hepatic dysfunction;
- 339 however, in patients with both hepatic dysfunction and significant renal disease,
- 340 Rocephin dosage should not exceed 2 gm daily without close monitoring of serum
- 341 concentrations.
- 342 Alterations in prothrombin times have occurred rarely in patients treated with Rocephin.
- Patients with impaired vitamin K synthesis or low vitamin K stores (eg, chronic hepatic
- 344 disease and malnutrition) may require monitoring of prothrombin time during Rocephin
- 345 treatment. Vitamin K administration (10 mg weekly) may be necessary if the
- 346 prothrombin time is prolonged before or during therapy.

- 347 Prolonged use of Rocephin may result in overgrowth of nonsusceptible organisms.
- 348 Careful observation of the patient is essential. If superinfection occurs during therapy,
- 349 appropriate measures should be taken.
- 350 Rocephin should be prescribed with caution in individuals with a history of
- 351 gastrointestinal disease, especially colitis.
- 352 There have been reports of sonographic abnormalities in the gallbladder of patients
- 353 treated with Rocephin; some of these patients also had symptoms of gallbladder
- 354 disease. These abnormalities appear on sonography as an echo without acoustical
- 355 shadowing suggesting sludge or as an echo with acoustical shadowing which may be
- 356 misinterpreted as gallstones. The chemical nature of the sonographically detected
- 357 material has been determined to be predominantly a ceftriaxone-calcium salt. The
- condition appears to be transient and reversible upon discontinuation of Rocephin
- and institution of conservative management. Therefore, Rocephin should be
- and institution of conscivative management. Therefore, Roccomm should be
- discontinued in patients who develop signs and symptoms suggestive of gallbladder
- 361 disease and/or the sonographic findings described above.
- 362 Information for Patients: Patients should be counseled that antibacterial drugs including
- 363 Rocephin should only be used to treat bacterial infections. They do not treat viral
- infections (eg, common cold). When Rocephin is prescribed to treat a bacterial infection,
- patients should be told that although it is common to feel better early in the course of
- 366 therapy, the medication should be taken exactly as directed. Skipping doses or not
- 367 completing the full course of therapy may (1) decrease the effectiveness of the immediate
- 368 treatment and (2) increase the likelihood that bacteria will develop resistance and will not
- be treatable by Rocephin or other antibacterial drugs in the future.
- 370 Diarrhea is a common problem caused by antibiotics which usually ends when the
- 371 antibiotic is discontinued. Sometimes after starting treatment with antibiotics, patients
- 372 can develop watery and bloody stools (with or without stomach cramps and fever) even
- as late as two or more months after having taken the last dose of the antibiotic. If this
- occurs, patients should contact their physician as soon as possible.
- 375 Carcinogenesis, Mutagenesis, Impairment of Fertility: Carcinogenesis: Considering the
- 376 maximum duration of treatment and the class of the compound, carcinogenicity studies
- 377 with ceftriaxone in animals have not been performed. The maximum duration of animal
- 378 toxicity studies was 6 months.
- 379 Mutagenesis: Genetic toxicology tests included the Ames test, a micronucleus test and a
- 380 test for chromosomal aberrations in human lymphocytes cultured in vitro with
- 381 ceftriaxone. Ceftriaxone showed no potential for mutagenic activity in these studies.
- 382 Impairment of Fertility: Ceftriaxone produced no impairment of fertility when given
- intravenously to rats at daily doses up to 586 mg/kg/day, approximately 20 times the
- recommended clinical dose of 2 gm/day.
- 385 Pregnancy: Teratogenic Effects: Pregnancy Category B. Reproductive studies have been
- 386 performed in mice and rats at doses up to 20 times the usual human dose and have no

- 387 evidence of embryotoxicity, fetotoxicity or teratogenicity. In primates, no embryotoxicity
- or teratogenicity was demonstrated at a dose approximately 3 times the human dose.
- 389 There are, however, no adequate and well-controlled studies in pregnant women. Because
- 390 animal reproductive studies are not always predictive of human response, this drug
- should be used during pregnancy only if clearly needed.
- 392 Nonteratogenic Effects: In rats, in the Segment I (fertility and general reproduction) and
- 393 Segment III (perinatal and postnatal) studies with intravenously administered ceftriaxone,
- 394 no adverse effects were noted on various reproductive parameters during gestation and
- 395 lactation, including postnatal growth, functional behavior and reproductive ability of the
- offspring, at doses of 586 mg/kg/day or less.
- 397 Nursing Mothers: Low concentrations of ceftriaxone are excreted in human milk.
- 398 Caution should be exercised when Rocephin is administered to a nursing woman.
- 399 Pediatric Use: Safety and effectiveness of Rocephin in neonates, infants and pediatric
- 400 patients have been established for the dosages described in the DOSAGE AND
- 401 ADMINISTRATION section. In vitro studies have shown that ceftriaxone, like some
- 402 other cephalosporins, can displace bilirubin from serum albumin. Rocephin should not be
- 403 administered to hyperbilirubinemic neonates, especially prematures (see
- 404 **CONTRAINDICATIONS**).
- 405 ADVERSE REACTIONS: Rocephin is generally well tolerated. In clinical trials, the
- 406 following adverse reactions, which were considered to be related to Rocephin therapy or
- 407 of uncertain etiology, were observed:
- 408 LOCAL REACTIONS—pain, induration and tenderness was 1% overall. Phlebitis was
- 409 reported in <1% after IV administration. The incidence of warmth, tightness or induration
- 410 was 17% (3/17) after IM administration of 350 mg/mL and 5% (1/20) after IM
- 411 administration of 250 mg/mL.
- 412 HYPERSENSITIVITY—rash (1.7%). Less frequently reported (<1%) were pruritus, fever
- 413 or chills.
- 414 HEMATOLOGIC—eosinophilia (6%), thrombocytosis (5.1%) and leukopenia (2.1%).
- 415 Less frequently reported (<1%) were anemia, hemolytic anemia, neutropenia,
- 416 lymphopenia, thrombocytopenia and prolongation of the prothrombin time.
- 417 GASTROINTESTINAL—diarrhea (2.7%). Less frequently reported (<1%) were nausea or
- 418 vomiting, and dysgeusia. The onset of pseudomembranous colitis symptoms may occur
- during or after antibacterial treatment (see WARNINGS).
- 420 HEPATIC—elevations of SGOT (3.1%) or SGPT (3.3%). Less frequently reported
- 421 (<1%) were elevations of alkaline phosphatase and bilirubin.
- 422 RENAL—elevations of the BUN (1.2%). Less frequently reported (<1%) were elevations
- of creatinine and the presence of casts in the urine.
- 424 CENTRAL NERVOUS SYSTEM—headache or dizziness were reported occasionally
- 425 (<1%).

- 426 GENITOURINARY—moniliasis or vaginitis were reported occasionally (<1%).
- 427 MISCELLANEOUS—diaphoresis and flushing were reported occasionally (<1%).
- 428 Other rarely observed adverse reactions (<0.1%) include abdominal pain,
- 429 agranulocytosis, allergic pneumonitis, anaphylaxis, basophilia, biliary lithiasis,
- 430 bronchospasm, colitis, dyspepsia, epistaxis, flatulence, gallbladder sludge, glycosuria,
- 431 hematuria, jaundice, leukocytosis, lymphocytosis, monocytosis, nephrolithiasis,
- 432 palpitations, a decrease in the prothrombin time, renal precipitations, seizures, and serum
- 433 sickness.
- 434 Cases of fatal reactions with calcium-ceftriaxone precipitates in lung and kidneys in
- 435 neonates and prematures have been described. In some cases the infusion lines and times
- 436 of administration of ceftriaxone and calcium-containing solutions differed.
- 437 **OVERDOSAGE:** In the case of overdosage, drug concentration would not be reduced
- 438 by hemodialysis or peritoneal dialysis. There is no specific antidote. Treatment of
- 439 overdosage should be symptomatic.
- 440 **DOSAGE AND ADMINISTRATION:** Rocephin may be administered intravenously or
- 441 intramuscularly.
- 442 ADULTS: The usual adult daily dose is 1 to 2 grams given once a day (or in equally
- 443 divided doses twice a day) depending on the type and severity of infection. The total
- daily dose should not exceed 4 grams.
- 445 If Chlamydia trachomatis is a suspected pathogen, appropriate antichlamydial coverage
- should be added, because ceftriaxone sodium has no activity against this organism.
- 447 For the treatment of uncomplicated gonococcal infections, a single intramuscular dose of
- 448 250 mg is recommended.
- 449 For preoperative use (surgical prophylaxis), a single dose of 1 gram administered
- intravenously 1/2 to 2 hours before surgery is recommended.
- 451 PEDIATRIC PATIENTS: For the treatment of skin and skin structure infections, the
- 452 recommended total daily dose is 50 to 75 mg/kg given once a day (or in equally divided
- doses twice a day). The total daily dose should not exceed 2 grams.
- 454 For the treatment of acute bacterial otitis media, a single intramuscular dose of 50 mg/kg
- 455 (not to exceed 1 gram) is recommended (see INDICATIONS AND USAGE).
- 456 For the treatment of serious miscellaneous infections other than meningitis, the
- recommended total daily dose is 50 to 75 mg/kg, given in divided doses every 12 hours.
- 458 The total daily dose should not exceed 2 grams.
- 459 In the treatment of meningitis, it is recommended that the initial therapeutic dose be 100
- 460 mg/kg (not to exceed 4 grams). Thereafter, a total daily dose of 100 mg/kg/day (not to
- 461 exceed 4 grams daily) is recommended. The daily dose may be administered once a day
- 462 (or in equally divided doses every 12 hours). The usual duration of therapy is 7 to 14
- 463 days.

- Generally, Rocephin therapy should be continued for at least 2 days after the signs and symptoms of infection have disappeared. The usual duration of therapy is 4 to 14 days; in
- 466 complicated infections, longer therapy may be required.
- When treating infections caused by Streptococcus pyogenes, therapy should be continued
- 468 for at least 10 days.
- 469 No dosage adjustment is necessary for patients with impairment of renal or hepatic
- 470 function; however, blood levels should be monitored in patients with severe renal
- 471 impairment (eg, dialysis patients) and in patients with both renal and hepatic
- 472 dysfunctions.
- 473 DIRECTIONS FOR USE: Intramuscular Administration: Reconstitute Rocephin powder
- with the appropriate diluent (see **COMPATIBILITY AND STABILITY**).
- 475 Inject diluent into vial, shake vial thoroughly to form solution. Withdraw entire contents
- 476 of vial into syringe to equal total labeled dose.
- 477 After reconstitution, each 1 mL of solution contains approximately 250 mg or 350 mg
- 478 equivalent of ceftriaxone according to the amount of diluent indicated below. If required,
- 479 more dilute solutions could be utilized. A 350 mg/mL concentration is not
- 480 recommended for the 250 mg vial since it may not be possible to withdraw the entire
- 481 contents
- 482 As with all intramuscular preparations, Rocephin should be injected well within the body
- 483 of a relatively large muscle; aspiration helps to avoid unintentional injection into a blood
- 484 vessel.

Vial Dosage Size	Amount of Dilu	ent to be Added
	250 mg/mL	350 mg/mL
250 mg	0.9 mL	. —
500 mg	1.8 mL	1.0 mL
1 gm	3.6 mL	2.1 mL
2 gm	7.2 mL	4.2 mL

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- Intravenous Administration: Rocephin should be administered intravenously by infusion
- 487 over a period of 30 minutes. Concentrations between 10 mg/mL and 40 mg/mL are
- 488 recommended; however, lower concentrations may be used if desired. Reconstitute vials
- with an appropriate IV diluent (see **COMPATIBILITY AND STABILITY**).

Vial Dosage Size	Amount of Diluent to be Added
250 mg	2.4 mL
500 mg	4.8 mL
1 gm	9.6 mL
2 gm	19.2 mL

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After reconstitution, each 1 mL of solution contains approximately 100 mg equivalent of ceftriaxone. Withdraw entire contents and dilute to the desired concentration with the appropriate IV diluent.

493 COMPATIBILITY AND STABILITY: Rocephin sterile powder should be stored at room 494 temperature—77°F (25°C)—or below and protected from light. After reconstitution, 495 protection from normal light is not necessary. The color of solutions ranges from light 496 yellow to amber, depending on the length of storage, concentration and diluent used.

Rocephin *intramuscular* solutions remain stable (loss of potency less than 10%) for the following time periods:

		Storage		
Diluent	Concentration	Room Temp.	Refrigerated	
	mg/ml	(25°C)	(4°C)	
Sterile Water for Injection	100	2 days	10 days	
	250, 350	24 hours	3 days	
0.9% Sodium Chloride	100	2 days	10 days	
Solution	250, 350	24 hours	3 days	
5% Dextrose Solution	100	2 days	10 days	
	250, 350	24 hours	3 days	
Bacteriostatic Water + 0.9%	100	24 hours	10 days	
Benzyl Alcohol	250, 350	24 hours	3 days	
1% Lidocaine Solution	100	24 hours	10 days	
(without epinephrine)	250, 350	24 hours	3 days	

Rocephin *intravenous*-solutions, at concentrations of 10, 20 and 40 mg/mL, remain stable (loss of potency less than 10%) for the following time periods stored in glass or PVC containers:

	Storage		
Diluent	Room Temp. (25°C)	Refrigerated (4°C)	
Sterile Water	2 days	10 days	
0.9% Sodium Chloride Solution	2 days	10 days	
5% Dextrose Solution	2 days	10 days	
10% Dextrose Solution	2 days	10 days	
5% Dextrose + 0.9% Sodium Chloride Solution*	2 days	Incompatible	
5% Dextrose + 0.45% Sodium Chloride Solution	2 days	Incompatible	

*Data available for 10 to 40 mg/mL concentrations in this diluent in PVC containers only.

The following *intravenous* Rocephin solutions are stable at room temperature (25°C) for 24 hours, at concentrations between 10 mg/mL and 40 mg/mL: Sodium Lactate (PVC container), 10% Invert Sugar (glass container), 5% Sodium Bicarbonate (glass container), Freamine III (glass container), Normosol-M in 5% Dextrose (glass and PVC containers), Ionosol-B in 5% Dextrose (glass container), 5% Mannitol (glass container), 10% Mannitol (glass container).

- 509 Ceftriaxone has been shown to be compatible with Flagyl®* IV (metronidazole
- 510 hydrochloride). The concentration should not exceed 5 to 7.5 mg/mL metronidazole
- 511 hydrochloride with ceftriaxone 10 mg/mL as an admixture. The admixture is stable for 24
- hours at room temperature only in 0.9% sodium chloride injection or 5% dextrose in water (D5W). No compatibility studies have been conducted with the Flagyl[®] IV RTU[®]
- water (D3W). No comparishing studies have been conducted with the Flagyr TV RTO
- 514 (metronidazole) formulation or using other diluents. Metronidazole at concentrations
- 515 greater than 8 mg/mL will precipitate. Do not refrigerate the admixture as precipitation
- 516 will occur.
- * Registered trademark of G.D. Searle & Co.
- Vancomycin and fluconazole are physically incompatible with ceftriaxone in admixtures.
- 519 When either of these drugs is to be administered concomitantly with ceftriaxone by
- 520 intermittent intravenous infusion, it is recommended that they be given sequentially, with
- thorough flushing of the intravenous lines (with one of the compatible fluids) between the
- 522 administrations.
- 523 After the indicated stability time periods, unused portions of solutions should be
- 524 discarded.
- 525 NOTE: Parenteral drug products should be inspected visually for particulate matter
- 526 before administration.
- 527 Rocephin reconstituted with 5% Dextrose or 0.9% Sodium Chloride solution at
- 528 concentrations between 10 mg/mL and 40 mg/mL, and then stored in frozen state (-20°C)
- 529 in PVC or polyolefin containers, remains stable for 26 weeks. Reconstituted ADD-
- Vantage units, however, should not be stored in a frozen state (-20°C).
- 531 Frozen solutions of Rocephin should be thawed at room temperature before use. After
- thawing, unused portions should be discarded. **DO NOT REFREEZE**.
- Rocephin solutions should not be physically mixed with or piggybacked into solutions
- 534 containing other antimicrobial drugs or into diluent solutions other than those listed
- above, due to possible incompatibility (see **WARNINGS**).
- 536 ANIMAL PHARMACOLOGY: Concretions consisting of the precipitated calcium salt
- 537 of ceftriaxone have been found in the gallbladder bile of dogs and baboons treated with
- 538 ceftriaxone
- These appeared as a gritty sediment in dogs that received 100 mg/kg/day for 4 weeks. A
- 540 similar phenomenon has been observed in baboons but only after a protracted dosing
- 541 period (6 months) at higher dose levels (335 mg/kg/day or more). The likelihood of this
- occurrence in humans is considered to be low, since ceftriaxone has a greater plasma
- 543 half-life in humans, the calcium salt of ceftriaxone is more soluble in human gallbladder
- bile and the calcium content of human gallbladder bile is relatively low.
- 545 **HOW SUPPLIED:** Rocephin is supplied as a sterile crystalline powder in glass vials.
- 546 The following packages are available:

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- Vials containing 250 mg equivalent of ceftriaxone. Box of 1 (NDC 0004-1962-02) and
- 548 box of 10 (NDC 0004-1962-01).
- Vials containing 500 mg equivalent of ceftriaxone. Box of 1 (NDC 0004-1963-02) and
- 550 box of 10 (NDC 0004-1963-01).
- Vials containing 1 gm equivalent of ceftriaxone. Box of 1 (NDC 0004-1964-04) and box
- 552 of 10 (NDC 0004-1964-01).
- Vials containing 2 gm equivalent of ceftriaxone. Box of 10 (NDC 0004-1965-01).
- Bulk pharmacy containers, containing 10 gm equivalent of ceftriaxone. Box of 1 (NDC
- 555 0004-1971-01). NOT FOR DIRECT ADMINISTRATION.
- 856 Rocephin is also supplied as a sterile crystalline powder in ADD-Vantage®* Vials as
- 557 follows:
- 558 ADD-Vantage Vials containing 1 gm equivalent of ceftriaxone. Box of 10 (NDC 0004-
- 559 1964-05).
- 560 ADD-Vantage Vials containing 2 gm equivalent of ceftriaxone. Box of 10 (NDC 0004-
- 561 1965-05).
- NOTE: Rocephin sterile powder should be stored at room temperature, 77°F (25°C) or
- below, and protected from light.
- *Registered trademark of Abbott Laboratories, Inc.
- 565 CLINICAL STUDIES: Clinical Trials in Pediatric Patients With Acute Bacterial Otitis
- 566 Media: In two adequate and well-controlled US clinical trials a single IM dose of
- 567 ceftriaxone was compared with a 10 day course of oral antibiotic in pediatric patients
- between the ages of 3 months and 6 years. The clinical cure rates and statistical outcome
- 569 appear in the table below:

	Clinical Efficacy in Evaluable Population			
Study Day	Ceftriaxone Single Dose	Comparator – 10 Days of Oral Therapy	95% Confidence Interval	Statistical Outcome
Study 1 – US		amoxicillin/clavulanate		
14	74% (220/296)	82% (247/302)	(-14.4%, -0.5%)	Ceftriaxone is lower than control at
28	58% (167/288)	67% (200/297)	(-17.5%, -1.2%)	study day 14 and 28.
Study 2 - US ⁵	**	TMP-SMZ		· · · · · · · · · · · · · · · · · · ·
14	54% (113/210)	60% (124/206)	(-16.4%, 3.6%)	Ceftriaxone is equivalent to
28	35% (73/206)	45% (93/205)	(-19.9%, 0.0%)	control at study day 14 and 28.

570 An open-label bacteriologic study of ceftriaxone without a comparator enrolled 108

571 pediatric patients, 79 of whom had positive baseline cultures for one or more of the

572 common pathogens. The results of this study are tabulated as follows:

REFERENCES:

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Week 2 and 4 Bacteriologic Eradication Rates in the Per Protocol Analysis in the Roche Bacteriologic Study by pathogen:

	Study Day 13-15	-	Study Day 30+2	
Organism	No. Analyzed	No. Erad. (%)	No. Analyzed	No. Erad. (%)
Streptococcus pneumoniae	38	32 (84)	35	25 (71)
Haemophilus influenzae	33	28 (85)	31	22 (71)
Moraxella catarrhalis	15	12 (80)	15	9 (60)

576 577 578 579	1.	National Committee for Clinical Laboratory Standards, <i>Methods for Dilution Antimicrobial Susceptibility Tests for Bacteria that Grow Aerobically;</i> Approved Standard-Fifth Edition. NCCLS document M7-A5 (ISBN 1-56238-309-9). NCCLS, Wayne, PA 19087-1898, 2000.
580 581 582	2.	National Committee for Clinical Laboratory Standards, Supplemental Tables. NCCLS document M100-S10(M7) (ISBN 1-56238-309-9). NCCLS, Wayne, PA 19087-1898, 2000.
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592 Trimethoprim-Sulfamethoxazole for Acute Otitis Media. Pediatrics. Vol. 99, No. 1, January 1997.

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