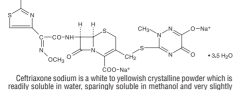
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Ceftriaxone for Injection, USP R_c only

To reduce the development of drug-resistant bacteria and maintain the effectiveness of ceftriaxone for injection, and other antibacterial drugs, ceftriaxone for injection should be used only to treat or prevent infections that are proven or strongly suspected to be caused by bacteria. DESCRIPTION

DESCRIPTION:
Ceftriaxone for injection, USP is a sterile, semisynthetic, broad-spectrum cephalosporin antibiotic for intravenous or intramuscular administration. Ceftriaxone sodium is (6*R*, 7*R*)-7-[2-(2-Amino-4-thiazolyl) glyoxylamido]-8-oxo-3-[[(1-2,5-6-terlaydhor-2-methyl-5-6-dioxo-as-triazin-3-y l)thio]methyl]-5-thia-1-azabicyclo[4.2 0]oct-2-ene-2-carboxylic acid, 7²-(2)-(0-methyloxime), disodium salt, sesquaterhydrate.
The chemical formula of ceftriaxone sodium is C₁₉H₁₆N₈N₈O₇S₃*3.5H₂O. It has a calculated molecular weight of 661.60 and the following structural formula:



Co-Na⁺
Ceftriaxone sodium is a white to yellowish 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 ceftriaxone sodium solutions ranges from light yellow to amber, depending on the length of storage, concentration and diluent used. Each vial contains ceftriaxone sodium equivalent to 250 mg, 500 mg, 1 gram or 2 grams of ceftriaxone activity. Ceftriaxone sodium contains approximately 83 mg (3.6 mEq) of sodium per gram of ceftriaxone activity. activity.

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 intramus-cular (IM) administration of a single 0.5 (250 mg/mL cor350 mg/mL con-centrations) or 1 gm dose in healthy subjects are presented in Table 1. Table 1. Ceftriaxone Plasma Concentrations After Single Dose Administration

| Average Plasma Concentrations (mcg/mL)
| Dose/Route | 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	
ND = Not dete	rmined.									
* IV doses we	* IV doses were infused at a constant rate over 30 minutes.									
Ceftriaxone	was com	plete	ly abs	orbed	follo	wing l	IM adm	inistrati	on with	

mean maximum plasma concentrations occurring between 2 and 3 hours post-dose. Multiple IV or IM doses ranging from 0.5 to 2 gm at 12- to 24-hour intervals resulted in 15% to 36% accumulation of ceftriaxone above single dose values.
Ceftriaxone concentrations in urine are shown in Table 2 Table 2. Urinary Concentrations of Ceftriaxone After Single Dose Administration

Average Urinary Concentrations (mcg/mL)

Dose/Route	0 to 2 hr	2 to 4 hr	4 to 8 hr	8 to 12 hr	12 to 24 hr	24 to 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
ND = Not det	ermined.					
Thirty-thre	anged drug	and the re	emainder w		in the b	oile and

utilimately found in the feces as microbiologically inactive compounds. After a 1 gm IV dose, average concentrations of ceftriaxone, determined from 1 to 3 hours after dosing, were 581 mcg/mL in the gallbladder bile, 788 mcg/mL in the common duct bile, 898 mcg/mL in the cystic duct bile, 78.2 mcg/mL in the plasma. Over a 0.15 to 3 gm dose range in healthy adult subjects, the values of elimination half-life ranged from 5.8 to 8.7 hours; apparent volume of distribution from 5.78 to 13.5 L; plasma clearance from 0.38 to 1.45 L/hour and renal clearance from 0.32 to 0.73 L/hour. Ceftriaxone is reversibly bound to human plasma proteins, and the binding decreased from a value of 95% bound at plasma concentrations of < 25 mcg/mL to a value of 85% bound at 300 mcg/mL. Ceftriaxone crosses the blood placenta harrier

barrier.

The average values of maximum plasma concentration, elimination halflife, plasma 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. Ceftriaxone penetrated the inflamed
meninges of infants and pediatric patients; CSF concentrations after a
50 mg/kg IV dose and after a 75 mg/kg IV dose are also shown in Table 3. Table 3. Average Pharmacokinetic Parameters of Cettriaxone in Pediatric Patients With Meningitis

50 mg/kg IV

75 mg/kg IV

	ou my/ky rv	/ 5 mg/kg rv
Maximum Plasma		
Concentrations (mcg/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 (mcg/mL)	5.6	6.4
Range (mcg/mL)	1.3 to 18.5	1.3 to 44
Time after dose (hr)	3.7 (± 1.6)	3.3 (± 1.4)
Compared to that in healthy adult sut triaxone were only minimally altered in a renal impairment or hepatic dysfunct adjustments are not necessary for these	elderly subjects ar tion (Table 4); t	nd in patients with herefore, dosage

adjustments are not necessary to these patients with certain according to the 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. Table 4. Average Pharmacokinetic Parameters of Ceftriaxone in Hum Elimination Plasma Volume of Distribution Clearance (L/hr) 0.58 to 1.45 Half-Life (L) 5.8 to 13.5 Subject Group

Healthy Subjects
Elderly Subjects
(mean age, 70.5 yr) 0.83 10.7 8.9

Patients With			
Renal Impairment			
Hemodialysis Patients			
(0 to 5 mL/min)*	14.7	0.65	13.7
Severe (5 to 15 mL/min)	15.7	0.56	12.5
Moderate (16 to 30 mL/min)	11.4	0.72	11.8
Mild (31 to 60 mL/min)	12.4	0.70	13.3
Patients With			
Liver Disease	8.8	1.1	13.6
* Creatinine clearance.			
The elimination of ceftriaxon administered with probenecid. Pharmacokinetics in the Middle		ed when cefti	riaxone is co-
In one study, total ceftriaxone c		(bound and u	nbound) were
measured in middle ear fluid obt	tained during	the insertion	of tympanos.
tomy tubes in 42 pediatric patient	ts with otitis r	nedia. Samplii	na times were

tomy 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 ceftriaxone. Mean (±SD) ceftriaxone levels in the middle ear reached a peak of 35 (±12) mcg/mL at 24 hours, and remained at 19 (±7) mcg/mL at 48 hours. Based on middle ear fluid cetriaxone concentrations in the 23 to 25 hour and the 46 to 50 hour sampling time intervals, a half-life of 25 hours was calculated. Cettriaxone is highly bound to plasma proteins. The extent of binding to proteins in the middle ear fluid is unknown. Interaction with Calcium:

Two in vitro studies, one using adult plasma and the other neonatal plasma from umbilical cord blood have been carried out to assess interplasma from umbilical cord blood have been carried out to assess inter-action of ceftriaxone and calcium. Ceftriaxone concentrations up to 1 mM (in excess of concentrations achieved *in vivo* following administration of 2 grams ceftriaxone infused over 30 minutes) were used in combination with calcium concentrations up to 12 mM (48 mg/dL). Recovery of cef-triaxone from plasma was reduced with calcium concentrations of 6 mM (24 mg/dL) or higher in adult plasma or 4 mM (16 mg/dL) or higher in neonatal plasma. This may be reflective of ceftriaxone-calcium precipitation. Microbiology: Microbiology: Mechanism of Action:

Ceftriaxone is a bactericidal agent that acts by inhibition of bacterial cell wall synthesis. Ceftriaxone has activity in the presence of some beta-lactamases, both penicillinases and cephalosporinases, of Gram-negative and Gram-positive bacteria. Mechanism of Resistance.

Resistance to ceftriaxone is primarily through hydrolysis by beta-lac-tamase, alteration of penicillin-binding proteins (PBPs), and decreased permeability. Interaction with Other Antimicrobials In an *in vitro* study antagonistic fects have been observed with the combination of chloramphenicol and ceftriaxone Ceftriaxone. Ceftriaxone has been shown to be active against most isolates of the following bacteria, both *in vitro* and in clinical infections as described in the INDICATIONS AND USAGE section:

Gram-negative bacteria
Acinetobacter calcoaceticus
Enterobacter aerogenes
Enterobacter cloacae Escherichia coli

Morganella morganii Neisseria gonorrhoeae Neisseria meningitidis Proteus mirabilis

Haemophilus influenzae Haemophilus parainfluenzae Klebsiella oxytoca Klebsiella pneumoniae Moraxella catarrhalis

Proteus vulgaris Pseudomonas aeruginosa Serratia marcescens Gram-positive bacteria Staphylococcus aureus Staphylococcus epidermidis

Staphylococcus epidermidis Streptococcus pneumoniae Streptococcus pyogenes Viridans group streptococci Anaerobic bacteria Bacteroides fragilis Clostridium species Peptostreptococcus species The following in vitro data a The following in vitro data are available, but their clinical signifi-cance is unknown. At least 90 percent of the following microorganisms exhibit an in vitro minimum inhibitory concentration (MIC) less than or equal to the susceptible breakpoint for certriaxone. However, the efficacy of certriaxone in treating clinical infections due to these microorgan-isms has not been established in adequate and well-controlled clinical trials.

Providencia species (including Providencia rettgeri) Salmonella species (including Salmonella typhi) Shigella species Gram-positive bacteria Streptococcus agalactiae Anaerobic bacteria
Porphyromonas (Bacteroides) melaninogenicus

Gram-negative bacteria Citrobacter diversus Citrobacter freundi

- Nuterioric Datcieria
Porphyromonas (Bacteroides) melaninogenicus
Prevotella (Bacteroides) bivius
Susceptibility Test Methods:
When available, the clinical microbiology laboratory should provide the results of in vitro susceptibility test results for antimicrobial drug products used in resident hospitals to the physician as periodic reports that describe the susceptibility profile of nosocomial and community-acquired pathogens. These reports should aid the physician in selecting an antibacterial drug product for treatment.

Dilution techniques: Quantitative methods are used to determine antimicrobial minimal inhibitory concentrations (MICs). These MICs provide estimates of the susceptibility of bacteria to antimicrobial compounds. The MICs should be determined using a standardized test method 1.3. The MIC values should be interpreted according to criteria provided in Table 5.

Diffusion techniques: Quantitative methods that require measurement

Diffusion techniques: Quantitative methods that require measurement of zone diameters also provide reproducible estimates of the susceptibility of bacteria to antimicrobial compounds. The zone size provides an estimate of the susceptibility of bacteria to antimicrobial compounds. The zone size should be determined using a standardized test method 2.3. This procedure uses paper disks impregnated with 30 mcg ceftriaxone to test the susceptibility of microorganisms to ceftriaxone. The disk diffusion interpretive criteria are provided in Table 5.

Anaerobic techniques: For anaerobic bacteria, the susceptibility of ceffriaxone as MICs can be determined by a standardized agar test method 3.4. The MIC values obtained should be interpreted according to the criteria provided in Table 5.

1

Table 5. Susceptibility Test Interpretive Criteria for Ceftriaxone. Minimum Inhibitory Disk Diffusion Zone

Pathogen		centration (mcg/ml)	ns	Diameters (mm)			
	(S) Sus- ceptible	(I) Inter- mediate	(R) Re- sistant	(S) Sus- ceptible	(I) Inter- mediate	(R) Re- sistant	
Enterobacteriaceae	≤ 1	2	≥4	≥ 23	20 to 22	≤19	
Haemophilus influenzae*	≤2	-	-	≥26	-	-	
Neisseria gonor- rhoeae*	≤ 0.25	-	-	≥ 35	-	-	
Neisseria meningi- tidis*	≤ 0.12	-	-	≥ 34	-	-	
Streptococcus pneumoniae† meningitis isolates	≤ 0.5	1	≥ 2	-	_	-	
Streptococcus pneumoniae† non- meningitis isolates	≤1	2	≥4	-	-	-	
Streptococcus species beta- hemolytic group*	≤0.5	-	-	≥ 24	-	-	
Viridans group streptococci	≤ 1	2	≥ 4	≥27	25 to 26	≤24	
Anaerobic bacteria	<16	32	>64	_	_	_	

[agar method] ≤16 32 ≥64 - - - Susceptibility of staphylococci to ceftriaxone may be deduced from testing only penicillin and either cefoxitin or oxacillin.

* The current absence of data on resistant isolates precludes defining any category other than 'Susceptible'. If isolates yield MIC results other than susceptible, they should be submitted to a reference laboratory for additional testing.

* Disc diffusion interpretive criteria for ceftriaxone discs against *Streptococcus pneumoniae* are not available, however, isolates of pneumococci with oxacillin zone diameters of ≥20 mm are susceptible (MIC s 0.06 mcg/mL) to penicillin and can be considered susceptible to ceftriaxone. *Streptococcus pneumoniae* isolates should not be reported as penicillin (ceftriaxone) resistant or intermediate base solely on an oxacillin zone diameter of ≤ 19 mm. The ceftriaxone MIC should be determined for those isolates with oxacillin zone diameters ≤ 19 mm.

A report of *Susceptible* indicates that the antimicrobial is likely to

A report of Susceptible indicates that the antimicrobial is likely to inhibit growth of the pathogen if the antimicrobial compound reaches the concentration at the infection site necessary to inhibit growth of the pathogen. A report of Intermediate indicates that the result should be considered equivocal, and if the mitcroorganism is not fully susceptible to alternative, clinically feasible drugs, the test should be repeated. This category implies possible clinical applicability in body sites where the drug is physiologically concentrated or in situations where a high dosage of drug can be used. This category also provides a buffer zone that prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of Resistant indicates that the antimicrobial is not likely to inhibit growth of the pathogen if the antimicrobial compound reaches the concentrations usually achievable at the infection site; other therapy should be selected.

Quality Control: Standardized susceptibility test procedures require the use of laboratory controls to monitor and ensure the accuracy and precision of supplies and reagents used in the assay, and the techniques of the individual performing the test1.2.3.4. Standard ceftriaxone powder should provide the following range of MIC values noted in Table 6. For the diffusion technique using the 30 mcg disk, the criteria in Table 6 should be achieved.

Table 6. Acceptable Quality Control Ranges for Ceftriaxone

Table 6. Acceptable Quality Control Ranges for Ceftriaxone Minimum Diff

QC Strain	Concentrations (mcg/mL)	Zone diam- eters (mm)
Escherichia coli ATCC 25922	0.03 to 0.12	29 to 35
Staphylococcus aureus ATCC 25923		22 to 28
Staphylococcus aureus ATCC 29213	1 to 8	
Haemophilus influenzae ATCC 49247	0.06 to 0.25	31 to 39
Neisseria gonorrhoeae ATCC 49226	0.004 to 0.015	39 to 51
Pseudomonas aeruginosa ATCC 27853	8 to 64	17 to 23
Streptococcus pneumoniae ATCC 49619	0.03 to 0.12	30 to 35
Bacteroides fragilis ATCC 25285 (agar method)	32 to 128	
Bacteroides thetaiotaomicron ATCC 29741 (agar method)	64 to 256	
INDICATIONS AND USAGE:	avone annronria	ta enaciman

INDICATIONS AND USAGE:
Before instituting treatment with ceftriaxone, appropriate specimens should be obtained for isolation of the causative organism and for determination of its susceptibility to the drug. Therapy may be instituted prior to obtaining results of susceptibility testing.

To reduce the development of drug-resistant bacteria and maintain the effectiveness of ceftriaxone for injection, USP and other antibacterial drugs, ceftriaxone for injection, USP should be used only to treat or prevent infections that are proven or strongly suspected to be caused by susceptible bacteria. When culture and susceptibility information are available, they should be considered in selecting or modifying antibacterial therapy. In the absence of such data, local epidemiology and susceptibility patterns may contribute to the empiric selection of therapy. Ceftriaxone for injection, USP is indicated for the treatment of the following infections when caused by susceptible organisms:

Lower Respiratory Tract Infections:

caused by Streptococcus pneumoniae, Staphylococcus aureus, Haemophilus influenzae, Haemophilus parainfluenzae, Klebsiella pneumoniae, Escherichia coli, Enterobacter aerogenes, Proteus mirabilis or Serratia marcescens.

Acute Bacterial Otitis Media:

caused by Streptococcus pneumoniae, Haemophilus influenzae (including beta-lactamase producing strains) or Moraxella catarrhalis (including beta-lactamase producing strains).

NOTE: In one study lower clinical cure rates were observed with a single dose of ceftriaxone comparator. The potentially lower clinical cure rate of ceftriaxone should be balanced against the potential advantages of parenteral therapy (see CLINICAL STUDIES).

Skin and Skin Structure Infections:

caused by Staphylococcus aureus, Staphylococcus epidermidis, Streptococcus pyogenes, Viridans group streptococci, Escherichia coli, Enterobacter cloacae, Klebsiella oxytoca, Klebsiella pneumoniae, Proteus mirabilis, Morganella morganii or Klebsiella oxytoca, Klebsiella pneumoniae, eaused by Staphyl

Escriericnia coli, Haemophiuis miuenzae of Kieosiella preumoniae.

Bone and Joint Infections:
caused by Staphylococcus aureus, Streptococcus pneumoniae,
Escharichia coli, Proteus mirabilis, Klebsiella pneumoniae or Enterobacter

species.

Intra-abdominal Infections:
caused by Escherichia coli, Klebsiella pneumoniae, Bacteroides fragilis, Clostridium species (Note: most strains of Clostridium difficile are resistant) or Peptostreptococcus species.

caused by Escherichia coli, Klebsiella pneumoniae, Bacteroides fragilis, Clostridium species (Note: most strains of Clostridium difficile are resistant) or Peptostreptococcus species.

Meningitis:
caused by Haemophilus influenzae, Neisseria meningitidis or Streptococcus pneumoniae. Cettriaxone has also been used successfully in a limited number of cases of meningitis and shunt infection caused by Staphylococcus epidermidis* and Escherichia coli.*

Surgical Prophylaxis:
The preoperative administration of a single 1 gm dose of cettriaxone may reduce the incidence of postoperative infections in patients undergoing surgical procedures classified as contaminated or potentially contaminated (e.g., vaginal or abdominal hysterectomy or cholecystectomy for chronic calculous cholecystitis in high-risk patients, such as those over 70 years of age, with acute cholecystitis not requiring therapeutic antimicrobials, obstructive jaundice or common duct bile stones) and in surjical patients for whom infection at the operative site would present serious risk (e.g., during coronary artery bypass surgery). Although ceftriaxone has been shown to have been as effective as cefazolin in the prevention of infection following coronary artery bypass surgery. When administered prior to surgical procedures for which it is indicated, a single 1 gm dose of ceftriaxone provides protection from most infections due to susceptible organisms throughout the course of the procedure.

Efficacy for this organism in this organ system was studied in fewer Infections due to susceptible organisms throughout the course of the procedure.

* Efficacy for this organism in this organ system was studied in fewer than ten infections.

CONTRAINDICATIONS:
Ceftriaxone for injection is contraindicated in patients with known allergy to the cephalosporin class of antibiotics.

Neonates (<28 days):
Hyperbillirubinemic neonates, especially prematures, should not be treated with ceftriaxone for injection. In vitro studies have shown that ceftriaxone and isplace bilirubin from its binding to serum albumin, leading to a possible risk of bilirubin encephalopathy in these patients.

Ceftriaxone is contraindicated in neonates if they require (or are expected to require) treatment with calcium-containing IV solutions, including continuous calcium-containing infusions such as parenteral nutrition because of the risk of precipitation of ceftriaxone-calcium (see CLINICAL PHARMACOLOGY, WARNINGS and DOSAGE AND ADMINISTRATION).

A small number of cases of fatal outcomes in which a crystalline material was observed in the lungs and kidneys at autopsy have been reported in neonates receiving ceftriaxone and calcium-containing fluids. In some of these cases, the same intravenous infusion line was used for both ceftriaxone and calcium-containing fluids. In some of these cases, the same intravenous infusion line was used for both ceftriaxone and calcium-containing fluids were administered at different time points via different intravenous lines; no crystalline material was observed at autopsy in this neonate. There have been no similar reports in patients other than neonates.

WARNINGS:

Hypersensitivity:

BEFORE THERAPY WITH CEFTRIAXONE IS INSTITUTED, CAREFUL INOUINY SHOULD BE MADE TO DETERMINE WHETHER THE PATIENT THAS HAD PREVIOUS HYPERSENSITIVITY PREATIONS TO CEPHALOSPORINS, PENICILLINS OR OTHER DRUGS. THIS PRODUCT SHOULD BE GIVEN CAUTIOUSLY TO PENICILLIN-SENSITIVE PATIENTS. ANTIBIOTICS SHOULD BE ADMINISTERED WITH CAUTION TO ANY PATIENT WHO HAS DEMONSTRATED SOME FORM OF ALLERGY, PARTICULARLY T procedure.

* Efficacy for this organism in this organ system was studied in fewer

have been reported, even if a patient is not known to be allergic or previously exposed. Interaction with Calcium-Containing Products:

Do not use diluents containing aclcium, such as Ringer's solution or Hartmann's solution, to reconstitute ceftriaxone vials or to further dilute a reconstituted vial for IV administration because a precipitate can form. Precipitation of ceftriaxone-calcium can also occur when ceftriaxone is mixed with calcium-containing solutions in the same IV administration line. Ceftriaxone must not be administered simultaneously with calcium-containing IV solutions, including continuous calcium-containing infusions such as parenteral nutrition via a Y-site. However, in patients other than neonates, ceftriaxone and calcium-containing solutions may be administered sequentially of one another if the infusion lines are thoroughly flushed between infusions with a compatible fluid. In vitro studies using adult and neonatal plasma from umbilical cord blood demonstrated that neonates have an increased risk of precipitation of ceftriaxone-calcium (see CLINICAL PHARMACOLOGY, CONTRAINDICATIONS and DOSAGE AND ADMINISTRATION).

Clostridium difficile:

PHARMACOLOGY, CONTRAINDICATIONS and DOSAGE AND ADMINISTRATION).

Clostridium difficile:

Clostridium difficile associated diarrhea (CDAD) has been reported with use of nearly all antibacterial agents, including ceftriaxone, and may range in severity from mild diarrhea to fatal collist. Treatment with antibacterial agents alters the normal flora of the colon leading to overgrowth of C. difficile produces toxins A and B which contribute to the development of CDAD. Hypertoxin producing strains of C. difficile cause increased morbidity and mortality, as these infections can be refractory to antimicrobial therapy and may require colectomy. CDAD must be considered in all patients who present with diarrhea following antibiotic use. Careful medical history is necessary since CDAD has been reported to occur over two months after the administration of antibacterial agents.

If CDAD is suspected or confirmed, ongoing antibiotic use not directed against C. difficile may need to be discontinued. Appropriate fluid and electrolyte management, protein supplementation, antibiotic treatment of C. difficile and surgical evaluation should be instituted as clinically indicated.

Hemolytic Anemia:

An immune mediated hemolytic anemia has been observed in patients receiving cephalosporin class antibacterials including ceftriaxone. Severe cases of hemolytic anemia, including fatalities, have been reported during treatment in both adults and children. If a patient develops anemia while on ceftriaxone, the diagnosis of a cephalosporin associated anemia should be considered and ceftriaxone stopped until the etiology is determined.

PRECAUTIONS:

PRECAUTIONS:
General:
Prescribing ceftriaxone for injection in the absence of a proven or strongly suspected bacterial infection or a prophylactic indication is unlikely to provide benefit to the patient and increases the risk of the development of drug-resistant bacteria.

Although transient elevations of BUN and serum creatinine have been observed, at the recommended dosages, the nephrotoxic potential of ceftriaxone is similar to that of other cephalosporins.
Ceftriaxone is excreted via both biliary and renal excretion (see CLIN-ICAL PHARMACOLOGY). Therefore, patients with renal failure normally require no adjustment in dosage when usual doses of ceftriaxone are administered.

require no adjustment in dosage when usual doses of ceftriaxone are administered.

Dosage adjustments should not be necessary in patients with hepatic dysfunction; however, in patients with both hepatic dysfunction and significant renal disease, caution should be exercised and the ceftriaxone dosage should not exceed 2 gm daily.

Alterations in prothrombin times have occurred rarely in patients treated with ceftriaxone. Patients with impaired vitamin K synthesis or low vitamin K stores (e.g., chronic hepatic disease and malnutrition) may require monitoring of prothrombin time during ceftriaxone treatment. Vitamin K administration (10 mg weekly) may be necessary if the prothrombin time is prolonged before or during therapy.

Prolonged use of ceftriaxone may result in overgrowth of nonsusceptible organisms. Careful observation of the patient is essential. If superinfection occurs during therapy, appropriate measures should be taken. Ceftriaxone for injection should be prescribed with caution in individuals with a history of gastrointestinal disease, especially colitis.

There have been reports of sonographic abnormalities in the gall-bladder of patients treated with ceftriaxone; some of these patients also had symptoms of gallbladder disease. These abnormalities appear on sonography as an echo without acoustical shadowing suggesting sludge or as an echo with acoustical shadowing be misinterpreted as (See Reverse)

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(See Reverse)



























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(Continued)

gallstones. The chemical nature of the sonographically detected material has been determined to be predominantly a ceftriaxone-calcium salt. The condition appears to be transient and reversible upon discontinuation of ceftriaxone for injection and institution of conservative management. Therefore, ceftriaxone should be discontinued in patients who develop signs and symptoms suggestive of gallbladder disease and/or the sonographic findings described above.

Cases of pancreatitis, possibly secondary to biliary obstruction, have been reported rarely in patients treated with ceftriaxone. Most patients presented with risk factors for biliary stasis and biliary sludge (preceding major therapy, severe illness, total parenteral nutrition). A cofactor role of ceftriaxone-related biliary precipitation cannot be ruled out.

Information for Patients:

Patients should be counseled that antibacterial drugs including ceftriaxone for injection should only be used to treat bacterial infections. They do not treat viral infections (e.g., the common cold). When ceftriaxone for injection is prescribed to treat a bacterial infection, patients should be told that although it is common to feel better early in the course of therapy, the medication should be taken exactly as directed. Skipping doses or not completing the full course of therapy may (1) decrease the effectiveness of the immediate treatment and (2) increase the likelihood that bacteria will develop resistance and will not be treatable by ceftriaxone for injection or other antibacterial drugs in the future.

Diarrhea is a common problem caused by antibiotics which usually ends when the antibiotic is discontinued. Sometimes after starting treatment with antibiotics, patients 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.

Carcinogenesis, Mutagenesis, Impairment of Fertility:

Carci

compound, carcinogenicity studies with ceftriaxone in animals have not been performed. The maximum duration of animal toxicity studies was Genetic toxicology tests included the Ames test, a micronucleus test and a test for chromosomal aberrations in human lymphocytes cultured in vitro with ceftriaxone. Ceftriaxone showed no potential for mutagenic

in value with central vote. Showed no potential for intragenic activity in these studies.
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.

Pregnancy:

Teratogenic Effects:

Reproductive studies have been performed in mice and rats at doses up to 20 times the usual human dose and have no evidence of embryotoxicity, fetotoxicity or teratogenicity. In primates, no embryotoxicity or teratogenicity was demonstrated at a dose approximately 3 times the human dose. There are, however, no adequate and well-controlled studies in preg-nant women. Because animal reproductive studies are not always predictive of human response, this drug should be used during pregnancy only if

clearly needed. Nonteratogenic Effects:

In rats, in the Segment I (fertility and general reproduction) and Segment III (perinatal and postnatal) studies with intravenously administered ceftriaxone, no adverse effects were noted on various reproductive parameters during gestation and lactation, including postnatal growth, functional behavior and reproductive ability of the offspring, at doses of 586 mg/kg/day or less.

Nursing Nothers:

Low conceptrations of ceftriaxone are excreted in human milk. Caution

Low concentrations of ceftriaxone are excreted in human milk. Caution should be exercised when ceftriaxone is administered to a nursing

should be exercised when ceftriaxone is administered to a nursing woman.

Pediatric Use:

Safety and effectiveness of ceftriaxone in neonates, infants and pediatric patients have been established for the dosages described in the DOSAGE AND ADMINISTRATION section. In vitro studies have shown that ceftriaxone, like some other cephalosporins, can displace bilirubin from serum albumin. Ceftriaxone should not be administered to hyperbilirubinemic neonates, especially prematures (see CONTRAINDICATIONS).

Geriatric Use:

Of the total number of subjects in clinical studies of ceftriaxone, 32% were 60 and over. No overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients, but greater sensitivity of some older individuals cannot be ruled out.

The pharmacokinetics of ceftriaxone were only minimally altered in geriatric patients compared to healthy adult subjects and dosage adjustments are not necessary for geriatric patients with ceftriaxone dosages up to 2 grams per day (see CLINICAL PHARMACOLOGY).

ADVERSE REACTIONS:

Ceftriaxone is generally well tolerated. In clinical trials, the following adverse reactions, which were considered to be related to ceftriaxone therapy or of uncertain etiology, were observed:

Local Reactions:

pain, induration and tenderness was 1% overall. Phlebitis was reported in <1% after IV administration. The incidence of warmth, tightness or induration was 17% (3/17) after IM administration of 350 mg/mL.

Hypersensitivity:

rash (1.7%). Less frequently reported (<1%) were pruritus, fever or

(1/20) after IM administration. C. 222

Hypersensitivity:
rash (1.7%). Less frequently reported (<1%) were pruritus, fever or Hematologic: eosinophilia (6%), thrombocytosis (5.1%) and leukopenia (2.1%). Less frequently reported (<1%) were anemia, hemolytic anemia, neutropenia, lymphopenia, thrombocytopenia and prolongation of the pro-

thrombin time. **Gastrointestinal**: diarrhea (2.7%). Less frequently reported (<1%) were nausea or vom-iting, and dysgeusia. The onset of pseudomembranous colitis symp-toms may occur during or after antibacterial treatment (see WARNINGS). elevations of SGOT (3.1%) or SGPT (3.3%). Less frequently reported (<1%) were elevations of alkaline phosphatase and bilirubin.

Renal:

Renal:
elevations of the BUN (1.2%). Less frequently reported (<1%) were elevations of creatinine and the presence of casts in the urine.

Central Nervous System:
headache or dizziness were reported occasionally (<1%).

Genitourinary:
moniliasis or vaginitis were reported occasionally (<1%).

Miscellaneus:

Miscellaneous:

Miscellaneous:

Miscellaneous:
diaphoresis and flushing were reported occasionally (<1%).
Miscellaneous:
diaphoresis and flushing were reported occasionally (<1%).
Other rarely observed adverse reactions (<0.1%) include abdominal pain, agranulocytosis, allergic pneumonitis, anaphylaxis, basophilia, biliary lithiasis, bronchospasm, colitis, dyspepsia, epistaxis, flatulence, gall-bladder sludge, glycosuria, hematuria, jaundice, leukocytosis, lymphocytosis, monocytosis, nephrolithiasis, palpitations, a decrease in the prothrombin time, renal precipitations, seizures, and serum sickness.
Postmarketing Experience:
In addition to the adverse reactions reported during clinical trials, the following adverse experiences have been reported during clinical practice in patients treated with ceftriaxone. Data are generally insufficient to allow an estimate of incidence or to establish causation.

A small number of cases of fatal outcomes in which a crystalline material was observed in the lungs and kidneys at autopsy have been reported in neonates receiving ceftriaxone and calcium-containing fluids. In some of these cases, the same intravenous infusion line was used for oth ceftriaxone and calcium-containing fluids were administered at different time points via different intravenous lines; no crystalline material was observed at autopsy in this neonate. There have been no similar reports in patients other than neonates.

Gastointestinal:

Gastrointestinal stomatitis and glossitis Genitourinary: oliguria. Dermatologic:

exanthema, allergic dermatitis, urticaria, edema. As with many med-ications, isolated cases of severe cutaneous adverse reactions (erythema multiforme, Stevens-Johnson syndrome or Lyell's syndrome/toxic epidermal necrolysis) have been reported.

Cephalosporin Class Adverse Reactions:
In addition to the adverse reactions listed above which have been observed in patients treated with ceftriaxone, the following adverse reactions and altered laboratory test results have been reported for considering adverse reactions.

cephalosporin class antibiotics:

**Adverse Reactions:*
Allergic reactions, drug fever, serum sickness-like reaction, renal dysfunction, toxic nephropathy, reversible hyperactivity, hypertonia, hepatic dysfunction including cholestasis, aplastic anemia, hemorrhage, and superinderities. dysfunction including cholestasis, aplastic anemia, hemorrhage, and superinfection.

Altered Laboratory Tests: Positive direct Coombs' test, false-positive test for urinary glucose, and elevated LDH.

Several cephalosporins have been implicated in triggering seizures, particularly in patients with renal impairment when the dosage was not reduced (see DOSAGE AND ADMINISTRATION). If seizures associated with drug therapy occur, the drug should be discontinued. Anticonvulsant therapy can be given if clinically indicated.

OVERIOSAGE:

In the case of overdosage, drug concentration would not be reduced.

In the case of overdosage, drug concentration would not be reduced by hemodialysis or peritoneal dialysis. There is no specific antidote. Treatment of overdosage should be symptomatic. DOSAGE AND ADMINISTRATION:

Ireatment of overdosage should be symptomatic.

DOSAGE AND ADMINISTRATION:

Ceftriaxone may be administered intravenously or intramuscularly.

Do not use diluents containing calcium, such as Ringer's solution

or Hartmann's solution, to reconstitute ceftriaxone vials or to further
dilute a reconstituted vial for IV administration because a precipitate
can form. Precipitation of ceftriaxone-calcium can also occur when ceftriaxone is mixed with calcium-containing solutions in the same IV
administration line. Ceftriaxone must not be administered simultaneously with calcium-containing IV solutions, including continuous calcium-containing infusions such as parenteral nutrition via a Y-site.
However, in patients other than neonates, ceftriaxone and calcium-containing solutions may be administered sequentially of one another if
the infusion lines are thoroughly flushed between infusions with a
compatible fluid (see WARNINGS).

There have been no reports of an interaction between ceftriaxone and
oral calcium-containing products or interaction between intramuscular ceftriaxone and calcium-containing products (IV or oral).

Neonates:

Hyperhility binemic peopates, especially prematures, should not be **Neonates:** Neonates:
Hyperbilirubinemic neonates, especially prematures, should not be treated with ceftriaxone for injection (see CONTRAINDICATIONS).
Ceftriaxone is contraindicated in neonates if they require (or are expected to require) treatment with calcium-containing IV solutions, including continuous calcium-containing infusions such as parenteral nutrition because of the risk of precipitation of ceftriaxone-calcium (see CONTRAINDICATIONS).
Pediatric Patients:

Pediatric Patients

14 days. Adults:

Directions for Use: Intramuscular Administration

Pediatric Patients:
For the treatment of skin and skin structure infections, the 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. For the treatment of acute bacterial otitis media, a single inframuscular dose of 50 mg/kg (not to exceed 1 gram) is recommended (see INDICATIONS AND USAGE).
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. The total daily dose should not exceed 2 grams. grams.

In the treatment of meningitis, it is recommended that the initial therapeutic dose be 100 mg/kg (not to exceed 4 grams). Thereafter, a total daily dose of 100 mg/kg/day (not to exceed 4 grams daily) is recommended. The daily dose may be administered once a day (or in equally divided doses every 12 hours). The usual duration of therapy is 7 to

14 days.

Adults:

The usual adult daily dose is 1 to 2 grams given once a day (or in equally divided doses twice a day) depending on the type and severity of infection. For infections caused by Staphylococcus aureus (MSSA), the recommended daily dose is 2 to 4 grams, in order to achieve >90% target attainment. The total daily dose should not exceed 4 grams.

If Chlamydia trachomatis is a suspected pathogen, appropriate antichlamydial coverage should be added, because ceftriaxone sodium has no activity against this organism.

For the treatment of uncomplicated gonococcal infections, a single intramuscular dose of 250 mg is recommended.

For preoperative use (surgical prophylaxis), a single dose of 1 gram administered intravenously 1/2 to 2 hours before surgery is recommended. Generally, ceftriaxone 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 complicated infections, longer therapy should be continued for at least 10 days.

No dosage adjustment is necessary for patients with impairment of renal or hepatic function.

Directions for Use:

Intramuscular Administration:
Reconstitute cettriaxone sodium powder with the appropriate diluent (see DOSAGE AND ADMINISTRATION: Compatibility and Stability). Inject diluent into vial, shake vial thoroughly to form solution. Withdraw entire contents of vial into syringe to equal total labeled dose.

After reconstitution, each 1 mL of solution contains approximately 250 mg or 350 mg equivalent of cettriaxone according to the amount of diluent indicated below. If required, more dilute solutions could be utilized. A 350 mg/mL concentration is not recommended for the 250 mg vial since it may not be possible to withdraw the entire contents. As with all intramuscular preparations, cettriaxone should be injected well within the body of a relatively large muscle; aspiration helps to avoid unintentional injection into a blood vessel. Amount of Diluent to be Added

Vial Dosage Size 250 mg/mL 350 mg/mL 250 mg 0.9 m 500 mg 1.8 ml 1 ml 1 gm 3.6 m 2.1 ml 4.2 ml 2 gm Intravenous Administration: Intravenous Administration:
Ceftriaxone should be administered intravenously by infusion over a period of 30 minutes. Concentrations between 10 mg/mL and 40 mg/mL are recommended; however, lower concentrations may be used if desired. Reconstitute vials with an appropriate IV diluent (see DOSAGE AND ADMINISTRATION: Compatibility and Stability).

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

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.

Compatibility and Stability:
Ceftriaxone has been shown to be compatible with Flagyl® IV (metronidazole hydrochloride). The concentration should not exceed 5 to 7.5 mg/mL metronidazole 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® (metronidazole) formulation or using other diluents. Metronidazole at concentrations greater than 8 mg/mL will precipitate. Do not refrigerate the admixture as precipitation will occur.

Vancomycin, amsacrine, aminoglycosides, and fluconazole are physically incompatible with ceftriaxone in admixtures. When any of these drugs

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are to be administered concomitantly with ceftriaxone by intermittent intra-venous infusion, it is recommended that they be given sequentially, with thorough flushing of the intravenous lines (with one of the compatible fluids) between the administrations.

Do not use diluents containing calcium, such as Ringer's solution or Hartmann's solution, to reconstitute ceftriaxone for injection or to further dilute a reconstituted vial for IV administration. Particulate for-

further dilute a reconstituted vial for IV administration. Particulate formation can result.

Ceftriaxone for injection solutions should not be physically mixed with or piggybacked into solutions containing other antimicrobial drugs or into diluent solutions other than those listed above, due to possible incompatibility (see WARNINGS).

Ceftriaxone sodium sterile powder should be stored at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature] and protected from light. After reconstitution, protection from normal light is not necessary. The color of solutions ranges from light yellow to amber, depending on the length of storage, concentration and diluent used.

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

| Concentration | St | Room Temp

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

time periods stored in glass or PVC containers. Storage Refrigerated Room Temp.

Diluent	(25 0)	(4 6)
Sterile Water	2 days	10 days
0.9% Sodium	0.1.	40.1
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 containers only. 	concentrations in	this diluent in PV
The following intravenous ceftria		

containers only.

The following intravenous ceftriaxone 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), S% Sodium Bicarbonate (glass container), Freamine III (glass container), Normosol-M in 5% Dextrose (glass and PVC containers), 10nosol-B in 5% Dextrose (glass container), 5% Mannitol (glass container), 10% Mannitol (glass container), 10% Mannitol (glass container), After the indicated stability time periods, unused portions of solutions should be discarded.

NOTE: Parenteral drup products should be inspected visually for particulate matter before administration.

Ceftriaxone reconstituted with 5% Dextrose or 0.9% Sodium Chloride solution at concentrations between 10 mg/mL and 40 mg/mL, and then stored in frozen state (-20°C) in PVC or polyolefin containers, remains stable for 26 weeks.

Frozen solutions of ceftriaxone for injection should be thawed at room temperature before use. After thawing, unused portions should be discarded. DO NOT REFREEZE.

ANIMAL PHARMACOLOGY:

Concretions consisting of the precipitated calcium salt of ceftriaxone have been found in the gallbladder bile of dogs and baboons treated with ceftriaxone.

These appeared as a gritty sediment in dogs that received 100 mg/kg/day for 4 weeks. A similar phenomenon has been observed in baboons brouly after a protracted dosing 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 half-life in humans, the calcium salt of ceftriaxone is more soluble in human gall-bladder bile and the calcium content of human gallbladder bile is relatively low.

Ceftriaxone for injection LISD is expelied as a stable caractalizate.

in humans, the calcium salt of certriaxone is more soluble in numan gall-bladder bile and the calcium content of human gallbladder bile is relatively low.

HOW SUPPLIED:
Ceftriaxone for injection, USP is supplied as a sterile crystalline powder in glass vials. The following packages are available:
Vials containing 250 mg equivalent to ceftriaxone. Package of 10 (0781-3206-95).
Vials containing 500 mg equivalent to ceftriaxone. Package of 10 (0781-3207-95).
Vials containing 1 gm equivalent to ceftriaxone. Package of 10 (0781-3208-95).
Vials containing 2 gm equivalent to ceftriaxone. Package of 10 (0781-3208-95).
Vials containing 250 mg equivalent to ceftriaxone. Package of 10 (0781-3208-95).
Vials containing 250 mg equivalent to ceftriaxone. Package of 1 (0781-3208-85).
Vials containing 1 gm equivalent to ceftriaxone. Package of 1 (0781-3208-85).
Storage Prior to Reconstitution:
Store at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature]. Protect from light.
CLINICAL STUDIES:
Clinical Trials in Pediatric Patients With Acute Bacterial Otitis Media: In two adequate and well-controlled US clinical trials a single IM dose of 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 appear in the table below.

Clinical Efficacy in Evaluable Population

Ceftriaxone Comparator- 95% Confidence Oral Therapy Interval Statistical Outcome Study Day Study 1– US amoxicillin Ceftriaxone is

		Ciavulaliato		Hower than
14	74% (220/296)	82% (247/302)	(-14.4%, -0.5%)	control at study day 14
28	58% (167/288)	67% (200/297)	(-17.5%, -1.2%)	and 28.
Study 2 – US ⁵		TMP-SMZ		Ceftriaxone is
14	54% (113/210)	60% (124/206)	(-16.4%, 3.6%)	equivalent to control at
28	35% (73/206)	45% (93/205)	(-19.9%, 0.0%)	study day 14 and 28.
enrolled 108 ped	diatric patients of the comm	s, 79 of whom I	nad positive	out a comparato baseline cultures of this study are
		ic Eradication e Bacteriologi		e Per Protocol Pathogen

Study Day 13 to 15 Study Day 30+2 Organism Analyzed 35 Analyzed (%) 25 (71) (%) 32 (84) eptococcus pneumonia

Moraxella catarrhalis	15	12 (80)	15	9 (60)
REFERENCES: 1. Clinical and Laboratory S. Antimicrobial Susceptib. Approved Standard - Nii and Laboratory Standard Wayne, Pennsylvania 19 2. Clinical and Laboratory Standards for Antimic. Informational Suppleme ment M100-S23, Clinic West Valley Road, Suit 2013.	ility Tests for nth Edition. s Institute, 9 087, USA, 2 v Standards robial Suscent, CLSI de al and Labe	or Bacteria CLSI docu 950 West V 2012. s Institute ceptibility ocument M oratory Sta	that Grow Ament M07- alley Road, (CLSI). Pe Testing; Tv 1100-S23. (andards Ins	Aerobically; A9, Clinical Suite 2500, erformance venty-third CLSI docu- stitute, 950

Haemophilus influenzae

West Valley Hoda, Sulte 250U, Wayne, Pennsylvania 19087, USA, 2013.
3. Clinical and Laboratory Standards Institute (CLSI). Performance Standards for Antimicrobial Disk Diffusion Susceptibility Tests: Approved Standard – Eleventh Edition CLSI document M02-A11, Clinical and Laboratory Standards Institute, 950 West Valley Road, Suite 2500, Wayne, Pennsylvania 19087, USA, 2012.
4. Clinical and Laboratory Standards Institute (CLSI). Methods for Antimicrobial Susceptibility Testing of Anaerobic Bacteria; Approved Standard – Eight Edition. CLSI document M11-A8. Clinical and Laboratory Standards Institute, 950 West Valley Road, Suite 2500, Wayne, PA 19087 USA, 2012
5. Barnett ED, Teele DW, Klein JO, et al. Comparison of Ceftriaxone and Trimethoprim-Sulfamethoxazole for Acute Otitis Media. Pediatrics. Vol. 99, No. 1, January 1997.
Flagy(®) is a registered trademark of G.D. Searle & Co. 05-2013M
46110817

Manufactured in Austria by Sandoz GmbH for Sandoz Inc., Princeton, NJ 08540









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K only Ceftriaxone for Injection, USP ZODNA2 &

To reduce the development of drug-resistant bacteria and maintain the effectiveness of ceftriaxone for injection, and other antibacterial drugs, ceftriaxone for injection should be used only to treat or prevent infections that are proven or strongly suspected to be caused by bacteria. DESCRIPTION:

DESCRIPTION:

Ceftriaxone for injection, USP is a sterile, semisynthetic, broad-spectrum cephalosporin antibiotic for intravenous or intramuscular administration. Ceftriaxone sodium is (6*R*, 7*R*)-7-[2-(2-Amino-4-thiazolyl) glyoxylamido]-8-oxo-3-[[1, 2, 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, 72-(2)-(0-methyloxime), disodium salt, sesquaterhydrate.

The chemical formula of ceftriaxone sodium is C₁₈H₁₆NgNa₂O₇S₃*3.5H₂O. It has a calculated molecular weight of 661.60 and the following structural formula:

CGOO-Na*

CGOO-N of sodium per gram of cettria 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									
	Average Plasma Concentrations (mcg/mL)								
0.5 hr	1 hr	2 hr	4 hr	6 hr	8 hr	12 hr	16 hr	24 hr	
82	59	48	37	29	23	15	10	5	
22	33	38	35	30	26	16	ND	5	
20	32	38	34	31	24	16	ND	5	
151	111	88	67	53	43	28	18	9	
40	68	76	68	56	44	29	ND	ND	
257	192	154	117	89	74	46	31	15	
ND = Not determined.									
	22 20 151 40 257	0.5 hr 1 hr 82 59 22 33 20 32 151 111 40 68 257 192	Seven Jerus Pictor 0.5 hr 1 hr 2 hr 2 hr 82 59 48 22 33 38 20 32 38 151 111 88 40 68 76 257 192 154	Section Sect	State	State	State	No. State State	

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

Ceftriaxone was completely absorbed following IM administration with mean maximum plasma concentrations occurring between 2 and 3 hours post-dose. Multiple IV or IM doses ranging from 0.5 to 2 gm at 12- to 24-hour intervals resulted in 15% to 36% accumulation of ceftriaxone above single dose

alues. Ceftriaxone concentrations in urine are shown in Table 2.

Table 2. Urinary Concentrations of Ceftriaxone After Single Dose

		Aumm	istration								
	Av	Average Urinary Concentrations (mcg/mL)									
Dose/Route	0 to 2 hr	2 to 4 hr	4 to 8 hr	8 to 12 hr	12 to 24 hr	24 to 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					
ND = Not determined.											

ND = Not determined.

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 ceftriaxone, determined from 1 to 3 hours after dosing, were 581 mcg/mL in the gallbladder bile, 788 mcg/mL in the common duct bile, 898 mcg/mL in the cystic duct bile, 78.2 mcg/gm in the gallbladder wall and 62.1 mcg/mL in the concurrent plasma.

Over a 0.15 to 3 gm dose range in healthy adult subjects, the values of elimination half-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. Ceftriaxone is reversibly bound to human plasma proteins, and the binding decreased from a value of 95% bound at 300 mcg/mL. Ceftriaxone concentrations of < 25 mcg/mL to a value of 85% bound at 300 mcg/mL. The average values of maximum plasma concentration, elimination half-life, plasma 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. Ceftriaxone penetrated the inflamed meninges of infants and pediatric patients; CSF concentrations after a 50 mg/kg IV dose and after a 75 mg/kg IV dose and as as oshown in Table 3.

Table 3. Average Pharmacokinetic Parameters of Ceftriaxone in Pediatric

Table 3. Average Pharmacokinetic Parameters of Ceftriaxone in Pediatric

Patients With Meningitis			
	50 mg/kg IV	75 mg/kg IV	
Maximum Plasma			
Concentrations (mcg/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 (mcg/mL)	5.6	6.4	
Range (mcg/mL)	1.3 to 18.5	1.3 to 44	
Time after dose (hr)	37 (+ 16)	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 ceftriax-

Table 4. Average Pharmacokinetic Parameters of Ceftriaxone in Humans				
Subject Group	Elimination Half-Life (hr)	Plasma Clearance (L/hr)	Volume of Distribution (L)	
Healthy Subjects	5.8 to 8.7	0.58 to 1.45	5.8 to 13.5	
Elderly Subjects (mean age, 70.5 yr)	8.9	0.83	10.7	
Patients With Renal Impairment Hemodialysis Patients				
(0 to 5 mL/min)*	14.7	0.65	13.7	
Severe (5 to 15 mL/min)	15.7	0.56	12.5	
Moderate (16 to 30 mL/min)	11.4	0.72	11.8	
Mild (31 to 60 mL/min)	12.4	0.70	13.3	
Patients With Liver Disease	8.8	1.1	13.6	

The elimination of ceftriaxone is not altered when ceftriaxone is co-admir

Pharmacokinetics in the Middle Ear Fluid:

1.

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 ceftriaxone. Mean (±5D) ceftriaxone levels in the middle ear reached a peak of 35 (±12) mcg/mL at 24 hours, and remained at 19 (±7) mcg/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 intervals, a half-life of 25 hours was calculated. Ceftriaxone is highly bound to plasma proteins. The extent of binding to proteins in the middle ear fluid is unknown. Interaction with Calcium:

Two in vitro studies, one using adult plasma and the other neonatal plasma from umbilical cord blood have been carried out to assess interaction of ceftriaxone and calcium. Ceftriaxone concentrations up to 1 mM (in excess of concentrations achieved in vivo following administration of 2 grams ceftriaxone infused over 30 minutes) were used in combination with calcium concentrations up to 1 mM (16 mg/dL). Recovery of ceftriaxone from plasma was reduced with calcium concentrations of 6 mM (24 mg/dL) or higher in adult plasma or 4 mM (16 mg/dL) or higher in neonatal plasma. This may be reflective of ceftriaxone-calcium precipitation.

crobiology: echanism of Action:

Ceftriaxo s a bactericidal agent that acts by inhibition of bacterial cell wall synthesis. Ceftriaxone has activity in the presence of some beta-lactamases, both penicillinases and cephalosporinases, of Gram-negative and Gram-pos-

nanism of Resistance:

Resistance to ceftriaxone is primarily through hydrolysis by beta-lacta-mase, alteration of penicillin-binding proteins (PBPs), and decreased

Interaction with Other Antimicrobials In an in vitro study antagonistic effects have been observed with the combination of observed and refrievens Ceftriaxone has been shown to be active against most isolates of the following bacteria, both *in vitro* and in clinical infections as described in the INDICATIONS AND USAGE section:

ID USAGE section: Gram-negative bacteria Acinetobacter calcoacet Enterobacter aerogenes Enterobacter cloacae Escherichia coli Haemophilus influenzae Haemophilus parainfluenzae Klebsiella oxytoca

Neusiella Oxylucia Klebsiella pneumoniae Moraxella catarrhalis Morganella morganii Neisseria gonorrhoeae Neisseria meningitidis Proteus mirabilis Proteus vulgaris Pseudomonas aeruginosa Serratia marrescens

Serratia marcescens Gram-positive bacteria
Staphylococcus aureus
Staphylococcus epidermidis

Viridans group streptococci Anaerobic bacteria Bacteroides fragilis

Clostridium species

Clostridium species Peptostreptococcus species The following in vitro data are available, but their clinical significance is unknown. At least 90 percent of the following microorganisms exhibit an in vitro minimum inhibitory concentration (MIC) less than or equal to the susceptible breakpoint for ceffriaxone. However, the efficacy of ceftriaxone in treating clinical infections due to these microorganisms has not been established in adequate and well-controlled clinical trials.

Gram-positive bacteria
Streptococcus agalactiae

Anaerobic bacteria
Porphyromonas (Bacteroides) melaninogenicus
Prevotella (Bacteroides) bivius

Prevotella (Bacteroides) bivius

Susceptibility Test Methods:

When available, the clinical microbiology laboratory should provide the results of in vitro susceptibility test results for antimicrobial drug products used in resident hospitals to the physician as periodic reports that describe the susceptibility profile of nosocomial and community-acquired pathogens. These reports should aid the physician in selecting an antibacterial drug product for treatment.

rejorts should aid the physician in selecting an antibacterial drug product for treatment.

Dilution techniques: Quantitative methods are used to determine antimicrobial minimal inhibitory concentrations (MiCs). These MiCs provide estimates of the susceptibility of bacteria to antimicrobial compounds. The MiCs should be determined using a standardized test method 1.3. The MiC values should be interpreted according to criteria provided in Table 5.

Diffusion techniques: Quantitative methods that require measurement of zone diameters also provide reproducible estimates of the susceptibility of bacteria to antimicrobial compounds. The zone size should be determined using a standardized test method 2.3. This procedure uses paper disks impregnated with 30 mcg ceftriaxone to test the susceptibility of microorganisms to ceftriaxone. The disk diffusion interpretive criteria are provided in Table 5.

Anaerobic techniques: For anaerobic bacteria, the susceptibility to ceftriaxone as MiCs can be determined by a standardized agar test method 3.4. The MiC values obtained should be interpreted according to the criteria provided in Table 5.

Table 5. Susceptibility Test Interpretive Criteria for Ceftriaxone.

Minimum Inhibitory Concentrations (mcg/ml)		Disk Diffusion Zone Diameters (mm)			
(S) Suscep- tible	(I) Interme- diate	(R) Resist- ant	(S) Suscep- tible	(I) Interme- diate	(R) Resist- ant
≤ 1	2	≥4	≥ 23	20 to 22	≤19
≤2	-	-	≥26	-	-
≤ 0.25	-	-	≥ 35	-	-
≤ 0.12	-	-	≥ 34	-	-
≤ 0.5	1	≥ 2	-	-	-
≤1	2	≥4	_	_	-
≤0.5	-	-	≥ 24	_	-
≤ 1	2	≥ 4	≥27	25 to 26	≤24
≤16	32	≥64	_	_	_
	Concent (S)				

penicillin and either cefoxitin or oxacillin.
* The current absence of data on resistant isolates precludes defining any category other than 'Susceptible'. If isolates yield MIC results other than susceptible, they should be submitted to a reference laboratory for additional testino.

testing.
† Disc diffusion interpretive criteria for ceftriaxone discs against *Streptococcus* pneumoniae are not available, however, isolates of pneumococci with oxacillin zone diameters of ±20 mm are susceptible (MIC < 0.06 mcg/mL) to penicillin and can be considered susceptible to cetriaxone. Streptococcus pneumoniae isolates should not be reported as penicillin (cetriaxone) preuninnae isolates should not be reported as penicimin (certifiakone resistant or intermediate based solely on an oxacillin zone diameter of \leq 16 mm. The ceftriaxone MIC should be determined for those isolates with oxacillin zone diameters \leq 19 mm.

A report of *Susceptible* indicates that the antimicrobial is likely to inhibit rowth of the pathogen if the antimicrobial compound reaches the concentration A report of Susceptible indicates that the antimicrobial is likely to inhibit growth of the pathogen if the antimicrobial compound reaches the concentration at the infection site necessary to inhibit growth of the pathogen. A report of Intermediate indicates that the result should be considered equivocal, and if the microorganism is not fully susceptible to alternative, clinically feasible drugs, the test should be repeated. This category implies possible clinical applicability in body sites where the drug is physiologically concentrated or in situations where a high dosage of drug can be used. This category also provides a buffer zone that prevents small uncontrolled technical factors from causing major discrepancies in interpretation. A report of Resistant indicates that the antimicrobial some of the pathogen if the antimicrobial compound reaches the concentrations usually achievable at the infection site; other therapy should be selected.

Quality Control: Standardized susceptibility test procedures require the use of laboratory controls to monitor and ensure the accuracy and precision of supplies and reagents used in the assay, and the techniques of the individual performing the test1-2.3-4. Standard ceftriaxone powder should provide the following range of MIC values noted in Table 6. For the diffusion technique using the 30 mcc disk, the criteria in Table 6 Should be achieved.

Table 6. Acceptable Quality Control Ranges for Ceftriaxone

Table 6. Acceptable Quality Control Ranges for Ceftriaxone

QC Strain	Inhibitory Concentrations (mcg/mL)	Diffusion Zone diame- ters (mm)
Escherichia coli ATCC 25922	0.03 to 0.12	29 to 35
Staphylococcus aureus ATCC 25923		22 to 28
Staphylococcus aureus ATCC 29213	1 to 8	
Haemophilus influenzae ATCC 49247	0.06 to 0.25	31 to 39
Neisseria gonorrhoeae ATCC 49226	0.004 to 0.015	39 to 51
Pseudomonas aeruginosa ATCC 27853	8 to 64	17 to 23
Streptococcus pneumoniae ATCC 49619	0.03 to 0.12	30 to 35
Bacteroides fragilis ATCC 25285 (agar method)	32 to 128	
Bacteroides thetaiotaomicron ATCC 29741 (agar method)	64 to 256	

INDICATIONS AND USAGE:

Before instituting treatment with ceftriaxone, appropriate specimens should be obtained for isolation of the causative organism and for determination of its susceptibility to the drug. Therapy may be instituted prior to obtaining results of susceptibility testing. ment with ceftriaxone, appropriate specimens should

its susceptibility to the drug. Therapy may be instituted prior to obtaining results of susceptibility testing.

To reduce the development of drug-resistant bacteria and maintain the effectiveness of ceftriaxone for injection, USP and other antibacterial drugs, ceftriaxone for injection, USP should be used only to treat or prevent infections that are proven or strongly suspected to be caused by susceptibile bacteria. When culture and susceptibility information are available, they should be considered in selecting or modifying antibacterial therapy. In the absence of such data, local epidemiology and susceptibility patterns may contribute to the empiric selection of therapy. Cettriaxone for injection, USP is indicated for the treatment of the following infections when caused by susceptible organisms:

Lower Respiratory Tract Infections:

caused by Streptococcus pneumoniae, Staphylococcus aureus, Haemophilus influenzae, Haemophilus parainfluenzae, Klebsiella pneumoniae, Escherichia coli, Enterobacter aerogenes, Proteus mirabilis or Serratia marcescens.

Acute Bacterial Otilis Media:

caused by Streptococcus pneumoniae, Haemophilus influenzae (including

Acute Bacterial Otitis Media:
caused by Streptococcus pneumoniae, Haemophilus influenzae (including
beta-lactamase producing strains) or Moraxella catarrhalis (including beta-lactamase producing strains).
NOTE: In one study lower clinical cure rates were observed with a single dose
of ceftriaxone compared to 10 days of oral therapy. In a second study comparable cure rates were observed between single dose ceftriaxone and the comparator. The potentially lower clinical cure rate of ceftriaxone should be balanced
against the potential advantages of parenteral therapy (see CLINICAL STUDIES).
Skin and Skin Structure Infections:

Skin and Skin Structure Infections:
caused by Staphylococcus aureus, Staphylococcus epidermidis, Streptococcus
pyogenes, Viridans group streptococci, Escherichia coli, Enterobacter cloacae, Klebsiella oxytoca, Klebsiella pneumoniae, Proteus mirabilis, Morganella
morganii, "Pseudomonas aeruginosa, Serratia marcescens, Acinetobacter calcoaceticus, Bacteroides fragilis" or Peptostreptococcus species.
Virinary Tract Infections (complicated and uncomplicated):
caused by Escherichia coli, Proteus mirabilis, Proteus vulgaris, Morganella
morganii or Klebsiella neumoniae

caused by Escherichia coli, Proteus mirabilis, Proteus vulgaris, Morganella morganii or Klebsiella pneumoniae.

Uncomplicated Gonorrhea (cervical/urethral and rectal):
caused by Meisseria gonorrhoeae, including both penicillinase- and non-penicillinase-producing strains, and pharyngeal gonorrhea caused by non-penicillinase-producing strains of Neisseria gonorrhoeae.

Pelvic Inflammatory Disease:
caused by Neisseria gonorrhoeae. Cettriaxone sodium, like other cephalosporins, has no activity against Chlamydia trachomatis. Therefore, when cephalosporins are used in the treatment of patients with pelvic inflammatory disease and Chlamydia trachomatis is one of the suspected pathogens, appropriate antichlamydial coverage should be added.

Bacterial Septicemia:
caused by Staphylococcus aureus, Streptococcus pneumoniae, Escherichia

Bacterial Septicemia:
caused by Staphylococcus aureus, Streptococcus pneumoniae, Escherichia
coli, Haemophilus influenzae or Klebsiella pneumoniae.
Sone and Joint Infections:
caused by Staphylococcus aureus, Streptococcus pneumoniae, Escherichia
coli, Proteus mirabilis, Klebsiella pneumoniae or Enterobacter species.

coli, Proteus mirabilis, ricusteria precinional.
Intra-abdominal Infections:
 caused by Escherichia coli, Klebsiella pneumoniae, Bacteroides fragilis,
Clostridium species (Note: most strains of Clostridium difficile are resistant)
or Peptostreptococcus species.

caused by Haemophilus influenzae, Neisseria meningitidis or Streptococcus neumoniae. Ceftriaxone has also been used successfully in a limited number f cases of meningitis and shunt infection caused by Staphylococcus epider-nidis* and Escherichia coli.* Surgical Prophylaxis:

midis* and Escherichia coli.*

Surgical Prophylaxis:
The preoperative administration of a single 1 gm dose of ceftriaxone may reduce the incidence of postoperative infections in patients undergoing surgical procedures classified as contaminated or potentially contaminated (e.g., vaginal or abdominal hysterectomy or cholecystectomy for chronic calculous cholecystitis in high-risk patients, such as those over 70 years of age, with acute cholecystitis not requiring therapeutic antimicrobials, obstructive jaundice or common duct bile stones) and in surgical patients for whom infection at the operative site would present serious risk (e.g., during coronary artery bypass surgery). Although ceftriaxone has been shown to have been as effective as cefazolin in the prevention of infection following coronary artery bypass surgery, no placebo-controlled trials have been conducted to evaluate any cephalosporin antibiotic in the prevention of infection following coronary artery bypass surgery. When administered prior to surgical procedures for which it is indicated, a single 1 gm dose of ceftriaxone provides protection from most infections due to susceptible organisms throughout the course of the procedure.

Efficacy for this organism in this organ system was studied in fewer than ten infections.

CONTRAINDICATIONS:

on is contraindicated in patients with known allergy to n class of antibiotics. neonates, especially prematures, should not be treated ection. *In vitro* studies have shown that ceftriaxone can

Hyperbilirubinemic neonates, especially prematures, should not be treated with ceftriaxone for injection. *In vitro* studies have shown that ceftriaxone can displace bilirubin from its binding to serum albumin, leading to a possible risk

displace bilirubin from its binding to serum albumin, leading to a possible risk of bilirubin encephalopathy in these patients.

Ceftriaxone is contraindicated in neonates if they require (or are expected to require) treatment with calcium-containing IV solutions, including continuous calcium-containing infusions such as parenteral nutrition because of the risk of precipitation of ceftriaxone-calcium (see CLINICAL PHARMA-COLOGY, WARNINGS and DOSAGE AND ADMINISTRATION).

A small number of cases of fatal outcomes in which a crystalline material was observed in the lungs and kidneys at autopsy have been reported in neonates receiving ceftriaxone and calcium-containing fluids. In some of these cases, the same intravenous infusion line was used for both ceftriaxone and calcium-containing fluids was observed in the intravenous infusion line. At least one fatality has been reported in a neonate in whom ceftriaxone and calcium-containing fluids were administered at different time points via different intravenous lines; no crystalline material was observed at autopsy in this neonate. There have been no similar reports in patients other than neonates.

WARNINGS:
WHENDERS TO THE STATE OF THE S

WARNINGS: Hypersensitivity: BEFORE THERAPY WITH CEFTRIAXONE IS INSTITUTED, CAREFUL INQUIRY SHOULD BE MADE TO DETERMINE WHETHER THE PATIENT HAS HAD PRE-

VIOUS HYPERSENSITIVITY REACTIONS TO CEPHALOSPORINS, PENICILLINS OR OTHER DRUGS. THIS PRODUCT SHOULD BE GIVEN CAUTIOUSLY TO PFNI-VIOUS HYPERSENSITIVITY REACTIONS TO CEPHALOSPORINS, PENICILLINS OR OTHER DRUGS. THIS PRODUCT SHOULD BE GIVEN CAUTIOUSLY TO PENICILLIN-SENSITIVE PATIENTS. ANTIBIOTICS SHOULD BE ADMINISTERED WITH CAUTION TO ANY PATIENT WHO HAS DEMONSTRATED SOME FORM OF ALLERGY, PARTICULARLY TO DRUGS. SERIOUS ACUTE HYPERSENSITIVITY REACTIONS MAY REQUIRE THE USE OF SUBCUTANEOUS EPINEPHRIBE AND OTHER EMERGENCY MEASURES.

AS with other exphalosporins, anaphylactic reactions with fatal outcome have been reported, even if a patient is not known to be allergic or previously

exposed.
Interaction with Calcium-Containing Products:

Interaction with Calcium-Containing Products:

Do not use diluents containing calcium, such as Ringer's solution or Hartmann's solution, to reconstitute etfriaxone vials or to further dilute a reconstituted vial for IV administration because a precipitate can form. Precipitation of ceftriaxone-calcium can also occur when ceftriaxone is mixed with calcium-containing solutions in the same IV administration line. Ceftriaxone must not be administered simultaneously with calcium-containing IV solutions, including continuous calcium-containing intusions such as parenteral nutrition via a Y-site. However, in patients other than neonates, ceftriaxone and calcium-containing solutions may be administered sequentially of one another if the infusion lines are thoroughly flushed between infusions with a compatible fluid. In vitro studies using adult and neonatal plasma from umbilical cord blood demonstrated that neonates have an increased risk of precipitation of ceftriaxone-calcium (see CLINICAL PHARMACOLOGY, CONTRAINDICATIONS and DOSAGE AND ADMINISTRATION).

Clostridium difficile:
Clostridium difficile associated diarrhea (CDAD) has been reported with use of nearly all antibacterial agents, including ceftriaxone, and may range in severity from mild diarrhea to fatal collitis. Treatment with antibacterial agents alters the normal flora of the colon leading to overgrowth of C.

difficile.

C. difficile produces toxins A and B which contribute to the development of CDAD. Hypertoxin producing strains of C. difficile cause increased morbidity and mortality, as these infections can be refractory to antimicrobial therapy and may require colectomy. CDAD must be considered in all patients who present with diarrhea following antibiotic use. Careful medical history is necessary since CDAD has been reported to occur over two months after the administration of antibacterial agents.

CDAD has been reported to occur over two months after the administration of antibacterial agents.

If CDAD is suspected or confirmed, ongoing antibiotic use not directed against C. difficile may need to be discontinued. Appropriate fluid and electrolyte management, protein supplementation, antibiotic treatment of C. difficile, and surgical evaluation should be instituted as clinically indicated. Hemolytic Anemia:

An immune mediated hemolytic anemia has been observed in patients receiving cephalosporin class antibacterials including ceftriaxone. Severe cases of hemolytic anemia, including flatalities, have been reported during treatment in both adults and children. If a patient develops anemia while on ceftriaxone, the diagnosis of a cephalosporin associated anemia should be considered and ceftriaxone stopped until the etiology is determined. PRECAUTIONS:

General:

Reneral:

Prescribing ceftriaxone for injection in the absence of a proven or strongly uspected bacterial infection or a prophylactic indication is unlikely to provide tenefit to the patient and increases the risk of the development of drug-resis-

benefit to the patient and increases the risk of the development of drug-resistant bacteria.

Although transient elevations of BUN and serum creatinine have been observed, at the recommended dosages, the nephrotoxic potential of ceftriaxone is similar to that of other cephalosporins.

Ceftriaxone is excreted via both biliary and renal excretion (see CLINICAL PHARMACULGY). Therefore, patients with renal failure normally require no adjustment in dosage when usual doses of ceftriaxone are administered.

Dosage adjustments should not be necessary in patients with hepatic dysfunction, however, in patients with both hepatic dysfunction and significant renal disease, caution should be exercised and the ceftriaxone dosage should not exceed 2 om daily.

disease, caution should be exercised and the certraxone obsage should not exceed 2 gm daily.

Alterations in prothrombin times have occurred rarely in patients treated with ceftriaxone. Patients with impaired vitamin K synthesis or low vitamin K of stores (e.g., chronic hepatic disease and malnutrition) may require monitoring of prothrombin time during ceftriaxone treatment. Vitamin K administration (10 mg weekly) may be necessary if the prothrombin time is prolonged before or during therapy. rapy.
onged use of ceftriaxone may result in overgrowth of nonsusceptible

weekly) may be necessary if the prothrombin time is prolonged before or during therapy.

Prolonged use of ceftriaxone may result in overgrowth of nonsusceptible organisms. Careful observation of the patient is essential. If superinfection occurs during therapy, appropriate measures should be taken.

Ceftriaxone for injection should be prescribed with caution in individuals with a history of gastrointestinal disease, especially colitis.

There have been reports of sonographic abnormalities in the gallbladder of patients treated with ceftriaxone; some of these patients also had symptoms of gallbladder disease. These abnormalities appear on sonography as an echo without acoustical shadowing suggesting sludge or as an echo with acoustical shadowing which may be misinterpreted as gallstones. The chemical nature of the sonographically detected material has been determined to be predominantly a ceftriaxone-calcium salt. The condition appears to be transient and reversible upon discontinuation of ceftriaxone for injection and institution of conservative management. Therefore, ceftriaxone should be discontinued in patients who develop signs and symptoms suggestive of gallbladder disease and/or the sonographic findings described above.

Cases of pancreatitis, possibly secondary to bilary obstruction, have been reported rarely in patients treated with ceftriaxone. Most patients presented with risk factors for biliary stasis and biliary sludge (preceding major therapy, severe illness, total parenteral nutrition). A cofactor role of ceftriaxone-related biliary precipitation cannot be ruled out.

Information for Patients:

Patients should be counseled that antibacterial drugs including ceftriaxone for injection should only be used to treat bacterial infections. They do not treat viral infections (e.g., the common cold). When ceftriaxone for injection is prescribed to treat a bacterial infection, patients should be told that atthough it is common to feel better early in the course of therapy, the medication should be taken ex

the future.

Diarrhea is a common problem caused by antibiotics which usually ends when bearinea is a common problem caused by antibiotics which usually ends when the antibiotic is discontinued. Sometimes after starting treatment with antibiotics, patients 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.

Carcinogenesis, Mutagenesis, Impairment of Fertility:

cinogenesis: considering the maximum duration of treatment and the class of the com-and, carcinogenicity studies with ceftriaxone in animals have not been per-med. The maximum duration of animal toxicity studies was 6 months.

Mulagenesis:

Genetic toxicology tests included the Ames test, a micronucleus test and a test for chromosomal aberrations in human lymphocytes cultured in vitro with ceftriaxone. Ceftriaxone showed no potential for mutagenic activity in these

studies. 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.

Pregnancy Category B:
Reproductive studies have been performed in mice and rats at doses up to 20 times the usual human dose and have no evidence of embryotoxicity, etc-toxicity or teratogenicity. In primates, no embryotoxicity or teratogenicity was demonstrated at a dose approximately 3 times the human dose.
There are, however, no adequate and well-controlled studies in pregnant women. Because animal reproductive studies are not always predictive of human response, this drug should be used during pregnancy only if clearly needed.

needed.

Nonteratogenic Effects:
In rats, in the Segment I (fertility and general reproduction) and Segment III (perinatal and postnatal) studies with intravenously administered ceftriaxone, no adverse effects were noted on various reproductive parameters during gestation and lactation, including postnatal growth, functional behavior and reproductive ability of the offspring, at doses of 586 mg/kg/day or less.

Nursing Mothers:
Low concentrations of ceftriouse.

tions of ceftriaxone are excreted in human milk. Caution

(See Reverse)

Reference ID: 3446318

(Continued)

Pediatric Use:

Safety and effectiveness of ceftriaxone in neonates, infants and pediatric patients have been established for the dosages described in the **DOSAGE AND ADMINISTRATION** section. *In vitro* studies have shown that ceftriaxone, like some other cephalosporins, can displace bilirubin from serum albumin.

like some other cephalosporins, can displace bilirubin from serum albumin. Ceftriaxone should not be administered to hyperbilirubinemic neonates, especially prematures (see CONTRAINDICATIONS).

Geriatric Use:

Of the total number of subjects in clinical studies of ceftriaxone, 32% were 60 and over. No overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients, but greater sensitivity of some older individuals cannot be ruled out.

The pharmacokinetics of ceftriaxone were only minimally altered in geriatric patients compared to healthy adult subjects and dosage adjustments are not necessary for geriatric patients with ceftriaxone dosages up to 2 grams per day (see CLINICAL PHARMACOLOGY).

Ceftriaxone is generally well tolerated. In clinical trials, the following adverse reactions, which were considered to be related to ceftriaxone therapy or of uncertain etiology, were observed:

Local Reactions:

nduration and tenderness was 1% overall. Phlebitis was reported in pain, induration and reincerness was 176 overall. I floating that 24 of 45 after I/M administration. The incidence of warmth, tightness or induration was 17% (3/17) after IM administration of 350 mg/mL and 5% (1/20) after IM administration of 250 mg/mL.

ash (1.7%). Less frequently reported (<1%) were pruritus, fever or chills.

Hematologic: eosinophilia (6%), thrombocytosis (5.1%) and leukopenia (2.1%). Less fre-quently reported (<1%) were anemia, hemolytic anemia, neutropenia, lym-phopenia, thrombocytopenia and prolongation of the prothrombin time.

diarrhea (2.7%). Less frequently reported (<1%) were nausea or vomiting, and dysgeusia. The onset of pseudomembranous collits symptoms may occur during or after antibacterial treatment (see WARNINGS).

elevations of SGOT (3.1%) or SGPT (3.3%). Less frequently reported (<1%)

were elevations of alkàline phosphatase and bilirubin **Renal**: vations of the BUN (1.2%). Less frequently reported (<1%) were eleva-of creatinine and the presence of casts in the urine.

Central Nervous System:
headache or dizziness were reported occasionally (<1%).
Geniturinary:
moniliasis or vaginitis were reported occasionally (<1%).

Miscellaneous:
diaphoresis and flushing were reported occasionally (<1%).
Other rarely observed adverse reactions (<0.1%) include abdominal pain, agranulocytosis, allergic pneumonitis, anaphylaxis, basophilia, biliary lithiasis, bronchospasm, colitis, dyspepsia, epistaxis, flatulence, galloladder sludge, glycosuria, hematuria, jaundice, leukocytosis, lymphocytosis, nenporolytosis, nephrolithiasis, palpitations, a decrease in the prothrombin time, renal precipitations, seizures, and serum sickness.
Postmarketing Experience:
In addition to the adverse reactions reported during clinical trials, the following adverse experiences have been reported during clinical practice in

In addition to the adverse reactions reported during clinical trials, the following adverse experiences have been reported during clinical practice in patients treated with ceftriaxone. Data are generally insufficient to allow an estimate of incidence or to establish causation.

A small number of cases of fatal outcomes in which a crystalline material was observed in the lungs and kidneys at autopsy have been reported in neonates receiving ceftriaxone and calcium-containing fluids. In some of these cases, the same intravenous infusion line was used for both ceftriaxone and calcium-containing fluids and in some a precipitate was observed in the intravenous infusion line. At least one fatality has been reported in a neonate in whom ceftriaxone and calcium-containing fluids were administered at different time points via different intravenous lines; no crystalline material was observed at autopsy in this neonate. There have been no similar reports in patients other than neonates.

Gastrointestinal:

stomatitis and glossitis.

Genitourinary:

exanthema, allergic dermatitis, urticaria, edema. As with many medications, isolated cases of severe cutaneous adverse reactions (erythema multi-forme, Stevens-Johnson syndrome or Lyell's syndrome/toxic epidermal necrol-ysis) have been reported.

Cephalosporin Class Adverse Reactions:

In addition to the adverse reactions: In addition to the adverse reactions listed above which have been observed in patients treated with ceftriaxone, the following adverse reactions and altered laboratory test results have been reported for cephalosporin class antibiotics: Adverse Reactions:

Adverse Reactions:
Allergic reactions, drug fever, serum sickness-like reaction, renal dysfunction, toxic nephropathy, reversible hyperactivity, hypertonia, hepatic dysfunction including cholestasis, aplastic anemia, hemorrhage, and superinfection.
Altered Laboratory Tests:
Positive direct Coombs' test, false-positive test for urinary glucose, and ele-

vated LDH

vated LDH.

Several cephalosporins have been implicated in triggering seizures, particularly in patients with renal impairment when the dosage was not reduced (see DOSAGE AND ADMINISTRATION). If seizures associated with drug therapy occur, the drug should be discontinued. Anticonvulsant therapy can be given OVERDOSAGE:

/ERDOSAGE: In the case of overdosage, drug concentration would not be reduced by modialysis or peritoneal dialysis. There is no specific antidote. Treatment of overdosage should be symptomat DOSAGE AND ADMINISTRATION:

overdosage should be symptomatic.

DOSAGE AND ADMINISTRATION:
Ceftriaxone may be administered intravenously or intramuscularly.
Do not use diluents containing calcium, such as Ringer's solution or Hartmann's solution, to reconstitute ceftriaxone vials or to further dilute a reconstituted vial for IV administration because a precipitate can form. Precipitation of ceftriaxone-calcium can also occur when ceftriaxone is mixed with calcium-containing solutions in the same IV administration line. Ceftriaxone must not be administered simultaneously with calcium-containing IV solutions, including continuous calcium-containing infusions such as parenteral nutrition via a Y-site. However, in patients other than neonates, ceftriaxone and calcium-containing solutions may be administered sequentially of one another if the infusion lines are thoroughly flushed between infusions with a compatible fluid (see WARNINGS).

There have been no reports of an interaction between ceftriaxone and calcium-containing products or interaction between intramuscular ceftriaxone and calcium-containing products (V or or oral).

Neonates:

Hyperbilirubinemic neonates, especially prematures, should not be treated with ceftriaxone for injection (see CONTRAINDICATIONS).

Ceftriaxone for injection (see CONTRAINDICATIONS).

Ceftriaxone ceftriaxone ceftriaxone-calcium (see CONTRAINDICATIONS).

Pediatric Patients:

For the treatment of skin and skin structure infections, the recommended

risk of precipitation of centracons section.

Pediatric Patients:
For the treatment of skin and skin structure infections, the 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.

For the treatment of acute bacterial otitis media, a single intramuscular dose for the treatment of acute bacterial otitis media, a single intramuscular dose

of 50 mg/kg (not to exceed 1 gram) is recommended (see INDICATIONS AND USAGE).

USAGE.

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. The total daily dose should not exceed 2 grams. In the treatment of meningitis, it is recommended that the initial therapeutic dose be 100 mg/kg (not to exceed 4 grams). Thereafter, a total daily dose of 100 mg/kg/day (not to exceed 4 grams). Thereafter, a total daily dose of 100 mg/kg/day (not to exceed 4 grams daily) is recommended. The daily dose may be administered once a day (or in equally divided doses every 12 hours). The usual duration of therapy is 7 to 14 days.

Adults:

Adults:

The usual adult daily dose is 1 to 2 grams given once a day (or in equally divided doses twice a day) depending on the type and severity of infection. For infections caused by Staphylococcus aureus (MSSA), the recommended daily dose is 2 to 4 grams, in order to achieve >90% target attainment. The total daily dose should not exceed 4 grams.

If Chlamydia trachomatis is a suspected pathogen, appropriate antichlamydial verage should be added, because ceftriaxone sodium has no activity against

coverage should be added, because ceftriaxone sodium has no activity against this organism.

For the treatment of uncomplicated gonococcal infections, a single intramuscular dose of 250 mg is recommended.

For preoperative use (surgical prophylaxis), a single dose of 1 gram administered intravenously 1/2 to 2 hours before surgery is recommended.

Generally, ceftriaxone 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 complicated infections, longer therapy may be required.

When treating infections caused by Streptococcus pyogenes, therapy should be continued for at least 10 days.

No dosage adjustment is necessary for patients with impairment of renal or hepatic function.

Intramuscular Administration:

Intramuscular Administration:
Reconstitute cetriaxone sodium powder with the appropriate diluent (see DOSAGE AND ADMINISTRATION: Compatibility and Stability).
Inject diluent into vial, shake vial thoroughly to form solution. Withdraw entire contents of vial into syringe to equal total labeled dose.
After reconstitution, each 1 mL of solution contains approximately 250 mg or 350 mg equivalent of ceftriaxone according to the amount of diluent indicated below. If required, more dilute solutions could be utilized. A 350 mg/mL concentration is not recommended for the 250 mg vial since it may not be possible to withdraw the entire contents. As with all intramuscular preparations, cettriaxone should be injected well within the body of a relatively large muscle; aspiration helps to avoid unintentional injection into a blood vessel.

	Amount o	Amount of Diluent to be Added		
Vial Dosage Size	250 mg/mL	350 mg/mL		
250 mg	0.9 mL	-		
500 mg	1.8 mL	1 mL		
1 gm	3.6 mL	2.1 mL		
2 gm	7.2 mL	4.2 mL		

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

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.

Compatibility and Stability:
Ceftriaxone has been shown to be compatible with Flagyl® IV (metronidazole hydrochloride). The concentration should not exceed 5 to 7.5 mg/mL metronidazole 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 (DSW). No compatibility studies have been conducted with the Flagyl® IV RTU® (metronidazole) for mulation or using other diluents. Metronidazole at concentrations greater than 8 mg/mL will precipitate. On ont refrigerate the admixture as precipitation will occur.

Vancomycin, amsacrine, aminoglycosides, and fluconazole are physically incompatible with ceftriaxone in admixture. When any of these drugs are to be administered concomitantly with ceftriaxone by 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

administrations. On to use diluents containing calcium, such as Ringer's solution or Dartmann's solution, to reconstitute ceftriaxone for injection or to further dilute a reconstituted vial for IV administration. Particulate formation can

result.

Ceftriaxone for injection solutions should *not* be physically mixed with or piggybacked into solutions containing other antimicrobial drugs or into diluent solutions other than those listed above, due to possible incompatibility (see

WARNINGS).
Ceftriaxone sodium sterile powder should be stored at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature] and protected from light. After reconstitution, protection from normal light is not necessary. The color of solutions ranges from light yellow to amber, depending on the length of storage, concentration and diluent used.
Ceftriaxone intramuscular solutions remain stable (loss of potency less than 10%) for the following time periods:

	Concentration	Sto	rage
Diluent	mg/mL	Room Temp. (25°C)	Refrigerated (4°C)
Sterile Water for Injection	250, 350	24 hours	3 days
100	2 days	10 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% Benzyl Alcohol	100	24 hours	10 days
	250, 350	24 hours	3 days
1% Lidocaine Solution (without epinephrine)	100	24 hours	10 days
	250, 350	24 hours	3 days

Ceftriaxone 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:

	Sto	rage
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* ceftriaxone solutions are stable at room temperature The following intravenous certinaxone solutions are stable at from 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), atter the indicated stability time periods, unused portions of solutions between the container of the container of

After the indicated stability units periods, unlock published be discarded.

NOTE: Parenteral drug products should be inspected visually for particulate matter before administration.

Ceftriaxone reconstituted with 5% Dextrose or 0.9% Sodium Chloride solution at concentrations between 10 mg/mL and 40 mg/mL, and then stored in frozen state (-20°C) in PVC or polyolefin containers, remains stable for 26 weeks.

Frozen solutions of ceftriaxone for injection should be thawed at room temperature before use. After thawing, unused portions should be discarded.

ceftriaxone is more soluble in human gallbladder bile and the calcium content of human gallbladder bile is relatively low.

IW SUPPLIED:
Ceftriaxone for injection, USP is supplied as a sterile crystalline powder in
iss vials. The following packages are available:
Vials containing 250 mg equivalent to ceftriaxone. Package of 10 (0781-3206-

. /ials containing 500 mg equivalent to ceftriaxone. Package of 10 (0781-3207-

95). Vials containing 1 gm equivalent to ceftriaxone. Package of 10 (0781-3208-

Vials containing 2 gm equivalent to ceftriaxone. Package of 10 (0781-3209

95). Vials containing 250 mg equivalent to ceftriaxone. Package of 1 (0781-3206-*).* Vials containing 500 mg equivalent to cettriaxone. Package of 1 (0781-3207:

85). Vials containing 1 gm equivalent to ceftriaxone. Package of 1 (0781-3208-

Viais containing 5 \$5). Storage Prior to Reconstitution: Store at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature].

Protect from light.

CLINICAL STUDIES:

Clinical Trials in Pediatric Patients With Acute Bacterial Otitis Media:

In two adequate and well-controlled US clinical trials a single IM dose of 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 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		Ceftriaxone is lower than
14	74% (220/296)	82% (247/302)	(-14.4%, -0.5%)	control at study day 14
28	58% (167/288)	67% (200/297)	(-17.5%, -1.2%)	and 28.
Study 2 – US5		TMP-SMZ		Ceftriaxone is
14	54% (113/210)	60% (124/206)		equivalent to
28	35% (73/206)	45% (93/205)	(-19.9%, 0.0%)	study day 14 and 28.

An open-label bacteriologic study of ceftriaxone without a comparator enrolled 108 pediatric patients, 79 of whom had positive baseline cultures for one or more of the common pathogens. The results of this study are tabulated as follows:

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

	Study Day 13 to 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)

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